DRUG FORMULARY
2016

Department of Health & Family Welfare
Government of Chhattisgarh
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2016

Department of Health & Family Welfare
Government of Chhattisgarh
# Contents

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Narayan Tripathi B Pharm and Mr Fidius Kerketta, MSc. in Public Health were the key organizing partners and handled all the logistics and support of this work. The Editorial team also places on record thanks to the services of HSS unit team members who from time to time provided their valuable inputs.

Note: Chhattisgarh Drug Formulary 2016 is not a regulatory document. Physicians are supposed to use their professional judgment. Inclusion/Exclusion of monographs in Chhattisgarh Drug Formulary 2016 is a dynamic process. The drugs contained in Chhattisgarh Drug Formulary 2016 have been chosen for rational and economic prescribing. Chhattisgarh Drug Formulary 2016 would serve as a guidance document to medical practitioners, rural medical practitioners (RMA), pharmacists, nurses, medical and pharmacy students, and other healthcare professionals and stakeholders in healthcare system. The feedback from stakeholders is invited.
Foreword

We are pleased to present the 2nd Edition of Chhattisgarh Drug Formulary 2016. It has materialized after a gap of one decade. During this period, there have been tremendous advancements in therapeutic strategies and newly available drugs. This edition incorporates the changes based on the current knowledge.

Valuable inputs that emerged during the meetings of the expert group meetings and the inputs received in response to the pre-print version circulated have given this edition a unique feature by incorporating value added information. The Department is greatly indebted to the Members of the Expert Group and the Subject Review Experts from diverse fields who consented to review the manuscript of the Formulary. The services of all these experts are appreciated.

The first edition of Chhattisgarh Drug Formulary (CDF) 2003 was based on the WHO Drug Formulary 2003. It was published in 2003 by the Department of Health, Govt. of Chhattisgarh. In the past one decade there has been vast expansion in the range of new drugs and their formulations. To address the need of publication of an updated version of Chhattisgarh Drug Formulary 2016, Department of Health and Family Welfare, Government of Chhattisgarh vide their Notification No. F. No.1-111/2013/9/17-1Raipur Dated 07/06/2013 State Health Resource Center, Raipur (SHRC), a Technical Support Agency to the Department of Health and Family Welfare, Government of Chhattisgarh took an initiative to update the Chhattisgarh Drug Formulary (CDF) 2016. For this purpose, SHRC called for expert advice/suggestion from different departments of Pt J.N.M. Medical College Raipur, Chhattisgarh Institute of Medical Sciences, Bilaspur and Government Medical College & Maharani Hospital, Jagdalpur.

Experts from following departments were involved in updating Chhattisgarh Drug Formulary (CDF) 2016.

- Medicine
- Obstetrics & Gynecology
- Cardiology
- Orthopedics
- Anesthesia
- Oncology
- Pharmacology
- Skin & V.D.
- Nephrology
- Psychiatry
- Ophthalmology
- Pulmonary Medicine
To fulfill the mandate of publishing the CDF, the following process has been adopted:

Policy Framework by SHRC

Drug Formulary 2003 based on WHO Model Formulary 2003 - taken as zero draft

Modification to Chhattisgarh Context suggested by Expert from Medical colleges in CG

Review by Expert Group

Pre-Print Version Comments/ Approval Department of Health, Govt. of Chhattisgarh

Review and Incorporation comments from of Department of Health, Govt. of Chhattisgarh

Adoption for CDF

Special thanks go to the expert group who prepared final draft of CDF. This Chhattisgarh Drug Formulary has been adopted from the Drug Formulary 2003 which was based on the WHO Model Formulary 2003 and thoroughly updated for its content, especially keeping in view the end user in Chhattisgarh for which we wish to thank profusely to the departments of Medicine, Obstetrics & Gynecology, Orthopedics, Anesthesia, Pharmacology, Psychiatry, Ophthalmology, Cardiology, Oncology, Nephrology and Pulmonary Medicine. With publication of this book another important Health Sector Reform milestone has been achieved by the Department of Health, Government of Chhattisgarh

Vikas Sheel
Secretary to Government of Chhattisgarh
Department of Health and Family Welfare
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मगहानदी भवन, नया रायपुर – 492002

क्र. / एक-1-111 / 2013 / नी / 17-1
रायपुर, दिनांक: 7 जुलाई 2013

प्रति,
अध्यक्ष, स्वास्थ्य सेवाएं, छोटागोमा
प्रबंध संचालक, छ. ग. मेडिकल सर्विसेस कार्यालय लिमिटेड, छोटागोमा
संघातक, स्वास्थ्य सेवाएं, छोटागोमा
संघातक, विकल्प सीमा, छोटागोमा

विषय —
शासकीय स्वास्थ्य संस्थाओं में नियुक्त जेनरिक दवा वितरण बाबु।

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गणतान्त्रिक मुख्यमंत्री जी द्वारा बनाई संगठन 2013-14 में सूचना की है कि स्वास्थ्य सेवाओं को बेहतर बनाने के उद्देश्य से आगामी वित्तीय वर्ष से शासकीय स्वास्थ्य संस्थाओं में सभी मरीजों के लिए नियुक्त जेनरिक दवाइयों उपलब्ध कराने हेतु उपरोक्त घोषणा के अनुसार स्वास्थ्य मंत्री, सामुदायिक स्वास्थ्य केंद्रों, स्थानिक अस्पतालों, जिला धिर्मितासभाओं एवं शासकीय धिर्मितासभा/स्थान धिर्मितासभा महाधिव्यासीयों से संबंधित अस्पतालों में सभी मरीजों को जेनरिक दवाओं का वितरण 15 अगस्त 2013 से सुनिश्चित किया जाना है। नीति के वितरण दिशा अनुसार निरूपित किये जाने हैं—

1. अनिवार्य औषधि सूची —

1.1 औषधियों का क्रम अनिवार्य औषधि सूची (Essential Drug List) के अधार पर किया जायेगा। इस सूची को राज्य सरकार पर प्रवेश 2 या 3 वर्षों में पुनर्दर्जित कर अधिति किया जायेगा। वर्तमान में, अनिवार्य औषधि सूची, 2013 लागू है।

1.2 ऐसी औषधियों जो अनिवार्य औषधि सूची में नहीं हैं, स्वास्थ्य सरकार पर आवश्यकतानुसार समस्त मुख्य धिर्मितासभा एवं स्वास्थ्य अधिकारी, स्थानिक सर्जन सह मुख्य अस्पताल अधीक्षक एवं अन्य अधीक्षक (अस्पताल) क्रम कर सकेंगे। समानान्तर यह राशि आप्रविजित बजट के 20 प्रतिशत से अधिक नहीं होगी।

2. क्रय, भण्डारण एवं स्वास्थ्य संस्थाओं के लिए वितरण व्यवस्था —

2.1 अनिवार्य औषधि सूची में शामिल दवाओं का क्रय छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय द्वारा किया जायेगा।
22 दस्तावेजों की संज्ञानकर्ता वार्षिक मांग मुख्य चिकित्सा एवं स्वास्थ्य अधिकारियों द्वारा संचालित, चिकित्सा सेवाओं के माध्यम से छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय को प्रस्तुत की जाएगी। चिकित्सा एवं वैद्यकीय गतिविधियों से संबंधित चिकित्सालय अपनी वार्षिक मांग संचालनालय, दिकित्सा शिक्षा के माध्यम से छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय को प्रस्तुत करेगी।

23 छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय द्वारा अनिवार्य दवा सूची (Essential Drug List) अनुसार दवा निर्णय (Rote Contract) तय किये जाएगे। दस्तावेजों के माध्यम एवं वितरण की समुच्चय व्यवस्था कार्यालय द्वारा की जाएगी।

24 छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय द्वारा तीन माह की आवश्यकता अनुसार दवाएं भण्डार में Buffer Stock के रूप में रखी जाएगी। साथ ही, कम से कम दो माह की आवश्यकता अनुसार दस्तावेजों का अधिग्रहण कार्यालय द्वारा किया जाएगा।

25 सभी स्वास्थ्य संस्थाओं के प्रमुख अनिवार्य औषधियों की एक पाश्चात्य सम्मानित करेंगे, जिसके आधार पर वे छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय को दवाओं की आवश्यक मात्रा हेतु Indent प्रस्तुत करेंगे।

26 किसी भी संस्था द्वारा अधिकतम कितनी मात्रा में दवाएं Indent की जा सकती हैं। इसकी तीन साल कार्यालय द्वारा, स्वास्थ्य संस्थाओं द्वारा उपलब्ध कराई जा रही सेवाओं एवं ओ.पी.डी. व आई.पी.डी. संस्थाओं के आधार पर निर्धारित की जाएगी।

27 छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय सभी प्राथमिक स्वास्थ्य केंद्रों, सामुदायिक स्वास्थ्य केंद्रों, रिजिल्ट अस्पतालों एवं विजय चिकित्सालयों तक दवाओं का परिवहन सुनिश्चित करेंगे। चिकित्सा/दंत चिकित्सा महाविद्यालयों से संबंधित चिकित्सालय एवं एथल स्वास्थ्य संस्थाओं को छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय के भण्डार से परिवहन का प्रबंधन स्वयं करना होगा।

28 स्वास्थ्य संस्थाओं में एक Stock Monitoring System स्थापित किया जाएगा जो कि छत्तीसगढ़ मेडिकल सर्विसेस कार्यालय के कंप्यूटर सिस्टम से जुड़ा (linked) होगा। इससे स्वास्थ्य संस्थाओं में अनिवार्य दवाओं की कंट्रोल में राहत रहेगी पहचान कर पूर्ण किया जा सकेगा।

29 स्वास्थ्य संस्थाओं के प्रमुखों एवं भण्डार प्रबंधकों को उपरोक्त नजीक दवा वितरण एवं आपूर्ति की व्यवस्था के बेहतर क्रियान्वयन हेतु कार्यालय द्वारा प्रशिक्षण प्रदान किया जाने का प्रबंध किया जाएगा।
3. मरीजों के लिए वितरण व्यवस्था –

3.1 मरीजों को दवा वितरण संस्था के फार्मासिस्ट द्वारा किया जाएगा। जहां फार्मासिस्ट उपलब्ध न हो वह व्यवस्था की जीवन दीव समिति स्थानीय स्तर पर फार्मासिस्ट की नियुक्ति संविदा के आधार पर करेंगी। इस व्यवस्था के होने तक दवा किसी संस्था के चिकित्सक, नर्तक अथवा ए.एन.एम. द्वारा किया जाएगा।

3.2 सभी मरीजों को, चाहे कोई विभाग रोगी हों अथवा लाभार्थी स्वास्थ्य कीमा योजना एवं मुख्यांशी स्वास्थ्य कीमा योजना के तहत स्वास्थ्य अधिकार जीवन, स्वास्थ्य जीवन देने वाले विशेषता दी जाती है।

3.3 यह सुनिश्चित किया जाना है कि किसी भी मरीज को दवा प्राप्त करने के लिए 10 मिनट से अधिक प्रवर्तक न करनी पड़े। जहां आवश्यक हो, सम्बन्धित जीवन दीव समिति इसके लिए स्थानीय स्तर पर अन्तरित कर फार्मासिस्ट की व्यवस्था संविदा नियुक्ति के आधार पर कर सकती है।

3.4 हर संस्था के लिए निष्कर्ष अनिवार्य औषधि सूची एवं प्रत्येक दवा के स्टाक का दैनिक स्थिति दवा वितरण केन्द्र पर अनिवार्य प्रशिक्षित की जायेगी।

4. पूरक गतिविधियाँ –

4.1 प्रदेश में सामान्य रोगों के उपचार हेतु 'मानक उपचार गार्डर' (Standard Treatment Guidelines) एवं उपयोग की जाने वाली दवाओं की फार्मुलरी (Formulary) उपलब्ध है। इन्हें अनिवार्य औषधि सूची के आधार पर पुनर्रीतिक्त (Revised) किया जाता है। प्रदेश का समस्त लाभार्थी स्वास्थ्य / चिकित्सा संस्था में उपचार द्वारा तदनुसार लागू मानक मार्गदर्शक एवं पुनर्रितिक्त फार्मुलरी (Formulary) प्रकाशन में रहेगा।

4.2 समस्त शासकीय जिला चिकित्सालयों एवं महाविद्यालयों से संबंधित चिकित्सालयों में ओढ़ित दवा के उपयोग (Rational Drug Use) को अन्तर्गत करते हुए “Drugs and Therapeutics Committee (DTC)” औषधि एवं चिकित्सा समिति की स्थापना की जायेगी। इस समिति के गृह गार्डर नियुक्तिकरित होंगे –

4.2.1 यह सुनिश्चित करना कि चिकित्सालय में कार्यान्वयन चिकित्सक एवं विशेषज्ञ जीवन दीव करेंगे।

4.2.2 यह सुनिश्चित करना कि सभी संबंधित कार्यकर्ता मानक उपचार गार्डर के अनुसार दवाओं की फार्मुलरी (Formulary) में प्रशिक्षित हों।

4.2.3 यह सुनिश्चित करना कि चिकित्सक /विशेषज्ञ के द्वारा लिखे जाने वाले दवाइयों के पर्यंत मानकों के आधार पर हो, समय—समय पर दवाइयों के
पत्र (Prescription) का निरीक्षण, अत्यधिक गुणवत्ता एवं दवाइयों की सही समय पर पहुँच (Logistic) पर मूल्यांकन।

4-2.4 प्रेस्क्रिप्शन आड्ट (Prescription Audit) के नक्सलों की समीक्षा करना एवं राज्य शासन की आवश्यक अनुमोदन करना।

उपरोक्त दिशा निर्देशों का पालन करने हुए निम्नलिखित जनेनरल द्वारा वितरण नीति का क्षेत्राभ्यास सुनिश्चित करें।

छत्तीसगढ़ के राज्यपाल के नाम से
राया आदेशानुसार

(उपच. शासन)

प्रमुख सचिव
छत्तीसगढ़ शासन
स्वास्थ्य एवं परिवार कल्याण विभाग

पुष्प. क्र. /एफ. Yi. 1-111 / 2013 / नी / 17-1

प्रतिलिपि :-
1. अधिकारिक, मुख्य सचिव छोटोगो शासन, कार्यालय, भंडारी छत्तीसगढ़, रायपुर की और सूचना।
2. समस्त अपर मुख्य अधिकारियों/प्रमुख सचिव/सचिव, छत्तीसगढ़, शासन।
3. मिशन संचालक, राष्ट्रीय ग्रामीण स्वास्थ्य मिशन, छत्तीसगढ़, रायपुर की और सूचना एवं पालनाध्यक्ष।
4. कार्यालय संचालक, राज्य स्वास्थ्य संस्थान केंद्र, छत्तीसगढ़, रायपुर की और सूचना एवं पालनाध्यक्ष।
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6. समस्त अधीक्षकाधीन/अधीक्षक विभाग /दंत विभाग स्वास्थ्य विभाग की और सूचनाएँ एवं पालनाध्यक्ष।
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9. समस्त मुख्य विभाग स्वास्थ्य एवं स्वास्थ्य अधीक्षक, छत्तीसगढ़ की और सूचना एवं पालनाध्यक्ष।
10. समस्त सिद्धित सर्जन सह मुख्य अस्पताल अधीक्षक, छत्तीसगढ़ रायपुर की और सूचना एवं पालनाध्यक्ष।

अधिकारिक
छत्तीसगढ़ शासन
स्वास्थ्य एवं परिवार कल्याण विभाग

XI
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# Color coded symbols for category of drugs for State Drug Formulary

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<th>Category</th>
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<td>Universal</td>
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<td>Primary Health Centers as well as Secondary and Tertiary care facilities</td>
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<td>District Hospital/Civil Hospital/ Community Health Centre/First referral unit/Sub-divisional Hospital as well as Tertiary care facilities</td>
</tr>
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<tr>
<td>Tertiary restricted</td>
<td>Anti-cancerous (Indicates use in centers having Cancer diagnostic and treatment facilities.</td>
</tr>
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</table>
Common Abbreviations

ACE  Angiotensin Converting Enzyme
ACE Inhibitors  Angiotensin Converting Enzyme Inhibitors
ADR  Adverse Drug Reaction
AE  Adverse Event
AIDS  Acquired Immuno Deficiency Syndrome
ARB  Angiotension Receptor Blocker
BCG  Bacillus Calmette Guerin
BNF  British National Formulary
BP  British Pharmacopoeia
BSA  Body Surface Area
CAPD  Continuous Ambulatory Peritoneal Dialysis
CD4  Cluster of Differentiation 4
CDF  Chhattisgarh Drug Formulary
CDSCO  Central Drugs Standards Control Organization
CIOMS  Council for International Organization of Medical Sciences
CMV  Cytomegalovirus
COPD  Chronic Obstructive Pulmonary Disease
COLD  Chronic Obstructive Lung Disease
CR  Controlled Release
CSF  Cerebrospinal Fluid
DCGI  Drugs Controller General (India)
DOHFW  Department of Health and Family Welfare
DMARDs  Disease Modifying Anti-Rheumatic Drugs
DOTS  Directly Observed Treatment Short course
DT  Dispersible Tablet/Diphtheria Tetanus
DPT  Diphtheria Pertussis Tetanus
EMEA  European Medicines Evaluation Agency
ER  Extended Release
FDA  Food and Drug Administration
FDC  Fixed Dose Combination
GCP  Good Clinical Practice
GERD  Gastroesophageal Reflux Disease
GFR  Glomerular Filtration Rate
G-6-PD  Glucose-6-Phosphate Dehydrogenase
HAART  Highly Active Anti-Retroviral Therapy
HD  Hemodialysis
HIV  Human Immunodeficiency Virus
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tr>
<td>HPA</td>
<td>Hypothalamic Pituitary Adrenal Axis</td>
</tr>
<tr>
<td>HT</td>
<td>Hormone Therapy</td>
</tr>
<tr>
<td>ICMR</td>
<td>Indian Council of Medical Research</td>
</tr>
<tr>
<td>INR</td>
<td>International Normalized Ratio</td>
</tr>
<tr>
<td>IP</td>
<td>Indian Pharmacopoeia</td>
</tr>
<tr>
<td>IPC</td>
<td>Indian Pharmacopoeia Commission</td>
</tr>
<tr>
<td>IPV</td>
<td>Inactivated Poliomyelitis Vaccine</td>
</tr>
<tr>
<td>IU</td>
<td>International Units</td>
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<tr>
<td>LFT</td>
<td>Liver Function Test</td>
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<tr>
<td>MAO</td>
<td>Mono Amine Oxidase</td>
</tr>
<tr>
<td>MD</td>
<td>Mouth Dissolving</td>
</tr>
<tr>
<td>MDR</td>
<td>Multi Drug Resistance</td>
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<tr>
<td>mEq</td>
<td>MilliEquivalent</td>
</tr>
<tr>
<td>MI</td>
<td>Myocardial Infarction</td>
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<tr>
<td>mMol</td>
<td>Millimole</td>
</tr>
<tr>
<td>MMR</td>
<td>Measles, Mumps and Rubella</td>
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<tr>
<td>MR</td>
<td>Modified Release</td>
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<tr>
<td>NLEM</td>
<td>National List of Essential Medicines</td>
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<td>NFI</td>
<td>National Formulary of India</td>
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<td>NRHM</td>
<td>National Rural Health Mission</td>
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<td>NSAIDs</td>
<td>Non-Steroidal Anti-Inflammatory Drugs</td>
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<td>NS</td>
<td>Normal Saline</td>
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<tr>
<td>ODT</td>
<td>Oral Dispersible Tablet</td>
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<tr>
<td>OPV</td>
<td>Oral Polio Vaccine</td>
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<tr>
<td>PFS</td>
<td>Pre-Filled Syringes</td>
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<tr>
<td>PK/PD</td>
<td>Pharmacokinetic/Pharmacodynamic</td>
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<tr>
<td>PD</td>
<td>Peritoneal Dialysis</td>
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<tr>
<td>PvPPI</td>
<td>Pharmacovigilance Programme of India</td>
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<tr>
<td>SC</td>
<td>Subcutaneous SL Sublingual</td>
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<tr>
<td>SLE</td>
<td>Systemic Lupus Erythematosus</td>
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<tr>
<td>SR</td>
<td>Sustained Release</td>
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<tr>
<td>SWI</td>
<td>Sterile Water for Injection</td>
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<tr>
<td>SSRI</td>
<td>Selective Serotonin Reuptake Inhibitor</td>
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<tr>
<td>TDM</td>
<td>Therapeutic Drug Monitoring</td>
</tr>
<tr>
<td>USP</td>
<td>United States Pharmacopoeia</td>
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<tr>
<td>WHO</td>
<td>World Health Organization</td>
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<tr>
<td>W/V</td>
<td>Weight/Volume</td>
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<td>WW</td>
<td>Weight/Weight</td>
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</table>
What Are Generic Drugs?

A generic drug (generic drugs, short: generics) is a drug defined as "a drug product that is comparable to brand/reference listed drug product in dosage form, strength, route of administration, quality and performance characteristics, and intended use." It has also been defined as a term referring to any drug marketed under its chemical name without advertising.

Generic drugs are usually sold for significantly lower prices than their branded equivalents. One reason for the relatively low price of generic medicines is that competition increases among producers when drugs no longer are protected by patents. Companies incur fewer costs in creating generic drugs (only the cost to manufacture, rather than the entire cost of development and testing) and are therefore able to maintain profitability at a lower price. The prices are low enough for users in many less-prosperous countries to afford them.

Health professionals and consumers can be assured that FDA approved generic drugs have met the same rigid standards as the innovator drug. To gain FDA approval, a generic drug must:

- contain the same active ingredients as the innovator drug (inactive ingredients may vary)
- be identical in strength, dosage form, and route of administration
- have the same use indications
- be bioequivalent
- meet the same batch requirements for identity, strength, purity, and quality
- be manufactured under the same strict standards of FDA's good manufacturing practice regulations required for innovator products

1. US Food and Drug Administration approves the use of generic medicine at public health facility.
2. As per the notification by the government of Chhattisgarh Generic medicine is to be prescribed in all public health facilities.
Guidance on Prescribing & Ethics

1. General guidance and ethics

Drugs should only be prescribed when they are necessary, and in all cases the benefit of administering the medicine should be considered in relation to the risks involved. Bad prescribing habits lead to ineffective and unsafe treatment, exacerbation or prolongation of illness, distress and harm to the patient, and higher cost. The Guide to Good Prescribing (WHO, Geneva; 1994) provides important tools for training in the process of rational prescribing.

This is particularly important during pregnancy, when the risk to both mother and fetus must be considered. It is important to discuss treatment options carefully with the patient to ensure that the patient is content to take the medicine as prescribed. In particular, the patient should be helped to distinguish the adverse effects of prescribed drugs from the effects of the medical disorder.

When the beneficial effects of the medicine are likely to be delayed, the patient should be advised of this.

Taking medicines to best effect

Difficulties in compliance with drug treatment occur regardless of age. Factors contributing to poor compliance with prescribed medicines include:

- prescription not collected or not dispensed;
- purpose of medicine not clear;
- perceived lack of efficacy;
- real or perceived side-effects;
- patients' perception of the risk and severity of side-effects may differ from that of the prescriber;
- instructions for administration not clear;
- physical difficulty in taking medicines (e.g. with swallowing the medicine, with handling small tablets, or with opening medicine containers);
- unattractive formulation (e.g. unpleasant taste);
- Complicated regimen.

The prescriber and the patient should agree on the health outcomes that the patient desires and on the strategy for achieving them ('concordance'). The prescriber should be sensitive to religious, cultural, and personal beliefs that can affect patients' acceptance of medicines. Taking the time to explain to the patient (and relatives) the rationale and the potential adverse effects of treatment may improve compliance. Reinforcement and elaboration of the physician's instructions by the pharmacist also helps. Advising the patient of the possibility of alternative treatments may encourage the patient to seek advice rather than merely abandon unacceptable treatment.

Simplifying the drug regimen may help; the need for frequent administration may reduce compliance, although there appears to be little difference in compliance between once-daily and twice-daily administration. Combination products reduce the number of drugs taken but this may be at the expense of the ability to titrate individual doses.
Health and safety

When handling chemical or biological materials particular attention should be given to the possibility of allergy, fire, explosion, radiation, or poisoning. Substances such as corticosteroids, some antimicrobials, phenothiazines, and many cytotoxics, are irritant or very potent and should be handled with caution. Contact with the skin and inhalation of dust should be avoided.

Safety in the home

Patients must be warned to keep all medicines out of the reach of children. All solid dose and all oral and external liquid preparations must be dispensed in a reclosable child-resistant container unless:

- the medicine is in an original pack or patient pack such as to make this inadvisable;
- the patient will have difficulty in opening a child-resistant container;
- a specific request is made that the product shall not be dispensed in a child-resistant container;
- no suitable child-resistant container exists for a particular liquid preparation.
- All patients should be advised to dispose of unwanted medicines by returning them to a supplier for destruction.

Name of medicine

The name of the medicine should appear on the label unless the prescriber indicates otherwise.

- The strength is also stated on the label in the case of tablets, capsules, and similar preparations that are available in different strengths.
- If it is the wish of the prescriber that a description such as 'The Sedative Tablets' should appear on the label, the prescriber should write the desired description on the prescription form.
- The name written on the label is that used by the prescriber on the prescription.
- When a prescription is written the name of the prescribed preparation will be stated on the label of the dispensed medicine unless the prescriber indicates otherwise.

Prescription writing

Shared care

In its guidelines on responsibility for prescribing between hospitals and general practitioners, the legal responsibility for prescribing lies with the doctor who signs the prescription.

Prescriptions should be written legibly in ink or otherwise so as to be indelible, should be dated, should state the full name and address of the patient, and should be signed in ink by the prescriber. The age and the date of birth of the patient should preferably be stated, and it is a legal requirement in the case of prescription for medicines to state the age for children under 12 years.
General Advice to Prescribers

1. Rational Approach to Therapeutics

Drugs should only be prescribed when they are necessary, and in all cases the benefit of administering the medicine should be considered in relation to the risks involved. Bad prescribing habits lead to ineffective and unsafe treatment, exacerbation or prolongation of illness, distress and harm to the patient, and higher cost. The Guide to Good Prescribing (WHO, Geneva; 1994) provides important tools for training in the process of rational prescribing. The following steps will help prescribers to follow the rational approach to therapeutics.

1 Define the Patient’s Problem
Whenever possible, making the right diagnosis is based on integrating many pieces of information: the complaint as described by the patient; a detailed history; physical examination; laboratory tests; X-rays and other investigations. This will help in rational prescribing, always bearing in mind that diseases are evolutionary processes.

2 Specify the Therapeutic Objective
Doctors must clearly state their therapeutic objectives based on the pathophysiology underlying the clinical situation. Very often physicians are required to select more than one therapeutic goal for each patient.

3 Selecting Therapeutic Strategies
The selected strategy should be agreed with the patient; this agreement on outcome, and how it may be achieved, is termed concordance. The selected treatment can be non-pharmacological and/or pharmacological; it also needs to take into account the total cost of all therapeutic options.

a. Non-Pharmacological Treatment
It is very important to bear in mind that the patient does not always need a medicine for treatment of the condition. Very often, health problems can be resolved by a change in lifestyle or diet, use of physiotherapy or exercise, provision of adequate psychological support, and other non-pharmacological treatments; these have the same importance as a prescription medicine, and instructions must be written, explained and monitored in the same way.

b. Pharmacological Treatment
Selecting the Correct Group of Drugs
Knowledge about the pathophysiology involved in the clinical situation of each patient, pharmacokinetics and pharmacodynamics of the chosen group of drugs, are fundamental principles for rational therapeutics.

Selecting the Medicine from the Chosen Group
The selection process must consider benefit/risk/cost information. This step is based on evidence about maximal clinical benefits of the medicine (efficacy) for a given indication with the minimum production of adverse effects (safety). It must be remembered that each medicine has adverse effects and it is estimated that up to 10% of hospital admissions in industrialized countries are due to adverse effects. Not all medicine-induced injury can be prevented but much of it is caused by inappropriate selection of drugs.
In cost comparison between drugs, the cost of the total treatment and not only the unit cost of the medicine must be considered.

**Verifying the Suitability of the Chosen Pharmaceutical Treatment for Each Patient**
The prescriber must check whether the active substance chosen, its dosage form, standard dosage schedule and standard duration of treatment are suitable for each patient. Medicine treatment should be individualized to the needs of each patient.

**Prescription Writing**
The prescription is the link between the prescriber, the pharmacist (or dispenser) and the patient so it is important for the successful management of the presenting medical condition.

**Giving Information, Instructions and Warnings**
This step is important to ensure patient compliance and is covered in detail in the following chapter (Refer 2.11. Adherence (compliance) with medicine treatment).

**Monitoring Treatment**
Evaluation of the follow up and the outcome of treatment allow the stopping of it (if the patient’s problem is solved) or to reformulate it when necessary. This step gives rise to important information about the effects of drugs contributing to building up the body of knowledge of pharmacovigilance, needed to promote the rational use of drugs.

**2. Factors Affecting Medicine Response**

**2.1. Variation in Dose**
Success and effectiveness of medicine therapy depends not only on the correct choice of medicine but also on the correct dose regimen. Unfortunately, treatment frequently fails because either the dose is too small or it is too large that it produces adverse effects amongst other factors. The concept of a standard or ‘average’ adult dose for every medicine is firmly rooted in the mind of most prescribers. After the initial ‘dose ranging studies on new drugs’, manufacturers recommend a dosage that appears to produce the desired response in the majority of subjects. These studies are usually done on healthy, young male volunteers, rather than on older men and women with illnesses and of different ethnic and environmental backgrounds. The use of standard doses in the marketing literature suggests that standard responses are the rule, but in reality there is considerable variation in medicine response. There are many reasons for this variation such as medicine formulation, body weight and age, variation in pharmacokinetics (absorption, distribution, metabolism and excretion), variation in pharmacodynamics, disease variables, environmental and genetic variables, adherence to instructions and adverse effects and interactions etc. Some of them are described below.

**2.2. Formulation**
The type of drug formulation is an important factor affecting its response, apart from its lipid solubility and so many other factors. Pharmaceutical dosage forms such as tablets, capsules, emulsions, ointments, injectables, liposomes etc provide a mechanism for safe, effective, accurate, and convenient delivery of drugs to the target site. Poorly formulated drugs may fail to disintegrate or dissolve. Enteric-coated drugs are particularly problematic, and have been known to pass through the gastrointestinal tract intact. Some drugs like digoxin or phenytoin have a track record of formulation problems, and dissolution profiles can vary not only from manufacturer to manufacturer but also from batch to batch manufactured by the same
manufacturer. Lately, biogeneric products (off patent biopharmaceuticals) have also been available in the pharmaceutical market. The production of biogenerics involves complex processes.

2.3. Body Weight and Age
Although the concept of varying the dose with the body weight or age of children has a long tradition, adult doses have been assumed to be the same irrespective of size or shape. Yet adult weights vary two to threefold, while a large fat mass can store large excess of highly lipid soluble drugs compared to lean patients of the same weight. Age changes are also important. Adolescents may oxidize some drugs relatively more rapidly than adults, while the elderly may have reduced renal function and eliminate some drugs more slowly.

2.4. Sex
Females usually require smaller doses than males. Iron preparations and other haematinics are exceptions to this rule because of the blood lost by women during menstruation. There is a possibility that males metabolize benzodiazepines, estrogen containing preparations and salicylate at a faster rate than females.

2.5. Route of Administration
It governs the speed and intensity of drug response. The indications for a drug may vary when route of administration varies. Example: Magnesium sulphate when administered orally acts as a purgative, when administered topically- decreases swelling on sprained joints, and when administered intravenously- CNS depression and hypotension occur.

2.6. Tolerance
The therapeutic effects of some medications are lessened in individuals over a prolonged period of use. Thus, a patient who has been using a drug for longer time, requires a higher dose so as to obtain the same therapeutic effect as produced by the drug when taken for the first time. This is called tolerance. Opioids, benzodiazepines, β2 agonists, caffeine, cocaine, amphetamines, and barbiturates fall into this category. Crosstolerance develops when the use of one drug causes a tolerance to another. Alcoholics, barbiturate and narcotic addicts develop a cross-tolerance to sedatives and anaesthetics. These individuals require very large amounts of anaesthetics before surgical anaesthesia can be attained.

2.7. Synergistic Effect
Several drugs when combined may show synergistic action in the form of either additive or supraadditive action or potentiation. A few examples are:
   a) Trimethoprim + Sulphamethoxazole.
   b) ACE inhibitor + Angiotensin Receptor blocker + Diuretic.
   c) Long acting β2 agonists + Inhaled steroids (Example-Salmeterol + Fluticasone)
2.8. Resistance

Development of resistance to drugs is a common problem with antimicrobial agents (antituberculosis drugs, antileprotic drugs, antimalarial drugs etc). Rational prescribing and in turn compliance by the user will prevent the emergence of resistance.

2.9. Pharmacokinetic Variables

2.9.1. Absorption

Absorption of a medicine is possible when it is present in solution form. Medicine absorption rates may vary widely between individuals and in the same individual at different times and in different physiological states. Drugs taken after a meal are delivered to the small intestine much more slowly than in the fasting state, leading to much lower medicine concentrations. In pregnancy gastric emptying is also delayed, while some drugs may increase or decrease gastric emptying and affect absorption of other drugs.

2.9.2 Distribution

Medicine distribution varies widely: fat soluble drugs are stored in adipose tissue, water soluble drugs are distributed chiefly in the extracellular space, acidic drugs bind strongly to plasma protein albumin and basic drugs to muscle cells. Hence variation in plasma albumin levels, fat content or muscle mass may all contribute to dose variation. With very highly albumin bound drugs like warfarin, a small change of albumin concentration can produce a big change in free medicine concentration and a dramatic change in therapeutic action of a medicine.

2.9.3. Metabolism

Medicine metabolic rates are determined both by genetic and environmental factors. Medicine acetylation shows genetic polymorphism, whereby individuals fall clearly into either fast or slow acetylator types. Medicine oxidation, however, is polygenic, and although a small proportion of the population can be classified as very slow oxidizers of some drugs, for most drugs and most subjects there is a normal distribution of medicine metabolizing capacity, and much of the variation is under environmental control. Also refer 2.10.2.

2.9.4. Excretion

Many drugs are eliminated by the kidneys without being metabolized. Renal disease or competitive tubular secretion of drugs can therefore slow down the excretion of certain drugs.

2.10. Pharmacodynamic Variables

There is significant variation in receptor response to some drugs, especially central nervous system responses, for example pain and sedation. Some of this is genetic, some due to tolerance, some due to interactions with other drugs and some due to addiction, for example, morphine and alcohol.

2.10.1. Disease Variables

Both liver and kidney disease can have major effects on medicine response, chiefly by the effect on metabolism and elimination respectively (increasing toxicity), but also by their effect on plasma albumin (increased free medicine also increasing toxicity). Heart failure can also affect metabolism of drugs with rapid hepatic clearance (for example lidocaine, propranolol). Respiratory disease and hypothyroidism can both impair medicine oxidation.
2.10.2. Environmental Factors and Genetic Factors (Pharmacogenetics)

Many drugs and environmental toxins can induce the hepatic microsomal enzyme oxidizing system (MEOS) or cytochrome P450 oxygenases, leading to more rapid metabolism and elimination and ineffective treatment. Environmental pollutants, carcinogens, tobacco smoke, alcohol, anaesthetic drugs and pesticides can also induce metabolism. Diet and nutritional status also have an impact on pharmacokinetics. For example, in infantile malnutrition and in malnourished elderly populations medicine oxidation rates are decreased, while high protein diets, charcoal cooked foods and certain other foods act as metabolizing enzyme inducers. Sedative and hypnotics induce sleep better in calm environment and when administered at night. Pharmacogenetic variation will affect the medicine response, by 4-6 fold among different individuals. All major determinants of medicine response such as transporters, metabolizing enzymes, and receptors are controlled genetically. These factors in certain cases may result in toxicity- for example toxicity caused by inhibitory effect of isoniazid on phenytoin metabolism seems to be more significant in slow acetylators of isoniazid than in those patients who metabolize the drug more rapidly. The Appendix 10 summarizes the pharmacogenetic variation, the frequency of occurrence, drugs involved and the outcome.

2.11. Adherence (Compliance) with Medicine Treatment

It is often assumed that once an appropriate medicine is chosen, the prescription correctly written and the medication correctly dispensed, that it will be taken correctly then the treatment will be successful. Unfortunately this is very often not the case, and physicians overlook one of the most important reasons for treatment failure that is poor adherence (compliance) with the treatment plan. There are sometimes valid reasons for poor adherence. The medicine may be poorly tolerated, may cause obvious adverse effects or may be prescribed in a toxic dose. Failure to adhere with such a prescription has been described as ‘intelligent non-compliance’. Bad prescribing or a dispensing error may also create a problem, and regarding which patients may have neither the insight nor the courage to question. Even with good prescribing, failure to adhere to treatment is common. Factors may be related to the patient, the disease, the doctor, the prescription, the pharmacist or the health system and can often be avoided. Low-cost strategies for improving adherence increase effectiveness of health interventions and reduce costs. Such strategies must be tailored to the individual patient. Health care providers should be familiar with techniques for improving adherence and they should employ systems to assess adherence and to determine what influences it.

2.11.1. Patient Reasons

In general, women tend to be more adherent than men, younger patients and the very elderly are less adherent, and people living alone are less adherent than those with partners or spouses. Specific education interventions have been shown to improve adherence. Patient disadvantages such as illiteracy, poor eyesight or cultural attitudes (for example preference for traditional or alternative drugs and suspicion of modern medicine) may be very important in some individuals or societies, as may economic factors. Such disabilities or attitudes need to be discussed and taken account of.
2.11.2 Disease Reasons
Conditions with a known worse prognosis (for example cancer) or painful conditions (for example rheumatoid arthritis) elicit better adherence rates than asymptomatic ‘perceived as benign’ conditions such as hypertension. Doctors should be aware that in most settings less than half of patients initiated on antihypertensive medicine treatment are still taking it a year later. Similarly, in epilepsy, where events may occur at long intervals, adherence is notoriously unsatisfactory.

2.11.3 Doctor Reasons
Doctors may cause poor adherence in many ways-by failing to inspire confidence in the treatment offered, by giving too little or no explanation, by thoughtlessly prescribing too many drugs, by making errors in prescribing, or by their overall attitude towards the patient.

2.11.4. The Doctor-Patient Interaction
There is considerable evidence that this is crucial to concordance. ‘Satisfaction with the interview’ is one of the best predictors of good adherence. Patients are often well informed and expect a greater say in their health care. If they are in doubt or dissatisfied they may turn to alternative options, including ‘complementary medicine’. There is no doubt that the medicine ‘doctor’ has a powerful effect to encourage confidence and perhaps contribute directly to the healing process.

2.11.5. Prescription Reasons
Many aspects of the prescription may lead to non-adherence (noncompliance). It may be illegible or inaccurate; it may get lost; it may not be refilled as intended or instructed for a chronic disease. Also, the prescription may be too complex; it has been shown that the greater the number of medications the poorer the adherence, while multiple doses also decrease adherence if more than two doses per day are given. Not surprisingly adverse effects like drowsiness, impotence or nausea reduce adherence and patients may not admit to the problem.

2.11.6. Pharmacist Reasons
The pharmacist’s behaviour and professionalism, like the doctor’s, may have a positive impact, supporting adherence, or a negative one, raising suspicions or concerns. This has been reported in relation to generic drugs when substituted for brand-name drugs. Pharmacist information and advice can be a valuable reinforcement, as long as it agrees with the doctor’s advice.

2.11.7. The Healthcare System
The healthcare system may be the biggest hindrance to adherence. Long waiting times, uncaring staff, uncomfortable environment, exhausted medicine supplies and so on, are all common problems in developing countries, and have a major impact on adherence. An important problem is the distance and accessibility of the clinic from the patient. Some studies have confirmed the obvious, that patients farthest from the clinic are least likely to adhere to treatment in the long term.
2.12. Adverse Effects and Interactions

An Adverse Drug Reaction (ADR) may be defined as ‘any response to a medicine which is noxious, unintended and occurs at doses normally used for prophylaxis, diagnosis, or therapy’. ADRs are therefore unwanted or unintended effects of a medicine, including idiosyncratic effects, which occur during its proper use. They differ from accidental to deliberate excessive dosage or medicine maladministration. ADRs may be directly linked to the properties of the medicine in use, the so-called ‘A’ type reactions. An example is hypoglycaemia induced by an antidiabetic medicine. ADRs may also be unrelated to the known pharmacology of the medicine, the ‘B’ type reactions including allergic effects, for example anaphylaxis with penicillins. Thalidomide marked the first recognized public health disaster related to the introduction of a new medicine. It is now recognized that clinical trials, however thorough, cannot be guaranteed to detect all adverse effects likely to be caused by a medicine and hence necessitating post-marketing surveillance. Health workers are thus encouraged to record and report to the National Pharmacovigilance Centre for any unexpected adverse effects with any medicine to achieve faster recognition of serious related problems. The National Regulatory Authority takes appropriate action on drugs showing serious ADRs.

2.12.1. Major Factors Predisposing to Adverse Effects

It is well known that different patients often respond differently to a given treatment regimen. For example, in a sample of 2422 patients who had been taking combinations of drugs known to interact, only 7 (0.3%) showed any clinical evidence of interactions. Therefore, in addition to the pharmaceutical properties of the medicine, the characteristics of the patients may be responsible for causing predisposition to ADRs.

2.12.2. Extremes of Age

The very old and the very young persons are more susceptible to ADRs. Drugs which commonly cause problems in the elderly include hypnotics, diuretics, non-steroidal anti-inflammatory drugs, antihypertensives, psychotropics, digoxin etc. All children, and particularly neonates, differ from adult in their response to drugs. Some drugs are likely to cause problems in neonates (for example morphine), but are generally tolerated in children. Valproic acid is associated with increased risk of ADRs in children of all ages. Other drugs associated with problems in children include chloramphenicol (grey baby syndrome), antiarrhythmics (worsening of arrhythmias), acetylsalicylic acid (Reye’s syndrome etc).

2.12.3. Intercurrent Illness

If besides the condition being treated the patient concomitantly suffers from another disease, such as kidney, liver or heart disease, special precautions may be necessary to prevent ADRs. Remember also that, apart from the above factors, the genetic make-up of the individual patient may also predispose to ADRs.

2.12.4. Drug Interactions

Interactions (see Appendix 6) may occur between drugs which compete for the same receptor or act on the same physiological system. They may also occur indirectly when a medicineinduced disease or a change in fluid or electrolyte balance alters the response to
another medicine. Interactions may occur when one medicine alters the absorption, distribution, metabolism or elimination of another medicine, such that the amount which reaches the site of action is increased or decreased. Medicine-medicine interactions are some of the commonest causes of adverse effects. When two drugs are administered to a patient, they may either act independent of each other, or interact with each other. Interactions may increase or decrease the effects of the drugs concerned and may cause unexpected toxicity. As newer and more potent drugs become available, the number of serious medicine interactions is likely to increase. Remember that interactions which modify the effects of a medicine may involve non-prescription drugs, non-medicinal chemical agents, and social drugs such as alcohol, marijuana, tobacco and traditional remedies, as well as certain types of food. The physiological changes in individual patients, caused by such factors as age and gender, also influence the predisposition to ADRs resulting from medicine interactions.

2.12.5. Pharmaceutical Interactions

Certain drugs, when added to intravenous fluids, may be inactivated by pH changes, by precipitation or by chemical reaction. Benzylpenicillin and ampicillin lose potency after 6-8 hours if added to dextrose solutions, due to the acidity of these solutions. Some drugs bind to plastic containers and tubing, for example diazepam and insulin. Aminoglycosides are incompatible with penicillins and heparin. Hydrocortisone is incompatible with heparin, tetracycline and chloramphenicol.

2.12.6. Adverse Effects Caused by Traditional Drugs

Patients who have been or are taking traditional herbal remedies may develop ADRs. It is not always easy to identify the responsible plant or plant constituent. For further details, refer to the Medicine and Toxicology Information Service if available and/or to suitable literature. Appendix 6d summarises the Drug Herbal/Food Interactions.

2.12.7. The Effect of Food on Medicine Absorption

Food delays gastric emptying and reduces the rate of absorption of many drugs; the total amount of medicine absorbed may or may not be reduced. However, some drugs are preferably taken with food, either to increase absorption or to decrease the irritant effect on the stomach. Appendix 6d summarises the Drug Food Interactions.

Recommendations

• Review the prescription to make sure that it is correct.
• Spend time explaining the health problem and the reason for the medicine.
• Counselling of patients.
• Establish good rapport with the patient.
• Explore problems, for example difficulty with reading the label or getting the prescription filled.
• Encourage patients to bring their medication to the clinic, so that tablet/capsule counts etc. can be done to monitor compliance.
• Encourage patients to learn the names of their drugs, and review their regimen with them. Write notes for them.
• Keep treatment regimens simple.
• Communicate with other health care professionals, to develop a team approach and to collaborate on helping and advising the patient.
• Involve the partner or another family member in eliciting clinical history of the patient and explaining the advice.
• Listen to the patient. Pharmacist plays and important role as a connecting link between the physician and patient.

**Drugs for Anaesthesia**

During the use of Anaesthetics special precautions and close monitoring of the patient are required. These drugs may be fatal if used inappropriately and should be used by non-specialized personnel only as a last resort. Irrespective of whether a general or conduction (regional or local) anaesthetic technique is used; it is essential that facilities for intubation and Mechanically assisted ventilation are available. A full preoperative assessment is required including; if necessary; appropriate fluid replacement.

**Long-Term Medication**

The risk of stopping long-term medication before surgery may be greater than the risk of continuing it. It is essential that the anaesthetist is told of all drugs that the patient is (or has been) taking; in case of oral anticoagulants; corticosteroids; hormonal contraceptives and diabetic patients.

**General Anaesthetics**

**Intravenous Agents:**

Intravenous anaesthetics may be used alone to produce anaesthesia for short surgical procedures but are more commonly used for induction only. They can produce apnoea and hypotension and thus facilities for adequate resuscitation must be available. Before intubation is attempted; a muscle relaxant must be given. Individual requirements vary considerably; lesser dosage is indicated in the elderly; debilitated or hypovolaemic patients.

Intravenous induction using thiopental is rapid and excitement does not usually occur. Anaesthesia persists for about 4–7 min; large or repeated doses severely depress respiration and delay recovery.

Anaesthesia with ketamine persists for up to 15 min after a single intravenous injection and is characterized by profound analgesia. It may be used as the sole agent for diagnostic and minor surgical interventions. Subanaesthetic concentrations of ketamine may be used to provide analgesia for painful procedures of short duration such as the dressing of burns; radiotherapeutic procedures; marrow sampling and minor orthopaedic procedures. Recovery from ketamine anaesthesia is associated with a high incidence of hallucinations and other emergence reactions. Ketamine is of particular value in children; in whom hallucinations are believed to be less significant.

**Volatile Inhalational Agents:**

One of the volatile anaesthetics; ether; halothane (with or without nitrous oxide); must be used for induction when intravenous agents are contraindicated and particularly when intubation is likely to be difficult. Full muscle relaxation is achieved in deep anaesthesia with ether. Excess
bronchial and salivary secretion can be avoided by premedication with atropine. Laryngeal spasm may occur during induction and intubation. Localized capillary bleeding can be troublesome and postoperative nausea and vomiting are frequent; recovery time is slow particularly after prolonged administration.

If intubation is likely to be difficult; halothane is preferred. It does not augment salivary or bronchial secretions and the incidence of postoperative nausea and vomiting is low. Severe hepatitis; which may be fatal; sometimes occurs; it is more likely in patients who are repeatedly anaesthetized with halothane within a short period of time.

Inhalational Gases:
Nitrous oxide is used for the maintenance of anaesthesia. It is too weak to be used alone; but it allows the dosage of other anaesthetic agents to be reduced. It has a strong analgesic action. Oxygen should be added routinely during anaesthesia with inhalational agents; even when air is used as the carrier gas; to protect against hypoxia. Oxygen is also used in the management of anaphylaxis; myocardial infarction and severe acute asthma.
SECTION - 1
ANAESTHETICS

General Anesthetics and Oxygen

Halothane

EDL – D253 Secondary hospital

AVAILABILITY
VOLATILE LIQUID 30, 50, 200, and 250 ml.

DOSE
Induction of anaesthesia using specially calibrated vaporiser; in oxygen or oxygen–nitrous oxide.
Introductory dose: 0.5 to 3%. Maintenance dose: 0.5 to 1.5%. Adult: Increase gradually 2 to 4%.
Child: 1.5 to 2%. Maintenance of anaesthesia using specially calibrated vaporiser; oxygen;
oxigen–nitrous oxide 0.5 to 2%.

INDICATION
Induction and maintenance of anaesthesia

CONTRAINDICATION
History of unexplained jaundice or pyrexia following previous exposure to halothane; family
history of malignant hyperthermia; raised cerebrospinal fluid pressure; porphyria; not
recommended for obstetrical anaesthesia, interactions

PRECAUTION
Anaesthetic history should be carefully taken to determine previous exposure and previous
reactions to halothane (at least 3 months should be allowed to elapse between each re-
exposure); avoid for dental procedures in patients under 18 years unless treated in hospital
(high risk of arrhythmias); pregnancy (Appendix 7c); lactation (Appendix 7b); renal failure;
hyperkalaemia.

ADVERSE EFFECTS
Arrhythmias; bradycardia; respiratory depression; hepatic damage; malignant hyperthermia;
cyanosis; post operative nausea and vomiting.

Isoflurane

EDL – D471 Tertiary

AVAILABILITY
Liquid 100 ml bottle

INDICATION
Same as Halothane

CONTRAINDICATION
Same as Halothane

DRUG INTERACTION
Same as Halothane

ADVERSE EFFECT
Trigger malignant hyperthermia. Since it is an irritant vapour it is less suitable for induction of
anaesthesia especially in children.

DOSE:
Adults induction: inhalation 1.5-3%
MAINTENANCE:
inhaletion 1-3.5%
CHILDREN:
dosage must be individualized.
Sevoflurane  
EDL-D471  
AVAILABILITY 
Liquid 250ml PEN bottle 
INDICATIONS 
Sevoflurane is non-irritant and is therefore often used for inhalational induction of anaesthesia. 
DOSE 
Induction of anaesthesia, using a specifically calibrated vaporiser, in oxygen or nitrous oxide–oxygen, adjusted according to response, ADULT up to 5%; CHILD 1 month–18 years up to 8% . 
Maintenance of anaesthesia, using a specifically calibrated vaporiser, in oxygen or nitrous oxide–oxygen, adjusted according to response, ADULT and CHILD over 1 month 0.5–3% Sevoflurane. 
PRECAUTIONS 
renal impairment; pregnancy ; interactions: (anaesthetics, general) 
CONTRAINDICATIONS 
Susceptibility to malignant hyperthermia Side-effects see notes above; also agitation in children; hepatitis and seizures also reported 

Ketamine hydrochloride  
EDL – D297,298 PHC  
AVAILABILITY 
INJECTIONS 2 ml ampoule (50 mg/ml); 10 ml vial (10 and 50 mg/ml). 
DOSE 
Intravenous injection Short Procedures: Initially 6.5 to 13 mg/kg adjusted according to response (10 mg/kg usually produces 12 to 25 min. of surgical anesthesia). Procedures not involving intense pain: initially 4 mg/kg; (usual dose is 1 to 4.5 mg/ kg). Short procedure over at least 60 min: initially 4 mg/kg (2 mg/kg usually produces 5 to 10 min. of surgical anesthesia). Longer Procedure: induction by intravenous injection using solution containing 1 mg/ml. Longer procedure: induction dose 0.5 to 2 mg/kg; maintenance 10 to 45 mg/kg/min. rate adjusted according to response. 
INDICATION 
Induction and maintenance of anaesthesia; analgesia for painful procedures of short duration especially for patients at the risk of hypotension and branchospasm. 
CONTRAINDICATION 
Thyrotoxicosis; hypertension (including pre-eclampsia); history of cerebrovascular accident; cerebral trauma; intracerebral mass or haemorrhage or other cause of raised intracranial pressure; open eye injury and increased intraocular pressure; psychiatric disorders; particularly hallucinations; hypersensitivity to the drug. 
PRECAUTION 
Supplementary analgesia often required in surgical procedures involving visceral pain pathways (morphine may be used but addition of nitrous oxide will often suffice); during recovery; patient must remain undisturbed but under observation; lactation; children; alcohol intoxicated patients; increased CSF pressure; cardiac decompensation; pregnancy (Appendix 7c). Warn patient not to perform skilled tasks; for example operating machinery or driving; for 24 h and also to avoid alcohol for 24 h. 
ADVERSE EFFECTS 
Hallucinations and other emergence reactions during recovery possibly accompanied by irrational behaviour (effects rarely, persist for more than few hour but can recur at any time
within 24 h); transient elevation of pulse rate and blood pressure common; arrhythmias have occurred; hypotension and bradycardia occasionally reported; confusion; delirium; mobilliform rash; transient erythema; diplopia; increased intraocular pressure; anorexia; nausea; vomiting; local pain and exanthema at injection site; apnoea; laryngospasm.

**Nitrous oxide**

**EDL-D372 Secondary hospital**

**AVAILABILITY**

INHALATIONAL GAS

**DOSE**

Maintenance of anesthesia using suitable equipment up to 66% in oxygen. Analgesic use: 50% in oxygen or according to patient’s need.

**INDICATION**

Maintenance of anaesthesia in combination with other anaesthetic agents (halothane; ether; or ketamine) and muscle relaxants; analgesia for obstetric practice; for emergency management of injuries; during postoperative physiotherapy and for refractory pain in terminal illness.

**CONTRAINDICATION**

Demonstrable collection of air in pleural; pericardial or peritoneal space; intestinal obstruction; occlusion of middle ear; arterial air embolism; decompression sickness; chronic obstructive airway disease; emphysema.

**PRECAUTION**

Minimize exposure of staff; interactions (Appendix 6c); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Nausea and vomiting; after prolonged administration megaloblastic anaemia; depressed white cell formation; peripheral neuropathy.

**Oxygen**

**EDL Category-D392 PHC**

**Indications**

To maintain an adequate oxygen tension in inhalation anaesthesia.

**Availability**

Inhalation gas

**Dose**

(Oxygen is treated as drug since it is prescribed for hypoxemic patient to support alveolar oxygen emergencies).

**Adverse Effect**

Concentration greater than 80% have a toxic effect on the lungs leading to pulmonary congestion; exudation and atelectasis.

**Storage**

Store under pressure to metal cylinder of the type conforming to appropriate safety regulation. Valves and taps should not be lubricated with oil or grease.

**Propofol**

**EDL Category-D439 PHC**

**AVAILABILITY**

INJECTIONS 10, 20 and 50 ml vials (10 mg/ml), 10 and 20 ml vials (20 mg/ml); Ampoule 12, 20 and 50 ml (1%)

**DOSE**

Intravenous Induction and maintenance of general anaesthesia:

Adult: Induction: 40 mg by injection or infusion every 10 seconds. Usual dose: 2-2.5 mg/kg. Maintenance: Infusion- 6-12 mg/kg/h, intermittent bolus injection - 20-50 mg as needed.
Child: >3 years: Induction dose of 2.5-3.5 mg/kg. Maintenance dose: 7.5-18 mg/kg/h by i.v infusion.

Elderly: Including debilitated patients: Infuse at a rate of 20 mg every 10 seconds. Maintenance: 3-6 mg/kg/h.

Sedation: Adult: In diagnostic and surgical procedures: Initially, 6-9 mg/kg/h by infusion given for 3-5 minutes or an alternative dose of 0.5 mg/kg by slow injection over 3-5 minutes. Maintenance: 1.5-4.5 mg/kg/h infusion. Reduce maintenance dose by 20% for high-risk patients needing sedation. For ventilated patients: 0.3 mg/kg/h by infusion, subsequent maintenance dose: 0.3 – 3 mg/kg/h.

INDICATION
Induction and maintenance of general anaesthesia, sedation.

CONTRAINDICATION
Sedation in children and adolescents ≤16 years, Known hypersensitivity to propofol.

PRECAUTION
Cardiac impairment; respiratory impairment; elderly; hypovolaemia; epilepsy; hypotension; patients with high intracranial pressure; monitor blood-lipid concentration if risk of fat overload or if sedation longer than 3 days; hepatic impairment; renal impairment, pregnancy (Appendix 7c), interactions (Appendix 6c).

ADVERSE EFFECTS
Apnoea, bradycardia, arrhythmias, hypotension, anaphylaxis, rash, pruritus, involuntary muscle movements, headache, pain, burning or stinging at injection site.

Thiopental

EDL-D503, 504 Secondary hospitals

INDICATIONS
Induction of anaesthesia prior to administration of inhalational anaesthetic; anaesthesia of short duration. Preferred if cerebral ischaemia is expected.

AVAILABILITY
STERILE POWDER 500 mg and 1g/vial.

DOSE
Slow intravenous injection
3 to 5 mg/kg as 2.5% solution.

Adult- over 18 year fit and premeditated: 10 to 150 mg.
Child- 4 to 7 mg/kg.

CONTRAINDICATIONS
Inability to maintain airway; hypersensitivity to barbiturates; cardiovascular disease; dyspnoea or obstructive respiratory disease; porphyria; hypotension or shock; Addison's disease; hepatic or renal dysfunction; increased blood urea; severe anaemia; asthma; myasthenia gravis.

PRECAUTIONS
Local extravasation can result in extensive tissue necrosis and sloughing; intra-arterial injection causes intense pain and may result in arteriospasm; hepatic impairment (Appendix 7a); interactions (Appendix 6a); pregnancy (Appendix 7c); patients with advanced cardiac disease; increased intracranial pressure; asthma; myasthenia gravis; endocrine insufficiency. Warn patient not to perform skilled tasks; for example operating machinery; driving for 24 h and also to avoid alcohol for 24 h.

ADVERSE EFFECTS
Respiratory depression; myocardial depression; cardiac arrhythmias; somnolence; bronchospasm; urticaria; vasodilation; apnoea; emergence delirium; headache; nausea; oedema.
Local Anaesthetics

Drugs used for conduction anaesthesia (also termed local or regional anaesthesia) act by causing a reversible block to conduction along nerve fibres. Local anaesthetics are used very widely in dental practice; for brief and superficial interventions; for obstetric procedures and for specialized techniques of regional anaesthesia calling for highly developed skills. Facilities and equipment for resuscitation should be readily available at all times. Local anaesthetic injections should be given slowly in order to detect inadvertent intravascular injection. Hypersensitivity testing should be done in all patients before administrations of local anaesthetics.

Local Infiltration

Many simple surgical procedures that neither involve the body cavities nor require muscle relaxation can be performed under local infiltration anaesthesia. Lower-segment caesarean section can also be performed under local infiltration anaesthesia. The local anaesthetic drug of choice is lidocaine 0.5% with or without epinephrine. No more than 4 mg/kg of plain lidocaine or 7 mg/kg of lidocaine with epinephrine should be administered on any one occasion. The addition of epinephrine (adrenaline) diminishes local blood flow; slows the rate of absorption of the local anaesthetic and prolongs its effect. Care is necessary when using epinephrine for this purpose since; in excess; it may produce ischaemic necrosis. It should not be added to injections used in digits or appendages.

Surface Anaesthesia

Topical preparations of lidocaine are available and topical eye drop solutions of tetracaine (chapter 19.2) are used for local anaesthesia of the cornea and conjunctiva.

Regional Block

A regional nerve block can provide safe and effective anaesthesia but its execution requires considerable training and practice. Nevertheless; where the necessary skills are available; techniques such as axillary or ankle blocks can be invaluable. Either lidocaine 1% or bupivacaine 0.5% is suitable. Bupivacaine has the advantage of a longer duration of action.

Spinal Anaesthesia

This is one of the most useful of all anaesthetic techniques and can be used widely for surgery of the abdomen and the lower limbs. It is a major procedure requiring considerable training and practice. Either lidocaine 5% in glucose or bupivacaine 0.5% in glucose can be used but the latter is often chosen because of its longer duration of action.

Bupivacaine hydrochloride

EDL-D80, 81, 82 PHC

AVAILABILITY

Injection 0.25%, 0.5% (hydrochloride) in vial. Injection for spinal anaesthesia: 0.5% (hydrochloride) in 4-ml ampoule to be mixed with 7.5% glucose solution.
DOSE
Adult-Local: Infiltration using 2.5 mg/ml solution (max. 60 ml). Peripheral nerve block: 2.5 mg/ml solution (max. 20 ml) or 5.0 mg/ml solution (max. 30 ml). Epidural block: Lubricant surgery 5 mg/ml solution (max. 20 ml). Sympathetic nerve block: 2.5 mg/ml solution (max. 50 ml).

INDICATION
Infiltration anaesthesia; peripheral and sympathetic nerve block; spinal anaesthesia; postoperative pain relief.

CONTRAINDICATION
Adjacent skin infection; inflamed skin; concomitant anticoagulant therapy; severe anaemia or heart disease; spinal or epidural anaesthesia in dehydrated or hypovolaemic patient.

PRECAUTION
Respiratory impairment; hepatic impairment (Appendix 7a); epilepsy; porphyria; myasthenia gravis; lactation; interactions (Appendix 6c); pregnancy (Appendix 7c).

ADVERSE EFFECTS
With excessive dosage or following intravascular injection; light-headedness; dizziness; blurred vision; restlessness; tremors and occasionally convulsions rapidly followed by drowsiness; unconsciousness and respiratory failure; cardiovascular toxicity includes hypotension; heart block and cardiac arrest; hypersensitivity and allergic reactions also occur; epidural anaesthesia occasionally complicated by urinary retention; faecal incontinence; headache; backache or loss of perineal sensation; transient paraesthesia and paraplegia very rare.

Lidocaine hydrochloride
EDL-D307,308,309,311 PHC D310 Secondary hospitals
INDICATIONS
Surface anaesthesia of mucous membranes; infiltration anaesthesia; peripheral and sympathetic nerve block; dental anaesthesia; spinal anaesthesia; intravenous regional anaesthesia; arrhythmias.

AVAILABILITY
INJECTIONS 30 ml vial (1%) and 10 ml (5%/2 ml) ampoule and 30 ml vial (2%); TOPICAL 30 ml vial (4%); 30 ml vial (lignocaine 20 mg and adrenaline 5 mg/ml); GEL 2% and 4%; Ointment 5% w/w; Spray 15% w/w.
INJECTIONS 2 ml ampoule (Lignocaine HCl 53.3 mg and Dextrose 75 mg/ml) for spinal anaesthesia.

DOSE
Induction of anaesthesia: By injection according to patient weight and nature of procedure. (max. 200 mg lignocaine or 500 mg with adrenaline).
Local application: Rub gently on the affected area.

CONTRAINDICATIONS
Adjacent skin infection; inflamed skin; concomitant anticoagulant therapy; severe anaemia or heart disease; spinal or epidural anaesthesia in dehydrated or hypovolaemic patient; hypersensitivity.

PRECAUTIONS
Respiratory impairment; hepatic impairment (Appendix 7a); epilepsy; porphyria; myasthenia gravis; avoid (or use with great care) solutions containing epinephrine (adrenaline) for ring block of digits or appendages (risk of ischaemic necrosis); lactation; pregnancy (Appendix 7c); interactions (Appendix 6c).

ADVERSE EFFECTS
Same as Bupivacaine (above).
Storage Store in a cool place.
Lidocaine hydrochloride + Epinephrine (adrenaline)  
EDL-D313, 314 Secondary hospitals  
AVAILABILITY  
- injection vial 30 ml (1, 2% w/v), 50 ml (21.3 mg/ml); 2%/50 ml; ampoule 5%/2 ml. JELLY 2% w/v  
OINTMENT 5% w/v  
DOSE  
Adult- Ventricular arrhythmias: loading dose of 50 to 100 mg (or 1 to 1.5 mg/kg) at a rate of 25 to 50 mg/min by intravenous injection, followed immediately by intravenous infusion of 1 to 4 mg/min, with ECG monitoring of all patients (reduce infusion dose if required for longer than 24 h).  
INDICATION  
- Ventricular arrhythmias (especially after myocardial infarction); local anaesthesia.  
CONTRAINDICATION  
- Sino-atrial disorder; any grade of atrioventricular block or any other type of conduction disturbances, severe myocardial depression, acute porphyria or hypovolaemia, bradycardia, cardiac decompensation.  
PRECAUTION  
- Lower dosage in congestive heart failure, bradycardia, ECG monitoring must during therapy, pediatrics; hypotension; renal impairment; porphyria; debilitated patients; hepatic impairment; marked hypoxia; severe respiratory depression; following cardiac surgery and in elderly; lactation; interactions; pregnancy  
ADVERSE EFFECTS  
- Dizziness; paraesthesia; drowsiness, confusion; apnoea, respiratory depression; coma; seizures and convulsions; hypotension, arrhythmias, heart block; cardiovascular collapse and bradycardia (may lead to cardiac arrest); nystagmus often an early sign of lidocaine overdosage; blurred vision, disorientation.  

Glycopyrrolate  
EDL-D248 Secondary hospitals  
INDICATION  
- Same as Atropine  
AVAILABILITY  
- Injection 200 mcg/mL, 1 mL amp, 3 mL amp.  
DOSE  
- For premedication it is given by IM or IV 10 mcg/kg, 200-400 mcg or 4-5 mcg/kg to a maximum of 400 mcg. For children it is given by IM or IV, 4-8 mcg/kg upto a maximum of 200 mcg. For intraoperative use it is given by IV injection as for premedication. For control of muscarinic side effects of neostigmine during reversal of competitive neuromuscular block it is given in a dose of 10 mcg/kg with 50 mcg/kg neostigmine.  
CONTRAINDICATIONS  
- Glaucoma, obstructive uropathy, myasthenia gravis, severe ulcerative colitis.  
PRECAUTIONS  
- Same as Atropine  
ADVERSE EFFECT  
- Same as Atropine  

Preoperative medication  

Pre-anaesthetic medication is often advisable prior to both conduction and general anaesthetic procedures. Sedatives improve the course of subsequent anaesthesia in apprehensive patients. Diazepam and promethazine are effective. Diazepam can be administered by mouth; by
rectum; or by intravenous injection. Promethazine; which has antihistaminic and antiemetic properties as well as a sedative effect; is of particular value in children.

A potent analgesic such as morphine should be administered preoperatively to patients in severe pain or for analgesia during and after surgery.

Anticholinergic (more correctly antimuscarinic) drugs such as atropine are also used before general anaesthesia. They inhibit excessive bronchial and salivary secretions induced; in particular; by ether and ketamine. Intramuscular administration is most effective; but oral administration is more convenient in children. Lower doses should be used in cardiovascular disease or hyperthyroidism.

**Atropine Sulphate**

**EDL-D56 Secondary hospitals**

**AVAILABILITY**

INJECTION 10 ml (0.6 mg/ml).

**DOSE**

Intravenous injection Adult- 0.3 to 0.6 mg immediately before induction of anaesthesia. Intra-operative bradycardia; 300 to 600 μg (longer dose in emergency). Inhibition of bradycardia; 0.4 to 1 mg. Reversal of neuromuscular block; 0.6 to 1.2 mg.

Child- Premedication: 20 μg/kg; Inhibition of bradycardia: 10 to 30 μg/kg. Reversal of neuromuscular block: 20 μg/kg.

Intramuscular route or subcutaneous

Premedication (30 to 60 min before induction of anaesthesia): 300 to 600 μg.

Child- 20 μg/kg (max. 60 μg). Intra operative bradycardia: (1 to 12 years) 10 to 20 μg/kg.

**INDICATION**

To inhibit salivary secretions; to inhibit arrhythmias resulting from excessive vagal stimulation; to block the parasympathomimetic effects of anticholinesterases such as neostigmine; organophosphate poisoning; antispasmodic; mydriasis and cycloplegia.

**CONTRAINDICATION**

Angle-closure glaucoma; myasthenia gravis; paralytic ileus; pyloric stenosis; prostatic enlargement.

**PRECAUTION**

colitis; diarrhoea; hyperthyroidism; heart failure; hypertension; patients with atrial fibrillation or flutter; lactation (Appendix 7b); interactions (Appendix 6a); pregnancy (Appendix 7c). Since atropine has a shorter duration of action than neostigmine; late unopposed bradycardia may result; close monitoring of the patient is necessary.

**ADVERSE EFFECTS**

Dry mouth; blurred vision; photophobia; flushing and dryness of skin; rash; difficulty in micturition; less commonly arrhythmias; tachycardia; palpitations; confusion (particularly in elderly); heat prostration and convulsions; ventricular fibrillation; hallucinations; dilated pupils; psychosis.

**Diazepam**

**EDL-D164, 165 PHC**

**AVAILABILITY**

TABLETS 2, 5 and 10 mg; CAPSULE 10 mg; INJECTION 2 ml ampoule (5 mg/ml).
DOSE
Adult- 5 mg on night before surgery or minor procedure; thereafter 5 mg for 2h before procedures. Elderly- Half of adult dose. Intravenous injection 10 to 20 mg over 2 to 4 min immediately before procedure. Premedication: 100 to 200 μg/kg. Child- 2.5 to 10 mg over 2 to 4 min. 0.1 to 0.3 mg/kg in divided doses over 24 h; every 4 to 8 h (adjust according to response).

INDICATION
Premedication before major or minor surgeries; sedation with amnesia for endoscopic procedures and surgeries under local anaesthesia; emergency reduction of fractures (in combination with pethidine when anaesthetics are not available); epilepsy; anxiety disorders.

CONTRAINDICATION
Respiratory depression; acute pulmonary insufficiency; sleep apnoea; severe hepatic impairment; myasthenia gravis; avoid injections containing benzyl alcohol in neonates, narrow angle glaucoma; hypersensitivity to benzodiazepine.

PRECAUTION
history of alcohol or drug abuse; marked personality disorder; elderly or debilitated patients (adverse effects more common in these groups); hepatic impairment (Appendix 7a) or renal failure; lactation (monitoring for adverse effects required Appendix 7b); porphyria; interactions (Appendix 6a, 6c); organic cerebral changes; epileptic patients. Warn patient not to perform skilled tasks; for example operating machinery; driving for 24 h.

ADVERSE EFFECTS
Drowsiness and lightheadedness the next day; confusion and ataxia (especially in the elderly); amnesia; dependence; paradoxical increase in aggression; muscle weakness; occasionally headache, vertigo, salivation changes, gastrointestinal disturbances, skin reactions, visual disturbances, dysarthria, tremors, incontinence, urinary retention; blood disorders and jaundice; hypotension and apnoea, pain and thrombophlebitis (with injection); increased appetite; weight gain.

Morphine (Sulphate or hydrochloride)
EDL-D358 Secondary hospitals

AVAILABILITY
INJECTION 10 ml ampoule (1 mg/ml, 10 mg/ml and 15 mg/ml); TABLETS 10, 20, 30 and 60 mg.

DOSE
Subcutaneous or intramuscular injection Adult- Preoperative medication before procedure: up to 10 mg; 60 to 90 min before procedure; 20 to 30 mg per 12 h depending on patient weight. Postoperative analgesia: 150 to 300 μg/kg every 4 h. Child- (By intramuscular injection) Preoperative medication before procedure: 150 μg/kg. Postoperative analgesia: 100 to 200 μg/kg. Intravenous injection

INDICATION
In severe pain (acute and chronic); myocardial infarction; acute pulmonary oedema; adjunct during major surgery and postoperative analgesia.

CONTRAINDICATION
Patients with acute respiratory depression and when there is risk of paralytic ileus; conditions associated with raised intracranial pressure and in head injury (they interfere with pupillary responses vital for neurological assessment); comatose patients; acute asthma; acute liver disease; acute alcoholism; pulmonary oedema; interactions; lactation; hepatic impairment

PRECAUTION
Patients with impaired respiratory function (avoid in chronic obstructive pulmonary disease) and asthma (avoid during an acute attack); hypotension; myasthenia gravis; prostatic hypertrophy
and hyperplasia; obstructive or inflammatory bowel disorders; disease of the biliary tract and convulsive disorders; pancreatitis; cardiac arrhythmias; hypothyroidism; head injury; circulatory shock; lactation; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea and vomiting (particularly in initial stages); constipation; dry mouth and biliary spasm; larger doses produce muscle rigidity; hypotension and respiratory depression; bradycardia; paralytic ileus; abdominal pain; anorexia; dyspepsia; exacerbation of pancreatitis; taste disturbance; hypertension; hypothermia; syncope; bronchospasm; inhibition of cough reflex; restlessness; seizures; paraesthesia; asthenia; malaise; disorientation; excitation; agitation; delirium; raised intracranial pressure; amenorrhoea; myoclonus; muscle fasciculation and rhabdomyolysis.

Promethazine Hydrochloride
EDL-D437 PHC
Premedication prior to surgery; antiemetic.

AVAILABILITY
TABLETS 10 and 25 mg; INJECTION 2 ml ampoule (25 mg/ml).

DOSE
Oral
Adult- Premedication: 25 mg at night and increase to 25 mg twice daily; if necessary; alternately 10 to 20 mg 2 to 3 times daily.
Child- 2 to 5 years: not recommended. 5 to 10 years: 20 to 25 mg.

Deep intramuscular route
Adult- 50 mg (max. 100 mg). Premedication: 25 to 60 mg 1 h before operation.
Child- 5 to 10 years: 6.25 to 12.5 mg. Premedication, 5 to 10 years: 6.5 to 12.5 mg.

Slow intravenous injection
In emergencies: 25 to 50 mg as solution containing 2.5 mg/ml in water for injection (max. 100 mg).

CONTRAINDICATIONS
Child under 2 year; impaired consciousness due to cerebral depressants or of other origin; porphyria.

PRECAUTIONS
Prostatic hypertrophy; urinary retention; glaucoma; epilepsy; hepatic impairment (Appendix 7a); lactation (Appendix 7b); interactions (Appendix 6a); pregnancy (Appendix 7c). Warn patient not to perform skilled tasks; for example operating machinery, driving for 24 h.

ADVERSE EFFECTS
Drowsiness (rarely, paradoxical stimulation in children); headache; anticholinergic effects such as dry mouth; blurred vision; urinary retention.

STORAGE
Store protected from light and moisture

Fentanyl citrate
EDL-D217 Tertiary

INDICATIONS
analgesia during operation, enhancement of anaesthesia; respiratory depressant in assisted respiration; analgesia in other situations.

DOSE
by intravenous injection, with spontaneous respiration, 50-200 micro-grams, and then 50 micrograms as required. CHILD 3-5 microgram/kg, and then 1 microgram/kg as required With assisted ventilation, 0.3-3.5 mg, then 100-200 micrograms as required. CHILD 15 micrograms/kg, then 1-3 micrograms/kg as required.
CONTRAINDICATIONS

Known hypersensitivity or intolerance to fentanyl or other opioid analgesics, Bronchial asthma, Head injuries and increased intracranial pressure. As for any opioid analgesic, Fentanyl should not be used in patients susceptible to respiratory depression, such as comatose patients who may have head injuries or a brain tumour. Fentanyl may obscure the clinical course of patients with head injury, Concomitant MAO inhibitors. Severe and unpredictable potentiation by MAO inhibitors has been reported with opioid analgesics and the use of Fentanyl in patients who have received MAO inhibitors within 14 days is not recommended. Myasthenia gravis. Fentanyl may cause muscle rigidity upon IV administration. Therefore, the need for reversal and muscle relaxants contraindicates its use in patients with a history of myasthenia gravis. Children two years of age or younger. Safe conditions for use have not been established.

PRECAUTIONS

It should be given with care since the respiratory depression can persist into the post-operative period and occasionally it may become apparent for the first time postoperatively when monitoring of the patient might be less intensive.

ADVERSE EFFECT

Its overdose may cause Narcosis (which may be preceded by marked skeletal muscle rigidity), cardiorespiratory depression accompanied by cyanosis, followed by a fall in body temperature, circulatory collapse, coma and possibly death.

Midazolam

EDL-D350 Tertiary

AVAILABILITY

INJECTION 1 ml ampoule (5 mg/ml); 5 and 10 ml vial (1 mg/ml).

DOSE

Slow intravenous injection Adult- Conscious sedation: approximately 2 mg/min; 5 to 10 min before procedure; initially 2 to 2.5 mg. Usual total dose 3.5 to 5 mg (Max. 7.5 mg). Elderly- 0.5 to 1.0 mg. Increase if necessary in steps of 1 mg. Intravenous injection (Over 2 to 3 min) Child- 6 months to 7 years: initially 50 to 100 μg/kg; increase if necessary in steps (max. total dose 6.0 mg). 6 to 12 years: initially 25 to 50 μg/kg increase in steps if necessary (max. total dose 10 mg). Intramuscular injection Adult- Sedation in combined anaesthesia: 30 to 100 μg/kg repeated as required by continuous intravenous infusion 30 to 100 μg/ kg/h (lower doses in elderly). Premedication: 70 to 100 μg/kg. 1 to 15 years: 50 to 150 μg/kg (max.1 mg). Elderly and debilitated- 25 to 50 μg/kg. (20 to 60 min induction).

INDICATION

Intravenous sedative administered before or during minor surgical procedures; sedative administered by intravenous route in intensive care induction of anaesthesia.

CONTRAINDICATION

Acute narrow angle glaucoma; comatose patients; shock; acute alcohol intoxication; for intrathecal and epidural use; acute pulmonary insufficiency; myasthenia gravis.

PRECAUTION

Chronic renal failure; cardiac disease; open angle glaucoma; respiratory disorders; neonates; prolonged use and abrupt withdrawal should be avoided; hepatic impairment; pregnancy (Appendix 7c) and lactation; interactions (Appendix 6a, 6c).

ADVERSE EFFECTS

Hypersensitivity; cardiac arrest; laryngospasm; apnoea; headache; hiccups; nausea; vomiting; cough; kernicterus; nystagmus; skin rash; CNS symptoms like euphoria; hallucination; ataxia.
SECTION - 2
ANALGESICS, ANTIPYRETICS, NONSTEROIDAL ANTI INFLAMMATORY MEDICINES, MEDICINES USED TO TREAT GOUT AND DISEASE MODIFYING AGENTS USED IN RHEUMATOID DISORDERS

Analgesics, Antipyretics, Non-Steroidal Anti-Inflammatory Drugs

Analgesics are used to relieve/reduce body pain and antipyretics are used to reduce elevated body temperature. Nonopioid analgesics are particularly suitable for relieving or management of pain in musculoskeletal conditions whereas the opioid analgesics are more suitable for moderate to severe visceral pain. Those non-opioid analgesics which also have anti-inflammatory actions include salicylates and NSAIDs; they can reduce both pain and inflammation of chronic inflammatory disorders such as rheumatoid arthritis, but they do not alter or modify the disease process itself. For the management of rheumatoid arthritis, DMARDs (disease-modifying antirheumatic drugs) may favourably influence the outcome of the disease. The pain and inflammation of an acute attack of gout is treated with a NSAID or colchicine; a xanthine oxidase inhibitor is used for long-term control of gout. Neurogenic pain generally responds poorly to conventional analgesics; treatment can be difficult and includes the use of carbamazepine for trigeminal neuralgia and amitriptyline for diabetic neuropathy and post-therapeutic neuralgia.

Non-Opioid, Non-Steroidal Anti-Inflammatory Drugs

Non-opioid analgesics with anti-inflammatory activity include salicylates such as acetylsalicylic acid and other nonsteroidal anti-inflammatory drugs such as ibuprofen. Non-opioid analgesics with little or no anti-inflammatory activity include paracetamol.

**Acetyl Salicylic Acid (ASA)**

**EDL-D2,3,4 PHC**

**AVAILABILITY**

TABLETS 50, 60, 75, 80, 150, 300 and 325 mg.

**DOSE**

Oral Adult- Analgesic and antipyretic including migraine attacks: 0.3 to 0.9g, 3 to 4 times a day (max. 4g daily). Acute Rheumatic fever: 4 to 6g or 75 to 100 mg/kg daily in divided doses. Antiplatelet: 75-325 mg/day. Child- Under 16 years: not recommended (can cause Reye’s syndrome).C3

**INDICATION**

Management of mild to moderate pain such as headache, acute migraine attacks, transient musculoskeletal pain, dysmenorrhoeal pain and for reducing fever; pain and inflammation of rheumatoid arthritis; antiplatelet agent for prophylaxis of myocardial infarction, stable angina pectoris; stroke prophylaxis.
CONTRAINDICATION
Hypersensitivity (including asthma; angioedema; urticaria or rhinitis) to acetylsalicylic acid or any other NSAID; children and adolescents under 16 years (may cause Reye’s syndrome); gastrointestinal ulceration; haemophilia and other bleeding disorders; not for treatment of gout; severe renal or hepatic impairment; lactation. It is known to cause haemolytic anaemia in people who have the genetic disease- G-6-PD-deficiency.

PRECAUTION
Asthma, allergic disease; impaired renal or hepatic function (Appendices 7d and 7a); lactation (Appendix 7b); pregnancy (Appendix 7c); elderly; G-6-PD-deficiency; dehydration; interactions (Appendix 6a, 6c, 6d).

ADVERSE EFFECTS
Bronchospasm; gastrointestinal haemorrhage (rarely, major); also other haemorrhage (for example subconjunctival); urticaria; hepatomegaly

Ibuprofen
EDL-D271,272,273 Primary

AVAILABILITY
TABLETS 200, 400 and 600 mg; Capsules 400 mg Plain, 300 mg SR; SUSPENSION 100 mg/5 ml.

DOSE
Child- 1-6 months: initially by intravenous injection (over atleast 5 min) 100-200 μg/kg then by continous infusion 10-30 μg/h. adjusted according to response. 6 months-12 years: initially by intravenous injection (over atleast 5 min) 100-200 μg/kg, adjusted according to response. Juvenile rheumatoid arthritis: 20 to 40 mg/ kg/day in 3 to 4 divided doses.

INDICATION
Pain and inflammation in rheumatic disease and other musculoskeletal disorders including juvenile arthritis; mild to moderate pain including dysmenorrhoeal pain, headache; pain in children; acute migraine attack.

CONTRAINDICATION
Hypersensitivity (including asthma; angioedema; urticaria or rhinitis) to acetylsalicylic acid or any other NSAID; active peptic ulceration; for treatment of pre-operative pain in the setting of coronary artery bypass graft surgery; neonates with congenital heart disease.

PRECAUTION
Renal and hepatic impairment (Appendix 7a); preferably avoid if history of peptic ulceration; cardiac disease; elderly; pregnancy (Appendix 7c); lactation (Appendix 7b); coagulation defects; allergic disorders; interactions (Appendix 6a, 6c, 6d)

ADVERSE EFFECTS
Gastrointestinal disturbances including nausea, diarrhoea, dyspepsia, gastrointestinal haemorrhage; hypersensitivity reactions including rash, angioedema; bronchospasm; headache; dizziness; nervousness; depression; drowsiness; insomnia; vertigo; tinnitus; photosensitivity; haematuria; renal failure; fluid retention (rarely, precipitating congestive heart failure in elderly), raised blood pressure; rarely, hepatic damage; alveolitis, pulmonary eosinophilia; pancreatitis; visual disturbances; erythema multiforme (Stevens- Johnson syndrome); toxic dermal necrolysis (Lyell’s syndrome); colitis; aseptic meningitis. Skin reactions like dermatitis.

Mefenamic Acid
Non-EDL Tertiary

INDICATIONS
Treatment of rheumatoid arthritis, osteoarthritis, dysmenorrhea, mild to moderate pain, inflammation, fever dental pain.
Availability
TABLETS 100 mg, 250 mg, 500 mg. CAPSULES 250 mg. SUSPENSION 50 mg/5 ml.

DOSE
Adult
Pain: 500 mg orally, followed by 250 mg every 6 hours as needed, not to exceed 7 days.
Dysmenorrhea: 500 mg orally, followed by 250 mg every 6 hours starting with the onset of menses.

Children
Pain: 14 to 18 years: 500 mg orally followed by 250 mg every 6 hours as needed, not to exceed 7 days.

CONTRAINDICATIONS
Known hypersensitivity to mefenamic acid; patients who have experienced asthma, urticaria, or allergic-type reactions after taking aspirin or other NSAIDs; peri-operative pain in the setting of coronary artery bypass graft (CABG) surgery, active ulceration or chronic inflammation of the gastrointestinal tract, pre-existing renal disease, pregnancy, interactions.

PRECAUTIONS
Hepatic effects: Borderline elevations of one or more liver function tests may occur. These laboratory abnormalities may progress, may remain unchanged, or may be transient with continuing therapy. A patient with symptoms and/or signs suggesting liver dysfunction, or in whom an abnormal liver test has occurred, should be evaluated for evidence of the development of a more severe hepatic reaction while on therapy. If clinical signs and symptoms consistent with liver disease develop, or if systemic manifestations occur (e.g., eosinophilia, rash, etc.), the drug should be discontinued.
Anaemia: Patients on long-term treatment should have their hemoglobin or hematocrit checked if they exhibit any signs or symptoms of anaemia.
Asthma: Mefenamic acid should not be administered to patients with aspirin sensitive asthma and should be used with caution in patients with preexisting asthma.

ADVERSE EFFECTS
Gastrointestinal experiences including abdominal pain, constipation, diarrhoea, dyspepsia, flatulence, gross bleeding/ perforation, heartburn, nausea, gastrointestinal ulcers, vomiting, abnormal renal function, bronchospasm, anaemia, dizziness, edema, elevated liver enzymes, headaches, increased bleeding time, pruritus, rashes, tinnitus.

Storage
Store protected from light and moisture.

Paracetamol
EDL- D395,396 Universal D397 PHC

AVAILABILITY
TABLETS 500 and 650 mg Plain; 750 mg DT; SYRUPS/SUSPENSION 125 and 250 mg/5 ml;
INJECTION 2 ml ampoule 125 mg/ml.; Intravenous infusion 500 mg and 1g.

DOSE
Oral Adult- 0.5 to 1g every 4 to 6 h (max. 4g, max 2g in alcoholics per day). Child- for post-immunisation pyrexia, up to 2 months: 60 mg. 3 month to 1 year: 60 to 120 mg every 4 to 6 h. 1 to 5 years: 120 to 250 mg every 4 to 6 h. 6 to 12 years: 250 to 500 mg every 4 to 6 h. Intramuscular injection Adult- 250 mg every 4 to 6 h or as required. Intravenous infusion Adult- 1g every 6 hours, maximum daily dose 4 g. Child- 15 mg/kg upto 4 times a day, maximum daily dose 60 mg/kg.
**INDICATION**
Mild to moderate pain including dysmenorrhoeal pain, headache; pain relief in osteoarthritis and soft tissue lesions; pyrexia including post-immunisation pyrexia; acute migraine attack

**PRECAUTION**
Hepatic impairment (Appendix 7a); renal impairment; alcohol dependence; lactation (Appendix 7b); pregnancy (Appendix 7c); overdosage: chapter 7.2; interactions (Appendix 6a); G-6-PD deficiency.

**ADVERSE EFFECTS**
Rare but rashes and blood disorders reported; important: liver damage (and less frequently renal damage) following overdosage; dyspepsia.

**Diclofenac sodium**
**EDL-D168,170,171 PHC**

**AVAILABILITY**
TABLETS 25 and 50 mg Plain; 75 and 100 mg CR; CAPSULES 100 mg, 100 mg CR; INJECTION 3 ml ampoule (25 mg/ml); Eye/Ear Drops 0.1% w/v; Suppositories 25, 50 and 100 mg; Gel 1%w/w.

**DOSE**
Oral 100 to 150 mg daily in 2 to 3 divided doses, (max 150 mg/day) maintenance by 50 to 100 mg in divided doses. Intramuscular injection 75 mg, 2 to 3 times daily. Topically Adult- Apply 1% w/w gel on to affected area 3 to 4 times daily. Instill to eye Post-operative ocular inflammation: Adult- as sodium (1% w/v), 4 times daily starting 24 h after surgery for up to 28 days. Rectal Post-operative pain. Adult- 75 to 150 mg daily in divided doses (max. 150 mg/day, inclusive of diclofenac administered through other routes). Child- 6 to 12 year: 1 to 2 mg/kg/day in divided doses for max. of 4 days.

**INDICATION**
Acute musculo-skeletal pain; arthritis; gout; spondylitis; migraine; post-operative pain

**CONTRAINDICATION**
Porphyria; avoid injections containing benzyl alcohol in neonates; history of gastric ulcers, bleeding or perforation. Additional contraindications include concomitant NSAID or anticoagulant use (including low-dose heparin); history of haemorrhagic diathesis; history of confirmed or suspected cerebrovascular bleeding; operations with high risk of haemorrhage; history of asthma; moderate or severe renal impairment; hypovolaemia; dehydration.

**PRECAUTION**
NSAIDs should be used with caution in the elderly (risk of serious side-effects and fatalities); interactions (Appendix 6a, 6c, 6d); pregnancy (Appendix 7c); patients with coagulation disorders; hepatic, renal and cardiac impairment; history of gastrointestinal lesions.

**ADVERSE EFFECTS**
Injection site reactions; transient epigastric pain, risk of thrombotic events; toxic epidermal necrolysis; Abnormality in kidney function.

**Diclofenac Potassium**
**EDL-D608 PHC**

**AVAILABILITY**
TABLETS 25 and 50 mg Plain; 75 and 100 mg CR; CAPSULES 100 mg, 100 mg CR; INJECTION 3 ml ampoule (25 mg/ml); Eye/Ear Drops 0.1% w/v; Suppositories 25, 50 and 100 mg; Gel 1%w/w.

**DOSE**
Oral 100 to 150 mg daily in 2 to 3 divided doses, (max 150 mg/day) maintenance by 50 to 100 mg in divided doses. Intramuscular injection 75 mg, 2 to 3 times daily. Topically Adult- Apply 1% w/w gel on to affected area 3 to 4 times daily. Instill to eye Post-operative ocular inflammation: Adult- as sodium (1% w/v), 4 times daily starting 24 h after surgery for up to 28 days. Rectal Post-operative pain. Adult- 75 to 150 mg daily in divided doses (max. 150 mg/day, inclusive of
diclofenac administered through other routes). Child- 6 to 12 year: 1 to 2 mg/kg/day in divided doses for max. of 4 days.

INDICATION
Acute musculo-skeletal pain; arthritis; gout; spondylitis; migraine; post-operative pain

CONTRAINDICATION
Porphyria; avoid injections containing benzyl alcohol in neonates; history of gastric ulcers, bleeding or perforation. Additional contraindications include concomitant NSAID or anticoagulant use (including low-dose heparin); history of haemorrhagic diathesis; history of confirmed or suspected cerebrovascular bleeding; operations with high risk of haemorrhage; history of asthma; moderate or severe renal impairment; hypovolaemia; dehydration.

PRECAUTION
NSAIDs should be used with caution in the elderly (risk of serious side-effects and fatalities); interactions (Appendix 6a, 6c, 6d); pregnancy (Appendix 7c); patients with coagulation disorders; hepatic, renal and cardiac impairment; history of gastrointestinal lesions.

ADVERSE EFFECTS
Injection site reactions; transient epigastric pain, risk of thrombotic events; toxic epidermal necrolysis; Abnormality in kidney function.

KETOROLAC
EDL-D675, 676 Secondary hospitals

INDICATION
Short term management of moderate to severe acute postoperative pain. Prophylaxis and reduction of inflammation and associated symptoms following ocular surgery

CONTRAINDICATION
History of hypersensitivity to aspirin or any other NSAIDs or to any ingredients of the formulation; children below 3 years; asthma, angioedema or bronchospasm, history of peptic ulcer; moderate to severe renal impairment, coagulation disorders, pregnancy and lactation.

PRECAUTIONS
Asthma, GI diseases, renal or hepatic disorder, allergy, haemostasis, children below 16 years. There is a potential for cross sensitivity to aspirin, phenylacetic acid derivatives and other NSAIDs, hence caution should be used when treating individuals who have previously exhibited sensitivities to these drugs; bleeding disorders.

ADVERSE EFFECT
Anaphylaxis; fluid retention, nausea, dyspepsia, abdominal discomfort, bowel changes, peptic ulceration; GI bleeding (elderly at greater risk), convulsions, myalgia, aseptic meningitis, hyponatraemia, hyperkalaemia, A raised blood urea and creatinine, urinary symptoms and acute renal failure, flushing or pallor, bradycardia, hypertension, purpura, thrombocytopenia, dyspnoea and pulmonary oedema, skin reactions (Stevens-Johnson & Lyell’s syndromes), post operative wound haemorrhage, haematoma, epistaxis, oedema, liver function changes. Theoretical risk of prolonged bleeding time, transient stinging and blurring of eyes on instillation

AVAILABILITY
Film coated tablets, 10mg; Injection, 30mg/mL, mL ampoules. Ophthalmic solution 0.5% w/v, 5ml.

DOSE
ADULT: Oral: 10mg every, 4-6 hours (elderly every 6-8 hours); max. 40mg daily, max. duration of treatment 7 days. I.M. or IM initially 10mg, then 10-30mg every 4-6 hours up to a max. of 90mg daily. ADULT: Instill 1 drop 3 times daily starting 24 hours pre-operatively and continuing for up to 3 weeks. CHILD: Not recommended under 16 years.
Opioid Analgesic

Morphine is effective in relieving moderate to severe pain, particularly of visceral origin; there is a large variation in patient response. Weaker opioids such as codeine are suitable for mild to moderate pain. Morphine remains the most valuable analgesic for severe pain. In addition to pain relief it confers a state of euphoria and mental detachment; repeated administration may cause dependence and tolerance, but this should not be a deterrent in the control of pain in terminal illness. Regular use may also be appropriate for certain cases of non-malignant pain, but specialist supervision is required. In normal doses common adverse effects include nausea, vomiting, constipation and drowsiness; larger doses produce respiratory depression and hypotension.

Codeine is an opioid analgesic much less potent than morphine and much less liable, in normal doses, to produce adverse effects including dependency. It is effective for mild to moderate pain but is too constipating for long-term use.

Pentazocine

EDL-D398 Secondary hospitals

AVAILABILITY

TABLETS 25 mg Plain, Combination: Paracetamol 500 mg + Pentazocine 15 mg; INJECTION 1 ml ampoule (30 mg/ml).

DOSE

Oral Adult- Pentazocine 50 mg every 3 to 4 h preferably after food (range 25 to 100 mg, max. 600 mg daily). Child- 6 to 12 years: 25 mg. Subcutaneous, intramuscular or intravenous injection

Adult- Moderate pain: 30 mg. Severe pain: 45 to 60 mg every 3 h to 4 h when necessary.

INDICATION

Moderate to severe pain; pre-anaesthetic medication; colic; trauma; surgical procedures; burns.

CONTRAINDICATION

Patients dependent on opioids; arterial or pulmonary hypertension; heart failure; narcotic dependence; hypersensitivity; ischaemia; myocardial infarction.

PRECAUTION

Avoid in porphyria; interactions (Appendix 6a); impaired respiratory function; pregnancy(Appendix 7c) ; renal or hepatic function; thyroid dysfunction; biliary tract impairment.

ADVERSE EFFECTS

Avoid in porphyria; interactions; impaired respiratory function; pregnancy ; renal or hepatic function; thyroid dysfunction; biliary tract impairment.

Tramadol

EDL-D753, 754 Secondary hospitals

AVAILABILITY

TABLETS 50 mg and 100 mg SR; CAPSULE 50 and 100 mg SR; INJECTION 1 and 2 ml ampoule (50 mg/ml).

DOSE

Adult- Moderate to severe pain: 50 to 100 mg, 4 to 6 hourly (max 400 mg/day). Post operative pain: 100 mg i.v. initially followed by 50 mg every 10 to 20 min upto max. of 250 mg in the 1st h. Maintenance dose 50 to 100 mg, 4 to 6 hourly (max 600 mg/day
INDICATION
Moderate or severe pain, post operative pain, in patients contraindicated to NSAIDs.

CONTRAINDICATION
Patients with suicidal tendency; raised intracranial pressure; severe renal impairment; acute alcoholism; lactation.

PRECAUTION
Renal or hepatic impairment; history of epilepsy; inflammatory or obstructive bowel disease; myasthenia gravis; hypothyroidism; adreno-cortical insufficiency; respiratory depression; prostatic hyperplasia; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Same as other opioids, however it has less addictive potential.

Morphine (Sulphate or hydrochloride)
EDL-D358 Tertiary

AVAILABILITY
INJECTION 10 ml ampoule (1 mg/ml, 10 mg/ml and 15 mg/ml); TABLETS 10, 20, 30 and 60 mg.

DOSE
Subcutaneous or intramuscular injection
Adult- Preoperative medication before procedure: up to 10 mg; 60 to 90 min before procedure; 20 to 30 mg per 12 h depending on patient weight. Postoperative analgesia: 150 to 300 μg/kg every 4 h. Child- (By intramuscular injection) Preoperative medication before procedure: 150 μg/kg. Postoperative analgesia: 100 to 200 μg/kg. Intravenous injection

INDICATION
In severe pain (acute and chronic); myocardial infarction; acute pulmonary oedema; adjunct during major surgery and postoperative analgesia.

CONTRAINDICATION
Patients with acute respiratory depression and when there is risk of paralytic ileus; conditions associated with raised intracranial pressure and in head injury (they interfere with pupillary responses vital for neurological assessment); comatose patients; acute asthma; acute liver disease; acute alcoholism; pulmonary oedema; interactions; lactation; hepatic impairment.

PRECAUTION
Patients with impaired respiratory function (avoid in chronic obstructive pulmonary disease) and asthma (avoid during an acute attack); hypotension; myasthenia gravis; prostatic hypertrophy and hyperplasia; obstructive or inflammatory bowel disorders; disease of the biliary tract and convulsive disorders; pancreatitis; cardiac arrhythmias; hypothyroidism; head injury; circulatory shock; lactation; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea and vomiting (particularly in initial stages); constipation; dry mouth and biliary spasm; larger doses produce muscle rigidity; hypotension and respiratory depression; bradycardia; paralytic ileus; abdominal pain; anorexia; dyspepsia; exacerbation of pancreatitis; taste disturbance; hypertension; hypothermia; syncope; bronchospasm; inhibition of cough reflex; restlessness; seizures; paraesthesia; asthenia; malaise; disorientation; excitation; agitation; delirium; raised intracranial pressure; amenorrhoea; myoclonus; muscle fasciculation and rhabdomyolysis.
Medicine used to treat gout

Acute Gout:
Acute attacks of gout are usually treated with high doses of a NSAID such as indomethacin (150-200 mg daily in divided doses); ibuprofen has weaker anti-inflammatory properties than other NSAIDs and is therefore less suitable for treatment of gout. Salicylates, including acetylsalicylic acid are also not suitable because they may increase plasma-urate concentrations. Colchicine is an alternative for those patients in whom NSAIDs are contraindicated. Its use is limited by toxicity with high doses. It does not induce fluid retention and can therefore be given to patients with heart failure; it can also be given to patients receiving anticoagulants.

Chronic Gout:
For long-term control of gout in patients who have frequent attacks, the xanthine oxidase inhibitor allopurinol may be used to reduce production of uric acid. It should not be used to treat an acute attack since it may prolong it indefinitely. Treatment for chronic gout should not be started until after an acute attack has completely subsided, usually 2-3 weeks. The initiation of allopurinol treatment may precipitate an acute attack therefore colchicine or a suitable NSAID should be used as a prophylactic and continued for at least one month after the hyperuricaemia has been corrected. If an acute attack develops during treatment for chronic gout, then allopurinol should continue at the same dosage and the acute attack should be treated in its own right. Treatment for chronic gout must be continued indefinitely to prevent further attacks of gout.

Allopurinol
EDL-D16 Tertiary

AVAILABILITY
 TABLET 100 mg.

DOSE
Oral Adult- Initially 100 mg daily after food, thereafter adjust according to uric acid concentration. (Usual maintenance dose in mild conditions: 100 to 200 mg daily, in moderately severe condition: 300 mg daily given in divided doses). Child- Neoplastic conditions and enzyme disorders: 10 to 20 mg/kg daily (max. 400 mg).

INDICATION
Prophylaxis of gout; prophylaxis of hyperuricaemia associated with cancer chemotherapy.

CONTRAINDICATION
Acute gout; if an acute attack occurs while receiving allopurinol; continue prophylaxis and treat attack separately

PRECAUTION
Ensure adequate fluid intake of 2-3 litres daily; lactation (Appendix 7b); renal and hepatic impairment (Appendices 7d and 7a); withdraw treatment if rash occurs; reintroduce if rash is mild but discontinue immediately if it recurs; interactions (Appendix 6c, 6d); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Rash (see precautions above); hypersensitivity reactions occur rarely, and include fever; lymphadenopathy; arthralgia; eosinophilia; erythema multiforme (Stevens-Johnson syndrome) or toxic epidermal necrolysis; vasculitis; hepatitis; renal impairment.
Disease modifying agents used in Rheumatoid disorders (DMARDs)

The process of cartilage and bone destruction which occurs in rheumatoid arthritis may be reduced by the use of a diverse group of drugs known as DMARDs (disease-modifying antirheumatic drugs). DMARDs include antimalarials (chloroquine, hydroxychloroquine), penicillamine, sulfasalazine, immunosuppressants (azathioprine, cyclophosphamide, methotrexate) and gold compounds.

Treatment should be started early in the course of the disease, before joint damage starts. Treatment is usually initiated with a NSAID when the diagnosis is uncertain and the disease course unpredictable. However, when the diagnosis, progression and severity of rheumatic disease have been confirmed, a DMARD should be introduced. DMARDs do not produce an immediate improvement but require 4-6 months of treatment for a full response. Their longterm use is limited by toxicity and loss of efficacy. If one drug does not lead to objective benefit within 6 months, it should be discontinued and another DMARD substituted. Adverse reactions with DMARDs occur frequently and may be life threatening; careful monitoring is needed to avoid severe toxicity. Blood disorders (bone marrow suppression) can occur during treatment with many DMARDs; blood counts should be carried out before and during treatment, and patients should be advised to report without delay any unexplained symptom such as bleeding, bruising, purpura, infection, sore throat or fever.

It has been suggested that combinations of DMARDs may be more effective than single drug but increased toxicity may be a problem; whether used alone or in combination, they should be prescribed only by specialists to ensure that they are used safely and to best advantage. The antimalarial chloroquine is less effective than most other DMARDs, but as it is generally better tolerated it may be preferred in the treatment of mild rheumatoid arthritis. Chloroquine should not be used for psoriatic arthritis. Because long-term therapy can result in retinopathy ophthalmological examinations should be conducted before and during treatment. Sulfasalazine has a beneficial anti-inflammatory effect and is considered by some rheumatologists to be a first-line DMARD, but it is poorly tolerated by about 25% of patients. Adverse reactions include blood disorders (bone marrow suppression), hepatotoxicity, skin reactions and gastrointestinal disturbances.

Methotrexate, an immunosuppressant, is considered to be a first-line DMARD; at the low doses used for rheumatoid arthritis it is well tolerated but there remains the risk of blood disorders (bone marrow suppression) and of hepatic and pulmonary toxicity. Other immunosuppressant drugs, including azathioprine, are generally reserved for use in patients with severe disease who have failed to respond to other DMARDs, especially in those with extra-cellular manifestations such as vasculitis. Immunosuppressants are used in psoriatic arthritis. Adverse reactions include blood disorders, alopecia, nausea and vomiting. Penicillamine is not a first-line drug and its use is limited by a significant incidence of adverse effects including blood disorders (bone marrow suppression), proteinuria and rash. Corticosteroids are potent anti-inflammatory drugs but their place in the treatment of rheumatoid arthritis remains controversial. Their usefulness is limited by adverse effects and their use should be controlled by specialists. Corticosteroids are usually reserved for use in patients with severe disease which has failed to respond to other antirheumatic drugs, or where there are severe extra-articular effects such as vasculitis.
Corticosteroids are also used to control disease activity during initial therapy with DMARDs. Although corticosteroids are associated with bone loss, this appears to be dose-related; recent studies have suggested that a low dose of a corticosteroid started during the first two years of moderate to severe rheumatoid arthritis may reduce the rate of joint destruction. The smallest effective dose should be used, such as oral prednisolone 7.5 mg daily for 2-4 years only, and at the end of treatment the dose should be tapered off slowly to avoid possible long-term adverse effects. Relatively high doses of a corticosteroid, with cyclophosphamide, may be needed to control vasculitis.

**Hydroxy Chloroquine (as phosphate or sulphate)**

**EDL-D266 Secondary hospitals**

**AVAILABILITY**
- TABLETS 250 and 500 mg;
- INJECTION 10 and 30 ml (40 mg/ml);
- SUSPENSION 50 mg/ml.

**DOSE**
- Oral Adult: Immediately 600 mg, after 6 h 300 mg followed by 300 mg daily for 2 days. Child: 10 mg/kg body weight followed by 5 mg/kg body weight after 6 h, thereafter once a day for 2 days.
- Intramuscular injection: Adult 10 ml followed by 5 ml after 6 h. Thereafter 5 ml daily for two days. Child 5 mg/kg body weight administered every 12 h followed by oral therapy.

**INDICATION**
- Treatment of acute malaria caused by P. malariae and susceptible P. falciparum; P. vivax and P. ovale (followed by primaquine to eliminate intrahepatic forms); prophylaxis of malaria for pregnant women and nonimmune individuals at risk; rheumatic disorders.

**CONTRAINDICATION**
- Severe hematologic distress or gastrointestinal distress; eye dysfunction; liver disease.

**PRECAUTION**
- If patient continues to deteriorate after chloroquine-suspect resistance and administer quinine intravenously as emergency measure; hepatic impairment; renal impairment; pregnancy (but in malaria, benefit considered to outweigh risk; lactation; may exacerbate psoriasis; neurological disorders (avoid for prophylaxis if history of epilepsy); may aggravate myasthenia gravis; severe gastrointestinal disorders; G-6-PD deficiency; avoid concurrent therapy with hepatotoxic drugs; interactions

**ADVERSE EFFECTS**
- Headache, gastrointestinal disturbances; also convulsions; visual disturbances (retinopathy associated with long-term, high dose therapy or inappropriate self-medication); depigmentation or loss of hair; rashes; pruritus—may become intolerable; bone-marrow suppression; hypersensitivity reactions such as urticaria and angioedema; atioventricular block (may be result of inappropriate self-medication); porphyria and psoriasis in susceptible individuals.

**Methotrexate Sodium**

**EDL-D335, D685Restricted**

**AVAILABILITY**
- TABLETS 2.5, 5.0 and 7.5 mg;
- INJECTION vial/ampoule 25 mg/ml and 100 mg/ml.

**DOSE**
- Oral Severe active rheumatoid arthritis: 7.5 mg once weekly, adjusted according to response (max. weekly dose 20 mg). Intramuscular, subcutaneous or intravenous route in severe attack under expert medical supervision at a dose of 7.5 mg once weekly.
INDICATION
Rheumatoid arthritis which has failed to respond to penicillamine or chloroquine; malignant
disease.

CONTRAINDICATION
Blood disorders (bone marrow suppression); liver damage; pulmonary toxicity; gastrointestinal
disturbances-if stomatitis and diarrhoea occur; stop treatment; renal failure; skin reactions;
alopecia; osteoporosis; arthralgia; myalgia; ocular irritation; precipitation of diabetes.

PRECAUTION
Monitor throughout treatment including blood counts and hepatic and renal function tests;
renal and hepatic impairment (avoid if severe; see also Appendices 7a); reduce dose or
withdraw if acute infection develops; for woman or man; during contraception and for at least 6
months after treatment; peptic ulceration; ulcerative colitis; diarrhoea; ulcerative stomatitis;
advise patient to avoid self-medication with salicylates or other NSAIDs; warn patient with
rheumatoid arthritis to report cough or dyspnoea; interactions (Appendix 6a, 6c, 6d). Patients
should be warned to report immediately any signs or symptoms of bone marrow suppression;
for example unexplained bruising or bleeding; purpura; infection; sore throat

ADVERSE EFFECTS
Blood disorders (bone marrow suppression); liver damage; pulmonary toxicity; gastrointestinal
disturbances-if stomatitis and diarrhoea occur; stop treatment; renal failure; skin reactions;
alopecia; osteoporosis; arthralgia; myalgia; ocular irritation; precipitation of diabetes.

Sulfasalazine
EDL-D493 Secondary hospitals

AVAILABILITY
TABLETS 500 and 1000 mg Enteric coated.

DOSE
Oral Acute rheumatoid arthritis: Adult- initially 500 mg daily increase by 500 mg at interval of
one week (max. 2 to 3g in divided doses). Child- 40-50 mg/kg/day.

INDICATION
Severe rheumatoid arthritis; ulcerative colitis; Crohn’s disease.

CONTRAINDICATION
Hypersensitivity to salicylates or sulfonamides; child under 2 years; porphyria; intestinal or
urinary obstruction; severe renal impairment; G-6-PD deficiency; blood dyscracias.

PRECAUTION
Monitor during first 3 months of treatment including blood counts and hepatic and renal function
tests; lactation (Appendix 7b); history of allergy; G-6-PD deficiency; slow acetylator
status; interactions (Appendix 6c, 6d); pregnancy (Appendix 7c).
Patients should be warned to report immediately any signs or symptoms of bone marrow
suppression; for example unexplained bruising or bleeding; purpura; infection; sore throat.

ADVERSE EFFECTS
Nausea, exacerbation of colitis; diarrhoea, loss of appetite, fever; blood disorders (including
Heinz body anaemia, megaloblastic anaemia, leukopenia, neutropenia, thrombocytopenia);
hypersensitivity reactions (including rash, urticaria, Stevens-Johnson syndrome (erythema
multiforme), exfoliative dermatitis, epidermal necrolysis, pruritus, photosensitization,
anaphylaxis, serum sickness, interstitial nephritis, lupus erythematosus- like syndrome); lung
complications (including eosinophilia, fibrosing alveolitis); ocular complications (including
periorbital oedema); stomatitis, parotitis; ataxia, aseptic meningitis, vertigo, tinnitus, alopecia,
peripheral neuropathy, insomnia, depression, headache, hallucinations; kidney reactions
(including proteinuria, crystalluria, haematuria); oligospermia; rarely, acute pancreatitis,
hepatitis; urine may be coloured orange; some soft contact lenses may be stained
SECTION - 3
ANTIALLERGICS AND MEDICINES USED IN ANAPHYLAXIS

Antihistamines are used to treat drug allergies, food allergies, insect stings and some of the symptoms of anaphylaxis and angioedema. Drug treatment and other supportive care should not be delayed in critically ill patients. Specific precipitants should be sought and if identified, further exposure avoided and desensitization considered.

Drowsiness and sedation are particular disadvantages of the older antihistamines and the patient should be warned against driving or operating machinery. Other central nervous system depressants, including alcohol, barbiturates, hypnotics, opioid analgesics, anxiolytics and neuroleptics, may enhance the sedative effects of antihistamines. Since antihistamines interfere with skin tests for allergy, they should be stopped at least one week before conducting a skin test.

Allergic reactions of limited duration and with mild symptoms, such as urticaria or allergic rhinitis, usually require no treatment. If on the other hand, symptoms become persistent, antihistamines constitute the mainstay of treatment. However, oral corticosteroids may be required for a few days in an acute attack of urticaria or for severe skin reactions. Oral corticosteroids are also used to relieve severe exacerbations in chronic urticaria, but long-term use should be avoided. Corticosteroids may be used topically to reduce inflammation in allergic rhinitis but should only be used systemically for this condition when symptoms are disabling.

Allergic Emergencies
Anaphylactic shock and conditions such as angioedema are medical emergencies that can result in cardiovascular collapse and/or death. They require prompt treatment of possible laryngeal oedema, bronchospasm or hypotension. Atopic individuals are particularly susceptible. Insect stings and certain foods including eggs, fish, cow's milk protein, peanuts and nuts are a risk for sensitized persons. Therapeutic substances particularly associated with anaphylaxis include blood products, vaccines, hyposensitizing (allergen) preparations, antibiotics (especially penicillins), iron injections, heparin and neuromuscular blocking drugs. Acetylsalicylic acid and other nonsteroidal anti-inflammatory drugs (NSAIDs) may cause bronchoconstriction in leukotriene sensitive patients. In the case of drug allergy, anaphylaxis is more likely to occur after parenteral administration. Resuscitation facilities should always be available while injecting a drug associated with risk of anaphylactic reactions.

First-line treatment of a severe allergic reaction includes administering epinephrine, keeping the airway open (with assisted respiration if necessary) and restoring blood pressure (laying the patient flat, raising the feet). Epinephrine should immediately be given by intramuscular injection to produce vasoconstriction and bronchodilation and injection should be repeated if necessary at 5-min intervals until blood pressure, pulse and respiratory function have stabilized. If there is cardiovascular shock with inadequate circulation, epinephrine must be given cautiously by slow intravenous injection of a dilute solution. Oxygen administration is also of primary importance. An antihistamine such as chlorpheniramine is a useful adjunctive treatment given after epinephrine injection and continued for 24 to 48 h to reduce the severity and duration of symptoms and to prevent relapse. An intravenous corticosteroid such as
hydrocortisone has an onset of action that is delayed by several hours but should be given to help prevent later deterioration in severely affected patients. Further treatment of anaphylaxis may include intravenous fluids, an intravenous vasopressor such as dopamine, intravenous aminophylline or injected or nebulized bronchodilator, such as salbutamol.

**Chlorpheniramine (hydrogen maleate)**

**EDL-D115, 116, 117 PHC**

**AVAILABILITY**

TABLETS 2, 4 and 6 mg; INJECTIONS 10 mg/10 ml, CAPSULE 8 mg; SYRUP 10 mg/50 ml, 100 mg/100 ml.

**DOSE**

Oral Adult- Allergic reactions: 4 mg every 4 to 6 h (max. 24 mg daily). Child- 1 to 2 years: 1 mg twice daily. 2 to 5 years: 1 mg every 4 to 6 h (max. 12 mg daily). 6 to 12 years: 2 mg every 4 to 6 h (max. 12 mg daily) Intramuscular or intravenous injection Adult- Allergic reactions: 10 to 20 mg, repeated if required (max. 40 mg in 24 h). Subcutaneous injection Child- Allergic reactions: 87.5 μg/kg, repeated if necessary up to 4 times daily. Intravenous injection (over 1 min). Adult- Anaphylaxis (adjunct): 10 to 20 mg. Child- Anaphylaxis (adjunct)- under 1 year: 250 μg/kg. 1 to 5 years: 2.5 to 5 mg. 6 to 12 years: 5 to 10 mg.

**INDICATION**

Symptomatic relief of allergy, allergic rhinitis (hay fever); conjunctivitis; urticaria; insect stings and pruritus of allergic origin; adjunct in the emergency treatment of anaphylactic shock and severe angioedema.

**CONTRAINDICATION**

Prostatic enlargement, urinary retention; ileus or pyloroduodenal obstruction; asthma; child under 1 year; hypersensitivity, narrow angle glaucoma, pregnancy lactation

**PRECAUTION**

Performing works requiring utmost alertness such as vehicle driving, operating machines etc within 24 h of taking the drug should be avoided. Lactation (Appendix 7b); renal and hepatic impairment (Appendix 7a); epilepsy; interactions (Appendix 6a); atropic gastritis, elderly.

**ADVERSE EFFECTS**

Drowsiness (rarely, paradoxical stimulation with high doses, or in children or elderly), hypotension, headache, palpitations, psychomotor impairment, urinary retention, dry mouth, blurred vision, gastrointestinal disturbances; liver dysfunction; blood disorders; also rash and photosensitivity reactions, hypersensitivity reactions (including bronchospasm, angioedema, anaphylaxis); sweating and tremor, injections may be irritant; flatulence, diarrhoea.

**Dexamethasone**

**EDL-D157 Universal**

**INDICATIONS**

Adjunct in the emergency treatment of anaphylaxis; short-term suppression of inflammation in allergic disorders; adrenocortical insufficiency, ocular inflammation, autoimmune disorders, rheumatic disorder, cerebral oedema, unresponsive shock, bacterial meningitis along with antibiotics.

**AVAILABILITY**

TABLETS 0.5 mg; INJECTION 2 ml vial (4 mg/ ml); CREAM 5 and 15 g (0.1% w/w).
**DOSE**

**Oral**
- Adult: 0.5 to 10 mg daily in divided doses, repeat if necessary.
- Child: 0.02 to 0.3 mg/kg in three or four divided doses daily.

**Intravenous injection**
- 4 to 10 mg every 6 h.

**CONTRAINDICATIONS**
- Untreated systemic infection (unless condition life-threatening); administration of live virus vaccines; renal failure, diabetes mellitus, psychosis, osteoporosis, pregnancy (Appendix 7c), CHF, tuberculosis, fungal infections of the eye.

**PRECAUTIONS**
- Increased susceptibility to and severity of infection; activation or exacerbation of tuberculosis, amoebiasis, strongyloidiasis; risk of severe chickenpox in non-immune patient (varicella-zoster immunoglobulin required if exposed to chickenpox); avoid exposure to measles (normal immunoglobulin possibly required if exposed); diabetes mellitus; peptic ulcer; hypertension; precautions relating to long-term use of corticosteroids; glaucoma, epilepsy; drug should not be abruptly withdrawn; interactions (Appendix 6c), lactation (Appendix 7b).

**ADVERSE EFFECTS**
- Nausea, dyspepsia, malaise, hiccups; hypersensitivity reactions including anaphylaxis; perineal irritation after intravenous administration; adverse effects associated with long-term corticosteroid treatment; hyperglycaemia, abdominal distension, angioedema, bradycardia, acne, erythema, Cushing’s syndrome, oropharyngeal candidiasis, hypothalamic pituitary adrenal axis suppression.

**STORAGE**
- Store protected from light at a temperature not exceeding 30°C.

**Epinephrine Hydrochloride (Adrenaline)**

**EDL-D199 PHC**

**AVAILABILITY**
- INJECTION 1 ml ampoule (1 mg/ml).

**DOSE**
- Intramuscular injection Anaphylaxis: preferable site is the midpoint in anterior thigh [1:1000 solution]. This route should be used by specialists only with extreme care. Slow intravenous injection When there is doubt regarding adequacy of circulation and absorption from the intramuscular site; slow intravenous injection of 1:10000 (10 mg/ml) solution be injected in severely ill patients only.

**INDICATION**
- Severe anaphylactic reaction; severe angioedema; cardiac arrest; hemostatic agent.

**CONTRAINICATION**
- Narrow angle glaucoma, organic brain damage, cardiac dilation, coronary insufficiency.

**PRECAUTION**
- Hyperthyroidism, hypertension, diabetes mellitus, heart disease, arrhythmias, cerebrovascular disease; second stage of labour; elderly; interactions (Appendix 6c); pregnancy (Appendix 7c); lactation (Appendix 7b);

**ADVERSE EFFECTS**
- “Epinephrine fastness”, tachycardia and arrhythmias, hypertension, tremor, anxiety, sweating, nausea, vomiting, weakness, hyperglycaemia, dizziness, pulmonary oedema have all been reported; headache common.
**Prednisolone**  
EDL-D427, 428,429 PHC

**INDICATIONS**  
Short-term as well as long term suppression of inflammation in allergic disorders; malignant disease; Autoimmune disease, bronchial asthma.

**AVAILABILITY**  
TABLETS 5, 10, 20, 30 and 40 mg; SYRUP 1 mg/ml and 3 mg/ml; EYE DROPS 1% w/v; INJECTION 2 ml vial (40 mg/ml).

**DOSE**  
**Oral**  
Adult and Child- Initially up to 10 to 20 mg daily in divided doses (severe diseases up to 60 mg), preferably after breakfast.  
**Intramuscular injection**  
Adult and Child- 25 mg to 100 mg once or twice weekly.

**CONTRAINDICATIONS**  
Untreated systemic infection; administration of live virus vaccines; hypersensitivity.

**PRECAUTIONS**  
Increased susceptibility to and severity of infection; activation or exacerbation of tuberculosis, amoebiasis, strongyloidiasis; risk of severe chickenpox in non-immune patient (varicella-zoster immunoglobulin required if exposed to chickenpox); avoid exposure to measles (normal immunoglobulin possibly required if exposed); diabetes mellitus; peptic ulcer; hypertension;  
Further precautions relating to long-term use of corticosteroids myasthenia gravis, congestive heart failure, renal insufficiency, pregnancy (Appendix 7c), osteoporosis, glaucoma, psychological disorders, diverticulitis, interactions (Appendix 6c, 6d), lactation (Appendix 7b), hepatic impairment (Appendix 7a).

**ADVERSE EFFECTS**  
Nausea, dyspepsia, malaise, hiccups; hypersensitivity reactions including anaphylaxis; supraclavicular lump, fragile skin.

**STORAGE**  
Store protected from light and moisture.

**Hydrocortisone**  
EDL-D261 PHC

**INDICATIONS**  
Adjunct in the emergency treatment of anaphylaxis; inflammatory skin conditions; inflammatory bowel disease; adrenocortical insufficiency; As acetate: rheumatology, neurology, episcleritis, sinusitis; Addison’s disease, Simmond’s disease, tuberculous meningitis; perineal trauma, joint inflammation, subaortic dermatitis.

**AVAILABILITY**  
TABLETS 5, 10 and 20 mg, CREAM 10g (1% w/w), OINTMENT 1%, 2.5% w/w INJECTION 100, 200 and 400 mg/vial, (25 mg/5 ml).

**DOSE**  
**Intramuscular injection or slow intravenous injection or intravenous infusion**  
Adult-100 mg to 500 mg, 3 to 4 times in 24 h or as required.  
**Slow intravenous injection**  
Child- Up to 1 year: 25 mg. 1 to 5 years: 50 mg.

**CONTRAINDICATIONS**  
Not relevant to emergency use but for contra-indications relating to long-term use; ulcers.
PRECAUTIONS
Not relevant to emergency use but for precautions relating to long-term use, interactions (Appendix 6b) lactation (Appendix 7b), pregnancy (Appendix 7c).

ADVERSE EFFECTS
Adverse effects associated with long-term corticosteroid treatment; opportunistic infections.

Methyl Prednisolone
EDL-D336 Secondary hospitals

AVAILABILITY
TABLETS 4, 8, 16 and 24 mg; INJECTION vials 40, 125, 500 and 1000 mg, 2 ml ampoule (80 mg/2 ml).

DOSE
Oral Adult- Asthma, allergies and dermatological conditions: 40 and 120 mg. Dose should be regulated in accordance with severity of condition; large joints- 20 to 80 mg; medium joints- 10 to 40 mg; small joints- 4 to 10 mg directly in bursae.

INDICATION
Corticosteroid responsive conditions such as severe allergic rhinitis, asthma, rheumatoid arthritis, osteoarthritis, collagen disease, dermatoses.

CONTRAINDICATION
Systemic fungal infection (unless specific antimicrobial therapy given); avoid live virus vaccines in those receiving immunosuppressive doses (serum antibody response diminished); hypersensitivity.

PRECAUTION
Refer notes above; interactions (Appendix 6c,6d); pregnancy (Appendix 7c)

ADVERSE EFFECTS
Besides the usual steroid side effects, acute hyperglycemia, hypokalemia, infections, and convulsions are more frequently encountered. Bolus injections may produce sudden cardiac death.

Cetirizine
EDL-D 583 PHC

INDICATIONS
symptomatic relief of allergy such as hay fever, chronic idiopathic urticaria

AVAILABILITY
Tablets 10 mg, 5mg, syrup 5 mg/5 mL

PRECAUTIONS
Caution may be required in epilepsy. Children and the elderly are more susceptible to sideeffects also renal impairment.

CONTRAINDICATIONS
Hypersensitivity, end-stage renal disease with creatinine clearance < 10 ml/min. Not recommended for lactating mothers or children below 6 months also pregnancy

DOSE
ADULT and CHILD over 6 years, 10 mg once daily or 5 mg twice daily; CHILD 1–2 years Child 1–2 years 250 micrograms/kg twice daily; for Children, 2–6 years, hay fever, 5 mg or 1 teaspoon syrup once daily or 2.5 mg (½ teaspoon) syrup twice daily.

ADVERSE EFFECTS
Somnolence, fatigue, dry mouth, nasopharyngitis have been reported in adults. Fever, cough, epistaxis and diarrhoea may occur in children <12 years.

Storage
Store protected from heat, light and moisture at a temperature not exceeding 30°C.
**Levocetirizine**  
**EDL-D683 PHC**

**INDICATIONS**  
Allergic rhinitis, chronic urticaria.

**AVAILABILITY**  
TABLETS 5 mg; SYRUP 2.5 mg/5 ml.

**Dose**  
Oral  
Rhinitis, chronic urticaria: Adult & children (>12 years) - 5 mg once daily in the evening.  
Children (6-12 yrs) - 2.5 mg once daily.  
Children (6 months - 5 yrs) – 1.25 mg once daily.

**CONTRAINDICATIONS**  
Hypersensitivity, end-stage renal disease with creatinine clearance < 10 ml/min. Not recommended for lactating mothers or children below 6 months. Precautions May impair the ability to drive or operate machinery, concurrent use of alcohol or CNS depressant drugs should be avoided, pregnancy, elderly, interactions.

**ADVERSE EFFECTS**  
Somnolence, fatigue, dry mouth, nasopharyngitis have been reported in adults. Fever, cough, epistaxis and diarrhoea may occur in children <12 years.

**Storage**  
Store protected from heat, light and moisture at a temperature not exceeding 30°C.

**Fluticasone**  
**EDL-D642 PHC**

**AVAILABILITY**  
Inhalation Aerosol- Formoterol + Fluticasone Propionate 6 μg + 125 μg 6 μg + 250 μg

**DOSE**  
Inhalation Asthma: Adults- 1-2 inhalations twice daily. Child- 1 rotacap twice daily. (Rotacaps to be used with a Rotahaler device only. Do not swallow the capsules). COPD: Adults- 2 inhalations twice daily. Not recommended for children below 4 years of age.

**INDICATION**  
Asthma, severe chronic obstructive pulmonary disease (COPD).

**CONTRAINICATION**  
Hypersensitivity, acute asthma symptoms.

**PRECAUTION**  
Severe cardiovascular disorders, cardiac rhythm abnormalities, seizure disorder, diabetes, thyrotoxicosis, hypokalemia, pulmonary tuberculosis, pregnancy (Appendix 7c), lactation, interactions

**ADVERSE EFFECTS**  
Headache, pharyngitis, throat irritation, upper respiratory tract infections, pneumonia, bronchitis, oral candidiasis, nausea, vomiting, diarrhea, chest pain, musculoskeletal pain, back pain, allergic reactions, wheezing, cough, skin rash, tremors, paradoxical bronchospasm, insomnia, adrenal suppression.

**Cinnarizine**  
**EDL-D121 Tertiary**

**INDICATION**  
Motion sickness, nausea, vomiting, vertigo and tinnitus associated with Meniere disease and other middle ear disorders, as a nootropic drug, adjunct therapy for symptoms of peripheral arterial disease.

**AVAILABILITY**  
TABLETS 25 & 75 mg Plain and 75 mg SR.

**DOSE**  
Oral: Motion sickness
Adult: 30 mg 2 hr before travel and 15 mg every 8 hr during travel if needed. **Vertigo**
Adult: 30 mg thrice daily.
Child: 5-12 year: half of adult dose.
**Peripheral circulatory disorders**
Adult: 75 mg tablets three times daily.

**CONTRAINDICATIONS**
Hypersensitivity, Parkinson's disease, children below 5 years.

**PRECAUTIONS**
Hypotension, patients should not drive or operate machinery, pregnancy (Appendix 7c), lactation, elderly, children and neonates, interactions (Appendix 6c).

**ADVERSE EFFECTS**
Drowsiness, rarely skin and hypersensitivity reactions, dry mouth, extrapyramidal symptoms sometimes associated with severe depression, muscular weakness, headache, euphoria, GI upsets, blurred vision, urinary difficulty or retention, constipation and increased gastric reflux, fatigue, hypolipidaemic effect.

**Fexofenadine**
Non- **EDL Tertiary**

**INDICATIONS**
Allergic rhinitis, urticaria.

**AVAILABILITY**
TABLETS 30, 60, 120 and 180 mg; Syrup 30 mg/5 ml.

**DOSE**
Allergic rhinitis: Adult- 120 mg once daily.
Child (6-11year)- 30 mg twice daily.
Urticaria and skin allergy: Adult-180 mg once daily.
Child- (6 month to 2 years): 15 mg twice daily, more than 2 years: 30 mg twice daily.

**CONTRAINDICATIONS**
Hypersensitivity.

**PRECAUTIONS**
Bradycardia, hypokalemia, preexisting long QT interval, renal impairment, pregnancy (Appendix 7c), lactation, interactions.

**ADVERSE EFFECTS**
Dizziness, stomach discomfort, pain in extremity, back pain, vomiting, diarrhoea, upper respiratory tract infection, headache, dysmenorrhoea.

**Hydroxyzine**
**EDL-D662 Secondary hospitals**
It has both antianxiety and antihistaminic activity.

**INDICATION**
Pruritus, acute and chronic urticaria and dermatosis, anxiety.

**CONTRAINDICATION**
Pregnancy, neonates, urinary and GI obstruction.

**PRECAUTION**
Renal impairment, lactation, peptic ulcer, BPH. (Benign Prostatic hypertrophy)

**ADVERSE EFFECT**
Tachycardia, arrhythmias, headache, blood dyscrasias, tinnitus.

**AVAILABILITY**
Tablet 10 mg, 25 mg Injection 25 mg/mL
These notes are only guidelines and it is strongly recommended that poisons information centres (Appendix 5) be consulted in cases where there is doubt about the degree of risk or about appropriate management.

**Non specific**

**General Care and Non-Specific Treatment:**

All patients who show features of poisoning should generally be admitted to hospital. Patients who have taken poisons with delayed actions should also be admitted, even if they appear well; delayed-action poisons include acetylsalicylic acid, iron, lithium, paracetamol, paraquat, tricyclic antidepressants and warfarin. The effects of modified-release or prolonged-release preparations are also delayed. However, it is often impossible to establish with certainty the identity of the poison and the size of the dose but information on the type and timing of poisoning may be useful for symptomatic management. Few patients require active removal of the poison. Most patients must be treated symptomatically and monitored. Particular care must be given to maintenance of respiration and blood pressure. Assisted ventilation may be required. Cardiac conduction defects and arrhythmias often respond to correction of underlying hypoxia, acidosis, or other biochemical abnormalities. Hypothermia which may develop in patients who have been unconscious for some hour is best treated by wrapping the patient in blankets to conserve body heat. Convulsions which are prolonged or recurrent may be controlled by intravenous diazepam. In some situations removal of the poison from the stomach by gastric lavage may be appropriate (see below). Activated charcoal can bind many poisons in the stomach and therefore prevent absorption. Active elimination techniques such as repeated administration of activated charcoal can enhance the elimination of some drugs after they have been absorbed (see below). Other techniques to enhance elimination of poisons after their absorption are only practical in hospital and are only suitable for a small number of patients and only to a limited number of poisons. Methods include haemodialysis and haemoperfusion. Alkalinization of urine can be used to increase the elimination of salicylates. Forced alkaline diuresis is no longer recommended.

**Gastric Lavage:**

The dangers of attempting to empty the stomach have to be balanced against the toxicity of the ingested poison, as assessed by the quantity ingested, the inherent toxicity of the poison and the time since ingestion. Gastric emptying is clearly unnecessary if the risk of toxicity is small or if the patient presents too late. Emptying the stomach may be of value if undertaken within 1-2 h after ingestion. The main risk is with inhalation of stomach contents and gastric lavage should not be undertaken in drowsy or comatose patients without assistance of an anaesthetist so that the airway can be protected by a cuffed endotracheal tube. Gastric lavage must not be attempted after corrosive poisoning or for hydrocarbon products which could be dangerous if aspirated.
**Emesis:**

Induction of emesis for the treatment of poisoning is not recommended. There is no evidence that it prevents absorption of the poison and it may increase the likelihood of aspiration. Furthermore, the effects of the emetic substance may complicate diagnosis.

**Prevention of Absorption:**

Given by mouth activated charcoal can bind many poisons in the gastrointestinal system, thereby reducing their absorption.

The sooner it is given, the more effective it is, but it may be effective for up to 1 hour after ingestion of the poison. It may be effective several hour after poisoning with modified-release preparations or drugs with anticholinergic (antimuscarinic) properties. It is relatively safe and particularly useful for prevention of absorption of poisons which are toxic in small amounts, for example, antidepressants. Furthermore, repeated doses of activated charcoal enhance the faecal elimination of some drugs (that undergo enterohepatic or enteroenteric recycling) several hours after ingestion and after they have been absorbed, for example phenobarbital, theophylline.

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**Active Charcoal**

**EDL-DS PHC**

**AVAILABILITY**

POWDER (for oral suspension), TABLETS 500mg.

**DOSE**

Oral Adult and child over 12years- 50g, 0.5g/kg may be repeated every 4-6 h for upto 12-24 h.

Child- Below 12years; 1g/kg (max 50g). May be repeated every 4 h.

**INDICATION**

Treatment of acute poisoning.

**CONTRAINDICATION**

Poisoning by hydrocarbons with high potential for harm if aspirated; poisoning by corrosive substances-may prevent visualization of lesions caused by poison.

**PRECAUTION**

Drowsy or unconscious patients-risk of aspiration (intubate before administration via nasogastric or gastric tube); not effective for poisoning with alcohols, clofenotane (dicophane, DDT), cyanides, malathion and metal salts including iron and lithium.

**ADVERSE EFFECTS**

Black stools; vomiting, constipation or diarrhoea; pneumonitis-due to aspiration

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**Calcium Disodium Edetate**

**Non- EDL Tertiary**

**INDICATIONS**

Lead poisoning (acute and chronic) and lead encephalopathy.

**AVAILABILITY**

AMPOULE 5 ml (200 mg/ml).

**DOSE**

Intravenous injection

Lead poisoning without encephalopathy: 1000 mg/m2/day as continous infusion for 5 days.
Lead encephalopathy: 1500 mg/m²/day by continuous intravenous infusion in 5% dextrose or 0.9% NaCl (Final Concentration of edentate < 500 mg/100 ml), starting 4 h after first dose of BAL and after an adequate urine flow is established. Infusion is continued for 5 days.
Intramuscular injection to be used if fluid overload is a concern. 1000 mg/m²/day divided into equal doses spaced 8 to 12 h apart.
Lignocaine or procaine should be added to the injection to minimize pain at the injection site.

CONTRAINDICATIONS
- Anuria; patients with active renal disease or hepatitis; pregnancy

PRECAUTIONS
- Ensure adequate urine output, pre-existing mild renal disease; patients with lead encephalopathy and cerebral edema may experience a lethal increase in intracranial pressure following intravenous infusion, the intramuscular route is preferred for these patients.

ADVERSE EFFECTS
- Renal tubular toxicity which may lead to acute renal failure, fever, chills, lacrimation, increased prothrombin time, pain at intramuscular injection site; hypotension; cardiac rhythm irregularities; thirst; headache; fatigue; malaise; urinary frequency; glycosuria; proteinuria; microscopic hematuria; histamine-like reactions.

Specific
**Paracetamol Overdosage:**
Paracetamol in a dose of 10-15g or 150 mg/kg of paracetamol taken within 24 h may cause severe hepatocellular necrosis and less frequently renal tubular necrosis. The only early features of poisoning, nausea and vomiting, usually settle within 24 h. Persistence beyond this time, often with the onset of right subcostal pain and tenderness, usually indicates the development of liver damage which is maximal 3-4 days after ingestion. In spite of a lack of significant early symptoms, patients who have taken an overdose of paracetamol should be transferred to hospital urgently.
Administration of activated charcoal should be considered if paracetamol in excess of 150 mg/kg or 12g, whichever is smaller, is thought to have been ingested within the previous hour. N-Acetylcysteine or N-methionine protect the liver if given within 10-12 h of ingesting paracetamol. Acetylcysteine, given intravenously is most effective within 8 h of overdosage, but is effective for up to and possibly beyond 24 h. Alternatively, methionine may be given by mouth provided the overdose was ingested within 10-12 h and the patient is not vomiting. However, acetylcysteine is the preferred treatment. Concurrent use of activated charcoal and specific oral antidotes should be avoided.
In remote areas methionine should be given, since administration of acetylcysteine outside hospital is not generally practicable. Once the patient is in hospital the need to continue antidote treatment can be assessed from plasma-paracetamol concentrations.

**Opioid Analgesic Overdosage:**
Opioids cause varying degrees of coma, respiratory depression and pinpoint pupils. Naloxone is a specific antidote indicated if there is coma or bradypnoea. Naloxone has a shorter duration of action than many opioids so close monitoring and repeated injections are required depending on respiratory rate and depth of coma; naloxone may alternatively be given by intravenous infusion. The effects of some opioids such as buprenorphine are only partially reversed by naloxone. Acute withdrawal syndromes may be precipitated by the use of naloxone in patients.
with a physical dependence on opioids or in overdosage with large doses; a withdrawal syndrome may occur in neonates of opioid-dependent mothers.

**Organophosphate and Carbamate Poisoning:**
Organophosphates are absorbed through the bronchi and intact skin as well as from the gastrointestinal tract. Initial treatment of organophosphate or carbamate poisoning includes prevention of further absorption by emptying the stomach by gastric lavage, moving patient to fresh air supply, removing contaminated clothing and washing contaminated skin. A clear airway must be maintained. Organophosphates inhibit cholinesterases and thus prolong the effects of acetylcholine. Toxicity depends on the particular compound involved and onset after ingestion, skin exposure may be delayed. Atropine will reverse the muscarinic effects of acetylcholine and is used (in conjunction with oximes such as pralidoxime) with additional symptomatic treatment. Additional treatment for carbamate poisoning is generally symptomatic and supportive. Atropine may be given but may not be required because of the rapidly reversible type of cholinesterase inhibition produced (oximes should not be given).

**Iron Poisoning and Iron and Aluminium Overload:**
Mortality from iron poisoning is reduced by specific therapy with desferrioxamine which chelates iron. Before administration of desferrioxamine the stomach should be emptied by gastric lavage (with a wide-bore tube) within 1 h of ingesting a significant quantity of iron or if radiography reveals tablets in the stomach. Desferrioxamine is also used to diagnose and treat chronic iron overload. It is used in the diagnosis of aluminium overload and to treat aluminium overload in patients with endstage renal failure undergoing maintenance haemodialysis.

**Heavy Metal Poisoning:**
Heavy metal poisoning may be treated with a range of antidotes including dimercaprol, penicillamine, potassium ferric hexacyanoferrate and Sodium calcium edetate. Penicillamine is also used to promote excretion of copper in Wilson’s disease.

**Methaemoglobinemia:**
Methylthioninium chloride can lower the levels of methaemoglobin in red blood cells and is used in the treatment of methaemoglobinemia. In large doses, it may cause methaemoglobinemia and therefore methaemoglobin levels should be monitored during treatment.

**Cyanide Poisoning:**
Cyanide poisoning may be treated with Sodium nitrite followed by Sodium thiosulphate.

**Pralidoxime**

**EDL-D426 PHC**

**AVAILABILITY**
Injection i.v infusion 500 mg/20 ml, 1g/20 ml (as chloride and iodide salt).

**DOSE**
For Chloride salt, 30 mg/kg i.v. over 15-20 minutes followed by infusion at 8-10 mg/kg/h. To be continued 12-24 hours after atropine is no longer required. For iodide salt, dose is about 30% higher than chloride salt. Child- 25 to 50 mg/kg, diluted to 5% concentration in NS and infused over 5-30 minutes. May be repeated after one h, then every 6 to 12 h. Severe poisoning: Adult- 500 mg/h via continuous infusion. max.- 12g/24 h. Child- 9 to 19 mg/kg/h. For anticholinesterase overdose in MG: Adult- 1-2g i.v. initially, then 250 mg every 5 minutes. Child (0-18 years)- 15-25 mg/kg by slow i.v (up to 1 g). Maintainance dose- (< 12 years) 15-50 mg/kg i.v every 5 minutes (up to 250 mg).
INDICATION
Adjunct to atropine in the treatment of organophosphate poisoning and anticholinesterase overdosage used in the treatment of myasthenia gravis (mg), respiratory depression or severe muscle weakness due to carbamate poisoning.

CONTRAINDICATION
Carbamate poisoning and organophosphates without anticholinesterase activity; hypersensitivity to the drug.

PRECAUTION
Impaired renal function; large doses can cause neuromuscular blockade, myasthenia gravis; atropinization occur faster on concurrent use with atropine; paediatrics; allergies; pregnancy (Appendix 7c)

ADVERSE EFFECTS
Headache, nausea; blurred vision, drowsiness, dizziness, impaired accommodation, tachycardia, hyperventilation, muscular weakness; transient elevation in SGOT and/ or SGPT levels; laryngospasm and rigidity.

Neostigmine Metilsulfate
EDL-D364 Secondary hospitals

AVAILABILITY
TABLET 15 mg; Injection 0.5 mg/ml

DOSE
Oral Adult- 15 mg every 3 to 4 hrs. Total daily dose 75 to 300 mg in divided doses. Child- 2 mg/kg daily in divided doses every 3 to 4 hrs. Total daily dose 15 to 90 mg. Neonate- 1 to 5 mg every 4 hour. Intramuscular Adult- 0.02 mg/kg as a single dose. Child- 0.04 mg/kg as a single dose. Intravenous Adult- 0.5 to 2.5 mg to a total daily dose of 5-20 mg. Child- 200 to 500 μg as single daily dose. Neonate- 50 to 250 μg every 4 hour.

INDICATION
Neostigmine is an anticholinesterase, which is particularly effective in postsynaptic neurotoxins such as those of cobra and is not useful against presynaptic neurotoxin i.e. common Krait and the Russell’s viper.15 Neostigmine test should be performed by administering 0.5–2 mg IV and if neurological improvement occurs, it should be continued 1/2 hourly over next 8 hours.

Treatment of Myasthenia gravis.

CONTRAINDICATION
Mechanical gastrointestinal or urinary tract obstruction; peritonitis.

PRECAUTION
Renal impairment; peptic ulcer; lactation ; heart blockage, slow heartbeat; bradycardia, hypotension; urinary tract infection; epilepsy; asthma; interactions ; pregnancy (Appendix 7c)

ADVERSE EFFECTS
Abdominal cramps, diarrhoea; pupil dilatation; excess saliva; headache; joint pain; severe allergic reactions; fainting; interrupted breathing; irregular heart beat; seizures; vision changes; anxiety.

Atropine
EDL-D56 PHC

INDICATIONS
Organophosphate and carbamate poisoning; premedication; antispasmodic; as mydriatic; cycloplegic refraction procedures.

AVAILABILITY
INJECTION 1 ml ampoules and 50 ml vial (0.6mg/ml).
DOSE

**Intramuscular and intravenous injection**

Adult: 1.8 - 3.0 mg intravenous bolus followed by doubling dose every 3 to 5 minutes depending upon response. End-point for atropinization include clear chest with no wheeze, systolic BP >80mm Hg, pulse >80 beats/min., pupils no longer pinpoint and dry axillae. Following that infusion of atropine at 10-20 % of total initial dose required/hour; may require boluses during infusion.

Child: 20-30 μg/kg initially with same schedule as above.

**CONTRAINDICATIONS**

In myasthenia gravis (but may be used to decrease muscarinic side-effects of anticholinesterases), paralytic ileus, pyloric stenosis and prostatic enlargement; reflux oesophagitis; unstable cardiac rhythm.

**PRECAUTIONS**

Elderly, Down syndrome; angle-closure glaucoma; myasthenia gravis; prostatic enlargement; pyrexia; lactation (Appendix 7b); interactions (Appendix 6a); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Constipation, transient bradycardia (followed by tachycardia, palpitation and arrhythmias), reduced bronchial secretions, urinary urgency and retention, dilatation of the pupils with loss of accommodation, photophobia, dry mouth, flushing and dryness of the skin. Occasionally, confusion (particularly in the elderly), nausea, vomiting and giddiness; very rarely, angle-closure glaucoma may occur.

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**Physostigmine**

**Non-EDL**

**INDICATION**

o.5-2 mg i.v. repeated as specific antidote for belladonna poisoning. It penetrates BBB and antagonizes both central and peripheral actions.

**PRECAUTIONS**

However it often induces hypotension and arrhythmias, it should be employed only as a last resort. Needs ECG availability. Care to be taken when administered into eye, pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Twitching lids, myopia, ocular and periocular pain, ciliary and conjunctival congestion.

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**Desferrioxamine Mesylate**

**Non- EDL Tertiary**

**INDICATIONS**

Acute iron poisoning; chronic iron overload; aluminium overload; primary hemochromatosis.

**AVAILABILITY**

INJECTION 5 ml and 10 ml vial (500 mg/vial).

**DOSE**

Continuous intravenous infusion

Adult and Child: Begin with 5 mg/kg/h, increasing over 15 minutes if tolerated to 15 mg/kg/h, to minimize the risk of hypotension. After 1 to 2 h reduce to 3-4 mg/kg/h for the next 22-23 hrs (max dose is 100 mg/kg over 24 hrs).

Patients with cardiovascular collapse: 5 mg/ kg/h (up to max. of 80 mg/kg in 24 h.) Chronic iron overload: Intramuscular 500 to 1000 mg daily, in addition 2g by intravenous infusion with each unit of blood transfused.

**CONTRAINDICATIONS**

Severe renal disease; pregnancy (Appendix 7C).
PRECAUTIONS
Renal impairment; eye and ear examinations before and at 3-month intervals during treatment; aluminium encephalopathy (may exacerbate neurological dysfunction); children under 3 years (may retard growth); lactation; interactions (Appendix 6C)

ADVERSE EFFECTS
Anaphylaxis; flushing, urticaria, hypotension, shock (especially if given by too rapid intravenous infusion); gastrointestinal disturbances; fever, headache, arthralgia, myalgia; arrhythmias; renal impairment; blood disorders; neurological disturbances including neuropathy, paraesthesia and dizziness; convulsions; Yersinia and mucormycosis infections; visual disturbances (including lens opacity and retinopathy) and hearing loss; rash; rarely, growth retardation (in young children); rarely, acute respiratory distress syndrome; pain on intramuscular or subcutaneous injection; local irritation on prolonged subcutaneous infusion; reddishbrown discolouration of urine.

Storage
Store protected from light in refrigerator (2-8°C). Do not freeze.

Dimercaprol (BAL)
Non-EDL Tertiary

INDICATIONS
Acute poisoning by antimony, arsenic, bismuth, copper gold, mercury and possibly thallium; adjunct (with sodium calcium edetate) in lead poisoning.

AVAILABILITY
OILY INJECTION 2 ml ampoule (50 mg/ml).

DOSE
Intramuscular injection
To be administered by deep intramuscular injection only
Lead poisoning: Adults-4 mg/kg every 4 h for 5 days. Child- 75 mg/m² every 5 h for 5 days.
Arsenic poisoning: 3 mg/kg every 4 h for 48 h and then twice a day for 7-10 days.
Mercury poisoning: 5 mg/kg followed by 2.5 mg/kg every 12-24 h for upto 10 days

CONTRAINDICATIONS
Not indicated for iron, selenium or cadmium poisoning; severe hepatic impairment (unless due to arsenic poisoning); hypertension; tellurium poisoning, peanut allergy, G-6-PD deficiency.

Precautions
Hypertension; renal impairment (discontinue or use with extreme caution if renal failure occurs during treatment); any abnormal reaction such as hyperpyrexia should be assessed; elderly; pregnancy (Appendix 6C); lactation, alkalinize urine to pH of 7.5-8.0 using sodium bicarbonate.

ADVERSE EFFECTS
Hypertension, tachycardia; malaise, nausea, vomiting, abdominal pain, salivation,lacrimation, sweating, burning sensation in the mouth, throat and eyes; feeling of constriction in throat and chest; headache, muscle spasms, tingling of the extremities; fever in children; local pain and abscess at injection site, iron toxicity potentiation.

STORAGE
Store protected from light.

D-Penicillamine
Non- EDL Tertiary

INDICATIONS
Poisoning by heavy metals, particularly lead and copper; Wilson’s disease; severe rheumatoid arthritis.

AVAILABILITY
CAPSULE/tablet 250 mg.
DOSE
Oral (given before food)
Adult- 1 to 2g daily in three divided doses starting with 250 mg OD and gradually increasing to full dose over 2-3 weeks.
Child- 20 mg/kg/day administered in 3-4 divided doses, initiating treatment at 25% of this dose and gradually increasing to full dose over 2-3 weeks to minimize adverse reactions.
Continue till blood lead levels <45 μg/dl.

CONTRAINDICATIONS
Hypersensitivity; lupus erythematosus; gold or antimalarial drug; penicillamine-induced agranulocytosis; aplastic anaemia; thrombocytopenia, pregnancy, lactation (for rheumatoid arthritis).

PRECAUTIONS
Monitor throughout treatment including blood counts and urine tests; renal impairment; immunosuppressive treatment; avoid oral iron within 2 h of a dose; hepatic impairment; pregnancy (Appendix 7c)
In Wilson’s disease, consider withdrawal if platelet count falls below 120 000/mm3 or white blood cells below 2500/mm3 or if 3 successive falls within reference range (can restart at reduced dose when counts return to reference range but permanent withdrawal necessary if neutropenia or thrombocytopenia recur).
In Wilson’s disease warn patient to tell doctor immediately if sore throat, fever, infection, non-specific illness, unexplained bleeding and bruising, purpura, mouth ulcers or rashes develop.

ADVERSE EFFECTS
Initially nausea (less of a problem if taken with food and on retiring), anorexia, fever; taste loss (mineral supplements not recommended); blood disorders including thrombocytopenia, neutropenia, agranulocytosis and aplastic anaemia; proteinuria, rarely, haematuria (withdraw immediately); haemolytic anaemia nephrotic syndrome, lupus erythematosus-like syndrome, myasthenia gravis-like syndrome, polymyositis (rarely, with cardiac involvement), dermatomyositis, mouth ulcers, stomatitis, alopecia, bronchiolitis and pneumonitis, pemphigus, Goodpasture syndrome and Stevens-Johnson syndrome also reported; male and female breast enlargement reported; rash early in treatment (usually allergic-may need temporary withdrawal), late rashes (reduce dose or withdraw treatment).

Flumazenil
Non- EDL Tertiary

INDICATIONS
Antidote for benzodiazepine overdose, reversal of sedative effects produced by benzodiazepenes administered during general anaesthesia or diagnostic or therapeutic procedures.

AVAILABILITY
Injection 0.1 mg/ml.
Dose Adult- 0.2 mg (2 ml) administered over 30 seconds, i.v, repeat 0.3 mg and 0.5 mg at 1-2 minute intervals. Not more than 3 mg over one hour.
Child- 10 μg/kg, i.v, for 2 doses.

CONTRAINDICATIONS
Epilepsy, neuromuscular blockade, hypersensitivity to benzodiazepines, patients of suspected tricyclic antidepressant overdose, raised intracranial pressure.

PRECAUTIONS
History of seizures, panic attack, alcohol drug dependence, bleeding disorder, liver disease, head injury, respiratory depression, pregnancy (Appendix 7c)
ADVERSE EFFECTS
Convulsions, fatigue, injection site pains, increased sweating, facial erythema, raised intracranial pressure, agitation, dizziness, abnormal vision, may cause complete heart block, flushing, transient increase in blood pressure and heart-rate.

Methylene Blue
Non-EDL Tertiary

INDICATIONS
Acute methaemoglobinaemia.

AVAILABILITY
INJECTION 10 mg/ml.

DOSE
Intravenous injection
Methaemoglobinaemia caused by high dosage of prilocaine infusion: 1-2 mg/kg intravenously over 5 minutes, followed immediately by a fluid flush of 15-30 ml to minimize local pain. May be repeated in 30-60 minutes. Maximum dose: 7 mg/kg.

CONTRAINDICATIONS
Severe renal impairment; methaemoglobinaemia due to chlorate or induced by sodium nitrite in treatment of cyanide poisoning; affects ability to drive machinery.

PRECAUTIONS
G-6-PD deficiency-may cause haemolytic anaemia; monitor blood methaemoglobin throughout treatment; pregnancy (Appendix 7c); lactation.

ADVERSE EFFECTS
Nausea, vomiting, abdominal pain, chest pain, headache, dizziness, confusion, profuse sweating; hypertension or hypotension reported; haemolytic anaemia in G-6-PD deficiency; methaemoglobinaemia with high dosage; bluish skin discolouration; blue saliva, urine and faeces.

STORAGE
Store protected from light in an airtight container.

Naloxone
Non-EDL Tertiary

INDICATIONS
Opioid overdosage; postoperative respiratory depression.

AVAILABILITY
INJECTION 0.4 mg/ml.

DOSE
Intravenous injection
Subcutaneous or intramuscular route (if i.v. route is not feasible but the dose is same, can be given oral as well).
Adult- Opioid poisoning: Start with 0.4 to 2 mg (at all ages) as intravenous bolus, repeat every 2 minutes if no response to a total of 10 mg. Once response occurs start infusion of naloxone at 2/3rd the total loading dose given every hour with continuous monitoring for recurrence of respiratory depression. May require additional bolus during infusion.
Child- Opioid poisoning: 10 μg/kg, followed by 100 μg/kg if there is no response.

CONTRAINDICATIONS
Hypersensitivity.

PRECAUTIONS
Physical dependence on opioids or other situations where acute withdrawal syndrome may be precipitated (see above); lactation; cardiovascular disease; pregnancy (Appendix 7c)

ADVERSE EFFECTS
Nausea, vomiting, sweating-may also be due to opioid withdrawal.
Sodium Nitrite
Non- EDL Tertiary

INDICATIONS
Cyanide poisoning (together with Sodium thiosulphate).

AVAILABILITY
Injection 30 mg/ml (10 ml).

DOSE
Intravenous injection (over 5 to 20 min)
Adult- 300 mg at 2.5-5.0 mg/minute.
Child- 4 to 10 mg/kg (max 300 mg) at 5 mg/ minute.
Note: Prepare as 3% solution of Sodium nitrite in Water for Injections (30 mg/ml) at the time of administration.

CONTRAINDICATIONS
Methaemoglobinaemia; hemolytic anaemia; G-6-PD deficiency.

PRECAUTIONS
Monitor plasma methaemoglobin levels; severe cardiovascular or cerebrovascular disease; hypotension; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea, vomiting and abdominal pain, vasodilatation resulting in syncope, hypotension, tachycardia, flushing, headache; methaemoglobinaemia; cyanosis, dyspnoea, tachypnoea.

Sodium Thiosulphate
Non- EDL Tertiary

INDICATIONS
Prophylactically with prolonged use of nitro prusside to prevent cyanide toxicity, cyanide poisoning (together with Sodium nitrite); pityriasis versicolor; skin disease.

AVAILABILITY
Injection 250 mg/ml; 500 mg/ml (50 ml).

DOSE
Intravenous injection (over 10 min). Adult- 12.5g intravenously over 10-30 minutes may be repeated at half the initial dose at 1-2 hours.
Child- 500 mg/kg intravenously over 10-30 minutes may be repeated at half the initial dose at 1-2 hours (12.5g maximum)

CONTRAINDICATIONS
Hypersensitivity; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Irritation; urticaria; hypotension; burning; stinging on application

Disulfiram
EDL-D609, 610 Secondary hospitals

INDICATION
Adjunct in the treatment of chronic alcohol dependence.

CONTRAINDICATION
Cardiac failure, coronary artery disease and history of cerebrovascular accident, hypertension, psychoses, pregnancy and breast-feeding.

PRECAUTION
Ensure that alcohol is not consumed for at least 24 hours before initiating treatment, hepatic and renal impairment, respiratory disease, diabetes mellitus, epilepsy.

ADVERSE EFFECT
Drowsiness and fatigue; nausea and vomiting, reduced libido, rarely psychotic reactions.
AVAILABILITY
Tablet 250 mg.

DOSE
1 g as a single dose on first day, reduced over 4 days to 0.75g to 0.25g od; should not be continued for longer than 6 months without review.

DRUG INTERACTION
Psychotic reaction with metronidazole, inhibition of metabolism of tricyclic antidepressants. Inhibition of metabolism of phenytoin. Inhibition of metabolism of benzodiazepines, leading to enhanced sedative effect.

**Calcium Gluconate**

**EDL-D84 Secondary hospitals**

INDICATIONS
Hypocalcaemic tetany; cardiopulmonary bypass.

AVAILABILITY
Tablets 250 and 500 mg; Injection 10 ml (1g/10 ml).

DOSE
Slow intravenous injection and continuous intravenous infusion
Adult- Hypocalcaemic tetany: 1g (2.2 mmol) by slow intravenous injection, followed by continuous intravenous infusion of about 4g (8.8 mmol) daily.

CONTRAINDICATIONS
Conditions associated with hypercalcaemia and hypercalciuria (for example some forms of malignant disease).

PRECAUTIONS
Monitor plasma calcium concentration; renal impairment; interactions (Appendix 6c); diarrhoea, parathyroid disease; stomach trouble.

ADVERSE EFFECTS
Mild gastrointestinal disturbances bradycardia, arrhythmias, hypotension; irritation at injection site; soft tissue calcification; nephrocalcinosis, renal calculi.

**Prazosin**

**EDL-D717, 718 Secondary hospitals**

INDICATION
It is used as Scorpion string.

CONTRAINDICATION
Heart failure due to mechanical obstruction like aortic stenosis.

PRECAUTIONS
May produce first dose hypotension and collapse. Withdraw diuretics if patient is already on diuretics. Reduce dose in renal impairment. Use with caution in pregnancy. The drug is preferably given at bed time.

ADVERSE EFFECT
Urinary frequency, incontinence, dizziness, headache, lack of energy, nausea, postural hypotension. It may cause increase in renin levels.

AVAILABILITY
Tablets 1 mg, 2 mg and 5 mg (sustained release)

DOSE
Start with 0.5 mg h.s. If no syncope or giddiness in the morning, gradually increase dose to 1 mg bd In the extended release form of prazosin containing 5 mg the first dose effect is not common.
DRUG INTERACTION
ACE inhibitors, alcohol, antidepressants, antipsychotics, anxiolytics, diuretics, beta-blockers and calcium channel blockers all potentiate the hypotensive action. Corticosteroids decrease the effect.

Deferiprone
EDL-D603 Tertiary hospitals

INDICATION
Iron chelation, it is an oral iron chelating drug

AVAILABILITY
Tablets 250 mg, 500 mg

DOSE
0.5 - 3 g daily (100 mg/kg bw) to be given 1 h before food, in three divided doses.

PRECAUTION
Pregnancy and lactation.

ADVERSE EFFECT
Agranulocytosis, arthralgias, arthritis, drug-induced lupus erythematosus, toxic overload of iron in the liver.
Control of Epilepsy:
Treatment of seizures should always be started with a single antiepileptic drug (AED), and the choice of an anticonvulsant should be made on an individual basis. The drug of choice will depend on the primary diagnosis, seizure type, efficacy of the drug and the patient’s tolerance of treatment. If a drug fails to control the seizures after it has been used in full therapeutic dosage for an adequate period, or if it is not tolerated, it should be gradually substituted with another drug, with the first drug being withdrawn only when the new regimen is established. If monotherapy is ineffective, next alternative drug should be started, and try to withdraw first drug if there was no response for that drug or continue with that if there was partial response for initial drug.

Initial dose of the drug of choice should be determined on the basis of the degree of urgency, the size and age of the patient. It should be increased gradually until an effective response is obtained. All antiepileptics commonly produce neurological adverse effects at higher dose ranges and patients should be monitored closely for adverse effects to help in accurate dose titration. Except for phenytoin, it is rarely useful to measure plasma-drug concentrations as an aid to dose adjustment. Non-compliance, inappropriate dosing and overdosing is a major impediment to effective antiepileptic treatment. Patients should ideally remain under supervision throughout treatment period.

Withdrawal:
Treatment is normally continued for a minimum of two years of seizure free period. In certain circumstances like in juvenile myoclonic epilepsy, antiepileptic drugs may need to be continued throughout life, because of the high relapse rate of seizure after AED withdrawal. Withdrawal should be extended over a period of several months because abrupt withdrawal can lead to recurrence of seizure and or/status epileptics. A general rule for duration of tapering is how many years patient had taken that particular drug, over a period of so many months it should be tapered. In patients receiving several antiepileptic drugs, only one drug should be withdrawn at a time. Many adult patients relapse once treatment is withdrawn and it may be justified to continue treatment indefinitely, particularly when the patient’s livelihood or lifestyle can be endangered by recurrence of a seizure.

Pregnancy and Lactation:
Untreated epilepsy during pregnancy may cause harm to the fetus; there is therefore no justification for abrupt withdrawal of treatment although withdrawal of therapy may be an option if the patient has been seizure-free for at least 2 years; resumption of treatment may be considered after the first trimester. If antiepileptics are continued in pregnancy, monotherapy with the lowest effective dose is preferred, with adjustment made to take account of changes in plasma levels associated with pregnancy. There is an increased risk of birth defects with the use of anticonvulsants, particularly carbamazepine, valproate and phenytoin. However, if there is good seizure control, there is probably no advantage in changing pregnant patients’ antiepileptic drugs. In view of the risks of neural tube and other defects, patients who may become pregnant should be informed of the risks and referred for advice and pregnant patients...
should be offered counseling and antenatal screening. To counteract the risk of neural tube defects, adequate folate supplements are advised for women before and during pregnancy. In view of the risk of neonatal bleeding associated with carbamazepine, Phenobarbital and phenytoin, prophylactic phytomenadione (vitamin K1) is recommended for the neonate and the mother before delivery. Antiepileptic drugs can be continued during lactation (see also Appendix 7b).

**Driving:**

Regulations are in place in many countries which may, for example, restrict driving by patients with epilepsy to those whose seizures are controlled. Further, antiepileptic drugs may cause CNS depression, particularly in the early stages of treatment and patients affected by adverse effects such as drowsiness or dizziness should not operate machinery or drive.

**Choice of Antiepileptic in Management of Convulsive Disorders**

**Generalized Tonic-Clonic Seizures:**

Phenobarbital, phenytoin and valproate are widely used in the treatment of these conditions. However, each of these drugs is associated with dose-related and idiosyncratic adverse effects and monitoring of haematological and hepatic function is routinely not advised.

**Simple Partial and Complex Partial Seizures:**

Carbamazepine, oxcarbamazepine, clobazam, lamotrigine and zonisamide are effective in partial epilepsy.

**Absence Seizures:**

Both ethosuximide and valproate are recommended in the treatment of absence seizures (petit mal) and are usually well tolerated. However, ethosuximide can, rarely, cause lupus erythematosus and psychoses which call for immediate, but cautious, discontinuation. Absence seizures are commonly associated with tonic-clonic seizures and valproate is preferred since it has a broad spectrum of activity.

**Tonic Seizures, Atonic Seizures and Atypical Absence Seizures:**

Phenobarbital or phenytoin is widely used for tonic seizures, valproate or clonazepam for atonic seizures and clonazepam for atypical absence seizures. However, tonic seizures most of the times are associated with multiple seizures types like Lennox-Gastaut syndrome (LGS), where phenytoin and phenobarbitone should be avoided as they can precipitate other type of seizures.

**Myoclonic Seizures:**

Valproate is widely used and most effective for juvenile myoclonic seizures. As juvenile myoclonic epilepsy is associated with a high relapse rate, it is often necessary to continue therapy indefinitely. Other myoclonic seizures are often resistant to treatment and some do not have an epileptic basis. Valproate or clonazepam can be of value in this case and other antiepileptic drugs may be useful in intractable cases. Both drugs are generally well accepted, although tolerance to clonazepam has been reported.

**Infantile Spasm (Infantile Myoclonic Epilepsy):**

Infantile spasms, which are often associated with severe brain damage, can be resistant to antiepileptic drugs. Drugs effective in this type of seizures are benzodiazepines (nitrazepam, clonazepam, clobazam), valproic acid, ACTH, vigabatrin, levetiracetam, topiramate, zonisamide, lamotrigine, and ketogenic diet.

**Febrile Convulsions:**

Sponging with tepid water and antipyretic such as paracetamol is effective in controlling the temperature. Recurrent febrile convulsions or prolonged convulsions (those lasting >5 min) are
treated with diazepam, either rectally in solution or by intravenous injection, or intranasal or buccal midazolam, to prevent possible brain damage. Intermittent prophylaxis, with diazepam (or clobazam) administered at the onset of fever, may prevent recurrence of febrile convulsions. Use of antiepileptics for continuous prophylaxis is controversial; it is probably indicated in only a small proportion of children including those who already have evident neurological abnormalities, or who have had previous prolonged or focal convulsions. Phenobarbital may be used for this purpose but careful clinical monitoring and dosage adjustment are necessary in order to minimize the risk of adverse effects. Valproate can also be used.

**Status Epilepticus:**

Status epilepticus is a medical emergency which carries a high mortality rate. Initial management includes positioning the patient to avoid injury, supporting respiration including provision of oxygen, maintaining blood pressure and the correction of any hypoglycaemia; hypocalcemia or any other electrolyte disturbance; maintenance of the airway and assisted ventilation are crucial even when the seizures are controlled, because the drugs used in its management may cause respiratory depression.

Intravenous lorazepam, midazolam are often effective in status epilepticus. Lorazepam, which acts rapidly, should be administered first and should be followed immediately by a loading dose of phenytoin which has a longer-acting effect. When cannulation is difficult or impossible, diazepam may be administered rectally as a solution (absorption from suppositories is too slow for treatment of status epilepticus). Intravenous phenobarbital is also effective but is more likely to cause respiratory depression; it is used in refractory cases but should be avoided in patients who have recently received oral phenobarbital. Rectal paraldehyde may also be used; it causes little respiratory depression and is therefore useful where facilities for resuscitation are poor. If seizures continue despite treatment, intravenous valproate, levetiracetam, midazolam infusion, propofol infusion, barbiturate coma and general anaesthesia may be required. The underlying cause must be identified and remedied in all cases.

**Carbamazepine**

EDL-D86,87,88 Secondary hospitals

**AVAILABILITY**

TABLETS 100, 200 and 400 mg Plain; 100 mg DT; 200, 300 and 400 mg CR and SR; SYRUP 100 ml (100 mg/5 ml).

**DOSE**

Oral Adult- Initially 100 and 200 mg 1 to 2 times daily increased slowly to usual dose of 400 mg to 1.2g daily in divided doses. In some cases 1.6 to 2g may be needed. Administer lower initial dose to elderly.Child- Start with 5 - 10 mg/kg/day in two to three divided doses then gradually increase at weekly intervals to a max. dose of 30-35 mg/kg/day.

**INDICATION**

Partial seizures with or without secondary generalisation; trigeminal neuralgia; bipolar disorder.

**CONTRAINDICATION**

Atrioventricular conduction abnormalities; history of bone-marrow depression; porphyria

**PRECAUTION**

Hepatic impairment (Appendix 7a), renal impairment; cardiac disease (see also Contraindications); skin reactions (see Adverse effects); history of blood disorders (blood counts before and during treatment); glaucoma; lactation(Appendix 7b), avoid sudden withdrawal; interactions (Appendix 6b, 6c, 6d), pregnancy (Appendix 7c), Patients or their caretakers should be told how to recognize signs of blood, liver or skin disorders and advised to seek immediate...
medical attention if symptoms such as fever, sore throat, rash, mouth ulcers, bruising or bleeding develop. Leukopenia which is severe, progressive and associated with clinical symptoms requires withdrawal (if necessary under cover of suitable alternative). May impair ability to perform skilled tasks, for example operating machinery, driving; see also notes above.

ADVERSE EFFECTS
Dizziness, drowsiness, headache, ataxia, blurred vision, diplopia (may be associated with high plasma levels); gastrointestinal intolerance including nausea and vomiting, anorexia, abdominal pain, dry mouth, diarrhoea or constipation; commonly, mild transient generalized erythematous rash (withdraw if worsens or is accompanied by other symptoms); leukopenia and other blood disorders (including thrombocytopenia, agranulocytosis and aplastic anaemia); cholestatic jaundice, hepatitis, acute renal failure, Stevens-Johnson syndrome (erythema multiforme), toxic epidermal necrolysis, alopecia, thromboembolism, arthralgia, fever, proteinuria, lymph node enlargement, arrhythmias, heart block and heart failure, dyskinesia, paraesthesia, depression, impotence, male infertility, gynaecomastia, galactorrhoea, aggression, activation of psychosis, photosensitivity, pulmonary hypersensitivity, hyponatraemia, oedema, disturbances of bone metabolism with osteomalacia also reported; confusion and agitation in elderly.

Magnesium Sulphate
EDL-D321 Universal

AVAILABILITY
INJECTION 500 mg/ml.

DOSE
Intravenous injection (concentration of magnesium sulphate should not exceed 20%) Prevention of seizure occurrence in eclampsia: initially 4g over 5 to 15 min, followed by infusion 1g/hr for at least 24 h after last seizure. If seizures recur, additional dose of 2g (or 4g if body weight is over 70 kg).

INDICATION
Prevention of recurrent seizures in eclampsia; prevention of seizures in pre-eclampsia; acute nephritis in children.

CONTRAINDICATION
Severe hepatic impairment; respiratory depression; acute narrow angle glaucoma; pregnancy, lactation.

PRECAUTION
Hepatic impairment(Appendix 7a); pregnancy(Appendix 7c); renal impairment; in severe hypomagnesaemia administer initially via controlled infusion device (preferably syringe pump); monitor blood pressure, respiratory rate, urinary output and for signs of overdosage (loss of patellar reflexes, weakness, nausea, sensation of warmth, flushing, drowsiness, double vision and slurred speech).

ADVERSE EFFECTS
Nausea and vomiting, dizziness; weakness; blurred vision; vertigo

Phenytoin Sodium
EDL-D408,409 PHC

AVAILABILITY
TABLETS 100, 150 and 200 mg Plain; 300 mg SR; CAPSULES 25 mg, 100 mg; INJECTION 2 ml ampoule (50 mg/ml); SUSPENSION 25 mg/ml.

DOSE
Oral or slow intravenous injection or infusion Adult- Status epilepticus: (with regular BP and ECG monitoring) 18 mg/kg at rate not exceeding 50 mg/min as loading dose, maintenance dose of about 100 mg should be given thereafter at an interval of 6 to 8 h (dose can be reduced
according to weight). Child - Status epilepticus: 20 mg/kg at a rate not exceeding 1 mg/kg/min, maintenance dose 4-7 mg/kg/day in 2 divided doses, max dose 300 mg/day.

INDICATION
Generalized tonic-clonic seizures; partial seizures; status epilepticus.

CONTRAINDICATION
Porphyria; avoid parenteral use in sinus bradycardia, sino-atrial block, second- and third-degree heart block, Stokes-Adams syndrome; pregnancy

PRECAUTION
Hepatic impairment (reduce dose (Appendix 7a); lactation (Appendix 7b); diabetes mellitus; monitor blood counts; hypotension and heart failure (caution with parenteral use); intravenous administration-resuscitation facilities must be available; injection solution alkaline (irritant to tissues); interactions (Appendix 6a,6b,6c); hypersensitivity; osteomalacia, it worsens myoclonus and absence seizures. Patients or their caretakers should be told how to recognize signs of blood or skin disorders and advised to seek immediate medical attention if symptoms such as sore throat, rash, mouth ulcers, bruising or bleeding develop. Leukopenia which is severe, progressive or associated with clinical symptoms requires withdrawal (if necessary under cover of suitable alternative). May impair ability to perform skilled tasks, for example operating machinery, driving; see notes above.

ADVERSE EFFECTS
Gastric intolerance, headache, sleeplessness, agitation (during initial phase); sedation, hallucinations, confusion; blurred vision, ataxia, nystagmus, diplopia; slurred speech, cerebellar-vestibular symptoms, behavioural disorders, hyperglycaemia (may be signs of overdosage); gingival hyperplasia, acne, coarse facies, hirsutism, fever; neurological changes (peripheral neuropathy, choreiform movements, impaired cognition, increased seizure frequency); osteomalacia, rickets (associated with reduced plasma calcium levels); lymph-node enlargement; rashes (discontinue; if mild re-introduce cautiously, but discontinue if recurrence); very rarely, Stevens-Johnson syndrome (erythema multiforme), systemic lupus erythematosus, toxic epidermal necrolysis; rarely, blood disorders including megaloblastic anaemia (may be treated with folic acid), leukopenia, thrombocytopenia, agranulocytosis with or without bone marrow depression; intravenous administration-cardiovascular and CNS depression (particularly if administered too rapidly) with arrhythmias, hypotension and cardiovascular collapse, alterations in respiratory function (including respiratory collapse); dyskinesia; hepatitis, hepatic failure.

Lorazepam
EDL-D313,319 Tertiary

AVAILABILITY
TABLETS 0.5, 1, 2, 2.5 and 3 mg INJECTIONS 2 ml ampoule (2 mg/ml).

DOSE
2 to 6 mg/day given in divided doses, initial dose of 2 to 3 mg/day given twice or thrice a day. Elderly or debilitated patients: Initial dosage of 1 to 2 mg/day in divided doses.

INDICATION
Anxiety disorders.

CONTRAINDICATION
Severe hepatic impairment; respiratory depression; acute narrow angle glaucoma; pregnancy, lactation.

PRECAUTION
Hepatic dysfunction; impaired ability to drive or operate machinery; interactions

ADVERSE EFFECTS
Nausea and vomiting, dizziness; weakness; blurred vision; vertigo.
Diazepam

EDL- D164 Secondary hospitals D165,166,167 PHC

INDICATIONS
Status epilepticus; emergency management of recurrent seizures; febrile convulsions; seizures associated with poisoning and medicine withdrawal; adjunct in acute alcohol withdrawal; premedication; anxiety disorders; psychosomatic behaviour disorder; spasticity.

AVAILABILITY
TABLETS 2, 5 and 10 mg; CAPSULE 10 mg; SUSPENSION 2 mg/ml; INJECTION 2 ml ampoule (5 mg/ml).

DOSE
Intravenous injection
Adult-Treatment of status epilepticus and convulsions due to poisoning: 10 mg at the rate of 1 ml/min (5 mg) repeated if necessary after 10 min.
Child-Under 12 years: 300 to 400 μg/kg, repeated after 10 min if necessary.

CONTRAINDICATIONS
Respiratory depression; acute pulmonary insufficiency; sleep apnoea; severe hepatic impairment; myasthenia gravis; avoid injections containing benzyl alcohol in neonates, narrow angle glaucoma; hypersensitivity to benzodiazepine.

PRECAUTIONS
Respiratory disease, muscle weakness, history of alcohol or drug abuse, marked personality disorder; pregnancy (Appendix 7c); lactation (Appendix 7b); reduce dose in elderly or debilitated patients and in hepatic impairment (avoid if severe, Appendix 7a), renal impairment; avoid prolonged use and abrupt withdrawal; when given intravenously, facilities for reversing respiratory depression with mechanical ventilation must be at hand (see below); porphyria; interactions (Appendix 6a, 6c); blood count test on prolonged treatment. Intravenous infusion of diazepam is potentially hazardous (especially if prolonged) calling for close and constant observation and best carried out in a speciality centre with intensive care facilities. Prolonged intravenous infusion may lead to accumulation and delay recovery. May impair ability to perform skilled tasks, for example operating machinery, driving; see also notes above.

ADVERSE EFFECTS
Drowsiness and lightheadedness the next day; confusion and ataxia (especially in the elderly); amnesia; dependence; paradoxical increase in aggression; muscle weakness; occasionally headache, vertigo, salivation changes, gastrointestinal disturbances, skin reactions, visual disturbances, dysarthria, tremors, incontinence, urinary retention; blood disorders and jaundice; hypotension and apnoea, pain and thrombophlebitis (with injection); increased appetite; weight gain.

STORAGE
Store protected from light.

Phenobarbitone

EDL-D402,403,404 PHC D 405 Tertiary

INDICATIONS
Generalized tonic-clonic seizures; partial seizures; neonatal seizures; febrile convulsions; status epilepticus; sedative, hypnotic, preanaesthetic.

AVAILABILITY
TABLETS 30 and 60 mg; INJECTION 1 ml ampoule (200 mg/ml); SYRUP 20 mg/ml.

DOSE
Slow intravenous injection
Status epileptics: (dilute injection 1 in 10 with water for injections), Adult- 10 mg/kg at a rate of not more than 100 mg/min (up to max. total dose of 1 g); Child- 10-20 mg/kg at a rate of not more than 30 mg/min.
Oral
Adult- 60-180 mg daily at night.
Child- 1 month-12 years: 1-1.5 mg/kg twice daily, maintenance dose 2.5-4 mg/kg once/twice daily. 12-18 years: Initially 60-180 mg twice daily, maintenance dose 60-180 mg once daily.

CONTRAINDICATIONS
Absence seizures; latent porphyria.

PRECAUTIONS
Elderly, debilitated, children (may cause behavioural changes); impaired renal function or hepatic function (Appendix 7a), respiratory depression (avoid if severe); pregnancy (see notes above; Appendix 7c); lactation (Appendix 7b); avoid sudden withdrawal; interactions (Appendix 6a, 6b, 6c); habit forming.

ADVERSE EFFECTS
Sedation, mental depression, agitation, hallucination, syncope; ataxia, nystagmus; allergic skin reactions including rarely, exfoliative dermatitis, toxic epidermal necrolysis, Steven’s-Johnson syndrome (erythema multiforme); paradoxical excitement, restlessness and confusion in the elderly; irritability and hyperactivity in children; megaloblastic anaemia (may be treated with folic acid); osteomalacia; status epilepticus (on treatment withdrawal); hypotension, bradycardia, shock; laryngospasm and apnoea (with intravenous injection); cognitive impairment; aplastic anaemia; hepatic failure; connective tissue disorder; hyperkinesias.

STORAGE
Store protected from moisture.

Sodium Valproate
EDL-D514, 515, 516 Secondary hospitals Tertiary

INDICATIONS
Generalized tonic-clonic seizures; partial seizures; atonic seizures; absence seizures; myoclonic seizures; acute mania; migraine.

AVAILABILITY
Tablets 125, 200, 250, 300 and 500 mg Plain; 200, 300 and 500 mg CR; Syrup 200 mg/5 ml; INJECTION 100 mg/vial, 5 ml ampoule (100 mg/5 ml) CR.

DOSE
Oral
Adult- 600 mg daily in two divided doses (preferably after food) thereafter increase by 200 mg at 3 days interval clinical response till desired.
Child- Initial dose 20 mg/kg/day, max. dose 60 mg/kg/day.

CONTRAINDICATIONS
Active liver disease, family history of severe hepatic dysfunction; pancreatitis; porphyria; hypersensitivity.

PRECAUTIONS
6 months of therapy (Appendix 7a), especially in patients at most risk (children under 3 years of age, those with metabolic disorders, degenerative disorders, organic brain disease or severe seizure disorders associated with mental retardation, or multiple antiepileptic therapy); ensure no undue potential for bleeding before starting and before major surgery or anticoagulant therapy; renal impairment; pregnancy (important see notes above, (neural tube screening)) (Appendix 7c); lactation (see notes above; Appendix 7b); systemic lupus erythematosus; false-positive urine tests for ketones; avoid sudden withdrawal; interactions (Appendix 6a, 6c, 6d); hyperammonemia.

ADVERSE EFFECTS
Gastrointestinal irritation, nausea, increased appetite and weight gain, hyperammonaemia; ataxia, tremor; transient hair loss (regrowth may be curly); oedema, thrombocytopenia, inhibition of platelet aggregation; impaired hepatic function and rarely, fatal hepatic failure (see
Precautions—withdraw treatment immediately if malaise, weakness, lethargy, oedema, abdominal pain, vomiting, anorexia, jaundice, drowsiness or loss of seizure control; sedation reported and also increased alertness; behavioural disturbances; rarely, pancreatitis (measure plasma amylase if acute abdominal pain), extrapyramidal symptoms, leukopenia, pancytopenia, red cell hypoplasia, fibrinogen reduction; irregular periods, amenorrhoea, gynaecomastia, hearing loss, Fanconi syndrome, dementia, toxic epidermal necrolysis, Steven’s–Johnson syndrome (erythema multiforme), vasculitis, hirsutism and acne reported; hallucinations; abnormal gut; pneumonia; headache; taste perversion; polycystic ovary.

STORAGE
Store protected from light.

Clobazam
Non-EDL Tertiary

INDICATIONS
Add-on for refractory partial, complex and generalized seizures, add-on in West syndrome, LGS, myoclonic epilepsy, absence seizures, to cover short period of increased seizure susceptibility addition of new AED examinations overnight travel catamnial epilepsy, intermittent prophylaxis in febrile seizures.

AVAILABILITY
TABLETS 5, 10 and 20 mg.

DOSE
Oral
0.3–2.9 mg/kg/day, (average 1 mg/kg/day)single at bed time or twice daily dose.

PRECAUTIONS
Pregnancy (Appendix 7c), interactions (Appendix 6c).

ADVERSE EFFECTS
Sedation, dizziness, hyperactivity, behavioural problem, irritability, drooling, weight gain, sleep disturbance, blurring, diplopia.

Clonazepam
Non-EDL Tertiary

INDICATIONS
Absence seizures, myoclonic seizures, akinetic seizures, panic disorder, subcortical myoclonus, adjuvant treatment of refractory epilepsy.

AVAILABILITY
TABLETS 0.25, 0.5, 1 and 2 mg.

DOSE
Adult- 0.5 - 5 mg thrice daily, initial dose should not exceed 1.5 mg/day, slow titration is recommended Maintenance dose 4-8 mg daily, Maximum dose 20 mg daily.
Infants and child: Initial dose 0.01-0.03 mg/ kg/day (not to exceed 0.05 mg/kg/day) given in 2-3 divided doses. Maintenance dose 0.1-0.2 mg/kg/day in 3 divided doses.
Panic disorder: Adult- Initial dose 0.25 mg twice daily, usual maintenance dose 1 mg/day, maximum dose 4 mg/day.

CONTRAINDICATIONS
Hypersensitivity to benzodiazepines, acute pulmonary insufficiency, acute narrow angle glaucoma.

PRECAUTIONS
Neonates, chronic pulmonary insufficiency, hepatic and renal dysfunction, porphyria, elderly, pregnancy (Appendix 7c), lactation (Appendix 7b), interactions (Appendix 6a, 6c); avoid sudden withdrawal.
ADVERSE EFFECTS
Sedation, dullness, CNS depression, ataxia, bronchial hypersecretion, abnormal eye movement, blood dyscrasias.

**Fosphenytoin**

**Non-EDL Tertiary**

**INDICATIONS**
Generalized tonic-clonic status epilepticus.

**AVAILABILITY**
INJECTION 2 ml vial (75 mg/ml).

**DOSE**
Adult- 15 mg/kg i.v. infusion at the rate of 100-150 mg/min.

**CONTRAINDICATIONS**
Porphyria.

**PRECAUTIONS**
Uremia, hypoalbuminemia, interactions (Appendix 6b, 6c); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
Cardiovascular collapse and/or central nervous system depression, nystagmus, dizziness, pruritus, paresthesia, headache, somnolence, ataxia, hypotension.

**Gabapentin**

**Non-EDL Tertiary**

**INDICATIONS**
Add-on drug in resistant partial seizures with or without secondary generalization, rolandic epilepsy- preferred for safety reason, first line in epilepsy patients with hepatic disease.

**AVAILABILITY**
TABLETS/CAPSULES 100 and 300 mg.

**DOSE**
Oral
Initially 10 mg/kg/day, increase 10 mg/kg/day to maintenance dose 30-100 mg/kg/day, in three divided doses.

**CONTRAINDICATIONS**
Pregnancy (Appendix 7c)

**ADVERSE EFFECTS**
Somnolence, dizziness, fatigue, nystagmus, behavioral changes (<10%)-aggression, hyperexcitability, tantrum, euphoria, weight gain.

**Lamotrigine**

**Non-EDL Tertiary**

**INDICATIONS**
Partial seizures and secondary generalized tonic-clonic seizures.

**AVAILABILITY**
TABLETS 25, 50 and 100 mg Plain; 5, 150 and 200 mg DT.

**DOSE**
Oral
Adult and Child over 12 years- 25 mg once daily for 2 weeks followed by 50 mg once daily for 2 weeks, increase by 50 to 100 mg every 1 to 2 weeks to maintenance dose of 100 to 200 mg daily.

Child- Monotherapy- Initial dose 2 mg/kg/day for 2 weeks then 5 mg/kg/day for 2 weeks. max. dose 5 - 15 mg/kg/day once or twice daily.

With valproic acid- Initial dose - 0.5 mg/kg/ day to max. dose of 1 - 5 mg/kg/day in single dose.

With enzyme inducer- 2 mg/kg/day for 2 weeks than 5 mg/kg/day for 2 weeks. Max. 5 - 15 mg/
kg/day once or twice daily, when valproic acid added to already regimen with lamotrigine, reduce dose of lamotrigine by 25 - 50%.

CONTRAINDICATIONS
Child less than 12 years; hypersensitivity; severe hepatic and renal impairment.

PRECAUTIONS
Monitoring of liver and renal function; abrupt withdrawal to be avoided; pregnancy (Appendix 7c) and lactation; avoid in patients who need to undertake task requiring mental alertness; patients taking sodium valproate.

ADVERSE EFFECTS
Skin eruptions; nausea; vomiting; headache; toxic epidermal necrosis; hepatotoxicity; leucopenia; thrombocytopenia; confusion; hallucination.

Levetiracetam
Non-EDL Tertiary

INDICATIONS
Good effect difficult-to-treat idiopathic focal epilepsies of childhood, including variations such as continuous spike and wave during sleep or Landau-Kleffner syndrome (LKS), photosensitivity and myoclonus- Generalised epilepsy with photosensitivity, idiopathic epilepsy– control of GTCS and Myoclonic, treatment of postanoxic and post-encephalitic myoclonic epilepsy, epileptic encephalopathies- LGS, West Syndrome, severe myoclonic epilepsy, absence seizure, rolandic epilepsy.

AVAILABILITY
TABLETS 250, 500 and 750 mg, SYRUP 100 mg/ml, INJECTION 5 ml ampoule (100 mg/ml).

DOSE
Oral
Initial dose- 10-20 mg/kg/day, increase by 10 mg/kg/day every 1-2 week upto 40-60 mg/kg/day in two divided doses.
Intravenous injection
20-30 mg/kg at the rate of 5 mg/kg/min.

CONTRAINDICATIONS
Hypersensitivity.

PRECAUTIONS
Renal disease; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Most frequent somnolence, asthenia (dose dependent); headache, hair loss, vertigo, nausea, infection; behavioral changes such as hostility aggression, apathy, anxiety, depression, psychosis.

Oxcarbamazepine
Non-EDL Tertiary

INDICATIONS
Monotherapy or adjunctive therapy in the treatment of partial seizures, secondary generalized seizure, substitution for carbamazepine can be made abruptly with an oxcarbamazepine-to-carbamazepine ratio of 300:200.

AVAILABILITY
TABLETS 150, 300, 450, 500 and 600 mg; SUSPENSIONS 300 mg/5 ml.

DOSE
Initial dose: 8-10 mg/kg/day, increasing by 8-10 mg/kg/day as tolerated at 3-7 day interval. Given in two divided doses. Maximum- 30 mg/kg.
PRECAUTIONS
Pregnancy (Appendix 7c) ; interactions (Appendix 6c).

ADVERSE EFFECTS
Less frequently than they do with carbamazepine (2.8% vs 6.5%), hyponatremia is more common but not clinically significant, rash, weight gain, alopecia, nausea, headache, somnolence.

Topiramate
Non-EDL Tertiary

INDICATIONS
Resistant partial seizures, LGS- I.S, Generalized Tonic-Clonic Seizures (GTCS), severe myoclonic epilepsy of infancy.

AVAILABILITY
TABLETS 25, 50, 100 and 200 mg.

DOSE
Oral
Initial dose: 0.5 - 1 mg/kg/day (two divided doses) increase by 0.5 - 1 mg/kg/day at 1 to 2 week intervals, maintenance dose usually 5 – 9 mg/kg/day, max.-24 mg/kg/day.
For prophylaxis of migraine headache: 100 mg/ day.

PRECAUTIONS
Pregnancy (Appendix 7c) ; interactions (Appendix 6c)

ADVERSE EFFECTS
Anorexia, weight loss, cognitive slowing and behavior changes, difficulty with memory, somnolence, dizziness, ataxia, fatigue, kidney stones (1.5%) 2-4 times higher than general population, paresthesias, liver functions- in 1% transient and mild enzymes, metabolic acidosis, Ac. myopia and sec, angle glaucoma- mostly at start, oligohydrosis, hyperthermia and sec rash, hyperammonemia and encephalopathy with concomitant valproic acid use, behavioral reactions (26%), Most frequent such as aggressiveness, hyperactivity, excitement, anxiety, obsessive behaviour, cognitive delay of various degree more in children than adults because of preexisting behavioral problems in children with drug resistant epilepsy.

Vigabatrin
Non-EDL Tertiary

INDICATIONS
Infantile spasms, refractory partial seizures with or without secondary generalization.

AVAILABILITY
CAPSULES 500 mg.

DOSE
Initial dose- 40 mg/kg/day in two divided doses, increase to 80-100 mg/kg/day. In infantile spasms- Initial dose 40-50 mg/ kg/day increase by 50 mg/kg/day till spasm control or to 150-200 mg/kg/day.

PRECAUTIONS
Pregnancy (Appendix 7c)

ADVERSE EFFECTS
Psychosis (5%), behavioral problems, hyperactivity (most common cause for discontinuation), confusion, fatigue, insomnia, ataxia, drowsiness, weight gain, facial oedema, GIT upset (dose related), no effect on cognition. Chronic toxicity-most serious: persistent ncentric visual field defects in 1/3rd cases (rarely, reversible with early withdrawal), many patients are asymptomatic.
Zonisamide  
Non-EDL Tertiary

INDICATIONS
Add-on in partial seizures, primary generalized tonic clonic seizures, myoclonic epilepsy, absence seizures, LGS, infantile spasms.

AVAILABILITY
CAPSULES 25, 50 and 100 mg; Tablet 100 mg.

DOSE
Oral
Child- Initial dose - 2-4 mg/kg/day divided twice daily, with increments at 2- week intervals to 6-8 mg/kg/day and a possible maximum of 12 mg/kg/day.

PRECAUTIONS
Pregnancy (Appendix 7c)

ADVERSE EFFECTS
Drowsiness, anorexia, ataxia, fatigue (dose related), photosensitivity; cognitive effects reversible psychotic effects, behavioral abnormalities, abnormal thinking, irritability (Do slow titration); weight loss, renal stones (mostly small); idiosyncratic-in 1.4% skin rash (including SJS, TEN), blood dyscrasias, hepatic failure; oligohidrosis and hyperthermia (more in children).
SECTION - 6
ANTI INFECTIVE DRUGS

Anti-helmenthis
Cestode Infections:
Cestode infections (tapeworms) include intestinal taeniasis and cysticercosis, hymenolepiasis (dwarf tapeworm), diphyllobothriasis and echinococcosis (hydatid disease). Cysticercosis is a systemic infection caused by the larval form (cysticercus) of *Taenia solium*. Neurocysticercosis occurs when the infection involves the brain. In man, echinococcosis is due to the larval stage of *Echinococcus granulosus* or *E. multilocularis*. The larvae (oncospheres) develop by expansion (cystic echinococcosis) or tumour-like infiltration (alveolar echinococcosis), respectively, in the liver, lungs, or other organs.

1. Diphyllobothriasis:
In diphyllobothriasis, niclosamide or praziquantel in a single dose is highly effective. Hydroxocobalamin and folic acid supplements may also be required.

2. Echinococcosis:
In echinococcosis, surgery (or, if this is not possible, a technique such as ‘puncture-aspiration-injection-reapiration’) is the treatment of choice for operable cystic disease due to *Echinococcus granulosus* but chemotherapy with benzimidazoles, such as mebendazole and albendazole, may be of value as adjunctive therapy. Alveolar echinococcosis due to *E. multilocularis* requires both surgery and long-term treatment with either mebendazole or albendazole to inhibit spread of the infection. In animal studies, albendazole and mebendazole have been found to be teratogenic. They are contraindicated for the treatment of cestode infections in pregnancy; pregnancy should be excluded before treatment with albendazole (non-hormonal contraception during and for 1 month after treatment). For single-dose or short-term use in pregnancy.

3. Hymenolepiasis:
In hymenolepiasis, praziquantel is more effective than niclosamide, although resistance to praziquantel has been reported. Repeated treatment may be necessary to cure intense infections or to eliminate the parasite within a family group or institution.

4. Taeniasis:
In taeniasis, praziquantel is well tolerated and extensively absorbed and kills adult intestinal taenia worms in a single dose. Praziquantel also kills *T. solium* cysticerci when taken for 14 days in high doses. It thus offers the prospect of a cure for neurocysticercosis, which has been treatable only by surgery, anti-inflammatory corticosteroids and anticonvulsants. However, because dying and disintegrating cysts may induce localized cerebral oedema, treatment with praziquantel must always be undertaken in a hospital setting. In addition, a corticosteroid is usually given to reduce the inflammatory response. Albendazole also kills neurocysticerci when given daily for one month; a corticosteroid or an antihistamine is also given to reduce any inflammatory reaction. The longer-established niclosamide acts only against the adult intestinal
worms. Cestode infections due to *T. solium*, occurring during pregnancy should always be treated immediately (with praziquantel or niclosamide, but not with albendazole) because of the risk of cysticercosis.

**Intestinal Nematode Infections:**

Intestinal nematode infections include ascariasis, capillariasis, enterobiasis, hookworm infection, strongyloidiasis, trichostrongyliasis and trichuriasis.

1. **Ascariasis:**

Ascariasis is an infection, usually of the small intestine, caused by *Ascaris lumbricoides* (roundworm). Single doses of levamisole or pyrantel are effective; the broad-spectrum anthelminthics, albendazole or mebendazole are also effective.

2. **Capillariasis:**

Capillariasis is caused by infection of the intestine with *Capillaria philippinensis*. Prolonged treatment with mebendazole or albendazole offers the only prospect of cure.

3. **Enterobiasis:**

Enterobiasis is an infection of the large intestine caused by *Enterobius vermicularis* (pinworm, threadworm). All household members should be treated concurrently with a single dose of mebendazole, albendazole or pyrantel. Since reinfection readily occurs, at least one further dose should be given 2-4 weeks later. Piperazine is also effective but must be taken regularly for at least 7 consecutive days.

4. **Hookworm Infections:**

Hookworm infections are caused by *Ancylostoma duodenale* (ancylostomiasis) and *Necator americanus* (necatoriasis); they are a major cause of iron-deficiency anaemia in the tropics and sub-tropics. Ideally all cases of hookworm infection should be treated. However, when this is impracticable, priority should be given to women in second- and third-trimester of pregnancy, children and debilitated patients. In hookworm, broad-spectrum anthelminthics are preferred wherever other nematode infections are endemic. Both mebendazole and albendazole are effective. In animal studies, albendazole and mebendazole have been found to be teratogenic. There is some evidence to suggest that the use of mebendazole in pregnancy is not associated with an increased incidence of adverse effects on the fetus. However, neither mebendazole nor albendazole should be used during the first trimester of pregnancy to treat nematode infections. Both drugs are contraindicated for the treatment of cestode infections in pregnancy. Levamisole is effective in the treatment of mixed Ascaris and hookworm infections and pyrantel has been highly effective in some community-based control programmes, although several doses are often needed to eliminate *Necator americanus* infection. Patients with iron-deficiency anaemia caused by hookworm infection require supplementary iron salts and should receive ferrous sulphate (200 mg daily for adults) for at least 3 months after the haemoglobin concentration of 12g/100 ml is obtained.

5. **Strongyloidiasis:**

Strongyloidiasis is an infection of the small intestine caused by *Strongyloides stercoralis*. All infected patients should be treated. Ivermectin in a single dose of 200 μg/kg or 200 μg/ kg/day
on two consecutive days is the treatment of choice for chronic strongyloidiasis but it may not be available in all countries. Albendazole 400 mg once or twice daily for 3 days is well tolerated by both adults and children aged over 2 years and it may eradicate up to 80% of infections. Mebendazole has also been used but, to be effective, it must be administered for longer periods as it has a limited effect on larvae and hence the prevention of autoinfection.

6. Trichostrongyliasis:
Trichostrongyliasis is an infection of the small intestine caused by *Trichostrongylus* spp. In symptomatic trichostrongyliasis, a single dose of pyrantel (10 mg/kg) or albendazole (400 mg) is effective.

7. Trichuriasis:
Trichuriasis is an infection of the large intestine caused by *Trichuris trichiura* ( whipworm). Chemotherapy is required whenever symptoms develop or when faecal samples are found to be heavily contaminated (up to 10,000 eggs per gram). A single dose of albendazole (400 mg) or mebendazole (500 mg) can be effective in mild to moderate infections; severe infections require a 3-day course.

**Tissue Nematode Infections:**

Tissue nematode infections include angiostrongyliasis, anisakiasis, cutaneous larva migrans, dracunculiasis, trichinellosis and visceral larva migrans.

1. Angiostrongyliasis:
Angiostrongyliasis is caused by infection with the larvae of the rat lungworm, *Parastrongylus cantonensis* (*Angiostrongylus cantonensis*). Symptomatic treatment pending spontaneous recovery is often all that is required.

2. Anisakiasis:
Anisakiasis is caused by infection with seafood containing larvae of *Anisakis, Contracaecum* or *Pseudoterranova* spp. In anisakiasis, anthelminthic treatment is rarely necessary. Prevention is dependent upon informing communities of the hazards of eating raw or inadequately prepared salt-water fish; and early evisceration of fish after capture and freezing of seafood at -20°C for at least 60 h before sale.

3. Cutaneous Larva Migrans:
Cutaneous larva migrans (creeping eruption) is caused by infection with larvae of animal hookworms, usually *Ancylostoma braziliense* and *A. caninum* which infect cats and dogs. Albendazole in a single dose of 400 mg is effective.

4. Dracunculiasis:
Dracunculiasis (dracontiasis, guinea-worm infection) is caused by infection with *Dracunculus medinensis*, acquired through drinking water containing larvae that develop in small freshwater crustaceans. Metronidazole (25 mg/kg daily for 10 days, with a daily max. of 750 mg for children) provides rapid symptomatic relief. It also weakens the anchorage of the worms in the subcutaneous tissues and they can then be removed by traction. However, since it has no effect on the larvae of pre-emergent worms, it does not immediately prevent transmission.
5. Trichinellosis:
Trichinellosis (trichinosis) is caused by infection with the larvae of *Trichinella spiralis*. Each case of confirmed or even suspected trichinellosis infection should be treated in order to prevent the continued production of larvae. In both adults and children, mebendazole (200 mg daily for 5 days), albendazole (400 mg daily for 3 days) and pyrantel (10 mg/kg daily for 5 days) are all effective. Prednisolone (40-60 mg daily) may be needed to alleviate the allergic and inflammatory symptoms.

6. Visceral Larva Migrans:
Visceral larva migrans (toxocariasis) is caused by infection with the larval forms of *Toxocara canis* and less commonly, *T. cati* (which infect dogs and cats). Treatment should be reserved for symptomatic infections. A 3 week oral course of diethylcarbamazine kills the larvae and arrests the disease, but established lesions are irreversible. To reduce the intensity of allergic reactions induced by dying larvae, dosage is commonly commenced at 1 mg/kg twice daily and raised progressively to 3 mg/kg twice daily (adults and children). Ocular larva migrans occurs when larvae invade the eye, causing a granuloma which may result in blindness. In order to suppress allergic inflammatory responses in patients with ophthalmic lesions, prednisolone should be administered concurrently, either topically or systemically.

**Albendazole**
EDL-D12,13 Universal

**AVAILABILITY**
- CHEWABLE/PLAIN TABLET 150, 200, 400 mg & 1.5g;
- CAPSULE 400 mg;
- ORAL SUSPENSION 200 mg/5 ml;
- SYRUP 200 mg/5 ml;
- DROPS 10 ml (200 mg/ml)

**DOSE**
- Oral Adult and child above 2 years- 400 mg daily as a single dose. Strongyloidiasis, taeniasis and H. nana infection: 400 mg once daily is given for 3 consecutive days. Hydatid disease: 400 mg twice daily with meals for 28 days (therapy may be repeated after 14 days in three cycles). Child 1 to 2 years: 200 mg as a single dose.

**INDICATION**
Echinococcus multilocularis and E. granulosus infections prior to or not amenable to surgery; neurocysticercosis; nematode infections; filariasis; ascariasis, hookworm infections, strongyloidiasis, enterobiasis, trichuriasis, trichostrongyliasis and capillariasis; cestode infections; tissue nematode infections.

**CONTRAINDICATION**
Pregnancy, adequate measures must be taken for non-hormonal contraceptive during and one month after therapy; hypersensitivity.

**PRECAUTION**
Pregnancy (see notes above and Appendix 7c); liver impairment, increased intracranial pressure; seizures; monitor blood count and liver function.

**ADVERSE EFFECTS**
Gastrointestinal discomfort; headache; adverse effects associated with use in cestode infections; reversible alopecia; leucopenia, neurocysticercosis; Steven’s Johnson syndrome.

**Mebendazole**
EDL-D324, 325 Secondary hospitals

**AVAILABILITY**
- TABLET 100 mg;
- ORAL SUSPENSION 100 mg/5 ml.
DOSE
Oral Adult and child over 2 years: Threadworm infection: 100 mg single dose. If re-infection occurs second dose may be needed after 2 weeks. Whip worm, roundworm and hookworm infection: 100 mg twice daily for 3 days.

INDICATION
Echinococcus granulosus and E. multilocularis infections before surgery or not amenable to surgery; nematode infections

CONTRAINDICATION
Pregnancy; lactation; hypersensitivity; patients with CNS disorders.

PRECAUTION
Pregnancy (Appendix 7c; see also notes above); lactation; interactions (Appendix 6c, 6d); expulsion of ascaris from mouth or nose; monitor blood count or hepatic function.

ADVERSE EFFECTS
Gastrointestinal disturbances; headache and dizziness; adverse effects associated with use in cestode infections; abdominal pain, diarrhoea; rashes, urticaria, angioedema.

Anti-filarials
Loiasis:
Loiasis is an infection with the filarial nematode Loa loa and is transmitted by the biting of tabanid fly Chrysops. Diethylcarbamazine is effective against both adult worms and larvae; a single weekly dose is normally effective as prophylaxis. During individual treatment, particularly of persons with heavy microfilaraemia (>50 000 microfilariae/ml blood), a condition simulating meningoencephalitis occasionally occurs. This probably results from sludging of moribund microfilariae within cerebral capillaries. The frequency of meningoencephalitis associated with diethylcarbamazine therapy of loiasis is reported as 1.25%, with a mortality rate of about 50% in affected patients; treatment with diethylcarbamazine should be stopped at the first sign of cerebral involvement (and specialist advice sought). Permanent cerebral damage is common among patients who survive and this possibility should be considered when deciding on treatment. Treatment of heavily infected patients should thus begin at low dosage and corticosteroid and antihistamine cover should be provided for the first 2 to 3 days.

Lymphatic Filariasis:
Lymphatic filariasis is caused by infection with Wuchereria bancrofti (bancroftian filariasis), Brugia malayi or B. timori (brugian filariasis). Occult filariasis (tropical pulmonary eosinophilia) is a clinical variant of W. bancrofti infection. Individual treatment with diethylcarbamazine which has both microfilaricidal and macrofilaricidal activity is effective. Total cumulative dosages of 72 mg/kg are generally recommended for Wuchereria bancrofti infections with half this dose used for Brugia malayi and B. timori infections. In all cases treatment is best initiated with smaller doses for 2-3 days to avoid the danger of immunological reactions. Rigorous hygiene to the affected limbs with adjunctive measures to minimize infection and promote lymph flow is important for reducing acute episodes of inflammation. In communities where filariasis is endemic, annual administration of single doses of albendazole 400 mg with either diethylcarbamazine (6 mg/kg) or ivermectin (200 μg/kg) is effective for interrupting transmission; this treatment is continued for at least 5 years. Trials in India and China have shown that the consistent use for 6-12 months of table salt containing diethylcarbamazine 0.1% can eliminate W. bancrofti; a concentration of 0.3% for 3-4 months may be required where B. malayi is endemic.
Diethylcarbamazine

**EDL-D174,175 PHC**

**INDICATIONS**
- Treatment of loiasis; prophylaxis of loiasis in temporary residents in endemic areas; tissue nematode infections; lymphatic filariasis; toxocariasis.

**AVAILABILITY**
- TABLETS 50 and 100 mg; SYRUP 5 mg/ml and 120 mg/5 ml.

**DOSE**
- Oral
- Adult and child- 11 mg/kg body weight daily in three divided doses on the first day. Thereafter increase gradually to 6 mg/kg body weight given after food daily for two to three days. Hookworm infection: treat for 21 days. Filariasis: 2 mg/kg body weight is given three times a day for 3 to 4 weeks. 1 mg/kg body weight for an adult of 50 kg. Treatment may be repeated once after 6 months.

**CONTRAINDICATIONS**
- Pregnancy (delay treatment until after delivery); infants, elderly, debilitated (usually excluded from mass treatment programmes; see also Precautions); cardiac disease, hypersensitivity, impaired renal function.

**PRECAUTIONS**
- Renal impairment; cardiac disorders; other severe acute diseases-delay diethylcarbamazine treatment until after recovery; risk of meningoencephalitis in severe infection (see notes above).

**Adverse Effects**
- Headache, dizziness, drowsiness, nausea and vomiting; immunological reactions, within a few hour of the first dose, subsiding by fifth day of treatment and including fever, headache, joint pain, dizziness, anorexia, malaise, nausea and vomiting, urticaria and asthma in asthmatics (similar to Mazzotti reaction), induced by disintegrating microfilariae; microencephalitis (with heavy microfilaraemia, see notes above); reversible proteinuria; enlargement of lymph nodes.

**STORAGE**
- Store protected from moisture.

Ivermectin

**EDL-D296 Tertiary**

**AVAILABILITY**
- TABLETS 3, 6, 9 and 12 mg; INJECTION 10 ml (0.1% w/v).

**DOSE**
- Oral Strongyloidosis: 200 μg/kg of body weight once daily for 1-2 days. Lymphatic filariasis: 400 μg/kg of body weight simple annual dose for 4-6 years. Scabies and pediculosis: 150-200 μg/kg of body weight single oral dose highly effective. Second dose may be required 7-10 days later.

**INDICATION**
- Nematodal infections such as ascariasis, trichuriasis, strongyloidiasis, enterbiasis, lymphatic filariasis, scabies and pediculosis.

**CONTRAINDICATION**
- Hypersensitivity, CNS disorders, pregnancy, meningitis, trypanosomiasis, seizures, contraindicated to children below the age of < 5 years old or under 15 kg body weight.

**PRECAUTION**
- Concurrent Loa Loa infection, impaired blood-brain barrier function, pregnancy (Appendix 7c); lactation, hepatic, cardiovascular, renal or pulmonary disease, anaemia, coagulation disorder, severe asthma, interactions (Appendix 6c)

**ADVERSE EFFECTS**
- Nausea, vomiting, constipation, abdominal pain and fatigue, rash, arthralgia, fever, myalgia, asthenia, hypotension, tachycardia, edema, lymphadenopathy, sore throat, cough, headache,
somnolence, transient eosinophilia, dizziness, diarrhoea, pruritus, orthostatic hypotension, lymph-node tenderness, rare but serious adverse effects such as marked disability and encephalopathies in patients coinfected with heavy burdens of Loa microfilaria.

**Anti-bacterials**

**Beta-Lactams:**
Beta-lactam antibiotics including penicillins, cephalosporins and carbapenems share a common structure; they are bactericidal, their mechanism of action resulting from inhibition of peptidoglycan, a mucopeptide in bacterial cell walls. Benzylpenicillin and phenoxyethylpenicillin are active against susceptible strains of Gram-positive bacteria and Gram-negative bacteria, spirochaetes and actinomycetes, but are inactivated by penicillinase and other beta-lactamases. Benzathine benzylpenicillin and procaine benzylpenicillin are long acting preparations which slowly release benzylpenicillin on injection. A range of penicillins with improved stability to gastric acid and penicillinases have been produced by substitution of the 6-amino position of 6-aminopenicillanic acid. Cloxacillin is an isoxazoyl penicillin which is resistant to staphylococcal penicillinase. Broad-spectrum penicillins such as ampicillin are acid-stable and active against Gram-positive and Gram-negative bacteria, but are inactivated by penicillinase. Beta-lactamase inhibitors such as clavulanic acid are often necessary to provide activity against beta-lactamases produced by a wide range of both Gram-negative and Grampositive bacteria.

Cephalosporins are classified by generation, with the first generation agents having Gram-positive and some Gram-negative activity; the second generation drugs have improved Gram-negative activity and the third generation cephalosporin have a wider spectrum of activity, although may be less active against Gram-positive bacteria than first generation drugs, but they are active against Gram-negative Enterobacteriaceae and *Pseudomonas aeruginosa*.

Carbapenems are semisynthetic derivatives of *Streptomyces cattleya*. They have a broad spectrum of activity and are stable to most penicillinases. They should be reserved for severe infections resistant to other antibiotics.

Penicillins may cause encephalopathy due to cerebral irritation. This rare, but serious adverse effect may result from very high doses or in severe renal failure. Penicillins should not be given by intrathecal injection because they can cause encephalopathy which may be fatal.

**Hypersensitivity:**

The most important adverse effect of penicillins is hypersensitivity which causes rashes and, occasionally anaphylaxis, which can be fatal. A careful history should be taken with regard to Anti-Infectives previous allergic reactions. If rash develops, another antimicrobial should be substituted. Allergic reactions to penicillins occur in 1-10% of exposed individuals, while anaphylactic reactions occur in fewer than 0.05% of treated patients. Individuals with a history of anaphylaxis, urticaria or rash immediately after penicillin administration are at risk of immediate hypersensitivity to penicillin. These individuals should not receive penicillin, rather a cephalosporins or another beta-lactam antibiotic may be used. Patients who are allergic to one penicillin will be allergic to them all because the hypersensitivity is related to the basic penicillin
structure and about 10% of penicillin-sensitive patients will be allergic to cephalosporins and other beta-lactams. Individuals with a history of a minor rash (a non-confluent rash restricted to a small area of the body) or a rash occurring more than 72 h after penicillin administration are possibly not allergic to penicillin and in these individuals a penicillin should not be withheld unnecessarily for a serious infection; however, the possibility of an allergic reaction should be borne in mind and facilities should be available for treating anaphylaxis.

**Ampicillin, Amoxycillin, Amoxycillin with Clavulanic Acid and Cloxacillin:**

**Ampicillin** is active against certain Gram-positive and Gram-negative organisms. It is used to treat a wide range of infections including otitis media, respiratory-tract and urinary-tract infections and gonorrhoea due to susceptible bacteria. However, ampicillin is inactivated by penicillinases including those produced by *Staphylococcus aureus* and by common Gram-negative bacilli such as *Escherichia coli*; many strains of *Haemophilus influenzae*, *Moraxella catarrhalis*, *Neisseria gonorrhoeae* and *Salmonella* and *Shigella* spp. are resistant. There are geographical variations in the incidence of resistance and an awareness of local patterns is important. In some areas, oral use should be restricted to treatment of *Shigella* infections; it is given in an oral dose of 1g every 6 h for 7-10 days.

**Amoxycillin** has a similar spectrum of activity to ampicillin, but is also inactivated by penicillinases. However, it is better absorbed after oral administration than ampicillin and higher plasma and tissue levels are achieved. Amoxycillin is preferred to ampicillin for the treatment of some infections including otitis media and respiratory-tract and urinary-tract infections.

**Clavulanic acid** is a beta-lactamase inhibitor. It has no significant antibacterial activity but in combination with Amoxycillin widens Amoxycillin’s spectrum of activity and allows its use against Amoxycillin-resistant strains of bacteria. It is used in respiratory-tract, genito-urinary and abdominal infections, cellulitis, animal bites and dental infections.

Cloxacillin is used to treat infections due to penicillinase-producing staphylococci which are resistant to benzylpenicillin. It is acid-stable and may therefore be given by mouth as well as by injection. These antibiotics may also be administered with an aminoglycoside to increase their spectrums of activity. The penicillin and aminoglycoside should not be mixed before or during administration, because loss of aminoglycoside activity can occur on mixing.

**Benzylpenicillin and Phenoxymethylpenicillin:**

Benzylpenicillin remains an important and useful antibiotic but it is inactivated by bacterial beta-lactamases. It is effective for many streptococcal (including pneumococcal), gonococcal and meningococcal infections and also for anthrax, diphtheria, gas gangrene, leptospirosis, tetanus and treatment of Lyme disease in children. Pneumococci, meningococci and gonococci often have decreased sensitivity to penicillin and benzylpenicillin is no longer the first choice for pneumococcal meningitis. Benzylpenicillin is given by injection as it is inactivated by gastric acid and absorption from the intestinal tract is low.

Depot preparations are used when therapeutic concentrations need to be sustained for several h. Benzathine benzylpenicillin or procaine benzylpenicillin provides a tissue depot from which the drug is slowly absorbed over a period of 12 hour to several days. They are the preferred choice for the treatment of syphilis or yaws. Phenoxymethylpenicillin is suitable for oral administration; it has a similar spectrum of activity but is less effective than benzylpenicillin. It
should not be used for serious infections because absorption can be unpredictable and plasma concentrations variable.

**Cephalosporins and Imipenem with Cilastatin:**

**Ceftazidime** and **ceftriaxone** are third generation cephalosporins. Ceftriaxone is used for serious infections such as septicaemia, pneumonia and meningitis; it is used as a reserve antimicrobial to treat meningitis due to *Streptococcus pneumoniae* in some areas where penicillin resistance is found. Ceftazidime is active against *Pseudomonas aeruginosa* and other Gram-negative bacteria; it is used in the treatment of pseudomonal infections and in some areas is restricted to use only where gentamicin resistance is high. **Imipenem** is a broad spectrum antibiotic. As it is partially inactivated by enzymatic activity in the kidney, it is administered with **cilastatin** which inhibits the renal metabolism of imipenem. It is active against many aerobic and anaerobic Gram-positive and Gram-negative bacteria; in some areas it is reserve agent for the treatment of infections due to *Acinetobacter* spp. and *P. aeruginosa*, which are resistant to other more usual treatments.

**Quinolones:**

Ciprofloxacin is active against both Gram-positive and Gram-negative bacteria. It is particularly active against salmonella, shigella, campylobacter, neisseria, Bacillus anthracis and pseudomonas. It is also active against chlamydia and some mycobacteria. Most anaerobic organisms are not susceptible. Ciprofloxacin is used with doxycycline and metronidazole to treat pelvic inflammatory disease. Nalidixic acid is an older quinolone effective in uncomplicated urinary-tract infections and, in the treatment of shigella in areas where it remains susceptible.

**Tetracyclines:**

Doxycycline is a tetracycline and is a broad-spectrum antibiotic effective for conditions caused by chlamydia, rickettsia, brucella and the spirochaete, Borrelia burgdorferi (Lyme disease). It is the preferred tetracycline since it has a more favourable pharmacokinetic profile than tetracycline. It is deposited in growing bone and teeth causing staining and occasionally dental hypoplasia. It should not be given to children under 8 years or pregnant women; in some countries, use in children under 12 years is contraindicated.

**Aminoglycosides:**

Aminoglycosides including gentamicin are bactericidal and active against some Gram-positive and many Gram-negative organisms including *Pseudomonas aeruginosa*. Aminoglycosides are not absorbed from the gut and must therefore be given by injection for systemic infections. Excretion is mainly by the kidney and accumulation occurs in renal impairment. Use of gentamicin should be restricted to trained health personnel and care must be taken to ensure correct dosage and duration of treatment are not exceeded, because most adverse effects are dose related. The most important adverse effects are ototoxicity and nephrotoxicity and they are most common in the elderly and in patients with renal impairment. These groups and, if possible, all patients should be monitored for ototoxicity by audiometry. If there is impairment of renal function the dose interval must be increased; in severe renal impairment, the dose should also be reduced. Serum concentration monitoring avoids both excessive and subtherapeutic concentrations and can prevent toxicity and ensure efficacy. If possible serum
concentrations should be monitored in all patients, but must be measured in infants, the elderly, in obesity, in cystic fibrosis, in high-dosage regimens, in renal impairment, or if treatment lasts for longer than 7 days.

For most infections, doses of up to 5 mg/kg daily in divided doses are used if renal function is normal; higher doses are used occasionally for serious infections. Loading and maintenance doses are based on the patient’s weight and renal function (for example, using a nomogram) with adjustments based on plasma gentamicin concentration.

**Chloramphenicol:**
 Chloramphenicol is a potent broad-spectrum antibiotic. It is associated with serious haematological adverse effects and should be reserved for the treatment of severe infections, particularly those caused by Haemophilus influenza and typhoid fever. The oily suspension should be reserved for use in situations of catastrophic epidemics of meningococcal meningitis occurring mainly in sub-Saharan Africa, during which the medical services are overwhelmed by the epidemic and in which the overwhelming scale of the epidemic precludes any other form of antimicrobial therapy.

**Macrolides:**

**Erythromycin** is a macrolide; it has an antibacterial spectrum that is similar but not identical to penicillin and is used as an alternative in penicillin-allergic patients. It is effective in respiratory infections, whooping cough, legionnaires’ disease and campylobacter enteritis.

**Azithromycin** is more active than erythromycin against some Gram-negative organisms such as *Chlamydia trachomatis*. The concentration and persistence of azithromycin is much higher in the tissue than in plasma; a single dose of azithromycin is used in the treatment of uncomplicated genital Chlamydia and trachoma. Azithromycin is not recommended if there is a possibility of gonorrhoea because macrolide resistance emerges rapidly when it is used in this setting.

**Metronidazole:**
 Metronidazole has high activity against anaerobic bacteria and protozoa. Metronidazole by the rectal route is an effective alternative to the intravenous route when oral administration is not possible.

**Nitrofurantoin:**
 Nitrofurantoin is bactericidal in vitro to most Gram-positive and Gram-negative urinary-tract pathogens and it is used to treat acute and recurrent urinary-tract infections. It is also used prophylactically in chronic urinary-tract infections. Sulfonamides and Trimethoprim: The usefulness of sulfonamides is limited by an increasing incidence of bacterial resistance. For many indications they have been replaced by antibiotics that are more active and safer. Sulfadiazine is used in the prevention of rheumatic fever recurrence. Sulfamethoxazole is used in combination with trimethoprim because of their synergistic activity. In some countries, indications for the use of this combination have been restricted. The treatment of Pneumocystis carinii infections must only be undertaken with specialist supervision where there are appropriate monitoring facilities. Trimethoprim is also used alone for respiratory-tract infections and, in particular, for urinary-tract infections.
Vancomycin:
Vancomycin is not significantly absorbed from the gastrointestinal tract and must be given intravenously for systemic infections which cannot be treated with other effective, less toxic antimicrobials. It is used to treat serious infections due to Gram-positive cocci including methicillin-resistant staphylococcal infections, brain abscess, staphylococcal meningitis and septicaemia.

Amoxicillin anhydrous
EDL-D28,29,30 PHC

AVAILABILITY
tablets 250 mg, 500 mg; KID TABLETS 125, 250 mg; CAPSULES 250, 500 mg; DRY SYRUP 125 and 250 mg per 5 ml; INJECTION 1 ml ampoule (100 mg/ml), 250 mg/vial; drop 10 ml (100 mg/ml).

DOSE
Oral Adult- 250 mg every 8 h, double in severe infection. Otitis media: 1g every 8 h. Enteric fever: 2 to 4g daily in divided doses for 14 to 21 days. Intramuscular injection 500 mg every 8 h. Intravenous injection or infusion 500 mg every 8 h, increase to 1g every 6 h in case of severe infection. Child up to 10 years- 125 mg every 8 h, double in severe infections. Otitis media: 40 mg/kg body weight daily in three divided doses. Enteric fever: 50 to 100 mg/kg body weight in three divided doses for 14 to 21 days. Intramuscular injection 50 to 100 mg/kg body weight in divided doses. Intravenous injection or infusion 50 to 100 mg/kg body weight in divided doses.

INDICATION
Urinary-tract infections, upper respiratory tract infections, bronchitis; pneumonia; otitis media; dental abscess; osteomyelitis; Lyme disease in children; endocarditis prophylaxis; post-splenectomy prophylaxis; gynaecological infections; gonorrhoea; Helicobacter pylori eradication.

CONTRAINDICATION
Hypersensitivity to penicillins

PRECAUTION
History of allergy; renal impairment; erythematous rashes common in glandular fever, chronic lymphatic leukaemia and possibly HIV infection; lactation (Appendix 7b); interactions (Appendix 6b, 6c, 6d); possibility of super infection with mycotic pathogens, mononucleosis, hepatic impairment (Appendix 7a); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea and vomiting, diarrhoea; rashes (hypersensitivity or toxic response, may be serious reaction-discontinue treatment); hypersensitivity reactions including Steven’s Johnson syndrome, urticaria, angioedema, anaphylaxis, serum sickness-like reactions, haemolytic anaemia, interstitial nephritis; rarely, antibiotic-associated colitis; neutropenia, thrombocytopenia, coagulation disorders; rarely, central nervous system disorders including convulsions associated with high doses or impaired renal function; mucocutaneous candidiasis, with discolouration; agitation

Ampicillin Sodium
EDL-D41 PHC

AVAILABILITY
TABLETS 125 and 250 mg; CAPSULES 250, 500 mg and 1g; DRY SYRUP 125 and 250 mg/5 ml; INJECTION 100, 250 and 500 mg/ vial.

DOSE
Oral Adult- 250 mg to 1g every 6 h at least 30 min before food. Urinary tract infection Adult- 500 mg every 8 h. Children under 10 years- Half of adult dose. Intramuscular and intravenous
injection or infusion 500 mg every 4 to 6 h. Listeria meningitis (in combination with antibiotics); by intravenous infusion 2g every 4h for 10 to 14 days. Child- Half of the adult dose. Listeria meningitis (in combination with antibiotics); infants 1 to 3 months; 50 to 100 mg/kg body weight every 6 h. 3 months to 12 years; 100 mg/kg body weight every 76 h (max 12g daily).

**INDICATION**
- Mastoiditis;
- gynaecological infections;
- septicaemia;
- endocarditis;
- meningitis;
- cholecystitis;
- osteomyelitis;
- respiratory tract infection.

**CONTRAINDICATION**
- Hypersensitivity to penicillins

**PRECAUTION**
- History of allergy (see notes above);
- renal impairment (Appendix 7d);
- erythematous rashes common in glandular fever, acute or chronic lymphocytic leukaemia and cytomegalovirus infection;
- lactation, pregnancy

**ADVERSE EFFECTS**
- Nausea and vomiting, diarrhoea; rashes, high fever (hypersensitivity or toxic response-may be serious reaction, discontinue treatment);
- hypersensitivity reactions including urticaria, angioedema, anaphylaxis, serum sicknesslike reaction, haemolytic anaemia, interstitial nephritis (see also notes above); rarely, antibiotic-associated colitis; neutropenia, thrombocytopenia,

**Benzathine benzyl penicillin**

**EDL-D66 PHC**

**AVAILABILITY**
- Injectable suspension- 1200,000 units/2 ml.

**DOSE**
- Streptococcal URTI: 1.2 million unit (>27 kg) single dose (deep IM inj) after sensitivity test (AST); 0.6 million unit (<27 kg) single dose (deep IM inj) AST. Secondary prophylaxis of Rheumatic fever: 1.2 million unit (>27 kg) single dose (deep IM inj) after sensitivity test (AST) every 21 days; 0.6 million unit (<27 kg) single dose (deep IM inj) AST every 15 days. Syphilis: Primary, secondary, or early latent: Single dose of 2.4 million Unit IM; Late latent (or latent of uncertain duration), cardiovascular, or benign tertiary: 2.4 million Unit IM weekly for 3 weeks.

**INDICATION**
- Mild to moderate infections of upper respiratory tract due to susceptible streptococci, Syphilis, prophylaxis of rheumatic fever.

**CONTRAINDICATION**
- Hypersensitivity, neurosyphilis.

**PRECAUTION**
- Hypersensitivity to cephalosporins or/ and penicillins, elderly, infants, asthma, kidney disease, lactation

**ADVERSE EFFECTS**
- Hypersensitivity reactions such as exfoliative dermatitis, pain at injection site, thrombophlebitis of injected vein, diarrhoea, nausea, joint pain, angioedema, serum sickness like reactions; haemolytic anaemia, interstitial nephritis

**Benzyl penicillin(sodium or potassium salt)**

**EDL-D71 PHC**

**AVAILABILITY**
- Injectable suspension - 6, 12, 24 Lac units; Injectable suspension - 1200,000 units/2 ml.

**DOSE**
- Streptococcal URTI: 1.2 million unit (>27 kg) single dose (deep IM inj) after sensitivity test (AST); 0.6 million unit (<27 kg) single dose (deep IM inj) AST. Secondary prophylaxis of Rheumatic fever: 1.2 million unit (>27 kg) single dose (deep IM inj) after sensitivity test (AST) every 21 days; 0.6 million unit (<27 kg) single dose (deep IM inj) AST every 15 days. Syphilis: Primary,
secondary, or early latent: Single dose of 2.4 million Unit IM; Late latent (or latent of uncertain duration), cardiovascular, or benign tertiary: 2.4 million Unit IM weekly for 3 weeks.

**INDICATION**
- Mild to moderate infections of upper respiratory tract due to susceptible streptococci, syphilis, prophylaxis of rheumatic fever.

**CONTRAINDICATION**
- Hypersensitivity, neurosyphilis.

**PRECAUTION**
- Hypersensitivity to cephalosporins or/ and penicillins, elderly, infants, asthma, renal impairment (Appendix 7d) Lactation (Appendix 7b) and pregnancy (Appendix 7c)

**ADVERSE EFFECTS**
- Hypersensitivity reactions such as exfoliative dermatitis, pain at injection site, thrombophlebitis of injected vein, diarrhoea, nausea, joint pain, angioedema, serum sickness like reactions; haemolytic anaemia, interstitial nephritis.

**Cloxacillin Sodium**
- **EDL-D133,135 PHC**

**AVAILABILITY**
- CAPSULES 250 and 500 mg; INJECTION 250 and 500 mg/vial; DRY SYRUP 125 mg/5 ml

**DOSE**
- Adult- 250-500 mg every 6 h at least 30 min. before food. Osteomyelitis; upto 8g daily in 2 to 3 divided doses. Surgical prophylaxis; 1 to 2g at induction thereafter up to 4 further doses each of 500 mg may be given every 6h. Slow intravenous injection or infusion Adult- Surgical prophylaxis; 1 to 2g at induction thereafter up to 4 further doses each of 500 mg may be given every 6 h. Child- High risk procedures; Under 2 years; quarter adult dose. 2 to 10 years; half adult dose.

**INDICATION**
- Multibacillary (MB) leprosy; type 2 lepra reactions; gram positive infection including resistant staphylococci

**CONTRAINDICATION**
- Hypersensitivity to penicillins

**PRECAUTION**
- History of allergy (see notes above); renal and hepatic impairment (Appendix 7a); heart failure; lactation (Appendix 7b); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
- Nausea and vomiting, diarrhoea; hypersensitivity reactions including urticaria, fever, joint pain, rashes, angioedema, anaphylaxis, serum sickness-like reactions, haemolytic anaemia, interstitial nephritis (see also notes above); neutropenia, thrombocytopenia, coagulation disorders; antibioticassociated colitis; hepatitis and cholestatic jaundice-may be delayed in onset; electrolyte disturbances; pain, inflammation, phlebitis or thrombophlebitis at injection sites.

**Cephalexin**
- **EDL-D95,96,97,98 Secondary hospital**

**INDICATIONS**
- Respiratory tract infections; otitis media; skin and skin structure infections; genitourinary tract infection; bone infection.

**AVAILABILITY**
- CAPSULES/TABLETS 125, 250 and 500 mg; 125 mg Kid tablets; 250 mg DT; DRY SYRUP 125 and 250 mg/5 ml.

**DOSE**
- To be given preferably on empty stomach.
Adult- 250 mg every 6 h or 500 mg every 8 to 12 h, increased to 1 to 1.5g every 6 to 8 h for severe infections. Prophylaxis of severe urinary tract infection: 125 mg at night.
Child- 25 mg/kg body weight daily in divided doses doubled for severe infections (max. 100 mg/kg body weight daily); Under 1 year: 125 mg every 12 h; 1 to 5 years: 125 mg every 8 h; 5 to 12 years: 250 mg every 8 h.

CONTRAINDICATIONS
Cephalosporin hypersensitivity.

PRECAUTIONS
Sensitivity to beta-lactam antibacterials (avoid if history of immediate hypersensitivity reaction, renal impairment; lactation; false positive urinary glucose (if tested for reducing substances) and false positive Coombs’ test; poor nutritional state; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Diarrhoea and rarely, antibiotic-associated colitis (more likely with higher doses), nausea and vomiting, abdominal discomfort, headache; allergic reactions including rashes, pruritus, urticaria, serum sickness-like reactions with rashes, fever and arthralgia and anaphylaxis; Stevens- Johnson syndrome, toxic epidermal necrolysis reported; disturbances in liver enzymes, transient hepatitis and cholestatic jaundice; other side-effects reported include eosinophilia and blood disorders (including thrombocytopenia, leucopenia, agranulocytosis, aplastic anaemia and haemolytic anaemia); reversible interstitial nephritis, hyperactivity, nervousness, sleep disturbances, hallucinations, confusion, hypertonia and dizziness; dyspnoea, colitis, increased blood urea, creatinine, alkaline phosphatase, bilirubin, LDH.

STORAGE
Store protected from light and moisture at a temperature not exceeding 30°C.

Ceftriaxone Sodium
EDL-D104,105 Secondary hospitals

INDICATIONS
Serious infections due to sensitive bacteria, including septicaemia, pneumonia and meningitis; surgical prophylaxis; prophylaxis of meningococcal meningitis; gonorrhoea; bone and joint infection.

AVAILABILITY
INJECTION 125, 250, 500 mg, 1g and 2g vial.

DOSE
Intramuscular and intravenous injection or infusion
Adult- Urinary tract infection, pneumonia, pelvic inflammatory disease, prophylaxis of surgical infections and meningitis: 4g initially once daily for 10 days or up to 72 h after fever disappears.
Typhoid: 4g daily for two days followed by 2g daily for next two days. 1 to 2g daily is used for any other type of condition.
Child- Meningitis: 75 to 100 mg/kg body weight for 7 to 9 days.
Typhoid: 5 mg/kg body weight for 7 days. 50 to 75 mg/kg body weight is used in case of any other condition (max 2g/day).

CONTRAINDICATIONS
Cephalosporin hypersensitivity; porphyria; neonates with jaundice, hypoalbuminaemia, acidosis or impaired bilirubin binding.

PRECAUTIONS
Penicillin sensitivity; severe renal impairment; hepatic impairment if accompanied by renal impairment (Appendix 7a); premature neonates; may displace bilirubin from serum albumin; treatment longer than 14 days, renal failure, dehydration or concomitant total parenteral nutrition-risk of ceftriaxone precipitation in gallbladder; lactation (but appropriate to use, see Appendix 7b); pregnancy (Appendix 7c); false positive urinary glucose (if tested for reducing substances) and false positive Coombs’ test; interactions (Appendix 6b, 6c); prophylactic
indication, patients with impaired vit K synthesis, monitoring of prothrombin time is recommended.

ADVERSE EFFECTS
Diarrhoea, nausea and vomiting, abdominal discomfort, headache; antibiotic-associated colitis (particularly with higher doses); allergic reactions including rashes, pruritus, urticaria, serum sickness-like reactions, fever and arthralgia and anaphylaxis; erythema multiforme, toxic epidermal necrolysis reported; transient hepatitis and cholestatic jaundice; elevation of SGOT and SGPT; eosinophilia and blood disorders (including thrombocytopenia, leukopenia, agranulocytosis, aplastic anaemia and haemolytic anaemia); reversible interstitial nephritis, hyperactivity, nervousness, sleep disturbances, confusion, hypertonia and dizziness; calcium ceftriaxone precipitates in urine (particularly in very young, dehydrated, or those who are immobilized) or in gall bladder-consider discontinuation if symptomatic; rarely, prolongation of prothrombin time, pancreatitis; local reaction, hypersensitivity.

STORAGE
Store protected from light at a temperature not exceeding 30°C.

Ceftazidime
EDL-D102,103 Tertiary

AVAILABILITY
INJECTION 250, 500 mg, 1g and 2g vial.

DOSE
Deep intramuscular and intravenous injection and infusion Adult- 1g every 8 h or 2g every 12 h. Severe infections: 2g every 12 h or 3g every 12 h (1g single dose by intravenous route). Immunocompromised or meningitis patients: 150 mg/kg body weight daily in 3 divided doses (max 6g daily) given by i.v route only. Elderly- Usual max dose of 3g daily. Child- Up to 2 months; 25 to 60g/kg body weight in two divided doses. Over 2 months: 30 to 100 mg/kg body weight in 2 to 3 divided doses.

INDICATION
Infections due to sensitive bacteria, especially those due to Pseudomonas spp. and including those resistant to aminoglycosides.

CONTRAINdICATION
Cephalosporin hypersensitivity; porphyria.

PRECAUTION
Penicillin sensitivity; renal impairment; lactation (Appendix 7b); false positive urinary glucose (if tested for reducing substances) and false positive Coombs’ test; interactions (Appendix 6b, 6c); pregnancy (Appendix 7c); fall in prothrombin activity, colitis.

ADVERSE EFFECTS
Diarrhoea, nausea, vomiting, abdominal discomfort, headache; rarely, antibiotic-associated colitis (particularly with higher doses); allergic reactions including rashes, pruritus, urticaria, serum sickness-like reaction, fever and arthralgia and anaphylaxis; erythema multiforme, toxic epidermal necrolysis reported; transient hepatitis, cholestatic jaundice; eosinophilia and blood disorders (including thrombocytopenia, leukopenia, agranulocytosis, aplastic anaemia and haemolytic anaemia); reversible interstitial nephritis; nervousness, sleep disturbances, confusion, hypertonia and dizziness; phlebitis, angioedema, myoclonia, candidiasis, transient elevation of blood urea and serum creatinine.
Imipenem Monohydrate + Cilastatin Sodium
EDL-D 274 Tertiary

AVAILABILITY
INJECTION
Imipenem + Cilastatin
125 mg + 125 mg vial
250 mg + 250 mg vial
500 mg + 500 mg vial
1g + 1g vial
2g + 2g vial

DOSE
Intravenous infusion in terms of imipenem
Adult- 2g daily in 2 to 3 divided doses. Less susceptible organism may be given up to 3 to 4 divided doses (max 4g daily).
Surgical prophylaxis: 1g for induction, repeated every three h, supplemented in high risk surgery by doses of 500 mg for 8 to 16 h.
Child- 3 months and older: 60 mg/kg body weight in fou

INDICATION
Severe aerobic and anaerobic Gram-positive and Gram-negative infections in hospital -acquired infections (not indicated for CNS infections), including infections caused by resistant Pseudomonas and Acinetobacter species.

CONTRAINDICATION
Hypersensitivity to beta-lactam antibiotics; local anaesthetics of the amide type and in patients with severe shock or heart block.

PRECAUTION
Renal impairment; CNS disorders, such as epilepsy; lactation (Appendix 7b); interactions (Appendix 6c); pregnancy (Appendix 7c)

ADVERSE EFFECTS
Nausea, vomiting, diarrhoea; antibiotic-associated colitis; taste disturbances; tooth or tongue discolouration, hearing loss; blood disorders, (decreased haematocrit, increased prothrombin time) positive Coombs’ test; allergic reactions including rash, pruritus, urticaria, erythema multiforme (Steven’s-Johnson syndrome), fever, anaphylactic reactions, rarely, toxic epidermal necrolysis, exfoliative dermatitis; myoclonic activity, convulsions, confusion and mental disturbances; slight increase in liver enzymes and bilirubin, rarely, hepatitis; increase in serum creatinine and blood urea; red coloration of urine in children; erythema, pain and induration and thrombophlebitis at injection sites; bone marrow depression.

Amoxicillin + Clavulanic Acid
EDL-D31,32,33,34,35,36,37 Secondary hospitals

AVAILABILITY
TABLETS
Amoxycillin + Clavulanic acid
500 mg + 125 mg
250 mg + 125 mg
875 mg + 125 mg
200 mg + 28.5 mg (DT)
CAPSULS
Amoxycillin + Clavulanic acid
500 mg + 125 mg
250 mg + 125 mg
SUSPENSION
Amoxycillin + Clavulanic acid
200 mg + 28.5 mg/5 ml
125 mg + 31.25 mg/5 ml
250 mg + 62.5 mg/5 ml
INJECTION
Amoxycillin + Clavulanic acid
250 mg + 50 mg
1g + 200 mg
125 mg + 25 mg
500 mg + 100 mg

DOSE
Oral Upper and lower respiratory tract infections, sinusitis, otitis media, skin and soft tissue infections, susceptible infections: Adult- 250-500 mg every 8 hours or 500-750 mg every 12 hours. Child- 125-250 mg every 8 hours; Children weighing <40 kg: 20-40 mg/kg/day in divided doses every 8 hours; Infants <3 months: up to 30 mg/kg/day in divided doses every 12 hours. Dental abscesses: Adult- 3 g as a single dose, followed by a second dose 8 hours later. Severe or recurrent respiratory tract infections: Adult-3 g twice daily. Child (2-6 years)- 5 ml twice daily; (7-12 years)- 10 ml twice daily before meals, upto 14 days (dose should be specified in terms of strength). Parenteral Susceptible infections and surgical prophylaxis: Adult- 500 mg every 8 hr. In severe infections, dose may be increased to 1 g every 6 hours, upto 14 days. Can be given via i.m or slow i.v over 3-4 minutes or i.v infusion over 30-60 minutes. Child: <10 years: 50-100 mg/kg/day in divided doses.

INDICATION
Treatment of infections caused by susceptible organisms, sinusitis, otitis media, dental abscesses, severe respiratory tract infections, urinary tract infections, skin and soft tissue infections, surgical prophylaxis.

CONTRAINDICATION
Hypersensitivity to penicillins, infectious mononucleosis, jaundice.

PRECAUTION
Renal impairment, hepatic dysfunction, patients on anticoagulant therapy, pregnancy (Appendix 7c), lactation, interactions (Appendix 6c).

ADVERSE EFFECTS
GI upset, mycosis, rash, nausea, vomiting, anaphylaxis, cholestatic jaundice, blood dyscracias, toxic epidermal necrolysis, convulsions, exfoliative dermatitis, Stevens Johnson syndrome, angioedema, hepatitis, tooth discolouration.

Amoxicillin
EDL-D560 PHC

AVAILABILITY
tablets 250 mg, 500 mg; KID TABLETS 125, 250 mg; CAPSULES 250, 500 mg; DRY SYRUP 125 and 250 mg per 5 ml; INJECTION 1 ml ampoule (100 mg/ml), 250 mg/vial; drop 10 ml (100 mg/ml).

DOSE
Oral Adult- 250 mg every 8 h, double in severe infection.Otitis media: 1g every 8 h. Enteric fever: 2 to 4g daily in divided doses for 14 to 21 days. Intramuscular injection 500 mg every 8 h. Intravenous injection or infusion 500 mg every 8 h, increase to 1g every 6 h in case of severe infection. Child up to 10 years- 125 mg every 8 h, double in severe infections. Otitis media: 40 mg/kg body weight daily in three divided doses. Enteric fever: 50 to 100 mg/kg body weight in
three divided doses for 14 to 21 days. Intramuscular injection 50 to 100 mg/kg body weight in divided doses. Intravenous injection or infusion 50 to 100 mg/kg body weight in divided doses.

INDICATION
Urinary-tract infections, upper respiratory tract infections, bronchitis; pneumonia; otitis media; dental abscess; osteomyelitis; Lyme disease in children; endocarditis prophylaxis; post-splenectomy prophylaxis; gynaecological infections; gonorrhoea; Helicobacter pylori eradication.

CONTRAINDICATION
Hypersensitivity to penicillins

PRECAUTION
History of allergy; renal impairment; erythematous rashes common in glandular fever, chronic lymphatic leukaemia and possibly HIV infection; lactation (Appendix 7b); interactions (Appendix 6b, 6c, 6d); possibility of super infection with mycotic pathogens, mononucleosis, hepatic impairment (Appendix 7a); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea and vomiting, diarrhoea; rashes (hypersensitivity or toxic response, may be serious reaction-discontinue treatment); hypersensitivity reactions including Steven’s Johnson syndrome, urticaria, angioedema, anaphylaxis, serum sickness-like reactions, haemolytic anaemia, interstitial nephritis; rarely, antibiotic-associated colitis; neutropenia, thrombocytopenia, coagulation disorders; rarely, central nervous system disorders including convulsions associated with high doses or impaired renal function; mucocutaneous candidiasis, with discolouration; agitation.
adolescents (see below); avoid exposure to excessive sunlight (discontinue if photosensitivity occurs); rarely, tendon damage-discontinue at first sign of pain or inflammation and rest affected limb; hepatic impairment; renal failure (Appendix 7d); avoid excessive alkalinity of urine and ensure adequate fluid intake as there is risk of crystalluria; interactions (Appendix 6c); cerebral arteriosclerosis, anxiety, paranoia, erythema, blistering. Use In Children. Ciprofloxacin causes arthropathy in the weight-bearing joints of immature animals and is therefore generally not recommended in children and growing adolescents. However, the significance of this effect in humans is uncertain and in some specific circumstances short-term use of ciprofloxacin in children may be justified. Ciprofloxacin is used for pseudomonal infections in cystic fibrosis (for children over 5 years) and for treatment and prophylaxis of anthrax. May impair ability to perform skilled tasks, for example operating machinery, driving.

ADVERSE EFFECTS
Nausea, vomiting, dyspepsia, abdominal pain, flatulence, diarrhoea (rarely, antibiotic-associated colitis), dysphagia, tremor, hyperglycaemia, headache, dizziness, sleep disorders, rash (rarely, erythema multiforme (Stevens-Johnson syndrome) and toxic epidermal necrolysis) and pruritus; vasculitis, erythema nodosum, petechiae, haemorrhagic bullae; less frequently anorexia, increase in blood urea and creatinine; drowsiness, restlessness, asthenia, depression, confusion, hallucinations, convulsions, paraesthesia; photosensitivity, hypersensitivity reactions including fever, urticaria, angioedema, arthralgia, myalgia and anaphylaxis; blood disorders (including eosinophilia, leukopenia, thrombocytopenia), altered prothrombin time; disturbances in vision, taste, hearing and smell, tinnitus; tenosynovitis; tachycardia, oedema, syncope, hot flushes and sweating; if psychiatric, neurological or hypersensitivity reactions (including severe rash) occur discontinue; arthralgia.

Doxycycline Hydrochloride
EDL-D192 PHC

AVAILABILITY
CAPSULES/TABLETS 50, 100, 150 and 200 mg; Syrup 25 mg/5 ml.

DOSE
Oral Adult- 200 mg on the first day then 100 mg daily. Severe infections including refractory urinary tract infection: 200 mg daily can be used. Early syphilis: 100 mg twice daily for 14 days and for latent syphilis 200 mg twice daily for 28 days is used. Uncomplicated genital Chlamydia, non gonococcal urethritis: 100 mg twice daily for 7 days. Child- Only if alternate antibacterial cannot be given 5 mg/kg body weight in two divided doses.

INDICATION
Supplement to quinine in treatment of multiple-medicine resistant P. falciparum malaria (where quinine resistance, in cases of hypersensitivity to sulfonamides); shortterm prophylaxis of multiple-medicine resistant P. falciparum malaria; bacterial infections.

CONTRAINDICATION
Cinchonism (tinnitus, headache, blurred vision, temporary blindness, altered auditory acuity, nausea, diarrhoea, hot and flushed skin, rashes, confusion); hypersensitivity reactions including angioedema; rarely, haemorrhage and asthma; hypoglycaemia (especially after parenteral administration); renal damage (culminating in acute renal failure and anuria); blood disorders; cardiovascular, gastrointestinal and CNS effects; very toxic in overdosage-immediate medical attention required; acute haemolytic anaemia.

PRECAUTION
Avoid exposure to sunlight or sunlampsphotosensitivity reported; renal impairment; hepatic impairment (Appendix 7a); lactation (Appendix 7b); interactions (Appendix 6a, 6b, 6c 6d); predisposition to candidiasis.
ADVERSE EFFECTS
Haemoglobinuria; optic neuritis; tinnitus; quinine resistant falciparum, pregnancy, lactation, prolonged QT interval.

**Erythromycin Stearate**

**EDL-D 202,203,204 PHC**

**AVAILABILITY**
TABLETS 125, 250 and 500 mg plain; 125 DT; SYRUP 125 mg/5 ml; ointment 2 and 3% w/w; cream 3% w/w.

**DOSE**
Oral Adult and child over 8 years: 250 to 500 mg every 6 h or 0.5 to 1g every 12 h up to 4g daily in severe infections. Child: 1 month to 2 years: 12.5 mg/kg body weight every 6 h; 2 to 8 years 250 mg every 6 h (doses doubled for severe infections). Early syphilis: 500 mg three times daily for 14 days.

**INDICATION**
Alternative to penicillin in hypersensitive patients; pneumonia; legionnaires' disease; syphilis; chancroid; chlamydia; nongonococcal urethritis; prostatitis; lymphogranuloma venereum; campylobacter enteritis; relapsing fever; diphtheria and whooping cough prophylaxis upper respiratory tract infection, acne vulgaris, sycosis, vulgaris.

**CONTRAINDICATION**
Hypersensitivity to erythromycin or other macrolides; porphyria; myasthenia gravis.

**PRECAUTION**
Hepatic impairment (Appendix 7a) and renal impairment (Appendix 7d); prolongation of the QT interval (ventricular tachycardia reported); pregnancy (Appendix 7c); (not known to be harmful); lactation (Appendix 7b); interactions

**ADVERSE EFFECTS**
Nausea, vomiting, abdominal discomfort, diarrhoea and (antibiotic-associated colitis); urticaria, rashes and other allergic reactions (rarely, anaphylaxis); reversible hearing loss after large doses; cholestatic jaundice, cardiac effects (including chest pain and arrhythmias), myasthenialike syndrome, erythema multiforme (Stevens-Johnson syndrome) and toxic epidermal necrolysis; burning sensation, itching, anorexia.

**Azithromycin**

**EDL-D60, 61 PHC**

**D62 Secondary hospitals**

**AVAILABILITY**
TABLETS 100, 250 and 500 mg; CAPSULES 250 and 500 mg; INJECTION 500 mg/vial DRY SYRUP 100, 200 mg/5 ml.

**DOSE**
Oral Adult: 500 mg once daily for 3 days or 500 mg on first day then 250 mg once daily for 4 days. Child: over 6 months: 10 mg/kg body weight once daily for three days. Body weight 15 to 20 kg: 200 mg once daily for 3 days; body weight 26 to 35 kg: 300 mg daily for 3 days. Uncomplicated genital chlamydia infection and non-gonococcal infection: 500 mg once daily for 7 days.

**INDICATION**
Uncomplicated genital chlamydial infections and trachoma.

**CONTRAINDICATION**
Hepatic impairment hypersensitivity to erythromycin.

**PRECAUTION**
Pregnancy (Appendix 7c) and lactation (Appendix 7b); renal impairment, prolongation of QT interval (ventricular tachycardia reported); interactions (Appendix 6c, 6d); exacerbation of symptoms of myasthenia gravis; impaired hepatic function.
ADVERSE EFFECTS
Fewer gastrointestinal effects as compared to erythromycin, also anorexia, dyspepsia, constipation; dizziness, headache, drowsiness; photosensitivity; hepatitis, interstitial nephritis, acute renal failure, asthma, paraesthesia, convulsions and mild neutropenia reported; rarely, tinnitus, hepatic necrosis, hepatic failure and taste disturbances; flatulence, somnolence, angioedema; eczema, pharyngitis; arthralgia, conjunctivitis.

Gentamicin Sulphate
EDL-D242,243 Universal

AVAILABILITY
eye drops 0.3% w/v, cream 15g (0.1% w/w); INJECTION 2 ml ampoule (40 mg/ml), 2 and 10 ml vials (40 mg/ml).

DOSE
Intravenous infusion Once daily dose regime; 5 to 7 mg/kg body weight, then adjust as per serum gentamicin concentration. Intramuscular or slow intravenous injection over at least 3 min. Multiple daily dose regimen: 3 mg/kg body weight divided into 8 hly doses.Child- 2 weeks to 12 years; 2 mg/kg body weight 8 hly.

INDICATION
Pneumonia; cholecystitis; peritonitis; septicaemia; acute pyelonephritis; prostatitis; skin infections; pelvic inflammatory disease; endocarditis; meningitis; listeriosis; tularaemia; brucellosis; plague; surgical prophylaxis; ocular bacterial infection

CONTRAINDICATION
Myasthenia gravis.

PRECAUTION
Renal impairment (Appendix 7d), infants and elderly (dosage adjustment and monitor renal, auditory and vestibular function and serum-gentamicin concentrations); avoid prolonged use; conditions characterized by muscular weakness; significant obesity (monitor serum-gentamicin concentration closely and possibly reduce dose); see notes above; interactions (Appendix 6c); purulent discharge, discontinue if pain/inflammation becomes aggravated; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Vestibular and auditory damage, nephrotoxicity; rarely, hypomagnesaemia on prolonged therapy; antibiotic-associated colitis, also nausea, vomiting, rash; bacterial/ fungal corneal ulcers, ocular burning or irritation, thrombocytopenia, joint pain.

Metronidazole
EDL-344,346 PHC D345 Universal

AVAILABILITY
TABLETS 200 and 400 mg; SUSPENSION 200 mg/5 ml; INJECTION 100 ml infusion (5 mg/ ml).

DOSE
Oral Adult- Amoebiasis: 400 to 800 mg every 8 h for 5 to 7 days. GiardiasisL: 200 mg three times a day for 7 to 10 days or intravenous injection 500 mg 8 hly for 7 days. Child- Amoebiasis: Below 12 years; 7.5 mg/ kg body weight. 12 years and above; 35 to 50 mg/kg body weight daily in three divided doses.

INDICATION
Anaerobic bacterial infections including gingivitis, pelvic inflammatory disease, tetanus, peritonitis, brain abscess, necrotizing pneumonia, antibiotic-associated colitis, leg ulcers and pressure sores and surgical prophylaxis; bacterial vaginosis; tissue nematode infections; trichomonal vaginitis, amoebiasis and giardiasis; Helicobacter pylori eradication.

CONTRAINDICATION
Chronic alcohol dependence; neurological disease, blood dyscrasias, first trimester of pregnancy.
PRECAUTION
Disulfiram-like reaction with alcohol; hepatic impairment and hepatic encephalopathy (Appendix 7a); lactation (Appendix 7b); clinical and laboratory monitoring in courses lasting longer than 10 days; interactions (Appendix 6a, 6c, 6d); pregnancy (Appendix 7c); phenobarbitone, history of blood dyscrasias.

ADVERSE EFFECTS
Nausea, vomiting, unpleasant metallic taste, furred tongue and gastrointestinal disturbances; rarely, headache, drowsiness, dizziness, ataxia, darkening of urine, erythema multiforme, pruritus, urticaria, angioedema and anaphylaxis; abnormal liver function tests, hepatitis, jaundice; thrombocytopenia, aplastic anaemia; myalgia, arthralgia; peripheral neuropathy, epileptiform seizures; leukopenia on prolonged or high dosage regimens; anorexia, glossitis, dryness of mouth.

Metronidazole Benzoate
EDL-D347 PHC

AVAILABILITY
TABLETS 200 and 400 mg; SUSPENSION 200 mg/5 ml; INJECTION 100 ml infusion (5 mg/ml).

DOSE
Oral Adult- Amoebiasis: 400 to 800 mg every 8 h for 5 to 7 days. GiardiasisL: 200 mg three times a day for 7 to 10 days or intravenous injection 500 mg 8 hly for 7 days. Child- Amoebiasis: Below 12 years; 7.5 mg/ kg body weight. 12 years and above; 35 to 50 mg/kg body weight daily in three divided doses.

INDICATION
Anaerobic bacterial infections including gingivitis, pelvic inflammatory disease, tetanus, peritonitis, brain abscess, necrotizing pneumonia, antibiotic-associated colitis, leg ulcers and pressure sores and surgical prophylaxis; bacterial vaginosis; tissue nematode infections; trichomonal vaginitis, amoebiasis and giardiasis; Helicobacter pylori eradication.

CONTRAINDICATION
Chronic alcohol dependence; neurological disease, blood dyscrasias, first trimester of pregnancy.

PRECAUTION
Disulfiram-like reaction with alcohol; hepatic impairment and hepatic encephalopathy; lactation; clinical and laboratory monitoring in courses lasting longer than 10 days; interactions; pregnancy; phenobarbitone, history of blood dyscrasias.

ADVERSE EFFECTS
Nausea, vomiting, unpleasant metallic taste, furred tongue and gastrointestinal disturbances; rarely, headache, drowsiness, dizziness, ataxia, darkening of urine, erythema multiforme, pruritus, urticaria, angioedema and anaphylaxis; abnormal liver function tests, hepatitis, jaundice; thrombocytopenia, aplastic anaemia; myalgia, arthralgia; peripheral neuropathy, epileptiform seizures; leukopenia on prolonged or high dosage regimens; anorexia, glossitis, dryness of mouth.

Sulphamethoxazole + Trimethoprim
EDL-D490,491,492 PHC

AVAILABILITY
TABLETS (TMP + SMZ) 80 mg + 400 mg and 160 mg + 800 mg; Suspenson 40 mg tmp + 200 mg SMZ/5 ml.

DOSE
Adult- 1 to 2 tablets twice daily for 7-14 days (160 + 800 mg). Child- Suspension 5 ml twice daily (40 + 200 mg). infant 2.5 ml.
INDICATION
Urinary-tract infections; respiratory-tract infections including bronchitis, pneumonia, infections in cystic fibrosis; melioidosis; listeriosis; brucellosis; granuloma inguinale; otitis media; skin infections; Pneumocystis carinii pneumonia.

CONTRAINDICATION
Hypersensitivity to sulfonamides or trimethoprim; porphyria; marked liver parenchymal damage, blood dyscrasias, severe renal insufficiency

PRECAUTION
Renal impairment; hepatic impairment (avoid if severe; Appendix 7a); maintain adequate fluid intake (to avoid crystalluria); avoid in blood disorders (unless under specialist supervision); monitor blood counts and discontinue immediately if blood disorder develops; rash-discontinue immediately; predisposition to folate deficiency, elderly; asthma; G-6-PD deficiency; lactation (Appendix 7b); avoid in infants under 6 weeks; elderly.; pregnancy (Appendix 7c); interactions (Appendix 6c).

ADVERSE EFFECTS
Nausea, vomiting, diarrhoea, headache; hypersensitivity reactions including rashes, pruritus, photosensitivity reactions, exfoliative dermatitis and erythema nodosum; rarely, erythema multiforme (Stevens-Johnson syndrome) and toxic epidermal necrolysis; systemic lupus erythematosus, myocarditis, serum sickness; crystalluriresulting in haematuria, oliguria, anuria; blood disorders including granulocytopenia, agranulocytosis, aplastic anaemia, purpura-discontinue immediately; also reported, liver damage, pancreatitis, antibiotic-associated colitis, eosinophilia, cough and shortness of breath, pulmonary infiltrates, aseptic meningitis, depression, convulsions, ataxia, tinnitus, vertigo, dizziness, hallucinations and electrolyte disturbances; megaloblastic anaemia due to trimethoprim; elevation of transaminase and bilirubin; skin rashes.

Amikacin
EDL-D 19,20,21 Tertiary

AVAILABILITY
INJECTION 10 ml vial (100 mg/2 ml), 2 ml vial (250 mg/2 ml), (500 mg/2 ml).

DOSE
Intramuscular or intravenous injection or infusion Adult- 15 mg/kg body weight daily in two divided doses, increased to 22.5 mg/kg body weight daily in three divided doses in severe infections. (max 1.5g daily for 10 days, max. cumulative dose is 15g). Child- 15 mg/kg body weight daily in two divided doses. Neonates- loading dose is 10 mg/kg body weight followed by 15 mg/kg body weight in two divided doses.

INDICATION
Short-term treatment of serious infections due to susceptible strains of Gram-negative bacteria, including Pseudomonas species, Escherichia coli, species of indole-positive and indole-negative Proteus, Providencia species, Klebsiella, Enterobacter, Serratia species and Acinetobacter (Mima-Herellea) species.

CONTRAINDICATION
Myasthenia gravis; hypersensitivity.

PRECAUTION
Pregnancy (Appendix 7c), renal impairment (Appendix 7d); neonates, infants and elderly; cross allergenicity.

ADVERSE EFFECTS
Vestibular and auditory damage, nephrotoxicity; rarely, hypomagnesaemia on prolonged therapy, antibiotic-associated colitis, stomatitis; also reported, nausea, vomiting, rash, blood disorders; acute muscular paralysis; albuminuria; azotemia.
Vancomycin Hydrochloride  
EDL-D517,518 Tertiary

AVAILABILITY  
TABLETS 500 mg; INJECTION 250 mg, 500 mg and 1g/vial; Capsule 125 and 250 mg.

DOSE  
Adult- 1 to 1.5g every 12 h. Elderly over 65 years; 500 mg every 12 h or 1g once daily. Child-  
Over 1 month; 15 mg/kg body weight every 8 h (max. 2g daily).

INDICATION  
Methicillin-resistant staphylococcal pneumonia; staphylococcal meningitis; endocarditis prophylaxis (with gentamicin).

CONTRAINDICATION  
Allergy to corn/corn products, hypersensitivity.

PRECAUTION  
Avoid rapid infusion (risk of anaphylactoid reactions, see Adverse effects); rotate infusion sites; renal impairment (Appendix 7d); elderly; history of deafness-avoid; plasma-vancomycin concentration measured after 3 or 4 doses (earlier if renal impairment), blood counts, urinalysis and renal function tests-use only in hospital setting; monitor auditory function and plasma-vancomycin concentrations in elderly or in renal impairment; lactation (Appendix 7b); pregnancy (Appendix 7c); interactions (Appendix 6c); Pseudomembranous colitis.

ADVERSE EFFECTS  
Nephrotoxicity including renal failure and interstitial nephritis; ototoxicity (discontinue if tinnitus occurs); blood disorders; nausea, chills, fever, eosinophilia, anaphylaxis, rashes, including exfoliative dermatitis, erythema multiforme (Stevens-Johnson syndrome), toxic epidermal necrolysis and vasculitis; phlebitis; on rapid infusion, severe hypotension (with shock, cardiac arrest), wheezing, dyspnoea, urticaaria, pruritus, flushing of the upper body (‘red man’ syndrome), pain and muscle spasm of back and chest; hypotension, pruritus, haematopoietic flebitis.

Cefixime  
EDL-D99,100,101 Tertiary

AVAILABILITY  
TABLETS 50, 100, 200 and 400 mg; CAPSULES 100 and 200 mg; SYRUP/SUSPENSION 50 mg/5 ml,  
100 mg/5 ml.

DOSE  
Adult- 200-400 mg/day as a single dose or in two divided doses. Child- (more than 6 months) 8 mg/kg/day as a single dose or two divided doses. Uncomplicated gonorrhea: Adult- 400 mg as a single dose.

INDICATION  
Otitis media, respiratory tract infections, uncomplicated UTIs, effective against infections caused by Enterobacteriaceae, H. influenza species.

CONTRAINDICATION  
Hypersensitivity to cephalosporins.

PRECAUTION  
History of allergy to penicillins, renal failure (Appendix 7d) or patients undergoing continuous ambulatory peritoneal dialysis (CAPD) and hemodialysis (HD), gastrointestinal disease, pregnancy

ADVERSE EFFECTS  
lactation, interactionsDiarrhoea, pseudomembranous colitis, loose or frequent stools, abdominal pain, nausea, dyspepsia; hypersensitivity reactions.
Meropenem
EDL-D 684 Secondary hospitals

AVAILABILITY
INJECTIONS 0.125, 0.250, 0.5, 1 g/vial.

DOSE
Adult- 0.5-2 g or 10-40 mg/kg by slow i.v injection 8 hourly. Neonate (less than 7 days)- 20 mg/kg 12 hourly. 7-28 days- 20 mg/kg 8 hourly. 1-3 months- 10 mg/kg 8 hourly. > 3 months- 10-20 mg/kg 8 hourly. Meningitis: Adult- 2g 8 hourly. Child- (> 3 months)- 40 mg/kg 8 hourly.

INDICATION
Nosocomial infection like septicemia, febrile neutropenia, intraabdominal and pelvic infection etc caused by cephalosporins resistant bacteria, meningitis, cystic fibrosis.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Renal insufficiency, neurological disorders, prolonged use of meropenem may result in overgrowth of nonsusceptible organisms, pregnancy, lactation, history of hypersensitivity to other β-lactam antibiotics; interactions

ADVERSE EFFECTS
Inflammation at the injection site; nausea, vomiting, headache, rash; diarrhoea, thrombophlebitis, anaphylaxis, pseudomembranous colitis, disturbances in LFTs.

Netilmicin
EDL-D 701,702 Secondary hospitals

INDICATION
Urinary tract infections, serious systemic infections (enterobacteriaceae, and gentamicin resistant pathogens), Klebsiella, staphylococci

AVAILABILITY
Injection 50mg/mL 2mL ampoule; 100 mg/mL, 2mL Ampoule

DOSE
Injection 4-6 mg/kg IM/IV as a single dose or in divided doses every 8 or 12 hours; in severe infections upto 7.5 mg daily in divided doses every 8 hours.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Neurotoxicity and nephrotoxicity

ADVERSE EFFECT
Same as other aminoglycosides, but milder.

DRUG INTERACTION
Same as aminoglycosides.

Antileprosy medicines

Leprosy is a chronic mycobacterial infection due to *Mycobacterium leprae*, which is a slow-growing intracellular bacillus that infiltrates the skin, peripheral nerves, the nasal and other mucosa and the eyes; it affects people of all ages and both sexes. The incubation period between infection and appearance of leprosy is normally between 2 to 10 years, but may be up to 20 years. It is transmitted from person-to-person when bacilli are shed from the nose; most individuals have natural immunity and symptoms are suppressed. For treatment purposes patients may be classified as having paucibacillary (PB) or multibacillary (MB) leprosy. The 2
forms may be distinguished by skin smears, but facilities are not always available to process them and their reliability is often doubtful. In practice, most leprosy programmes classify and choose a regimen based on number of skin lesions; these are PB leprosy (1-5 skin lesions) and MB leprosy (more than 5 skin lesions).

Drugs used in the treatment of leprosy should always be used in combination; this is essential to prevent the emergence of resistance. Rifampicin is now combined with dapsone to treat PB leprosy and rifampicin and clofazimine are now combined with dapsone to treat MB leprosy. The WHO Programme for the Elimination of Leprosy currently provides, free of charge, oral multidrug therapy in colour-coded blister packs (MDT blister packs) to improve patients’ adherence to treatment. Any patient with a positive skin smear should be treated with the MDT regimen for MB leprosy. The regimen for PB leprosy should never be given to a patient with MB leprosy. If diagnosis classification in a particular patient is not possible the MDT regimen for MB leprosy must be used.

Lepra reactions are episodes of sudden increase in the activity of leprosy and are often accompanied by neuritis; reactions must always be treated promptly to prevent permanent nerve damage and disability. Leprosy multidrug therapy should continue during a lepra reaction without interruption. This reduces the frequency and severity of lepra reactions.

Type 1 lepra reactions, or reversal reactions, are delayed hypersensitivity reactions and may occur in either PB or MB leprosy. If there is no nerve damage, type 1 reactions may be treated with analgesics such as acetylsalicylic acid or paracetamol. If there is nerve involvement corticosteroids, such as oral prednisolone should be used in addition to analgesics.

The type 2 lepra reaction, also known as erythema nodosum leprosum (ENL), is an antibody response to dead leprosy bacteria and occurs only in MB leprosy. Therapy for type 2 reactions may include analgesics, such as acetylsalicylic acid or paracetamol and a corticosteroid, such as oral prednisolone. In patients not responding to a corticosteroid, clofazimine may be used. Severe type 2 lepra reactions should be treated under medical supervision in hospital.

If a patient does not respond to lepra reaction treatment within 6 weeks or seems to become worse, the patient must be sent immediately to the nearest specialist centre. Neuritis may occur during or independently of lepra reactions. It can be successfully treated with a 12-week course of oral prednisolone; if patients do not respond, specialist centre treatment is required.

**Treatment Regimens:**

The recommended regimen for paucibacillary leprosy in adults (50-70 kg) is rifampicin 600 mg once monthly and dapsone 100 mg daily. Children aged 10-14 years may be given rifampicin 450 mg once monthly and dapsone 50 mg daily. Appropriate dose adjustments are required for younger children. For example, dapsone 25 mg daily and rifampicin 300 mg once a month. Treatment is continued for 6 months for PB leprosy. The recommended regimen for MB leprosy in adults (50-70 kg) is rifampicin 600 mg and clofazimine 300 mg, both given once a month together with clofazimine 50 mg and dapsone 100 mg, both daily. Children aged 10-14 years may be given rifampicin 450 mg and clofazimine 150 mg, both once a month together with clofazimine 50 mg every other day and dapsone 50 mg daily. Appropriate dosage adjustments
are required for younger children. For example, dapsone 25 mg daily, clofazimine 50 mg twice a week and clofazimine 100 mg and rifampicin 300 mg once a month. Treatment is continued for 12 months for MB leprosy. For patients who cannot take rifampicin because of allergy, other diseases, or rifampicin-resistant leprosy and for patients who refuse to take clofazimine, there are alternative regimens which incorporate ofloxacin and minocycline

**Clofazimine**

**EDL-D128,129 PHC**

**AVAILABILITY**

tablets 25, 50, 100 mg; CAPSULES 50 and 100 mg.

**DOSE**

Oral Adult- 300 mg spread over a week. Sulfone resistant cases: 600 mg weekly preferably after meal. Lepra reaction: 200 mg daily for 3 weeks or as required. Child- 1 to 2 mg/kg body weight daily or

**INDICATION**

MB leprosy; type 2 lepra reactions.

**CONTRAINDICATION**

Pregnancy, lactation, renal and hepatic impairment

**PRECAUTION**

Pre-existing gastrointestinal symptoms (reduce dose, increase dose interval or discontinue if symptoms develop during treatment); liver and renal impairment; may discolour soft contact lenses; paediatrics, elderly, interactions (Appendix 6d)

**ADVERSE EFFECTS**

Reversible discolouration of skin, hair, cornea, conjunctiva, tears, sweat, sputum, faeces and urine; dose-related gastrointestinal symptoms including pain, nausea, vomiting and diarrhoea; severe mucosal and submucosal oedema, with prolonged treatment with high doses—may be severe enough to cause subacute small-bowel obstruction (see also Precautions); pruritus, ichthyosis, elevated blood sugar, diminished vision, dizziness, eosinophilic enteropathy.

**Dapsone**

**EDL-D147,148,149 PHC**

**AVAILABILITY**

TABLETS 25, 50 and 100 mg; gel 5% w/w.

**DOSE**

Adult- Leprosy: 50 to 100 mg daily depending upon body weight. Dermatitis herpetiformis: start with 50 mg daily and increase up to 400 mg till full response is obtained; dose reduced to minimum maintenance level as soon as possible. Child- 1 to 2 mg/kg body weight as minimum dose to start with, increased weekly so that at the end of 7th week patient is receiving max. dose.

**INDICATION**

PB and MB leprosy; acne vulgaris, dermatitis, pneumocystic pneumonia.

**CONTRAINDICATION**

Hypersensitivity to sulfones; severe anaemia; porphyria.

**PRECAUTION**

Anaemia (treat severe anaemia before therapy and monitor blood counts during treatment); susceptibility to haemolysis including G-6-PD deficiency (including lactation affected infants); lactation (Appendix 7b); porphyria; interactions (Appendix 6c); hyperbilirubinemia, methaemoglobinemia; renal impairment (Appendix 7d); pregnancy (Appendix 7c). On long-term treatment patients and their caretakers should be told how to recognize blood disorders and
advise to seek immediate medical attention if symptoms such as fever, sore throat, rash, mouth ulcers, purpura, bruising or bleeding develop.

ADVERSE EFFECTS
Haemolysis and methaemoglobinemia; allergic dermatitis (rarely, including toxic epidermal necrolysis and the Stevens-Johnson syndrome); rarely, hepatitis and agranulocytosis; 'dapsone syndrome' resembling mononucleosis-rare hypersensitivity reaction with symptoms including rash, fever, jaundice and eosinophilia; gastrointestinal irritation; tachycardia, headache, nervousness, insomnia, blurred vision, paraesthesia, reversible peripheral neuropathy and psychoses reported; increase in reticulocytes, vertigo; pancreatitis; renal papillary necrosis; anorexia

Rifampicin
EDL-D 456,457,458 PHC

AVAILABILITY
CAPSULES 150, 300, 450 and 600 mg; tablets 150, 300, 350, 450, 500, 600 and 750 mg; Syrup 100 mg/5 ml.

DOSE
Oral Adult- 450 to 600 mg single dose before breakfast. Child- 10 to 20 mg/kg body weight daily.

INDICATION
PB leprosy; MB leprosy; tuberculosis.

CONTRAINDICATION
Hypersensitivity; jaundice; patients with earlier drug induced liver disease.

PRECAUTION
(Appendix 7a); liver function tests and blood counts required in liver disorders, alcohol dependency, elderly and on prolonged therapy; renal impairment (if dose above 600 mg daily); lactation; porphyria; discolors soft contact lenses; advise patients on oral contraceptives to use additional means; interactions (Appendix 6b, 6c, 6d); pregnancy (Appendix 7c).

Note: Resumption of rifampicin treatment after a long interval may cause serious immunological reactions, resulting in renal impairment, haemolysis, or thrombocytopenia-discontinue permanently if serious adverse effects occur.

Patients or their caretakers should be told how to recognize signs of liver disorders and advised to discontinue treatment and seek immediate medical attention if symptoms such as persistent nausea, vomiting, malaise or jaundice develop.

ADVERSE EFFECTS
Severe gastrointestinal disturbances including anorexia, nausea, vomiting and diarrhoea (antibiotic-associated colitis reported); headache, drowsiness; rashes, fever, influenza-like syndrome and respiratory symptoms, collapse, shock, haemolytic anaemia, acute renal failure and thrombocytopenic purpura-m ore frequent with intermittent therapy; alterations of liver function-jaundice and potentially fatal hepatitis (dose-related, do not exceed max. daily dose of 600 mg); oedema, muscular weakness and myopathy, exfoliative dermatitis, toxic epidermal necrolysis, pemphigoid reactions, leukopenia, eosinophilia and menstrual disturbances; urine, tears, saliva and sputum coloured orange-red; cerebral haemorrhage, visual disturbances.

Ofloxacin
EDL-D 378,379 Secondary hospitals

AVAILABILITY
TABLETS 100, 200 and 400 mg; SYRUP 30 ml (50 mg/5 ml, 100 mg/5 ml); INJECTION 100 ml (2 mg/ml); EYE DROPS 0.3% w/v.

DOSE
Oral Community acquired pneumonia: Adult- 400 mg twice daily for 10 days. Pelvic inflammatory disease: Adult- 400 mg twice daily for 14 days. Complicated UTI: Adult- 200 mg
twice daily for 10 days.

**Parenteral Complicated UTI:** Adult - 200 mg daily by i.v infusion over at least 30 minutes, max. 400 mg twice infused over at least 1 h. Septicaemia, lower respiratory tract infection: Adult - 200 mg twice daily by i.v infusion over at least 30 minutes, max. 400 mg twice daily infused over at least 1 h. Bacterial corneal ulcer: Adult - 0.3%, 1-2 drops every 30 minutes. Ophthalmic Bacterial conjunctivitis: Adult - 0.3%, 1-2 drops every 2-4 h. Child - >1 year, 1-2 drops every 2-4 h.

**INDICATION**
Acute uncomplicated cystitis, community acquired pneumonia, acute exacerbation of chronic bronchitis.

**CONTRAINDICATION**
Hypersensitivity.

**PRECAUTION**
Patients with epilepsy, kidney disease, tendon problem, nervous system problem, liver disease (Appendix 7a), limit alcohol intake, pregnancy (Appendix 7c); lactation (Appendix 7b).

**ADVERSE EFFECTS**
Sinus tachycardia, hallucination, Steven’s Johnson syndrome, seizure; dizziness, headache, nausea, vomiting, diarrhoea; insomnia, pruritus, photosensitivity.

**Antituberculosis medicines**
Tuberculosis is a chronic infectious disease caused primarily by *Mycobacterium tuberculosis* or sometimes by *M. bovis*. Infection is usually due to inhalation of infected droplet nuclei with the lung generally being the first organ affected, but the primary infection is usually asymptomatic. Infection and inflammatory responses resolve with the development of acquired immunity. Surviving bacteria may become dormant or in susceptible patients, progress to active primary disease; dormant organisms may produce disease and this often occurs if immune status is altered.

Tuberculosis is the most prevalent infectious disease of adults and causes 26% of avoidable adult deaths in the developing world. More than 80% of tuberculosis cases are pulmonary (PTB). At least 30% of patients who are infected with HIV will also develop active tuberculosis. The increase in resistant strains and poor compliance of dosage regimen which may contribute to resistance and treatment failure has led to the development of regimens with directly supervised treatment. Directly observed treatment short-course (DOTS) therapy which lasts for 6 or 8 months, given under direct observation is one of the most important components of the WHO strategy against tuberculosis. Simplified drug regimens and intermittent therapy have been introduced to improve compliance. WHO does not generally recommend twice weekly regimens. If a patient receiving a twice weekly regimen misses a dose of tablets, the missed dose represents a bigger fraction of the total number of treatment doses than if the patient was receiving a three times weekly or daily dose regimen. Therefore, there is a greater risk of treatment failure with twice weekly regimens. Fixeddose combination tablets incorporating 2 or more drugs are also used to improve compliance and decrease medication errors; they should be used unless one of the components cannot be given because of resistance or intolerance.

Modern short-course therapy is usually in 2 phases. The initial phase (2 months) involves the concurrent use of at least 3 drugs to reduce the bacterial population rapidly and prevent drug-resistant bacteria emerging. The second continuation phase (4-6 months) involves fewer drugs...
and is used to eliminate any remaining bacteria and prevent recurrence. Direct observation of therapy is considered essential to ensure compliance in the initial phase and also useful in the continuation phase if patients are receiving rifampicin. Five antituberculosis drugs, isoniazid, rifampicin, pyrazinamide, streptomycin (which are bactericidal) and ethambutol (which is bacteriostatic) are used in various combinations as part of WHO-recommended treatment regimens; thiacetazone is used only if ethambutol cannot be used. In supervised regimens change of drug regimen should be considered only if the patient fails to respond after 5 months of DOTS.

Isoniazid, rifampicin and pyrazinamide are components of all antituberculosis drug regimens currently recommended by WHO. Unsupervised and alternative regimens as set out in the following tables may be administered as specified. Additional reserve antituberculosis drugs (amikacin, p-aminosalicylic acid, capreomycin, ciprofloxacin, cycloserine, ethionamide, kanamycin, levofloxacin and ofloxacin) for the treatment of multidrug-resistant tuberculosis should be used in specialized centres adhering to WHO standards for TB control.

Worldwide, an important predisposing cause of immunosuppression leading to tuberculosis is human immunodeficiency virus (HIV) infection; it increases susceptibility to primary infection and increases the reactivation rate of tuberculosis. Preventative antituberculosis therapy of such persons is recommended.

Chemoprophylaxis with isoniazid can prevent the development of clinically apparent disease in persons in close contact with infectious patients and also prevent the reactivation of previously dormant disease in other persons at high risk particularly those who are immunodeficient.

Where the disease remains highly prevalent routine immunization of infants within the first year of age with BCG vaccine is cost-effective. However, there is no evidence that BCG will protect children older than 15 years of age. Infants born to HIV-positive mothers should be vaccinated during the first year of life, provided they have no clinical signs suggestive of HIV.

The tuberculin test has limited diagnostic value. A positive tuberculin test indicates previous exposure to mycobacterial antigens through infection with one of the tubercle bacilli, or BCG vaccination. The tuberculin test does not distinguish between tuberculosis and other mycobacterial infection, between active and quiescent disease, or between acquired infection and seroconversion induced by BCG vaccination.

**Recommended 6-Month Treatment Regimens for Tuberculosis**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Initial phase (2 months)</th>
<th>Continuation phase (4 months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid</td>
<td>5 mg/kg daily</td>
<td>5 mg/kg daily</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>10 mg/kg daily</td>
<td>10 mg/kg daily</td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td>25 mg/kg daily</td>
<td></td>
</tr>
<tr>
<td>together with</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Streptomycin</td>
<td>15 mg/kg daily</td>
<td></td>
</tr>
<tr>
<td>or Ethambutol</td>
<td>15 mg/kg daily</td>
<td></td>
</tr>
</tbody>
</table>

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93
Isoniazid 10 mg/kg 3 times weekly  
Rifampicin 10 mg/kg 3 times weekly  
Pyrazinamide 35 mg/kg 3 times weekly  
**together with**  
Streptomycin 15 mg/kg 3 times weekly  
or  
Ethambutol 30 mg/kg 3 times weekly

1Unless otherwise indicated, doses are suitable for both adults and children  
2Not suitable for children

**Recommended 8-month treatment regimen for tuberculosis**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Initial phase (2 months)</th>
<th>Continuation phase (6 months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid</td>
<td>5 mg/kg daily</td>
<td>5 mg/kg daily</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>10 mg/kg daily</td>
<td></td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td>25 mg/kg daily</td>
<td></td>
</tr>
<tr>
<td><strong>together with</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| Ethambutol | 15 mg/kg daily | 15 mg/kg daily  
| **or** |  |  |
| Streptomycin | 15 mg/kg daily |  |

1Unless otherwise indicated, doses are suitable for both adults and children  
2Streptomycin always replaces ethambutol in meningeal TB  
3Not suitable for children under 5 years  
4Thiacetazone (2.5 mg/kg daily) may be used (only if ethambutol cannot be given) in combination with isoniazid in the continuation phase; risk of severe toxicity, particularly in HIV-infected individuals

**Category I:** New pulmonary disease (smear-positive or smear-negative with extensive involvement of parenchyma), concomitant severe HIV disease and new severe extra-pulmonary Disease.

Initial phase (antibacterials administered daily or 3 times weekly): isoniazid + rifampicin + pyrazinamide + ethambutol (or streptomycin) for 2 months Continuation phase (antibacterials administered daily or 3 times weekly): isoniazid + rifampicin for 4 months (or isoniazid + ethambutol for 6 months but less effective than isoniazid + rifampicin)

**Category II:** Previously treated smear-positive pulmonary disease which has relapsed, or failed to respond, or if treatment was interrupted.

Initial phase (antibacterials administered daily or 3 times weekly): isoniazid + rifampicin + pyrazinamide + ethambutol + streptomycin for 2 months then: isoniazid + rifampicin + pyrazinamide + ethambutol for 1 month Continuation phase (antibacterials administered daily or 3 times weekly): isoniazid + rifampicin + ethambutol for 5 months.

**Category III:** New smear-negative pulmonary disease (other than in Category I) and less severe extra-pulmonary disease Initial phase (antibacterials administered daily or 3 times weekly): isoniazid + rifampicin + pyrazinamide + ethambutol for 2 months Continuation phase (antibacterials administered daily or 3 times weekly): isoniazid + rifampicin for 4 months (or isoniazid + ethambutol for 6 months but less effective than isoniazid + rifampicin).
Category IV: Chronic and multi-drug-resistant tuberculosis (MDR-TB) (smear-positive despite supervised re-treatment) specially designed standardized or individualized regimens recommended.

Treatment regimens by category of tuberculosis diagnosis

1. Drug intake should be directly observed in patients who are smear positive during the initial phase and always when rifampicin is given.

2. Drug sensitivity testing recommended before prescribing Category II treatment in failure cases; patients with MDR-TB should be prescribed Category IV regimen.

3. Omit ethambutol in initial phase if disease is not complicated by cavitary disease or concomitant HIV disease and in patients infected with fully susceptible bacilli or young children with primary tuberculosis.

4. Early culture and sensitivity testing recommended for contacts of patients with MDR-TB.

Ethambutol Hydrochloride

**EDL-D205, 630 PHC**

**AVAILABILITY**

TABLETS 200, 400, 600, 800 mg and 1g.

**DOSE**

Oral Adult- 15 mg/kg body weight as a single dose, retreatment with 25 mg/kg body weight as a single dose for two months, thereafter reduce to 15 mg/kg body weight. Given as combination therapy with other anti-tubercular drugs. Child- Same as for Adult. Do not use under 3 years.

**INDICATION**

Tuberculosis, in combination with other drugs.

**CONTRAINDICATION**

Optic neuritis; children under 5 years-unable to report symptomatic visual disturbances; severe renal impairment; hypersensitivity.

**PRECAUTION**

Visual disturbances-ocular examination recommended before and during treatment (see note below); reduce dose in renal impairment (Appendix 7d) and monitor plasma concentration; elderly; pregnancy (Appendix 7c) (not known to be harmful); lactation.

Note: Patients should report visual disturbances immediately and discontinue treatment; children who are incapable of reporting symptomatic visual changes accurately should be given alternative therapy, as should, if possible, any patient who cannot understand warnings about visual adverse effects.

**ADVERSE EFFECTS**

Optic neuritis-reduced visual acuity and red/ green colour blindness (early changes usually reversible, prompt withdrawal may prevent blindness); peripheral neuritis-especially in legs; gout; rarely, rash, pruritus, urticaria, thrombocytopenia; pulmonary infiltrates gastrointestinal upset.

Isoniazid

**EDL-D 287,288 PHC**

**AVAILABILITY**

TABLETS 100 and 300 mg.

**DOSE**

Adult- 3 to 5 mg/kg body weight up to 300 mg as single dose daily. Child- 10 to 15 mg/kg body weight as a single dose, not to exceed 300 mg/day.

**INDICATION**

Tuberculosis, in combination with other drugs; tuberculosis prophylaxis also.
CONTRAINDICATION
Pregnancy

PRECAUTION
Hepatic impairment (monitor hepatic function; Appendix 7a); malnutrition, chronic alcohol dependence, chronic renal failure (Appendix 7d); diabetes mellitus and HIV infection—prophylactic pyridoxine 10 mg daily required because risk of peripheral neuritis; epilepsy; slow acetylator status (increased risk of adverse effects); history of psychosis; pregnancy (Appendix 7c) (not known to be harmful); lactation (Appendix 7b); porphyria; interactions (Appendix 6a, 6c, 6d). Patients or their caretakers should be told how to recognize signs of liver disorder and advised to discontinue treatment and seek immediate medical attention if symptoms such as nausea, vomiting, malaise or jaundice develop.

ADVERSE EFFECTS
Combined preparation not suitable for use in children; see Rifampicin, Isoniazid and Pyrazinamide; pregnancy

**Pyrazinamide**

EDL-445,446 PHC

AVAILABILITY
TABLETS 300, 500 and 750 mg; 1 and 1.5g; suspension 100 ml (5%).

DOSE
Oral Adult and Child- 20 to 35 mg/kg body weight as a single dose (max. 3g daily).

INDICATION
Tuberculosis, in combination with other drugs.

CONTRAINDICATION
Severe hepatic impairment; porphyria.

PRECAUTION
Hepatic impairment (monitor hepatic function; (Appendix 7a); renal impairment (Appendix 7d); diabetes mellitus (monitor blood glucose—may change suddenly); gout; pregnancy (Appendix 7c) and lactation; hypouricemia. Patients or their caretakers should be told how to recognize signs of liver disorder and advised to discontinue treatment and seek immediate medical attention if symptoms such as persistent nausea, vomiting, malaise or jaundice develop.

ADVERSE EFFECTS
Hepatotoxicity including fever, anorexia, hepatomegaly, splenomegaly, jaundice, liver failure; nausea, vomiting; arthralgia; gout; sideroblastic anaemia; rash, photosensitivity; porphyria, dysuria, thrombocytopenia, hyperplasia, myalgia.

**Rifampicin**

EDL-D 456,457,458 PHC

AVAILABILITY
CAPSULES 150, 300, 450 and 600 mg; tablets 150, 300, 350, 450, 500, 600 and 750 mg; Syrup 100 mg/5 ml.

DOSE
Oral Adult- 450 to 600 mg single dose before breakfast. Child- 10 to 20 mg/kg body weight daily.

INDICATION
PB leprosy; MB leprosy; tuberculosis.

CONTRAINDICATION
Hypersensitivity; jaundice; patients with earlier drug induced liver disease.

PRECAUTION
(Appendix 7a); liver function tests and blood counts required in liver disorders, alcohol
dependency, elderly and on prolonged therapy; renal impairment (if dose above 600 mg daily); lactation; porphyria; discours soft contact lenses; advise patients on oral contraceptives to use additional means; interactions (Appendix 6b, 6c, 6d); pregnancy (Appendix 7c).

Note: Resumption of rifampicin treatment after a long interval may cause serious immunological reactions, resulting in renal impairment, haemolysis, or thrombocytopenia-discontinue permanently if serious adverse effects occur. Patients or their caretakers should be told how to recognize signs of liver disorders and advised to discontinue treatment and seek immediate medical attention if symptoms such as persistent nausea, vomiting, malaise or jaundice develop.

ADVERSE EFFECTS
Severe gastrointestinal disturbances including anorexia, nausea, vomiting and diarrhoea (antibiotic-associated colitis reported); headache, drowsiness; rashes, fever, influenza-like syndrome and respiratory symptoms, collapse, shock, haemolytic anaemia, acute renal failure and thrombocytopenic purpura-m ore frequent with intermittent therapy; alterations of liver function-jaundice and potentially fatal hepatitis (dose-related, do not exceed max. daily dose of 600 mg); oedema, muscular weakness and myopathy, exfoliative dermatitis, toxic epidermal necrolysis, pemphigoid reactions, leukopenia, eosinophilia and menstrual disturbances; urine, tears, saliva and sputum coloured orange-red; cerebral haemorrhage, visual disturbances.

Streptomycin Sulphate
EDL-D485 PHC
AVAILABILITY
INJECTION vial 750 mg and 1g.
DOSE
Deep intramuscular injection. Adult- 0.75g to 1g daily. Elderly- 0.5g daily. Child- 20 to 40 mg/kg body weight daily.
INDICATION
Tuberculosis, in combination with other drugs.
CONTRAINDICATION
Hearing disorders; myasthenia gravis; pregnancy
PRECAUTION
Children-painful injection, avoid use if possible; renal impairment (Appendix 7d), infants and elderly (dosage adjustment and monitor renal, auditory and vestibular function and plasma streptomycin concentrations); interactions(Appendix 6c)
ADVERSE EFFECTS
Vestibular and auditory damage, nephrotoxicity; hypersensitivity reactions-withdraw treatment; parasthenia of mouth; rarely, hypomagnesaemia on prolonged therapy; antibiotic-associated colitis; also, nausea, vomiting, rash; rarely, haemolytic anaemia, aplastic anaemia, agranulocytosis, thrombocytopenia; pain and abscess at injection site

Cycloserine
EDL-D 596 PHC
AVAILABILITY
CAPSULE/TABLET 250 mg.
DOSE
Oral Adult- Initially 250 mg every 12 h for 2 weeks, increase according to blood concentration and response to 500 mg every 2 h. Child- Initially 10 mg/kg body weight daily adjusted to blood concentration and response.
INDICATION
Tuberculosis resistant to first-line drugs.
CONTRAINDICATION
Severe renal impairment; epilepsy; depression, severe anxiety, psychotic states, alcohol dependence; porphyria; hypersensitivity

PRECAUTION
Reduce dose in renal impairment (avoid if severe); monitor haematological, renal and hepatic function; lactation; discontinue or reduce dose if allergic skin reactions or CNS toxicity occur, pregnancy (Appendix 7c)

ADVERSE EFFECTS
Mainly neurological, including headache, dizziness, vertigo, drowsiness, tremor, convulsions, confusion, psychosis, depression (discontinue or reduce dose if symptoms of CNS toxicity); rashes, allergic dermatitis (discontinue or reduce dose); megaloblastic anaemia; changes in liver function tests; heart failure at high doses reported.

Kanamycin
EDL-D 673,674 PHC

AVAILABILITY
INJECTION Vial 500, 750 mg and 1g.

DOSE
Intramuscular and intravenous injection Adult- 1g daily as a single dose. Child- 6 to 15 mg/kg body weight daily in divided doses, 8 to 12 h (slow injection), usual duration of therapy 7 to 10 days

INDICATION
Tuberculosis; hepatic coma; penicillin resistant gonorrhoea, chronic bacterial infections

CONTRAINDICATION
Lactation; pregnancy (Appendix 7c); hypersensitivity; renal impairment.

PRECAUTION
Myasthenia gravis; renal impairment; elderly patients with neuromuscular disorder

ADVERSE EFFECTS
Nephrotoxicity; ototoxicity; skin rash; urticaria; neuromuscular blockade; malabsorption syndrome.

Antifungal medicines
Fungal infections can be superficial or systemic. Superficial infections affect only the skin, hair, nails or mucous membranes whereas systemic fungal infections affect the body as a whole. Systemic fungal infections are sometimes caused by inhalation, ingestion or inoculation of primary pathogens and sometimes by opportunistic invasion of commensals in patients with lowered host resistance. They are increasing in prevalence not only because of the pandemic of HIV infection, but also because of the rise in illicit intravenous drug use in many countries and greater use of broad spectrum antibiotics and invasive medical procedures. In immunodeficient patients systemic fungal infections are often disseminated.

Amphotericin B is a lipophilic polyene antibiotic; it is fungistatic against a broad spectrum of pathogenic fungi, including Candida spp., Aspergillus spp., Cryptococcus neoformans, Histoplasma capsulatum, Blastomyces dermatitidis, Coccidioides immitis, Paracoccidioides brasiliensis, Mucor, Absidia and Phicopes spp.; it is active against algal Prototheca spp. And against the Leishmania protozoa. It is used for the empirical treatment of serious fungal infections and is used in conjunction with flucytosine to treat cryptococcal meningitis and systemic candidosis.
Amphotericin B has to be administered parenterally as there is little or no absorption from the gastrointestinal tract; amphotericin B is liable to cause nephrotoxicity. Duration of therapy varies with the initial severity of the infection and the clinical response of the patient. In some infections a satisfactory response is only obtained after several months of continuous treatment. Intrathecal infusion has been used successfully in patients with meningeal coccidioidomycosis.

Fluconazole, an orally active synthetic imidazole derivative, possesses fungistatic activity against dermatophytes, yeasts and other pathogenic fungi. It is widely used in the treatment of serious gastrointestinal and systemic mycoses as well as in the management of superficial infections. Fluconazole is also used to prevent fungal infections in immunocompromised patients.

Flucytosine, is a synthetic fluorinated pyrimidine with a narrow spectrum of antifungal activity, particularly against Cryptococcus and Candida spp. In susceptible fungi, it is converted to 5-fluorouracil by cytosine deaminase. Flucytosine is myelosuppressive and plasma concentrations above 75 μg/ml are associated with myelotoxicity.

Griseofulvin is a fungistatic antibiotic derived from Penicillium griseofulvum with selective activity against the dermatophytes causing ringworm, Microsporum canis, Trichophyton rubrum and T. verrucosum. It has no activity against pityriasis versicolor or candida infections. Griseofulvin is deposited selectively in keratin precursor cells of skin, hair and nails where it disrupts the mitotic apparatus of fungal cells thus preventing fungal invasion of newly-formed cells. It is unsuitable for prophylactic use. Close attention should be given to hygiene and to possible reservoirs of reinfection in clothing, footwear and bedding.

Nystatin, a polyene antifungal antibiotic derived from Streptomyces noursei, is effective against infections caused by a wide range of yeasts and yeast-like fungi. It is poorly absorbed from the gastrointestinal tract and it is not absorbed from the skin or mucous membranes when applied topically. It is used for the prophylaxis and treatment of candidosis.

Potassium iodide aqueous oral solution is a clear liquid with a characteristic, strong salty taste. It is effective against sporotrichosis and subcutaneous phycomycosis, which are fungal infections caused by Sporothrix schenckii and Basidiobolus haptosporus respectively. In subcutaneous sporotrichosis, amphotericin B is often effective in patients unable to tolerate iodides. Itraconazole, by mouth has been tried as an alternative to potassium iodide in both cutaneous and extracutaneous sporotrichosis. In phycomycosis, fluconazole may be effective.

Amphotericin B lipoholised
EDL-D 38 Tertiary

AVAILABILITY
VIALS 10, 25, 50 and 100 mg plain, 50 mg/ vial (liposomal).

DOSE
Intravenous infusion (plain) Adult- Systemic fungal infection: 250 μg/kg body weight daily, increase gradually 1 mg/ kg body weight if tolerated (max 1.5 mg/kg body weight daily) or alternate days. Child- Same as for Adult based on body weight. Intravenous (liposomal) For fever in neutropenic patients: 3 mg/kg/ day, max. dose 5 mg/kg/day i.v. For cryptococcal meningitis:
3-4 mg/kg, max. 6 mg/kg, i.v. once daily. Visceral leismaniasis: Immunocompetent patients: 3 mg/kg. Immunocompromized patients: 4 mg/kg.

**INDICATION**
Life-threatening fungal infections including histoplasmosis, coccidioidomycosis, paracoccidioidomycosis, blastomycosis, aspergillosis, cryptoacoccus, mucormycosis, sporotrichosis and candidiasis; visceral and mucocutaneous leishmaniasis unresponsive to pentavalent antimony compounds; severe meningitis, perioral candidiasis.

**CONTRAINDICATION**
Toxic effects must be weighed against benefits. Regular kidney, liver function tests and blood counts must be conducted; lactation; antineoplastic therapy.

**PRECAUTION**
Close medical supervision throughout treatment and initial test dose required (see note, below); renal impairment (Appendix 7d); pregnancy (Appendix 7c); hepatic and renal function tests; blood counts and plasma electrolyte monitoring; corticosteroids (avoid, except to control reactions); lactation; avoid rapid infusion (risk of arrhythmias); interactions (Appendix 6c); geriatric use. Anaphylaxis occurs rarely, with intravenous amphotericin B and a test dose is advisable before the first infusion. The patient should be observed for about 30 min after the test dose.

**ADVERSE EFFECTS**
Fever, headache, anorexia, weight loss, nausea and vomiting, malaise, diarrhoea, muscle and joint pain, dyspepsia and epigastric pain; renal function disturbances including hypokalaemia, hypomagnesaemia and renal toxicity; blood disorders; cardiovascular toxicity (including arrhythmias); neurological disorders (including peripheral neuropathy); abnormal liver function (discontinue treatment); rash; anaphylactoid reactions (see above); pain and thrombophlebitis at injection site; respiratory failure.

**Fluconazole**
**EDL-D 225,226,227,228 PHC**

**AVAILABILITY**
TABLETS/CAPSULES 50, 100, 150 and 200 mg; EYE DROPS 5 ml (0.3% w/v).

**DOSE**
Adult- Mucosal: 50 to 100 mg daily for 14 to 30 days. Vaginal: 150 mg as a single dose. Oral: systemic loading dose of 400 mg on first day and thereafter 200 to 400 mg once daily for at least 28 days. Prophylaxis of fungal infection: 50 to 100 mg once daily

**INDICATION**
Systemic mycosis including histoplasmosis, non-meningal coccidioidomycosis, paracoccidioidomycosis and blastomycosis treatment and, in AIDS and other immunosuppressed patients, prophylaxis of cryptococcal meningitis; oesophageal and oropharyngeal candidiasis, vaginal candidiasis and systemic candidiasis.

**CONTRAINDICATION**
Sensitivity to primaquine; infants below 1 year of age; alcohol; coadministration of cisapride, terfenadine.

**PRECAUTION**
Renal impairment (Appendix 7d); lactation (Appendix 7b); monitor liver function, discontinue if signs or symptoms of hepatic disease (risk of hepatic necrosis; Appendix 7a); interactions (Appendix 6b, 6c); pregnancy (Appendix 7c); immunocompromised patients.

**ADVERSE EFFECTS**
Nausea, vomiting, abdominal pain; flatulence, diarrhoea; headache, taste disturbance, hepatic disorders, dizziness, seizures, alopecia, pruritus; rash (withdraw treatment); angioedema,
anaphylaxis, bullous lesions, toxic epidermal necrolysis and erythema multiforme (Stevens-Johnson syndrome) reported (skin reactions more common in AIDS); hyperlipidaemia, leukopenia, thrombocytopenia, hypokalaemia.

**Nystatin**

**EDL-376,377 Tertiary**

**AVAILABILITY**

TABLETS 5,00,000 units; ointment 3g (100000 IU).

**DOSE**

Oral Adult- Intestinal candidiasis: 5,00,000 units every six h, doubled in severe infections. Child-1 month to 12 years: 1,00,000 units 4 times daily, immunocompromised children may require higher doses up to 5,00,000 units. Topical application Dissolve one tablet in glycerine and apply locally 3 to 4 times. Intravaginal Insert one tablet deep into vagina before bed time once at night.

**INDICATION**

Oral, oesophageal, intestinal, vaginal and cutaneous candidiasis

**CONTRAINDICATION**

Hypersensitivity.

**PRECAUTION**

Lactation; discontinue if sensitivity develops, teratogenic effect, should not be used for the treatment of systemic, oral, intravaginal or ophthalmic infections; pregnancy(Appendix 7c)

**ADVERSE EFFECTS**

Nausea, vomiting, diarrhoea at high doses; oral irritation and sensitization; rash and rarely, erythema multiforme (Steven’s- Johnson syndrome); eczema, burning.

**Povidone Iodine**

**EDL-D 425 Secondary hospitals**

**AVAILABILITY**

SOLUTIONS 100 and 500 ml (5% w/v), 500 ml (7.5% w/v and 10% w/v); OINTMENT 15g (5% w/w).

**DOSE**

Adult and Child- Pre- and post-operative skin disinfection: apply undiluted. Antiseptic (minor wounds and burns): apply twice daily.

**INDICATION**

Antiseptic; skin disinfection; Mouth wash.

**CONTRAINDICATION**

Avoid regular or prolonged use in patients with thyroid disorders or those taking lithium; avoid regular use in neonates; avoid in very low birthweight infants; burn covering large surface area; hypersensitivity to iodine.

**PRECAUTION**

Pregnancy (Appendix 7c) ; lactation(Appendix 7b) ; broken skin ; renal impairment; avoid contact with eyes; neonates. The application of povidone iodine to large wounds or severe burns may produce systemic adverse effects such as metabolic acidosis; hypernatraemia; and impairment of renal function.

**ADVERSE EFFECTS**

Irritation of skin and mucous membranes; may interfere with thyroid function tests; systemic effects

**Clotrimazole**

**EDL-D 130,131,132 Secondary hospitals**

**AVAILABILITY**

PESSARIES/VAGINAL TABLETS 100 and 200 mg; CREAM 1% w/w; powder 75g; Lotion 50 ml.
DOSE
Adult- Pessaries/vaginal tablets: 100 mg pessary/vaginal tablet to be inserted into vagina at night before going to bed as deep as possible for consecutive 6 to 7 days or 200 mg for 3 consecutive night before going to bed or 500 mg single dose. Child- Pessaries/vaginal tablets: not recommended. Cream: Rub on affected area 2 to 3 times by applying in thin layer and rubbing, continue for 14 days after healing.

INDICATION
Vulvo-vaginal candidiasis, trichomoniasis, vaginitis, non-specific vaginitis, mixed vaginal infection, Gram-positive and Gram-negative bacterial infection, infective leucorrhoeas; prevention of athletes foot and ringworm disease of skin folds.

CONTRAINDICATION
Ophthalmic use; hypersensitivity.

PRECAUTION
Avoid contact with eyes, pregnancy (Appendix 7c) and lactation(Appendix 7b)

ADVERSE EFFECTS
Local irritation, burning sensation and itching, abnormal liver function, unpleasant mouth sensation.

BENZOIC ACID COMPOUND (BENZOIC ACID + SALICYLIC ACID)

EDL-D 67 TERTIARY

AVAILABILITY
CREAM 25 and 50g (Aluminium tubes, jars).

DOSE
Fungal skin infections: apply twice daily until the infected skin is shed (usually at least 4 weeks)

INDICATION
Mild dermatophyte infections, particularly caused by Tinea pedis and Tinea corporis.

PRECAUTION
Avoid contact with eye, nose and mouth, pregnancy (Appendix 7c)

ADVERSE EFFECTS
Occasionally localized; mild inflammatory reaction; swelling of face, lips and tongue; difficulty in breathing.

CLINDAMYCIN

EDL-D 578 PRIMARY

INDICATIONS
Respiratory tract infections, penicillin resistant staphylococcal infections and many anaerobes such as bacteroides, skin, soft tissue and dental infections.

AVAILABILITY
TABLETS/CAPSULES 150 & 300 mg; SYRUP 4 ml (150 mg/ml); INJECTION 2 ml (150 mg/ml); CREAM/GEL/OINTMENT 10g (1%w/w); LOTION 25 ml (1%w/v).

DOSE
Oral
Serious anaerobic infections
Adult: 150-300 mg 6 every hr; for more severe infection: 300 to 450 mg every 6 hr.
Child: 2.4 mg/kg every 6 hr; for more severe infection: 3-6 mg/kg every 6 hr; 10 kg: 37.5 mg every 8hr.
Prophylaxis of endocarditis 600 mg 1 hr before dental procedure.
Intravenous/Intramuscular
Serious anaerobic infections
Adult: 0.6-2.7 g/day in 3-4 divided doses, up to 4.8 g/day for severe infections.
Child: 20-40 mg/kg daily in 3-4 divided dose.
Neonate: 15-20 mg/kg daily in 3-4 divided dose
Toxic shock syndrome
Adult: 900 mg every 8 hr along with penicillin G or ceftriaxone.
Pelvic inflammatory disease
Adult: 900 mg every 8 hr along with gentamicin.
Vaginal
Bacterial vaginosis
As pessary or 2% cream: 100 mg once nightly for 3-7 days.
Topical
Acne
As 1% preparation: Apply twice daily.

CONTRAINDICATIONS
Hypersensitivity, meningitis as it has less penetration into CNS, pseudomembranous colitis.

PRECAUTIONS
Hepatic and renal impairment, pregnancy and lactation, GI disease, elderly, atopic patients, regular monitoring of blood counts, in conjuction with antibiotic therapy, pregnancy (Appendix 7c), interactions (Appendix 6c).

ADVERSE EFFECTS
Urticaria, rashes, contact dermatitis, exfoliative and vesiculous dermatitis, local irritation abdominal pain, oesophagitis, nausea, vomiting, diarrhoea, jaundice and liver abnormalities, eosinophilia, erythema multiforme, thrombophlebitis, gasping syndrome (premature infants and neonates) due to preservative benzoyl alcohol in parenteral formulation, pseudomembranous colitis, azotemia, oliguria, proteinuria.

STORAGE
Store protected from moisture

Antiviral medicines
Antiherpes medicines
Herpes and Cytomegalovirus Infections:

Herpes Simplex Virus (HSV):

Acyclovir is active against herpes viruses but does not eradicate them. It is only effective if started at onset of infection; it is also used for prevention of recurrence in the immunocompromised patients. Genital lesions, oesophagitis and proctitis may be treated with oral Acyclovir. HSV encephalitis or pneumonitis should be treated with intravenous Acyclovir. Valacyclovir, a prodrug of Acyclovir, can be given by mouth as an alternative treatment for herpes simplex infections of the skin and mucous membranes (including initial and recurrent genital herpes).

Herpes Zoster Virus:

While most HIV positive patients with zoster experience only one self-limiting course, some will experience repeated episodes. Treatment should be reserved for debilitating disease and when there is high risk of serious complications, such as in advanced HIV disease. Acyclovir is the treatment of choice and it can be administered in high oral dose or in the case of lack of response to oral therapy or CNS involvement, it should be given intravenously.

Cytomegalovirus (CMV):

Parenteral antiviral ganciclovir arrests retinochoroiditis and enteritis caused by CMV in HIV infected patients. Maintenance therapy with oral ganciclovir should be given to prevent relapse of retinitis. Alternative therapy with intravenous foscarnet can be used if necessary.
Acyclovir  
EDL-D 6,7,9,10,559 Secondary hospitals

AVAILABILITY
TABLETS Plain/DT 200, 400 and 800 mg; SUSPENSION 400 mg/5 ml; INFUSION 100 ml (after reconstitution) (250 mg); Ointment 5g (3%w/w); Drops 5 ml (3% w/w); cream 5g (5% w/w).

DOSE
Adult- Non-genital herpes simplex treatment, 200 mg five times daily usually for 5 days, longer if new lesions appear during treatment or if healing is incomplete. 400 mg for immunocompromised patients or if absorption is impaired. Genital herpes simplex treatment; 200 mg 5 times daily for 5 days or 400 mg three times daily for three days. Longer if new lesions appear or healing is incomplete. Immunocompromised or HIV positive patients; 400 mg is given five times daily for 7 to 10 days during first episode or 400 mg three times a day for 5 to 10 days during recurrent injection. Herpes simplex prevention of recurrence; 200 mg 4 times daily or 400 mg twice daily reduced to 200 mg two or three times daily interrupted every 6 to 12 months. Varicella and herpes zoster; 800 mg five times daily for 7 days. Chicken pox; 800 mg five times daily for 7 to 10 days. Intravenous infusion Severe initial genital herpes, Varicella zoster, Herpes simplex infection; 5 mg/kg body weight every 8 h for five days. Child- Under 2 years; half dose. Above 2 years; adult dose.Varicella and herpes zoster; 20 mg/kg body weight (max. 800 mg) four times daily for 5 days, under 2 years 200 mg four times daily, for 2 to 5 years; 400 mg four times daily. Over 6 years; 800 mg four times daily. Chicken pox; 20 mg/kg body weight (max 800 mg) four times daily for 5 days.

INDICATION
Treatment of primary genital herpes; disseminated Varicella-zoster in immunocompromised patients; Herpes simplex encephalitis; chicken pox.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Maintain adequate hydration; renal impairment (Appendix 7d); lactation (Appendix 7b); pregnancy (Appendix 7c); paediatrics.

ADVERSE EFFECTS
Nausea, vomiting, abdominal pain, diarrhoea, headache, fatigue, rash, urticaria, pruritus, photosensitivity; very rarely, hepatitis, jaundice; dyspnoea; neurological reactions (including dizziness, confusion, hallucinations, convulsions and drowsiness); acute renal failure; anaemia, thrombocytopenia and leucopenia; on intravenous infusion; severe local inflammation (sometimes leading to ulceration), and very rarely, agitation, tremors; psychosis and fever; increase in blood urea and creatinine, encephalopathy; seizures; anorexia, tremors.

Antiretroviral medicines
Antiretroviral drugs do not cure HIV (human immunodeficiency virus) infection; they only temporarily suppress viral replication and improve symptoms. Patients receiving these drugs require careful monitoring by appropriately trained health professionals in an adequately resourced setting. Rigorous promotion of measures to prevent new infections remains essential and its need is not diminished by the availability of antiretroviral drugs. Effective therapy requires the simultaneous use of 3 or 4 drugs; alternative regimens are necessary to meet specific requirements at start-up, to substitute for first-line regimens in cases of intolerance, or to replace failing regimens. The use of a 3- or 4-drug combination as specified in the WHO treatment guidelines is recommended. The use of fixed-dose preparations for these combinations is also recommended if the pharmaceutical quality is assured and
interchangeability with the single products is demonstrated as specified by the relevant drug regulatory authority.

Selection of 2 or 3 protease inhibitors from the Model List will need to be determined by each country after consideration of local treatment guidelines and experience, as well as comparative costs of available products. Low-dose ritonavir is used in combination with indinavir, lopinavir or saquinavir as a ‘booster’; ritonavir is not recommended as a drug in its own right.

**Principles of Treatment:**

Treatment is aimed at reducing the plasma viral load as much as possible and for as long as possible; it should be started before the immune system is irreversibly damaged. The need for early drug treatment should, however, be balanced against the development of toxicity. Commitment to treatment and strict adherence over many years are required; the regimen chosen should take into account convenience and the patient’s tolerance of it. The development of resistance is reduced by using a combination of 3 or 4 drugs; such combinations should have additive or synergistic activity while ensuring that their toxicity is not additive. Testing for resistance to antiviral drugs, particularly in therapeutic failure, should be considered. Women of childbearing age receiving antiretroviral therapy must have available effective contraceptive methods to prevent unintended pregnancy. Women who are taking nonnucleoside reverse transcriptase inhibitors or protease inhibitors which can lower blood concentration of hormonal oral contraceptives, should be advised to use additional or alternative contraceptives.

**Drugs used to treat HIV Infection:**

Zidovudine, a nucleoside reverse transcriptase inhibitor (or ‘nucleoside analogue’), was the first anti-HIV drug introduced. Other nucleoside reverse transcriptase inhibitors include abacavir, didanosine, lamivudine, stavudine and zalcitabine.

The protease inhibitors include amprenavir, indinavir, lopinavir, nelfinavir, ritonavir and saquinavir. Ritonavir in low doses is used in combination with indinavir, lopinavir or saquinavir as a booster. The small amount of ritonavir in such combinations has no intrinsic antiviral activity but it increases the antiviral activity of the other protease inhibitors by reducing their metabolism. Indinavir, nelfinavir, ritonavir and possibly saquinavir inhibit the cytochrome P450 enzyme system and therefore have a potential for significant drug interactions. Protease inhibitors are associated with lipodystrophy and metabolic effects (see below).

The non-nucleoside reverse transcriptase inhibitors include efavirenz and nevirapine. They interact with a number of drugs metabolized in the liver; the doses of protease inhibitors may need to be increased when they are given with efavirenz or nevirapine. Nevirapine is associated with a high incidence of rash (including Stevens-Johnson syndrome) and occasionally fatal hepatitis. Rash is also associated with efavirenz but it is usually milder. Efavirenz treatment has also been associated with an increased plasma cholesterol concentration.

**Initiation of Treatment**

The time for initiating antiviral treatment is determined by the clinical stage of the HIV infection as indicated by symptoms and where available, by the CD4-cell count or total lymphocyte count; the plasma viral load, if available, is also a valuable guide for staging the disease (see Monitoring, below). Recommended initial treatment with a combination of drugs (‘highly active antiretroviral therapy’, HAART) includes: 2 nucleoside reverse transcriptase inhibitors plus a
non-nucleoside reverse transcriptase inhibitor or a third nucleoside reverse transcriptase inhibitor or a protease inhibitor which may be combined with ritonavir as booster.

**Monitoring:**
In resource-limited settings the basic clinical assessment before initiating antiretroviral therapy includes documentation of past medical history, identification of current and past HIV-related illnesses, identification of co-existing medical conditions that may influence the choice of therapy (for example, pregnancy or tuberculosis) as well as current symptoms and physical signs.
The absolute minimum laboratory tests before initiating antiretroviral therapy are an HIV antibody test (in patients over 18 months of age) and a haemoglobin or haematocrit measurement.

Additional basic testing should include:
- white blood cell count;
- differential cell count (to identify a decline in neutrophils and the possibility of neutropenia);
- total lymphocyte count;
- serum alanine or aspartate aminotransferase concentration to assess the possibility of hepatitis co-infection and to monitor for hepatotoxicity;
- serum creatinine and/or blood urea nitrogen to assess baseline renal function;
- serum glucose;
- pregnancy tests for women.

Desirable supplemental tests include measurement of bilirubin, amylase and serum lipids. CD4-cell determinations are, of course, very desirable and efforts should be made to make these widely available. Viral load testing is currently considered optional because of constraints on resources.

**Changing Therapy:**
Deterioration of the condition (including clinical and virological changes) usually calls for replacement of the failing drugs. Intolerance to adverse effects and drug-induced organ dysfunction usually require change in therapy.
The choice of an alternative regimen depends on factors such as the response to previous treatment, tolerance and the possibility of cross-resistance. If treatment fails, a new second-line regimen will be needed. If toxicity occurs, either a new second-line regimen is indicated or, if the toxicity is related to an identifiable drug in the regimen, the offending drug can be replaced with another drug that does not have the same adverse effects.

**Pregnancy:**
Treatment of HIV infection in pregnancy aims to:
- minimize the viral load and disease progression in the mother;
- reduce the risk of toxicity to the fetus (although the teratogenic potential of most antiretroviral drugs is unknown);
- prevent transmission of infection to the neonate.

In pregnant women, it may be desirable to initiate antiretroviral therapy after the first trimester, although for pregnant women who are severely ill, the benefit of early therapy outweighs the potential risk to the fetus. All treatment options require careful assessment by a specialist.
The use of zidovudine, lamivudine, nevirapine, nelfinavir and saquinavir are recommended for women of child-bearing potential or who are pregnant. Efavirenz should be avoided because of its potential teratogenic effect on the fetus in the first trimester. First-line treatment in pregnant women should when possible include zidovudine and lamivudine. Monotherapy with either zidovudine or with nevirapine reduces transmission of infection to the neonate (see also below), but combination antiretroviral therapy maximizes the chance of preventing transmission and represents optimal therapy for the mother. Low-dose ritonavir is required if either indinavir or saquinavir is used in pregnancy because adequate drug concentration is achieved only with ritonavir boosting. Information is lacking on the use of lopinavir with ritonavir in pregnancy.

Lactic acidosis and hepatic steatosis associated with nucleoside reverse transcriptase inhibitors may be more frequent in pregnant women and therefore the combination of stavudine and didanosine should be used in pregnancy only when no alternatives are available. Protease inhibitors have been associated with glucose intolerance and pregnant women should be instructed to recognize symptoms of hyperglycaemia and to seek health care advice if they occur.

Various regimens have been used to specifically prevent the transmission of HIV from mother to the neonate at term. More information is available in New Data on the Prevention of Mother-to-Child Transmission of HIV and their Policy Implications: Conclusions and Recommendations (WHO/ RHR/01.28), which reflects an inter-agency consultation, held on 11-13 October 2000.

**Lactation:**
Antiretroviral drugs may be present in breastmilk and may reduce viral load in breastmilk and reduce the risk of transmission through lactation. However, the concentration of antiretroviral drugs in breastmilk may not be adequate to prevent viral replication and there is therefore the possibility of promoting the development of drug-resistant virus which could be transmitted to the infant.

Women with HIV infection should be counselled about the risks of lactation and, where possible, they should limit or avoid lactation; in particular, lactation should be avoided where replacement feeding is acceptable, affordable, sustainable and safe. HIV-infected women should be counselled on infant feeding options and they should be supported in their choice.

**Post-Exposure Prophylaxis:**
Treatment with antiretroviral drugs may be appropriate following occupational exposure to HIV-contaminated material. Immediate expert advice should be sought in such cases; national guidelines on post-exposure prophylaxis for healthcare workers have been developed and local ones may also be available.

**Lipodystrophy and Metabolic Effects:**
Combination antiretroviral therapy, including regimens containing a protease inhibitor, is associated with redistribution of body fat in some patients (for example, decreased fat under the skin, increased abdominal fat, ‘buffalo humps’ and breast enlargement). Protease inhibitors are also associated with metabolic abnormalities such as hyperlipidaemia, insulin resistance and hyperglycaemia. Clinical examination should include an evaluation of fat distribution; measurement of serum lipids and blood glucose should be considered.
Nucleoside/Nucleotide Reverse Transcriptase Inhibitors

In some settings it may not be possible to carry out full monitoring described under each drug entry; in such cases the level of monitoring should be determined by local guidelines (see also notes above).

Zidovudine

**ESL-D 536,537,538,539,540 Secondary hospitals**

**AVAILABILITY**

TABLETS 30, 40, 100 and 300 mg; capsules 100 and 300 mg; Syrup 50 mg/5 ml.

**DOSE**

Oral  HIV infection  
**Adult**- 600 mg daily in divided doses in combination with other antiretroviral drugs.  
**Child**- 6 weeks to 12 years: 160 mg/m2 every 8 hour, max. dose 200 mg every 8 hour.  
Prevention of maternal-foetal HIV transmission.  
**Adult**- 100 mg five times daily or 200 mg thrice daily or 300 mg twice daily, start treatment after 14th week of gestation until the start of labour.  
Prevention of HIV transmission in neonates.  
**Child**- neonates- 2 mg/kg every 6 hour for first 6 weeks of life, starting with 12 hour after birth.

**INDICATION**

HIV infection in combination with at least two other antiretroviral drugs; monotherapy for prevention of maternal-fetal HIV transmission

**CONTRAINDICATION**

Abnormally low neutrophil counts or haemoglobin; neonates either with hyperbilirubinaemia requiring treatment other than phototherapy or with raised transaminase; life threatening allergic reactions.

**PRECAUTION**

Haematological toxicity; vitamin B1 deficiency (increased risk of neutropenia); reduce dose or interrupt treatment if anaemia or myelosuppression; renal impairment (Appendix 7d); hepatic impairment (Appendix 7a); risk of lactic acidosis; elderly; lactation (Appendix 7b); interactions (Appendix 6c, 6d); pregnancy (Appendix 7c); myopathy, use with interferon and ribavirin based regimens in HIV/HCV coinfected patients, immune reconstitution syndrome.

**ADVERSE EFFECTS**

Anaemia (may require transfusion), neutropenia and leukopenia (all more frequent with high dose and advanced disease); also nausea and vomiting, abdominal pain, dyspepsia, diarrhoea, flatulence, taste disturbance, pancreatitis, liver disorders including fatty change and raised bilirubin and liver enzymes (see hepatic disease, above); chest pain, dyspnoea, cough; influenza-like symptoms; headache; fever; paraesthesia, neuropathy; convulsions; dizziness; somnolence, insomnia; anxiety; depression; malaise; anorexia; asthenia; myopathy; myalgia; pancytopenia; thrombocytopenia; gynaecomastia; urinary frequency; rash, pruritus, pigmentation of nail, skin and oral mucosa.

Lamivudine

**EDL-D737 Secondary hospitals**

**INDICATIONS**

HIV infection in combination with at least two other antiretroviral drugs.

**AVAILABILITY**

TABLETS 100, 150 and 300 mg; ORAL SOLUTION 50 mg/ml.

**DOSE**

Oral  
**Adult**- 150 mg twice daily administered with zidovudine. 
**Child**- 3 months to 12 years: 4 mg/kg body weight twice a day (max. 150 mg twice daily).
CONTRAINDICATIONS
Pregnancy (Appendix 7c); lactation (Appendix 7b); hepatic dysfunction (Appendix 7a); renal disease (Appendix 7d).

PRECAUTIONS
Renal impairment (Appendix 7d); hepatic disease (see below); pregnancy (Appendix 7c) and lactation (Appendix 7b) (see notes above); interactions (Appendix 6c). Potentially life-threatening lactic acidosis and severe hepatomegaly with steatosis reported therefore caution (particularly in obese women) in liver disease, liver enzyme abnormalities, or risk factors for liver disease; suspend or discontinue if deterioration in liver function tests, hepatic steatosis, progressive hepatomegaly or unexplained lactic acidosis. Recurrent hepatitis in patients with chronic hepatitis B may occur on discontinuation of lamivudine.

ADVERSE EFFECTS
Nausea, vomiting, diarrhoea, abdominal pain; cough; headache, fatigue, insomnia; malaise, fever, rash, alopecia, muscle disorders; nasal symptoms; peripheral neuropathy reported; rarely, pancreatitis (discontinue); neutropenia, anaemia, thrombocytopenia and red-cell aplasia; lactic acidosis; raised liver enzymes and serum amylase.

STORAGE
Store protected from moisture.

Stavudine
EDL-D737 Secondary hospitals

INDICATIONS
HIV infection in combination with at least two other antiretroviral drugs.

AVAILABILITY
Tablets /CAPSULES 30 and 40 mg.

DOSE
Oral
Adult- Under 60 kg: 30 mg every 12 h preferably at least 1 h before food. 60 kg and over: 40 mg every 12 h.
Neonate under 2 weeks- 500 μg/kg body weight.
Child- over 2 weeks and body weight under 30 kg: 1 mg/kg body weight every 12 h. 30 kg and over: 30 mg every 12 h.

CONTRAINDICATIONS
Hypersensitivity.

PRECAUTIONS
History of peripheral neuropathy (see below); history of pancreatitis or concomitant use with other drugs associated with pancreatitis; hepatic disease (see below); renal impairment; pregnancy (Appendix 7c) and lactation (Appendix 7b) (see notes above); fat redistribution, immune reconstitution syndrome. Suspend if peripheral neuropathy develops characterized by persistent numbness, tingling or pain in feet or hands; if symptoms resolve satisfactorily on withdrawal and if stavudine needs to be continued, resume treatment at half previous dose. Potentially life-threatening lactic acidosis and severe hepatomegaly with steatosis reported therefore caution in liver disease, liver enzyme abnormalities, or risk factors for liver disease (particularly in obese women); suspend or discontinue if deterioration in liver function tests, hepatic steatosis, progressive hepatomegaly or unexplained lactic acidosis.

ADVERSE EFFECTS
Peripheral neuropathy (dose-related, see above); pancreatitis; nausea, vomiting, diarrhoea, constipation, anorexia, abdominal discomfort; chest pain; dyspnoea; headache, dizziness, insomnia, mood changes; asthenia, musculoskeletal pain; influenza-like symptoms, rash and other allergic reactions; lymphadenopathy; neoplasms; elevated liver enzymes (see hepatic disease, above) and serum amylase; neutropenia, thrombocytopenia.
STORAGE
Store protected from moisture at a temperature not exceeding 30°C.

**Lamivudine + Nevirapine + Stavudine**
Non-EDL Secondary hospitals

INDICATIONS
HIV infection.

AVAILABILITY
TABLETS
Lamivudine + Nevirapine + Stavudine
40 mg + 10 mg + 70 mg
150 mg + 40 mg + 200 mg
150 mg + 30 mg + 200 mg
100 mg + 30 mg + 200 mg

DOSE
Adult- One tablet twice daily. Patients with body weight less than 50 kg, 2 mg/kg body weight two times a day.
Child- 3 months to 12 years; half adult dose is given two times a day.

PRECAUTIONS
Pregnancy (Appendix 7c).

STORAGE
Store protected from moisture at a temperature not exceeding 25°C for DT.

**Lamivudine + Zidovudine**
Non-EDL Secondary hospitals

INDICATIONS
HIV infection.

AVAILABILITY
TABLET lamivudine + zidovudine
150 mg + 300 mg.

DOSE
Adult- 2 tablets three times a day or as prescribed.
Child- Half the adult dose.

PRECAUTIONS
Pregnancy (Appendix 7c).

STORAGE
Store protected from moisture.

**Zidovudine + Lamivudine + Nevirapine**
EDL-D 762 Secondary hospitals

INDICATIONS
HIV infection.

AVAILABILITY
TABLETS Zidovudine 300 mg + Lamivudine 150 mg + Nevirapine 200 mg.

DOSE
Adult- 2 tablets three times a day.
Child- Half adult dose.
Other Antivirals

Oseltamivir

EDL-D 388,389,390,391,707 Tertiary

AVAILABILITY
CAPSULES 30, 45 and 75 mg.

DOSE
Oral Adult and adolescent- Prevention of influenza, over 13 years: 75 mg once daily for 10 days for post exposure prophylaxis, for up to 6 weeks in epidemics. Treatment of influenza, over 13 years: 75 mg every 12 h for 5 days. Child- Prevention of influenza: body weight under 15 kg: 30 mg once daily; 15 to 23 kg: 45 mg once daily; 23 to 40 kg: 60 mg once daily: above 40 kg: adult dose. Treatment of influenza: body weight under 15 kg: 39 mg every 12 h for 5 days; 15 to 23 kg: 45 mg every 12 h for 5 days; 23 to 40 kg: 60 mg every 12 h for 5 days; above 40 kg: adult dose.

INDICATION
Influenza A, B and its subtypes like swine flu.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Hepatic impairment; pregnancy (Appendix 7c); lactation; renal impairment.

ADVERSE EFFECTS
Nausea, vomiting, abdominal pain, dyspepsia, diarrhoea; headache, fatigue, insomnia, dizziness; conjunctivitis, epistaxis; rash; very rarely, hepatitis, Stevens-Johnson syndrome and toxic epidermal necrolysis; neuropsychiatric disorders also reported (in children); cough, bronchitis, eczema, seizures, aggravation of diabetes.

Nevirapine

EDL-D 363,364 Tertiary

AVAILABILITY
TABLET/Capsule 200 mg; ORAL SUSPENSION 100 mg/5 ml.

DOSE
Oral Adult- 200 mg once a day for 14 days, if tolerated and no rash is observed then increase to 200 mg two times a day. Child- 2 months to 8 years: 4 mg/kg body weight once a day for 14 days, if tolerated and no rash is observed increase to 4 mg/kg body weight two times a day.

INDICATION
HIV infection, in combination with at least two other antiretroviral drugs; prevention of mother-to-child transmission in HIV-infected patients.

CONTRAINDICATION
Acute porphyria; severe hepatic impairment; post-exposure prophylaxis; breast feeding.

PRECAUTION
Hepatic impairment; history of chronic hepatitis (greater risk of hepatic adverse effects), pregnancy and lactation; interactions. Potentially life-threatening hepatotoxicity including fatal fulminant hepatitis reported usually occurring in first 8 weeks; monitor liver function before long-term treatment then every 2 weeks for 2 months then after 1 month and then every 3-6 months; discontinue permanently if abnormalities in liver function tests accompanied by hypersensitivity reaction (rash, fever, arthralgia, myalgia, lymphadenopathy, hepatitis, renal impairment, eosinophilia, granulocytopenia); suspend if severe abnormalities in liver function tests but no hypersensitivity reaction-discontinue permanently if significant liver function abnormalities recur; monitor patient closely if mild to moderate abnormalities in liver function
tests with no hypersensitivity reaction. Rash, usually in first 8 weeks, is most common adverse effect; incidence reduced if introduced at low dose and dose increased gradually; discontinue permanently if severe rash or if rash accompanied by blistering, oral lesions, conjunctivitis, swelling, general malaise or hypersensitivity reactions; if rash mild or moderate may continue without interruption but dose should not be increased until rash resolves. Patients should be told how to recognize hypersensitivity reactions and advised to seek immediate medical attention if symptoms develop.

ADVERSE EFFECTS
Rash including Stevens-Johnson syndrome and rarely, toxic epidermal necrolysis; hepatitis or jaundice reported; nausea, vomiting, abdominal pain, diarrhoea, headache, drowsiness, fatigue, fever; hypersensitivity reactions; anaphylaxis, angioedema, urticaria also reported; granulocytopenia.

Antiprotozoal medicines

Amoebiasis:
Amoebic dysentery is caused by *Entamoeba histolytica*. It is transmitted by the faeco-oral route and infection is usually caused by ingestion of cysts from contaminated food and drink. Asymptomatic carriers are common in endemic areas. In non-endemic areas, symptomatic carriers should be treated with a luminal amoebicide which will reduce the risk of transmission and protect the patient from invasive amoebiasis. Diloxanide furoate is most widely used, but other compounds, including clefamide, etofamide and teclozan, are also effective. Treatment with diloxanide furoate is regarded as successful if stools are free of *E. histolytica* for one month. Several specimens should be examined in evaluating response to treatment.

Symptomatic (invasive) amoebiasis may be classified as intestinal or extra-intestinal. Intestinal amoebiasis is either amoebic dysentery or non-dysenteric amoebic colitis. Extraintestinal amoebiasis most commonly involves the liver, but may involve the skin, genito-urinary tract, lung and brain. Invasive amoebiasis is more likely in malnutrition, immunosuppression and pregnancy. Amoebic dysentery may take a fulminating course in late pregnancy and the puerperium; treatment with metronidazole may be life saving. In less severe infection, metronidazole should, if possible, be avoided in the first trimester. All patients with invasive amoebiasis require treatment with a systemically active compound such as metronidazole, ornidazole and tinidazole followed by a luminal amoebicide in order to eliminate any surviving organisms in the colon. Combined preparations are useful. In severe cases of amoebic dysentery, tetracycline given in combination with a systemic amoebicide lessens the risk of superinfection, intestinal perforation and peritonitis. Hepatic abscesses should be lanced by needle aspiration.

Giardiasis:
Giardiasis is caused by Giardia intestinalis and is acquired by oral ingestion of Giardia cysts. Giardiasis can be treated with tinidazole in a single dose or with another 5-nitroimidazole such as metronidazole; both are highly effective and should be offered when practicable to all infected patients. Family and institutional contacts should also be treated. Larger epidemics are difficult to eradicate because of the high proportion of symptomatic carriers and because excreted cysts can survive for long periods outside the human host.

Trichomoniasis:
Trichomoniasis is an infection of the genito-urinary tract caused by Trichomonas vaginalis and transmission is usually sexual. In women it causes vaginitis although some are asymptomatic. It
is usually asymptomatic in men but may cause urethritis. Patients and their sexual partners should be treated with metronidazole or other nitroimidazole.

**Tinidazole**

**EDL-D 507,508 Secondary hospitals**

**AVAILABILITY**

TABLETS 300 and 500 mg, 1g; INJECTION 400 ml infusion (2 mg/ml); Suspension 75 mg/5 ml, 150 mg/5 ml.

**DOSE**

Oral Anaerobic infections: Adult- 2g on first day, followed by 1g daily or 0.5g twice daily for 5-6 days. Amoebiasis: Adult- 1.5 - 2g daily as a single dose for 3 - 6 days. Child- 30-50 mg/kg daily as a single dose for 3 days. Trichomoniasis and giardiasis: Adult- 2g as a single dose. Child- 50 to 75 mg/kg as a single dose. Parenteral Bacterial vaginosis and ulcerative gingivitis: Adult- 2g as a single dose parenterally. Anaerobic infections: Adult- Initially 800 mg/400 ml infused i.v. at a rate of 10 ml/minute followed by 800 mg daily. Abdominal surgical prophylaxis: Adult- 2.0g as single i.v. infusion 12 h prior to surgery.

**INDICATION**

Amoebiasis, trichomoniasis and giardiasis, anaerobic infections, necrotising ulcerative gingivitis, bacterial vaginosis, H. pylori associated peptic ulcers, abdominal surgery prophylaxis.

**CONTRAINDICATION**

Hypersensitivity to nitroimidazole derivatives, first trimester of pregnancy, lactation, blood dyscrasias, porphyria; interactions

**PRECAUTION**

Seizures, peripheral neuropathy, CNS disease, disulfiram-like reaction with alcohol

**ADVERSE EFFECTS**

Similar to metronidazole.

**Metronidazole**

**EDL-D 344,345,346 PHC**

**AVAILABILITY**

TABLETS 200 and 400 mg; SUSPENSION 200 mg/5 ml; INJECTION 100 ml infusion (5 mg/ ml).

**DOSE**

Oral Adult- Amoebiasis: 400 to 800 mg every 8 h for 5 to 7 days. GiardiasisL: 200 mg three times a day for 7 to 10 days or intravenous injection 500 mg 8 hly for 7 days. Child- Amoebiasis: Below 12 years; 7.5 mg/ kg body weight. 12 years and above; 35 to 50 mg/kg body weight daily in three divided doses.

**INDICATION**

Anaerobic bacterial infections including gingivitis, pelvic inflammatory disease, tetanus, peritonitis, brain abscess, necrotizing pneumonia, antibiotic-associated colitis, leg ulcers and pressure sores and surgical prophylaxis; bacterial vaginosis; tissue nematode infections; trichomonal vaginitis, amoebiasis and giardiasis; Helicobacter pylori eradication.

**CONTRAINDICATION**

Chronic alcohol dependence; neurological disease, blood dyscrasias, first trimester of pregnancy.

**PRECAUTION**

Disulfiram-like reaction with alcohol; hepatic impairment and hepatic encephalopathy (Appendix 7a); lactation (Appendix 7b); clinical and laboratory monitoring in courses lasting longer than 10 days; interactions (Appendix 6a, 6c, 6d); pregnancy (Appendix 7c); phenobarbitone, history of blood dyscrasias.
ADVERSE EFFECTS
Nausea, vomiting, unpleasant metallic taste, furred tongue and gastrointestinal disturbances; rarely, headache, drowsiness, dizziness, ataxia, darkening of urine, erythema multiforme, pruritus, urticaria, angioedema and anaphylaxis; abnormal liver function tests, hepatitis, jaundice; thrombocytopenia, aplastic anaemia; myalgia, arthralgia; peripheral neuropathy, epileptiform seizures; leukopenia on prolonged or high dosage regimens; anorexia, glossitis, dryness of mouth.

Metronidazole Benzoate
EDL-D 347 PHC
AVAILABILITY
TABLETS 200 and 400 mg; SUSPENSION 200 mg/5 ml; INJECTION 100 ml infusion (5 mg/ ml).
DOSE
Oral Adult- Amoebiasis: 400 to 800 mg every 8 h for 5 to 7 days. GiardiasisL: 200 mg three times a day for 7 to 10 days or intravenous injection 500 mg 8 hly for 7 days. Child- Amoebiasis: Below 12 years; 7.5 mg/ kg body weight. 12 years and above; 35 to 50 mg/kg body weight daily in three divided doses.

INDICATION
Anaerobic bacterial infections including gingivitis, pelvic inflammatory disease, tetanus, peritonitis, brain abscess, necrotizing pneumonia, antibiotic-associated colitis, leg ulcers and pressure sores and surgical prophylaxis; bacterial vaginosis; tissue nematode infections; trichomonal vaginitis, amoebiasis and giardiasis; Helicobacter pylori eradication.

CONTRAINDICATION
Chronic alcohol dependence; neurological disease, blood dyscrasias, first trimester of pregnancy.

PRECAUTION
Disulfiram-like reaction with alcohol; hepatic impairment and hepatic encephalopathy; lactation; clinical and laboratory monitoring in courses lasting longer than 10 days; interactions; pregnancy phenobarbitone, history of blood dyscrasias.

ADVERSE EFFECTS
Nausea, vomiting, unpleasant metallic taste, furred tongue and gastrointestinal disturbances; rarely, headache, drowsiness, dizziness, ataxia, darkening of urine, erythema multiforme, pruritus, urticaria, angioedema and anaphylaxis; abnormal liver function tests, hepatitis, jaundice; thrombocytopenia, aplastic anaemia; myalgia, arthralgia; peripheral neuropathy, epileptiform seizures; leukopenia on prolonged or high dosage regimens; anorexia, glossitis, dryness of mouth.

Antimalarial medicines
Human malaria, which is transmitted by female anopheline mosquitoes (and rarely, by congenital transmission, transfusion of infected blood or use of contaminated syringes among drug addicts), is caused by four species of plasmodial parasites. *Plasmodium vivax* is the most extensively distributed and causes much debilitating disease. *P. falciparum* is also widespread and causes the most severe infections which are responsible for nearly all malaria-related deaths. *P. ovale* is mainly confined to Africa and is less prevalent, while *P. malariae*, which causes the least severe but most persistent infections, also occurs widely.

Certain tissue forms of *P. vivax* and *P. ovale* which persist in the liver for many months and even years are responsible for the relapses characteristic of malaria. Such latent forms are not
generated by *P. falciparum* or *P. malariae*. Recrudescence of these infections results from persistent blood forms in inadequately treated or untreated patients.

**Treatment of Malaria:**

Blood schizonticides, which suppress malaria by destroying the asexual blood forms of the parasites, are the mainstay of the treatment of acute malaria and some are used for prophylaxis. They include the 4-aminoquinolines (example amodiaquine and chloroquine), the related arylaminoalcohols (example mefloquine and quinine) and artemisinin and its derivatives (example artesunate and artesunate). Blood schizonticides are not active against intrahepatic forms and therefore they do not eliminate infections by *P. vivax* and *P. ovale*.

Some antimetabolites act synergistically when given in combination. For example, pyrimethamine in combination with a sulfonamide (sulfadoxine) or sulfone and some antibiotics (for example doxycycline) are blood schizonticides. Because they act more slowly, these substances are of little value when used alone. The tetracyclines are used primarily as adjuncts to quinine where multiple-drug-resistant *P. falciparum* is prevalent. Chloroquine, a rapidly acting schizonticide, is well tolerated, safe and inexpensive. It should be used to treat malaria wherever the parasites remain susceptible. *P. malariae* and *P. ovale* remain fully sensitive to chloroquine.

A 3-day course of chloroquine by mouth is sufficient to eliminate susceptible *P. falciparum* infections because effective plasma-chloroquine concentration is sustained for several weeks.

If subsequent relapse occurs in *P. ovale* and *P. vivax* infections primaquine should be administered, after a second course of chloroquine, to eliminate the intrahepatic infection. Amodiaquine is an alternative to chloroquine for the treatment of uncomplicated *P. falciparum* infection; but crossresistance with chloroquine exists in some areas. It should preferably be used as part of combination therapy with other antimalarials, for example artesunate. Hepatitis and blood disorders were reported when amodiaquine was used for prophylaxis of malaria; patients should be told how to recognize the symptoms of these conditions and advised to seek medical help if they occur. The combination of sulfadoxine with pyrimethamine is recommended for the treatment of malaria only in areas of high chloroquine resistance. A single dose of sulfadoxine with pyrimethamine is usually sufficient to eliminate infection; quinine should also be given for 3 days in patients in whom quinine may accelerate reduction of parasitaemia and in those at risk of fulminating disease. Because sulfonamides are associated with a risk of haemolysis and methaemoglobinaemia in the newborn, quinine is preferred to treat chloroquine-resistant malaria during pregnancy.

Mefloquine is generally well tolerated, although, some adverse effects have been reported (see notes). However, because of the danger of the emergence of mefloquine-resistant strains of *P. falciparum* and because of its potential toxicity, it should be used only following either microscopic or careful clinical diagnosis of *P. falciparum* infections that are known or strongly suspected to be resistant to chloroquine or sulfadoxine with pyrimethamine.

Quinine, given orally, should be reserved for *P. falciparum* infections likely to be unresponsive to other drugs. Doxycycline, which is an effective oral schizonticide, should be given in combination with quinine except in pregnant women and children under 8 years.
In multi-drug resistant malaria, preparations of artemisinin or its derivatives (artemether or artesunate) offer the only prospect of cure. They should not be used in the first trimester of pregnancy.

For the treatment of multi-drug resistant falciparum malaria oral artesunate may be an effective antimalarial. It should always be given in combination with mefloquine. Parenteral artemether or artesunate, whose use is restricted, are effective alternatives to quinine for the treatment of severe falciparum malaria and are preferred in areas where decreased efficacy of quinine has been documented. To ensure radical cure following parenteral treatment with artemether or oral treatment with artesunate, a full therapeutic dose of mefloquine should be given. A fixed-dose oral formulation of artemether with lumefantrine has recently become available and is recommended for the treatment of uncomplicated falciparum malaria in areas with significant resistance. The combination is not for use in pregnancy or lactation.

**Prophylaxis Against Malaria:**

No drug regimen gives assured protection to everybody and indiscriminate use of antimalarials can increase the risk of inducing resistance. Chloroquine, which is usually well tolerated at the required dosage, is preferred where *P. falciparum* remains fully sensitive. The combination of proguanil with chloroquine may overcome mild chloroquine resistance. Chloroquine must be started 1 week before exposure and be continued in pregnant women until after delivery and for at least 4 weeks after the last risk of exposure in the case of non-immune individuals. This is sufficient to ensure elimination of *P. falciparum* and *P. malariae*, but not of *P. vivax* and *P. ovale*, whose residual hepatic forms survive.

Mefloquine may be used for prophylaxis in areas of high risk or where multiple-drug resistance has been reported. Where possible prophylaxis should be started 2-3 weeks before travel to enable any adverse reactions to be identified before exposure (over three-quarters of adverse reactions occur by the third dose) and should be continued for 4 weeks after last exposure. Mefloquine may be used for prophylaxis during the second and third trimesters. It should be used in early pregnancy only if alternative drugs are either not available or unlikely to be effective and when it is impracticable for the woman to leave the endemic area.

Proguanil, a predominantly tissue schizonticide with little blood schizonticidal activity, is a causal prophylactic agent since it is active against pre-erythrocytic intrahepatic forms, particularly of *P. falciparum*. The latent persistent liver forms of *P. ovale* and *P. vivax* are unresponsive. However, there is evidence that it may be effective against *P. vivax* only immediately after the initial infection. *P. falciparum* resistance to proguanil or related compounds may occur in malaria endemic areas and particularly where it has been employed in mass prophylaxis. Proguanil is used for prophylaxis with chloroquine in areas where there is resistance to chloroquine but a low risk of infection as it may give some protection against and may alleviate symptoms if an attack occurs. Proguanil and chloroquine may also be used prophylactically in areas of high risk or multi-drug resistance as a second choice where mefloquine is not appropriate.

There is no evidence that proguanil is harmful in prophylactic doses during pregnancy. Because of the vulnerability of pregnant women to falciparum malaria, it should be used at full
prophylactic dosage wherever the disease is prevalent and likely to be responsive to proguanil, if chloroquine is not available or with chloroquine, if the latter alone is unlikely to be effective.

**Artemether**  
**EDL-D 47 Secondary hospitals**  
**AVAILABILITY**  
CAPSULE 40 mg; INJECTION 1 ml ampoule (80 mg/ml, 160 mg/2 ml).  
**DOSE**  
Oral Adult- 160 mg in two divided doses on first day followed by 80 mg once a day for next four days. Intramuscular injection Adult- 80 mg twice a day for 3 days. Child- 1.6 mg/kg body weight twice a day followed by 1.6 mg/kg body weight once a day for 4 days, alternatively 1.6 mg/kg body weight twice a day for 3 days.  
**INDICATION**  
Treatment of severe P. falciparum malaria in areas where evidence is there that quinine is ineffective; multi drug resistant malaria.  
**CONTRAINDICATION**  
First trimester of pregnancy and hypersensitivity.  
**PRECAUTION**  
Electrolyte disturbances, concomitant use with other drugs known to cause QT-interval prolongation; hepatic impairment; renal impairment; monitor patients unable to take food (greater risk of recrudescence); interactions (Appendix 6c); lactation (Appendix 7b). Dizziness may impair ability to perform skilled tasks, for example operating machinery, driving.  
**ADVERSE EFFECTS**  
Headache, nausea, vomiting, abdominal pain, diarrhoea; dizziness, tinnitus, neutropenia, elevated liver enzyme values; cardiotoxicity (after high doses); neurotoxicity-in animal studies; decrease in reticulocyte count.

**Artesunate**  
**EDL-D 50 PHC**  
**AVAILABILITY**  
TABLET 25, 50 & 60 mg; INJECTION 50, 60, 1000 & 2000 mg/vial.  
**DOSE**  
Oral Adult- total oral dose 600 mg can be divided into two 50 mg tablets twice a day on first day thereafter 50 mg twice a day for next 4 days. Child- half adult dose. Intramuscular injection 60 mg twice daily.  
**INDICATION**  
Treatment of uncomplicated P. falciparum malaria in areas of multiple drug resistance.  
**CONTRAINDICATION**  
First trimester of pregnancy and hypersensitivity.  
**PRECAUTION**  
Risk of recurrence if used alone in nonimmune patients; hepatic/renal insufficiency, pregnancy (Appendix 7c), lactation, paediatrics. Dizziness may impair ability to perform skilled tasks, for example operating machinery, driving.  
**ADVERSE EFFECTS**  
Headache, nausea, vomiting, abdominal pain, diarrhoea, dizziness, tinnitus, neutropenia, elevated liver enzyme values; ECG abnormalities, including prolongation of QT interval;
temporary suppression of reticulocyte response and induction of blackwater fever reported; neurotoxicity-in animal studies

Chloroquine

**EDL-D 112,113,114 Universal**

**INDICATIONS**

Treatment of acute malaria caused by *P. malariae* and susceptible *P. falciparum*; *P. vivax* and *P. ovale* (followed by primaquine to eliminate intrahepatic forms); prophylaxis of malaria for pregnant women and nonimmune individuals at risk; rheumatic disorders.

**AVAILABILITY**

TABLETS 250 and 500 mg; INJECTION 10 and 30 ml (40 mg/ml); SUSPENSION 50 mg/ml.

**DOSE**

**Oral**

**Adult**- Immediately 600 mg, after 6 h 300 mg followed by 300 mg daily for 2 days.

**Child**- 10 mg/kg body weight followed by 5 mg/kg body weight after 6 h, thereafter once a day for 2 days.

**Intramuscular injection**

**Adult**- 10 ml followed by 5 ml after 6 h. Thereafter 5 ml daily for two days.

**Child**- 5 mg/kg body weight administered every 12 h followed by oral therapy.

**CONTRAINDICATIONS**

Severe haematologic distress or gastrointestinal distress; eye dysfunction; liver disease.

**PRECAUTIONS**

If patient continues to deteriorate after chloroquine-suspect resistance and administer quinine intravenously as emergency measure; hepatic impairment; renal impairment (Appendix 7d); pregnancy (but in malaria, benefit considered to outweigh risk; Appendix 7c); lactation (Appendix 7b); may exacerbate psoriasis; neurological disorders (avoid for prophylaxis if history of epilepsy); may aggravate myasthenia gravis; severe gastrointestinal disorders; G-6-PD deficiency; avoid concurrent therapy with hepatotoxic drugs; interactions (Appendix 6c, 6d).

**ADVERSE EFFECTS**

Headache, gastrointestinal disturbances; also convulsions; visual disturbances (retinopathy associated with long-term, high dose therapy or inappropriate selfmedication); depigmentation or loss of hair; rashes; pruritus-may become intolerable; bone-marrow suppression; hypersensitivity reactions such as urticaria and angioedema; atrioventricular block (may be result of inappropriate self-medication); porphyria and psoriasis in susceptible individuals.

**STORAGE**

Store protected from light.

Primaquine Diphosphate

**EDL-D 432,433 Universal**

**AVAILABILITY**

TABLETS 2.5, 7.5 and 15 mg.

**DOSE**

**Radical treatment**

**Adult**- 15 mg daily for 14 days, may be increased to higher dose. **Child**- 250 μg/kg daily for 14 days. Malaria prophylaxis **Adult**- 30 mg once daily; **Child**- 0.5 mg/kg once daily (to be started 1-2 days before travel and continue for 7 days after departure from malaria endemic area). Gametocidal treatment of *P. falciparum* malaria (after standard blood schizontocide therapy). **Adult** and **Child**- 500–50 μg/kg as a single dose.

**INDICATION**

Radical cure of *P. vivax* and *P. ovale* malaria (after chloroquine therapy to eradicate erythrocytic forms), elimination of gametocytes of *P. Falciparum*, malaria prophylaxis.

**CONTRAINDICATION**

Hypersensitivity, granulocytopenia, pregnancy, lactation, children below 1 year.
PRECAUTION
Patients with history of granulocytosis/ methaemoglobinemia, G-6-PD deficiency, monitor Hb levels, blood counts routinely and withdraw if signs of haemolysis or methaemoglobinemia occur; lactation(Appendix 7b)

ADVERSE EFFECTS
Nausea, vomiting, abdominal cramps, haemolytic anaemia in G-6-PD deficient patients; rarely, leukopenia, agranulocytosis, leukocytosis, methaemoglobinemia and cardiac arrhythmias.

Quinine (bisulphate or sulphate)
EDL-D 448,449 PHC

AVAILABILITY
TABLETS 100, 150, 300 and 600 mg; SUSPENSION 150 mg/5 ml; INJECTION 1 and 2 ml ampoule (300 mg/ml).

DOSE
Oral Adult- 300 to 600 mg every 8 h in divided doses for 5 to 7 days. Child- 25 mg/kg body weight every 8 h in divided doses for 5 to 7 days. Intravenous infusion for patients unable to swallow tablets Loading dose 900 mg to 1.4g infused over 4 h, then 300 to 600 mg every 8 h infused over 4 h.

INDICATION
Multiple drug resistant P. falciparum malaria.

CONTRAINDICATION
Haemoglobinuria; optic neuritis; tinnitus; quinine resistant falciparum, pregnancy, lactation, prolonged QT interval

PRECAUTION
Atrial fibrillation, conduction defects, heart block; monitor for signs of cardiac toxicity and blood glucose levels (with intravenous use); renal impairment (Appendix 7d); G-6-PD deficiency; may aggravate myasthenia gravis; interactions(Appendix 6d)

ADVERSE EFFECTS
Cinchonism (tinnitus, headache, blurred vision, temporary blindness, altered auditory acuity, nausea, diarrhoea, hot and flushed skin, rashes, confusion); hypersensitivity reactions including angioedema; rarely, haemorrhage and asthma; hypoglycaemia (especially after parenteral administration); renal damage (culminating in acute renal failure and anuria); blood disorders; cardiovascular, gastrointestinal and CNS effects; very toxic in overdosage-immediate medical attention required; acute haemolytic anaemia.

ACT combi bister pack
EDL-D 450,555,558,554,556,557 Universal

INDICATIONS
Treatment of malaria due to susceptible P. falciparum in areas of high chloroquine resistance and in patients who have not responded to chloroquine; additionally quinine may be given for 3 days.

DOSE
(PINK COLOUR) ORAL
infant less than 1 year
Total dose of Artesunate – 75 mg divided over three days, Sulphadoxine pyremethamine (250 mg + 12.5 mg) single dose

Each Combi Blister Pack: containing 3 tablet of Artesunate (each tablet of Artesunate 25mg strength) and 1 tablet of Sulphadoxine Pyremethamine (250 mg + 12.5 mg)

Each row – No. of tablets:
First Row (Day 1): One tablet of Artesunate (25 mg) and one tablet of Sulphadoxine-Pyremethamine ((250 mg + 12.5 mg)
Second Row (Day 2): one tablet of Artesunate (25 mg)
Third Row (Day 3): one tablet of Artesunate (25 mg)

CAUTION:
The blister should be superscribed that SP is not to be given to children under 5 months and should be treated with alternate ACTs.

DOSE (YELLOW COLOUR) ORAL
CHILD 1-4 YEARS

Total dose of Artesunate – 150 mg divided over three days, Sulphadoxine pyremethamine – (500+25)mg single dose

Each Combi Blister Pack: containing 3 tablets of Artesunate (50 mg each) and 1 tablet of Sulphadoxine Pyremethamine (500+25)mg

Each row – No. of tablets:
First Row (Day 1): One tablet of Artesunate (50 mg) and One tablet of Sulphadoxine Pyremethamine (500+25)mg
Second Row (Day 2): One tablet of Artesunate (50 mg)
Third Row (Day 3): One tablet of Artesunate (50 mg)

DOSE (GREEN COLOUR) ORAL
CHILD 5-8 YEARS

Total dose of Artesunate – 300 mg divided over three days, Sulphadoxine pyremethamine (750 +37.5) mg single dose

Each Combi Blister Pack containing 3 tablets of Artesunate (100mg each) and 1 tablet of Sulphadoxine Pyremethamine (750+37.5) mg

Each row – No. of tablets:
First Row (Day 1): one tablet of Artesunate (100mg) and one tablet of sulphadoxine Pyremethamine (750mg+ 37.5) mg
Second Row (Day 2): one tablet of Artesunate (100mg)
Third Row (Day 3): one tablet of Artesunate (100mg)

DOSE (RED COLOUR)
ORAL
CHILD 9-14 YEARS
Total dose of Artesunate – 450 mg divided over three days, Sulphadoxine pyremethamine (1000 +50) mg single dose
Each Combi Blister Pack containing 3 tablets of Artesunate 150 mg and 2 tablets of Sulfadoxine Pyremethamine (500mg+ 25mg)
Each row – No. of tablets
First Row (Day 1): One tablets of Artesunate (150 mg) and two tablets of Sulphadoxine Pyremethamine 500+25 mg) mg each
Second Row (Day 2): One tablet of Artesunate (150 g)
Third Row (Day 3): One tablet of Artesunate (150 mg)

DOSE (WHITE COLOUR)
ORAL ADULTS

Total dose of Artesunate – 600 mg divided over three days, Sulphadoxine pyremethamine – (1500 +75)mg single dose
Each Combi Blister Pack containing 3 tablets of Artesunate (each 200 mg) and 2 tablets of Sulphadoxine Pyremethamine (750+37.5)mg each or 3 tablets of Sulphadoxine Pyremethamine (500+25) mg each
Each row – No. of tablets:
First Row (Day 1): one tablet of Artesunate (200 mg) and two tablets of Sulphadoxine pyremethamine (750+37.5) mg each or three tablets of Sulphadoxine Pyremethamine (500+25) mg each
Second Row (Day 2) one tablet of Artesunate (200mg)
Third Row (Day 3): one tablet of Artesunate (200 mg)

CONTRAINDICATIONS
Hypersensitivity to sulfonamides or pyrimethamine; severe hepatic or renal impairment (except where no alternative treatment available); blood dyscrasias, neonate megaloblastic anaemia and folate deficiency.

PRECAUTIONS
Avoid in blood disorders-unless specialist supervision; discontinue immediately if blood disorder occurs; rash, sore throat, mouth ulcers, or shortness of breath withdraw treatment; G-6-PD deficiency; predisposition to folate deficiency; hepatic impairment (Appendix 7a); pregnancy (Appendix 7c); lactation (Appendix 7b); interactions (Append 6c).

ADVERSE EFFECTS
Rashes, pruritus, slight hair loss; rarely, erythema multiforme (Stevens-Johnson syndrom and toxic epidermal necrolysis; gastrointestinal disturbances including nausea, vomitin stomatitis; rarely, hepatitis, leukopenia, thrombocytopenia, megaloblastic anaemia or purpurawithdraw treatment; fatigue, headache, fever, polyneuritis, also report pulmonary infiltrates such as eosinophilic or allergic alveolitis-if symptoms of cough or shortness of breath-withdraw treatment.

STORAGE
Store protected from light and moisture.

Arteether
EDL D 562 PHC

INDICATIONS
Complicated falciparum malaria; chloroquine resistant malaria; cerebral malaria.

AVAILABILITY
INJECTION 2 ml ampoule (150 mg/2 ml).
Arteether is an ethyl derivative of dihydroartemisinin. It is a mixture of α and β arteether in a 30:70 ratio.

**DOSE**
Adult- 150 mg daily i.m. injection, once daily for 3 consecutive days.

**CONTRAINDICATIONS**
Hypersensitivity to artemisinin derivatives; pregnancy (Appendix 7c).

**ADVERSE REACTIONS**
It is clinically very well tolerated without any significant side effects; neurological or biochemical.

**STORAGE**
Store protected from light in tamper evident container so as to avoid contamination by micro-organisms.
Chronic recurrent headache is associated with many disorders, both somatic and psychogenic. An accurate diagnosis must consequently be made before appropriate treatment can be initiated for migraine. Untreated migraine attacks last for several hours and sometimes for as long as 3 days.

Migraine headache is frequently accompanied by episodes of gastrointestinal disturbance including nausea and vomiting. The headache may be preceded or accompanied by aura (classical migraine) which is characterised by visual disturbances such as flickering lines and fragmented vision or sensory disturbances such as tingling or numbness; rarely, hemiparesis or impaired consciousness may occur. Migraine without aura (common migraine) is the more common form occurring in about 75% of patients who experience migraine.

Emotional or physical stress, lack of or excess sleep, missed meals, menstruation, alcohol and specific foods including cheese and chocolate are often identified as precipitating factors; oral contraceptives may increase the frequency of attacks. Avoidance of such precipitating factors can be of great benefit in preventing or reducing the frequency of attacks and should be addressed in detail. Women taking combined oral contraceptives who experience an onset or increase in frequency of headaches should be advised of other contraceptive measures.

The two principal strategies of migraine management are treatment of acute attacks and prophylactic treatment.

**FOR ACUTE ATTACK**

Treatment of acute attacks may be non-specific using simple analgesics, or specific using an ergot alkaloid such as ergotamine. If nausea and vomiting are features of the attack, an antiemetic drug may be given. Treatment is generally by mouth; some drugs are available as suppositories which may be administered if the oral route is not effective (poor oral bioavailability, or absorption from the gut impaired by vomiting) or not practicable (patient unable to take drugs orally).

Simple analgesics including NSAIDs (nonsteroidal anti-inflammatory drugs) can be effective in mild to moderate forms of migraine if taken early in the attack; most migraine headaches respond to paracetamol, acetylsalicylic acid or NSAID such as ibuprofen or naproxen sodium. Peristalsis is often reduced during migraine attacks and, if available, a dispersible or effervescent preparation of the drug is preferred because of enhanced absorption compared with a conventional tablet. The risk of Reye syndrome due to acetylsalicylic acid in children can be avoided by giving paracetamol instead. Frequent and prolonged use of analgesics by migraine sufferers may lead to analgesic-induced headache.
**Ergotamine** should be considered only when attacks are unresponsive to non-opioid analgesics. It is poorly absorbed when taken orally or sublingually. Rectal suppositories may offer an advantage when other routes of administration are unsatisfactory. To be fully effective, ergotamine must be taken in adequate amounts as early as possible during each attack. Adverse effects limit how much ergotamine can be used in a single attack and consequently the recommended dosage should never be exceeded and at least four days should elapse between successive treatments. Even normal dosage can lead to dependence, tolerance to adverse effects and to a withdrawal syndrome on discontinuing the drug. To avoid dependence the frequency of administration should be limited to no more than twice a month. Adverse effects include nausea, vomiting, diarrhoea and vertigo; chronic ergotism is characterized by severe peripheral vasoconstriction which can lead to gangrene in the extremities. The severity of adverse effects prevents the use of ergotamine for migraine prophylaxis.

An antiemetic such as **metoclopramide**, given as a single dose orally or by intramuscular injection at the onset of a migraine attack, preferably 10-15 min before the analgesic or ergotamine, is useful not only in relieving nausea but also in restoring gastric motility, thus improving absorption of the antimigraine drug. Products which contain barbiturates or codeine are undesirable, particularly in combination with ergotamine, since they may cause physical dependence and withdrawal headaches.

**Acetylsalicylic Acid**

**EDL-D4 PHC**

**Indications**
- Management of mild to moderate pain such as headache, acute migraine attacks, transient musculoskeletal pain, dysmenorrheal pain and for reducing fever; pain and inflammation of rheumatoid arthritis; antiplatelet agent for prophylaxis of myocardial infarction, stable angina pectoris; stroke prophylaxis.

**AVAILABILITY**
- TABLETS 50, 60, 75, 80, 150, 300 and 325 mg.

**DOSE**
- **Oral**
  - Adult - Analgesic and antipyretic including migraine attacks: 0.3 to 0.9g, 3 to 4 times a day (max. 4g daily). Acute Rheumatic fever: 4 to 6g or 75 to 100 mg/kg daily in divided doses. Antiplatelet: 75-325 mg/day.
  - Child - Under 16 years: not recommended (can cause Reye’s syndrome).

**CONTRAINDICATIONS**
- Hypersensitivity (including asthma; angioedema; urticaria or rhinitis) to acetylsalicylic Acidor any other NSAID; children and adolescents under 16 years (may cause Reye’s syndrome); gastrointestinal ulceration; haemophilia and other bleeding disorders; not for treatment of gout; severe renal or hepatic impairment; lactation. It is known to cause haemolytic anaemia in people who have the genetic disease- G-6-PD-deficiency.

**PRECAUTIONS**
Asthma, allergic disease; impaired renal or hepatic function (Appendices 7d and 7a); lactation (Appendix 7b); pregnancy (Appendix 7c); elderly; G-6-PD-deficiency; dehydration; interactions (Appendix 6a, 6c, 6d).

ADVERSE EFFECTS
Generally mild and infrequent for lower doses, but common with anti-inflammatory doses; gastrointestinal discomfort or nausea, ulceration with occult bleeding (occasionally major haemorrhage); also other haemorrhage (including subconjunctival); hearing disturbances such as tinnitus (rarely, deafness); vertigo; confusion; hypersensitivity reactions (angioedema; bronchospasm and rash); increased bleeding time, blood disorders (particularly thrombocytopenia); rarely, oedema; myocarditis; Reye’s syndrome.

STORAGE
Store protected from moisture at a temperature not exceeding 30°C.

Paracetamol
EDL-D 395 PHC

INDICATIONS
Mild to moderate pain including dysmenorrhoeal pain, headache; pain relief in osteoarthritis and soft tissue lesions; pyrexia including post-immunisation pyrexia; acute migraine attack.

AVAILABILITY
TABLETS 500 and 650 mg Plain; 750 mg DT; SYRUPS/SUSPENSION 125 and 250 mg/5 ml;
INJECTION 2 ml ampoule 125 mg/ml.; Intravenous infusion 500 mg and 1g.

DOSE
Oral
Adult- 0.5 to 1g every 4 to 6 h (max. 4g, max 2g in alcoholics per day).
Child- for post-immunisation pyrexia, up to 2 months: 60 mg. 3 month to 1 year: 60 to 120 mg every 4 to 6 h. 1 to 5 years: 120 to 250 mg every 4 to 6 h. 6 to 12 years: 250 to 500 mg every 4 to 6 h.
Intramuscular injection
Adult- 250 mg every 4 to 6 h or as required.
Intravenous infusion
Adult- 1g every 6 hours, maximum daily dose 4 g.
Child- 15 mg/kg upto 4 times a day, maximum daily dose 60 mg/kg.

PRECAUTIONS
Hepatic impairment (Appendix 7a); renal impairment; alcohol dependence; lactation (Appendix 7b); pregnancy (Appendix 7c); overdosage: chapter 7.2; interactions (Appendix 6a); G-6-PD deficiency.

ADVERSE EFFECTS
Rare but rashes and blood disorders reported; important: liver damage (and less frequently renal damage) following overdosage; dyspepsia.

STORAGE
Store protected from light and moisture.

Propranolol
EDL-D 440 Secondary hospitals

Indications
Prophylaxis of migraine.
**Dose**

**Oral**

Initially 40 mg 2 to 3 times a day. Maintenance dose 80 to 160 mg daily.

Child- 2-4 mg/kg/day

**CONTRAINDICATIONS**

Asthma or history of obstructive airway disease; uncontrolled heart failure; Prinzmetal angina, marked bradycardia, hypotension; sick sinus syndrome, second- or third-degree atrioventricular block, cardiogenic shock, metabolic acidosis, severe peripheral arterial disease; pheochromocytoma.

**PRECAUTIONS**

First-degree atrioventricular block; renal impairment; liver disease; pregnancy (Appendix 7c); lactation (Appendix 7b); portal hypertension; diabetes mellitus; myasthenia gravis; history of hypersensitivity (increased reaction to allergens, also reduced response to epinephrine (adrenaline); interactions (Appendix 6a, 6b, 6d).

**ADVERSE EFFECTS**

Bradycardia, heart failure, hypotension, conduction disorders, bronchospasm, peripheral vasoconstriction, exacerbation of intermittent claudication and Raynaud phenomenon; gastrointestinal disturbances, fatigue, sleep disturbances including nightmares; rarely; rash, dry eyes (reversible); exacerbation of psoriasis.

**STORAGE**

Store protected from light and moisture.

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**Dihydroergotamine**

**EDL-D 441 Secondary hospitals**

**INDICATIONS**

Acute treatment of migraine headaches with or without aura and acute treatment of cluster headache episodes.

**AVAILABILITY**

TABLET 1 mg; INJECTION 1 ml Ampoule (1 mg/ml).

**DOSE**

Usually in combination with other analgesics e.g. caffeine.

Adult and child over 12 years- 1 to 2 tablets at onset (max. 4 tablets in 24 h), not to be repeated at intervals of less than 4 days.

**Intravenous infusion**

Termination of an acute attack of cluster headache, migraine:

Adult- 0.5 to 1 mg, 1 dose (Max: 3 mg/day or 6 mg/week).

**CONTRAINDICATIONS**

Peripheral vascular disease, coronary heart disease, obliterative vascular disease and Raynaud’s syndrome, temporal arteritis; hepatic impairment, renal impairment, sepsis; severe or inadequately controlled hypertension, hyperthyroidism, pregnancy (Appendix 7c); lactation; porphyria, ischaemic heart disease; angina pectoris.

**PRECAUTIONS**

Risk of peripheral vasospasm; elderly; it should not be used for migraine prophylaxis; interactions (Appendix 6c). Warn patient to stop treatment immediately if numbness or tingling of extremities develops and to contact doctor, compromised circulation; hypertension.
ADVERSE EFFECTS
Nausea, vomiting, vertigo, abdominal pain, diarrhoea, muscle cramps and occasionally headache provoked (usually because of prolonged excessive dosage or abrupt withdrawal); precordial pain, myocardial and intestinal ischaemia, rarely, myocardial infarction; repeated high dosage may cause ergotism with gangrene and confusion; pleural, peritoneal and heart-valve fibrosis may occur with excessive use; coronary artery vasospasm; ventricular tachycardia; altered sense of taste; rhinitis.

STORAGE
Store protected from light.

Sumatriptan

Non-EDL Secondary hospitals

Indications
Acute treatment of migraine.

AVAILABILITY
TABLET 25, 50 and 100 mg; INJECTION 0.5 ml ampoule (6 mg/ml).

DOSE
Oral
The recommended oral dose is 25-100 mg, repeatable after 2 hours up to a total dose of 200 mg over a 24 hour period.
Parenteral
6 mg at onset subcutaneously, may be repeated once after 1 h for maximum of 2 doses in 24 hours.

CONTRAINDICATIONS
Ischaemic heart disease, hypertension; pregnancy (Appendix 7c); renal impairment.

PRECAUTIONS
Ischaemic heart disease; hepatic impairment.

ADVERSE EFFECTS
Tightness in head and chest, paraesthesia in limbs, dizziness; rise in BP, bradycardia, sudden death, seizures.

FOR PROPHYLAXIS
Prophylactic treatment should be considered for patients in whom treatment of acute migraine attacks with analgesics or ergotamine is ineffective, or in whom attacks occur more than once a month, or for those with less frequent but severe or prolonged attacks. Prophylaxis can reduce the severity and frequency of attacks but does not eliminate them completely; additional symptomatic treatment is still needed. However, long-term prophylaxis is undesirable and treatment should be reviewed at 6-monthly intervals. Of the many drugs that have been advocated beta-adrenoceptor antagonists (betablockers) are most frequently used. Propranolol, a non-selective beta-blocker and other related compounds with similar profile such as atenolol are generally preferred. The potential for beta-blockers to interact with ergotamine should be borne in mind. Tricyclic antidepressants, such as amitriptyline or calcium-channel blocking drugs such as flunarizine or verapamil may be of value.
Flunarizine  
EDL-D 637 Secondary hospitals

INDICATIONS  
Prophylaxis of migraine.

AVAILABILITY  
TABLETS/CAPSULES 5 and 10 mg.

DOSE  
Oral  
Adults- 10 mg at night.  
Child < 40 kg- 5 mg at night.

CONTRAINDICATIONS  
Pregnancy (Appendix 7c); lactation.

PRECAUTIONS  
Patient may have drowsiness, should not operate hazardous machines.

ADVERSE EFFECTS  
Drowsiness; weight gain; depression; gastric pain, dry mouth; insomnia; extrapyramidal side effects.
Antineoplastics

Note: Who advises that adequate resources and specialist supervision are a prerequisite for the introduction of this class of drugs. Specific expertise, diagnostic precision, individualization of dosage or special equipment are required for their proper use.

The treatment of cancer with drugs, radiotherapy and surgery is complex and should only be undertaken by an oncologist. For this reason, the following information is provided merely as a guide. Chemotherapy may be curative or used to alleviate symptoms or to prolong life. Where the condition can no longer be managed with cytotoxic therapy, alternative palliative treatment should be considered.

For some tumours, single-drug chemotherapy may be adequate, but for many malignancies a combination of drugs provides the best response. Examples of combination therapy include:

- ‘CHOP’ (cyclophosphamide, doxorubicin, vincristine, prednisolone) for non-Hodgkin’s disease;
- ‘ABVD’ (doxorubicin, bleomycin, vinblastine, dacarbazine) for Hodgkin’s disease;
- ‘MOPP’ (chlormethine, vincristine, procarbazine, prednisolone) for Hodgkin’s disease.

Cytotoxic drugs are often combined with other classes of drugs in the treatment of malignant conditions. Such drugs include hormone agonists and antagonists, corticosteroids and immunostimulant drugs. Combinations are, however, more toxic than single drugs.

Precautions and Contraindications

Treatment with cytotoxic drugs should be initiated only after baseline tests of liver and kidney function have been performed and baseline blood counts established. It may be necessary to modify or delay treatment in certain circumstances. The patient should also be monitored regularly during chemotherapy and cytotoxic drugs withheld if there is significant deterioration in bone-marrow, liver or kidney function.

Many cytotoxic drugs are teratogenic and should not be administered during pregnancy especially in the first trimester. Contraceptive measures are required during therapy and possibly for a period after therapy has ended. Cytotoxic drugs are also contraindicated during lactation.

Cytotoxic drugs should be administered with care to avoid undue toxicity to the patient or exposure during handling by the health care provider. All waste, including patient’s body fluids and excreta (and any material contaminated by them) should be treated as hazardous.
Extravasation of intravenously administered cytotoxic drugs can result in severe pain and necrosis of surrounding tissue. If extravasation occurs, aspiration of the drug should first be attempted, then the affected limb is elevated and warm compresses applied to speed and dilute the infusion or it is localized by applying cold compresses until the inflammation subsides; in severe cases, hydrocortisone cream may be applied topically to the site of inflammation. The manufacturer’s literature should also be consulted for more specific information.

**Adverse Effects**

Cytotoxic drugs have a considerable potential to damage normal tissue. Specific adverse effects apply, but a number of effects are common to all cytotoxics such as bone-marrow and immunological suppression. Furthermore, the concomitant use of immunosuppressive drugs will enhance susceptibility to infections. Fever associated with neutropenia or immunosuppression requires immediate treatment with antibiotics.

Nausea and vomiting: Nausea and vomiting following administration of cytotoxic drugs and abdominal radiotherapy are often distressing and may compromise further treatment. Symptoms may be acute (occurring within 24 h of treatment), delayed (first occurring more than 24 h after treatment), or anticipatory (occurring before subsequent doses). Delayed and anticipatory symptoms are more difficult to control than acute symptoms and require different management.

Cytotoxic drugs associated with a low risk of emesis include etoposide, 5-fluorouracil, low-dose methotrexate and the vinca alkaloids; those with an intermediate risk include low-dose cyclophosphamide, doxorubicin and high-dose methotrexate; and the highest risk is with cisplatin, high-dose cyclophosphamide and dacarbazine.

For patients at a low risk of emesis, pretreatment with an oral phenothiazine (for example chlorpromazine), continued for up to 24 h after chemotherapy, is often helpful. For patients at a higher risk dexamethasone 6-10 mg by mouth may be added before chemotherapy. For patients at a high risk of emesis or when other therapies are ineffective, high doses of intravenous metoclopramide may be used.

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**Note:** High doses of metoclopramide are preferably given by continuous intravenous infusion: an initial dose of 2-4 mg/kg is given over 15 to 20 min, followed by a maintenance dose of 3-5 mg/kg over 8 to 12 h; the total dose should not exceed 10 mg/kg in 24 h.

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Dexamethasone is the drug of choice for the prevention of delayed symptoms; it is used alone or with metoclopramide. Good symptom control is the best way to prevent anticipatory symptoms and the addition of diazepam to antiemetic therapy is helpful because of its sedative, anxiolytic and amnesic effects.

Hyperuricaemia: Hyperuricaemia may complicate treatment of conditions such as non-Hodgkin’s lymphomas and leukaemia. Renal damage may result from the formation of uric acid crystals. Patients should be adequately hydrated and hyperuricaemia may be managed with allopurinol initiated 24 h before cytotoxic treatment and continued for 7 to 10 days afterwards.
Alopecia: Alopecia is common during treatment with cytotoxic drugs. There is no drug treatment, but the condition often reverses spontaneously once treatment has stopped.

**Alkylating Drugs:**
Alkylating drugs are among the most widely used drugs in cancer chemotherapy. They act by damaging DNA and therefore interfering with cell replication. However, there are two complications. Firstly, they affect gametogenesis and may cause permanent male sterility; in women, the reproductive span may be shortened by the onset of a premature menopause. Secondly, they are associated with a marked increase in the incidence of acute non-lymphocytic leukaemia, in particular when combined with extensive radiation therapy.

Cyclophosphamide requires hepatic activation; it can therefore be given orally and is not vesicant when given intravenously. Like all alkylating drugs its major toxic effects are myelosuppression, alopecia, nausea and vomiting. It can also cause haemorrhagic cystitis; an increased fluid intake for 24 to 48 h will help to avoid this complication. Cyclophosphamide is used either as part of treatment or as an adjuvant in Non- Hodgkin’s lymphomas, breast cancer, childhood leukaemia and ovarian cancer. It is also used in several palliative regimens.

Chlorambucil is used to treat chronic lymphocytic leukaemia, Non-Hodgkin’s lymphomas, Hodgkin’s disease, ovarian cancer and Waldenstrom (primary) macroglobulinaemia. Adverse effects, apart from bone marrow suppression, are uncommon. However, severe widespread rash can develop and may progress to Stevens-Johnson syndrome or toxic epidermal necrolysis. If a rash occurs, further treatment with chlorambucil is contraindicated.

Chlormethine (mustine) forms part of the regimen for treatment of advanced Hodgkin’s disease and malignant lymphomas. Its toxicity includes myelosuppression, severe nausea and vomiting, alopecia and thrombophlebitis due to vesicant effect.

**Cytotoxic Antibiotics:**
Bleomycin is used in regimens for the treatment of Hodgkin’s disease and testicular cancer. It has several antineoplastic drug toxicities; it is known to cause dose-related pneumonitis and fibrosis which can be fatal and is associated with rare acute hypersensitivity reactions. Cutaneous toxicity has also been reported.

Doxorubicin is the most widely used anthracycline antibiotic. It is used for acute leukaemias although other anthracyclines are more commonly used in these circumstances. Doxorubicin also plays a palliative role in the treatment of other malignancies. The primary toxic effects are myelosuppression, alopecia, nausea, vomiting and dose-related cardiomyopathy. It is also vesicant and can cause severe skin ulceration on extravasation. Daunorubicin is used in acute leukaemias. Its toxicity is similar to that of doxorubicin, but it is not cardiotoxic. Daunorubicin is used in acute leukaemias. Its toxicity is similar to that of doxorubicin.

**Antimetabolites and Related Therapy:**
Cytarabine is used in the treatment of acute leukaemia; children may tolerate high doses better than adults. Its effects are highly dependent upon the schedule of administration. It causes myelosuppression, mucositis and in high doses, central neurotoxicity.
5-Fluorouracil is primarily used in the adjuvant treatment of colorectal and breast cancer. It is also employed in the palliative treatment of other malignancies. It causes myelosuppression and the palmar-plantar syndrome (erythema and painful desquamation of the hands and feet). When its action is modified by other drugs (such as calcium folinate), its toxicity profile can change; mucositis and diarrhoea may be significant problems. Central neurotoxicity can also occur.

6-Mercaptopurine is frequently used in the therapy of childhood leukaemia. It can be administered orally and myelosuppression and nausea are the only important toxic effects. Methotrexate is used to treat a variety of malignancies and it plays a major role as an adjuvant for the treatment of breast cancer. Like 5-fluorouracil, methotrexate is myelotoxic, but nausea and vomiting are minimal. It also causes mucositis. Renal impairment reduces methotrexate excretion and can exacerbate toxicity.

Calcium folinate is used to counteract the folate-antagonist action of methotrexate and thus speeds recovery from methotrexate-induced mucositis or myelosuppression. Calcium folinate also enhances the effects of 5-fluorouracil when the two are used together for metastatic colorectal cancer.

Vinca Alkaloids and Etoposide:

The vinca alkaloids, vinblastine and vincristine, are primarily used in the treatment of acute leukaemias. Vinblastine is also used for Hodgkin’s disease and some solid tumours. Vincristine is also used in the management of Non-Hodgkin’s lymphomas. Both can cause neurotoxicity, but this is more of a problem with vincristine. Myelosuppression is more common with vinblastine.

Etoposide is an important component of the treatment of testicular carcinoma and is also used in several regimens for lung cancers and lymphomas. It causes myelosuppression and alopecia and it can cause hypotension during infusion. It does not produce significant nausea and vomiting.

Other Antineoplastic Drugs:

The enzyme asparaginase is an important component in the management of childhood leukaemia, but is not used in any other malignancy. Its toxicity profile is broad and the drug must be carefully administered because of the risk of anaphylaxis.

Cisplatin is a platinum compound used in the treatment of ovarian and testicular malignancies. It is also a component of regimens used in non-small cell and small cell lung cancer and plays a palliative role in other malignancies. Cisplatin is myelosuppressive and also produces slight alopecia. However, it causes severe dose-related nausea and vomiting. It is also nephrotoxic and neurotoxic. Nephrotoxicity can be reduced by maintaining high urine output during cisplatin administration and immediately afterwards, but neurotoxicity is often dose-limiting.

Dacarbazine, thought to act as an alkylating drug, is a component of a regimen for Hodgkin’s disease. It is also used in the palliative therapy of metastatic malignant melanoma. Its major toxic effects are myelosuppression and intense nausea and vomiting.

Levamisole is an anthelmintic with immunostimulating properties; it is used in combination with 5-fluorouracil as adjuvant therapy for colorectal cancer following
resection of the tumour. Its major toxic effects are a variety of CNS symptoms, nausea, dermatitis and hypersensitivity reactions. **Procarbazine** is used in the treatment of advanced Hodgkin’s disease. Toxic effects include myelosuppression, nausea, vomiting, CNS symptoms and depression. Procarbazine possesses a weak monoamine oxidase inhibitory effect but dietary restriction is not necessary.

**Immunosuppressives**

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*Note: who advises that this class of drugs is for use only when adequate resources and specialist care are available. Specific expertise, diagnostic precision, individualization of dosage or special equipment are required for their proper use.*

Immunosuppressive drugs are used in organ transplant recipients to suppress rejection; they are also used as second-line drugs in chronic inflammatory conditions. Treatment should only be initiated by a specialist. Careful monitoring of blood counts is required in patients receiving immunosuppressive drugs and the dose should be adjusted to prevent bone marrow toxicity. Immunosuppressed patients are particularly prone to atypical infections.

**Azathioprine** is the most widely used drug in transplant recipients. It is useful when corticosteroid therapy alone has proven inadequate or for other conditions when a reduction in the dose of concurrently administered corticosteroids is required. It is metabolized to 6-mercaptopurine and, as with mercaptopurine, doses need to be reduced when given with allopurinol. The predominant toxic effect is myelosuppression, although hepatic toxicity also occurs.

**Cyclosporine** is a potent immunosuppressant which is virtually free of myelotoxic effects, but is markedly nephrotoxic. It is particularly useful for the prevention of graft rejection and for the prophylaxis of graft-versus-host disease. The dose is adjusted according to plasma-cyclosporine concentrations and renal function. Dose-related increases in serum creatinine and blood urea nitrogen (BUN) during the first few weeks may necessitate dose reduction. Corticosteroids such as prednisolone have significant immunosuppressant activity and can also be used to prevent rejection of organ transplants.

**Melphalan**

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**EDL-328 D 328 Tertiary restricted**

**AVAILABILITY**

TABLETS 2 and 5 mg; INJECTION 50 mg/vial.

**DOSE**

Oral Adult- Multiple myeloma: usual dose 6 mg/day. Maintenance dose 2 mg/day. Alternatively 10 mg daily for 7 days (total dose 70 mg), repeat if required after blood counts particularly neutrophils and platelets. Ovarian carcinoma: 0.2 mg/kg body weight daily for 5 days, repeat after 4 to 5 weeks. Child- 0.15 mg/kg body weight daily for 7 days. Maintenance dose is 0.05 mg/kg body weight daily when platelet count is rising. Intravenous injection
INDICATION
Breast carcinoma, multiple myeloma, advanced ovarian carcinoma, malignant melanoma, polycythaemia vera.

CONTRAINDICATION
Pregnancy; hypersensitivity; myelosuppression; lactation.

PRECAUTION
Hepatic impairment; renal impairment; interactions (Appendix 6d).

ADVERSE EFFECTS
Nausea, vomiting, oral mucositis, hyperuricaemia, bone marrow suppression, alopecia, thromboembolism, leucopenia; menstrual irregularities; haemolytic anaemia.

Mercaptopurine

EDL –D 332 Tertiary restricted

AVAILABILITY
INJECTION 2 ml ampoule (200 mg/2 ml) TABLET 50 mg.

DOSE
Oral Leukaemia in children (maintenance): 2.5 mg/ kg body weight in continuation with other drugs daily.

INDICATION
Acute leukaemias; Chronic granulocytic leukaemia; choreocarcinoma.

CONTRAINDICATION
hypersensitivity; pregnancy and lactation.

PRECAUTION
See notes above and consult literature; monitor blood count; uric acid levels; renal impairment and hepatic impairment (Appendix 7a) ; interactions(Appendix 6c)

ADVERSE EFFECTS
Hepatotoxicity; anorexia; nausea; hyperuricaemia; ulcers.

Chlorambucil

EDL- D 107 Tertiary restricted

AVAILABILITY
TABLETS 2 and 5 mg.

DOSE
Adult- Chronic lymphocytic leukaemia: initially 150 μg/kg body weight daily until leucocyte count sufficiently reduced. Maintenance (started 4 weeks after first course) 100 μg/kg body weight. Waldarstrom’s macroglobulinaemia: 6 to 12 mg daily until leucopenia occurs, then reduce to 2 to 8 mg daily. Child- Not recommended.

INDICATION
Chronic lymphocytic leukaemia; some non- Hodgkin’s lymphomas; Hodgkin’s disease, ovarian cancer and Waldenstrom (primary) macroglobulinaemia.

CONTRAINDICATION
hypersensitivity; porphyria; pregnancy and lactation.

PRECAUTION
See notes above and consult literature; renal impairment; hepatic impairment (Appendix 7a).

ADVERSE EFFECTS
Hepatotoxicity; peripheral neuropathy; cystitis; seizures; pulmonary fibrosis.
Cisplatin

**EDL-D 127,586 Tertiary restricted**

**AVAILABILITY**
INJECTION 10 ml (10 mg) and 50 ml (50 mg) vials.

**DOSE**
Intravenous injection (use syringes devoid of aluminium component) Ovarian tumor: 50 mg/m² of body surface area once every three weeks. Bladder cancer: 50 to 70 mg/m² once every 3 to 4 weeks. Testicular tumor: 20 mg/m² for 5 days every 3 weeks for 3 courses.

**INDICATION**
Metastatic testicular tumours, metastatic ovarian tumours, advanced bladder carcinoma and other solid tumours.

**CONTRAINDICATION**
See notes above and consult literature; hypersensitivity; renal impairment; pregnancy and lactation

**PRECAUTION**
See notes above and consult literature; hyperuraemia; hypomagnesaemia; hypocalcaemia; interactions (Appendix 6c).

**ADVERSE EFFECTS**
Tinnitus; neuropathy.

Cyclophosphamide

**EDL-D 144 Tertiary restricted**

**AVAILABILITY**
TABLET 50 mg; INJECTION 15 ml (200 mg), 30 ml (500 mg) and 50 ml (1g) vials; dry powder to be reconstituted before administration.

**DOSE**
Intravenous injection Malignancy: 40 to 50 mg/kg body weight in divided doses over 2 to 5 days. Alternatively 10 to 15 mg/kg body weight every 7 to 10 days or 3 to 5 mg/kg body weight twice a week. Oral 1 to 5 mg/kg body weight. Minimal change nephrotic syndrome: 2.5 to 3 mg/kg body weight.

**INDICATION**
Malignant lymphomas including Non- Hodgkin’s lymphomas, lymphocytic lymphoma, Burkitt’s lymphoma; multiple myeloma; leukaemias, mycosis fungoides; neuroblastoma; adenocarcinoma of the ovary; retinoblastoma; breast cancer.

**CONTRAINDICATION**
bladder haemorrhage; thrombocytopenia; severe bone marrow depression; pregnancy and lactation

**PRECAUTION**
See notes above and consult literature; renal impairment (Appendix 7d), hepatic impairment (Appendix 7a), interaction (Appendix 7c).

**ADVERSE EFFECTS**
Haemorrhagic cystitis; colitis; cardiac toxicity; anorexia; thrombocytopenia; dermatitis.

Mitomycin

**EDL-D 355 Tertiary restricted**

**AVAILABILITY**
INJECTIONS vial 2 and 10 mg (dry powder to be reconstituted before administration).
DOSE
Intravenous injection Adult- 6 to 10 mg twice a week, alternatively 0.5 mg/kg body weight daily for 5 days, repeat after 2 weeks.

INDICATION
Adrenocarcinoma, lymphosarcoma and seminoma, superficial bladder cancer (adjuvant therapy).

CONTRAINDICATION
Pregnancy; bone marrow depression; severe anaemia; thrombocytopenia; lactation.

PRECAUTION
It causes delayed bone-marrow toxicity and therefore it is usually administered at 6-weekly intervals. Caution in handling because it is irritant to tissues, thrombocytopenia; necrosis; leucopenia.

ADVERSE EFFECTS
Prolonged use may result in permanent bone-marrow damage. It may also cause lung fibrosis and renal damage; dyspnea.

Monteleukast

Non-EDL Tertiary restricted

AVAILABILITY
Oral Adult- 10 mg once a day. Child- 2-5yrs: 4 mg once daily; 6-14 yrs: 5 mg once daily; ≥ 15 yrs: 10 mg once daily.

DOSE
TABLETS 5 and 10 mg.

INDICATION
Prophylaxis of mild to moderate asthma

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
History of liver disease, pregnancy (Appendix 7c).

ADVERSE EFFECTS
Headache; rashes; eosinophilia; neuropathy; Churg-strauss syndrome.

Procarbazine

EDL-D 434 Tertiary restricted

AVAILABILITY
TABLETS 50 mg.

DOSE
Oral 50 mg daily to start with initially, increased to 250 to 300 mg individual doses. Maintenance (on remission): 50 to 100 mg daily to cumulative total of at least 6g.

INDICATION
Part of MOPP regimen in Hodgkin’s and Non- Hodgkin’s lymphomas.

CONTRAINDICATION
Hypersensitivity; pregnancy(Appendix 7c) and lactation(Appendix 7b).

PRECAUTION
See notes above and consult literature; ulceration; haemorrhage; leucopenia: renal and hepatic impairment(Appendix 7a).; interactions(Appendix 6a).

ADVERSE EFFECTS
Leucopenia; anaemia; thrombocytopenia; hypotension; retinal haemorrhage.
Busulfan

**EDL-D83 Tertiary restricted**

**AVAILABILITY**
- TABLET 2 mg.

**DOSE**
- Oral Chronic myeloid leukaemia, induction of remission: 60 μg/kg body weight daily (max 4 mg) maintenance dose 0.5 to 2 mg daily

**INDICATION**
- Chronic granulocytic leukaemia, chronic myelogenous leukaemia, polycythaemia vera, myelofibrosis, thrombocytopenia.

**CONTRAINDICATION**
- Pregnancy (Appendix 7c) bone marrow suppression; chronic lymphocytic leukaemia; lactation.

**PRECAUTION**
- Monitor cardiac function; pregnancy; lactation previous radiation therapy; avoid in porphyria, hepatic impairment; interactions (Appendix 6c).

**ADVERSE EFFECTS**
- Hepatotoxicity (including hepatic venoocclusive disease, hyperbilirubinaemia, jaundice and fibrosis); cardiac tamponad at high doses in thalassaeic patients; pneumonia; skin hyperpigmentation; hyperuricaemia; pulmonary fibrosis.

Cytarabine

**EDL-D 145 Tertiary restricted**

**AVAILABILITY**
- INJECTION vials 1 ml (100 mg), 5 ml (500 mg) and 10 ml (1g).

**DOSE**
- Intravenous injection Adult- 100 mg/m2 body surface area every 12 h for seven days. Child- 100 mg/m2 body surface area twice daily by rapid injection or 100 mg/m2 body surface area daily by continuous infusion given by 5 to 10 days

**INDICATION**
- Acute lymphoblastic leukaemia; chronic myeloid leukaemia; meningeal leukaemia; erythroleukaemia; Non-Hodgkin’s lymphomas; lymphosarcoma.

**CONTRAINDICATION**
- hypersensitivity; pregnancy and lactation

**PRECAUTION**
- See notes above and consult literature; uric acid level monitoring recommended; hepatic impairment

**ADVERSE EFFECTS**
- GIT Disturbances

Florouracil

**EDL-D224 Tertiary restricted**

**AVAILABILITY**
- INJECTION 5 and 10 ml ampoule (50 mg/ml). TABLETS 50 mg.

**DOSE**
- Intravenous injection Initially 12 mg/kg body weight once a day for 4 days, max. daily dose 800 mg. If tolerated well without toxicity 6 mg/kg body weight can be given on 6th, 8th, 10th and 12th day. Discontinue on 12th day. Maintenance dose 10 to 15 mg/kg body weight every week (max dose 1g/week).
INDICATION
Carcinomas of the colorectum, breast, stomach, pancreas, cervix, prostate, ovary and endometrium; liver tumours; head and neck tumours; actinic keratosis

CONTRAINDICATION
See notes above and consult literature; bone marrow depression; pregnancy (Appendix 7c) and lactation (Appendix 7b).

PRECAUTION
See notes above and consult literature; pelvic irradiation; renal impairment; hepatic impairment (Appendix 7a); interactions (Appendix 6c).

ADVERSE EFFECTS
Cardiac toxicity; tachycardia; dermatitis; diarrhoea.

Bleomycin

EDL-D 79 Tertiary restricted

AVAILABILITY
INJECTION 15 and 30 mg/vial.

DOSE
Intramuscular and subcutaneous injection 30 mg twice a week, dose can also vary from 15 mg daily to 15 mg weekly; total 300 to 400 mg. Small cell cancer; 0.25 to 0.5 mg/kg body weight once or twice a week.

INDICATION
Adjunct to surgery and radiotherapy in palliative treatment of Hodgkin’s and non-Hodgkin’s lymphomas; reticulum cell sarcoma and lymphoma; carcinomas of the head, neck, larynx, cervix, penis, skin, vulva, testicles including embryonal cell carcinoma, choriocarcinoma and teratoma; malignant effusions.

CONTRAINDICATION
See notes above and literature; preexisting lung disease; pregnancy and lactation

PRECAUTION
renal impairment (Appendix 6c).

ADVERSE EFFECTS
Dermatitis; nephrotoxicity; hepatotoxicity.

Doxorubicin

EDL-D 191,616,617 Tertiary restricted

AVAILABILITY
INJECTION 10 & 50 mg lyophilized powder/vial, 2 mg/ml solution LIPOSOMAL injection 10 ml vial (2 mg/ml).

DOSE
Intravenous 50-75 mg/m2 body surface area by slow i.v injection every 3 weeks. AIDS-related Kaposi's sarcoma: Adult: As pegylated liposome: 20 mg/m2 body surface area infused over 1 hr once every 3 weeks. Ovarian carcinoma: Adult: As pegylated liposome: 50 mg/m2 BSA infused over 1 hr once every 4 weeks.

INDICATION

CONTRAINDICATION
Known hypersensitivity, cardiac disease, pregnancy (Appendix 7c), lactation, neonates.
PRECAUTION
Avoid extravasation, monitor ECG changes, arrhythmias, blood counts, hypotension or congestive heart failure, hepatic impairment, interactions (Appendix 6c), Liposomal and non-liposomal preparations are not interchangeable.

ADVERSE EFFECTS
Infusion reactions, cardiotoxicity, bone marrow suppression, liver impairment, nausea and vomiting, reversible alopecia, stomatitis, conjunctivitis, keratitis, mucositis, discoloration of body fluids, local skin reactions and tissue damage, secondary leukemias.

Dactinomycin

EDL-D 146 Tertiary restricted

AVAILABILITY
INJECTION Vial 500 mg.

DOSE
Intravenous injection Adult and child above 6 months- 15 μg/kg/ day. Principally used to treat paediatric cancers.

INDICATION
Trophoblastic tumours, Wilm’s tumour, Ewing’s sarcoma, rhabdomyosarcoma.

CONTRAINDICATION
hypersensitivity; lactation; infection with children; herpes zoster; pregnancy (Appendix 7c) and lactation.

PRECAUTION
See notes above and consult literature.

ADVERSE EFFECTS
Hair loss; nausea; vomiting; mouth sores; diarrhoea.

NOTE: Irritant to tissues

Cyclophosphamide

EDL-D 595 Tertiary restricted

INDICATIONS
Malignant lymphomas including Non- Hodgkin’s lymphomas, lymphocytic lymphoma, Burkitt’s lymphoma; multiple myeloma; leukaemias, mycosis fungoides; neuroblastoma; adenocarcinoma of the ovary; retinoblastoma; breast cancer.

AVAILABILITY
TABLET 50 mg; INJECTION 15 ml (200 mg), 30 ml (500 mg) and 50 ml (1g) vials; dry powder to be reconstituted before administration.

DOSE
Intravenous injection
Malignancy: 40 to 50 mg/kg body weight in divided doses over 2 to 5 days. Alternatively 10 to 15 mg/kg body weight every 7 to 10 days or 3 to 5 mg/kg body weight twice a week.
Oral
1 to 5 mg/kg body weight. Minimal change nephrotic syndrome: 2.5 to 3 mg/kg body weight.

CONTRAINDICATIONS
See notes above and consult literature; bladder haemorrhage; thrombocytopenia; severe bone marrow depression; pregnancy (Appendix 7c) and lactation (Appendix 7b).

PRECAUTIONS
See notes above and consult literature; renal impairment (Appendix 7d) and hepatic impairment (Appendix 7a); interactions (Appendix 6c).
ADVERSE EFFECTS
See notes above and consult literature. Haemorrhagic cystitis; colitis; cardiac toxicity; anorexia; thrombocytopenia; dermatitis.

Storage
Injection: Store in refrigerator (2 to 8°C). Avoid long exposure to temperature above 30°C. The solution should be used immediately after preparation as it deteriorates on storage. Tablet: Store at a temperature not exceeding 30°C.

Cytosine Arabinoside (Cytarabine)

EDL-D 597 Tertiary restricted

INDICATIONS
Acute lymphoblastic leukaemia; chronic myeloid leukaemia; meningeal leukaemia; erythroleukaemia; Non-Hodgkin’s lymphomas; lymphosarcoma.

AVAILABILITY
INJECTION vials 1 ml (100 mg), 5 ml (500 mg) and 10 ml (1g).

DOSE
Intravenous injection
Adult- 100 mg/m² body surface area every 12 h for seven days.
Child- 100 mg/m² body surface area twice daily by rapid injection or 100 mg/m² body surface area daily by continuous infusion given by 5 to 10 days.

CONTRAINDICATIONS
See notes above and consult literature; hypersensitivity; pregnancy (Appendix 7c) and lactation (Appendix 7b).

PRECAUTIONS
See notes above and consult literature; uric acid level monitoring recommended; hepatic impairment (Appendix 7a).

ADVERSE EFFECTS
See notes above and consult literature; g.i.t. disturbances.

STORAGE
Store protected from light.

Erythropoietin

EDL-D 624 Tertiary restricted

Indications

AVAILABILITY
INJECTIONS 1000, 2000, 3000, 4000, 5000, 6000, 10000, 20000 and 40000 IU/Vial

DOSE
Parenteral
Anaemia of chronic renal failure
Adult: As epoetin alfa: Initially, 50 U/kg subcutaneous/intravenous 3 times weekly for predialysis and haemodialysis patients and 50 U/kg twice weekly for peritoneal dialysis patients, dose may be increased according to response in steps of 25 U/kg 3 times weekly at 4 weekly intervals.
Child: As epoetin alfa: Initially, 50 U/kg 3 times weekly. Dose may be increased at 4 weekly intervals in increments of 25 U/kg 3 times weekly until a target haemoglobin concentration of 9.5-11 g/100 ml is reached.
Usual maintenance dose: <10 kg: 225-450 U/kg/week; 10-30 kg: 180-450 U/kg/week and >30 kg: 90-300 U/kg/week.

Anaemia in zidovudine-treated HIV-infected patients
Adult: As epoetin alfa: Initially, 100 U/kg subcutaneous/intravenous thrice weekly for 8 weeks; increase every 4-8 week by 50-100 U/kg according to response. Max: 300 U/kg thrice weekly.

Subcutaneous

Anaemia related to non-myeloid malignant disease chemotherapy
Adult: As epoetin alfa or zeta: Initially, 150 U/kg 3 times weekly. Dose may be increased at 4-8 week intervals to 300 U/kg 3 times weekly. Stop treatment if response is still inadequate after 4 week of treatment using this higher dose.

Intravenous
Increase yield of autologous blood
Adult: As epoetin alfa or zeta: 600 U/kg over 2 minutes twice weekly for 3 week before surgery; in conjunction with iron, folate and B12 supplementation.

Contraindications
Hypersensitivity to mammalian cell products and human albumin, uncontrolled hypertension.

PRECAUTIONS
Ischaemic heart diseases, chronic renal failure, hypertension, seizures, liver dysfunction, pregnancy (Appendix 7c) and lactation, interactions (Appendix 6c).

ADVERSE EFFECTS
Nausea, vomiting, increased risk of hypertension, myalgia, arthralgia, rashes and urticaria, headache, confusion, generalized seizures, thrombosis specifically during dialysis, fever, diarrhoea, tissue swelling, flulike syndrome, paraesthesia, constipation, nasal or chest congestion, immunogenicity leading to Pure Red Cell Aplasia.

STORAGE
Store in an air tight container at a temperature below -20°C. Avoid repeated freezing and thawing.

L-Asparaginase

EDL-D 678 Tertiary restricted

INDICATIONS
Acute lymphoblastic leukaemia.

AVAILABILITY
INJECTION 5,000, 6,000 and 10,000 IU (for reconstitution before administration).

DOSE
Intramuscular, intravenous or subcutaneous injection
Exclusively in acute lymphoblastic leukaemia. Careful monitoring is required. Urine is tested for glucose because of risk of hyperglycaemia.

CONTRAINDICATIONS
See notes above and consult literature; pregnancy (Appendix 7c) and lactation (Appendix 7b).

PRECAUTIONS
See notes above and consult literature.

ADVERSE EFFECTS
See notes above and consult literature.
Etoposide

EDL-D 213,214 Tertiary restricted

AVAILABILITY
CAPSULES 25, 50 and 100 mg; INJECTION vial 100 mg/5 ml.

DOSE
Intramuscular injection Adult- Initially 50 to 100 mg/m² body surface area daily by infusing over 30 to 60 min. Thereafter, no injection for 3 to 4 weeks is given. Small cell lung cancer: 350 mg/m² daily. Oral Adult- 100 to 200 mg/m² body surface area from day 1 to 5 taken on empty stomach, thereafter no treatment for 3 to 4 weeks.

INDICATION
Refractory testicular tumours; acute leukaemia; malignant lymphoma; lung cancer

CONTRAINDICATION
See notes above and consult literature; hypersensitivity; severe liver dysfunction; pregnancy (Appendix 7c) and lactation (Appendix 7b).

PRECAUTION
hepatic impairment (Appendix 7a); interactions (Appendix 6c); renal impairment (Appendix 7d).

ADVERSE EFFECTS
Alopecia; gastrointestinal disturbances; thrombophlebitis; neuritis

Vinblastine

EDL-D 522 Tertiary restricted

AVAILABILITY
VIAL 10 ml (1 mg/ml).

DOSE
Intravenous injection only 3.7 mg/m² body surface surface area in single dose. Increase on weekly intervals depending on WBC count (max 18.5 mg/m² body area).

INDICATION
Disseminated Hodgkin’s and Non-Hodgkin’s lymphomas; advanced testicular carcinoma, breast carcinoma; palliative treatment of Kaposi’s sarcoma; trophoblastic tumours; Letterer-Siwe disease; Histolytic lymphoma.

CONTRAINDICATION
hypersensitivity; severe granulocytopenia; lactation (Appendix 7b). Intrathecal injection is contraindicated.

PRECAUTION
neurotoxicity; ischaemic heart disease; hepatic impairment (Appendix 7a); interactions (Appendix 6c); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Stomatitis; leucopenia; constipation; bone pain.

Vincristine

EDL-D 523 Tertiary restricted

AVAILABILITY
VIAL 10 ml (1 mg/ml).

DOSE
Intravenous injection only 3.7 mg/m² body surface surface area in single dose. Increase on weekly intervals depending on WBC count (max 18.5 mg/m² body area).
INDICATION
Disseminated Hodgkin’s and Non-Hodgkin’s lymphomas; advanced testicular carcinoma, breast carcinoma; palliative treatment of Kaposi’s sarcoma; trophoblastic tumours; Letterer-Siwe disease; Histolytic lymphoma.

CONTRAINDICATIONS
lactation (Appendix 7b).
Intrathecal injection is contraindicated.

PRECAUTIONS
See notes above and consult literature; uric acid neuropathy; branchospasm; hepatic impairment (Appendix 7a); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Stomatitis; leucopenia; constipation; bone pain.

Fluorouracil (5FU) IP

EDL-D 639, 640 Tertiary restricted

AVAILABILITY
INJECTION 5 and 10 ml ampoule (50 mg/ml). TABLETS 50 mg.

DOSE
Intravenous injection Initially 12 mg/kg body weight once a day for 4 days, max. daily dose 800 mg. If tolerated well without toxicity 6 mg/kg body weight can be given on 6th, 8th, 10th and 12th day. Discontinue on 12th day. Maintenance dose 10 to 15 mg/kg body weight every week (max dose 1g/week).

INDICATION
Carcinomas of the colorectum, breast, stomach, pancreas, cervix, prostate, ovary and endometrium; liver tumours; head and neck tumours; actinic keratosis.

CONTRAINDICATIONS
See notes above and consult literature; bone marrow depression; pregnancy (Appendix 7c) and lactation (Appendix 7b).

PRECAUTIONS
See notes above and consult literature; lactation; pelvic irradiation; renal impairment; hepatic impairment (Appendix 7a); interactions (Appendix 6c).

ADVERSE EFFECTS
Cardiac toxicity; tachycardia; dermatitis; diarrhoea.

Gemcitabine IP

EDL-D tertiary restricted 649,650

AVAILABILITY
INJECTION Vial 200 mg and 1g (dry powder to be reconstituted before administration).

DOSE
1g/m2 body surface area for over 30 min once a week for up to 7 weeks, if not tolerated reduce or withhold. After one week rest administer by infusion once weekly for three weeks, withhold for 4th week before repeating.

INDICATION
Adenocarcinoma of pancreas

CONTRAINDICATION
Pregnancy (Appendix 7C) concurrent radial radiotherapy; hypersensitivity; lactation.

PRECAUTION
Gemcitabine is not recommended for patients who can have potentially curative surgery. There is insufficient evidence about its use for second-line treatment of pancreatic adenocarcinoma, hepatic impairment; renal impairment, interactions (Appendix 6c)
ADVERSE EFFECTS
Nausea, vomiting, oral mucositis, hyperuricaemia, bone marrow suppression, alopecia, thromboembolism, flu like syndrome; edema; thrombocytopenia; somnolence; hematuria; dyspnoea; loss of appetite.

Methotrexate IP (preservative free)

EDL-D 688,686 Tertiary restricted

AVAILABILITY
TABLETS 2.5, 5.0 and 7.5 mg; INJECTION vial/ampoule 25 mg/ml and 100 mg/ml.

DOSE
Oral Severe active rheumatoid arthritis: 7.5 mg once weekly, adjusted according to response (max. weekly dose 20 mg). Intramuscular, subcutaneous or intravenous route in severe attack under expert medical supervision at a dose of 7.5 mg once weekly.

INDICATION
Rheumatoid arthritis which has failed to respond to penicillamine or chloroquine; malignant disease.

CONTRAINDICATION
Blood disorders (bone marrow suppression); liver damage; pulmonary toxicity; gastrointestinal disturbances-if stomatitis and diarrhoea occur; stop treatment; renal failure; skin reactions; alopecia; osteoporosis; arthralgia; myalgia; ocular irritation; precipitation of diabetes.

PRECAUTION
Monitor throughout treatment including blood counts and hepatic and renal function tests; renal and hepatic impairment (avoid if severe; see also Appendices 7a); reduce dose or withdraw if acute infection develops; for woman or man; during contraception and for at least 6 months after treatment; peptic ulceration; ulcerative colitis; diarrhoea; ulcerative stomatitis; advise patient to avoid self-medication with salicylates or other NSAIDs; warn patient with rheumatoid arthritis to report cough or dyspnoea; interactions (Appendix 6a, 6c, 6d). Patients should be warned to report immediately any signs or symptoms of bone marrow suppression; for example unexplained bruising or bleeding; purpura; infection; sore throat

ADVERSE EFFECTS
Blood disorders (bone marrow suppression); liver damage; pulmonary toxicity; gastrointestinal disturbances-if stomatitis and diarrhoea occur; stop treatment; renal failure; skin reactions; alopecia; osteoporosis; arthralgia; myalgia; ocular irritation; precipitation of diabetes.

Paclitaxel IP

EDL-D 710 tertiary restricted

AVAILABILITY
injection vial 30, 100, 200, 260 and 300 mg (dry powder to be reconstituted before administration).

DOSE
Intravenous infusion Adult- 175 mg/m² body surface area over 3 h, repeat every 3 weeks. Antihistamines, corticosteroids or H2 antagonist may be required during treatment. Child- Not recommended.

INDICATION
Metastatic ovarian and breast cancer.
CONTRAINDICATION
Hypersensitivity; severe hepatic impairment; lactation; pregnancy (Appendix 7C)

ADVERSE EFFECTS
Myelosuppression, peripheral neuropathy and cardiac conduction defects with arrhythmias (which are nearly always asymptomatic); alopecia, muscle pain; nausea and vomiting is mild to moderate, hypersensitivity reactions; myalgia; arthralgia.

Imatinib

**EDL-D 667,668 tertiary restricted**

**INDICATIONS**
Chronic myeloid leukaemia, Philadelphia chromosome positive acute lymphoblastic leukaemia, Gastrointestinal stromal tumor.

**AVAILABILITY**
CAPSULES 100 and 400 mg.

**DOSE**
400-600 mg/day.

**CONTRAINDICATIONS**
Hypersensitivity.

**PRECAUTIONS**
Interactions (Appendix 6c); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
Acute-nausea and vomiting; chronic fluid retention with ankle and periorbital edema, diarrhoea, myalgias, congestive heart failure.

Thalidomide

**EDL-D 748 Tertiary restricted**

**AVAILABILITY**
CAPSULES 50 and 100 mg.

**DOSE**
Oral Multiple myeloma Adult: The dose is 200 mg administered orally once daily with water, preferably at bedtime and at least 1-hour after the evening meal. Thalidomide is administered in combination with dexamethasone in 28-day treatment cycles. Dexamethasone is 40 mg daily administered orally on days 1-4, 9-12, and 17-20 every 28 days. Erythema nodosum leprosum (ENL) Adult: For cutaneous ENL, thalidomide dosing should be initiated at 100 to 300 mg/day, administered once daily with water, preferably at bedtime and at least 1 hour after the evening meal. Not for monotherapy if moderate or severe neuritis present. Max: 400 mg/day. Patients < 50 kg: Initially, 100 mg daily. Dosing with thalidomide should continue until signs and symptoms of active reaction have subsided, usually a period of at least 2 weeks. Patients may then be tapered off medication in 50 mg decrements every 2 to 4 weeks. Patients who have a history of requiring prolonged maintenance treatment to prevent the recurrence of cutaneous ENL or who flare during tapering, should be maintained on the minimum dose necessary to control the reaction. Tapering off medication should be attempted every 3 to 6 months, in decrements of 50 mg every 2 to 4 weeks.

**INDICATION**
Multiple myeloma, erythema nodosum leprosum
CONTRAINDICATION
Hypersensitivity, pregnancy (Appendix 7C) and lactation, interactions (Appendix 6c)

PRECAUTION
During the period of treatment both males and females should take adequate means of contraception before, during and after (atleast 4 weeks) the therapy, therapy to be stopped immediately if pregnancy occurs, no blood or sperm donation during therapy, signs and symptoms of hypersensitivity include the occurrence of erythematous macular rash, possibly associated with fever, tachycardia, and hypotension, and if severe, may necessitate interruption of therapy. Serious dermatologic reactions including Stevens-Johnson syndrome and toxic epidermal necrolysis, seizures, impairment of mental and/or physical abilities required for the performance of hazardous tasks such as driving a car or operating complex machinery, potentiation of somnolence caused by alcohol, peripheral neuropathy, thromboembolism reported.

ADVERSE EFFECTS
Teratogenicity, Drowsiness/somnolence, peripheral neuropathy, constipation, dizziness, bradycardia, orthostatic hypotension, hypersensitivity, and neutropenia.

Gemcitabine

EDL-D 648 Tertiary restricted
INDICATIONS
Adenocarcinoma of pancreas.

AVAILABILITY
INJECTION Vial 200 mg and 1g (dry powder to be reconstituted before administration).

DOSE
1g/m2 body surface area for over 30 min once a week for up to 7 weeks, if not tolerated reduce or withhold. After one week rest administer by infusion once weekly for three weeks, withhold for 4th week before repeating.

CONTRAINDICATIONS
Pregnancy (Appendix 7c); concurrent radial radiotherapy; hypersensitivity; lactation.

PRECAUTIONS
Gemcitabine is not recommended for patients who can have potentially curative surgery. There is insufficient evidence about its use for second-line treatment of pancreatic adenocarcinoma, hepatic impairment; renal impairment, interactions (Appendix 6c).

ADVERSE EFFECTS
Nausea, vomiting, oral mucositis, hyperuricaemia, bone marrow suppression, alopecia, thromboembolism, flu like syndrome; edema; thrombocythemia; somnolence; hematuria; dyspnoea; loss of appetite.

STORAGE
Store in a sterile, airtight, tamperproof container.

Tamoxifen

EDL-D 742 Tertiary restricted
AVAILABILITY
Tablets 10, 20, 25, 40 and 100 mg.
DOSE
Breast cancer: Adult- 20 mg daily as a single dose or in 2 divided doses. max. 40 mg/day.
Anovulatory infertility: Adult- 20 mg daily on second- fifth day of the menstrual cycle. max.- 80 mg/day.

INDICATION
Adjuvant treatment for estrogen receptor positive breast cancer, metastatic breast cancer, male infertility, anovulatory infertility

CONTRAINDICATION
Hypersensitivity, deep vein thrombosis, pulmonary embolism, pregnancy (exclude before treatment and advise non-hormonal contraception if appropriate), lactation

PRECAUTION
If patient experiences swelling around ankles or legs, decrease salt intake, cystic ovarian swellings in premenopausal woman.

ADVERSE EFFECTS
Hypersensitivity reactions such as angioedema, Steven’s Johnson syndrome and bullous pemphigoid. Hot flushes, nausea, vomiting; vaginal discharge and bleeding, menstrual irregularities, increased risk of venous thromboembolism; distaste of food; depression; hair thinning; hypercalcaemia; peripheral oedema; decreased platelet count; increased pain and hypercalcaemia with bony maetastasis; tumor flare; liver enzyme changes (rarely, cholestasis); hepatitis; hepatic necrosis; hypertriglyceridaemia (sometimes with pancreatitis).

5-Fluorouracil

EDL-D 224 Tertiary restricted

INDICATIONS
Carcinomas of the colorectum, breast, stomach, pancreas, cervix, prostate, ovary and endometrium; liver tumours; head and neck tumours; actinic keratosis.

AVAILABILITY
INJECTION 5 and 10 ml ampoule (50 mg/ml).
TABLETS 50 mg.

DOSE
Intravenous injection
Initially 12 mg/kg body weight once a day for 4 days, max. daily dose 800 mg. If tolerated well without toxicity 6 mg/kg body weight can be given on 6th, 8th, 10th and 12th day. Discontinue on 12th day.
Maintenance dose 10 to 15 mg/kg body weight every week (max dose 1g/week).

CONTRAINDICATIONS
See notes above and consult literature; bone marrow depression; pregnancy (Appendix 7c) and lactation (Appendix 7b).

PRECAUTIONS
See notes above and consult literature; lactation; pelvic irradiation; renal impairment; hepatic impairment (Appendix 7a); interactions (Appendix 6c).

ADVERSE EFFECTS
See notes above and consult literature. Cardiac toxicity; tachycardia; dermatitis; diarrhoea.

STORAGE
Store protected from light in single dose container at a temperature not exceeding 30°C. The injection should not be allowed to freeze.
The use of pharmacotherapy will depend upon the degree of incapacity of the patient and is generally not justified until symptoms compromise working ability and social relationships; although levodopa is used in the early stages in some patients. Close supervision is then needed to ensure that treatment regimens are tolerated and that appropriate changes are made to the regimen as the disease progresses.

The most effective form of therapy is a combination of levodopa and a peripheral dopa-decarboxylase inhibitor, such as carbidopa. The response to levodopa with carbidopa is a compromise between increased mobility and adverse effects. Dyskinesias may be dose limiting and increasingly frequent with increased duration of treatment. Many factors including tolerance and progression of the disease may result in complications after 2-5 years of treatment. ‘End-of-dose’ deterioration occurs when there is a reduced duration of benefit from a dose, resulting in disability and dystonias. The ‘on-off’ phenomenon is characterized by sudden swings from mobility to episodes of akinesia, tremor and rigidity lasting from a few minutes to several hours. Amelioration of these effects can sometimes be achieved by administering levodopa in a sustained-release preparation or in a greater number of fractionated doses throughout the day. Psychiatric symptoms inducing disruption of sleep, vivid dreams and hallucinations are characteristic adverse effects that may occur at any time, especially in the elderly and may require dose reduction or withdrawal of levodopa.

Treatment for idiopathic parkinsonism is often initiated with a dopamine receptor agonist such as bromocriptine. Supplementary use of amantadine, bromocriptine or the monoamine-oxidase-B inhibitor, selegiline can be of value either to enhance the effect of levodopa or to reduce ‘end-of-dose’ fluctuations and ‘on-off’ effects.

Anticholinergic (more correctly termed antimuscarinic) drugs such as biperiden are usually sufficient in drug-induced parkinsonism.

**Drugs Used in Essential Tremor and Related Disorders:**

**Essential Tremor:** It can be treated with β-blockers such as propranolol (120 mg daily) (chapter 13.4) which may be of value if the tremor results in physical or social disability.

**Dystonias:** If no identifiable cause is found and the patient does not go into spontaneous remission, a trial of levodopa should be given to determine whether the patient has dopaminergic responsive dystonia. If there is no response within three months, the drug should be withdrawn and small doses of an anticholinergic drug such as biperiden should be given. The dosage may be increased gradually and up to 16 mg daily may be tolerated. In patients who fail to respond to either levodopa or an anticholinergic, other drugs including diazepam, baclofen, carbamazepine or phenothiazines may be of value. Psychological treatments have also been used successfully in the management of dyskinesias.
**Chorea:**
Choreiform movements can be induced by certain drugs including levodopa, phenytoin and antipsychotic drugs. Huntington’s disease is the most common of the hereditary choreas. Drug treatment is symptomatic and does not alter the progression of the disease. The aim of therapy is to reduce dopaminergic transmission which results from excessive or enhanced cholinergic activity. Antipsychotic drugs antagonize dopamine and usually lessen the chorea temporarily. Tetrabenzaine, the dopamine-depleting drug, is used to control movement disorders in Huntington’s chorea and related disorders.

**Tics:**
Tics which resemble choreiform movements are commonly associated with anxiety. However, in the more complex multiple tic disorder, Tourette syndrome, treatment with antipsychotic drugs may be required.

**Tardive Dyskinesia:**
It is associated with chronic administration of antipsychotic drugs. It is characterized by involuntary, repetitive, choreiform movement of the cheek, mouth and fingers. The first step of treatment should always be discontinuation of the antipsychotic drug or dosage reduction if the underlying psychotic disorder permits.

**Trihexyphenidyl Hydrochloride**

**EDL-D 511 Secondary hospitals**

**AVAILABILITY**
TABLETS 2 and 5 mg; INJECTION vial 2 mg/ ml

**DOSE**
1 mg daily, increased gradually; usual maintenance dose 5 to 15 mg daily in 3 to 4 divided doses (max. 20 mg daily); elderly preferably lower end of range.

**INDICATION**
All forms of parkinsonism other than medicine-induced, control of extrapyramidal disorders caused by CNS drugs.

**CONTRAINDICATION**
Avoided in gastrointestinal obstruction and myasthenia gravis; closed angle glaucoma; chronic pulmonary disease; sick sinus syndrome; thyrotoxicosis; tachycardia.

**PRECAUTION**
Use with caution in cardiovascular disease, hypertension, psychotic disorders, prostatic hypertrophy, pyrexia, in those susceptible to angle-closure glaucoma and in the elderly. It should not be withdrawn abruptly in patients receiving long-term treatment. Antimuscarinics are liable to abuse. Elderly males with possible prostate hypertrophy; tardive dyskinesia; neuroleptic malignant syndrome. Use with caution in renal impairment and hepatic impairment, lactation and interactions.

**ADVERSE EFFECTS**
Constipation, dry mouth, nausea, vomiting, tachycardia, dizziness, confusion, euphoria, hallucinations, impaired memory, anxiety, restlessness, urinary retention, blurred vision and rash. Angle-closure glaucoma may occur very rarely, paralytic ileus; dilation of colon.
Levodopa + Carbidopa  
EDL-D 301,302 Secondary hospitals  
AVAILABILITY  
TABLETS Levodopa 100 mg + Carbidopa 10 mg; Levodopa 100 mg + Carbidopa 25 mg; Levodopa 200 mg + Carbidopa 50 mg; Levodopa 250 mg + Carbidopa 25 mg.  
DOSE  
Oral Adult- Parkinsonism: expressed in terms of levodopa, initially 100 mg (with carbidopa 10 mg) twice daily, increased by 100 mg (with carbidopa 10 mg) every few days as necessary, to a max. of 1.5g. Optimum daily dose must be determined for each patient by careful monitoring and be taken after meals.  
INDICATION  
All forms of parkinsonism other than medicine-induced  
CONTRAINDICATION  
Concurrent use of monoamine oxidase inhibitors; undiagnosed chin lesion; lactation; psychosis; decompensated endocrine; angleclosure glaucoma; confirmed or suspected malignant melanoma.  
PRECAUTION  
Pulmonary disease, peptic ulceration, cardiovascular disease (including previous myocardial infarction); diabetes mellitus, osteomalacia, open-angle glaucoma, history of melanoma (risk of activation), psychiatric illness (avoid if severe); close monitoring of hepatic, haematological, psychiatric, cardiovascular and renal function required in long-term therapy; elderly: avoid rapid dose increases; warn patients to resume normal activities gradually; avoid abrupt withdrawal; pregnancy (toxicity in animals)(Appendix 7c), lactation; interactions (Appendix 6c).  
ADVERSE EFFECTS  
Nausea, anorexia and vomiting, particularly at the start of treatment; postural hypotension at the start of treatment, particularly in elderly and those receiving antihypertensives; excessive drowsiness and sudden onset of sleep (warn patient of these effects); confusion, vivid dreams, dizziness, tachycardia, arrhythmias; reddish discolouration of body fluids; insomnia, headache, flushing, gastrointestinal bleeding, peripheral neuropathy; taste disturbances, pruritus, rash, liver enzyme changes; psychiatric symptoms including psychosis, depression, hallucinations, delusions and neurological disturbances including dyskinesias may be dose-limiting; painful dystonic spasms ('end-of-dose' effects) and ('on-off' effects) after prolonged treatment (see notes above); neuroleptic malignant syndrome, on sudden withdrawal; rarely, hypersensitivity, dyspnoea; upper respiratory infection.
Antianemia medicines
Iron-Deficiency Anaemia:

Anaemia has many different aetiologies. It occurs when the haemoglobin concentration falls below the normal range for the age and sex of the individual. It is essential that a correct diagnosis is made before initiating therapy.

Any serious underlying cause of iron-deficiency anaemia, including gastric erosion and colonic carcinoma, should be excluded before giving iron replacement. Prophylaxis with iron salts in pregnancy should be given to women who have additional factors for iron-deficiency; low-dose iron and folic acid preparations are used for the prophylaxis of megaloblastic anaemia in pregnancy.

Ferrous salts should be given orally wherever possible. They differ only marginally in efficiency of absorption and thus the choice of preparation is usually decided by incidence of adverse effects and cost. Ferric salts are much less well absorbed. The oral dose of elemental iron for treatment of iron-deficiency anaemia in adults should be 100-200 mg daily with meals.

The approximate elemental iron content of various ferrous salts is: ferrous fumarate 200 mg (65 mg iron), ferrous gluconate 300 mg (35 mg iron), ferrous succinate 100 mg (35 mg iron), ferrous sulphate 300 mg (60 mg iron) and dried ferrous sulphate 200 mg (65 mg iron).

The haemoglobin concentration should rise by about 100-200 mg/100 ml per day or 2 g/100 ml over 3-4 weeks. After the haemoglobin has risen to normal, treatment should be continued for a further 3 months to replenish the iron stores.

Iron intake in the evening has been reported to improve its absorption. Iron intake with meals may reduce bioavailability but improve tolerability and adherence.

If adverse effects arise with one salt, dosage can be reduced or a change made to an alternative iron salt but an improvement in tolerance may be due to lower content of elemental iron. Gastrointestinal irritation may occur with iron salts. Nausea and epigastric pain are dose-related. Iron preparations taken orally may be constipating, particularly in the elderly, occasionally leading to faecal impaction. Oral iron may exacerbate diarrhoea in patients with inflammatory bowel disease but care is also needed in patients with intestinal strictures and diverticula. Iron as iron dextran (a complex of ferric hydroxide with dextrans) should be given parenterally only if the patient cannot tolerate oral iron, or does not take it reliably or there is continuing severe blood loss or malabsorption. Many patients with chronic renal failure who are receiving haemodialysis (and some on peritoneal dialysis) require intravenous iron on a regular basis. Parenteral iron may cause more harm than benefit. With the exception of patients on haemodialysis the haemoglobin response is not significantly faster with the parenteral route than the oral route.
**Megaloblastic Anaemia:**
Megaloblastic anaemias result from a lack of either vitamin B12 (hydroxocobalamin) or folate or both. The clinical features of folate-deficient megaloblastic anaemia are similar to those of vitamin B12 deficiency except that the accompanying severe neuropathy does not occur; it is essential to establish the underlying cause in every case. Hydroxocobalamin is used to treat vitamin B12 deficiency whether due to dietary deficiency or malabsorption including pernicious anaemia (due to a lack of intrinsic factor, which is essential for vitamin B12 absorption).

Folate deficiency due to poor nutrition, pregnancy, antiepileptics or malabsorption is treated with folic acid but this should never be administered without vitamin B12 in undiagnosed megaloblastic anaemia because of the risk of precipitating neurological changes due to vitamin B12 deficiency.

Preparations containing a ferrous salt and folic acid are used for the prevention of megaloblastic anaemia in pregnancy. The low doses of folic acid in these preparations are inadequate for the treatment of megaloblastic anaemias.

**Prevention of Neural Tube Defects:**
An adequate intake of folic acid before conception and during early pregnancy reduces the risk of neural tube defects in babies. Therefore, women planning a pregnancy should receive sufficient folic acid before conception and in the first 12 weeks of pregnancy; folic acid may be given as a food or a medicinal supplement in a dose of 400-500 μg daily. A woman who has not received supplementary folic acid and suspects that she might be pregnant should start taking folic acid at once and continue until 12th week of pregnancy.

Women at increased risk of giving birth to a baby with neural tube defects (for example history of neural tube defect in a previous child) should receive a higher dose of folic acid of approximately 5 mg daily, starting before conception and continuing for 12 weeks after conception. Women taking antiepileptic medication should be counselled by their doctor before starting folic acid.

**Ferrous Gluconate, Ferrous Sulphate**  
**Iron Sucrose, Sodium Ferric Gluconate**  
**Iron Dextran**

**AVAILABILITY**
- TABLETS (sugar coated, film coated) Ferrous sulphate 200 mg, Ferrous fumarate 200 mg, Ferrous gluconate 300 mg. (all equivalent to 65 mg elemental iron). In women, folic acid may also be given. SYRUPS also available. CAPSULES Iron sulfate 60-150 mg (20% Iron), Iron fumarate 200-300 mg (33% Iron). INJECTIONS Iron dextran 50 mg/ml, Iron sucrose 20, 50 and 100 mg/ml, Sodium ferric gluconate 12.5 mg/ml.

**DOSE**
- Oral Adult - Iron-deficiency anaemia: elemental iron 100 to 200 mg daily in divided doses.
- Prevention of iron deficiency anaemia (in those at particular risk): for woman elemental iron 60 mg daily. Child- under 5 years: elemental iron 2 mg/kg (max. 30 mg) daily. Over 5 years: elemental iron 30 mg daily. Over 5 years: folic acid may also be given. Parenteral Total dose (ml) = 0.0442 (desired haemoglobin - observed haemoglobin) x LBW + (0.26 x LBW) [Note: LBW = Lean Body Weight (Kg)] Total dose may be given in divided doses in a daily or twice weekly basis via
IM inj. (into the upper quadrant of the buttock); may also be given intravenously by total-dose infusion or as divided inj. A-Z track technique (displacement of the skin laterally prior to injection) is recommended to avoid injection or leakage into subcutaneous tissue

INDICATION
Iron-deficiency anaemia.

CONTRAINDICATION
Haemosiderosis, haemochromatosis; any form of anaemia not caused by iron deficiency; evidence of iron overload; patients receiving repeated blood transfusions; parenteral iron therapy.

PRECAUTION
A test dose of 0.5 ml should be given observe patient for at least 1 hour for signs of hypersensitivity, respiratory distress, tachycardia or back/chest pain; should not be administered for longer than 6 months; pregnancy (Appendix 7c); peptic ulcer; hypotension; regional enteritis, ulcerative colitis, intestinal strictures, diverticula; interactions (Appendix 6c, 6d).

ADVERSE EFFECTS
Nausea, vomiting, metallic taste; constipation, diarrhoea, dark stools, epigastric pain, gastrointestinal irritation; long-term or excessive administration may cause haemosiderosis; allergic reaction; back pain; staining of teeth. Parenteral: Pain at injection site, sterile abscess.

Hydroxyurea
EDL-D268, 661 Secondary hospitals

INDICATION
Myeloproliferative disorders (primarily polycythemia vera) not responding to venepuncture and essential thrombocytosis), Sickle Cell Disease (Breaks down cells that are prone to sickle, as well as increasing content), Second Line treatment for Psoriasis

DOSE
10-30 mg/kg per day

ADVERSE EFFECT
Drowsiness, nausea, vomiting and diarrhea, mucositis, (which may take 7-21 days to recover after the drug has been discontinued), skin changes

Folic Acid
EDL-D232,233 Universal

INDICATIONS
Treatment of folate-deficiency megaloblastic anaemia; prevention of neural tube defect in pregnancy.

AVAILABILITY
TABLETS 1, 5 and 10 mg.

DOSE
Oral
Adult- Treatment of folate-deficiency, megaloblastic anaemia: 5 mg daily for 4 months (up to 15 mg daily may be necessary in malabsorption states). Prevention of first occurrence of neural tube defect: 400 to 500 μg daily before conception and during the first twelve weeks of pregnancy. Prevention of recurrence of neural tube defect: 5 mg daily (reduced to 4 mg daily, if suitable preparation available) from at least 4 weeks before conception until twelfth week of pregnancy.

CONTRAINDICATIONS
Should never be given without vitamin B12 in undiagnosed megaloblastic anaemia or other vitamin B12 deficiency states because risk of precipitating subacute combined degeneration of the spinal cord; folatedependent malignant disease.
PRECAUTIONS
Women receiving antiepileptic therapy need counselling before starting folic acid; pernicious anaemia; folate dependent tumor; interactions (Appendix 6c); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Neuropathy; bronchospasm; skin eruption; anorexia; skin rash; status epilepticus.

STORAGE
Store protected from light.

Ferrous Sulphate + Folic Acid
EDL-D 221,222,223 Universal

AVAILABILITY
TABLETS (sugar coated, film coated) Ferrous sulphate 200 mg, Ferrous fumarate 200 mg, Ferrous gluconate 300 mg. (all equivalent to 65 mg elemental iron). In women, folic acid may also be given. SYRUPS also available. CAPSULES Iron sulfate 60-150 mg (20% Iron), Iron fumarate 200-300 mg (33% Iron). INJECTIONS Iron dextran 50 mg/ml, Iron sucrose 20, 50 and 100 mg/ml, Sodium ferric gluconate 12.5 mg/ml.

DOSE
Oral Adult- Iron-deficiency anaemia: elemental iron 100 to 200 mg daily in divided doses. Prevention of iron deficiency anaemia (in those at particular risk): for woman elemental iron 60 mg daily. Child- under 5 years: elemental iron 2 mg/kg (max. 30 mg) daily. Over 5 years: elemental iron 30 mg daily. Over 5 years: folic acid may also be given. Parenteral Total dose (ml) = 0.0442 (desired haemoglobin- observed haemoglobin) x LBW + (0.26 x LBW) [Note: LBW = Lean Body Weight (Kg)] Total dose may be given in divided doses in a daily or twice weekly basis via IM inj. (into the upper quadrant of the buttock); may also be given intravenously by total-dose infusion or as divided inj. A-Z track technique (displacement of the skin laterally prior to injection) is recommended to avoid injection or leakage into subcutaneous tissue

INDICATION
Iron-deficiency anaemia.

CONTRAINDICATION
Haemosiderosis, haemochromatosis; any form of anaemia not caused by iron deficiency; evidence of iron overload; patients receiving repeated blood transfusions; parenteral iron therapy.

PRECAUTION
A test dose of 0.5 ml should be given & observe patient for at least 1 hour for signs of hypersensitivity, respiratory distress, tachycardia or back/chest pain; should not be administered for longer than 6 months; pregnancy ; peptic ulcer; hypotension; regional enteritis, ulcerative colitis, intestinal strictures, diverticula; interactions

ADVERSE EFFECTS
Nausea, vomiting, metallic taste; constipation, diarrhoea, dark stools, epigastric pain, gastrointestinal irritation; long-term or excessive administration may cause haemosiderosis; allergic reaction; back pain; staining of teeth. Parenteral: Pain at injection site, sterile abscess.

Folic Acid (Sodium salt)
EDL-D234 Secondary hospitals

AVAILABILITY
TABLETS 1, 5 and 10 mg.

DOSE
Oral Adult- Treatment of folate-deficiency, megaloblastic anaemia: 5 mg daily for 4 months (up to 15 mg daily may be necessary in malabsorption states). Prevention of first occurrence of neural tube defect: 400 to 500 μg daily before conception and during the first twelve weeks of
pregnancy. Prevention of recurrence of neural tube defect: 5 mg daily (reduced to 4 mg daily, if suitable preparation available) from at least 4 weeks before conception until twelfth week of pregnancy.

**INDICATION**
Treatment of folate-deficiency megaloblastic anaemia; prevention of neural tube defect in pregnancy.

**CONTRAINDICATION**
Should never be given without vitamin B12 in undiagnosed megaloblastic anaemia or other vitamin B12 deficiency states because risk of precipitating subacute combined degeneration of the spinal cord; folate-dependent malignant disease.

**PRECAUTION**
Women receiving antiepileptic therapy need counselling before starting folic acid; pernicious anaemia; folate dependent tumor; interactions (Appendix 6c); pregnancy (Appendix 7c)

**ADVERSE EFFECTS**
Neuropathy; bronchospasm; skin eruption; anorexia; skin rash; status epilepticus.

**Iron Dextran**

**Non-EDL  Secondary category**

**INDICATIONS**
Iron deficiency anaemia, prevention of iron deficiency before, during or after pregnancy, to make up iron deficiency after pregnancy and during lactation.

**AVAILABILITY**
INJECTION (iron as iron dextran) 1.5 ml ampoule (50 mg/ml).

**DOSE**
Deep intramuscular injection into the gluteal muscle or slow intravenous injection or intravenous infusion.

Adult- Calculated according to body-weight and iron deficit. While deciding on parenteral therapy, oral therapy should be stopped at least 24 h before. Urine may darken on starting.

Child- Under 14 years: not recommended.

**CONTRAINDICATIONS**
History of allergic disorders including asthma and eczema; infection; active rheumatoid arthritis; liver disease.

**PRECAUTIONS**
Oral iron not to be given until 5 days after last injection; hepatic impairment; renal impairment; pregnancy (Appendix 7c); interactions (Appendix 6d). Anaphylactic reactions can occur with parenteral iron and a test dose is recommended before each dose; the patient should be carefully observed for 60 min after the first test dose and for 15 min after subsequent test doses (subsequent test doses not necessary for intramuscular administration). Facilities for cardiopulmonary resuscitation must be at hand; risk of allergic reactions increased in immune or inflammatory conditions.

**ADVERSE EFFECTS**
Less commonly nausea, vomiting, abdominal pain, flushing, dyspnoea, anaphylactic reactions (see Anaphylaxis above), numbness, cramps, blurred vision, pruritus and rash; rarely, diarrhoea, chest pain, hypotension, angioedema, arrhythmias, tachycardia; dizziness, restlessness, fatigue; seizures, tremor, impaired consciousness, myalgia, arthralgia and sweating; injection-site reactions also reported, thrombophlebitis; peripheral vascular flushing; taste disturbances; syncope.
Hydroxocobalamin
EDL-D 265 Secondary hospitals

AVAILABILITY
INJECTION 1 ml (1 mg/ml).

DOSE
Intramuscular injection Adult and Child- Megaloblastic anaemia without neurological involvement: initially 1 mg 3 times a week for 2 weeks, then 1 mg every 3 months. Megaloblastic anaemia with neurological involvement: initially 1 mg on alternate days until no further improvement occurs, then 1 mg every 2 months. Prophylaxis of macrocytic anaemias: 1 mg every 2 to 3 months. Tobacco amblyopia and Leber optic atrophy: 1 mg daily for 2 weeks, then 1 mg twice weekly until no further improvement, then 1 mg every 1 to 3 months.

INDICATION
Megaloblastic anaemia due to vitamin B12 deficiency, congenital intrinsic factor disease.

CONTRAINDICATION
Anaphylactic reaction.

PRECAUTION
Except in emergencies, should not be given before diagnosis confirmed; monitor serum potassium levels-arrhythmias secondary to hypokalaemia in early therapy; pregnancy (Appendix 7c)

ADVERSE EFFECTS
Itching, exanthema, fever, chills, hot flushes, nausea, dizziness; rarely, acneiform and bullous eruptions, anaphylaxis; hypersensitivity; headache; diarrhea.

Medicine affecting coagulation
Anticoagulants are used to prevent thrombus formation or extension of an existing thrombus in the slower-moving venous side of the circulation, where the thrombus consists of a fibrin web enmeshed with platelets and red cells. They are therefore used widely in the prevention and treatment of deep vein thrombosis in the legs, prophylaxis of embolization in rheumatic heart disease and atrial fibrillation and to prevent thrombi forming on prosthetic heart valves.

Heparin is a parenteral anticoagulant that initiates anticoagulation rapidly but has a short duration of action. The low molecular weight heparins have a longer duration of action.

For the treatment of deep venous thrombosis and pulmonary embolism heparin is given as an intravenous loading dose followed by continuous intravenous infusion (using an infusion pump) or by intermittent subcutaneous injection. An oral anticoagulant is started at the same time as heparin. The heparin needs to be continued for at least 5 days, until the oral anticoagulant has taken effect and the INR (international normalized ratio) has been in the therapeutic range for 2 consecutive days. Laboratory monitoring is essential, on a daily basis. Heparin is also used in regimens for the management of myocardial infarction, the management of unstable angina, acute peripheral arterial occlusion and in dialysis.

In patients undergoing general surgery, low-dose heparin by subcutaneous injection is used to prevent postoperative deep-vein thrombosis and pulmonary embolism in high risk patients (those with obesity, malignant disease, history of deep-vein thrombosis or pulmonary embolism, patients over 40 years, those with an established thrombophilic disorder or those undergoing major or complicated surgery). It is also of value in high-risk medical patients, for example obesity, heart failure, when confined to bed.
If haemorrhage occurs it is usually sufficient to withdraw heparin, but if rapid reversal of the effects of heparin is required, protamine sulphate is a specific antidote.

Oral anticoagulants take at least 48-72 h for the anticoagulant effect to develop fully; if an immediate effect is needed, heparin must be given concomitantly. Warfarin is indicated in deep vein thrombosis, pulmonary embolism, for patients with atrial fibrillation who are at risk of embolization and for those with mechanical prosthetic heart valves (to prevent emboli developing on the valves); oral anticoagulants should not be used in cerebral thrombosis or peripheral arterial occlusion as firstline therapy. The main adverse effect of oral anticoagulants is haemorrhage. Prothrombin time (usually reported as INR, international normalized ratio) should be checked on a daily basis initially then at longer intervals depending on response. If severe haemorrhage occurs, stop warfarin and give phytomenadione (vitamin K) by slow intravenous injection.

**Anticoagulants in Pregnancy:**

Oral anticoagulants are teratogenic and should not be given in the first trimester of pregnancy. Women at risk of pregnancy should be warned of this danger since stopping warfarin before the sixth week of gestation may largely avoid the risk of fetal abnormality. Oral anticoagulants cross the placenta with the risk of placental or fetal haemorrhage, especially during the last few weeks of pregnancy and at delivery. Therefore, if at all possible, oral anticoagulants should be avoided in pregnancy, especially in the first and third trimester. Difficult decisions may have to be made, particularly in women with prosthetic heart valves or with a history of recurrent venous thrombosis or pulmonary embolism.

**Haemophilia:**

Desmopressin by injection may aid haemostasis and be useful in mild forms of haemophilia. For minor procedures including dental surgery, it may circumvent the need for factor VIII.

**Heparin Sodium**

**EDL-D 254,255,256 Secondary hospitals**

**AVAILABILITY**

INJECTION vials 1000, 5000 and 25,000 IU/ml

**DOSE**

Intravenous injection

Adult-Treatment of deep-vein thrombosis and pulmonary embolism: loading dose of 5000 units (10,000 units in severe pulmonary embolism) followed by continuous intravenous infusion of 15 to 25 units/kg/h. Child- 50 to 100U/kg every 4 to 6 h. Subcutaneous injection 15,000 units every 12 h; laboratory monitoring is essential, preferably on a daily basis and dose adjusted accordingly. Prophylaxis in general surgery: 5,000 units 2 h before surgery, then every 8 to 12 h for 7 days or until patient is ambulant (monitoring not needed); during pregnancy (with monitoring) 5,000-10,000 units every 12 h. Note: Not intended to cover prosthetic heart valve management in pregnancy, which requires specialist management. Child- 250 units/kg every 12 h. Intravenous injection and continuous intravenous infusion. Child- By intravenous injection: lower loading dose, then by continuous intravenous infusion; 15 to 25 units/kg/h.
INDICATION
Treatment and prophylaxis of deep-vein thrombosis and pulmonary embolism; atrial fibrillation with embolism; treatment and prophylaxis of peripheral arterial embolism; prophylaxis of deep vein thrombosis in major surgery; lipemia clearing.

CONTRAINDICATION
Hypersensitivity to heparin; haemophilia and other haemorrhagic disorders; thrombocytopenia; peptic ulcer; recent cerebral haemorrhage; severe hypertension; severe liver or renal disease; after major trauma or recent surgery (especially to eye or nervous system); threatened abortion; piles; bacterial endocarditis; large malignancies; tuberculosis; lumbar puncture; chronic alcoholics; acetylsalicylic acid and other antiplatelet drugs.

PRECAUTION
Hepatic impairment (Appendix 7a) and renal failure; hypersensitivity to low molecular weight heparins; spinal or epidural anaesthesia-risk of spinal haematoma; diabetes mellitus; acidosis; concomitant potassium-sparing drugs-increased risk of hyperkalaemia; lactation; paediatrics; elderly; interactions (Appendix 6c); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Immune-mediated thrombocytopenia usually developing 6 to 10 days after commencement of therapy (requires immediate withdrawal of heparin); haemorrhage; skin necrosis; hypersensitivity reactions including urticaria; angioedema and anaphylaxis; osteoporosis after prolonged use and rarely, alopecia; bleeding due to overdose.

Phytomenadione
EDL-D 410,411 Secondary hospitals

AVAILABILITY
TABLETS 5 and 10 mg; INJECTION 10 mg/ml.

DOSE
Slow intravenous injection Adult- Warfarin-induced hypoprothrombinaemia, no bleeding or minor bleeding: 500 μg. Oral For vitamin K deficiency: 10 to 40 mg daily. Warfarin-induced hypoprothrombinaemia, no bleeding or minor bleeding: 5 mg. Oral or intramuscular injection Less severe haemorrhage: 10 to 20 mg. Slow intravenous injection Severe haemorrhage: 2.5 to 5 mg; very rarely, up to 50 mg (but risk of over correction with high dosage). Intravenous or intramuscular injection Child- Neonates: Haemorrhagic disease of the newborn (treatment): 1 mg with further doses if necessary at 8 h intervals (prophylaxis). Intramuscular injection Child-0.5 to 1 mg as single dose. Oral Child- 2 mg followed by a second dose after 4 to 7 days and for breastfed babies a third dose after 1 month.

INDICATION
Antagonist to warfarin; prophylaxis against haemorrhagic disease of the newborn; vit K deficiency, hematuria, menorrhagia.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Reduce dose in elderly; hepatic impairment; not an antidote to heparin; can cause haemolysis in patients with G-6-PD; increased risk of severe haemolytic anaemia in neonates after large doses; premature neonates weighing < 2.5 kg; pregnancy.

ADVERSE EFFECTS
Hypersensitivity reactions including flushing; dyspnoea; bronchospasm; dizziness; hypotension and respiratory or circulatory collapse which may be due to polyethoxylated castor oil surfactant in some injection formulations rather than due to phytomenadione.
Protamine Sulphate  
EDL-D 411 Secondary hospitals  

AVAILABILITY
SOLUTION 5 ml (1%); Injection 5 ml ampoule (10 mg/ml).

DOSE
Intravenous injection Heparin overdose, over approximately 10 min; 1 mg neutralizes 80 to 100 units heparin when given within 15 min, if longer time, less protamine needed as heparin is rapidly excreted. 1 ml neutralises the effect of 1000 ml i.u. of circulating heparin; max. single dose 50 mg (5 ml).

INDICATION
Antidote to overdosage with heparin; antidote for heparin in controlled bleeding

PRECAUTION
If used in excess protamine has an anticoagulant effect; allergic reactions increased in persons at risk including previous treatment with protamine or protamine insulin; fish allergies; men who are infertile or who have had a vasectomy; pregnancy (Appendix 7c); lactation; children.

ADVERSE EFFECTS
Nausea; vomiting; lassitude; flushing; hypotension; bradycardia; dyspnoea; allergic reactions (including angioedema; anaphylaxis); allergy specially if previous exposure to protamine insulin; fish allergy; infertile or vasectomised men.

Warfarin Sodium  
EDL-D 530,531 Secondary hospitals  

AVAILABILITY
TABLETS 1, 2 and 5 mg.

DOSE
Oral Adult- Prophylaxis and treatment of thromboembolic disorders; usual induction dose is 10 mg daily for 2 days, according to the individual patient; the subsequent dose depends upon the prothrombin time; the usual daily maintenance dose is 3 to 9 mg administered at the same time each day.Given as slow injection over 1 to 2 minutes into peripheral vein, initially 5 mg daily. For rapid anticoagulation: initially 10 mg daily for 2 days, maintenance dose 2 to 10 mg daily.

INDICATION
Prophylaxis of embolisation in rheumatic heart disease and atrial fibrillation; prophylaxis after insertion of prosthetic heart valve; prophylaxis and treatment of venous thrombosis and pulmonary embolism; transient ischaemic attacks; myocardial infarction; vulvular heart disease.

CONTRAINDICATIONS
Pregnancy (Appendix 7c); peptic ulcer; severe hypertension; bacterial endocarditis; hypersensitivity; blood dyscrasias; recent surgery; psychosis; pericardial effusion; cerebrovascular disorder; alcoholism; senility; aneurysm.

PRECAUTIONS
Heparin induced thrombocytopenia; surgery or trauma; Vit C, K; lactation; alcoholics; purple toes syndrome; discontinue if necrosis develops; elderly; hepatic impairment (Appendix 7a) or renal failure; recent surgery; lactation (Appendix 7b); interactions (Appendix 6a, 6b, 6c, 6d).

ADVERSE EFFECTS
Heparin induced thrombocytopenia; surgery or trauma; Vit C, K; lactation; alcoholics; purple toes syndrome; discontinue if necrosis develops; elderly; hepatic impairment or renal failure; recent surgery; lactation; interactions
Streptokinase
EDL-D 484 Tertiary

AVAILABILITY
INJECTION (Powder for solution for injection) 7,50,000 and 15,00,000 units vial.

DOSE
Intravenous infusion. Adult- Thrombosis: 2,50,000 units over 30 min, followed by 1,00,000 units every h for 12 to 72 h according to condition with monitoring of clotting parameters. Myocardial infarction: 15,00,000 units over 60 min.

INDICATION
Life-threatening deep-vein thrombosis, pulmonary embolism, acute arterial thromboembolism; thrombosed arteriovenous shunts; acute myocardial infarction.

CONTRAINDICATION
Recent haemorrhage; surgery (including dental); parturition; trauma; heavy vaginal bleeding; haemorrhagic stroke; history of cerebrovascular disease (especially recent or if residual disability); coma; severe hypertension; coagulation defects; bleeding diatheses; aortic dissection; risk of gastrointestinal bleeding such as recent history of peptic ulcer; oesophageal varices; ulcerative colitis; acute pancreatitis; severe liver disease; acute pulmonary disease with cavitation; previous allergic reactions; pregnancy

PRECAUTION
Risk of bleeding from any invasive procedure; including injection; external chest compression; abdominal aneurysm or where thrombolysis may give rise to embolic complications such as enlarged left atrium with atrial fibrillation (risk of dissolution of clot and subsequent embolization); diabetic retinopathy (small risk of retinal haemorrhage); recent or concurrent anticoagulant treatment; platelet count; fibrinogen level; thrombin and prothrombin time

ADVERSE EFFECTS
Nausea and vomiting; bleeding; usually limited to site of injection but internal bleeding including intracranial haemorrhage may occur (if serious bleeding occurs; discontinue infusion-coagulation factors may be required); hypotension; arrhythmias (particularly in myocardial infarction); allergic reactions including rash; flushing; uveitis; anaphylaxis; fever; chills; back or abdominal pain; Guillain-Barré syndrome reported rarely
**Plasma substitute**

Dextran 70 and polygeline are macromolecular substances which are metabolized slowly; they may be used to expand and maintain blood volume in shock arising from conditions such as burns or septicaemia. They are rarely needed when shock is due to sodium and water depletion as, in these circumstances, the shock responds to water and electrolyte repletion.

Plasma substitutes should not be used to maintain plasma volume in conditions such as burns or peritonitis where there is loss of plasma protein, water and electrolytes over periods of several days. In these situations, plasma or plasma protein fractions containing large amounts of albumin should be given.

Plasma substitutes may be used as an immediate short-term measure to treat massive haemorrhage until blood is available, but large volumes of some plasma substitutes can increase the risk of bleeding by depleting coagulation factors. Dextran may interfere with blood group cross-matching or biochemical measurements and these should be carried out before the infusion is started.

**Hydroxy Ethyl Starch**

**EDL-D 267 Secondary hospitals**

**INDICATIONS**

Therapy for hypovolaemia, shock in surgery, trauma and infection to improve haemodynamics, macrocirculation, microcirculation and oxygen supply; improve organ function in blood loss.

**AVAILABILITY**

INFUSION 300 and 500 ml.

**DOSE**

Intravenous infusion 500 to 1000 ml (daily max. 1500 ml).

**CONTRAINDICATIONS**

Renal failure; haemorrhage; coagulation disorders; anuria; oliguria.

**Precautions**

Should be used with caution in patients with cardiac disease; liver disease; or renal impairment; urine output should be monitored. Care should be taken to avoid haematocrit concentration from falling below 25-30% and the patient should be monitored for hypersensitivity reactions; bleeding disorder; sufficient fluid should be administered to avoid dehydration; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Hypersensitivity reactions may occur including; rarely, severe anaphylactoid reactions; transient increase in bleeding time may occur; headache; tachycardia; itching; fall in blood pressure.

**Tranexamic Acid**

**EDL-D 509,510 Tertiary**

**AVAILABILITY**

TABLETS- 250 mg, 500 mg and 1g. INJECTION- 100 mg/ml, 500 mg/5 ml.

**DOSE**

Dental extraction in Hemophiliacs: Immediately before tooth extraction, 10 mg/ kg intravenously. Following tooth extraction, intravenous therapy, at a dose of 10 mg/kg body weight three to four times daily, may be used for 2 to 8 days. Menorrhagia: 1300 mg orally 3 times daily up to 5 days during menstruation. Cone biopsy: 1000-1500 mg 2-3 times daily for 12 days postop.
eratively. Epistaxis: 1000 mg 3 times daily for 7 days. Hyphema: 1000-1500 mg 2-3 times daily for 7 days. Hereditary angioedema: 1000-1500 mg 2-3 times daily.

INDICATION
Prevention of hemorrhage due to dental procedures in hemophilics, cyclic heavy menstrual bleeding, hereditary angioedema, cone biopsy, epistaxis, traumatic hyphema

CONTRAINdICATION
Hypersensitivity, acquired defective colour vision, subarachnoid hemorrhage, active intravascular clotting, pregnancy (Appendix 7c), interactions (Appendix 6c).

PRECAUTION
Renal impairment, disseminated intravascular coagulation, thromboembolic history, coadministration with hormonal contraceptives may increase risk of thrombosis, stroke, or myocardial infarction; women using hormonal contraception should take tranexamic acid only if there is a strong medical need, and if the benefit of treatment outweighs risks. Ligneous conjunctivitis has been reported. Retinal venous and arterial occlusion has been reported in patients using tranexamic acid. Patients should be instructed to report visual and ocular symptoms promptly.

ADVERSE EFFECTS
Nausea, vomiting, diarrhoea, disturbances in colour vision (discontinue), thromboembolic events, allergic skin reactions; giddiness and hypotension on rapid intravenous injection, headache, backache, musculoskeletal pain

**Plasma fraction for specific –T**
Factor VIII is essential for blood clotting and the maintenance of effective haemostasis; von Willebrand factor is a mediator in platelet aggregation and also acts as a carrier for factor VIII. Blood coagulation factors VII, IX and X are essential for the conversion of factor II (prothrombin) to thrombin. Deficiency in any of these factors results in haemophilia. Bleeding episodes in haemophilia require prompt treatment with replacement therapy. Factor VIII, used for the treatment of haemophilia A, is a sterile freeze-dried powder containing the blood coagulation factor VIII fraction prepared from pooled human venous plasma. Standard factor VIII preparations also contain von Willebrand factor and may be used to treat von Willebrand disease. Highly purified preparations, including recombinant factor VIII, are available; they are indicated for the treatment of haemophilia A but do not contain sufficient von Willebrand factor for use in the management of von Willebrand disease.
Factor IX Complex is a sterile freeze-dried concentrate of blood coagulation factors II, VII, IX and X derived from fresh venous plasma. Factor IX complex which is used for the treatment of haemophilia B may also be used for the treatment of bleeding due to deficiencies of factor II, VII and X. High purity preparations of factor IX which do not contain clinically effective amounts of factor II, VII and X are available. A recombinant factor IX preparation is also available.

**Albumin**

EDL-D 14, 15 Tertiary

AVAILABILITY
Solution 5%, 10%, 20%.

DOSE
Intravenous infusion For hypovolemia: Adult- 25g, Child- 1g/kg. Max.- 2g of 20%/kg body weight. For hypoproteinaemia: Adult- 2g/kg daily. Usual rates of infusion: up to 5 ml/min (5%) or 1 to 2 ml/min (20%).

INDICATION
Burns, hypoproteinaemia, shock, hypovolemia, acute liver failure, dialysis.
CONTRAINDICATION
Congestive heart failure, severe anaemia, history of allergic reactions to human albumin; pregnancy (Appendix 7c)

PRECAUTION
If dehydration is present additional fluid must follow the administration of albumin. Administration of albumin should be supplemented or replaced by packed red blood cells, history of cardiac or circulatory disease, increased capillary permeability.

ADVERSE EFFECTS
Allergic (or) pyrogenic reactions, tachycardia, rash, anaphylactic shock, increased salivation.

Factor IX Complex (Coagulation factors II, VII, IX X)
EDL-D 215 Tertiary

AVAILABILITY
INFUSION (Powder for solution for infusion), factor II, VII, IX and X 500 to 1500 units.

DOSE
Slow intravenous infusion Adult and child- Haemophilia B: according to patient’s needs. Treatment of bleeding due to deficiencies in factor II, VII or X as well as IX: according to patient’s need.

INDICATION
Replacement therapy for factor IX deficiency in haemophilia; bleeding due to deficiencies of factors II, VII or X.

CONTRAINDICATION
Disseminated intravascular coagulation; hypersensitivity to any component of the product.

PRECAUTION
Risk of thrombosis (probably less risk with highly purified preparations); pregnancy(Appendix 7c) ; preexisting disease; check heart rate; interactions

ADVERSE EFFECTS
Allergic reactions including chills; fever; hepatitis; pulmonary embolism; disseminated intravascular coagulation.

Factor VIII Concentrate
EDL-D Tertiary

INDICATIONS
Control of haemorrhage in haemophilia A.

AVAILABILITY
INFUSION (Powder for solution for infusion), factor VIII 250 to 1500 units.

DOSE
Slow intravenous infusion Adult and child- Haemophilia A; according to patient’s needs.

CONTRAINDICATIONS
Hypersensitivity to any component of the product.

PRECAUTIONS
Intravascular haemolysis after large or frequently repeated doses in patients with blood groups A; B; or AB (less likely with high potency; highly purified concentrates); pregnancy (Appendix 7c); check heart rate.

ADVERSE EFFECTS
Allergic reactions including chills; fever; hepatitis; anaphylaxis; fulminating hepatitis.

STORAGE
Store protected from light.
Antianginal medicines

The three main types of angina are:

- **Stable angina** (angina of effort), where atherosclerosis restricts blood flow in the coronary vessels; attacks are usually caused by exertion and relieved by rest
- **Unstable angina** (acute coronary insufficiency), which is considered to be an intermediate stage between stable angina and myocardial infarction
- **Prinzmetal angina** (variant angina), caused by coronary vasospasm, in which attacks occur at rest.

Management depends on the type of angina and may include drug treatment, coronary artery bypass surgery, or percutaneous transluminal coronary angioplasty.

**Stable Angina:**

Drugs are used both for the relief of acute pain and for prophylaxis to reduce further attacks; they include organic nitrates, beta-adrenoceptor antagonists (beta-blockers) and calcium-channel blockers.

**Nitrates:**

Organic nitrates have a vasodilating effect; they are sometimes used alone, especially in elderly patients with infrequent symptoms. Tolerance leading to reduced antianginal effect is often seen in patients taking prolonged-action nitrate formulations. Evidence suggests that patients should have a ‘nitrate-free’ interval to prevent the development of tolerance. Adverse effects such as flushing, headache and postural hypotension may limit nitrate therapy but tolerance to these effects also soon develops. The short-acting sublingual formulation of glyceryl trinitrate is used both for prevention of angina before exercise or other stress and for rapid treatment of chest pain. A sublingual tablet of isosorbide dinitrate is more stable in storage than glyceryl trinitrate and is useful in patients who require nitrates infrequently; it has a slower onset of action, but effects persist for several hours.

**Beta-Blockers:**

Beta-adrenoceptor antagonists (beta-blockers), such as atenolol, block beta-adrenergic receptors in the heart and thereby decrease heart rate and myocardial contractility and oxygen consumption, particularly during exercise. Beta-blockers are first-line therapy for patients with effort-induced chronic stable angina; they improve exercise tolerance, relieve symptoms, reduce the severity and frequency of angina attacks and increase the anginal threshold. Beta-blockers should be withdrawn gradually to avoid precipitating an anginal attack; they should not be used in patients with underlying coronary vasospasm (Prinzmetal’s angina).

Beta-blockers may precipitate asthma and should not be used in patients with asthma or a history of obstructive airways disease. Some, including atenolol, have less effect on β2 (bronchial) receptors and are therefore relatively cardioselective. Although they have less effect on airways resistance they are not free of this effect and should be avoided.
Beta-blockers slow the heart and may induce myocardial depression, rarely, precipitating heart failure. They should not be given to patients who have incipient ventricular failure, second-or third-degree atrioventricular block, or peripheral vascular disease. Beta-blockers should be used with caution in diabetes since they may mask the symptoms of hypoglycaemia, such as rapid heart rate. Beta-blockers enhance the hypoglycaemic effect of insulin and may precipitate hypoglycaemia.

**Calcium-Channel Blockers:**
A calcium-channel blocker, such as verapamil, is used as an alternative to a beta-blocker to treat stable angina. Calcium-channel blockers interfere with the inward movement of calcium ions through the slow channels in heart and vascular smooth muscle cell membranes, leading to relaxation of vascular smooth muscle. Myocardial contractility may be reduced, the formation and propagation of electrical impulses within the heart may be depressed and coronary or systemic vascular tone may be diminished. Calcium-channel blockers are used to improve exercise tolerance in patients with chronic stable angina due to coronary atherosclerosis or with abnormally small coronary arteries and limited vasodilator reserve. Calcium-channel blockers can also be used in patients with unstable angina with a vasospastic origin, such as Prinzmetal’s angina and in patients in whom alterations in cardiac tone may influence the angina threshold.

**Unstable Angina:**
Unstable angina requires prompt aggressive treatment to prevent progression to myocardial infarction. Initial treatment is with acetylsalicylic acid to inhibit platelet aggregation, followed by heparin. Nitrates and beta-blockers are given to relieve ischaemia; if beta-blockers are contraindicated, verapamil is an alternative, provided left ventricular function is adequate. **Prinzmetal’s Angina:** Treatment is similar to that for unstable angina, except that a calcium-channel blocker is used instead of a beta-blocker.

**Isosorbide Dinitrate**

**EDL-D289,290,291 Secondary hospitals**

**AVAILABILITY**
TABLETS 10, 20, 40, 50 and 60 mg; TABLETS (SR) 50 mg and 60 mg; CAPSULE 30, 40 and 60 mg.

**DOSE**
Oral 20 mg 2 to 3 times a day initially, or 40 mg twice daily (max 120 mg daily individual dose).

**INDICATION**
Prophylaxis and treatment of angina, congestive heart failure.

**CONTRAINDICATION**
Hypersensitivity to nitrates; hypotension; hypovolaemia; myocardial infarction; hypertrophic obstructive cardiomyopathy, aortic stenosis, cardiac tamponade, constrictive pericarditis, mitral stenosis; marked anaemia; head trauma; cerebral haemorrhage; angle-closure glaucoma.

**PRECAUTION**
Severe hepatic or renal impairment; hypothyroidism; malnutrition; hypothermia; recent history of myocardial infarction; interactions (Appendix 6a, 6b, 6c, 6d); pregnancy (Appendix 7c). Patients taking isosorbide dinitrate for the long-term management of angina may often develop tolerance to the antianginal effect; this can be avoided by giving the second of 2 daily doses of longer-acting oral presentations after an 8-h rather than a 12-h interval, thus ensuring a nitrate-free interval each day.
ADVERSE EFFECTS
Throbbing headache; flushing; dizziness, postural hypotension, tachycardia (paradoxical bradycardia also reported); palpitation, decreased cardiac output; confusion; increased intracranial pressure.

Isosorbide Mononitrate
EDL-D 292, Secondary hospitals

AVAILABILITY
TABLETS 10, 20, 40, 50 and 60 mg; TABLETS (SR) 50 mg and 60 mg; CAPSULE 30, 40 and 60 mg.

DOSE
Oral 20 mg 2 to 3 times a day initially, or 40 mg twice daily (max 120 mg daily individual dose).

INDICATION
Prophylaxis and treatment of angina, congestive heart failure.

CONTRAINDICATION
Hypersensitivity to nitrates; hypotensive conditions and hypovolaemia; hypertrophic cardiomyopathy; aortic stenosis; cardiac tamponade; constrictive pericarditis; mitral stenosis; marked anaemia; glaucoma; obstructive cardiomyopathy; raised intracranial pressure.

PRECAUTION
Hypothyroidism; malnutrition; hypothermia; head trauma; cerebral haemorrhage; gastrointestinal disease; recent history of myocardial infarction; hypoxaemia or other ventilation and perfusion abnormalities; susceptibility to angle-closure glaucoma; metal-containing transdermal systems should be removed before cardioversion or diathermy; avoid abrupt withdrawal; tolerance; severe hepatic impairment; severe renal impairment; pregnancy (Appendix 7c); lactation; interactions (Appendix 6a)

ADVERSE EFFECTS
Postural hypotension, tachycardia (but paradoxical bradycardia also reported); throbbing headache; dizziness; less commonly nausea; vomiting; heartburn; flushing; temporary hypoxaemia; rash; application site reactions with transdermal patches; very rarely, angle-closure glaucoma; decreased cardiac output; urinary and faecal incontinence. Specific side-effects following injection (particularly if given too rapidly) include severe hypotension, diaphoresis, apprehension, restlessness, muscle twitching, retrosternal discomfort, palpitation, abdominal pain, syncope; prolonged administration has been associated with methaemoglobinaemia.

Acetylsalicylic Acid
EDL-D2, 3 PHC

INDICATIONS
As an antiplatelet agent for prophylaxis of myocardial infarction, stable angina; stable angina pectoris; stroke prophylaxis.

AVAILABILITY
TABLETS 50, 60, 75, 80, 150, 300 and 325 mg.

DOSE
Oral
Adult- Analgesic and antipyretic including migraine attacks: 0.3 to 0.9g, 3 to 4 times a day (max. 4g daily). Acute Rheumatic fever: 4 to 6g or 75 to 100 mg/kg daily in divided doses. Antiplatelet: 75-325 mg/day.
Child- Under 16 years: not recommended (can cause Reye’s syndrome).

CONTRAINDICATIONS
Hypersensitivity (including asthma; angioedema; urticaria or rhinitis) to acetylsalicylic acid or any other NSAID; children and adolescents under 16 years (may cause Reye’s syndrome);
gastrointestinal ulceration; haemophilia and other bleeding disorders; not for treatment of  
gout; severe renal or hepatic impairment; lactation. It is known to cause haemolytic anaemia in 
people who have the genetic disease- G-6-PD-deficiency.

PRECAUTIONS
Asthma, allergic disease; impaired renal or hepatic function (Appendices 7d and 7a);  
lactation (Appendix 7b); pregnancy (Appendix 7c); elderly; G-6-PD-deficiency; dehydration;  
interactions (Appendix 6a, 6c, 6d).

ADVERSE EFFECTS
Generally mild and infrequent for lower doses, but common with anti-inflammatory doses;  
gastrointestinal discomfort or nausea, ulceration with occult bleeding (occasionally major  
haemorrhage); also other haemorrhage (including subconjunctival); hearing disturbances such  
as tinnitus (rarely, deafness); vertigo; confusion; hypersensitivity reactions (angioedema;  
bronchospasm and rash); increased bleeding time, blood disorders (particularly thrombocytopenia); rarely, oedema; myocarditis; Reye’s syndrome.

STORAGE
Store protected from moisture at a temperature not exceeding 30°C

Verapamil Hydrochloride
EDL-D 520 Tertiary △

AVAILABILITY
TABLETS 40, 80, 120 and 240 mg (SR); INJECTION 2 ml (5 mg/2 ml).

DOSE
Oral Adult- 80 to 120 mg 3 times daily (120 mg 3 times daily usually required in Prinzmetal  
angina). Supraventricular arrhythmias: 40 to 120 mg 3 times daily. Intravenous injection Adult-  
Supraventricular arrhythmias: 5 to 10 mg over 2 min (preferably with ECG monitoring). Elderly-  
Paroxysmal tachyarrhythmias: 5 to 10 mg over 3 min, further 5 mg may be given after 5 to 10  
min if required.

INDICATION
Angina, including stable, unstable and Prinzmetal angina; arrhythmias; ischaemic heart disease;  
migraine.

CONTRAINDICATION
Hypotension, bradycardia, second- and third-degree atroventricular block, sinoatrical block, sick  
sinus syndrome; cardiogenic shock; history of heart failure or significantly impaired left  
ventricular function (even if controlled by therapy); atrial flutter or fibrillation complicating  
Wolff-Parkinson- White syndrome; porphyria; platelet dysfunction

PRECAUTION
First-degree atroventricular block; kidney impairment; cirrhosis patients; acute phase of  
myocardial infarction (avoid if bradycardia, hypotension, left ventricular failure); hepatic  
impairment (Appendix 7a); children (specialist advice only); lactation; pregnancy (Appendix 7c);  
interactions (Appendix 6b, 6c).

ADVERSE EFFECTS
Constipation; less commonly nausea, vomiting, flushing, headache, dizziness, fatigue, ankle  
oedema; rarely, allergic reactions (erythema, pruritus, urticaria, angioedema, Stevens-Johnson  
syndrome); myalgia; arthralgia, paraesthesia, increased prolactin concentration; gynaecomastia  
and gingival hyperplasia on long-term treatment; with high doses, hypotension, heart failure,  
bradycardia, heart block and asystole (due to negative inotropic effect), impotence;  
hepatotoxicity; hyperprolactinemia; myoclonic dystonia.
Glyceryl Trinitrate  
EDL-D 371Tertiary  
D654,655 Secondary hospitals  
AVAILABILITY  
TABLETS 0.5, 2.6 and 6.4 mg; CAPSULES 2.5 and 6.4 mg; INJECTION 5 and 10 ml (5 mg/ml);  
SUBLINGUAL TAB 500 µg. SPRAY 0.4 mg/puff (200 mdi)  
DOSE  
Sublingual Adult- 0.5 to 1 mg, repeated as required. Intravenous infusion 10 to 200 µg/min.  
INDICATION  
Prophylaxis and treatment of angina, myocardial infarction; post operative hypertension; cardio-pulmonary edema.  
CONTRAINDICATION  
Hypersensitivity to nitrates; hypotension; hypovolaemia; raised intracranial pressure; hypertrophic obstructive cardiomyopathy, aortic stenosis, cardiac tamponade, constrictive pericarditis, mitral stenosis; marked anaemia; head trauma; cerebral haemorrhage; angle-closure glaucoma.  
PRECAUTION  
Severe hepatic or renal impairment; hypothyroidism; malnutrition; gastrointestinal hypermotility; malabsorption syndrome; hypothermia; recent history of myocardial infarction; interactions (Appendix 6b,6c)  
ADVERSE EFFECTS  
Throbbing headache; flushing; dizziness, postural hypotension; tachycardia (paradoxical bradycardia also reported); abdominal pain; collapse; neurological deficit.  

Clopidogrel  
EDL-D 589 Secondary hospitals  
AVAILABILITY  
TABLETS 75 and 150 mg.  
DOSE  
Adult- 75 mg once daily. Non-ST segment elevation myocardial infarction: loading dose 300 mg followed by 75 mg once daily.  
INDICATION  
Prophylaxis in thromboembolic disorders including myocardial infarction, peripheral arterial disease and stroke, acute coronary syndrome.  
CONTRAINDICATION  
Hypersensitivity, active pathological bleeding such as peptic ulcer or intracranial hemorrhage, coagulation disorders, lactation  
PRECAUTION  
Patient with increased risk of bleeding from trauma, surgery or other pathological conditions, ulcers, renal impairment, hepatic impairment, history of bleeding or haemostatic disorder, pregnancy (Appendix 7c); interactions (Appendix 6c)  
ADVERSE EFFECTS  
Bleeding, neutropenia, thrombocytopenia, other bone marrow toxicity, diarrhoea, epigastric pain, rashes, paraesthesia, vertigo.
Antiarrhythmic medicines

Treatment of arrhythmias requires precise diagnosis of the type of arrhythmia and electrocardiography is essential; underlying causes such as heart failure require appropriate treatment. Antiarrhythmic drugs must be used cautiously since most drugs that are effective in treating arrhythmias can provoke them in some circumstances; this arrhythmogenic effect is often enhanced by hypokalaemia. When antiarrhythmic drugs are used in combination, their cumulative negative inotropic effects may be significant, particularly if myocardial function is impaired.

Atrial Fibrillation:
The increased ventricular rate in atrial fibrillation can be controlled with a beta-adrenoceptor antagonist (beta-blocker) or verapamil. Digoxin is often effective for controlling the rate at rest; it is also appropriate if atrial fibrillation is accompanied by congestive heart failure. Intravenous digoxin is occasionally required if the ventricular rate needs rapid control. If adequate control at rest or during exercise cannot be achieved readily verapamil may be introduced with digoxin, but it should be used with caution if ventricular function is impaired. Anticoagulants are indicated especially in valvular or myocardial disease and in the elderly. Warfarin is preferred to acetylsalicylic acid in preventing emboli. If atrial fibrillation began within the previous 48 h and there does not appear to be a danger of thromboembolism, antiarrhythmic drugs, such as procainamide or quinidine, may be used to terminate the fibrillation or to maintain sinus rhythm after cardioversion.

Atrial Flutter:
Digoxin will sometimes slow the ventricular rate at rest. Reversion to sinus rhythm is best achieved by direct current electrical shock. If the arrhythmia is long-standing, treatment with an anticoagulant should be considered before cardioversion to prevent emboli. Intravenous verapamil reduces ventricular fibrillation during paroxysmal (sudden onset and intermittent) attacks of atrial flutter. An initial intravenous dose may be followed by oral treatment; hypotension may occur with high doses. It should not be used for tachyarrhythmias where the QRS complex is wide unless a supraventricular origin has been established beyond doubt. If the flutter cannot be restored to sinus rhythm, antiarrhythmics such as quinidine can be used.

Paroxysmal Supraventricular Tachycardia:
In most patients this remits spontaneously or can revert to sinus rhythm by reflex vagal stimulation. Failing this, intravenous injection of a beta-adrenoceptor antagonist (beta-blocker) or verapamil may be effective. Verapamil and a beta-blocker should never be administered concomitantly because of the risk of hypotension and asystole.

Ventricular Tachycardia:
Very rapid ventricular fibrillation causes profound circulatory collapse and must be treated immediately with direct current shock. In more stable patients intravenous lidocaine or procainamide may be used. After sinus rhythm is restored, drug therapy to prevent recurrence of ventricular tachycardia should be considered; a beta-adrenoceptor antagonist (beta-blocker) or verapamil may be effective.
Torsades de pointes is a special form of ventricular tachycardia associated with prolongation of the QT interval. Initial treatment with intravenous infusion of magnesium sulphate (usual dose 2g over 10-15 min, repeated once if necessary) together with temporary pacing is usually effective; alternatively, isoprenaline infusion may be given with extreme caution until pacing can be instituted. Isoprenaline is an inotropic sympathomimetic; it increases the heart rate and therefore shortens the QT interval, but given alone it may induce arrhythmias.

Bradyarrhythmias:
Sinus bradycardia (less than 50 beats/min) associated with acute myocardial infarction may be treated with atropine. Temporary pacing may be required in unresponsive patients. Drugs are of limited value for increasing the sinus rate long term in the presence of intrinsic sinus node disease and permanent pacing is usually required.

Cardiac Arrest:
In cardiac arrest, epinephrine (adrenaline) is given by intravenous injection in a dose of 1 mg (10 ml of 1 in 10,000 solution) as part of the procedure for cardiopulmonary resuscitation.

Atenolol
EDL-D 52 PHC
AVAILABILITY
TABLETS 12.5, 25, 50, and 100 mg; INJECTION ampoule 5 mg/ml (10 ml).
DOSE
Oral Adult- 50 mg once daily, increased if necessary to 50 mg twice daily or 100 mg once daily. Angina: 50 mg daily administered alone or with a diuretic, dose can be increased to 100 mg (over 100 mg has no added advantage). May also be administered in combination with a mlodipine besylate 2.5 or 5 mg. Child- 1 to 1.3 mg/kg body weight once daily or divided every 12 h. Intravenous injection 2.5 mg at a rate of 1 mg/min, repeat at 5 min interval to a max. 10 mg.
INDICATION
Angina and myocardial infarction; arrhythmias; hypertension; migraine prophylaxis.
CONTRAINDICATION
Hypersensitivity; sinoarterial node disease, atrioventricular node disease
PRECAUTION
Avoid abrupt withdrawal especially in angina; may precipitate or worsen heart failure; acute myocardial infarction, pregnancy (Appendix 7c), thyrotoxicosis, pheochromocytoma; lactation (Appendix 7b); first-degree atrioventricular block; liver function deteriorates in portal hypertension; reduce dose in renal impairment; diabetes mellitus (small decrease in glucose tolerance, masking of symptoms of hypoglycaemia); history of hypersensitivity (increased reaction to allergens, also reduced response to epinephrine (adrenaline); myasthenia gravis; interactions (Appendix 6a, 6b, 6c).
ADVERSE EFFECTS
Dry mouth; sedation; dizziness; nausea; nocturnal restlessness; occasionally rashes; cardiac arrhythmias; systemic lupus erythmatosus; anxiety; constipation; abdominal pain; hallucination; impotence and depression.

Digoxin
EDL-D178,179,180 Secondary hospitals
AVAILABILITY
TABLET 0.25 mg; INJECTION 2 ml (0.5 mg/2 ml); ELIXIR 0.05 mg/ml (paediatric use); SYRUP 1.5 mg/30 ml.
DOSE
Oral Adult- Atrial fibrillation and heart failure: 1 to 1.5 mg in divided doses over 24 h for rapid digitalization or 250 μg 1 to 2 times daily if digitalization less urgent; maintenance 62.5 to 500 μg daily (higher dose may be divided), according to renal function and heart rate response; usual range 125 to 250 μg daily. Elderly- Lower dose more appropriate. Intravenous infusion Emergency control of atrial fibrillation, over at least 2 h: 0.75 to 1 mg. Emergency loading dose for heart failure, over at least 2 h: 0.75 to 1 mg.

INDICATION
Supraventricular arrhythmias, particularly atrial fibrillation; heart failure.

CONTRAINDICATION
Hypertrophic obstructive cardiomyopathy (unless also atrial fibrillation and heart failure); ventricular tachycardia; hypokalaemia; digitalis toxicity; arrhythmias; Wolff-Parkinson-White syndrome or other accessory pathway, particularly if accompanied by atrial fibrillation; intermittent complete heart block; seconddegree atrioventricular block.

PRECAUTION
Recent myocardial infarction; sick sinus syndrome; severe pulmonary disease; thyroid disease; congestive cardiac myopathy; hypercalcaemia; aortic valve disease, heart block, cardiac dysrhythmias; elderly (reduce dose); renal impairment (Appendix 7d); avoid hypokalaemia; avoid rapid intravenous administration (nausea and risk of arrhythmias); lactation; interactions (Appendix 6c, 6d); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Usually associated with excessive dosage and include anorexia, nausea, vomiting, diarrhoea, abdominal pain; visual disturbances, headache, fatigue, drowsiness, confusion, delirium, hallucinations, depression; arrhythmias, heart block; rarely, rash, intestinal ischaemia; gynaecomastia on long-term use; thrombocytopenia reported; sinus bradycardia; apathy; psychosis; malaise.

Lidocaine hydrochloride
EDL-D 312 Secondary hospitals

AVAILABILITY
injection vial 30 ml (1, 2%w/v), 50 ml (21.3 mg/ml); 2%/50 ml; ampoule 5%/2 ml. JELLY 2% w/v

OINTMENT 5% w/v

DOSE
Adult- Ventricular arrhythmias: loading dose of 50 to 100 mg (or 1 to 1.5 mg/kg) at a rate of 25 to 50 mg/min by intravenous injection, followed immediately by intravenous infusion of 1 to 4 mg/min, with ECG monitoring of all patients (reduce infusion dose if required for longer than 24 h).

INDICATION
Ventricular arrhythmias (especially after myocardial infarction); local anaesthesia.

CONTRAINDICATION
Sino-atrial disorder; any grade of atrioventricular block or any other type of conduction disturbances, severe myocardial depression, acute porphyria or hypovolaemia, bradycardia, cardiac decompensation.

PRECAUTION
Lower dosage in congestive heart failure, bradycardia, ECG monitoring must during therapy, pediatrics; hypotension; renal impairment; porphyria; debilitated patients; hepatic impairment (Appendix 7a); marked hypoxia; severe respiratory depression; following cardiac surgery and in elderly; lactation; interactions (Appendix 6c); pregnancy (Appendix 7c).
ADVERSE EFFECTS
Dizziness; paraesthesia; drowsiness, confusion; apnoea, respiratory depression; coma; seizures and convulsions; hypotension, arrhythmias, heart block; cardiovascular collapse and bradycardia (may lead to cardiac arrest); nystagmus often an early sign of lidocaine overdosage; blurred vision, disorientation.

Amiodarone
EDL-D 23,24 Tertiary

AVAILABILITY
TABLETS 100 and 200 mg; INJECTION 3 ml ampoule (50 mg/ml).

DOSE
Oral 200 mg three times a day for one week, reduced to 200 mg twice daily for further one week. Maintenance 200 mg daily or reduced to minimum required to control arrhythmia. Intravenous infusion (with central venous catheter). Initially 5 mg/kg body weight over 20 to 120 min with ECG monitoring, subsequent infusion given if necessary according to response (up to max 1.2g in 24 h).

INDICATION
Severe rhythmic disorder where other therapies cannot be used including tachyarrhythmia associated with Wolff-Parkinson-White syndrome, atrial flutter and fibrillation; all types of paroxysmal tachycardia.

CONTRAINDICATION
Sinus bradycardia, sino-atrial heart block; unless pacemaker fitted avoid in severe conduction disturbances or sinus node disease; thyroid dysfunction; iodine sensitivity; goitre; avoid intravenous use in severe respiratory failure, circulatory collapse, severe arterial hypotension, avoid bolus injection in congestive heart failure or cardiomyopathy; lactation; pregnancy (Appendix 7c)

PRECAUTION
Liver-function and thyroid-function tests required before treatment and then every 6 months; hypokalaemia (measure serum potassium concentration before treatment); chest x-ray required before treatment; heart failure; elderly; severe bradycardia and conduction disturbances in excessive dosage; intravenous use may cause moderate and transient fall in blood pressure (circulatory collapse precipitated by rapid administration or overdosage) or severe hepatocellular toxicity (monitor transaminases closely); ECG monitoring and resuscitation facilities must be available during intravenous use; porphyria, interactions(Appendix 6d)

ADVERSE EFFECTS
Nausea, vomiting, taste disturbances, raised serum transaminases (may require dose reduction or withdrawal if accompanied by acute liver disorders), jaundice; bradycardia; pulmonary toxicity (including pneumonitis and fibrosis); tremor, sleep disorders; hypothyroidism, hyperthyroidism; reversible corneal microdeposits (sometimes with night glare); phototoxicity, persistent slate-grey skin discolouration; less commonly onset or worsening of arrhythmia, conduction disturbances, peripheral neuropathy and myopathy (usually reversible on withdrawal); very rarely, chronic liver disease including cirrhosis, sinus arrest, bronchospasm (in patients with severe respiratory failure), ataxia, benign intracranial hypertension, headache, vertigo, epididymo-orchitis, impotence, haemolytic or aplastic anaemia, thrombocytopenia, rash (including exfoliative dermatitis), hypersensitivity including vasculitis, alopecia, impaired vision due to optic neuritis or optic neuropathy (including blindness), anaphylaxis on rapid injection, also hypotension, respiratory distress syndrome, sweating and hot flushes.
Diltiazem  
EDL-D 182,183 Tertiary  
AVAILABILITY 
TABLETS/TABLETS (SR) 30, 60, 90, 120, 180 and 240 mg; CAPSULE 60, 90, 120, 180 and 240 mg; INJECTION 5 ml (25 mg/5 ml).

DOSE 
Oral Adult-30 mg 2 to 5 times a day before food and at night (bed time), increase gradually to 240 mg in 3 to 4 divided doses daily. Child- Not recommended. Cardiac arrhythmia Adult-Initially 250 μg/kg by i.v. bolus over 2 min.

INDICATION 
Angina pectoris due to coronary artery spasm; chronic stable angina; cardiac arrhythmia.

CONTRAINDICATION 
Severe bradycardia; left ventricular failure with pulmonary congestion; second- or third-degree AV block (unless pacemaker fitted); sick sinus syndrome; lactation.

PRECAUTION 
Reduce dose in hepatic and renal impairment; heart failure or significantly impaired left ventricular function; bradycardia (avoid if severe); first degree AV block; or prolonged PR interval; interactions (Appendix 6c); sinoatrial nodal dysfunction; pregnancy (Appendix 7c).

ADVERSE EFFECTS 
Bradycardia, sino-atrial block, AV block; palpitation; dizziness; hypotension, malaise; asthenia; headache; hot flushes; gastrointestinal disturbances; oedema (notably of ankles); rarely, rashes (including erythema multiforme and exfoliative dermatitis), photosensitivity; hepatitis; gynaecomastia; gum hyperplasia; extrapyramidal symptoms; depression reported; gastrointestinal haemorrhage; sinus arrest.

Adenosine  
EDL-D 11 Tertiary  
AVAILABILITY 
TABLETS 40, 80 and 120 mg (DT); INJECTION 2 ml ampoule (3 mg/ml).

DOSE 
Oral 40 to 80 mg, 3 to 4 times daily (Max. 480 mg/ day). Rapid intravenous injection (into central or large peripheral vein) 3 mg every 2 seconds with regular cardiac monitoring, if necessary, followed by 6 mg every 1 to 2 min. Increment should not be given if higher level AV block occurs at any particular dose.

INDICATION 
Coronary vasodilator; paroxysmal supraventricular tachycardia; cardiac imaging for coronary artery disease; angina pectoris.

CONTRAINDICATION 
Second- or third-degree AV block and sick sinus syndrome (unless pacemaker fitted), acute myocardial infarction, cardiovascular shock; asthma.

PRECAUTION 
Atrial fibrillation or flutter with accessory pathway (conduction down anomalous pathway may increase); heart transplant; pregnancy (Appendix 7c).

ADVERSE EFFECTS 
Transient facial flush, chest pain, dyspnoea, bronchospasm, choking sensation, nausea, light-headedness; severe bradycardia reported (requiring temporary pacing); ECG may show transient rhythm disturbances; edema; constipation.
Antihypertensive medicines

Management of Hypertension:

Treatment of hypertension should be integrated into an overall programme to manage factors that increase the risk of cardiovascular events (such as stroke and myocardial infarction). Treatment is often life-long. Hypertension was formerly classified as mild, moderate or severe, but a grading system is now preferred. Grade 1 hypertension is defined as 140-159 mmHg systolic blood pressure and 90-99 mmHg diastolic blood pressure, Grade 2 hypertension 160-179 mmHg systolic and 100-109 mmHg diastolic and Grade 3 hypertension more than 180 mmHg systolic and more than 110 mmHg diastolic. The goal of treatment is to obtain the maximum tolerated reduction in blood pressure.

Lifestyle changes should be introduced for all patients; they include weight reduction, reduction in alcohol intake, reduction of dietary Sodium, stopping tobacco smoking and reduction in saturated fat intake. The patient should eat a healthy nutritious diet including adequate fruit and vegetables and should exercise regularly. These measures alone may be sufficient in mild hypertension, but patients with moderate to severe hypertension will also require specific antihypertensive therapy.

Drug Treatment of Hypertension:

Three classes of drug are used for first-line treatment of hypertension: thiazide diuretics, beta-adrenoceptor antagonists (beta-blockers) and angiotensin-converting enzyme (ACE) inhibitors. Calcium-channel blockers are considered first-line in specific populations only e.g. Africans or the elderly. Other classes of drugs may be used in certain situations.

Thiazide diuretics, such as hydrochlorothiazide, have been used as first-line antihypertensive therapy and are particularly indicated in the elderly. They have few adverse effects in low doses, but in large doses they may cause a variety of unwanted metabolic effects (principally potassium depletion), reduced glucose tolerance, ventricular ectopic beats and impotence; they should be avoided in gout. These effects can be reduced by keeping the dose as low as possible; higher doses do not produce an increased reduction in blood pressure. Thiazides are inexpensive and, when used in combination, can enhance the effectiveness of many other classes of antihypertensive drugs.

Beta-adrenoceptor antagonists (beta-blockers) such as atenolol are effective in all grades of hypertension and are particularly useful in angina and following myocardial infarction; they should be avoided in asthma, chronic obstructive pulmonary disease and heart block.

Angiotensin-converting enzyme inhibitors (ACE inhibitors) such as enalapril are effective and well tolerated by most patients. They can be used in heart failure, left ventricular dysfunction and diabetic nephropathy, but should be avoided in renovascular disease and in pregnancy. The most common adverse effect is a dry persistent cough.
Dihydropyridine calcium-channel blockers such as nifedipine are useful for isolated systolic hypertension, in populations unresponsive to other antihypertensives (e.g. Africans) and in the elderly when thiazides cannot be used. Short-acting formulations of nifedipine should be avoided as they may evoke reflex tachycardia and cause large variations in blood pressure.

Drugs acting on the central nervous system are also effective antihypertensive drugs. In particular, methyldopa is effective in the treatment of hypertension in pregnancy. A single antihypertensive drug is often not adequate and other antihypertensive drugs are usually added in a stepwise manner until blood pressure is controlled.

**Hypertensive Emergencies**

In situations where immediate reduction of blood pressure is essential and treatment by mouth is not possible, intravenous infusion of Sodium nitroprusside is effective. Over-rapid reduction in blood pressure is hazardous and can lead to reduced organ perfusion and cerebral infarction.

**Atenolol**

**EDL-D52,53 PHC**

**AVAILABILITY**

TABLETS 12.5, 25, 50, and 100 mg; INJECTION ampoule 5 mg/ml (10 ml).

**DOSE**

Oral Adult: 50 mg once daily, increased if necessary to 50 mg twice daily or 100 mg once daily.

Angina: 50 mg daily administered alone or with a diuretic, dose can be increased to 100 mg (over 100 mg has no added advantage). May also be administered in combination with amlodipine besylate 2.5 or 5 mg. Child: 1 to 1.3 mg/kg body weight once daily or divided every 12 h. Intravenous injection 2.5 mg at a rate of 1 mg/min, repeat at 5 min interval to a max. 10 mg.

**INDICATION**

Angina and myocardial infarction; arrhythmias; hypertension; migraine prophylaxis.

**CONTRAINDICATION**

Hypersensitivity; sinoarterial node disease, atrioventricular node disease

**PRECAUTION**

Avoid abrupt withdrawal especially in angina; may precipitate or worsen heart failure; acute myocardial infarction, pregnancy (Appendix 7c), thyrotoxicosis, pheochromocytoma; lactation (Appendix 7b); first-degree atrioventricular block; liver function deteriorates in portal hypertension; reduce dose in renal impairment; diabetes mellitus (small decrease in glucose tolerance, masking of symptoms of hypoglycaemia); history of hypersensitivity (increased reaction to allergens, also reduced response to epinephrine (adrenaline); myasthenia gravis; interactions (Appendix 6a, 6b, 6c).

**ADVERSE EFFECTS**

Dry mouth; sedation; dizziness; nausea; nocturnal restlessness; occasionally rashes; cardiac arrhythmias; systemic lupus erythmatosus; anxiety; constipation; abdominal pain; hallucination; impotence and depression.

**Methyldopa**

**EDL-D337 Secondary hospitals**

**INDICATIONS**

Hypertension in pregnancy.

**AVAILABILITY**

TABLET 250 mg.
DOSE

Oral

Adult- Hypertension in pregnancy: initially 250 mg 2 to 3 times daily; if necessary, gradually increased at intervals of 2 or more days (max 3g daily).

CONTRAINDICATIONS

Depression; active liver disease; hypersensitivity; therapy with MAO inhibitors; pheochromocytoma; porphyria.

PRECAUTIONS

History of hepatic impairment (Appendix 7a); renal impairment; blood counts and liverfunction tests advised; history of depression; positive direct Coomb test in up to 20% of patients (affects blood cross-matching); interference with laboratory tests; lactation; pregnancy (Appendix 7c); interactions (Appendix 6b, 6c). May impair ability to perform skilled tasks; for example operating machinery; driving.

ADVERSE EFFECTS

Tend to be transient and reversible including sedation; dizziness; lightheadedness; postural hypotension; weakness; fatigue; headache; fluid retention and oedema; sexual dysfunction; impaired concentration and memory; depression; mild psychosis; disturbed sleep and nightmares; drug fever; influenza-like syndrome; nausea; vomiting; constipation; diarrhoea; dry mouth; stomatitis; sialadenitis; liver function impairment; hepatitis; jaundice; rarely, fatal hepatic necrosis; bone marrow depression; haemolytic anaemia; leukopenia; thrombocytopenia; eosinophilia; parkinsonism; rash (including toxic epidermal necrolysis); nasal congestion; black or sore tongue; bradycardia; exacerbation of angina; myalgia; arthralgia; paraesthesia Bell palsy; pancreatitis; hypersensitivity reactions including lupus erythematosus-like syndrome; myocarditis; pericarditis; gynaecomastia; hyperprolactinaemia; amenorrhoea; urine darkens on standing.

Sodium Nitroprusside

EDL-D 735 Tertiary

INDICATIONS

Hypertensive crisis (when treatment by mouth not possible), congestive heart failure.

AVAILABILITY

INJECTION ampoule/vial 5 ml (50 mg/ml).

DOSE

Intravenous infusion

Adult- Hypertensive crisis: initially 0.3 μg/kg/min; usual maintenance dose 0.5 to 6 μg/kg/min; max. dose 8 μg/kg/min; stop infusion if response is unsatisfactory after 10 min at max. dose; lower doses in patients already being treated with antihypertensives.

CONTRAINDICATIONS

Compensatory hypertension; severe vitamin B12 deficiency; Leber optic atrophy; arterial venous shunting; patients with acute CHF associated with reduced peripheral vascular resistance.

PRECAUTIONS

Impaired pulmonary function; hypothyroidism; renal impairment; ischaemic heart disease; impaired cerebral circulation; hyponatraemia; raised intracranial pressure; elderly; hypothermia; monitor blood pressure and blood-cyanide concentration; also blood-thiocyanate concentration if given for more than 3 days; avoid sudden withdrawal (reduce infusion over 15-30 min to avoid rebound effects); pregnancy (Appendix 7c); lactation; interactions (Appendix 6b); hepatic impairment (Appendix 7a).

ADVERSE EFFECTS

Severe hypotension; effects associated with over-rapid reduction in blood pressure include headache; dizziness; retching; abdominal pain; perspiration; palpitations; apprehension;
retrosternal discomfort; rarely, reduced platelet count; acute transient phlebitis; muscle twitching; hypothyroidism; increased anaerobic metabolism.

Adverse effects associated with excessive concentration of cyanide metabolite include tachycardia; sweating; hyperventilation; arrhythmias; marked metabolic acidosis (discontinue infusion and give antidote).

**STORAGE**
Store protected from light.

**Enalapril**
**EDL-D 197,198 Secondary hospitals**

**AVAILABILITY**
TABLETS 2.5, 5 and 10 mg; INJECTION 1 ml ampoule (1.25 mg/ml).

**DOSE**
Oral Adult- Hypertension: initially 5 mg once daily; if used in addition to diuretic. Heart failure, asymptomatic left ventricular dysfunction: initially 2.5 mg daily under close medical supervision; usual maintenance dose 20 mg daily in 1 to 2 divided doses. Elderly- Renal impairment: initially 2.5 mg daily. Usual maintenance dose 10 to 20 mg once daily; In severe hypertension may be increased to max. 40 mg once daily.

**INDICATION**
Heart failure (with a diuretic); prevention of symptomatic heart failure and prevention of coronary ischaemic events in patients with left ventricular dysfunction; hypertension; renal hypertension.

**CONTRAINDICATION**
Renal failure with anuria; precomatose states associated with liver cirrhosis; severe sodium and water depletion; hypersensitivity to sulphonamides and furosemide; hypokalaemia; addison’s disease; lactation.

**PRECAUTION**
Use with diuretics; hypotension with first doses; especially in patients on diuretics; on a low-Sodium diet; on dialysis; if dehydrated; or with heart failure; peripheral vascular disease or generalized atherosclerosis (risk of clinically silent renovascular disease); use with great care in severe or symptomatic aortic stenosis; monitor renal function before and during treatment; renal impairment (reduce dose; liver impairment (Appendix 7a); possibly increased risk of agranulocytosis in collagen vascular disease; history of idiopathic or hereditary angioedema (use with care or avoid); lactation; interactions; hypervolemia; patients with immunosuppression; hyperkalemia. Risk of very rapid fall in blood pressure in volume-depleted patients; treatment should therefore be initiated with very low doses. High-dose diuretic therapy (furosemide dose greater than 80 mg) should be discontinued, or dose significantly reduced, at least 24 h before starting enalapril (may not be possible in heart failure-risk of pulmonary oedema). If high-dose diuretic cannot be stopped, medical supervision advised for at least 2 h after administration or until blood pressure stable. Avoid enalapril during dialysis with high-flux polyacrylonitrile membranes and during lowdensity lipoprotein apheresis with dextran sulphate ; also withhold before desensitization with wasp or bee venom.

**ADVERSE EFFECTS**
Hypokalaemia; hypomagnesaemia; hyponatraemia; hypochloraemic alkalosis (for symptoms of fluid and electrolyte imbalance; see introductory notes); increased calcium excretion; hypovolaemia; hyperglycaemia (but less often than with thiazide diuretics); temporary increase in plasma cholesterol and triglyceride concentration; less commonly hyperuricaemia and gout; rarely, rash; photosensitivity; bone marrow depression (withdraw treatment); pancreatitis (with large parenteral doses); tinnitus and deafness (with rapid administration of large parenteral doses and in renal impairment; deafness may be permanent if other ototoxic drugs taken);
gastrointestinal upset; malaise; blood dyscrasias; vertigo; orthostatic hypotension; jaundice; tinnitus; renal calcification in premature infants.

**Hydrochlorothiazide**

**EDL-D 259,260 PHC**

**AVAILABILITY**

TABLETS 12.5, 25 and 50 mg

**DOSE**

Oral Adult- Hypertension: 12.5 to 25 mg daily. Heart failure: initially 25 mg daily on waking up, increasing to 50 mg daily if necessary. Elderly- Initially 12.5 mg daily for hypertension as well as heart failure.

**INDICATION**

Alone in mild hypertension and in combination with other drugs in moderate to severe hypertension; heart failure; oedema; diabetes insipidus

**CONTRAINDICATION**

Severe renal or severe hepatic impairment; hyponatraemia; hypercalcaemia; refractory hypokalaemia; symptomatic hyperuricaemia; Addison’s disease; gout; diabetes mellitus; persisting hypercalcaemia; anuria; sulphonamide allergy.

**PRECAUTION**

Renal and hepatic impairment (Appendix 7a); lactation (Appendix 7b); elderly (reduce dose); may cause hypokalaemia; may aggravate diabetes mellitus and gout; may exacerbate systemic lupus erythematosus; porphyria; severe heart failure; edema; hyperlipidemia; interactions (Appendix 6a, 6b, 6c); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Fluid and electrolyte imbalance leading to dry mouth; thirst; gastrointestinal disturbances (including nausea; vomiting); weakness; lethargy; drowsiness; seizures; headache; muscle pains or cramps; hypotension (including postural hypotension); arrhythmias; hypokalaemia; oliguria; hypomagnesaemia; hyponatraemia; hypochloraeemic alkalosis; hypercalcaemia; hyperglycaemia; hyperuricaemia; gout; rash; photosensitivity; altered plasma lipid concentration; rarely, impotence (reversible); blood disorders (including neutropenia; thrombocytopenia); pancreatitis; intrahepatic cholestasis; acute renal failure; hypersensitivity reactions (pneumonitis; pulmonary oedema; severe skin reactions); increased heart rate and ventricular ectopic activity

**Nifedipine**

**EDL-D 366, 367 Secondary hospitals**

**D 368 PHC**

**AVAILABILITY**

TABLETS 5, 10, 20 and 30 mg plain and SR; CAPSULES 5, 10, 20 and 30 mg.

**DOSE**

Oral Adult- Hypertension (as sustained-release tablets): usual range 20 to 100 mg daily in 1 to 2 divided doses.

**INDICATION**

Hypertension; angina prophylaxis; heart failure; Raynaud’s phenomenon.

**CONTRAINDICATION**

Cardiogenic shock, advanced aortic stenosis, within 1 month of myocardial infarction, unstable or acute attacks of angina, porphyria; hypersensitivity.

**PRECAUTION**

Stop if ischaemic pain occurs or existing pain worsens shortly after starting treatment; poor cardiac reserve; heart failure or significantly impaired left ventricular function; monitor drug response in cirrhosis patients; blood pressure monitoring; calcium channel blockers; reduce dose in hepatic impairment; diabetes mellitus; may inhibit labour; lactation (Appendix 7b); pregnancy (Appendix 7c); interactions (Appendix 6b, 6c).
ADVERSE EFFECTS
Headache; flushing; dizziness; lethargy; tachycardia; palpitations; gravitational oedema (only partly responsive to diuretics); rash (erythema multiforme reported); pruritus; urticaria; nausea; constipation or diarrhoea; increased frequency of micturition; eye pain; visual disturbances; gum hyperplasia; paraesthesia; myalgia; tremor; impotence; gynaecomastia; depression; telangiectasis; cholestasis; jaundice; exacerbated angina; cardiovascular collapse; ankle swelling; gastrointestinal upset; reversible gingival hyperplasia.

Nor adrenaline
EDL-D 373 PHC

AVAILABILITY
INJECTIONS Vials (4 mg/ml, 4 mg/2 ml and 2 mg/2 ml)

DOSE
Parenteral Intravenous Acute hypotension Adult: 8-12 μg/minute, up to 8-30 μg/minute in refractory shock. Infused using a solution of 4 μg/ml in glucose 5%, or sodium chloride 0.9% and glucose 5% at a rate of 2-3 ml/minute. Adjust according to blood pressure response. Average maintenance dose: 0.5-1 ml/minute (2-4 μg/minute). Infuse via a central venous catheter or into a large vein. Child: Administer at a rate of 0.5 μg/minute. Alternatively, 2 μg/m2/minute. Adjust rate according to BP response and perfusion. Elderly: Initial dose should be at low end of dose range. Upper gastrointestinal haemorrhage Adult: 8 mg in 250 ml of 0.9% sodium chloride injection via intraperitoneal route. Alternatively, instill 8 mg in 100 ml of 0.9% sodium chloride solution through a nasogastric tube every hr for 6-8 hrs, then every 2 hrs for 4-6 hrs. Withdraw drug gradually. Reconstitution Dilute with 5% glucose injection, with or without sodium chloride; dilution with sodium chloride injection alone is not recommended.

INDICATION
Acute hypotension, adjunct in cardiac arrest, upper gastrointestinal haemorrhage.

CONTRAINDICATION
Hypertension, pregnancy (Appendix 7c), patients with peripheral or mesenteric vascular thrombosis unless necessary as a life-saving procedure. During cyclopropane and halothane anaesthesia, noradrenaline is considered contraindicated because of the risk of producing ventricular tachycardia or fibrillation

PRECAUTION
Monitor BP frequently during infusion, Use large vein for infusion to avoid skin necrosis, interactions

ADVERSE EFFECTS
Elevation of blood pressure, bradycardia, peripheral ischemia, arrhythmias, anxiety, transient headache, respiratory difficulty, extravasation necrosis at injection site.

Metoprolol
EDL-D 341,342 Secondary hospitals 343 Tertiary

AVAILABILITY
TABLETS 10, 25, 50 and 100 mg; CAPSULE 12.5, 25, 50 and 100 mg; INJECTION 100 mg/2 ml, 250 mg/2 ml, 500 mg/2 ml.

DOSE
Oral Heart failure: Initiating dose 12.5 - 25 mg once a day, Maximum dose: 200 mg once a day; Hypertension: initially 100 mg daily, increase if required to 200 mg in two divided doses (max 400 mg daily). Angina: 50 mg daily, up to 300 mg daily in 2 to 3 divided doses if necessary. Intravenous injection Arrhythmia: up to 5 mg at a rate of 1 to 2 mg per min, repeated after 5 min if necessary (max dose 10 to 15 mg). Arrhythmia developing during anaesthesia: 2 to 4 mg during induction.

INDICATION
Supraventricular arrhythmia, angina pectoris, hypertension, myocardial infarction; migraine prophylaxis; hyperthyroidism, heart failure
CONTRAINDICATION
Asthma (important: see Bronchospasm below), uncontrolled heart failure, Prinzmetal’s angina, marked bradycardia, hypotension, sick sinus syndrome, secondor third- degree AV block, cardiogenic shock, metabolic acidosis, severe peripheral arterial disease; pheochromocytoma (apart from specific use with alpha-blockers). Beta-blockers, including those considered to be cardioselective, should not be given to patients with a history of asthma or bronchospasm. However, in rare situations where there is no alternative a cardioselective beta-blocker is given to these patients with extreme caution and under specialist supervision.

PRECAUTION
Avoid abrupt withdrawal especially in ischaemic heart disease, first-degree AV block, portal hypertension (risk of deterioration in liver function); diabetes; history of obstructive airways disease (introduce cautiously and monitor lung function); myasthenia gravis; symptoms of hypoglycaemia and thyrotoxicosis may be masked; history of hypersensitivity-may increase sensitivity to allergens and result in more serious hypersensitivity response; also may reduce response to adrenaline (epinephrine); reduce dose of oral propranolol in hepatic impairment; renal impairment; lactation; pregnancy (Appendix 7c)

ADVERSE EFFECTS
Gastro-intestinal disturbances; bradycardia, heart failure, hypotension, conduction disorders; peripheral vasoconstriction (including exacerbation of intermittent claudication and Raynaud’s phenomenon); bronchospasm; dyspnoea; headache; fatigue; sleep disturbances; paraesthesia; dizziness; vertigo; psychosis; sexual dysfunction; purpura; thrombocytopenia; visual disturbances; exacerbation of psoriasis; alopecia; rarely, rashes and dry eyes (reversible on withdrawal); on infusion venous irritation and thrombophlebitis; agranulocytosis; hypoglycemia; myocardial depression

Labetolol
EDL-D299, 300 Secondary hospitals

INDICATION
Systemic hypertension, hypertensive emergencies, phaeochromo-cytoma.

AVAILABILITY
Tablets. 50 mg, 100 mg, 200 mg.

DOSE
50 mg b.d., increased to 100 - 200 mg b.d.

CONTRAINDICATION
AV Block, bronchospasm, cardiogenic shock.

PRECAUTION
CHF, diabetes mellitus, liver dysfunction, postural hypotension.

ADVERSE EFFECT
Headache, hallucination, impotence.

DRUG INTERACTION
Action of oral hypoglycemic agents increased, with anaesthetic agents may cause myocardial depression.

Telmisartan
EDL-D743,744 Secondary hospitals

AVAILABILITY
TABLETS 20, 40 and 80 mg.

DOSE
Adult- 40-80 mg once daily

INDICATION
Hypertension.

CONTRAINDICATION
Hypersensitivity,Renal artery stenosis, pregnancy (Appendix 7c) , hyperkalemia.
PRECAUTION
   Interactions (Appendix 6c)
ADVERSE EFFECTS
   Dizziness, drowsiness, fatigue, dyspnoea, blurred vision, postural hypotension, asthenia, nasal congestion, miosis, chest pain, urinary frequency, weight gain, thrombocytopenia, decreased libido, back pain and pain in extremities.

S-Amlodipine
   EDL-D 730,731 Secondary hospitals
AVAILABILITY
   TABLETS 1.25, 2.5, 5, 7.5, 10 and 20 mg.
DOSE
   Oral Angina: Adult- Initially 5 mg once daily, increased if necessary; max. 10 mg once daily.
   Hypertension: Adult- Initially 5 mg once daily, increased if necessary; max. 10 mg once daily.
   Elderly- Initial dose- 2.5 mg once daily.
INDICATION
   Angina, hypertension, coronary artery disease.
CONTRAINDICATION
   Significant aortic stenosis, sinoatrial node disease, hypersensitivity to dihydropyridines, cardiogenic shock, unstable angina; interactions (Appendix 6d)
PRECAUTION
   Hypotension, myocardial infarction, impaired renal function sick-sinus syndrome, severe ventricular dysfunction, hypertrophic cardiomyopathy, severe aortic stenosis, elderly, children, pregnancy (Appendix 7c); lactation; hepatic impairment (Appendix 7a)
ADVERSE EFFECTS
   Arrhythmias, postural hypotension; dizziness, ankle edema, hypoesthesia, flatulence, dizziness, blurred vision, facial flushing, dyspnoea, asthenia, muscle cramps, conduction system delay, abdominal pain, headache; sleep disturbances, fatigue.

Medicines used in heart failure
Treatment of heart failure aims to relieve symptoms, improve exercise tolerance, reduce incidence of acute exacerbations and reduce mortality. Drugs used to treat heart failure due to left ventricular systolic dysfunction include ACE inhibitors, diuretics, β-blockers (metoprolol, carvedilol and bisoprolol), cardiac glycosides and vasodilators. In addition, measures such as weight reduction, moderate salt restriction and appropriate exercise should be introduced. The primary treatment of heart failure is with ACE inhibitors such as enalapril which can be used in all stages of chronic heart failure to prevent further deterioration and progression of heart disease.
A thiazide diuretic such as hydrochlorothiazide is used in the management of mild to moderate heart failure when the patient has mild fluid retention and severe pulmonary oedema is not present; however thiazides are ineffective if renal function is poor. In these patients and in more severe fluid retention, a loop diuretic such as furosemide is required. In severe fluid retention, intravenous furosemide produces relief from breathlessness and reduces preload sooner than would be expected from the time of onset of diuresis. Hypokalaemia may develop, but is less likely with the shorter-acting loop diuretics than with the thiazides; care is needed to avoid hypotension.
A combination of a thiazide and a loop diuretic may be required to treat refractory oedema. The combination often produces a synergistic effect on solute and water excretion, which relieves symptoms in the diuretic-resistant heart failure patient. However, the combination may produce excessive intravascular volume depletion and electrolyte disturbances including potentially life-threatening hypokalaemia.

The aldosterone antagonist spironolactone may be considered for patients with severe heart failure who are already receiving an ACE inhibitor and a diuretic; a low dose of spironolactone (usually 25 mg daily) reduces symptoms and mortality rate in these patients. Close monitoring of serum creatinine and potassium is necessary with any change in treatment or in the patient’s clinical condition.

Digoxin, a cardiac glycoside, increases the strength of cardiac muscle contractions and increases cardiac output. In mild heart failure, digoxin inhibits the sympathetic nervous system and produces arterial vasodilation. It produces symptomatic improvement, increases exercise tolerance and reduces hospitalization, but it does not reduce mortality. It is considered for patients with atrial fibrillation and those who remain symptomatic despite treatment with an ACE inhibitor, a diuretic and a suitable beta-blocker.

Vasodilators are used in heart failure to reduce systemic vascular resistance. Isosorbide dinitrate produces mainly venous dilatation, which reduces left ventricular preload, leading to a reduction in pulmonary congestion and dyspnoea. Hydralazine produces mainly arterial vasodilation, which reduces left ventricular afterload and increases stroke volume and cardiac output. Isosorbide dinitrate and hydralazine can be used in combination when an ACE inhibitor cannot be used. Dopamine, an inotropic sympathomimetic, may be given for short periods in the treatment of severe heart failure. Dosage is critical; at low doses it stimulates myocardial contractility and increases cardiac output, however, higher doses (more than 5 μg/kg per min) cause vasoconstriction, with a worsening of heart failure.

**Enalapril**

**EDL-D 197,198 Secondary hospitals**

**AVAILABILITY**

- TABLETS 2.5, 5 and 10 mg; INJECTION 1 ml ampoule (1.25 mg/ml).

**DOSE**

- Oral Adult- Hypertension: initially 5 mg once daily; if used in addition to diuretic. Heart failure, asymptomatic left ventricular dysfunction: initially 2.5 mg daily under close medical supervision; usual maintenance dose 20 mg daily in 1 to 2 divided doses. Elderly- Renal impairment: initially 2.5 mg daily. Usual maintenance dose 10 to 20 mg once daily; In severe hypertension may be increased to max. 40 mg once daily.

**INDICATION**

- Heart failure (with a diuretic); prevention of symptomatic heart failure and prevention of coronary ischaemic events in patients with left ventricular dysfunction; hypertension; renal hypertension.

**CONTRAINDICATION**

- Renal failure with anuria; precomatose states associated with liver cirrhosis; severe sodium and water depletion; hypersensitivity to sulphonamides and furosemide; hypokalaemia; addison’s disease; lactation.
PRECAUTION
Use with diuretics; hypotension with first doses; especially in patients on diuretics; on a low-Sodium diet; on dialysis; if dehydrated; or with heart failure; peripheral vascular disease or generalized atherosclerosis (risk of clinically silent renovascular disease); use with great care in severe or symptomatic aortic stenosis; monitor renal function before and during treatment; renal impairment (reduce dose; liver impairment (Appendix 7a); possibly increased risk of agranulocytosis in collagen vascular disease; history of idiopathic or hereditary angioedema (use with care or avoid); lactation; interactions; hypervolemia; patients with immunosuppression; hyperkalemia. Risk of very rapid fall in blood pressure in volume-depleted patients; treatment should therefore be initiated with very low doses. High-dose diuretic therapy (furosemide dose greater than 80 mg) should be discontinued, or dose significantly reduced, at least 24 h before starting enalapril (may not be possible in heart failure-risk of pulmonary oedema). If high-dose diuretic cannot be stopped, medical supervision advised for at least 2 h after administration or until blood pressure stable. Avoid enalapril during dialysis with high-flux polycrilonitrile membranes and during low-density lipoprotein apheresis with dextran sulphate; also withhold before desensitization with wasp or bee venom.

ADVERSE EFFECTS
Hypokalaemia; hypomagnesaemia; hyponatraemia; hypochloraemic alkalosis (for symptoms of fluid and electrolyte imbalance; see introductory notes); increased calcium excretion; hypovolaemia; hyperglycaemia (but less often than with thiazide diuretics); temporary increase in plasma cholesterol and triglyceride concentration; less commonly hyperuricaemia and gout; rarely, rash; photosensitivity; bone marrow depression (withdraw treatment); pancreatitis (with large parenteral doses); tinnitus and deafness (with rapid administration of large parenteral doses and in renal impairment; deafness may be permanent if other ototoxic drugs taken); gastrointestinal upset; malaise; blood dyscrasias; vertigo; orthostatic hypotension; jaundice; tinnitus; renal calcification in premature infants.

Dobutamine
EDL-D 187 Secondary hospitals

AVAILABILITY
INJECTION 250 mg/20 ml, 40 mg/ml, 12.5 mg/ml, 5 ml ampoule (50 mg/ml), vial 250 mg/20 ml, 50 mg/4 ml; 250 mg dry sterile lyophilised powder.

DOSE
2.5 to 10 μg/kg/min which can be titrated to 40 μg/kg/min as per the individual requirement.

INDICATION
Acute heart failure; acute myocardial infarction; cardiogenic shock following cardiac surgery; specific shock; acute decompensation of chronic CHF.

CONTRAINDICATION
Hypersensitivity; idiopathic hypertrophic subaortic stenosis

PRECAUTION
Interactions (Appendix 6c); pregnancy (Appendix 7c); monitor heart rate and rhythm; arterial BP and infusion rate closely; correct hypovolemia prior to treatment; elderly; neonates; risk of rapid ventricular response

ADVERSE EFFECTS
Interactions; pregnancy; monitor heart rate and rhythm; arterial BP and infusion rate closely; correct hypovolemia prior to treatment; elderly; neonates; risk of rapid ventricular response in patients with atrial fibrillation; children.
Furosemide

EDL-D 239,240 PHC

AVAILABILITY
TABLETS 40, 100 and 500 mg; injection ampoule 20 mg/ml, 10 mg/2 ml, 250 mg/25 ml, 20 mg/2 ml.

DOSE
Oral Adult- Oedema: initially 40 mg daily on waking up. Maintenance. 20 to 40 mg daily; may be increased to 80 mg daily or more in resistant oedema: max 600 mg daily in severe cases. Child- 1 to 3 mg/kg daily (max. 40 mg daily). Slow intravenous injection Adult- Acute pulmonary oedema: 20 to 50 mg, if necessary increase by 20 mg step-bystep every 2 h; if effective single dose is more than 50 mg, at a rate not exceeding 4 mg/ min. Child- 0.5 to 1.5 mg/kg daily (max. 20 mg daily). Slow intravenous infusion Adult- Oliguria (glomerular filtration rate less than 20 ml/min): at a rate not exceeding 4 mg/min, initially 250 mg over 1 h. If urine output not satisfactory during the h after first dose, infuse 500 mg over 2 h then; if no satisfactory response is there in an h after second dose, infuse 1g over 4 h. If no response is there after third dose, dialysis is probably necessary.

INDICATION
Oedema; mild to moderate hypertension.

CONTRAINDICATION
Renal failure with anuria; precomatose states associated with liver cirrhosis; hypersensitivity.

PRECAUTION
Monitor electrolytes particularly potassium and Sodium; hypotension; elderly (reduce dose); pregnancy (Appendix 7c); lactation correct hypovolaemia before using in oliguria; renal impairment; hepatic impairment (Appendix 7a); prostatic enlargement; porphyria; interactions (Appendix 6b, 6c); gout; impaired micturition; infusion rate should not exceed 4 mg/min to reduce the risk of ototoxicity; monitor serum levels for calcium or magnesium (may be lowered).

ADVERSE EFFECTS
Hypokalaemia; hypomagnesaemia; hyponatraemia; hypochloraemic alkalosis (for symptoms of fluid and electrolyte imbalance; see introductory notes); increased calcium excretion; hypovolaemia; hyperglycaemia (but less often than with thiazide diuretics); temporary increase in plasma cholesterol and triglyceride concentration; less commonly hyperuricaemia and gout; rarely, rash; photosensitivity; bone marrow depression (withdraw treatment); pancreatitis (with large parenteral doses); tinnitus and deafness (with rapid administration of large parenteral doses and in renal impairment; deafness may be permanent if other ototoxic drugs taken); hepatic encephalopathy, anorexia, orthostatic hypotension.

Ramipril

EDL-D 726,727 Secondary hospitals

AVAILABILITY
TABLETS AND CAPSULES 1.25, 2.5, 5 and 10 mg.

DOSE
Reduction in risk of myocardial infarction, stroke, and death from cardiovascular causes: Initial dose of 2.5 mg, once a day for 1 week, 5 mg, once a day for the next 3 weeks, and then increased as tolerated, to a maintenance dose of 10 mg once a day. Hypertension: The recommended initial dose for patients not receiving a diuretic is 2.5 mg once a day. The usual maintenance dosage range is 2.5 to 20 mg per day administered as a single dose or in two equally divided doses. Heart failure post myocardial infarction: Initial dose is 2.5 mg twice daily, after one week at the starting dose titrate to ( if tolerated) toward a target dose of 5 mg twice daily, with dosage increases being about 3 weeks apart.
**INDICATION**
Reduction in risk of myocardial infarction, stroke and death from cardiovascular causes; hypertension; heart failure post myocardial infarction

**CONTRAINDICATION**
Hypersensitivity to ramipril or any other ACE inhibitor, bilateral renal artery stenosis or a single kidney with unilateral renal artery stenosis

**PRECAUTION**
Impaired renal function, impaired liver function, diabetes mellitus (increased risk of hyperkalemia), patients undergoing surgery, history of angioedema; symptomatic hypotension is most likely to occur in patients who have been volume- and/or salt-depleted as a result of prolonged diuretic therapy, dietary salt restriction, dialysis, diarrhoea, or vomiting. Volume and/or salt depletion should be corrected before initiating therapy; monitoring of white blood cell counts should be considered in patients with collagenvascular disease, especially if the disease is associated with impaired renal function; administration during pregnancy (Appendix 7c) can cause fetal/neonatal morbidity and death; when pregnancy is detected ACE inhibitors should be discontinued as soon as possible, interactions (Appendix 6a and 6c).

**ADVERSE EFFECTS**
Hypotension, cough, asthenia, dizziness, headache, angioneurotic edema, hypersensitivity reactions, erythema multiforme, toxic epidermal necrolysis, Stevens Johnson syndrome, hepatic necrosis, pancreatitis, pancytopenia, thrombocytopenia.

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**Dopamine**

**EDL-D 190 Secondary hospitals**

**INDICATIONS**
Cardiogenic shock in myocardial infarction or cardiac surgery; acute heart failure.

**AVAILABILITY**
Injection 5 ml vial (40 mg/ml), 5 and 10 ml ampoule (200 mg/5 ml).

**DOSE**
Intravenous infusion
Adult- Cardiogenic shock: into large vein, initially 2 to 5 μg/kg/min; gradually increased by 5 to 10 μg/kg/min according to blood pressure, cardiac output and urine output; seriously ill patients up to 20 to 50 μg/kg/ min. By intravenous route initially 1 to 5 μg/ kg/min can be increased gradually to 5 to 10 μg/kg/min. max 20 to 50 μg/kg/min in serious patients.

**CONTRAINDICATIONS**
Hypersensitivity; tachyarrhythmias, ventricular fibrillation, ischaemic heart disease; pheochromocytoma; hyperthyroidism.

**PRECAUTIONS**
Correct hypovolaemia before and maintain blood volume during treatment; correct hypoxia; hypercapnia and metabolic acidosis before or at same time as starting treatment; low dose in shock due to myocardial infarction; history of peripheral vascular disease (increased risk of ischaemia of extremities); elderly; interactions (Appendix 6c); history of atherosclerosis; Raynaud’s disease; diabetic endocarditis; disproportionate increase in diastolic pressure; pregnancy (Appendix 7c); lactation; paediatrics. Dopamine must be diluted before i.v. administration.

**ADVERSE EFFECTS**
Nausea and vomiting; peripheral vasoconstriction; hypotension with dizziness; fainting; flushing; tachycardia; ectopic beats; palpitations; anginal pain; headache; dyspnoea; hypertension particularly in overdosage.

**STORAGE**
Store in an airtight container protected from light.
**Digoxin**  
*EDL-D 178,179,180 Secondary hospitals*

**AVAILABILITY**  
TABLET 0.25 mg; INJECTION 2 ml (0.5 mg/2 ml); ELIXIR 0.05 mg/ml (paediatric use); SYRUP 1.5 mg/30 ml.

**DOSE**  
Oral Adult - Atrial fibrillation and heart failure: 1 to 1.5 mg in divided doses over 24 h for rapid digitalization or 250 μg 1 to 2 times daily if digitalization less urgent; maintenance 62.5 to 500 μg daily (higher dose may be divided), according to renal function and heart rate response; usual range 125 to 250 μg daily. Elderly - Lower dose more appropriate. Intravenous infusion Emergency control of atrial fibrillation, over at least 2 h: 0.75 to 1 mg. Emergency loading dose for heart failure, over at least 2 h: 0.75 to 1 mg.

**INDICATION**  
Supraventricular arrhythmias, particularly atrial fibrillation; heart failure.

**CONTRAINDICATION**  
Hypertrophic obstructive cardiomyopathy (unless also atrial fibrillation and heart failure); ventricular tachycardia; hypokalaemia; digitalis toxicity; arrhythmias; Wolff-Parkinson-White syndrome or other accessory pathway, particularly if accompanied by atrial fibrillation; intermittent complete heart block; seconddegree atrioventricular block.

**PRECAUTION**  
Recent myocardial infarction; sick sinus syndrome; severe pulmonary disease; thyroid disease; congestive cardiac myopathy; hypercalcaemia; aortic valve disease, heart block, cardiac dysrythmias; elderly (reduce dose); renal impairment (Appendix 7d); avoid hypokalaemia; avoid rapid intravenous administration (nausea and risk of arrhythmias); lactation; interactions (Appendix 6c, 6d); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**  
Usually associated with excessive dosage and include anorexia, nausea, vomiting, diarrhoea, abdominal pain; visual disturbances, headache, fatigue, drowsiness, confusion, delirium, hallucinations, depression; arrhythmias, heart block; rarely, rash, intestinal ischaemia; gynaecomastia on long-term use; thrombocytopenia reported; sinus bradycardia; apathy; psychosis; malaise.

**Hydrochlorothiazide**  
*EDL-D 259,260 PHC*

**AVAILABILITY**  
TABLETS 12.5, 25 and 50 mg

**DOSE**  
Oral Adult - Hypertension: 12.5 to 25 mg daily. Heart failure: initially 25 mg daily on waking up, increasing to 50 mg daily if necessary. Elderly - Initially 12.5 mg daily for hypertension as well as heart failure.

**INDICATION**  
Alone in mild hypertension and in combination with other drugs in moderate to severe hypertension; heart failure; oedema; diabetes insipidus

**CONTRAINDICATION**  
Severe renal or severe hepatic impairment; hyponatraemia; hypercalcaemia; refractory hypokalaemia; symptomatic hyperuricaemia; Addison’s disease; gout; diabetes mellitus; persisting hypercalcaemia; anuria; sulphonamide allergy.

**PRECAUTION**  
Renal and hepatic impairment (Appendix 7a); lactation (Appendix 7b); elderly (reduce dose); may cause hypokalaemia; may aggravate diabetes mellitus and gout; may exacerbate systemic
lupus erythematosus; porphyria; severe heart failure; edema; hyperlipidemia; interactions (Appendix 6a, 6b, 6c); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Fluid and electrolyte imbalance leading to dry mouth; thirst; gastrointestinal disturbances (including nausea; vomiting); weakness; lethargy; drowsiness; seizures; headache; muscle pains or cramps; hypotension (including postural hypotension); arrhythmias; hypokalaemia; oliguria; hypomagnesaemia; hypochloraemia; hyperchloremia; hyperglycaemia; hyperuricaemia; gout; rash; photosensitivity; altered plasma lipid concentration; rarely, impotence (reversible); blood disorders (including neutropenia; thrombocytopenia); pancreatitis; intrahepatic cholestasis; acute renal failure; hypersensitivity reactions (pneumonitis; pulmonary oedema; severe skin reactions); increased heart rate and ventricular ectopic activity.

**Drugs used in Eclampsia**

**Hypertension in Pregnancy**

This is defined as a sustained diastolic blood pressure of 90mmHg or more. Drug therapy for chronic hypertension during pregnancy remains controversial. If diastolic blood pressure is greater than 95 mmHg, methyldopa is the safest drug. Betablockers should be used with caution in early pregnancy, since they may retard fetal growth; they are effective and safe in the third trimester. ACE inhibitors are contraindicated in pregnancy since they may damage fetal and neonatal blood pressure control and renal function. Women who are taking these drugs and become pregnant should have their antihypertensive therapy changed immediately.

Pre-eclampsia and eclampsia: If pre-eclampsia or severe hypertension occurs beyond the 36th week of pregnancy, delivery is the treatment of choice. For acute severe hypertension in preeclampsia or eclampsia, intravenous hydralazine can be used. Magnesium sulphate is the treatment of choice to prevent eclamptic convulsions in eclampsia and severe pre-eclampsia.

**Magnesium Sulphate**

**EDL-D 321 Universal**

**INDICATIONS**

Prevention of recurrent seizures in eclampsia; prevention of seizures in pre-eclampsia; acute nephritis in children.

**AVAILABILITY**

**INJECTION** 500 mg/ml.

**DOSE**

Intravenous injection (concentration of magnesium sulphate should not exceed 20%) Prevention of seizure occurrence in eclampsia: initially 4g over 5 to 15 min, followed by infusion 1g/hr for at least 24 h after last seizure. If seizures recur, additional dose of 2g (or 4g if body weight is over 70 kg).

**CONTRAINDICATIONS**

Not to be injected parenterally in patients with heart block or myocardial damage.

**PRECAUTIONS**

Hepatic impairment (Appendix 7a); pregnancy (Appendix 7c); renal impairment; in severe hypomagnesaemia administer initially via controlled infusion device (preferably syringe pump);
monitor blood pressure, respiratory rate, urinary output and for signs of overdosage (loss of patellar reflexes, weakness, nausea, sensation of warmth, flushing, drowsiness, double vision and slurred speech).

ADVERSE EFFECTS
Generally associated with hypermagnesaemia, nausea, vomiting, thirst, flushing of skin, hypotension, arrhythmias, coma, respiratory depression, drowsiness and confusion, loss of tendon reflexes, muscle weakness; colic and diarrhoea following oral administration; hypothermia; stupor.

**Methyldopa**  
**EDL-D 337 PHC**

**INDICATIONS**  
Hypertension in pregnancy.

**AVAILABILITY**  
TABLET 250 mg.

**DOSE**  
Oral  
Adult- Hypertension in pregnancy: initially 250 mg 2 to 3 times daily; if necessary, gradually increased at intervals of 2 or more days (max 3g daily).

**CONTRAINDICATIONS**  
Depression; active liver disease; hypersensitivity; therapy with MAO inhibitors; pheochromocytoma; porphyria.

**PRECAUTIONS**  
History of hepatic impairment (Appendix 7a); renal impairment; blood counts and liverfunction tests advised; history of depression; positive direct Coomb test in up to 20% of patients (affects blood cross-matching); interference with laboratory tests; lactation; pregnancy (Appendix 7c); interactions (Appendix 6b, 6c). May impair ability to perform skilled tasks; for example operating machinery; driving.

**ADVERSE EFFECTS**  
Tend to be transient and reversible including sedation; dizziness; lightheadedness; postural hypotension; weakness; fatigue; headache; fluid retention and oedema; sexual dysfunction; impaired concentration and memory; depression; mild psychosis; disturbed sleep and nightmares; drug fever; influenza-like syndrome; nausea; vomiting; constipation; diarrhoea; dry mouth; stomatitis; sialadenitis; liver function impairment; hepatitis; jaundice; rarely, fatal hepatic necrosis; bonemarrow depression; haemolytic anaemia; leukopenia; thrombocytopenia; eosinophilia; parkinsonism; rash (including toxic epidermal necrolysis); nasal congestion; black or sore tongue; bradycardia; exacerbation of angina; myalgia; arthralgia; paraesthesia Bell palsy; pancreatitis; hypersensitivity reactions including lupus erythematosuslike syndrome; myocarditis; pericarditis; gynaecomastia; hyperprolactinaemia; amenorrhoea; urine darkens on standing.
Lipid lowering agent
Drug therapy to lower plasma lipids should be used in addition to dietary management and correction of other modifiable cardiovascular risk factors. Studies indicate that, 1% drop in serum cholesterol reduces the risk for Coronary heart disease (CHD) by 2%.
Various classes of drugs used as lipid lowering drugs are-

**H mg-CoA reductase inhibitors**
They are the most efficacious and tolerable drugs like simvastatin, pravastatin, atorvastatin etc. They are primarily indicated in secondary prevention of myocardial infarction and stroke in patients who have symptomatic atherosclerotic disease following acute myocardial infarction or stroke and in primary prevention of arterial disease in patients who are at high risk because of elevated serum cholesterol concentration. Common adverse effects include mild gastrointestinal disturbances, rhabdomyolysis etc.

**Fibric acid derivatives**
This class of drugs including fenofibrate, gemfibrozil etc are indicated in patients with mixed dyslipidemia (i.e. raised serum triglycerides and cholesterol), low high density lipoprotein (HDL) and high risk of atheromatous disease (often type 2 diabetic patients), and in severe treatment-resistant dyslipidemia. Major adverse effect include rhabdomyolysis and myoglobulinuria. Fibrates are better avoided in alcoholics.

**Bile acid sequestrants**
Drugs like cholestyramine, colestipol though are not clinically popular because of interference with absorption of many drugs like digoxin, warfarin etc and poor patient acceptability, but can be indicated in heterozygous familiar hypercholesterolemia. Adverse effects include nausea, abdominal bloating, constipation or diarrhoea.

**Nicotinic acid**
Nicotinic acid reduces serum cholesterol and triglycerides levels in types II, III, IV, and V hyperlipoproteinemias. Adverse effects include flushing, palpitations and gastrointestinal tract disturbances.

**Atorvastatin**

EDL-D 563,564 Secondary hospitals

**AVAILABILITY**
TABLETS 5, 10, 20, 40 and 80 mg.

**DOSE**
Oral Adult- 10 mg daily, increased at 4 weeks interval. Max dose 80 mg.

**INDICATION**
Primary and secondary hypercholesterolemia, prevention of cerebrovascular accidents, primary prevention of coronary heart disease.

**CONTRAINDICATION**
Hypersensitivity; active liver diseases or unexplained persistent elevation of serum transaminase; pregnancy (Appendix 7c), lactation.

**PRECAUTION**
Patients who consume substantial quantities of alcohol and have a history of liver diseases, Children below 10 years, premenarcheal females; interactions (Appendix 6a, 6c)

**ADVERSE EFFECTS**
Myopathy is the serious adverse effect; headache; infrequent elevation of creatinine phosphokinase; rhabdomyolysis; insomnia; dizziness; abdominal pain, constipation, diarrhoea, dyspepsia, flatulence and nausea.
Fenofibrate
   EDL-D 635 Secondary hospitals

AVAILABILITY
   CAPSULES 67 and 200 mg, TABLETS 145 and 160 mg INJECTIONS 20, 40 and 60 mg/vial.

DOSE
   Hyperlipidemia: Adult- Initial dose 67 mg 2-4 times a day (micronized) or 200 mg/day in divided doses (non-micronized). Child- 5 mg/kg daily.

INDICATION
   Hypercholesterolemia, hypertriglyceridemia.

CONTRAINDICATION
   Hypersensitivity, severe renal and hepatic impairment, preexisting gall bladder disease, primary biliary cirrhosis, pregnancy (Appendix 7c), lactation.

PRECAUTION
   Pancreatitis; skeletal muscle effects; renal and hepatic impairment; monitor for LFT and blood counts regularly; interactions (Appendix 6c)

ADVERSE EFFECTS
   Myalgia; hepatitis; rashes; cholelithiasis, rhabdomyolysis; increased SGPT and SGOT, abdominal pain, photosensitivity; rhinitis; sinusitis
Antifungal medicines

Ringworm:
Benzoic acid and methylrosanilinium chloride (gentian violet) solution are inexpensive and effective fungistatic compounds for the treatment of dermatophyte infections such as ringworm. Minor skin lesions due to ringworm can be cleared with repeated applications of compound benzoic acid ointment (Whitfield ointment), which combines the fungistatic action of benzoic acid with the keratolytic action of salicylic acid. However, the most effective topical treatment for dermatophyte infections is a cream containing an imidazole such as miconazole, which is effective for long-established lesions but is more expensive than compound benzoic acid ointment. Extensive and generalized infections of the skin, nails and scalp should be treated systemically for several weeks with griseofulvin or fluconazole.

Scalp ringworm (Tinea capitis) typically appears as a patch of scaling alopecia, or a swollen inflammatory area (Tinea kerion). Mild forms may remit spontaneously at puberty. Inflamed lesions should be treated systemically with griseofulvin. Application of miconazole cream may accelerate healing of scaly lesions.

Ringworm on the body (Tinea corporis) can also be cleared with compound benzoic acid ointment or a topical imidazole such as miconazole. In resistant cases a 4-week course of oral griseofulvin is required.

Foot ringworm (Tinea pedis or athlete’s foot) is usually treated topically. Compound benzoic acid ointment should be applied twice daily to all infected areas and all toe clefts for at least 4 weeks. Systemic therapy with griseofulvin or fluconazole may be required if the foot is extensively infected. Tinea pedis commonly recurs and may be treated with miconazole cream. Severe weeping lesions respond to frequent soaking in solutions of 1:10,000 potassium permanganate and systemic antifungals may also be needed.

Nail infections (onychomycosis, tinea unguium) are difficult to treat; fingernails may require 6 months treatment with oral griseofulvin and toenails may require 12 months or more of this treatment. Approximately 60% of nail infections either do not respond or relapse after treatment with griseofulvin.

Ringworm of the groin (Tinea cruris) is usually limited to the skin of the inner thigh in contact with the scrotum. Flexural eczema, often superinfected with candida or bacteria, occurs in the same site. The latter is frequently treated with combined antifungal/corticosteroid preparations, but must not be treated with a corticosteroid alone, which will worsen the condition. An imidazole cream such as miconazole applied daily for 2 weeks is usually effective. Lesions unresponsive to topical preparations can usually be cleared with a 4-week course of griseofulvin.

Candidosis:
Candida can infect the oral cavity, the vagina or the skin. Cutaneous lesions tend to occur in patients with diabetes mellitus and some chronic debilitating conditions, including
hypoparathyroidism and various congenital disorders of the immune system. The most severe infections of candida are now seen in patients with HIV infection.

Cutaneous candidosis usually responds to miconazole cream as a twice daily application. Chronic candida paronychia, which can result ultimately in nail dystrophy, is more difficult to treat. Treatment should be based on determination of the underlying cause and its reduction or elimination; hands and folds of the nail must be kept dry and daily application of an imidazole cream for several months may be required, ensuring penetration of the cleft between the nail plate and the swollen skin around the nail.

**Pityriasis Versicolor:**

Pityriasis (tinea) versicolor is caused by a commensal yeast. Application of Sodium thiosulfate twice daily for 4 weeks is usually effective although areas of depigmentation on darker skins remain after completion of treatment. Relapses can be frequent, however, probably because much of the infected area may appear normal and be left untreated. Better results have been reported with topical applications of miconazole or selenium sulphide.

**Miconazole**

**EDL-D349 PHC**

**INDICATIONS**

Superficial fungal infections due to dermatophytes and yeasts, and secondary infections caused by Gram-positive cocci, including ringworm, intertrigo, candida napkin rash, paronychia, and pityriasis versicolor.

**AVAILABILITY**

CREAM 10 and 15g (2%); gel 2% w/w.

**DOSE**

Skin infections: apply twice daily to clean dry lesions, continuing for at least 10 days after the condition has cleared. Nail infections: apply 1 to 2 times daily.

**PRECAUTIONS**

Contact with eyes and mucous membranes should be avoided; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Occasional local irritation and burning; also contact dermatitis; discontinue if sensitization occurs.

**STORAGE**

Store protected from light and moisture at a temperature not exceeding 30°C. If it is packed in aluminates; the inner surface of tubes should be coated with suitable lacquer.
Anti-infective medicines
Staphylococcal infections of the skin such as impetigo, folliculitis, and furunculi and streptococcal infections such as cellulitis and erysipelas are very common where the climate is hot and humid, where standards of hygiene are compromised, and in immunodeficient patients.
In all skin infections, an important part of treatment is cleansing and thorough drying. Washing with soap and water will often help to prevent infection. Light localized infections can often be treated effectively with an antiseptic solution such as chlorhexidine.

Superficial crusts should be gently washed with soap and water or a weak solution of aluminium acetate or a 0.01% solution of potassium permanganate. Infected burns should be treated with silver sulfadiazine, which is bactericidal against both Gram-positive and Gram-negative organisms. An ointment containing 2% mupirocin, which is active against Gram-positive bacteria, is of value, particularly in impetigo. To prevent the development of resistance, mupirocin should not be used for more than 10 days. Topical preparations containing neomycin and bacitracin are also widely used but these carry a risk of sensitization particularly with continued or repeated use.

Topical use of preparations containing antimicrobials which are widely used systemically should be avoided. These include penicillins, sulfonamides, streptomycin and gentamicin, which should be reserved for the systemic treatment of infections because of the possibility of inducing sensitivity and favouring the emergence of resistant organisms. Only widespread superficial or deep-seated infections associated with fever require treatment with a systemic antibiotic. Whenever possible, the choice of an antimicrobial should be based on the results of sensitivity tests.

Acyclovir
EDL-D 8 Secondary hospitals

INDICATIONS
Treatment of Herpes simplex keratitis; long term suppression of skin infections in Herpes simplex as well as mucous membrane, prophylaxis in immunocompromised patients; Herpes zoster treatment.

AVAILABILITY
Ointment 5g (3% w/w); drops 5 ml (3% w/w).

DOSE
Adult- Herpes simplex keratitis: apply 3% w/w ointment 5 times daily for 3 days.

CONTRAINDICATIONS
Hypersensitivity.

PRECAUTIONS
Maintain adequate hydration (especially with infusion or high doses); monitor neutrophil count at least twice weekly in neonates; renal impairment (Appendix 7d); lactation (Appendix 7b); pregnancy (Appendix 7c); not to be applied on mucous membrane.

ADVERSE EFFECTS
Nausea, vomiting, abdominal pain, diarrhoea, headache, fatigue, rash, urticaria, pruritus, photosensitivity; very rarely, hepatitis, jaundice; dyspnoea; neurological reactions (including dizziness, confusion, hallucinations, convulsions and drowsiness); acute renal failure; anaemia, thrombocytopenia and leucopenia; on intravenous infusion; severe local inflammation (sometimes leading to ulceration), and very rarely, agitation, tremors; psychosis and fever; increase in blood urea and creatinine, encephalopathy; seizures; anorexia, tremors.
Framycetin Sulphate  
EDL-D 235 Universal  
AVAILABILITY 
Cream 1% - 5, 15 and 40g; Drops 5 ml (0.5%); Dressing 1%; Powder 15g.  
DOSE  
Topical Skin infections: Adult- as 1% dressing. Ophthalmic Blepharitis along with conjunctivitis: Adult- as 0.5 % ointment, apply 2-3 times daily. Otitis externa Adult- 0.5% drops.  
INDICATION  
Bacterial skin infections, burns, ENT infections, surgical infections, traumatic injury, conjunctivitis, blepharitis.  
CONTRAINDICATION  
Tuberculosis, glaucoma, perforated tympanic membrane, fungal, viral or resistant bacterial infections of eye, hypersensitivity.  
PRECAUTION  
Pregnancy, ototoxicity due to systemic absorption may occur if applied on large areas in children, elderly and patients with renal failure, avoid prolonged use, interactions  
ADVERSE EFFECTS  
Ototoxicity, gastrointestinal symptoms, inflammation, transient irritation, contact dermatitis, burning sensation, pruritus.  

Povidone Iodine  
EDL-D422,423 Universal  
INDICATIONS  
Antiseptic; skin disinfection; Mouth wash.  
AVAILABILITY  
SOLUTIONS 100 and 500 ml (5% w/v), 500 ml (7.5% w/v and 10% w/v); OINTMENT 15g (5% w/w).  
DOSE  
Adult and Child- Pre- and post-operative skin disinfection: apply undiluted. Antiseptic (minor wounds and burns): apply twice daily.  
CONTRAINDICATIONS  
Avoid regular or prolonged use in patients with thyroid disorders or those taking lithium; avoid regular use in neonates; avoid in very low birthweight infants; burn covering large surface area; hypersensitivity to iodine.  
PRECAUTIONS  
Pregnancy (Appendix 7c); lactation (Appendix 7b); broken skin (see below); renal impairment; avoid contact with eyes; neonates. The application of povidone iodine to large wounds or severe burns may produce systemic adverse effects such as metabolic acidosis; hypernatraemia; and impairment of renal function.  
ADVERSE EFFECTS  
Irritation of skin and mucous membranes; may interfere with thyroid function tests; systemic effects (see under Precautions).  
STORAGE  
Store protected from light.  

Silver Sulfadiazine  
EDL-D 474 Universal  
INDICATIONS  
Prophylaxis and treatment of infection in burns.
AVAILABILITY
CREAM 1%w/w.

DOSE
Infection in burns: apply using aseptic technique daily (more frequently if volume of exudate is large) whilst there is a possibility of infection, or until healing is complete. Contraindications
Hypersensitivity to sulfonamides; neonates; premature infants.

PRECAUTIONS
Renal or hepatic impairment; G-6-PD deficiency; lactation (Appendix 7b); monitor serum sulphadiazine concentration and check urine for sulpha crystals; pregnancy (Appendix 7c).

Adverse Effects
Allergic reactions include rashes; burning and itching; argyria and sulfonamideinduced systemic toxicity; including blood disorders following application to large areas or prolonged use; transient leukopenia; skin necrosis; skin discolouration.

STORAGE
Store protected from light.

Neomycin + Bacitracin
EDL-D 360 PHC

INDICATIONS
Superficial bacterial infections of the skin due to staphylococci and streptococci.

AVAILABILITY
CREAM 5, 10 and 15g (Aluminium tubes).

DOSE
Adult and child- Bacterial skin infections over 2 years: apply as a thin layer 3 times daily.

CONTRAINDICATIONS
Neonates

PRECAUTIONS
Avoid application to substantial areas of skin or to broken skin (risk of significant systemic absorption); overgrowth of resistant organisms on prolonged use; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Sensitization; especially to neomycin; causing reddening and scaling; systemic absorption leading to irreversible ototoxicity; particularly in children; elderly; and in renal impairment; pregnancy (Appendix 7c).

STORAGE
Store protected from light at temperature not exceeding 30°C.

Linezolid
Non-EDL Tertiary

Effective for treatment of resistant g+ve coccal(aerobic and anaerobic)and bacillary infections;MRSA, VRSA and VRE.G-ve bacteria not affected

INDICATION
Pneumonia, complicated skin and soft tissue infections caused by Gram+ve bacteria

AVAILABILITY
Tablets, 600mg;200 mg/100 ml infusion

DOSE
Adult Oral/I.V, 600mg twice daily for 10-14 days; upto 28 days in vancomycin-resistant cases

CONTRAINDICATION
Hypersensitivity to the drug, breast feeding.
PRECAUTION
Hepatic impairment; renal impairment; pregnancy; monitor full blood count including platelet count, avoid in uncontrolled hypertension, phaeochromocytoma, carcinoid tumor, thyrotoxicosis, bipolar depression, schizophrenia, or acute confusional states, concomitant use of other MAO inhibitors.

ADVERSE EFFECT
GI disturbances, Rash, pruritus, headache, thirst, dry mouth, glossitis, stomatitis, tongue discoloration, oral and vaginal candidiasis, leucopenia, thrombocytopenia

DRUG INTERACTION
Being MAO Inhibiter it interacts with adrenergic/serotonergic drugs

Benzoyl peroxide
EDL-D68, 69 Secondary hospitals

INDICATION
Mild to moderate acne and as an adjunct to oral therapy in more severe cases.

AVAILABILITY
Gel, 2.5% w/w, 5% w/w Cream, benzoyl peroxide 10% Lotion (cutaneous suspension), benzoyl peroxide 5%

PRECAUTIONS
avoid contact with eyes, mouth, and mucous membranes; avoid use of occlusive dressings

ADMINISTRATION
Initially apply to clean skin on alternate days, increasing frequency to 1–2 times daily as tolerance to irritant effect develops.

ADVERSE EFFECTS
initial irritation common but subsides with continued use; rarely, contact sensitivity occurs, occasionally even 1 application can cause severe irritation; may bleach fabrics, hair and skin

PSORIASIS. Psoriasis, which affects people of all ages in all countries, is one of the most common chronic dermatoses in industrialized regions, and is characterized by epidermal thickening and scaling. It needs specialisation to treat and the drugs for it are reserved for higher referral centres.

Podophyllum resin
EDL-D 616 Secondary Hospitals
An example of an application to treat warts. Various drugs can serve as alternatives Solution (cutaneous solution), podophyllum resin 10–25%.

INDICATION
external anogenital warts; plantar warts.

ADMINISTRATION
Medical supervision required; apply carefully to warts, avoiding contact with normal tissue; rinse off after 1–4 hours; may be repeated at weekly intervals but no more than 4 times in all; only few warts to be treated at any one time.

CONTRAINDICATIONS
pregnancy (Appendix 2); breastfeeding; children.

PRECAUTIONS
avoid use on large areas, mucous membranes; irritant to eyes; avoid contact with normal skin.

ADVERSE EFFECTS
systemic effects resulting from cutaneous absorption include nausea, vomiting, abdominal pain and diarrhoea; also transient leukopenia and thrombocytopenia; delayed neurotoxicity including visual and auditory hallucinations, delusions, disorientation, confusion and delirium following excessive application.
Permethrin
   EDL-D714,715 Secondary hospitals
INDICATION
Scabies; head and body lice
PRECAUTION
Do not use on inflamed or broken skin; avoid contact with eyes; breastfeeding (withhold during treatment)
ADVERSE EFFECT
Local irritation; rarely rashes and oedema
PRECAUTION
Cream: 5%; Lotion: 1% (Head lice); Lotion 5%
DOSE
Scabies and body lice apply cream over whole body and wash off after 8–12 hours; if hands washed with soap within 8 hours of application, treat again; repeat application after 7 days
Head lice, apply lotion to clean damp hair and rinse off after 10 minutes all family members have to be treated. Pediculosis: 1% lotion to be applied for 7 minutes

Fusidic acid
   EDL-D 645 PHC
INDICATION
Primary and secondary pyodermas caused by Gram positive organisms.
CONTRAINDICATION
Known hypersensitivity.
PRECAUTION
Hepatic disease, Neonates, pregnancy lactation.

Anti-inflammatory and antipruritic medicines

Contact Dermatitis:
Contact dermatitis can result from an allergic or irritant skin reaction. Removal of the substance provoking the reaction is the first step in treating this condition. Mild cases of contact dermatitis can be treated with topical hydrocortisone which suppresses inflammation. A short course of oral prednisolone or a topical corticosteroid such as betamethasone should be considered for more severe cases and for suppression of severe acute reactions associated with blistering, exudation and oedema. Soaking in clean water or mild saline solution is recommended in the acute stages of severe dermatitis.

Pruritus:
Pruritus or itching is a common symptom of many skin diseases. However, contact with certain substances, conditions that dry the skin, stress, and extremes of temperature may also be a cause. Thus, an important part of treatment is to eliminate or minimize the reason for the irritation. Corticosteroids, such as hydrocortisone or betamethasone applied topically, can give relief. Soothing baths or the application of an emollient cream may also be helpful. Systemic antihistamines, such as oral chlorpheniramine, may relieve generalized pruritus.

Atopic Dermatitis:
Atopic dermatitis (or eczema) is a common skin disorder, which mainly occurs in infants and children; it is associated with intense itching, with areas of red skin. Pruritus may be partially relieved by applying astringent aluminium acetate lotion to exudative lesions and emollients to lichenified plaques. Topical hydrocortisone should be applied in short courses of 1–2 weeks to treat even mild areas of involvement. The use of betamethasone should be considered in the treatment of persistent localized dermatitis in adults. Topical antihistamines are not effective
and should be avoided because of the risk of sensitization. However, a sedative antihistamine can be given at night to calm pruritus and facilitate sleep. A secondary infection, often involving *Staphylococcus aureus*, may be responsible for exacerbations; in such cases, an oral antibiotic such as erythromycin can be given for 7-10 days.

**Seborrhoeic Dermatitis:**
Use of a keratolytic shampoo and exposure to ultraviolet light reduce both the inflammation and the scaling resulting from seborrhoeic dermatitis of the scalp (dandruff). The shampoo should be massaged into the scalp, immediately rinsed off and then reapplied until a foam is produced, leaving the second application in contact with the scalp for at least 5 min. Selenium sulfide, which has both antifungal and keratolytic properties, is widely used in many proprietary shampoos. A combination of sulphur and salicylic acid, which has an additional antimicrobial action, is also effective.

**Ichthyosis:**
In ichthyosis, emollients such as aqueous creams and emulsifying creams should be applied daily (or more frequently in severe cases) to affected skin. The addition of a keratolytic, such as salicylic acid 5% can be helpful.

**Lichen Planus:**
Lichen planus is a chronic, papular, pruritic skin eruption that occurs typically in middle age and later life; the condition is often mild and may need no treatment. In more severe cases, when the underlying cause cannot be identified, a topical corticosteroid offers the only prospect of remission.

**Pityriasis Rosea:**
In pityriasis rosea, a common self-limiting dermatosis that is probably of infective origin, calamine lotion helps to relieve pruritus in most cases. If it does not, topical application of hydrocortisone in a concentration not exceeding 1% is worth trying.

**Salicylic Acid**

**EDL-D 468 Secondary hospitals**

**INDICATIONS**
Hyperkeratotic conditions.

**AVAILABILITY**
TOPICAL SOLUTION 2%W/W, OINTMENT 6%, 12%W/W.

**DOSE**
Hyperkeratotic skin disorders: apply once daily, starting with lower strength preparations; gradually increase strength until satisfactory response obtained.

**CONTRAINDICATIONS**
Broken or inflamed skin; children under 2 years.

**PRECAUTIONS**
Diabetes mellitus or if peripheral blood circulation impaired; avoid contact with eyes; mouth; and mucous membranes; avoid application to large areas; irritated; loose/ infected skin; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
Local irritation; dermatitis; salicylism on excessive application or treatment of large areas; particularly in children; salicylic acid poisoning; confusion; dizziness; headache; rapid breathing; ringing/buzzing in ears.

**STORAGE**
Store protected from light.
Betamethasone  
EDL-D 73 PHC

AVAILABILITY  
TABLETS 0.5 mg; injection 1 ml ampoule (4 mg/ml); CREAM 0.1%; OINTMENT 0.1%.

DOSE  
Adult and child- Inflammatory skin conditions, over 2 years of age: apply small quantity to the affected area 1 to 2 times daily until improvement occurs, then less frequently.

INDICATION  
Severe inflammatory skin conditions including contact dermatitis, atopic dermatitis (eczema), seborrhoeic dermatitis, lichen planus, psoriasis of the scalp, hands and feet, intractable pruritus; Addison’s disease, Simmond’s disease, bursitis.

CONTRAINDICATION  
Untreated skin infections or broken skin; rosacea; acne; perioral dermatitis; systemic infections unless specific anti-infective therapy is employed

PRECAUTION  
Children (avoid prolonged use); adrenal suppression if used on a large area of the body or for a long time; particularly with an occlusive dressing or on broken skin; avoid use on the face for more than 7 days; secondary infection requires treatment with an appropriate antimicrobial; may impair the ability to resist and counteract infections; diabetes mellitus; pregnancy (Appendix 7c), elderly; lactation (Appendix 7b)

ADVERSE EFFECTS  
Exacerbation of local infection; local atrophic changes particularly on the face and in skinfolds; characterized by thinning of the dermis; depigmentation; dilatation of superficial blood vessels and formation of striae; perioral dermatitis; acne at site of application; suppression of the hypothalamic-pituitary-adrenal axis with prolonged or widespread use (particularly under occlusion); subcapsular cataract; osteoporosis; glaucoma; intracranial hypertension; psychic instability

Hydrocortisone Acetate  
EDL-D 262 PHC

AVAILABILITY  
CREAM 10 and 15g (1%).

DOSE  
Inflammatory skin conditions: apply a small quantity to the affected area 1 to 2 times daily until improvement occurs, then less frequently.

INDICATION  
Contact dermatitis, atopic dermatitis (eczema), lichen planus; intractable pruritus and phototoxic reactions, including polymorphic light eruptions and actinic prurigo; short-term treatment of psoriasis of the face and flexures; ulcerative colitis.

CONTRAINDICATION  
Untreated skin infections or broken skin; rosacea; acne; perioral dermatitis.

PRECAUTION  
Children (avoid prolonged use); occlusive dressings increase penetration into keratinized lesions (use occlusive dressings only at night and for no longer than 2 days; avoid use on weeping lesions); secondary infection requires treatment with an appropriate antimicrobial; latent peptic ulcer; hypertension; hypothyroidism; psychic derangement; lactation (Appendix 7b); interactions (Appendix 6d); pregnancy (Appendix 7c).

ADVERSE EFFECTS  
Exacerbation of local infection; atrophic changes (see under Betamethasone) less likely with mild corticosteroids; but infants and children particularly susceptible; fluid retention;
hypokalaemia; osteoporosis; impaired wound healing; increased intracranial and intraocular pressure; negative nitrogen balance.

**Benzyl Benzoate**  
**Non-EDL Secondary hospitals**

**INDICATIONS**  
Scabies; head, body and pubic lice; pediculosis.

**AVAILABILITY**  
LOTION 100 ml (25% w/v); ointment 25% w/w (25g).

**DOSE**  
Adult- Scabies: apply from neck down at night for 2 nights; on each occasion wash off after at least 24 h.  
Pediculosis: apply to affected area and wash off 24 h later; further applications possibly needed after 7 and 14 days.

**CONTRAINDICATIONS**  
Irritated skin; neonates; pregnancy.

**PRECAUTIONS**  
Do not use on inflamed or broken skin; avoid contact with eyes and mucous membranes; not recommended for children; lactation (withhold during treatment); apply below neck only; elderly.

**ADVERSE EFFECTS**  
Local irritation; particularly in children.

**STORAGE**  
Store protected from light and air in well filled containers.

**Gamma Benzene Hexachloride**  
**Non-EDL Secondary hospitals**

**INDICATIONS**  
Pediculosis (but use for head lice is restricted by resistance), scabies.

**AVAILABILITY**  
Lotion 1%w/v; Ointment 1%w/w; Cream 1%w/w; Shampo 1%w/v.

**DOSE**  
For pediculosis: As 1% preparation, apply to scalp and hair (taking care not to enter eyes), it should be massaged for 4 minutes and rinsed thoroughly.  
For scabies: Take a proper bath and dry your skin then apply lotion in a thin layer below the neck upto the sole of feet. Leave it for 8-12 hour and then take bath.

**CONTRAINDICATIONS**  
Seizure; hypersensitivity; skin inflammation; broken skin; premature infants; lactation; pregnancy (Appendix 7c).

**PRECAUTIONS**  
Seizure disorder; open wound or sores; neonates, infants below 2 years; avoid contact with face, eyes; mucus membranes urethral meatus, psoriasis, elderly.

**ADVERSE EFFECTS**  
Insomnia; paresthesia; giddiness, agranulocytosis, aplastic anaemia, skin irritation, contact dermatitis; ataxia; alopecia; severe neurologic toxicities; symptoms of acute poisoning include nausea, vomiting, tremors, coma, convulsions and respiratory failure. Liver, kidney and myocardial toxicity have been reported.

**STORAGE**  
Store protected from light.
Scabicides and pediculicides

Scabies:
Scabies is caused by a mite, Sarcoptes scabiei, that burrows into the skin. It is readily transmitted from person to person; therefore the entire household must be treated at the same time to prevent reinfection. It is not necessary to take a bath before treatment with an acaricide, but all clothing and bedding should be washed to prevent reinfection. Benzyl benzoate is an inexpensive scabicide. It must be applied to all skin surfaces, from the scalp to the soles of the feet, avoiding contact with the eyes; it is too irritant for use on children. Permethrin is less irritant and more effective than benzyl benzoate, but also more expensive; it may be used on children. Young infants can be treated with a cream containing precipitated sulphur 6-10% applied once daily for one week.

Pediculosis:
Pediculosis of the head and body is caused by Pediculus humanus capitis and Pediculus humanus corporis respectively; pubic lice (crab lice) infestations are caused by Pthirus pubis, which may also affect the eye lashes and brows. All are transmitted by person to person contact, and may also contaminate clothing and bedding. All members of the affected household (and sexual contacts) must be treated at the same time, and clothing and bedding should be washed or exposed to the air; in head lice infestations, hair brushes and combs should also be disinfected. Head and body lice are readily treated with permethrin; malathion is effective against pubic lice. Benzyl benzoate may be used for all lice infestations.

Ivermectin

EDL-D 296 Tertiary

INDICATIONS
Nematodal infections such as ascariasis, trichuriasis, strongyloidiasis, enterbiasis, lymphatic filariasis, scabies and pediculosis.

AVAILABILITY
TABLETS 3, 6, 9 and 12 mg; INJECTION 10 ml (0.1% w/v).

DOSE
Oral
Strongyloidiasis: 200 μg/kg of body weight once daily for 1-2 days.
Lymphatic filariasis: 400 μg/kg of body weight simple annual dose for 4-6 years.
Scabies and pediculosis: 150-200 μg/kg of body weight single oral dose highly effective.
Second dose may be required 7-10 days later.

CONTRAINDICATIONS
Hypersensitivity, CNS disorders, pregnancy, meningitis, trypanosomiasis, seizures, contraindicated to children below the age of < 5 years old or under 15 kg body weight.

PRECAUTIONS
Concurrent Loa Loa infection, impaired blood-brain barrier function, pregnancy (Appendix 7c), lactation, hepatic, cardiovascular, renal or pulmonary disease, anaemia, coagulation disorder, severe asthma, interactions (Appendix 6c).

ADVERSE EFFECTS
Nausea, vomiting, constipation, abdominal pain and fatigue, rash, arthralgia, fever, myalgia, asthenia, hypotension, tachycardia, edema, lymphadenopathy, sore throat, cough, headache, somnolence, transient eosinophilia, dizziness, diarrhoea, pruritus, orthostatic hypotension, lymph-node tenderness, rare but serious adverse effects such as marked disability and encephalopathies in patients coinfected with heavy burdens of Loa microfilaria.
Antiseptic

An antiseptic destroys or inhibits growth of micro-organisms on living tissues without causing injurious effects when applied to surfaces of the body or to exposed tissues. Some antiseptics are applied to the unbroken skin or mucous membranes, to burns and to open wounds to prevent sepsis by removing or excluding microbes from these areas. Iodine has been modified for use as an antiseptic. The iodophore, povidoneiodine, is effective against bacteria, fungi, viruses, protozoa, cysts and spores and significantly reduces surgical wound infections. The solution of povidone iodine releases iodine on contact with the skin. Chlorhexidine has a wide spectrum of bactericidal and bacteriostatic activity and is effective against both Gram-positive and Gram-negative bacteria although it is less effective against some species of *Pseudomonas* and *Proteus* and relatively inactive against mycobacteria. It is not active against bacterial spores. Chlorhexidine is incompatible with soaps and other anionic materials, such as bicarbonates, chlorides, and phosphates, forming salts of low solubility which may precipitate out of solution. Ethanol has bactericidal activity and is used to disinfect skin prior to injection, venepuncture or surgical procedures.

**Chlorhexidine**  
**EDL-D 111 PHC**

**AVAILABILITY**  
SOLUTION 100 ml (2% and 4% w/v); Mouth Wash 100 ml (0.2%, w/v).

**DOSE**  
Antiseptic (pre-operative skin disinfection and hand washing): use solution in alcohol (70%). Antiseptic (wounds, burns and other skin damage): apply 0.05% aqueous solution. Disinfection of clean instruments: immerse for at least 30 min in 0.05% solution containing Sodium nitrite 0.1% (to inhibit metal corrosion). Emergency disinfection of clean instruments: immerse for 2 min in 0.5% solution in alcohol (70%).

**INDICATION**  
Antiseptic; disinfection of clean instruments; gingivitis.

**CONTRAINDICATION**  
Meningitis; middle ear surgery; sensitive tissues.

**ADVERSE EFFECTS**  
Occasional skin sensitivity and irritation; Upper respiratory tract infection

**Ethyl Alcohol**  
**EDL-D 206 PHC**

**INDICATIONS**  
Disinfection of skin prior to injection venepuncture or surgical procedures.

**AVAILABILITY**  
Regulated by state excise, license is required.

**DOSE**  
Apply undiluted solution.

**PRECAUTIONS**  
Flammable; avoid broken skin; patients have suffered severe burns when diathermy has been preceded by application of alcoholic skin disinfectants; lactation (Appendix 7b).
ADVERSE EFFECTS
Skin dryness and irritation with frequent application.

STORAGE
Store in a tightly closed container at temperature not exceeding 30°C, away from fire and protected from moisture.

Povidone Iodine
EDL D 412,422 Universal

INDICATIONS
Antiseptic; skin disinfection; Mouth wash.

AVAILABILITY
SOLUTIONS 100 and 500 ml (5% w/v), 500 ml (7.5% w/v and 10% w/v); OINTMENT 15g (5% w/w).

DOSE
Adult and Child- Pre- and post-operative skin disinfection: apply undiluted. Antiseptic (minor wounds and burns): apply twice daily.

CONTRAINDICATIONS
Avoid regular or prolonged use in patients with thyroid disorders or those taking lithium; avoid regular use in neonates; avoid in very low birthweight infants; burn covering large surface area; hypersensitivity to iodine.

PRECAUTIONS
Pregnancy (Appendix 7c); lactation (Appendix 7b); broken skin (see below); renal impairment; avoid contact with eyes; neonates. The application of povidone iodine to large wounds or severe burns may produce systemic adverse effects such as metabolic acidosis; hypernatraemia; and impairment of renal function.

ADVERSE EFFECTS
Irritation of skin and mucous membranes; may interfere with thyroid function tests; systemic effects (see under Precautions).

STORAGE
Store protected from light.

Disinfectants

A disinfectant is a chemical agent, which destroys or inhibits growth of pathogenic microorganisms in the non-sporing or vegetative state. Disinfectants do not necessarily kill all organisms but reduce them to a level, which does not harm health or the quality of perishable goods. Disinfectants are applied to inanimate objects and materials such as instruments and surfaces to control and prevent infection. They may also be used to disinfect skin and other tissues prior to surgery (see also Antiseptics, above).

Disinfection of water can be either physical or chemical. Physical methods include boiling, filtration and ultraviolet irradiation. Chemical methods include the addition of chlorine releasing compounds, such as Sodium hypochlorite solution, chloramines T powder, or Sodium dichloroisocyanurate (NaDCC) powder or tablets. Where water is not disinfected at source it may be disinfected by boiling or by chemical means for drinking, cleaning teeth and food preparation.

Chlorine is a hazardous substance. It is highly corrosive in concentrated solution and splashes can cause burns and damage the eyes. Appropriate precautions must be taken when concentrated chlorine solutions or powders are handled.
The chlorinated phenolic compound, chloroxylenol, is effective against a wide range of Gram-positive bacteria. It is less effective against staphylococci and Gram-negative bacteria; it is often ineffective against *Pseudomonas* spp. and inactive against spores.

The aldehyde bactericidal disinfectant, glutaraldehyde, is strongly active against both Gram-positive and Gram-negative bacteria. It is active against the tuberculosis bacillus, fungi such as *Candida albicans*, and viruses such as HIV and hepatitis B. A 2% w/v aqueous alkaline (buffered to pH 8) glutaral solution can be used to sterilize heat-sensitive pre-cleansed instruments and other equipments.

**Sodium Hypochlorite (Bleaching Powder)**

**EDL-D 480 PHC**

**INDICATIONS**
Disinfection of surfaces, equipments, water.

**AVAILABILITY**
POWDER FOR SOLUTION 1g chlorine/litre (1000 parts per million; 0.1%).

**DOSE**
Surface disinfection (minor contamination): apply solutions containing 1000 parts per million. Instrument disinfection: soak in solution containing 1000 parts per million for a minimum of 15 min; to avoid corrosion do not soak for more than 30 min; rinse with sterile water.

**STORAGE**
Store protected from moisture in a tightly closed container.

**Glutaraldehyde**

**EDL-D 247 Secondary hospitals**

**AVAILABILITY**
SOLUTIONS 1, 2 and 5 Litre. (2% aqueous alkaline (pH 8) solution).

**DOSE**
Disinfection of clean instruments - immerse in undiluted solution for 10 to 20 min; up to 2 h may be required for certain instruments (for example bronchoscopes with possible mycobacterial contamination); rinse with sterile water or alcohol after disinfection. Sterilization of clean instruments - Immers in undiluted solution for up to 10 h; rinse with sterile water or alcohol after disinfection.

**INDICATION**
Disinfection and sterilization of instruments and surfaces; conditions like warts and hyperhidrosis of palms and soles.

**CONTRAINDICATION**
Damaged skin

**PRECAUTION**
Minimize occupational exposure by adequate skin protection and measures to avoid inhalation of vapour; lung damage; oral and nasal lesions, if swallowed do not induce vomiting.

**ADVERSE EFFECTS**
Nausea (occupational exposure); headache; airway obstruction; asthma; rhinitis; eye irritation and dermatitis and skin discolouration.
SECTION - 15
DIURETICS

Diuretics increase urinary excretion of water and electrolytes and are used to relieve oedema associated with heart failure, nephrotic syndrome or hepatic cirrhosis. Some diuretics are used at lower doses to reduce raised blood pressure. Osmotic diuretics are mainly used to treat cerebral oedema, and also to lower raised intraocular pressure.

Most diuretics increase urine volume by inhibiting the reabsorption of Sodium and chloride ions in the renal tubule; they also modify renal handling of potassium, calcium, magnesium and urate. Osmotic diuretics act differently; they cause an increase in urine volume by an osmotic effect.

Although loop diuretics are the most potent their duration of action is relatively short, whilst thiazide diuretics are moderately potent but produce diuresis for a longer period. Potassium-sparing diuretics are relatively weak. Carbonic anhydrase inhibitors are weak diuretics which are rarely used for their diuretic effect and are principally used to lower intraocular pressure in glaucoma.

Electrolyte Imbalance:
The adverse effects of diuretic therapy are mainly due to the fluid and electrolyte imbalance induced by the drugs. Hyponatraemia is an adverse effect of all diuretics. The risk of hypokalaemia, which may occur with both thiazide and loop diuretics, depends more on the duration of action than on potency and is thus greater with thiazides than with loop diuretics (when given in equipotent doses). Potassium-sparing diuretics can cause hyperkalaemia. Other electrolyte disturbances include hypercalcaemia (thiazides), hypocalcaemia (loop diuretics) and hypomagnesaemia (thiazide and loop diuretics).

Symptoms of fluid and electrolyte imbalance include dry mouth, thirst, gastrointestinal disturbances (including nausea, vomiting), weakness, lethargy, drowsiness, restlessness, seizures, confusion, headache, muscle pains or cramps, hypotension (including postural hypotension), oliguria, arrhythmias.

Elderly:
The elderly are more susceptible to electrolyte imbalance than younger patients. Treatment should begin with a lower initial dose of the diuretic (commonly about 50% of the adult dose) and then adjusted carefully according to renal function, plasma electrolytes and diuretic response.

Thiazide Diuretics:
Thiazide diuretics, such as hydrochlorothiazide, are moderately potent and act by inhibiting Sodium and chloride reabsorption at the beginning of the distal convoluted tubule. They produce diuresis within 1-2 h of oral administration and most have a duration of action of 12-24 h.

Thiazide diuretics are used in the management of oedema associated with mild to moderate congestive heart failure, renal dysfunction or hepatic disease; however, thiazides are not effective in patients with poor renal function (creatinine clearance of less than 30 ml per min). In severe fluid retention a loop diuretic may be necessary.
In hypertension, a thiazide diuretic is used at a low dose to lower blood pressure with very little biochemical disturbance; the max. therapeutic effect may not be seen for several weeks. Higher doses should not be used because they do not necessarily increase the hypotensive response but may cause marked changes in plasma potassium, magnesium, uric acid, glucose and lipids. If a thiazide alone does not lower blood pressure adequately, it may be used in combination with another antihypertensive such as a beta-adrenoceptor antagonist. Urinary excretion of calcium is reduced by thiazide diuretics and this property is occasionally utilized in the treatment of idiopathic hypercalciuria in patients with calcium-containing calculi. Paradoxically, thiazide diuretics are used in the treatment of diabetes insipidus, since in this disease they reduce urine volume.

Thiazide diuretics, especially in high doses, produce a marked increase in potassium excretion which may cause hypokalaemia; this is dangerous in patients with severe coronary artery disease and those being treated with cardiac glycosides. In hepatic failure hypokalaemia can precipitate encephalopathy, particularly in alcoholic cirrhosis. Potassium-sparing diuretics are used as a more effective alternative to potassium supplements for prevention of hypokalaemia induced by thiazide diuretics; however supplementation with potassium in any form is seldom necessary with the smaller doses of diuretics used to treat hypertension.

**Loop Diuretics:**
Loop diuretics, or high-ceiling diuretics, such as furosemide, are the most potent and rapidly produce an intense dose-dependent diuresis of relatively short duration. Oral furosemide produces diuresis within 30-60 min of administration, with the max. diuretic effect in 1-2 h. The diuretic action lasts for 4-6 h. Intravenous furosemide produces diuresis within 5 min, with the max. Diuretic effect in 20-60 min and diuresis completes within 2 h.

Loop diuretics inhibit reabsorption from the ascending loop of Henlé in the renal tubule and are useful, particularly in situations where rapid and effective diuresis is needed such as reduction of acute pulmonary oedema due to left ventricular failure. They are also used to treat oedema associated with renal and hepatic disorders and are used in high doses in the management of oliguria due to chronic renal insufficiency. Loop diuretics may be effective in patients unresponsive to thiazide diuretics.

Because of their shorter duration of action, the risk of hypokalaemia may be less with loop diuretics than with thiazide diuretics; if required, potassium-sparing diuretics may be used for prevention of hypokalaemia. Loop diuretics may cause hypovolaemia and excessive use can produce severe dehydration with the possibility of circulatory collapse. Furosemide may cause hyperuricaemia and precipitate attacks of gout. Rapid high-dose injection or infusion of furosemide may cause tinnitus and even permanent deafness.

**Potassium-Sparing Diuretics:**
Potassium-sparing diuretics include amiloride and spironolactone; they are weak diuretics and reduce potassium excretion and increase Sodium excretion in the distal tubule. Amiloride acts about 2 h after oral administration, reaching a peak in 6-10 h and persisting for about 24 h. Spironolactone, which acts by antagonising aldosterone, has a relatively slow onset of action requiring 2-3 days to achieve max. diuretic effect, and a similar period of 2-3 days for diuresis to cease after discontinuation of treatment.
Amiloride may be used alone, but its principal use is in combination with a thiazide or a loop diuretic to conserve potassium during treatment of congestive heart failure or hepatic cirrhosis with ascites.

Spironolactone is used in the treatment of refractory oedema due to heart failure, hepatic cirrhosis (with or without ascites), nephrotic syndrome and ascites associated with malignancy. It is frequently given with a thiazide or a loop diuretic, helping to conserve potassium in those at risk from hypokalaemia. A low dose of spironolactone is beneficial in severe heart failure in patients who are already taking an ACE inhibitor and a diuretic. Spironolactone is used in the diagnosis and treatment of primary hyperaldosteronism; presumptive evidence for diagnosis is provided by correction of hypokalaemia and of hypertension.

The most dangerous adverse effect of potassium-sparing diuretics, such as amiloride or spironolactone, is hyperkalaemia, which can be life-threatening. These diuretics are thus best avoided or used very carefully in patients who have or may develop hyperkalaemia, such as those with renal failure, patients receiving other potassium-sparing diuretics and patients taking ACE inhibitors or potassium supplements.

**Osmotic Diuretics:**

Osmotic diuretics, such as mannitol, are administered in sufficiently large doses to raise the osmolality of plasma and renal tubular fluid. Osmotic diuretics are used to reduce or prevent cerebral oedema, to reduce raised intraocular pressure or to treat disequilibrium syndrome. Mannitol is also used to control intraocular pressure during acute attacks of glaucoma. Reduction of cerebrospinal and intraocular fluid pressure occurs within 15 min of the start of infusion and lasts for 3-8 h after the infusion has been discontinued; diuresis occurs after 1-3 h.

Circulatory overload due to expansion of extracellular fluid is a serious adverse effect of mannitol; as a consequence, pulmonary oedema can be precipitated in patients with diminished cardiac reserve, and acute water intoxication may occur in patients with inadequate urine flow.

**Acetazolamide**

**EDL-D 1 Secondary hospitals**

**AVAILABILITY**

Tablet 250 mg; capsule 250 mg.

**DOSE**

Oral Adult- 0.25 to 1g daily in divided doses.

**INDICATION**

As an adjunct in the treatment of chronic open-angle glaucoma; secondary glaucoma; as part of pre-operative treatment of acute angle-closure glaucoma.

**CONTRAINDICATION**

Hypersensitivity to sulfonamides; chronic angle-closure glaucoma (may mask deterioration); hypokalaemia, hyponatraemia, hyperchlorae mic acidosis; renal impairment, severe hepatic impairment; renal hyperchloremic acidosis, addison’s disease.

**PRECAUTION**

Elderly; lactation; diabetes mellitus; pulmonary obstruction; monitor blood count and electrolytes if used for long periods; interactions (Appendix 6b, 6c); pregnancy (Appendix 7c);
severe respiratory acidosis. May impair ability to perform skilled tasks, for example operating machinery, driving.

ADVERSE EFFECTS
Stinging, burning, pain, itching, erythema, transient dryness, allergic blepharitis, transient conjunctivitis, keratitis, decreased corneal sensitivity, diplopia, ptosis; systemic effects; particularly on the pulmonary, cardiovascular and central nervous systems, may follow absorption; blurred vision; headache.

Amiloride
Non-EDL Secondary hospitals

INDICATIONS
Oedema associated with heart failure or hepatic cirrhosis (with ascites), usually with thiazide or loop diuretic; hypertension.

AVAILABILITY
TABLETS 5 mg (Amiloride) + 50 mg (Hydrochlorothiazide), 5 mg (Amiloride) + 40 mg (furosemide).

DOSE
Oral
Oedema: used alone initially 10 mg daily in 1 or 2 divided doses, adjusted according to response (max. 20 mg daily). Combined with a thiazide or a loop diuretic: initially 5 mg daily, increasing to 10 mg if necessary (max. 20 mg daily).

CONTRAINDICATIONS
Hyperkalaemia; renal failure; potassium supplementation.

PRECAUTIONS
Monitor electrolytes; particularly potassium; hypocholeremia, hepatic cirrhosis, renal impairment (Appendix 7d); diabetes mellitus; elderly (reduce dose); lactation; interactions (Appendix 6b, 6c); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Hyperkalaemia; hyponatremia (for symptoms of fluid and electrolyte imbalance see introductory notes); diarrhoea; constipation; anorexia; paraesthesia; dizziness; minor psychiatric or visual disturbances; rash; pruritus; rise in blood urea nitrogen; headache; abdominal pain, flatulence.

STORAGE
Store protected from light.

Furosemide
EDL-D 238,239,240,241 PHC

INDICATIONS
Oedema; oliguria due to renal failure; pulmonary oedema; hypertension.

AVAILABILITY
TABLETS 40, 100, 200 and 500 mg; INJECTION 2 ml (20 mg/ml).

DOSE
Oral
Adult- Oedema: initially 40 mg daily on waking up; maintenance dose 20 to 40 mg daily; may be increased to 80 mg daily or more in resistant oedema.
Child- 1 to 3 mg/kg daily (max. 40 mg daily).

CONTRAINDICATIONS
Renal failure with anuria; precomatose states associated with liver cirrhosis; hypersensitivity.

PRECAUTIONS
Monitor electrolytes particularly potassium and Sodium; hypotension; asymptomatic hyperuricaemia, systemic lupus erythmatosus, elderly (reduce dose); pregnancy (Appendix 7c); lactation; correct hypovolaemia before using in oliguria; renal impairment; hepatic impairment (Appendix 7a); prostatic enlargement; porphyria; interactions (Appendix 6b, 6c).
ADVERSE EFFECTS
Hypokalaemia; hypomagnesaemia; hyponatraemia; hypochloraemic alkalosis (for symptoms of fluid and electrolyte imbalance; see introductory notes); increased calcium excretion hypovolaemia; hyperglycaemia (but less often than with thiazide diuretics); temporary increase in plasma cholesterol and triglyceride concentration; less commonly hyperuricaemia and gout; rarely, rash; photosensitivity; bone marrow depression (withdraw treatment); pancreatitis (with large parenteral doses); tinnitus and deafness (with rapid administration of large parenteral doses and in renal impairment; deafness may be permanent if other ototoxic drugs taken); hepatic encephalopathy, anorexia, orthostatic hypotension.

Hydrochlorothiazide
EDL-D 260 PHC
INDICATIONS
Oedema; diabetes insipidus; hypertension; heart failure.

DOSE
Oral
Adult - Hypertension: 12.5 to 25 mg daily.
Oedema: initially 25 mg daily on waking up, increased to 50 mg daily if necessary.
Severe oedema in patients unable to tolerate loop diuretics: up to 100 mg either daily or on alternate days (max. 100 mg daily).
Nephrogenic diabetes insipidus: initially up to 100 mg daily.
Elderly - Hypertension: initially 12.5 mg daily.
Oedema: initially 12.5 mg daily.

ADVERSE EFFECTS
Hypokalaemia; hypomagnesaemia; hyponatraemia; hypochloraemic alkalosis (for symptoms of fluid and electrolyte imbalance see introductory notes); hypercalcaemia; hyperglycaemia; hyperuricaemia; gout; rash; photosensitivity; altered plasma lipid concentration; rarely, impotence (reversible); blood disorders (including neutropenia; thrombocytopenia); pancreatitis; intrahepatic cholestasis and hypersensitivity reactions (including pneumonitis; pulmonary oedema; severe skin reactions) also reported; acute renal failure.

Mannitol
Non –EDL Secondary hospitals
INDICATIONS
Cerebral edema, impending acute renal failure, acute poisonings, raised intraocular pressure (emergency treatment or before surgery).

AVAILABILITY
Infusion 5, 10 and 20%.

DOSE
Test dose (if patient is oliguric or if renal function is inadequate), By intravenous infusion as a 20% solution infused over 3–5 minutes, Adult and Child- 200 mg/kg; repeat test dose if urine output is less than 30–50 ml/h; if response is inadequate after a second test dose, re-evaluate the patient. Raised intracranial or intraocular pressure: By i.v infusion as a 20% solution infused over 30–60 minutes, Adult- 0.25–2g/kg; Child- 0.5–1.5g/kg.
Cerebral oedema: By i.v infusion as a 20% solution infused rapidly, Adult and Child- 1g/kg.

CONTRAINDICATIONS
Acidosis, congestive heart failure, pulmonary oedema (particularly in diminished cardiac reserve), dehydration, inadequate urine flow, acute tubular necrosis, anuria, acute left ventricular failure, intracranial bleeding.

PRECAUTIONS
Patients with cardiovascular disease; hypervolemia; urinary tract obstruction; should not be given with whole blood; pregnancy (Appendix 7c).
ADVERSE EFFECTS
Headache, nausea, vomiting, dehydration, edema, hypernatraemia, inflammation, skin necrosis, urticaria, chills, convulsions, fluid and electrolyte imbalance, acidosis, circulatory overload, visual disturbance.

STORAGE
Store at temperatures between 20° and 30°C. Exposure to lower temperatures may cause the deposition of crystals, which should be dissolved by warming before use.

Spironolactone
EDL-D 482 Secondary hospitals D 483 Tertiary

INDICATIONS
Refractory oedema in congestive heart failure; adjunct to ACE inhibitor and loop or thiazide diuretic in severe congestive heart failure; nephrotic syndrome; hepatic cirrhosis with ascites and oedema; ascites associated with malignancy; primary hyperaldosteronism.

AVAILABILITY
TABLETS 25 and 100 mg.

DOSE
Oral
Adult- Oedema: 100 to 200 mg daily, increased if necessary to 400 mg daily in resistant oedema; usual maintenance dose 75-200 mg daily.
Primary hyperaldosteronism (diagnosis): 400 mg daily for 3 to 4 weeks. Preoperative management: 100 to 400 mg daily. If not suitable for surgery; lowest effective dose for long-term maintenance.
Adjunct in severe heart failure: usually 25 mg daily.
Child- Initially 3 mg/kg daily in divided doses.

CONTRAINDICATIONS
Pregnancy (Appendix 7c); lactation; hyperkalaemia; hyponatraemia; severe renal impairment; Addison’s disease; anuria.

PRECAUTIONS
Monitor blood urea nitrogen and plasma electrolytes (discontinue if hyperkalaemia); concomitant administration of potassium sparing diuretics and its inhibitors and NSAIDs, elderly (reduce dose); diabetes mellitus; renal impairment; hepatic impairment; porphyria; high doses carcinogenic in rodents; interactions (Appendix 6b, 6d).

ADVERSE EFFECTS
Hyperkalaemia; hyponatraemia; hyperchloraemic acidosis; dehydration (for symptoms of fluid and electrolyte imbalance see introductory notes); transient increase in blood urea nitrogen; diarrhoea; gynaecomastia; menstrual irregularities; impotence; hirsutism; deepening of voice; rash; ataxia; fever; hepatotoxicity; gastric bleeding, ulceration; agranulocytosis.

STORAGE
Store protected from light and moisture.

Torasemide (Torsemid)
EDL-D 751,752 Secondary hospitals

Similar to Furosemide but 3 times more potent.

INDICATION
Oedema and hypertension

AVAILABILITY
5mg, 10mg, 20mg.

DOSE
Hypertension 2.5-5mg OD; Oedema 5-20 mg/day; Renal failure upto 100 mg daily

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**Antacids and other anti ulcer medicines**

Antacids (usually containing aluminium or magnesium compounds) can often relieve symptoms in ulcer dyspepsia and in non-erosive gastro-oesophageal reflux; they are also sometimes used in non-ulcer dyspepsia but the evidence of benefit is uncertain. Antacids are best given when symptoms occur or are expected, usually between meals and at bedtime, **Liquid preparations are more effective than solids.**

Aluminium- and magnesium-containing antacids (for example aluminium hydroxide and magnesium hydroxide), being relatively insoluble in water, are long-acting if retained in the stomach. They are suitable antacids for most purposes. **Magnesium-containing antacids have a laxative effect whereas aluminium-containing antacids may be constipating.**

H$_2$-receptor antagonists heal gastric and duodenal ulcers by reducing the secretion of gastric acid as a result of histamine H$_2$-receptor blockade; they can also relieve gastro-oesophageal reflux disease. High doses of H$_2$-receptor antagonists have been used in the Zollinger-Ellison syndrome, but a proton-pump inhibitor is now preferred.

Maintenance treatment with low doses has largely been replaced in *Helicobacter pylori* positive patients by eradication regimens. Maintenance treatment may occasionally be used for those with frequent severe recurrences and for the elderly who suffer ulcer complications.

Treatment of undiagnosed dyspepsia with H$_2$-receptor antagonists may be acceptable in younger patients but care is required in older patients because their symptoms may be caused by gastric cancer. **H$_2$-receptor antagonist therapy can promote healing of NSAID-associated ulcers (particularly duodenal).** Treatment also reduces the risk of acid aspiration in obstetric patients at delivery (Mendelson syndrome).

**Peptic Ulcer**

Ulcer disease is caused by peptic ulceration that involves the stomach, duodenum and lower oesophagus. General and inexpensive measures like introducing healthy life-style, stopping smoking and taking antacids should be promoted. The possibility of malignant disease should be considered in all patients over the age of 40 years who are suspected of having an ulcer.

Gastric and duodenal ulcers are healed by 4-8 weeks treatment with H2-receptor antagonists but there is a high rate of relapse (greater than 70% over 2 years) requiring maintenance therapy. Relapses can be prevented very successfully by eradicating Helicobacter pylori which is causally associated with most peptic ulcers (except those related to NSAID use). Eradication of H. pylori reduces the relapse rate to about 4-8%. This is undoubtedly cost-effective compared to the alternatives of long-term maintenance therapy with low-dose H2-receptor antagonists or repeated treatment of recurrent ulcers. It is recommended that the presence of H. pylori is confirmed before starting eradication treatment, particularly for gastric ulcers. The urea breath test is used widely to test for H. pylori, but it may produce false negative results if used soon
after proton-pump inhibitors or antibacterials. Eradication regimens are based on a combination of an acid-reducing (‘antisecretory’) drug and antibiotics.

The following model eradication regimen is suggested on the basis of its efficacy and simplicity (only doses suitable for adults are shown):

- Omeprazole 40 mg daily for 1 week plus
- Metronidazole 400 mg thrice daily for 1 week plus
- Amoxycillin 500 mg thrice daily for 1 week

The decision on choosing an eradication regimen should take into account local resistance to antibacterials, cost and availability of the necessary drugs.

**NSAID -Associated Ulcers**

Gastrointestinal bleeding and ulceration may occur with NSAID use. To avoid this, emphasis should be on stopping NSAID use but this is not always possible. A proton-pump inhibitor may be considered for protection against NSAID-associated gastric and duodenal ulcers. An H2-receptor antagonist may be effective for protection against NSAID-associated duodenal ulcers only.

Patients who must continue NSAID therapy after ulcer development may take high-dose H2-receptor antagonists concomitantly, but ulcers tend to heal more slowly with H2-receptor antagonists if NSAIDs are continued. A proton-pump inhibitor such as omeprazole is more effective but it is also more expensive.

In patients who can discontinue NSAID therapy after ulcer development, treatment with an H2-receptor antagonist is effective, but a treatment period of up to 8 weeks may be necessary. A proton-pump inhibitor usually produces the most rapid healing. After healing, continued prophylaxis is required.

**Dyspepsia**

Dyspepsia covers pain, fullness, early satiety, bloating, or nausea. It can occur with gastric and duodenal ulceration and gastric cancer but most commonly it is of uncertain origin.

Patients with non-ulcer dyspepsia should be advised to avoid smoking, alcohol and aggravating foods and to eat small regular meals to aid digestion. Non-ulcer dyspepsia tends to be self-limiting but antacids and H2-receptor antagonists are often used to suppress gastric acid. Effective treatment is important in the presence of severe oesophageal ulceration to prevent longer term complications such as oesophageal stricture and carcinoma.

**Gastro-Esophageal Reflux Disease (GERD)**

GERD (including non-erosive gastro-esophageal reflux and erosive esophagitis) is characterized by symptoms which include heartburn, acid regurgitation and sometimes difficulty in swallowing (dysphagia); esophageal inflammation (esophagitis), ulceration and stricture formation may occur and there is an association with asthma.

The management of GERD includes drug treatment, lifestyle changes and, in some cases, surgery. Initial treatment is guided by the severity of symptoms and treatment is then adjusted according to response.

For mild symptoms of GERD, initial management may include the use of antacids. H2-receptor antagonists suppress acid secretion and they may relieve symptoms and permit reduction in antacid consumption. Severe symptoms initially require a short-course of a proton-pump inhibitor.
Zollinger-Ellison Syndrome
Management of Zollinger-Ellison syndrome requires high dose H2-receptor antagonist treatment. The proton pump inhibitors are more effective particularly for cases resistant to other treatment but they are more expensive.

Aluminium Hydroxide+ Magnesium Hydroxide + Active Dimethicon/ Simethicon
EDL-D 17,18 Universal

DOSE
adults and children 12 years and older: take 2 to 4 teaspoonsful (10-20 mL) four times a day or as directed by a physician, do not take more than 16 teaspoonsful in 24 hours or use the maximum dosage for more than 2 weeks, children under 12 years: consult a physician

INDICATION
acid indigestion, heartburn, sour stomach, upset stomach associated with these symptoms, pressure and bloating commonly referred to as gas

CONTRAINDICATION
Hypophosphataemia; undiagnosed gastrointestinal or rectal bleeding; appendicitis; porphyria; hypersensitivity to aluminium salts.

PRECAUTION
do not exceed 16 teaspoonsful (80 mL) in a 24-hour period, or use the maximum dosage for more than 2 weeks, unless directed by a doctor

ADVERSE EFFECTS
Constipation, intestinal obstruction (large doses); hypophosphataemia with increased bone resorption, hypercalciuria and risk of osteomalacia (patients on low phosphate diet or prolonged therapy); hyperaluminaemia resulting in osteomalacia, encephalopathy, dementia, microcytic anaemia (in chronic renal failure treated with aluminium hydroxide as phosphate-binding agent); loss of appetite.

Ranitidine
EDL-D 452 PHC D 453 Secondary hospitals

AVAILABILITY
TABLETS 150 and 300 mg. INJECTION 2 ml ampoule (25 mg/ml), SYRUP 375 mg/5 ml.

DOSE
Adult- Benign gastric and duodenal ulceration: 150 mg twice daily or 300 mg at night for 4 to 8 weeks, up to 6 weeks in chronic episodic dyspepsia and up to 8 weeks in NSAID-associated ulceration (in duodenal ulcer 300 mg can be given twice daily for 4 weeks to achieve a higher healing rate); maintenance, 150 mg at night. Prophylaxis of NSAID-induced duodenal ulcer: 150 mg twice daily. Reflux oesophagitis: 150 mg twice daily or 300 mg at night for up to 8 weeks, or if necessary 12 weeks (moderate to severe, 150 mg 4 times daily for up to 12 weeks). Long-term treatment of healed oesophagitis: 150 mg twice daily. Zollinger-Ellison syndrome: 150 mg 3 times daily (up to 6g daily in divided doses has been used). Gastric acid reduction (prophylaxis of acid aspiration) in obstetrics: 150 mg at onset of labour, then every 6 h. Surgical procedures: 150 mg 2 h before induction of anaesthesia and also, when possible on the preceding evening. Child-Peptic ulcer: 2 to 4 mg/kg twice daily (max. 300 mg daily). Intramuscular injection Adult- Benign gastric and duodenal ulceration, reflux oesophagitis, Zollinger-Ellison syndrome: 50 mg every 6 to 8 h. Surgical procedures: 50 mg 45 to 60 min before induction of anaesthesia. Slow intravenous injection Benign gastric and duodenal ulceration, reflux oesophagitis, Zollinger-Ellison syndrome: 50 mg diluted to 20 ml and given over at least 2 min, may be repeated every 6 to 8 h. Surgical procedures: 50 mg 45 to 60 min before induction of anaesthesia (intravenous injection diluted to 20 ml and given over at least 2 min). Intravenous infusion Benign gastric and duodenal ulceration, reflux oesophagitis, Zollinger-Ellison syndrome: 25 mg/h for 2 h, may be repeated every 6 to 8 h. Prophylaxis of stress ulceration: initial slow intravenous injection of 50
mg diluted to 20 ml and given over at least 2 min then by continuous intravenous infusion, 125-250 μg/kg per h (may be followed by 150 mg twice daily by mouth when oral feeding commences).

INDICATION
Benign gastric and duodenal ulceration, GERD, Zollinger-Ellison syndrome, other conditions where gastric acid reduction is beneficial. Prophylaxis during NSAIDs treatment in patients with high risk for peptic ulceration, eradication of H.pylori, as preoperative medication, systemic mastocytosis

CONTRAINDICATION
Porphyria.

PRECAUTION
Hepatic impairment (Appendix 7a); renal impairment; lactation (Appendix 7b); middleaged or older patients and those whose symptoms change-may mask gastric cancer; interactions (Appendix 6a); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Diarrhoea and other gastrointestinal disturbances; headache; dizziness; rash; tiredness; acute pancreatitis; bradycardia, tachycardia; AV block, confusion; depression; rarely, hallucinations (particularly in the elderly or the very ill); hypersensitivity reactions (including fever, arthralgia, myalgia, anaphylaxis); blood disorders (including agranulocytosis, leukopenia, pancytopenia, thrombocytopenia); hepatitis; agitation; visual disturbances; erythema multiforme; alopecia; gynaecomastia and impotence; malaise; somnolence

Omeprazole
EDL-D 382 PHC

AVAILABILITY
TABLETS 20 and 40 mg; INJECTION 10 ml vial (40 mg/10 ml); CAPSULES 10, 20 and 40 mg.

DOSE
Oral Benign gastric and duodenal ulcers: 20 mg once a day for 4 weeks in duodenal ulcers, for 8 weeks in gastric ulcers, Increase to 40 mg in severe case. Maintenance for recurrent duodenal ulcers: 20 mg once daily. Prevention of relapse: 10 mg daily. NSAIDs associated gastric or duodenal ulcers or gastro-duodenal erosions: 20 mg daily for 4 weeks. Prophylaxis in case of history associated with gastric/duodenal ulcers or dyspepsia: 20 mg daily. Zollinger-Ellison syndrome: 60 mg to 120 mg/day or more, into divided doses. Gastric acid reduction during gastric surgery: 40 mg on preceding evening then 40 mg 2 to 6 h before surgery.

INDICATION
Benign gastric and duodenal ulcers; Zollinger Ellison syndrome; gastric acid reduction during gastric surgery; GERD, NSAID- induced ulcer, prophylaxis during NSAIDs treatment in patients with high risk for peptic ulceration, eradication of H.pylori, as preoperative medication, systemic mastocytosis and in patients not responsive to H2 blockers.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Interactions (Appendix 6c, 6d); pregnancy(Appendix 7c) ; concomitant gastric malignancy.

ADVERSE EFFECTS
Nausea, abdominal pain, constipation, flatulence, diarrhoea, headache, skin rashes, subacute, myopathy, arthralgias, increased risk of hip fractures, decreased B12 absorption, hypergastrinemainia, respiratory and Clostridium difficile infections, hepatic dysfunction.

Pantoprazole
EDL-D 400 Secondary hospitals

AVAILABILITY
TABLETS 20 and 40 mg, INJECTIONS 20 and 40 mg/vial, CAPSULES 20 and 40 mg.
DOSE
Oral Adult- 40 mg once daily up to 8 weeks. Intravenous Adult- 40 mg twice daily.

INDICATION
Duodenal ulcer, gastric ulcer, GERD, erosive esophagitis.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Hepatic impairment; monitor liver function; pregnancy(Appendix 7c); cyanocobalamin deficiency; tumorogenicity

ADVERSE EFFECTS
Diarrhoea; pruritus; dizziness; pyrexia; blurred vision; vertigo.

Antiemetic medicines
Antiemetics are drugs effective against nausea and vomiting. They are typically used to treat motion sickness and the side effects of opioid analgesics, general anaesthetics and chemotherapy induced nausea and vomiting in cancer patients either alone or in combination. They act on the brain by preventing the stimulation of the vomiting centre (chemoreceptor trigger zone-CTZ). Some medications act on the gut by speeding up the rate at which the stomach empties and help to facilitate the quick transit of food through intestine (prokinetic action).

Classification:
• 5-HT3 receptor antagonists block serotonin receptors in the central nervous system and gastrointestinal tract: Ondansetron, Granisetron, Dolasetron etc.
• Dopamine D2-receptor antagonists act in the brain: Domperidone, Metoclopramide, Mosapride etc.
• Antihistamines or H1- histamine receptor antagonists: Diphenhydramine, Promethazine etc.
• Benzodiazepines: Midazolam, Lorazepam etc.
• Anticholinergics: Scopolamine, Hyoscine, Dicyclomine etc.
• Steroids: Dexamethasone etc.

Metoclopramide has antiemetic properties and also stimulates upper gastrointestinal motility. It is effective against nausea and vomiting associated with gastrointestinal disorders or migraine, following surgery and chemotherapy and is also effective against radiation-induced nausea and vomiting. Combining metoclopramide with corticosteroids (such as dexamethasone) can improve its antiemetic effect in chemotherapy- induced nausea and vomiting. Metoclopramide may be useful in the management of gastro-oesophageal reflux and gastroparesis, as well as preoperatively in the prevention of aspiration syndromes. It is also used to facilitate intubation of the small bowel during radiographic examinations. It is not effective in the prevention or treatment of motion sickness.

Metoclopramide may cause acute dystonic reactions with facial and skeletal muscle spasms and oculogyric crisis. These reactions are most common in the young (especially girls and young women) and the elderly; they occur shortly after the start of treatment and subside within 24 h of drug withdrawal. Promethazine is a phenothiazine derivative. In addition to D2 dopaminergic blockade it has pronounced histamine H1 and muscarinic receptor blocking properties. It is effective in the prevention and treatment of vertigo and motion sickness.
Promethazine may be useful in the prevention and treatment of postoperative and drug-induced nausea and vomiting. It has limited effect on chemotherapy-induced mild to moderate emesis.

**Metoclopramide Hydrochloride**

**EDL-D 339,340 PHC**

**AVAILABILITY**

TABLETS 10 and 15 mg; INJECTION 2 ml ampoule (5 mg/ml); SYRUP 30 ml (1 mg/ml).

**DOSE**

Oral or intramuscular injection or Slow intravenous injection

- Adult- Nausea and vomiting, gastroesophageal reflux, gastroparesis: (over 1 to 2 min for slow intravenous injection), 10 mg 3 times daily. 15 to 19 years (under 60 kg) 5 mg 3 times daily. Aid to gastrointestinal intubation: 20 mg as a single dose 5 to 10 min before examination; Adolescent (15 to 19 years), 10 mg. Child-
  - Up to 1 year (up to 10 kg) 1 mg twice daily; 1 to 3 years (10 to 14 kg) 1 mg 2 to 3 times daily; 3 to 5 years (15 to 19 kg) 2 mg 2 to 3 times daily; 5 to 9 years (20 to 29 kg) 2.5 mg 3 times daily; 9 to 14 years (30 kg and over) 5 mg 3 times daily (usual max. 500 µg/kg daily, particularly for children and young adult). Slow intravenous injection only Adult- Premedication: 10 mg as a single dose.

**INDICATION**

Nausea and vomiting in gastrointestinal disorders and treatment with cytotoxics or radiotherapy; gastro-oesophageal reflux disease; gastroparesis; premedication and postoperatively; aid to gastrointestinal intubation; nausea and vomiting in migraine; diabetic gastric stasis.

**CONTRAINDICATION**

Gastrointestinal obstruction, haemorrhage or perforation, 3-4 days after gastrointestinal surgery; convulsive disorders; pheochromocytoma; hypersensitivity.

**PRECAUTION**

Elderly, children and young adults; hepatic impairment (Appendix 7a); renal impairment (Appendix 7d); pregnancy (Appendix 7c); may mask underlying disorders such as cerebral irritation; avoid for 3-4 days after gastrointestinal surgery; lactation (Appendix 7b); interactions (Appendix 6a); Parkinson’s disease; epilepsy; depression; porphyria; driving or operating machines; hypertension; cirrhosis; congestive heart failure.

**ADVERSE EFFECTS**

Extrapyramidal symptoms (especially in children and young adults; see notes above); tardive dyskinesias on prolonged use; hyperprolactinaemia; drowsiness, restlessness, dizziness, headache, diarrhoea, depression, hypotension and hypertension reported; rarely, neuroleptic malignant syndrome; rashes, pruritus, oedema; cardiac conduction abnormalities following intravenous administration; rarely, methaemoglobinemia (more severe in G-6-PD deficiency); galactorrhoea; amenorrhoea; bradykinesia; gynaecomastia; insomnia.

**Domperidone**

**EDI-D 188,189 PHC**

**AVAILABILITY**

TABLETS 5 and 10 mg; SYRUP 30 ml (1 mg/ml); Capsule 30 mg.

**DOSE**

Oral Adult- 10 to 20 mg 3 to 4 times a day Child- 0.3 to 0.6 mg/kg TDS.

**INDICATION**

Nausea and vomiting from any cause in adult, epigastric senses of fullness; upper abdominal distress; non ulcer dyspepsia; migraine.
CONTRAINDICATION
Hypersensitivity; prolactinoma, hepatic impairment; where increased gastrointestinal motility harmful; pregnancy; gastrointestinal haemorrhage; intestinal obstruction.

PRECAUTION
Children; renal impairment, interactions(Appendix 6c); history of breast cancer; allergies; pheochromocytoma; i.v. administration can lead to hypokalaemia and cardiac arrhythmias.

ADVERSE EFFECTS
Rarely, gastrointestinal disturbances (including cramps) and hyperprolactinaemia; very rarely, extrapyramidal effects and rashes; headache; dizziness; dry mouth; nervousness; flushing.

Prochlorperazine
EDL-D 438 PHC

AVAILABILITY
TABLETS 3 and 5 mg; INJECTION 1 ml ampoule (2.5 mg/ml).

DOSE
Oral and intravenous injection Adult- Nausea, vomiting acute attack: initially 20 mg then 20 mg every 2 h. Prevention; 5 to 10 mg 2 to 3 times daily. Child- (over 10 kg only). Oral: 0.4 mg/kg/day in 3-4 divided doses. Intravenous injection: 0.13 mg/kg/day in 3-4 divided doses. Adult- Labyrinthine disorder: 5 mg 3 times daily increased to 30 mg daily in divided doses that decrease after meal to 5 to 10 mg daily. Child- Labyrinthine disorder Not recommended. Intravenous injection: 0.13 mg/kg/day in 3-4 divided doses.

INDICATION
Nausea and vomiting.

CONTRAINDICATION
Comatose states, CNS depression and pheochromocytoma. Most antipsychotics are best avoided during pregnancy; hypersensitivity; prolactin dependant tumors.

PRECAUTION
Patients with hepatic impairment, renal impairment, cardiovascular disease, Parkinson’s disease (may be exacerbated by antipsychotics), epilepsy (and conditions predisposing to epilepsy), depression, myasthenia gravis, prostatic hypertrophy, or a susceptibility to angle-closure glaucoma. Caution is also required in severe respiratory disease and in patients with a history of jaundice or who have blood dyscrasias (perform blood counts if unexplained infection or fever develops). Caution should be taken in elderly, who are particularly susceptible to postural hypotension and to hyper- or hypothermia in very hot or cold weather. Serious consideration should be given before prescribing these drugs for elderly patients. As photosensitisation may occur with higher dosages, patients should avoid direct sunlight; extrapyramidal syndrome; pregnancy(Appendix 7c); interactions(Appendix 6a).

ADVERSE EFFECTS
Less sedating; extrapyramidal symptoms, particularly dystonias, more frequent; respiratory depression may occur in susceptible patients; amenorrhoea; blurred vision; cholestatic jaundice; neuroleptic malignant syndrome; leucopenia; agranulocytosis.

Ondansetron
EDL-D 384, 385 Secondary hospitals D 706 PHC

AVAILABILITY
TABLETS 4 and 8 mg; INJECTION 2 and 4 ml ampoule (2 mg/ml); DROPS 2 mg/5 ml; SYRUP 2 mg/5 ml; SUSPENSION 1 mg/5 ml.

DOSE
Oral Prevention of post-operative nausea and vomiting: Adult 16 mg, 1 h before induction of anaesthesia. Nausea and vomiting associated with cancer chemotherapy: Adult- 24 mg as a
single dose taken 30 min before start of single day chemotherapy. Child (4-11 yrs)- 4 mg tablets 3 times a day; continue for 1-2 days after completion of chemotherapy. Parenteral Postoperative nausea and vomiting: Adult- 4 mg by i.m or slow i.v as a single dose. Prevention of chemotherapy-induced nausea and vomiting: Adult- single 32 mg i.v dose infused over 15 min begining 30 min before start of emetogenic chemotherapy.

INDICATION
Postoperative nausea and vomiting, chemotherapy and/or radiotherapy induced nausea and vomiting.

CONTRAINDICATION
Hypersensitivity.

PRECAUTION
Moderate to severe liver impairment; pregnancy (Appendix 7c), lactation; hypersensitivity to other selective 5-HT3 - receptor antagonists, subacute intestinal obstruction; cardiac disease, electrolyte abnormalities, QT interval prolongation (avoid concomitant administration of drugs that prolong QT interval), interactions (Appendix 6a).

ADVERSE EFFECTS
Headache, constipation or diarrhoea, dizziness; flushing, hypersensitivity reaction, anaphylaxis/anaphylactoid reactions, angioedema; bronchospasm, hypotension, laryngeal edema, urticaria, hiccups, oculogyric crisis.

Promethazine
EDL-D 721 PHC

AVAILABILITY
TABLETS 10 and 25 mg; INJECTION 2 ml ampoule (25 mg/ml).

DOSE
Oral Nausea and vomiting (including postoperative): 12.5 to 25 mg, repeated at intervals of not less than 4 h (usual max., 100 mg in 24 h). Motion sickness, prevention: 20 to 25 mg at bedtime on night before travel, repeated on day of travel if necessary. Child- Motion sickness, prevention; 2 to 5 years: 5 mg at night and on day of travel, if necessary. 5 to 10 years: 10 mg at night and on day of travel, if necessary.
Intramuscular injection or Slow intravenous injection
Nausea and vomiting (including postoperative); (diluted to 2.5 mg/ml in water for injection); 12.5 to 25 mg, repeated at intervals of not less than 4 h (usual max., 100 mg in 24 h).

INDICATION
Nausea, vomiting, labyrinthine disorders, motion sickness; premedication; allergic rhinitis; vasomotor rhinitis.

CONTRAINDICATION
Porphyria; hypersensitivity; coma; hypokalaemia

PRECAUTION
Prostatic hypertrophy; urinary retention; glaucoma; hepatic disease (Appendix 7a); epilepsy; elderly and children (more susceptible to adverse effects); lactation (Appendix 7b); pregnancy (Appendix 7c); interactions (Appendix 6a).
May impair ability to perform skilled tasks, for example operating machinery, driving.

ADVERSE EFFECTS
Drowsiness, dizziness, sedation (but paradoxical stimulation may occur, especially with high doses or in children and elderly); headache, psychomotor impairment; urinary retention, dry mouth, blurred vision, gastrointestinal disturbances; hypersensitivity reactions, rashes, photosensitivity reactions; jaundice; blood disorders; cardiovascular adverse effects-after injection; venous thrombosis at site of intravenous injection; pain on intramuscular injection; somnolence; torticollis; tinnitus; leucopenia; thrombocytopenia, agranulcytosis; apnoea; angioneurotic edema.
Antiaemorrhoidal medicines Local anaesthetic, 
Astringent and Anti-inflammatory drug

Ulcerative colitis and Crohn’s disease are inflammatory diseases of the intestinal tract. 

**Ulcerative Colitis:**
Acute attacks of ulcerative colitis require treatment with local corticosteroids such as hydrocortisone in the form of suppositories or retention enemas. Because of the risk of intestinal perforation, rectal administration of hydrocortisone must be used with extreme caution in patients with severe ulcerative disease and should not be given to such patients without conducting a thorough proctological examination. More extensive disease requires oral corticosteroid treatment and severe extensive or fulminant disease needs hospital admission and intravenous corticosteroid administration; other therapy may include intravenous fluid and electrolyte replacement, blood transfusion and possibly parenteral nutrition and antibiotics.

The aminosalicylate sulfasalazine is useful in the treatment of symptomatic disease. It also has value in the maintenance of remission in ulcerative colitis for which corticosteroid treatment is unsuitable because of adverse effects. In resistant or frequently relapsing cases azathioprine 2–2.5 mg/kg daily (chapter 12.1) given under close supervision may be helpful. Laxatives are required to facilitate bowel movement when proctitis is present. Antimotility drugs such as codeine and antispasmodic drugs should not be used in active ulcerative colitis because they can precipitate paralytic ileus and megacolon. Diarrhoea resulting from reduced bile salt absorption may improve with cholestyramine. General nutritional care and appropriate supplements are essential. High-fibre or low-residue diets should be used as appropriate.

Irritable bowel syndrome during remission of ulcerative colitis requires avoidance of a high-fibre diet and possibly treatment with an antispasmodic.

**Crohn’s Disease:**

Treatment of Crohn’s disease of the colon is similar to that of ulcerative colitis. In small bowel disease sulfasalazine may have marginal benefit. Symptoms and inflammation associated with disease exacerbation are suppressed by oral corticosteroids such as prednisolone. Metronidazole may be beneficial in the treatment of active Crohn disease particularly with perianal involvement, possibly through its antibacterial activity. Other antibacterials should be given if specifically indicated (for example, sepsis associated with fistulas and perianal disease) and for managing bacterial overgrowth in the small bowel. General nutritional care and appropriate supplements are essential.
Betamethasone Dipropionate + Phenylephrine + Lignocaine  
**EDL-D 75 PHC**

**AVAILABILITY**
- TABLETS 0.5 mg; injection 1 ml ampoule (4 mg/ml); CREAM 0.1%; OINTMENT 0.1%.

**DOSE**
Adult and child - Inflammatory skin conditions, over 2 years of age: apply small quantity to the affected area 1 to 2 times daily until improvement occurs, then less frequently.

**INDICATION**
- Severe inflammatory skin conditions including contact dermatitis, atopic dermatitis (eczema), seborrhoeic dermatitis, lichen planus, psoriasis of the scalp, hands and feet, intractable pruritus; Addison’s disease, Simmond’s disease, bursitis.

**CONTRAINDICATION**
- Untreated skin infections or broken skin; rosacea; acne; perioral dermatitis; systemic infections unless specific anti-infective therapy is employed.

**PRECAUTION**
- Children (avoid prolonged use); adrenal suppression if used on a large area of the body or for a long time; particularly with an occlusive dressing or on broken skin; avoid use on the face for more than 7 days; secondary infection requires treatment with an appropriate antimicrobial; may impair the ability to resist and counteract infections; diabetes mellitus; pregnancy (Appendix 7c), elderly; lactation.

**ADVERSE EFFECTS**
- Exacerbation of local infection; local atrophic changes particularly on the face and in skinfolds; characterized by thinning of the dermis; depigmentation; dilatation of superficial blood vessels and formation of striae; perioral dermatitis; acne at site of application; suppression of the hypothalamic-pituitary-adrenal axis with prolonged or widespread use (particularly under occlusion); subcapsular cataract; osteoporosis; glaucoma; intracranial hypertension; psychic instability.

**Liquid paraffin**  
**EDL-D 315 PHC**

**INDICATION**
- Constipation.

**CONTRAINDICATION**
- Children less than 3 years of age.

**PRECAUTION**
- Avoid prolonged use.

**ADVERSE EFFECT**
- Anal seepage of paraffin and consequent anal irritation after prolonged use. Granulomatous reactions caused by absorption of small quantities of liquid paraffin.

**AVAILABILITY**
- Oral emulsion, also combinations are available-Cremaffin.

**DOSE**
- 10 - 30 mL hs.

**DRUG INTERACTION**
- Not reported.
Hydrocortisone  
EDL-D 263 Secondary hospitals  
Schedule H  
INDICATIONS  
Ulcerative colitis, proctitis, proctosigmoiditis; anaphylaxis; skin; adrenocortical insufficiency.  
AVAILABILITY  
INJECTION 100 mg/vial; SUPPOSITORIES 25 mg; RETENTION ENEMA 60 ml (Rectal solution 100 mg/60 ml); cream 0.1% w/w.  
DOSE  
Rectal (suppositories)  
Adult- Ulcerative colitis, proctitis: 25 mg twice daily for 2 weeks; may be increased to 25 mg 3 times daily or 50 mg twice daily in severe cases; in factitial proctitis treatment may be required for 6 to 8 weeks.  
Rectal (retention enema)  
Adult- Ulcerative colitis, ulcerative proctitis, ulcerative proctosigmoiditis: 100 mg at night for 21 days or until clinical and proctological remission; if no clinical and proctological improvement after 21 days, discontinue; treatment for 2 to 3 months may be required for proctological remission; when used for more than 21 days, discontinue gradually using 100 mg every other night for 2 to 3 weeks.  
CONTRAINDICATIONS  
Use of enemas in bowel obstruction, bowel perforation, or extensive fistulas; untreated infections.  
PRECAUTIONS  
Proctological examination required before treatment; systemic absorption may occur; prolonged use should be avoided; lactation (Appendix 7b); interactions (Appendix 6d); pregnancy (Appendix 7c).  
ADVERSE EFFECTS  
Local pain or burning sensation; rectal bleeding (reported with use of enema); exacerbation of untreated infections; suppositories may stain fabrics; systemic adverse effects.  
STORAGE  
Store protected from light.  
Sulfasalazine  
EDL-D 493 Secondary hospitals  
INDICATIONS  
Ulcerative colitis; Crohn’s disease; severe rheumatoid arthritis; inflammatory bowel disease.  
AVAILABILITY  
TABLETS 500 and 1000 mg.  
Dose Oral  
Adult- Ulcerative colitis: 1 to 2g 4 times daily in acute attack until remission, reducing to maintenance dose of 500 mg 4 times daily. Active Crohn’s disease: 1 to 2g four times daily in acute attack until remission occurs.  
Child- Ulcerative colitis: over 2 years; 40 to 60 mg/kg daily in acute attack, reducing to maintenance dose of 20–30 mg/kg daily. Active Crohn disease: over 2 years, 40–60 mg/kg daily in acute attack.  
CONTRAINDICATIONS  
Hypersensitivity to salicylates or sulfonamides; child under 2 years; porphyria; intestinal or urinary obstruction; severe renal impairment; G-6-PD deficiency; blood dyscracias.  
PRECAUTIONS  
Renal impairment; hepatic impairment; G-6-PD deficiency; slow acetylator status; monitor blood counts and liver function initially and at monthly intervals for first 3 months; monitor kidney
function initially and at intervals during treatment; history of allergy; lactation (Appendix 7b); interactions (Appendix 6c, 6d). Patients should be advised to report any unexplained bleeding, bruising, purpura, sore throat, fever or malaise occurring during treatment; blood count should be performed and sulfasalazine stopped immediately if there is suspicion or evidence of blood disorder; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea, exacerbation of colitis; diarrhoea, loss of appetite, fever; blood disorders (including Heinz body anaemia, megaloblastic anaemia, leukopenia, neutropenia, thrombocytopenia); hypersensitivity reactions (including rash, urticaria, Stevens-Johnson syndrome (erythema multiforme), exfoliative dermatitis, epidermal necrolysis, pruritus, photosensitization, anaphylaxis, serum sickness, interstitial nephritis, lupus erythematosus-like syndrome); lung complications (including eosinophilia, fibrosing alveolitis); ocular complications (including periorbital oedema); stomatitis, parotitis; ataxia, aseptic meningitis, vertigo, tinnitus, alopecia, peripheral neuropathy, insomnia, depression, headache, hallucinations; kidney reactions (including proteinuria, crystalluria, haematuria); oligospermia; rarely, acute pancreatitis, hepatitis; urine may be coloured orange; some soft contact lenses may be stained.

STORAGE
Store protected from light.

**Antispasmodic medicines**

**Dicyclomine Hydrochloride**

**EDL-D 172,173 PHC**

**AVAILABILITY**
TABLETS 10 and 20 mg. DROPS 10 mg/ml; INJECTION 10 mg/ml.

**DOSE**
Oral Adult- 10-20 mg three times a day. Parenteral IM injection: 80 mg daily in 4 divided doses.

**INDICATION**
Infantile colic, gastrointestinal tract spasm.

**CONTRAINDICATION**
Glaucema, reflux oesophagitis, myasthenia gravis, lactation, intestinal obstruction

**PRECAUTION**
Patients with mental depression and mental disturbances, hepatic or kidney disease, angle closure glaucoma, hyperthyroidism, CHF, elderly, pregnancy, may impair the ability to drive or operate machinery; interactions(Appendix 6c); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
Dry mouth; nausea; vomiting; constipation; taste loss; anorexia; dizziness; dyskinesia; lethargy, respiratory arrest; drowsiness; photophobia, blurred vision; increased ocular pressure; tachycardia; urinary retention.

**Hyosine butyl bromide**

**EDL-D 269,270 PHC**

**INDICATION**
Intestinal, biliary and ureteric colics, spasmodic dysmenorrhoea, preparatory regimen for special radiological investigations such as hypotonic duodenography and for GI endoscopy.

**CONTRAINDICATION**
Intestinal obstruction, glaucoma, hepatic or renal failure, pregnancy, and lactation.

**PRECAUTION**
Avoid driving or operating machinery.

**ADVERSE EFFECT**
Dry mouth, thirst, increased intraocular pressure, flushing, palpitations followed by arrhythmias, constipation and difficulty in micturition, rashes.
AVAILABILITY
Tablets 10mg; Inj 20mg/ml Injection 20mg/ml
DOSE
Oral: 10 mg t.d.s
Parenteral route: 10 - 20 mg IM or IV 8 h.

DRUG INTERACTION
Other anticholinergic drugs and tricyclic antidepressants and alcohol potentiate the effects of hyoscine butyl bromide.

Laxatives
A balanced diet, including adequate fluid intake and fibre is of value in preventing constipation. Before prescribing laxatives, it is important to be sure that the patient is constipated and that the constipation is not secondary to an underlying undiagnosed complaint. It is also important that the patient understands that bowel habit can vary considerably in frequency without doing harm. For example, some people consider themselves constipated if they do not have a bowel movement each day. A useful definition of constipation is the passage of hard stools less frequently than the patient’s own normal pattern and this should be explained to the patient since misconceptions about bowel habits have led to excessive laxative use which in turn has led to hypokalaemia and an atonic non-functioning colon.

Laxatives should generally be avoided except where straining will exacerbate a condition such as angina or increase the risk of rectal bleeding as in haemorrhoids. Laxatives are of value in drug-induced constipation, for the expulsion of parasites after anthelminthic treatment and to clear the alimentary tract before surgery and radiological procedures. Prolonged treatment of constipation is rarely necessary except occasionally in the elderly.

There are many different laxatives. These include bulk-forming laxatives which relieve constipation by increasing faecal mass and stimulating peristalsis, stimulant laxatives which increase intestinal motility and often cause abdominal cramp, faecal softeners which lubricate and soften impacted faeces and osmotic laxatives which act by retaining fluid in the bowel by osmosis. Bowel cleansing solutions are used before colonic surgery, colonoscopy or radiological examination to ensure that the bowel is free of solid contents; they are not a treatment for constipation.

Bisacodyl
EDL-D 78 PHC D 570 Secondary hospitals
AVAILABILITY
TABLETS 5 mg; suppositories 5 and 10mg.
DOSE
Oral/Rectal Adult and child over 10 years- 5 to 10 mg daily at night. Before radiological procedure and surgery: 16 to 20 mg at night before procedure.
INDICATION
Constipation.
CONTRAINDICATION
Intestinal obstruction (causes abdominal cramps), acute surgical abdominal conditions, acute inflammatory bowel disease, severe dehydration; faecal impaction, chronic use
PRECAUTION
Excessive use of stimulant laxatives can cause diarrhoea and related effects such as hypokalaemia; however, prolonged use may be justifiable in some circumstances; don’t give antacid within 1 hour, pregnancy (Appendix 7c), inflammatory bowel disease, pre-existing heart disease or bowel disease, allergies, interactions
ADVERSE EFFECTS
Tablets- griping; suppositories-local irritation; fainting, dizziness, soreness in anal region due to suppository leakage; abdominal discomfort, electrolyte imbalance, hypokalaemia.

Ispaghula
Non-EDL PHC
INDICATIONS
Constipation; irritable colon syndrome.
AVAILABILITY
GRANULES (flavoured and sweetened) 37.5 and 100g.
DOSE
Oral
Adult- 6 teaspoonful of water or milk at night before bed time.
Child- 1-3 teaspoonful in water or milk before bed time.
CONTRAINDICATIONS
Intestinal obstruction; colonic atony; difficulty in swallowing.
PRECAUTIONS
Salt restriction; interactions (Appendix 6c).
ADVERSE EFFECTS
Abdominal discomfort, flatulence, gastrointestinal obstruction.
STORAGE
Store protected from light and moisture.

Lactulose
EDL- D 550 PHC
AVAILABILITY
SOLUTION/SYRUP 3.35g/5 ml.
DOSE
10 to 20g (15 to 20 ml/day, max 45 ml/day).
INDICATION
Constipation, hepatic encephalopathy.
CONTRAINDICATION
Galactosemia, intestinal obstruction, patients on low galactose diet.
PRECAUTION
Lactose intolerance, diabetes mellitus.
ADVERSE EFFECTS
Diarrhoea (dose related), nausea, vomiting, hypokalaemia; dehydration; hypernatremia; bloating and abdominal cramps.
Medicines used in diarrhoea

Acute diarrhoeal diseases are a leading cause of childhood morbidity and mortality; frail and elderly patients are also at risk. In adults acute diarrhoea is the most frequent health problem of travellers and is increasingly common among HIV-infected persons. Assessment and correction of dehydration and electrolyte disturbance is the priority in all cases of acute diarrhoea. Symptomatic relief in adults may be warranted in some cases but antidiarrhoeals should never be used in children since they do not reduce fluid and electrolyte loss and may cause adverse effects.

Diarrhoea persisting for longer than a month is known as chronic diarrhoea. A mild malabsorption syndrome, tropical enteropathy, is apparent in most healthy indigenous populations of tropical countries. However the majority of cases of chronic diarrhoea have non-infectious causes including gluten-sensitivity, inherited metabolic disorders or inflammatory bowel disease.

Bloody diarrhoea is usually a sign of invasive enteric infection and should be treated with an appropriate anti-infective agent.

Antidiarrhoeal Symptomatic Drugs in Adult

Codeine

Non-EDL Secondary hospitals

INDICATIONS
Short-term symptomatic relief of acute diarrhoea in adult; pain.

AVAILABILITY
TABLET 30 mg.

DOSE
Oral
Adult- Symptomatic relief of acute diarrhoea:
30 mg 3 to 4 times daily.
Child- (1-12 years) 500 μg/kg 4-6 times daily.

CONTRAINDICATIONS
Conditions where inhibition of peristalsis should be avoided; abdominal distension; acute diarrhoeal conditions such as ulcerative colitis or antibiotic-associated colitis; acute respiratory depression.

PRECAUTIONS
Tolerance or dependence may occur with prolonged use; elderly and debilitated patients; hepatic impairment (Appendix 7a); renal impairment; lactation; overdosage: see chapter 7.2; interactions (Appendix 6c); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea, vomiting, constipation, drowsiness; respiratory depression and hypotension (large doses); dependence; difficulty with micturition; ureteric or biliary spasm; dry mouth, sweating, headache, facial flushing, vertigo, bradycardia, tachycardia, palpitations, hypothermia, hallucinations, dysphoria, mood changes, miosis, decreased libido or potency, rash, urticaria, pruritus; convulsions (large doses).
**Furazolidone**  
Non-EDL Secondary hospitals

**INDICATIONS**  
Giardiasis; cholera; gastrointestinal infections; protozoal or bacterial diarrhea and enteritis; food poisoning.

**AVAILABILITY**  
TABLETS 100 mg; CAPSULE 100 mg; SUSPENSION 25 mg/5 ml.

**DOSE**  
Oral  
Adult- 100 mg 3 to 4 times a day.  
Child- 5 mg/kg body weight daily in 4 divided doses.

**CONTRAINDICATIONS**  
Hypersensitivity; alcoholics; primaquine sensitivity.

**PRECAUTIONS**  
Urine colour changes to yellow after administration; orthostatic hypotension; hypoglycaemia; pregnancy (Appendix 7c); interactions (Appendix 6a, 6c).

**ADVERSE EFFECTS**  
Nausea, vomiting, headache; hypotension; urticaria; dyspnea; dizziness.

**STORAGE**  
Store protected from light at temperature not exceeding 30°C.

**Loperamide**  
Non-EDL Secondary hospitals

**INDICATIONS**  
For the control and symptomatic relief of acute nonspecific diarrhoea and chronic diarrhoea associated with inflammatory bowel disease or gastroenteritis; for reducing the volume of discharge from ileostomies.

**AVAILABILITY**  
TABLET/CAPSULE 2 mg; LIQUID 1 mg/5 ml.

**DOSE**  
Oral  
Adult- 4 mg initially thereafter 2 mg after every motion.  
Child- 2 mg followed by 2 mg after every motion.

**CONTRAINDICATIONS**  
Conditions where inhibition of peristalsis should be avoided, where abdominal distension develops, or in conditions such as active ulcerative colitis or antibiotic-associated colitis.

**PRECAUTIONS**  
Liver disease; pregnancy: (Appendix 7c); interactions (Appendix 6c); glaucoma; Crohn’s disease; urinary bladder obstruction.

**ADVERSE EFFECTS**  
Abdominal cramps, dizziness, drowsiness and skin reactions including urticaria; paralytic ileus and abdominal bloating also reported; constipation; headache; meteorism; nausea; dry mouth; urinary retention.

**STORAGE**  
Store protected from light and moisture.
Oral Rehydration

Acute diarrhoea in children should always be treated with oral rehydration solution according to plan A, B or C as shown. Severely dehydrated patients must be treated initially with intravenous fluids until they are able to take fluids by mouth. For oral rehydration it is important to administer the solution in small amounts at regular intervals as indicated below.

Treatment of Dehydration:
Who Recommendations

According to the degree of dehydration, health professionals are advised to follow one of the three management plans.

Plan A: No dehydration: Nutritional advice and increased fluid intake are sufficient (soup, rice, water and yoghurt, or even water). For infants aged under 6 months who have not yet started taking solids, oral rehydration solution must be presented before offering milk. Mother’s milk or dried milk must be given without any particular restrictions. In the case of mixed breast-milk/formula feeding, the contribution of lactation must be increased.

Plan B: Moderate dehydration: Whatever the child’s age, a 4-h treatment plan is applied to avoid short-term problems. Feeding should not therefore be envisaged initially. It is recommended that parents are shown how to give approximately 75 ml/kg of oral rehydration solution with a spoon over a 4-h period and it is suggested that parents should be watched to see how they cope at the beginning of the treatment. A larger amount of solution can be given if the child continues to have frequent stools. In case of vomiting, rehydration must be discontinued for 10 min and then resumed at a slower rate (about one teaspoonful every 2 min). The child’s status must be re-assessed after 4 h to decide on the most appropriate subsequent treatment. Oral rehydration solution should continue to be offered once dehydration has been controlled, for as long as the child continues to have diarrhoea.

Plan C: Severe dehydration: Hospitalization is necessary, but the most urgent priority is to start rehydration. In hospital (or elsewhere), if the child can drink, oral rehydration solution must be given pending, and even during intravenous infusion (20 ml/kg every h by mouth before infusion, then 5 ml/kg every h by mouth during intravenous rehydration). For intravenous supplementation, it is recommended that compound solution of sodium lactate (see chapter 28.2) is administered at a rate adapted to the child’s age (infant under 12 months: 30 ml/kg over 1 h then 70 ml/kg over 5 h; child over 12 months:

**Oral Rehydration Salts**

**EDL-D 386,387 Universal**

**AVAILABILITY**

GLUCOSE SALT SOLUTION 5 and 37.5g. Sodium chloride 2.6 g/litre of water Sodium citrate 2.9 g/litre of water Potassium chloride 1.5 g/litre of water Glucose (anhydrous) 13.5 g/litre of water When glucose and sodium citrate are not available, they may be replaced by Sucrose (common sugar) 27 g/litre of water Sodium bicarbonate 2.5 g/litre of water In cases of cholera, oral rehydration salts containing a higher concentration of sodium may be required to prevent hyponatraemia.
DOSE
Oral 5g (single use): dissolve in water and drink; 37.5g: to reconstitute it with 1 litre of clean water. Adult- Fluid and electrolyte loss in acute diarrhoea; 200 to 400 ml solution after every loose motion.

INDICATION
Dehydration from acute diarrhoea.

PRECAUTION
Renal impairment.

ADVERSE EFFECTS
Vomiting- may indicate too rapid administration; hypernatraemia and hyperkalaemia may result from overdose in renal impairment or administration of too concentrated a solution.

Zinc sulfate
EDL-D 541Universal

INDICATION
Adjunct to oral rehydration therapy in acute diarrhea, acrodermatitis enteropathica, Wilson’s disease

ORAL LIQUID : In 10 mg and 20mg per unit dosage forms.
TABLET: in 10 mg and 20mg per unit dosage forms.
Daily requirements: 5-15mg/day

DOSE
in acute diarrhoea, infant under 6 months, 10 mg (elemental zinc) daily for 10–14 days; child 6 months–5years, 20 mg (elemental zinc) daily for 10–14 days, 25mg tid in Wilson’s disease.

ADVERSE EFFECTS
GI upset, copper deficiency
HORMONES, OTHER ENDOCRINE MEDICINES AND CONTRACEPTIVES

Adrenal hormones and synthetic substitutes
Corticosteroids include hormones secreted by the adrenal cortex and synthetic analogues of these hormones. The adrenal cortex normally secretes hydrocortisone which has glucocorticoid activity and weak mineralocorticoid activity. It also secretes the mineralocorticoid aldosterone. Synthetic glucocorticoids include betamethasone, dexamethasone and prednisolone. Fludrocortisone has glucocorticoid properties but it has potent mineralocorticoid properties and is used for its mineralocorticoid effects.

Pharmacology of the corticosteroids is complex and their actions are wide-ranging. In physiological (low) doses, they replace deficient endogenous hormones. In pharmacological (high) doses, glucocorticoids decrease inflammation and suppress the immune response.

In therapeutic doses glucocorticoids suppress release of corticotrophin (adrenocorticotropic hormone, ACTH) from the pituitary thus the adrenal cortex ceases secretion of endogenous corticosteroids. If suppressive doses are given for prolonged periods, the adrenal cortex may undergo atrophy and this leads to a deficiency on sudden withdrawal or dosage reduction or situations such as stress or trauma where corticosteroid requirements are increased. After high dosage or prolonged therapy, withdrawal should be gradual, the rate depending on various factors including patient response, corticosteroid dose, duration of treatment and disease state. The suppressive action of a corticosteroid on cortisol secretion is least when given in the morning. Corticosteroids should normally be given in a single morning dose to attempt to minimize pituitary-adrenal suppression. Because the therapeutic effects of corticosteroids are of longer duration than the metabolic effects, intermittent therapy may allow the body's normal metabolic rhythm and the therapeutic effects to be maintained. Alternate day dosing is, however, suitable only in certain disease states and with corticosteroids with small mineralocorticoid effects and a relatively short duration of action.

Hydrocortisone is used in adrenal replacement therapy and on a short-term basis by intravenous injection for the emergency management of some conditions. Its mineralocorticoid activity is too high for it to be used on a long-term basis for disease suppression. The mineralocorticoid activity of fludrocortisones is also high and its anti-inflammatory activity is of no clinical relevance. It is used together with glucocorticoids in adrenal insufficiency. Prednisolone has predominantly gluco corticoid activity and is the corticosteroid most commonly administered for long-term disease suppression. It is the active metabolite of prednisone, conversion of which is variable and prednisone should not be used interchangeably with prednisolone. Dexamethasone has very high glucocorticoid activity in conjunction with insignificant mineralocorticoid activity making it particularly suitable for high-dose therapy in conditions where water retention would be a disadvantage such as cerebral oedema. It also has a long duration of action and this, together with its lack of mineralocorticoid activity makes it particularly suitable for conditions requiring suppression of corticotrophin secretion such as congenital adrenal hyperplasia.
Adverse Effects of Corticosteroids:
Overdosage or prolonged use may exaggerate some of the normal physiological actions of corticosteroids leading to mineralocorticoid and glucocorticoid adverse effects.

Mineralocorticoid adverse effects include hypertension, sodium and water retention and potassium loss. These effects are most marked with fludrocortisone but are significant with hydrocortisone, occur slightly with prednisolone and are negligible with dexamethasone. Glucocorticoid adverse effects include diabetes mellitus and osteoporosis which is of particular importance in the elderly since it may result in osteoporotic fractures of the hip or vertebral. High doses may also be associated with avascular necrosis of the femoral neck. Muscle wasting may also occur and there is a weak link with peptic ulceration. Mental disturbances can occur, including serious paranoid state or depression with risk of suicide, particularly in patients with a history of mental disorders; euphoria is also common. High doses may cause Cushing syndrome (typical moon face, striae and acne), which is usually reversible on withdrawal of treatment, but this should always be tapered gradually to avoid symptoms of acute adrenal insufficiency (see also Withdrawal). In children, corticosteroids may result in suppression of growth and corticosteroids administered during pregnancy can affect adrenal development in the fetus. Any adrenal suppression in the neonate following prenatal exposure usually resolves spontaneously after birth and is rarely, clinically important. Healing of wounds may be impaired and infections and thinning of the skin may occur; spread of infections may result from modification of tissue reactions.

Adrenal Suppression
Adrenal suppression occurs during prolonged therapy with corticosteroids, with development of adrenal atrophy which may persist for years after stopping. Abrupt withdrawal after a prolonged period may lead to acute adrenal insufficiency, hypotension or death (see Withdrawal of Systemic Corticosteroids, below). Withdrawal may also be associated with fever, myalgia, arthralgia, rhinitis, conjunctivitis, painful itchy skin nodules and weight loss.

Corticosteroid Cover During Stress:
To compensate for a diminished adrenocortical response caused by prolonged corticosteroid treatment, any significant intercurrent illness, trauma, or surgery requires a temporary increase in corticosteroid dose, or if already stopped, a temporary re-introduction of corticosteroid treatment. Anaesthetists must therefore know whether a patient is taking or has been taking a corticosteroid, to avoid a precipitous fall in blood pressure during anaesthesia or in the immediate postoperative period. A suitable regimen for corticosteroid replacement, in patients who have taken more than 10 mg prednisolone daily (or equivalent) within 3 months of surgery, is:

- Minor surgery under general anaesthesia-usual oral corticosteroid dose on the morning of surgery or hydrocortisone 25-50 mg intravenously at induction; the usual oral corticosteroid dose is recommenced after surgery.
- Moderate or major surgery-usual oral corticosteroid dose on the morning of surgery and hydrocortisone 25-50 mg intravenously at induction, followed by hydrocortisone 25-50 mg 3 times a day by intravenous injection for 24 h after moderate surgery or for 48-72 h after major surgery; the usual preoperative oral corticosteroid dose is recommenced on stopping hydrocortisone injections.
**Infections:**

Prolonged courses of corticosteroids increase susceptibility to infections and increase their severity; clinical presentation of infections may also be atypical. Serious infections, for example septicaemia and tuberculosis, may reach an advanced stage before being recognised, and amoebiasis or strongyloidiasis may be activated or exacerbated (exclude before initiating a corticosteroid in those at risk or with suggestive symptoms). Fungal or viral ocular infections may also be exacerbated.

**Chickenpox**

Unless they have had chickenpox, patients receiving oral or parenteral corticosteroids for purposes other than replacement should be regarded as being at risk of severe chickenpox on exposure. Manifestations of fulminant illness include pneumonia, hepatitis and disseminated intravascular coagulation; rash is not necessarily a prominent feature.

Passive immunization with varicella-zoster immunoglobulin is needed for exposed non-immune patients receiving systemic corticosteroids or for those who have used them within the previous 3 months; varicella-zoster immunoglobulin should preferably be given within 3 days of exposure and no later than 10 days. Confirmed chickenpox warrants specialist care and urgent treatment. Corticosteroids should not be stopped and dosage may need to be increased. Topical, inhaled or rectal corticosteroids are less likely to be associated with an increased risk of severe chickenpox.

**Measles**

Patients taking corticosteroids should be advised to take particular care to avoid exposure to measles and to seek immediate medical advice if exposure occurs. Prophylaxis with intramuscular normal immunoglobulin may be needed.

**Dosage and Administration:**

Adverse effects of systemic glucocorticoids, including suppression of the Hypothalamo-Pituitary-Adrenal (HPA) axis, are doseand duration-dependent; thus patients should be given treatment for the shortest period at the lowest dose that is clinically necessary. Patient response is variable and doses should therefore be individualized. In life-threatening diseases, high doses may be needed because the complications of therapy are likely to be less serious than the disease. In long-term therapy in relatively benign chronic conditions such as rheumatoid arthritis, adverse effects often outweigh the advantages. In order to minimize the adverse effects, the maintenance dose should be kept as low as possible and if possible, single morning doses or alternate day therapy should be used. Glucocorticoids can improve the prognosis of serious conditions such as systemic lupus erythematosus, temporal arteritis and polyarteritis nodosa; in such disorders the effects of the disease process may be suppressed and symptoms relieved but the underlying condition is not cured.

Glucocorticoids are used both topically and systemically. In emergency situations, hydrocortisone may be given intravenously; in the treatment of asthma, inhalation therapy with beclomethasone may be used (chapter 20.1). Whenever possible, local treatment with creams, intra-articular injections, inhalations, eye-drops or enemas should be used in preference to systemic therapy.
Withdrawal of Systemic Corticosteroids:
The rate of withdrawal of systemic glucocorticoids is dependent upon several factors including size of dose, duration of treatment, individual patient’s response and the likelihood of relapse of the underlying disease. If there is uncertainty about suppression of the HPA axis, withdrawal should be gradual to enable the adrenal gland to recover. Patients should be advised not to stop taking glucocorticoids abruptly unless permitted by their doctor. Gradual withdrawal should be considered in those whose disease is unlikely to relapse and who have:

- recently received repeated courses (particularly if taken for longer than 3 weeks)
- taken a short course within 1 year of stopping longterm therapy
- other possible causes of adrenal suppression
- received more than 40 mg daily prednisolone (or equivalent)
- been given repeat doses in the evening
- received more than 3 weeks’ treatment

Abrupt withdrawal may be considered in those whose disease is unlikely to relapse and who have received treatment for 3 weeks or less and who are not included in the patient groups described above.

During corticosteroid withdrawal the dose may be reduced rapidly down to the physiological dosage (equivalent to 7.5 mg prednisolone daily) and then reduced more slowly. Assessment of the disease may be needed during withdrawal to ensure that relapse does not occur.

Hydrocortisone Sodium Succinate

**INDICATION**
Adrenocortical insufficiency, shock, hypersensitivity reactions (anaphylactic shock and angioedema), acute severe asthma.

**CONTRAINDICATION**
As for prednisolone (pg no.270)

**PRECAUTION**
As for prednisolone (pg no.270)

**ADVERSE EFFECTS**
As for Prednisolone (pg no.270)

**AVAILABILITY**
Tablets (as hydrocortisone), 5mg, 10mg, 20mg.
Injection (as sodium succinate), 100mg/vial (powder for reconstitution).

**DOSE**
ADULT: As hydrocortisone, I.M/IV 100-500mg 3-4 times in 24 hours; or as required.
CHILD: Oral, 4mg/m2 three times daily. Slow IV upto 1 year 25mg; 1-5 years, 50mg; 6-12 years 100mg.

Prednisolone

**INDICATIONS**
Suppression of inflammatory and allergic reactions; with antineoplastic drugs for acute leukaemias and lymphomas; asthma; rheumatic disorder; hematologic disorder.

**AVAILABILITY**
Tablets 5, 10, 20 and 40 mg; Injection 1 ml vial (40 mg/ml); SYRUP 60 ml (5 mg/5 ml and 15 mg/5 ml).

**DOSE**
Oral
Adult- Suppression of inflammatory and allergic disorders: initially up to 10 to 20 mg daily (severe disease, up to 60 mg daily), preferably taken in the morning after breakfast; dose can often be reduced within a few days, but may need to be continued for several weeks or months. Maintenance dose 2.5 to 15 mg daily or higher; cushingoid features are increasingly likely with doses above 7.5 mg daily.

Myasthenia gravis: initially 10 mg on alternate days, increased in steps of 10 mg on alternate days to 1-1.5 mg/kg (max. 100 mg) on alternate days or initially 5 mg daily increased in steps of 5 mg daily to usual dose of 60-80 mg daily (0.75-1 mg/kg daily).

Child- Fractions of adult dose may be used (At 1 year: 25% of adult dose; at 7 years: 50%; and at 12 years: 75%) but clinical factors must be given due weight.

CONTRAINDICATIONS
See notes above; systemic infection (unless life-threatening or specific antimicrobial therapy given); avoid live virus vaccines in those receiving immunosuppressive doses (serum antibody response diminished).

PRECAUTIONS
Refer notes above; hepatic impairment (Appendix 7a); lactation (Appendix 7b); interactions (Appendix 6c, 6d); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Refer Adverse effects of Corticosteroids.

Methyl Prednisolone
EDL-D 336 Secondary hospitals

INDICATIONS
Corticosteroid responsive conditions such as severe allergic rhinitis, asthma, rheumatoid arthritis, osteoarthritis, collagen disease, dermatoses.

AVAILABILITY
TABLETS 4, 8, 16 and 24 mg; INJECTION vials 40, 125, 500 and 1000 mg, 2 ml ampoule (80 mg/2 ml).

DOSE
Oral
Adult- Asthma, allergies and dermatological conditions: 40 and 120 mg.
Dose should be regulated in accordance with severity of condition; large joints- 20 to 80 mg; medium joints- 10 to 40 mg; small joints- 4 to 10 mg directly in bursae.

CONTRAINDICATIONS
Systemic fungal infection (unless specific antimicrobial therapy given); avoid live virus vaccines in those receiving immunosuppressive doses (serum antibody response diminished); hypersensitivity.

PRECAUTIONS
Refer notes above; interactions (Appendix 6c, 6d); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Refer adverse effects of corticosteroids.

STORAGE
Store protected from light at a temperature not exceeding 30°C. The injection should not be allowed to freeze.

Contraceptives
Oral Hormonal Contraceptives
Hormonal contraception is one of the most effective methods of reversible fertility control.
**Combined Oral Contraceptives:**

Estrogen plus progestogen combinations are the most widely used hormonal contraceptives. They produce a contraceptive effect mainly by suppressing the hypothalamic-pituitary system resulting in prevention of ovulation; in addition, changes in the endometrium make it unreceptive to implantation.

Endometrial proliferation is usually followed by thinning or regression of the endometrium resulting in reduced menstrual flow. Ovulation usually resumes within three menstrual cycles after oral contraception has been discontinued; anovulation and amenorrhea persisting for six months or longer requires investigation and appropriate treatment if necessary.

Potential non-contraceptive benefits of combined oral contraceptives include improved regularity of the menstrual cycle, decreased blood loss, less iron-deficiency anaemia and significant decrease in dysmenorrhoea. Long-term use is associated with reduced risk of endometrial and ovarian cancer and of some pelvic infections.

An association between the amount of estrogen and progestogen in oral contraceptives and an increased risk of adverse cardiovascular effects has been observed. The use of oral contraceptive combinations containing the progestogens, desogestrel or gestodene are associated with a slightly increased risk of venous thromboembolism compared with oral contraceptives containing the progestogens, levonorgestrel or norethisterone.

**Risk Factors for Venous Thromboembolism or Arterial Disease:**

Risk factors for venous thromboembolism include family history of venous thromboembolism in first-degree relative aged under 45 years, obesity, long-term immobilization and varicose veins.

Risk factors for arterial disease include family history of arterial disease in first-degree relative aged under 45 years, diabetes mellitus, hypertension, smoking, age over 35 years (avoid if over 50 years), obesity and migraine.

If any one of the factors is present, combined oral contraceptives should be used with caution; if 2 or more factors for either venous thromboembolism or arterial disease are present, combined oral contraceptives should be avoided. Combined oral contraceptives are contraindicated in migraine with aura, in severe migraine without aura regularly lasting over 72 h despite treatment and in migraine treated with ergot derivatives.

**Surgery:**

Estrogen-containing oral contraceptives should preferably be discontinued (and adequate alternative contraceptive arrangements made) 4 weeks before major elective surgery and all surgery to the legs or surgery which involves prolonged immobilization of a lower limb. They should normally be restarted at the first menses occuring at least 2 weeks after full mobilization. When discontinuation is not possible thromboprophylaxis (with heparin and graduated compression hosiery) is advised.

**Reasons to Stop Combined Oral Contraceptives Immediately:**

Combined estrogen-containing oral contraceptives should be stopped immediately if any of the following symptoms occur and resumed only after consultation with a health care provider:

- Sudden severe chest pain (even if not radiating to left arm);
• Sudden breathlessness (or cough with blood-stained sputum);
• Severe pain in calf of one leg;
• Severe stomach pain;
• Serious neurological effects including unusual, severe, prolonged headache especially if first time or getting progressively worse or sudden partial or complete loss of vision or sudden disturbance of hearing or other perceptual disorders or dysphagia or bad fainting attack or collapse or first unexplained epileptic seizure or weakness, motor disturbances, very marked numbness suddenly affecting one side or one part of body;
• Hepatitis, jaundice, liver enlargement;
• Blood pressure above 160 mmHg systolic and 100 mmHg diastolic;
• Detection of 2 or more risk factors for venous thromboembolism or arterial disease.

Progestogen-Only Contraceptives:
Progestogen-only contraceptives, such as oral levonorgestrel may offer a suitable alternative when estrogens are contraindicated but the oral progestogen-only preparations do not prevent ovulation in all cycles and have a higher failure rate than combined estrogen-containing preparations. Progestogenonly contraceptives carry less risk of thromboembolic and cardiovascular disease than combined oral contraceptives and are preferable for women at increased risk of such complications, for example smokers over 35 years. They can be used as an alternative to estrogen-containing combined preparations prior to major surgery. Oral progestogen-only contraceptives may be started 3 weeks after birth; lactation women should preferably start at least 6 weeks after birth. Menstrual irregularities (oligomenorrhoea, menorrhagia, amenorrhoea) are common. Injectable preparations of medroxyprogesterone acetate or norethisterone enantate may be given intramuscularly. They have prolonged action and should only be given with full counselling and manufacturer’s information leaflet.

Emergency Contraception:
Levonorgestrel is used for emergency contraception. Levonorgestrel 1.5 mg should be taken as a single dose within 72 h of unprotected intercourse; alternatively, levonorgestrel 750 μg can be taken within 72 h of unprotected intercourse followed 12 h later by another 750 μg. Under these circumstances levonorgestrel prevents about 86% of pregnancies that would have occurred if no treatment had been given. Adverse effects include nausea, vomiting, headache, dizziness, breast discomfort, and menstrual irregularities. If vomiting occurs within 2-3 h of taking the tablets, replacement tablets can be given with an antiemetic. It should be explained to the woman that her next period may be early or late; that she needs to use a barrier contraceptive method until her next period, and that she should return promptly if she has any lower abdominal pain or if the subsequent menstrual bleed is abnormally light, heavy, brief or absent. There is no evidence of harmful effects to the fetus if pregnancy should occur.

Hormone Releasing IUD
Non-EDL PHC

INDICATIONS
• For contraception.

AVAILABILITY
At Family Welfare clinics or specialty centres.
DOSE

For contraception, the device can be inserted at any time between day 4 and day 12 after the start of menstrual bleeding; not to be fitted during heavy menstrual bleeding. Emergency contraception, the device may be inserted up to 120 h (5 days) after unprotected intercourse, at any time of menstrual cycle; if intercourse has occurred more than 5 days previously, device can still be inserted up to 5 days after the earliest likely calculated day of ovulation; device can be removed at the beginning of menstruation if no longer required.

CONTRAINDICATIONS
Abnormal pap smear or abnormal vaginal bleeding.

ADVERSE EFFECTS
Heavy bleeding, perforation of uterus; cramps.

IUD Containing Copper
EDL-D 141 PHC

INDICATIONS
Contraception; emergency contraception.

AVAILABILITY
Single IUD in pouch pack.

DOSE

For contraception, the device can be inserted at any time between day 4 and day 12 after the start of menstrual bleeding; not to be fitted during heavy menstrual bleeding. Emergency contraception, the device may be inserted up to 120 h (5 days) after unprotected intercourse, at any time of menstrual cycle; if intercourse has occurred more than 5 days previously, device can still be inserted up to 5 days after the earliest likely calculated day of ovulation; device can be removed at the beginning of menstruation if no longer required.

CONTRAINDICATIONS
Pregnancy; 48h-4 weeks post partum; puerperal sepsis; postseptic abortion; cervical or endometrial cancer; pelvic inflammatory disease; recent sexually transmitted disease (if not fully investigated and treated); pelvic tuberculosis; unexplained uterine bleeding; malignant gestational trophoblastic disease; distorted or small uterine cavity; copper allergy; Wilson’s disease; medical diathermy; abnormal pap smear or abnormal vaginal bleeding.

PRECAUTIONS
Anaemia; heavy menstrual bleeding, endometriosis, severe primary dysmenorrhoea, history of pelvic inflammatory disease, history of ectopic pregnancy or tubal surgery, fertility problems, nulliparity and young age, severely scarred uterus or severe cervical stenosis, valvular heart disease (requires antibacterial cover)-avoid if prosthetic valve or history of endocarditis; HIV infection or immunosuppressive therapy (risk of infectionavoid if marked immunosuppression); joint and other prostheses; increased risk of expulsion if inserted before uterine involution; gynaecological examination before insertion and 4-6 weeks afterwards-counsel women to see doctor promptly if significant symptoms such as pain; anticoagulant therapy; remove if pregnancy occurs (consider possibility of ectopic pregnancy).

ADVERSE EFFECTS
Uterine or cervical perforation, displacement, expulsion; pelvic infection exacerbated; heavy menstrual bleeding; dysmenorrhoea; pain and bleeding and occasionally epileptic seizure or vasovagal attack on insertion.
EDL-D 207, 208 Secondary hospitals

AVAILABILITY
TABLETS 0.01, 0.05 and 1 mg; INJECTION 1 ml ampoule (10 mg/ml).

DOSE
Oral  Adult- Hormone replacement: 10 to 20 μg daily. Palliation in breast cancer in postmenopausal women: 0.1 to 1 mg 3 times daily.

INDICATION
Hormone replacement for menopausal symptoms; osteoporosis prophylaxis; palliation in breast cancer in men and postmenopausal women; contraception in combination with a progestogen; dysfunctional uterine bleeding, prostatic carcinoma

CONTRAINDICATION
Use within 3 weeks of birth; lactation until weaning or for first 6 months after birth (Appendix 7b); personal history of 2 or more risk factors for venous or arterial thrombosis (see notes above); heart disease associated with pulmonary hypertension or risk of embolism; migraine (see below); history of sub-acute bacterial endocarditis; ischaemic cerebrovascular disease; liver disease (Appendix 7a), including disorders of hepatic secretion such as Dubin-Johnson or Rotor syndromes, infectious hepatitis (until liver function normal); porphyria; systemic lupus erythematosus; liver adenoma; history of cholestasis with oral contraceptives; gallstones; estrogen-dependent neoplasms; neoplasms of breast or genital tract; undiagnosed vaginal bleeding; history of pruritus during pregnancy, chorea, herpes, deteriorating otosclerosis, cholestatic jaundice; diabetes mellitus (if either retinopathy, neuropathy or if more than 20 years duration); after evacuation of hydatidiform mole (until return to normal of urine and plasma gonadotrophin values). Migraine with typical focal aura; migraine without aura regularly lasting over 72 h duration despite treatment; migraine treated with ergot derivatives; migraine without focal aura or controlled with 5-HT1 agonist.

PRECAUTION
Pregnancy (Appendix 7c); estrogendependent cancer; active thrombophlebitis or thromboembolic disorders or history of recent venous thromboembolism (unless already on anticoagulant therapy); undiagnosed vaginal bleeding; lactation ; liver disease (where liver function tests have failed to return to normal), Dubin-Johnson and Rotor syndromes (or monitor closely); jaundice; cerebrovascular disease; hepatic carcinoma; CV disease; estrogen dependent cancer

ADVERSE EFFECTS
Nausea, vomiting, headache; breast tenderness; increase in body weight; thrombosis; changes in libido; depression; chorea; skin reactions; chloasma; hypertension; impairment of liver function; ‘spotting’ in early cycles; absence of withdrawal bleeding; breast cancer (small increase in risk of breast cancer during use which reduces during the 10 years after stopping; risk factor seems related to age at which contraceptive is stopped rather than total duration of use; small increase in risk of breast cancer should be weighed against the protective effect against cancers of the ovary and endometrium which persists after stopping); dizziness, stomach upset, bloating, mental and mood changes.

Ethinylestradiol + Levonorgestrel
EDL-D 209, 210 PHC

AVAILABILITY
TABLETS Levonorgestrel + Ethinylestradiol 0.15 mg + 0.03 mg 0.25 mg + 0.05 mg Levonorgestrel 0.15 mg + Ethinylestradiol 0.03 mg + Ferrous fumarate 60 mg. Norethisterone + Ethinylestradiol 0.5 mg + 0.03 mg 1.0 mg + 0.03 mg
DOSE
Oral Adult- Contraception: 1 tablet (pill) daily for 21 days; subsequent courses repeated after 7-day pill-free interval (during which withdrawal bleeding occurs). Each tablet (pill) should be taken at approximately the same time each day; if delayed by longer than 24 h contraceptive protection may be lost. It is important to bear in mind that the critical time for loss of protection is when a pill is omitted at the beginning or end of a cycle (which lengthens the pill-free interval).

INDICATION
Contraception; menstrual symptoms; endometriosis

CONTRAINDICATION
Use within 3 weeks of birth; lactation until weaning or for first 6 months after birth (Appendix 7b); personal history of 2 or more risk factors for venous or arterial thrombosis (see notes above); heart disease associated with pulmonary hypertension or risk of embolism; migraine (see below); history of sub-acute bacterial endocarditis; ischaemic cerebrovascular disease; liver disease (Appendix 7a), including disorders of hepatic secretion such as Dubin-Johnson or Rotor syndromes, infectious hepatitis (until liver function normal); porphyria; systemic lupus erythematosus; liver adenoma; history of cholestasis with oral contraceptives; gallstones; estrogen-dependent neoplasms; neoplasms of breast or genital tract; undiagnosed vaginal bleeding; history of pruritus during pregnancy, chorea, herpes, deteriorating otosclerosis, cholestatic jaundice; diabetes mellitus (if either retinopathy, neuropathy or if more than 20 years duration); after evacuation of hydatidiform mole (until return to normal of urine and plasma gonadotrophin values). Migraine with typical focal aura; migraine without aura regularly lasting over 72 h duration despite treatment; migraine treated with ergot derivatives; migraine without focal aura or controlled with 5-HT1 agonist.

PRECAUTION
Risk factors for venous thromboembolism and arterial disease (see notes above); migraine (see below); hyperprolactinaemia (seek specialist advice); some types of hyperlipidaemia; gallbladder disease; severe depression; long-term immobilization (see also Travel below); sickle-cell disease; inflammatory bowel disease including Crohn's disease, interactions (Appendix 6c, 6d). Patients should report any increase in headache frequency or onset of focal symptoms (discontinue immediately and refer urgently to neurology expert if focal neurological symptoms not typical of aura persist for more than one hour). Women taking oral contraceptives may be at increased risk of deep-vein thrombosis during travel involving long periods of immobility (over 5 h). The risk may be reduced by appropriate exercise during the journey and possibly by wearing elastic hosiery; pregnancy

ADVERSE EFFECTS
Nausea, vomiting, headache; breast tenderness; increase in body weight; thrombosis; changes in libido; depression; chorea; skin reactions; chloasma; hypertension; impairment of liver function; ‘spotting’ in early cycles; absence of withdrawal bleeding; breast cancer (small increase in risk of breast cancer during use which reduces during the 10 years after stopping; risk factor seems related to age at which contraceptive is stopped rather than total duration of use; small increase in risk of breast cancer should be weighed against the protective effect against cancers of the ovary and endometrium which persists after stopping); dizziness, stomach upset, bloating, mental and mood changes.

Levonorgestrel
EDL-D 303 PHC

INDICATIONS
Emergency hormonal contraception.

AVAILABILITY
TABLETS 0.75 and 1.5 mg.
DOSE

Oral

Adult- Contraception: 1 tablet (‘pill’) (30 μg) daily, starting on the first day of the cycle and then continuously.

CONTRAINDICATIONS

Progestogen-only oral contraceptives; undiagnosed vaginal bleeding; severe arterial disease; liver tumours; breast cancer; thromboembolic disorders; sickle cell anaemia; porphyria; after evacuation of hydatidiform mole (until return to normal urine and plasma gonadotrophin values); progestogen-only emergency hormonal contraceptives; severe liver disease.

PRECAUTIONS

Possible small increase in risk of breast cancer; cardiac disease; sex-steroid dependent cancer; past ectopic pregnancy; malabsorption syndrome; ovarian cysts; active liver disease, recurrent cholestatic jaundice, history of jaundice in pregnancy (Appendix 7c); increase in frequency or severity of headache (discontinue pending investigation); lactation (Appendix 7b); pregnancy (Appendix 7c).

ADVERSE EFFECTS

Menstrual irregularities (including oligomenorrhoea and menorrhagia); nausea, vomiting, headache, dizziness; breast discomfort, depression; skin disorders; disturbances of appetite; weight increase; change in libido.

Insulin and other antidiabetic agents

Diabetes mellitus is characterized by hyperglycaemia and disturbances of carbohydrate, fat and protein metabolism. There are 2 principal types of diabetes.

Type-1 diabetes or insulin-dependent diabetes mellitus is due to a deficiency of insulin caused by autoimmune destruction of pancreatic β-cells. Patients require administration of insulin.

Type-2 diabetes or non-insulin dependent diabetes mellitus is due to reduced secretion of insulin or to peripheral resistance to the action of insulin. Patients may be controlled by diet alone, but often require administration of oral antidiabetic drugs or insulin. The energy and carbohydrate intake must be adequate but obesity should be avoided. In type 2 diabetes, obesity is one of the factors associated with insulin resistance. Diets high in complex carbohydrate and fibre and low in fat are beneficial. Emphasis should be placed on exercise and increased activity.

The aim of treatment is to achieve the best possible control of plasma glucose concentration and prevent or minimize complications including microvascular complications (retinopathy, albuminuria, neuropathy). Diabetes mellitus is a strong risk factor for cardiovascular disease; other risk factors such as smoking, hypertension, obesity and hyperlipidaemia should also be addressed.

Insulin

Appropriate insulin regimens should be worked out for each patient. Insulin requirements may be affected by variations in lifestyle (diet and exercise)-drugs such as corticosteroids, infections, stress, accidental or surgical trauma, puberty and pregnancy (second and third trimesters) may increase insulin requirements; renal or hepatic impairment and some endocrine disorders (for
example Addison’s disease, hypopituitarism) or coeliac disease may reduce requirements. In pregnancy insulin requirements should be monitored frequently.

If possible patients should monitor their own blood-glucose concentration using blood glucose strips. Since blood-glucose concentration varies throughout the day, patients should aim to maintain blood-glucose concentration between 4 and 9 mmol/litre (4-7 mmol/L before meals, <9 mmol/L) for most of the day while accepting that on occasions it will be higher; strenuous efforts should be made to prevent blood-glucose concentrations falling below 4 mmol/litre because of the risk of hypoglycaemia. Patients should be advised to look for troughs and peaks of blood glucose and to adjust their insulin dosage only once or twice a week. Insulin doses are determined on an individual basis, by gradually increasing the dose to optimise blood-glucose concentration while avoiding hypoglycaemia.

In the absence of blood-glucose monitoring strips, urine-glucose monitoring strips can be used; in fact this is the method of personal choice for many patients with Type 2 diabetes mellitus. It is less reliable than blood glucose but is easier and costs much less. All patients should monitor either blood- or urine-glucose concentration daily.

Hypoglycaemia is a potential complication in all patients treated with insulin or oral hypoglycaemic agents. The consequences of hypoglycaemia include confusion, seizures, coma and cerebral infarction.

Loss of warning of hypoglycaemia is common among insulintreated patients and can be a serious hazard especially for drivers and those in dangerous occupations. Very tight control lowers the blood glucose concentration needed to trigger hypoglycaemic symptoms; increase in the frequency of hypoglycaemic episodes reduces the warning symptoms experienced by patients. Beta-blockers can also blunt hypoglycaemic awareness (and delay recovery). Some patients report loss of hypoglycaemic warning after transfer to human insulin. Clinical studies do not confirm that human insulin decreases hypoglycaemic awareness. If a patient believes that human insulin is responsible for loss of warning it is reasonable to revert to animal insulin. To restore warning signs, episodes of hypoglycaemia must be reduced to a minimum; this involves appropriate adjustment of insulin dose and frequency, and suitable timing and quantity of meals and snacks.

Drivers need to be particularly careful to avoid hypoglycaemia. They should check their blood-glucose concentration before driving and, on long journeys, at intervals of approximately two hour; they should ensure that a supply of sugar is always readily available. If hypoglycaemia occurs, the driver should stop the vehicle in a safe place, ingest a suitable sugar supply and wait until recovery is complete (may be 15 min or longer). Driving is particularly hazardous when hypoglycaemic awareness is impaired.

For sporadic physical activity, extra carbohydrate may need to be taken to avert hypoglycaemia. Blood glucose should be monitored before, during and after exercise. Hypoglycaemia can develop in patients taking oral antidiabetics, notably the sulfonylureas, but this is uncommon and usually indicates excessive dosage. Sulfonylurea-induced hypoglycaemia may persist for several hour and must be treated in hospital.
Diabetic ketoacidosis is a potentially lethal condition caused by an absolute or relative lack of insulin; it commonly occurs when adjustments to insulin dosage fail to compensate for increases in insulin requirements, for example during severe infection or major intercurrent illness. Diabetic ketoacidosis occurs mostly in patients with Type 1 diabetes mellitus. It also occurs in Type 2 diabetics who have a temporary need for insulin. Diabetic ketoacidosis is characterized by hyperglycaemia, hyperketonaemia and acidaemia with dehydration and electrolyte disturbances. It is essential that soluble insulin (and intravenous fluids) is readily available for its treatment.

Infections are more likely to develop in patients with poorly controlled diabetes mellitus. These should be treated promptly and effectively to avoid diabetic ketoacidosis.

Surgery: Particular attention should be paid to insulin requirements when a patient with diabetes undergoes surgery that is likely to need an intravenous infusion of insulin for longer than 12 h. Soluble insulin should be given in intravenous infusion of glucose and potassium chloride (provided the patient is not hyperkalaemic), and adjusted to provide a blood-glucose concentration of between 7 and 12 mmol/litre. The duration of action of intravenous insulin is only a few min therefore the infusion must not be stopped unless the patient becomes frankly hypoglycaemic. For non-insulin dependent diabetics, insulin treatment is almost always required during surgery (oral hypoglycaemic drugs having been omitted).

Insulin must be given by injection because it is inactivated by gastrointestinal enzymes. Generally, insulin is given by subcutaneous injection into the upper arms, thighs, buttocks, or abdomen. There may be increased absorption from a limb, if the limb is used in strenuous exercise following the injection. It is essential to use only syringes calibrated for the particular concentration of insulin administered.

There are three main types of insulin preparations, classified according to duration of action after subcutaneous injection:

- those of short duration which have a relatively rapid onset of action, for example soluble or neutral insulin;
- those with an intermediate action, for example isophane insulin and insulin zinc suspension;
- those with a relatively slow onset and long duration of action, for example crystalline insulin zinc suspension.

Soluble insulin, when injected subcutaneously, has a rapid onset of action (after 30-60 min), a peak action between 2 and 4 h, and a duration of action up to 8 h. Soluble insulin by the intravenous route is reserved for urgent treatment and fine control in serious illness and perioperative state. When injected intravenously, soluble insulin has a very short half-life of only about 5 min.

When injected subcutaneously, intermediate-acting insulins have an onset of action of approximately 1-2 h, a maximal effect at 4-12 h and a duration of action of 16-24 h. They can be given twice daily together with short-acting insulin or once daily, particularly in elderly patients. They can be mixed with soluble insulin in the syringe, essentially retaining properties of each component.
The duration of action of different insulin preparations varies considerably from one patient to another and this needs to be assessed for every individual. The type of insulin used and its dose and frequency of administration depend on the needs of each patient. For patients with acute onset diabetes mellitus, treatment should be started with soluble insulin given 3 times daily with medium-acting insulin at bedtime. For those less seriously ill, treatment is usually started with a mixture of pre-mixed short- and medium-acting insulins (for example 30% soluble insulin with 70% isophane insulin) given twice daily. The proportions of soluble insulin can be increased in patients with excessive post-prandial hyperglycaemia.

**Oral Antidiabetic Drugs**

Oral antidiabetic (hypoglycaemic) drugs are used for non-insulin-dependent diabetes mellitus in patients who do not respond to dietary adjustment and an increase in physical exercise. They are used to supplement the effect of diet and exercise. There are various types of oral antidiabetic agents. The most commonly used are the sulfonylureas and the biguanide, metformin.

Sulfonylureas act mainly by augmenting insulin secretion and are therefore only effective if there is some residual pancreatic beta-cell activity. They may occasionally lead to hypoglycaemia 4 h or more after food. This may be dose-related and usually indicates excessive dose and it occurs more frequently with long-acting sulfonylureas such as glibenclamide and occurs particularly in the elderly. The sulfonylureas have the disadvantage that they may encourage weight gain. They should not be used during lactation and caution is required in the elderly and those with renal or hepatic insufficiency because of the risk of hypoglycaemia. Insulin therapy is generally required during intercurrent illness such as myocardial infarction, coma, infection, and trauma, during surgery and also during pregnancy.

Metformin exerts its effect by decreasing gluconeogenesis and by increasing peripheral utilization of glucose. Metformin can only act in the presence of endogenous insulin therefore is effective only in diabetics with some residual functioning pancreatic islet cells. It is used as a first-line treatment in overweight non-insulin-dependent diabetic patients and in others when strict dieting and sulfonylureas have failed to control the disease. Gastrointestinal adverse effects are common on initial treatment and may persist, particularly when very high doses (such as 3g daily) are given. In order to reduce gastrointestinal effects, treatment should be initiated with a low dose which may be gradually increased. Metformin may provoke lactic acidosis which is most likely to occur in patients with renal impairment; it should not be used in patients with even mild renal impairment.

One major advantage of metformin is that it does not usually cause hypoglycaemia. It may be used together with insulin (but weight gain and hypoglycaemia can be a problem) or sulfonylureas (but possibility of increased adverse effects with such combinations). During medical and surgical emergencies insulin treatment is almost always required; insulin should be substituted for metformin before elective surgery and in pregnancy.
**Glibenclamide**  
EDL-D 245,246 Secondary hospitals  

**AVAILABILITY**  
TABLETS 1.25, 2.5 and 5 mg.

**DOSE**  
Oral Adult- initially 5 mg once daily with or immediately after breakfast; max. 15 mg daily. Elderly- 2.5 mg, but it should preferably be avoided, adjusted according to response (max. 15 mg daily).

**INDICATION**  
Type II diabetes mellitus.

**CONTRAINDICATION**  
Ketoacidosis; porphyria; lactation

**PRECAUTION**  
Renal impairment; hepatic impairment (Appendix 7a); elderly; substitute insulin during severe infection, trauma, surgery (see notes above); interactions (Appendix 6b, 6c); diabetic coma; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**  
Mild and infrequent, including gastrointestinal disturbances and headache; liver disorders; hypersensitivity reactions usually in first 6-8 weeks; rarely; erythema multiforme, exfoliative dermatitis, fever and jaundice; hypoglycaemia, particularly in the elderly; rarely, blood disorders including leukopenia, thrombocytopenia, agranulocytosis, pancytopenia, haemolytic anaemia, and aplastic anaemia; cholestatic jaundice.

**Metformin Hydrochloride**  
EDL-D 333 Secondary hospitals  

**AVAILABILITY**  
TABLETS 250, 500, 850 mg, and 1g.

**DOSE**  
Oral Adult- Diabetes mellitus: initially 500 mg with breakfast for at least 1 week, then 500 mg with breakfast and evening meal for at least 1 week, then 500 mg with breakfast, lunch and evening meal or 850 mg every 12 h with or after food (max. 2g daily in divided doses).

**INDICATION**  
Diabetes mellitus

**CONTRAINDICATION**  
Renal impairment (withdraw if renal impairment suspected ; withdraw if tissue hypoxia likely (for example sepsis, respiratory failure, recent myocardial infarction, hepatic impairment), use of iodine-containing X-ray contrast media (do not restart metformin until renal function returns to normal) and use of general anaesthesia (suspend metformin 2 days beforehand and restart when renal function returns to normal); alcohol dependence; pregnancy; anaemia; ketosis.

**PRECAUTION**  
Measure serum creatinine before treatment and once or twice annually during treatment; substitute insulin during severe infection; trauma, surgery (see notes above and contraindications); lactation (Appendix 7b); interactions (Appendix 6a, 6b, 6c); hepatic or renal disease (Appendix 7a); heart disease.

**ADVERSE EFFECTS**  
Anorexia, nausea and vomiting, diarrhoea (usually transient), abdominal pain, metallic taste; lactic acidosis most likely in patients with renal impairment (discontinue); decreased vitamin B12 absorption.
**Glipizide**  
**EDL-D 547.548,549 Secondary hospitals**  
**AVAILABILITY**  
TABLETS 2.5, 5, 7.5 and 10 mg.  
**DOSE**  
2.5-20 mg once or twice daily. Maximum 40 mg daily.  
**INDICATION**  
Type II diabetes mellitus.  
**CONTRAINDICATION**  
Hypersensitivity; type I diabetes mellitus, ketoacidosis with or without coma; severe hepatic or renal insufficiency; pregnancy, lactation  
**PRECAUTION**  
Stress; fever; trauma; infection or surgery; elderly; thyroid impairment; monitor blood glucose concentration.  
**ADVERSE EFFECTS**  
Hypoglycemia, nausea, diarrhoea, allergic skin reactions, thrombocytopenia, leucopenia, agranulocytosis, jaundice, hemolytic anaemia.  

**Gliclazide**  
**Non-EDL Secondary hospitals**  
**INDICATIONS**  
Type II diabetes mellitus.  
**AVAILABILITY**  
TABLETS 20, 30, 40, 80 and 160 mg; MODIFIED RELEASE TABLETS 30 and 60 mg; CAPSULES 30, 40, 60 and 80 mg.  
**DOSE**  
40-320 mg daily, doses >160 mg daily may be given in 2 divided doses.  
Modified release tablets 30-120 mg daily.  
**CONTRAINDICATIONS**  
Type I diabetes mellitus, severe renal and hepatic impairment, diabetic ketoacidosis, pregnancy (Appendix 7c), lactation.  
**PRECAUTIONS**  
Monitor blood glucose concentration, increased risk of hypoglycaemia in elderly; debilitated patients; renal and hepatic impairment, metabolic stressful situations; interactions (Appendix 6c).  
**ADVERSE EFFECTS**  
Cutaneous reactions; blood dyscrasias, gastrointestinal disturbances; cholestatic jaundice.  

**Glimepiride**  
**EDL-D 545,546 Secondary hospitals**  
**INDICATIONS**  
Type II diabetes mellitus.  
**AVAILABILITY**  
TABLETS 1, 2, 3 and 4 mg.  
**DOSE**  
Adult 1-2 mg daily. Max dose 8 mg daily.  
**CONTRAINDICATIONS**  
Hypersensitivity; pregnancy (Appendix 7c); diabetic ketoacidosis.
PRECAUTIONS
Elderly; hepatic and renal impairment; interactions (Appendix 6b, 6c); monitor blood-glucose concentration; lactation.

ADVERSE EFFECTS
Hypoglycaemia; weight gain.

STORAGE
Store protected from moisture at temperature not exceeding 30°C.

Glucagon
Non-EDL Tertiary

INDICATIONS
Severe hypoglycaemia and radiological examination of gastrointestinal tract.

AVAILABILITY
Injection (powder for reconstitution) - 1 mg vial with pre-filled syringe containing water for injection.

DOSE
Parenteral Severe hypoglycaemia:
Adult and child over 8 years (or body weight over 25 kg) - 1 mg by s.c, i.m or i.v route.
Child under 8 years (or body weight under 25 kg) - 500 μg, if no response within 10 minutes i.v glucose must be given.
As diagnostic aid in gastrointestinal examination: Adult - 1-2 mg by i.m or 0.2-2 mg by i.v. injection.
Diagnosis of pheochromocytoma: 1 mg i.v.

CONTRAINDICATIONS
Pheochromocytoma; hypersensitivity.

PRECAUTIONS
Patients with insulinoma, glucagonoma, monitor prothrombin time, starvation and adrenal insufficiency, ineffective in chronic hypoglycaemia, alcohol-induced hypoglycaemia, pregnancy (Appendix 7c), lactation, interactions (Appendix 6b, 6c).

ADVERSE EFFECTS
Hypokalemia; nausea, vomiting, abdominal pain; rarely, hypersensitivity.

Insulin
EDL D 276, 277 Secondary hospitals

INDICATIONS
Diabetes mellitus; diabetic emergencies and at surgery; diabetic ketoacidosis or coma.

AVAILABILITY
INJECTION (multi-dose vials/prefilled syringes/cartridges) - 40 and 100 IU/ml.

DOSE
Subcutaneous, intramuscular, intravenous injection or intravenous infusion.
Adult and Child- Diabetes mellitus: according to individuals requirement.

PRECAUTIONS
See notes above; reduce dose in renal impairment, lactations; interactions (Appendix 6a, 6b, 6c); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Hypoglycaemia in overdose; localized, and rarely, generalized allergic reactions; lipodystrophy at injection site.

STORAGE
Store in multi dose container in a refrigerator (2 to 8°C). It should not be allowed to freeze.
Intermediate Acting Insulin Insulin Zinc
EDL-D 278, 279 Secondary hospitals

INDICATIONS
Diabetes mellitus.

AVAILABILITY
INJECTION 40 and 80 IU/ml.

Dose
Subcutaneous injection
Adult and Child- Diabetes mellitus: according to individuals requirement.

PRECAUTIONS
See notes above; reduce dose in renal impairment; lactation.

ADVERSE EFFECTS
Hypoglycaemia in overdose; localized, and rarely, generalized allergic reactions; lipodystrophy at injection site.

STORAGE
Store in multi dose containers in a refrigerator (2 to 8°C). It should not be allowed to freeze.

Isophane Insulin
Non-EDL Tertiary

INDICATIONS
Diabetes mellitus.

AVAILABILITY
INJECTION 40 and 80 IU/ml.

DOSE
Subcutaneous injection
Adult and Child- Diabetes mellitus: according to individual’s requirement.

PRECAUTIONS
See notes above; reduce dose in renal impairment; lactation.

ADVERSE EFFECTS
Hypoglycaemia in overdose; localized and rarely, generalized allergic reactions; lipodystrophy at injection site.

STORAGE
Store in multi dose containers in a refrigerator (2 to 8°C). It should not be allowed to freeze.

Pioglitazone
Non-EDL Tertiary

INDICATIONS
Type 2 diabetes mellitus.

AVAILABILITY
TABLETS 15 and 30 mg.

DOSE
Oral
Type 2 diabetes mellitus: Adult- 15-30 mg once daily.
Max. dose- 45 mg per day.

CONTRAINDICATIONS
Hypersensitivity, type 1 diabetes, diabetic ketoacidosis, symptomatic or history of heart failure, children, lactation.

PRECAUTIONS
Oedema, congestive heart failure, hepatic dysfunction, anaemia, concomitant oral contraceptives and hormone replacement therapy, pregnancy (Appendix 7c), interactions (Appendix 6c).
ADVERSE EFFECTS
Oedema, headache, upper respiratory tract infection, GI disturbances, nausea, shortness of breath, weight gain, blurred vision, dizziness, arthralgia, impotence.

STORAGE
Store protected from heat, light and moisture at a temperature not exceeding 30°C.

Progestogens
Progestosterone is a hormone secreted by the corpus luteum whose actions include induction of secretory changes in the endometrium, relaxation of uterine smooth muscle and production of changes in the vaginal epithelium. Progesterone is relatively inactive following oral administration and produces local reactions at site of injection. This has led to the development of synthetic progestogens including levonorgestrel, norethisterone and medroxyprogesterone. Where endometriosis requires drug treatment, it may respond to synthetic progestogens on a continuous basis. They may also be used for the treatment of severe dysmenorrhoea. In postmenopausal women receiving long-term estrogen therapy for hormone replacement, a progestogen needs to be added for women with an intact uterus to prevent hyperplasia of the endometrium.

Progestogens are also used in combined oral contraceptives and progestogen-only contraceptives.

Medroxy Progesterone Acetate
EDL-D 326,327 Universal

AVAILABILITY
TABLETS 2.5, 5 and 10 mg; INJECTION 150 mg (1 ml VIAL/PREFILLED SYRINGE).

DOSE
Deep intramuscular injection Adult- Contraception (short-term): 150 mg within first 5 days of cycle or within first 5 days after parturition (delay until 6 weeks after parturition if lactating). Contraception (long-term); as for short-term, repeated every 3 months. Mild to moderate endometriosis: 10 mg 3 times daily for 90 consecutive days, beginning on day 1 of cycle. Dysfunctional uterine bleeding; 2.5 to 10 mg daily for 5 to 10 days beginning on day 16 to 21 of cycle for 2 cycles. Secondary amenorrhoea; 5 to 10 mg daily for 5 to 10 days beginning on day 16 to 21 of cycle for 3 cycles. If interval between injections is greater than 3 months and 14 days, exclude pregnancy before next injection and advise patient to use additional contraceptive measures (for example barrier) for 7 days after the injection.

INDICATION
Parenteral progestogen-only contraception (short- term or long-term); menstrual symptoms and endometriosis; dysmenorrhoea.

CONTRAINDICATION
Pregnancy ; hormonedefendent breast or genital neoplasms; undiagnosed vaginal bleeding; hepatic impairment or active liver disease; severe arterial disease; porphyria; active thrombophlebitis; lactation .

PRECAUTION
Small increase in possible risk of breast cancer; migraine; liver disease; thromboembolic or coronary vascular disease; diabetes mellitus; trophoblastic disease; hypertension; renal disease; fluid retention, CNS disorder and convulsions.

ADVERSE EFFECTS
Menstrual irregularities; delayed return to fertility; reduction in bone mineral density; weight gain; depression; rarely, anaphylaxis; abdominal pain, asthenia, breast pain, bloating, insomnia, vaginitis.
Norethisterone
EDL-D 374 Secondary hospitals

AVAILABILITY
TABLET 5 mg; injection 1 ml ampoule (200 mg/ml).

DOSE
Oral Adult- Endometriosis: 10 mg daily starting on fifth day of cycle (increased if spotting occurs to 20 to 25 mg daily, reduce once bleeding has stopped). Menorrhagia: 5 mg three times daily for 10 days to stop bleeding; to prevent bleeding 5 mg twice daily from day 19 to 26 of cycle. Dysmenorrhoea: 5 mg, 2 to 3 times daily from day 5 to 24 for 3 to 4 cycles.

INDICATION
Endometriosis; menorrhagia; severe dysmenorrhoea; contraception; premenstrual tension.

CONTRAINDICATION
Pregnancy ; breast or endometrial cancer; severe liver disease (Dubin-Johnson or Rotor’s syndromes); history of jaundice, pruritus, herpes or of deteriorating otosclerosis during pregnancy; severe diabetes mellitus with vascular changes; hypertension; 12 weeks before planned surgery and during immobilization; thromboembolic disease; disturbances of lipid metabolism; undiagnosed vaginal bleeding; porphyria; epilepsy, hepatitis, amenorrhoea, herpes gestation.

PRECAUTION
Epilepsy; migraine; diabetes mellitus; hypertension; cardiac or renal disease and those susceptible to thromboembolism; depression; lactation (Appendix 7b).

ADVERSE EFFECTS
Bloating; breast discomfort; headache; dizziness, depression; nausea; menstrual irregularities; rarely; weight gain; hepatitis; cataract; optic neuritis; mental discomfort.

Thyroid hormones and antithyroid medicines

Thyroid Drugs:
Thyroid agents are natural or synthetic agents containing levothyroxine (thyroxine) or liothyronine (tri-iodothyronine). The principal effect is to increase the metabolic rate. They also exert a cardiotimulatory effect which may be the result of a direct action on the heart. Thyroid hormones are used in hypothyroidism (myxoedema) and also in diffuse non-toxic goitre, Hashimoto thyroiditis (lymphadenoid goitre) and thyroid carcinoma. Neonatal hypothyroidism requires prompt treatment for normal development.

Levothyroxine Sodium (thyroxine Sodium) is the treatment of choice for maintenance therapy. It is almost completely absorbed from the gastrointestinal tract but the full effects are not seen for up to 1 to 3 weeks after beginning therapy; there is a slow response to dose change and effects may persist for several weeks after withdrawal. Dosage of levothyroxine in infants and children for congenital hypothyroidism and juvenile myxoedema should be titrated according to clinical response, growth assessment and measurement of plasma thyroxine and thyroid-stimulating hormone.

Antithyroid Drugs:
Antithyroid drugs such as propylthiouracil and carbimazole are used in the management of thyrotoxicosis. They are also used to prepare the patient for thyroidectomy. They are usually well tolerated, with mild leukopenia or rashes developing in a few percent of cases, usually during the first 6-8 weeks of therapy. During this time the blood count should be checked every
2 weeks or if a sore throat or other signs of infection develop. The drugs are generally given in a high dose in the first instance until the patient becomes euthyroid, the dose may then be gradually reduced to a maintenance dose which is continued for 12-18 months, followed by monitoring to identify relapse. There is a lag time of some 2 weeks between the achievement of biochemical euthyroidism and clinical euthyroidism. Betaadrenoceptor antagonists (beta-blockers) (usually propranolol) may be used as a short-term adjunct to antithyroid drugs to control symptoms but their use in heart failure associated with thyrotoxicosis is controversial. Treatment can be given, if necessary, in pregnancy but antithyroid drugs cross the placenta and in high doses may cause fetal goitre and hypothyroidism. The lowest dose that will control the hyperthyroid state should be used (requirements in Graves disease tend to fall during pregnancy). Propylthiouracil appears in breast milk but does not

Levothyroxine Sodium

EDL-D 304,305,306 Secondary hospitals

AVAILABILITY
Tablets 50 and 100 μg

DOSE
Oral Adult- Hypothyroidism: Initially 50 to 100 μg daily (25 to 50 μg for those over 50 years) before breakfast, increased by 25 to 50 μg every 3 to 4 weeks until normal metabolism maintained (usual maintenance dose, 100 to 200 μg daily); where there is cardiac disease, initially 25 μg daily or 50 μg on alternate days, adjusted in steps of 25 μg every 4 weeks. Child- Congenital hypothyroidism and juvenile myxoedema; Up to 1 month: initially 5 to 10 μg/kg daily. Over 1 month: initially 5 μg/kg daily, adjusted in steps of 25 μg every 2 to 4 weeks, until mild toxic symptoms appear, then reduce dose slightly.

INDICATION
Hypothyroidism.

CONTRAINDICATION
Thyrotoxicosis.

PRECAUTION
Cardiovascular disorders (myocardial insufficiency or ECG evidence of myocardial infarction); hypopituitarism or predisposition to adrenal insufficiency (must be corrected by corticosteroid prior to initial levothyroxine); elderly; long-standing hypothyroidism, diabetes insipidus, diabetes mellitus (may need to increase dose of insulin or oral antidiabetic drug); pregnancy (Appendix 7c), lactation; interactions (Appendix 6c, 6d).

ADVERSE EFFECTS
Anginal pain, arrhythmias, palpitations, tachycardia, skeletal muscle cramps; diarrhoea, vomiting; tremors; restlessness excitability, insomnia, headache, flushing, sweating; excessive loss of weight and muscular weakness; heat intolerance

Potassium iodide

EDL-D 420 Secondary hospitals

AVAILABILITY
Tablets, potassium iodide 60 mg.

INDICATION
Thyrotoxicosis (pre-operative treatment); sporotrichosis, subcutaneous phycomycosis (section 6.3).

CONTRAINDICATIONS
Breastfeeding (Appendix 3); long-term treatment.
PRECAUTIONS
pregnancy (Appendix 2), children.

DOSE
Pre-operative management of thyrotoxicosis, by mouth, ADULT 60–180 mg daily

ADVERSE EFFECTS
Hypersensitivity reactions including coryza like symptoms, headache, lacrimation, conjunctivitis, pain in salivary glands, laryngitis, bronchitis, rashes; on prolonged treatment, depression, insomnia, impotence, goitre in infants of mothers taking iodides.

Propyl thiouracil
EDL-D 442,443 Secondary hospitals

INDICATION
Hyperthyroidism.

AVAILABILITY
Tablets 50 mg, 100 mg

DOSE
Hyperthyroidism - 300 to 900 mg/ day in divided doses till patient becomes euthyroid.
Maintenance - 50 to 600 mg/ day in divided doses.

CONTRAINDICATION
Hypersensitivity, pregnancy, lactation.

PRECAUTION
Same as for carbimazole (pg no.250).

ADVERSE EFFECT
Fever, leukopenia, agranulocytosis, peripheral neuropathy, nephritis, renal vasculitis, changes in menstrual period, headache, nausea and vomiting.

DRUG INTERACTION
Decreased response to propyl thiouracil on concomitant use with iodine or potassium iodide, response to oral anticoagulants may be decreased, increased risk of digitalis toxicity.

Note: Though anti thyroid drugs are specific agents to reduce the levels of circulating thyroid hormones, other auxiliary treatment are often required since the oral antithyroid drugs produce their full effect only within 2 - 3 weeks. Tachycardia and cardiac irritability can be controlled by propranolol in a dose of 10 — 40 mg/day orally. Anxiety and excitement can be controlled by anxiolytic drugs like diazepam. In atleast a few cases hyperthyroidism is associated with abnormalities of serum potassium. This has to be monitored and appropriate steps taken.

Carbimazole
EDL-D 89 Secondary hospitals

AVAILABILITY
TABLETS 5 and 10 mg.

DOSE
Oral Initially 15 to 45 mg daily in 4 divided doses depending upon severity. Maintenance dose 25 to 50 mg for 1 year.

INDICATION
Thyrotoxicosis; Grave’s disease.

CONTRAINDICATION
Liver disorders; pregnancy, lactation; neutropenia.

PRECAUTION
Liver disorders; pregnancy (Appendix 7c), lactation; neutropenia.

ADVERSE EFFECTS
Nausea, mild gastro-intestinal disturbances; headache; rashes and pruritus, arthralgia; rarely, myopathy, alopecia, bone marrow suppression (including pancytopenia and agranulocytosis); vasculitis; cholestatic jaundice, hepatic necrosis.

**Iodine**

**EDL-D 280 Secondary hospitals**

**INDICATIONS**

Hypothyroidism; sporotrichosis.

**AVAILABLE**

COLLOIDAL IODINE 8 mg/5 ml.

**DOSE**

5 to 10 ml diluted in water 3 times a day.

**CONTRAINDICATIONS**

Lactation (Appendix 7b), tuberculosis, bronchitis, asthma, hyperkalaemia, acne vulgaris.

**PRECAUTIONS**

Pregnancy (Appendix 7c), children; not for long-term treatment; cardiac disease, interactions (Appendix 6c).

**ADVERSE EFFECTS**

Hypersensitivity reactions including coryzalike symptoms; headache; lacrimation; conjunctivitis, pain in salivary glands; laryngitis, bronchitis, rashes; on prolonged treatment depression, insomnia, impotence; goitre in infants of mothers taking iodides; eosinophilia, hypothyroidism, abdominal pain, arrhythmia.

**STORAGE**

Store in ground glass stoppered container or earthenware container with waxed bungs.
SECTION - 18
IMMUNOLOGICALS

Active Immunity:
Active immunity may be induced by the administration of micro-organisms or their products which act as antigens to induce antibodies to confer a protective immune response in the host. Vaccination may consist of (a) a live attenuated form of a virus or bacteria, (b) inactivated preparations of the virus or bacteria, or (c) extracts of or detoxified exotoxins. Live attenuated vaccines usually confer immunity with a single dose which is of long duration. Inactivated vaccines may require a series of injections in the first instance to produce an adequate antibody response and in most cases, require reinforcing (booster) doses. The duration of immunity varies from months to many years. Extracts of or detoxified exotoxins require a primary series of injections followed by reinforcing doses.

Passive Immunity:
Passive immunity is conferred by injecting preparations made from the plasma of immune individuals with adequate levels of antibody to the disease for which protection is sought. Treatment has to be given soon after exposure to be effective. This immunity lasts only a few weeks but passive immunization can be repeated where necessary.

Sera and Immunoglobulins
Antibodies of human origin are usually termed immunoglobulins. Material prepared from animals is called antiserum. Because of serum sickness and other allergic-type reactions that may follow injections of antisera, this therapy has been replaced wherever possible by the use of immunoglobulins. All immunoglobulins and antisera should comply with WHO requirements for blood and plasma products.

Contraindications and Precautions
Anaphylaxis, although rare, can occur and epinephrine (adrenaline) must always be immediately available during immunization. Immunoglobulins may interfere with the immune response to live virus vaccines which should normally be given either at least 3 weeks before or at least 3 months after the administration of the immunoglobulin.

Adverse Reactions:
Intramuscular injection; Local reactions including pain and tenderness may occur at the injection site. Hypersensitivity reactions may occur including, rarely, anaphylaxis. Intravenous injection; Systemic reactions including fever, chills, facial flushing, headache and nausea may occur, particularly following high rates of infusion. Hypersensitivity reactions may occur including, rarely, anaphylaxis.

Anti-D Immunoglobulin (Human):
Anti-D immunoglobulin is prepared from plasma with a high titre of anti-D antibody. It is available to prevent a rhesusnegative mother from forming antibodies to fetal rhesuspositive cells which may pass into the maternal circulation. The aim is to protect any subsequent child from the hazard of haemolytic disease of the newborn. It should be administered following any potentially sensitizing episode (for example abortion, miscarriage, still-birth) immediately or within 72 h of the episode but even if a longer period has elapsed it may still give protection and should be used. The dose of anti-D immunoglobulin given depends on the level of exposure to rhesus-positive blood. The injection of anti-D immunoglobulin is not effective once the mother has formed anti-D antibodies. It is also given following Rh0 (D) incompatible blood.
Antitetanus Immunoglobulin (Human):
Antitetanus immunoglobulin of human origin is a preparation containing immunoglobulins derived from the plasma of adults immunized with tetanus toxoid. It is used for the management of tetanus-prone wounds in addition to wound toilet and if appropriate antibacterial prophylaxis and adsorbed tetanus vaccine.

Diphtheria Antitoxin:
Diphtheria antitoxin is prepared from the plasma or serum of healthy horses immunized against diphtheria toxin or diphtheria toxoid. It is used for passive immunization in suspected cases of diphtheria without waiting for bacterial confirmation of the infection. A test dose should be given initially to exclude hypersensitivity. Diphtheria antitoxin is not used for prophylaxis of diphtheria because of the risk of hypersensitivity.

Rabies Immunoglobulin (Human):
Rabies immunoglobulin is a preparation containing immunoglobulins derived from the plasma of adults immunized with rabies vaccine. It is used as part of the management of potential rabies following exposure of an unimmunized individual to an animal in or from a high-risk country. It should be administered as soon as possible after exposure without waiting for confirmation that the animal is rapid. The site of the bite should be washed with soapy water and the rabies immunoglobulin should be infiltrated round the site of the bite and also given intramuscularly. In addition, rabies vaccine should be administered at a different site.

Diagnostic agents

Immunologicals
The tuberculin test has limited diagnostic value. A positive tuberculin test indicates previous exposure to mycobacterial antigens through infection with one of the tubercle bacilli, or BCG vaccination. The tuberculin test does not distinguish between tuberculosis and other mycobacterial infection, between active and quiescent disease, or between acquired infection and seroconversion induced by BCG vaccination.

Tuberculin, Purified Protein Derivative (PPD)

EDL-D 513 PHC

AVAILABILITY
Available at special clinics or by specialists

DOSE
Intradermal injection Adult and Child- 5 to 10 units (1 unit may be used in hypersensitive patients or if tuberculosis is suspected)

INDICATION
Used for test of hypersensitivity to tuberculoprotein.

CONTRAINDICATION
Should not be used within 3 weeks of receiving a live viral vaccine.

PRECAUTION
Elderly; malnutrition; viral or bacterial infections (including HIV and severe tuberculosis), malignant disease, corticosteroid or immunosuppressant therapy-diminished sensitivity to tuberculin; avoid contact with open cuts, abraded or diseased skin, eyes or mouth
ADVERSE EFFECTS
Occasionally nausea, headache, malaise, rash; immediate local reactions (more common in atopic patients); rarely, vesicular or ulcerating local reactions, regional adenopathy and fever; necrosis, pruritis, pain.

Sera and immunoglobulins
Antivenom Sera:
The snake bite may cause local and systemic effects. Local effects include pain, swelling, bruising and tender enlargement of regional lymph nodes. Wounds should be cleaned and pain may be relieved by analgesics. If significant amounts of toxin are absorbed after a snake bite, this may result in early anaphylactoid symptoms such as transient hypotension, angioedema, abdominal colic, diarrhea and vomiting, followed by persistent or recurrent hypotension and ECG abnormalities. Spontaneous systemic bleeding, coagulopathy, adult respiratory distress syndrome and acute renal failure may occur. Early anaphylactoid symptoms may be treated with epinephrine (adrenaline). Snake antivenom sera are the only specific treatment available but they can produce severe adverse reactions. They are generally only used if there is a clear indication of systemic involvement or severe local involvement or, if supplies are not limited, in patients at high risk of systemic or severe local involvement.

Spider bites may cause either necrotic or neurotoxic syndromes depending on the species involved. Supportive and symptomatic treatment is required and in the case of necrotic syndrome, surgical repair may be necessary. Spider antivenom sera, suitable for the species involved, may prevent symptoms if administered as soon as possible after envenomation.

Typhoid Vaccine

EDL-D 759 Tertiary Restricted
AVAILABILITY
Injection 30 μg/vial and 150 μg/vial.

DOSE
Oral Adult- Each dose given on days 0, 2 and 4 (total of 3 doses), with reinforcing doses every year for travellers to disease-endemic countries and every 3 years for those living in disease-endemic areas. Child- Over 6 years: each dose given on days 0, 2 and 4 (total of 3 doses), with reinforcing doses every year for travellers to disease-endemic countries and every 3 years for those living in disease-endemic areas. Intramuscular or deep subcutaneous injection Adult- 0.5 ml, with reinforcing doses every 3 years for those at continued risk. Child- 0.5 ml, with reinforcing doses every 3 years for those at continued risk.

INDICATION
Active immunization against typhoid.

CONTRAINICATION
hypersensitivity.

PRECAUTION
See introductory notes and notes above; illness, infection, allergy, radiation therapy, interactions (Appendix 6d); pregnancy (Appendix 7c).

ADVERSE EFFECTS
anaphylactoid reaction; nausea; vomiting; fever; redness; itching; abdominal pain

Antitetanus Immunoglobulin (Human)

EDL-D 46 Secondary hospitals
AVAILABILITY
INJECTIONS 250, 500 and 1000 I.U/vial.
DOSE
Intramuscular injection Adult and Child-250 units, increased to 500 units if wound older than 12 h or there is risk of heavy contamination or if patient weighs more than 90 kg. Second dose of 250 μg given after 3 to 4 weeks if patient is immunosuppressed or if active immunisation with tetanus vaccine contraindicated.

INDICATION
Passive immunisation against tetanus as part of the management of tetanus-prone wounds.

CONTRAINDICATION
Anaphylaxis, although rare, can occur and epinephrine (adrenaline) must always be immediately available during immunization. Immunoglobulins may interfere with the immune response to live virus vaccines which should normally be given either at least 3 weeks before or at least 3 months after the administration of the immunoglobulin.

PRECAUTION
See introductory notes; must not be administered i.v and patient must be observed for 20 min after administration; pregnancy

ADVERSE EFFECTS
Intramuscular injection; Local reactions including pain and tenderness may occur at the injection site. Hypersensitivity reactions may occur including, rarely, anaphylaxis. Intravenous injection; Systemic reactions including fever, chills, facial flushing, headache and nausea may occur, particularly following high rates of infusion. Hypersensitivity reactions may occur including, rarely, anaphylaxis

Anti-D Immunoglobulin Polyclonal (Thiomersal free)
EDL-D 44,45 Secondary hospitals
AVAILABILITY
INJECTIONS 150 μg/vial and 300 μg/vial.

DOSE
Intramuscular injection Adult and Child- Following birth of a rhesuspositive infant in rhesus-negative mother: 250 μg immediately or within 72 h. Following any potentially sensitizing episode like amniocentesis, still birth, up to 20 weeks gestation: 250 μg per episode, after 20 weeks: 500 μg immediately or within 72 h. Following Rho (D) incompatible blood transfusion: 10 to 20 μg/ml transfused rhesus-positive blood.

INDICATION
Prevention of formation of antibodies to rhesus-positive blood cells in rhesus-negative patients.

CONTRAINDICATION
Known Hypersensitivity

PRECAUTION
See introductory notes; caution in rhesuspositive patients for treatment of blood disorders; caution in rhesus-negative patients with anti-D antibodies in their serum; patients should be observed for 20 min after injection.

ADVERSE EFFECTS
caution in rhesuspositive patients for treatment of blood disorders; caution in rhesus-negative patients with anti-D antibodies in their serum; patients should be observed for 20 min after injection.

Polyvalent Snake Antivenom Serum
EDL-D 418 PHC
AVAILABILITY
INJECTION 10 ml ampoule.
DOSE
60-100 ml in 5% dextrose or normal saline intravenously over one hour; start at 1 ml of diluted solution per minute initially, watching for reaction. Skin sensitivity test is not recommended; In hemotoxic snake bites, may repeat a second dose at 6 h. if bleeding/ clotting abnormalities continue, or whole blood clotting time is still prolonged at 6 h; In neurotoxic snake bites, may repeat at 1-2 h.

INDICATION
Treatment of snake bites.

PRECAUTION
Resuscitation facilities should be immediately available; antihistamine and treatment for anaphylactic shock should be kept ready

ADVERSE EFFECTS
Serum sickness; anaphylaxis with hypotension, dyspnoea, urticaria and shock.

Rabies Immunoglobulin Equine
EDL-D 725 Secondary hospitals

AVAILABILITY
INJECTION 150 IU/2 ml; 300 IU/ml; 200-400 IU/5 ml; 1000 IU/ml; 1000 IU/5 ml; 1500 IU/5 ml.

DOSE
Intramuscular injection and wound infiltration Adult and Child- 20 units/kg (half by intramuscular injection and half by wound infiltration).

INDICATION
Passive immunisation either post-exposure or in suspected exposure to rabies in highrisk countries in unimmunised individuals (in conjunction with rabies vaccine).

CONTRAINDICATION
avoid repeat doses after vaccine treatment initiated; intravenous administration.

PRECAUTION
Epinephrine should be available for management of anaphylactic reaction. If schedule requires rabies vaccine and rabies immunoglobulin to be administered at the same time, they should be administered using separate syringes and separate sites; pregnancy(Appendix 7c)

ADVERSE EFFECTS
soreness at injection site; fever; chest pain; tremor; dyspnoea.

BCG Vaccine
EDL-D 65 PHC

AVAILABILITY
Between 1x106 and 33x106 CFU/ml INJECTION (1 million units of BCG Strain/dose).

DOSE
Intradermal injection Adult- 0.1 ml. Child- Over 3 months: 0.1 ml; Infant, up to 3 months: 0.05 ml.

INDICATION
Active immunisation against tuberculosis

CONTRAINDICATION
generalized oedema; antimycobacterial treatment; HIV infection, febrile illness, burn patients; hypersensitivity.

PRECAUTION
Eczema, scabies-vaccine site must be lesionfree; severly immunocompromised patients; pregnancy
ADVERSE EFFECTS
See introductory notes; lymphadenitis and keloid formation; osteitis and localized necrotic ulceration; rarely, disseminated BCG infection in immunodeficient patients; rarely, anaphylaxis; bladder irritation, anorexia, weight loss.

Diphtheria, Pertussis and Tetanus combined (DPT) vaccine
EDL-D 186 PHC
AVAILABILITY
INJECTIONS 0.5 ml/ampoule and 5 ml/vial
DOSE
Intramuscular injection Child- Infant; 0.5 ml at 6, 10 and 14 weeks.

INDICATION
Active immunisation against diphtheria, tetanus and pertussis.

CONTRAINDICATION
Hypersensitivity, do not administer i.v., seizures, hypotension, hyporeactivity syndrome

PRECAUTION
In cases of severe reaction, the pertussis component should be omitted and the primary course of immunization completed with diphtheria and tetanus vaccine; postpone vaccination if fever, acute disease.

ADVERSE EFFECTS
tetanus component rarely, associated with peripheral neuropathy; pertussis component rarely, associated with convulsions and encephalopathy; induction, oedema.

Hepatitis B Vaccine
EDL-D 257 Secondary hospitals
AVAILABILITY
INJECTIONS 2 ml/vial (10 μg/0.5 ml) 10 μg/ml, 10 μg/2 ml, 10 μg/5 ml, 10 μg/10 ml, 100 μg/10 ml, and multidose vial 10 ml/vial (20 μg/ml).

DOSE
Intramuscular injection Adult- Immunisation of unimmunised and high risk persons: 3 doses of 1 ml with an interval of 1 month between the first and second dose and 5 months between the second and third doses. Child- Immunisation of children, Infant: 0.5 ml either (Scheme A) at birth and at 6 and 14 weeks of age or (Scheme B) at 6, 10 and 14 weeks of age. Immunisation of unimmunised and high risk children, over 15 years: 3 doses of 1 ml with an interval of 1 month between the first and second dose and 5 months between the second and third doses. Under 15 years; 0.5 ml.

INDICATION
Active immunisation against hepatitis B.

CONTRAINDICATION
Acute febrile illness

PRECAUTION
Abdominal pain and gastrointestinal disturbances; muscle and joint pain, dizziness and sleep disturbance; occasionally cardiovascular effects; convulsions, neuropathy, meningitis, paralysis, syncope.

ADVERSE EFFECTS
Abdominal pain and gastrointestinal disturbances; muscle and joint pain, dizziness and sleep disturbance; occasionally cardiovascular effects; convulsions, neuropathy, meningitis, paralysis, syncope.
Measles Vaccine  
EDL-D 323 PHC
AVAILABILITY  
INJECTION 0.5 ml ampoule containing at least 1000 CCID - 50.  
DOSE  
Intramuscular or deep subcutaneous injection  
Child: For immunisation of children against measles; Infant, at 9 months: 0.5 ml. Prophylaxis in susceptible children after exposure to measles; over 9 months: 0.5 ml within 72 h of contact.  
INDICATION  
Active immunization against measles.  
CONTRAINDICATION  
hypersensitivity to any antibiotic present in vaccine (consult literature); hypersensitivity to egg or gelatin; respiratory tract infection, tuberculosis, AIDS.  
PRECAUTION  
See introductory notes; febrile seizures, cerebral injury, pregnancy (Appendix 7c)  
ADVERSE EFFECTS  
rashes sometimes accompanied by convulsions; rarely, encephalitis and thrombocytopenia; headache, pruritus, purpuraea.

Poliomyelitis Vaccine  
EDL-D 417 PHC
AVAILABILITY  
INJECTION 0.5 ml/vial.  
DOSE  
Oral Primary immunisation of unimmunised adult: 3 doses each of 3 drops with an interval of at least 4 weeks between each dose. Reinforcing immunisation of unimmunised adult: 3 doses after 10 years of completion of primary course. Intramuscular injection 2 booster doses of injection first before school entry and second at leaving school. Further booster doses may be required to adults at special risk of polio endemic areas. Child- Primary immunisation: 3 drops at birth and at 6, 10 and 14 weeks of age. Reinforcing immunisation of children: 3 drops at least 3 years after completion of primary course and a further 3 drops at 15 to 19 years of age.  
INDICATION  
Active immunisation against poliomyelitis.  
CONTRAINDICATION  
primary immunodeficiency or immunosuppression; not to be taken with food which contains a preservative; hypersensitivity to any antibiotic present in vaccine  
ADVERSE EFFECTS  
rarely, vaccine-associated poliomyelitis in recipients of vaccine and contacts of recipients; paralytic poliomyelitis.

Tetanus Vaccine (Tetanus Toxoid)  
EDL-D 499 PHC
AVAILABILITY  
INJECTION 0.5 ml ampoule  
DOSE  
Intramuscular injection  
Adult- Primary immunisation: 3 doses each of 0.5 ml with an interval of 4 weeks between each dose. Reinforcing immunisation: 2 doses each of 0.5 ml, the first dose; 10 years after completion of primary course and the second dose; after 10 years. Immunisation of women of child-bearing age against tetanus: 3 primary doses each of 0.5 ml with an interval of not less than 4 weeks between the first and second doses and 6 months between the second and third dose. 2 reinforcing doses each of 0.5 ml, the first dose; 1 year after completion of the primary course and the second dose; 1 year later. Unimmunised pregnant women: 2 doses each of 0.5 ml with an interval of 4 weeks between each dose (second dose at least 2 weeks before
delivery) and 1 dose during each of subsequent 3 pregnancies (max. 5 doses). Intramuscular or deep subcutaneous injection Adult - Management of tetanus-prone wounds and clean wounds: 0.5 ml, the dose schedule being dependent upon the immune status of the patient and the level of contamination of the wound.

**INDICATION**
Active immunisation against tetanus and neonatal tetanus; wound management (tetanus-prone wounds and clean wounds).

**CONTRAINDICATION**
anaphylactic reaction, hypersensitivity

**PRECAUTION**
See introductory notes and notes above; mild cold. If schedule requires tetanus vaccine and antitetanus immunoglobulin to be administered at the same time, they should be administered using separate syringes and separate sites, pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
tetanus component rarely, associated with peripheral neuropathy; seizures, neurological disturbance, fever, loss of appetite.

**Pneumococcal vaccine (Polyvalent)**

**EDL-D 716 Tertiary Restricted**

**INDICATION**
Active immunization of those at risk from streptococcal infection

**DOSE**
Adult: SC/IM, single dose of 0.5 mL

**COINCDICATION**
Hypersensitivity, pregnancy, breast feeding

**PRECAUTION**
Multiple myeloma, Hodgkins and NonHodgkins Lymphomas especially during treatment and in chronic alcohols; chemotherapy or radiation; should be given at least 10 days before starting immunosuppressive therapy or be delayed until at least 6 months after completion of therapy

**ADVERSE EFFECT**
Hypersensitivity reactions, local reaction at injection site

Child: Not recommended in children < 2 years
The muscle relaxants are mainly of two types—peripherally acting and centrally acting and are used along with general anaesthetics for carrying out surgical procedures and to control painful muscle spasms and spastic neurological conditions. They should never be given until it is certain that general anaesthesia has been established and ventilation must be mechanically assisted until they have been completely inactivated. The most common adverse effects of muscle relaxants are vision changes, such as double vision or blurred vision, dizziness, light headness, drowsiness, dry mouth etc.

**Pancuronium**

**EDL-D 394 Tertiary**

**AVAILABILITY**

INJECTION 2 ml ampoule (2 mg/ml).

**DOSE**

Intravenous injection Adult- Initially 50 to 100 μg/kg body weight, 10-20 μg/kg body weight for maintenance dose. Neonates- 30-40 μg/kg body weight.

**INDICATION**

Adjunct to general anaesthesia to facilitate tracheal intubation and to provide skeletal muscle relaxation during surgery or mechanical ventilation.

**CONTRAINDICATION**

Anuria, reduced airway control, lactation.

**PRECAUTION**

Refer Atracurium; hypermagnesemia, hypercalcemia, hyperkalemia, hypoproteinemia; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**

Skin flushing, hypotension, tachycardia, bronchospasm, anaphylactoid reactions, acute myopathy have also been reported after prolonged use in intensive care.

**Atracurium**

**EDL-D 54 Tertiary**

**AVAILABILITY**

Injection 2.5 and 5 ml ampoule (10 mg/ml).

**DOSE**

Intravenous injection Adult and child over 1month- Surgery or intubation: 300 to 600 μg/kg body weight, maintenance by 100 to 200 μg as required. Intensive care: 300 to 600 μg/kg body weight. Intravenous infusion Adult and child- 5 to 10 μg/kg body weight/ min (300 to 600 μg/kg/h).

**INDICATION**

Used adjunctively in anaesthesia to facilitate endotracheal intubation and to provide skeletal muscle relaxation during surgery or mechanical ventilation.

**CONTRAINDICATION**

Hypersensitivity.

**PRECAUTION**

Allergic cross-reactivity between neuromuscular blocking agents has been reported; caution is advised in cases of hypersensitivity. Their activity is prolonged in patients with myasthenia gravis and in hypothermia, therefore lower doses are required. Nondepolarising muscle relaxants
should be used with great care in those with other neuromuscular disorders and those with fluid and electrolyte disturbances, as response is unpredictable. Resistance may develop in patients with burns who may require increased doses; low plasma cholinesterase activity in these patients requires dose titration for mivacurium, renal or hepatic dysfunction, severe electrolyte disturbance, smoking, pregnancy (Appendix 7c), lactation, peripheral neuropathy, demyelinating lesions, denervations. Neonates, severe CVS disorder, severe electrolyte imbalance, respiratory insufficiency or pulmonary disease.

ADVERSE EFFECTS
Skin flushing; hypotension, tachycardia; bronchospasm and very rarely; anaphylactoid reactions, acute myopathy have also been reported after prolonged use in intensive care; prolonged musculoskeletal block, wheezing or bronchial secretion, erythema, dyspnoea.

Baclofen
Non-EDL Tertiary

INDICATIONS
Severe chronic spasticity.

AVAILABILITY
TABLETS 10 and 25 mg;
INJECTIONS 20 ml ampoule (50 μg/ml).

DOSE
Oral
Adult- 5-10 mg thrice daily to 25 mg thrice daily Max. 100 mg daily.
Child- 0.75-2.0 mg/kg daily.

Intrathecal
Adult
Screening dose: 50 μg administered into intrathecal space by barbotage over not less than 1 minute, observe for 4-8 hours. Increase dose by 25 μg not more often every 24 hours until appropriate response is obtained or a maximum dose of 100 μg is achieved. Non responders to 100 μg dose are not suitable for intrathecal baclofen therapy. For responders with response lasting >8-12 hours the screening dose can be given as 24 hour infusion, if response lasted ≤ 8-12 hours then a dose equivalent to twice the screening dose is given as 24 hour infusion.
Child- Similar to that of adult except that the screening dose to be started initially is 25 μg.

CONTRAINDICATIONS
Hypersensitivity; active peptic ulcer disease.

PRECAUTIONS
Cerebrovascular disorder; epilepsy; severe psychotic disorder; respiratory depression; hepatic or renal impairment; elderly; pregnancy (Appendix 7c), avoid sudden withdrawal; interactions (Appendix 6c, 6d).

ADVERSE EFFECTS
Drowsiness; mental confusion; weakness; ataxia; rise in serum transaminases, sudden withdrawal after chronic use may cause hallucinations; tachycardia and seizures, respiratory or cardiovascular depression.

Pyridostigmine
Non-EDL Tertiary

INDICATIONS
Myasthenia gravis.

AVAILABILITY
TABLETS 30, 60 and 180 mg.
DOSE
Oral
Adult- Myasthenia gravis: initially 30 to 120 mg at suitable intervals throughout the day, gradually increased until desired response obtained, total daily dose within range 0.3 to 1.2g, taken at appropriate intervals when max. strength required.
Note: Doses above 450 mg daily not usually advisable in order to avoid acetylcholine receptor downregulation.
Child- Up to 6 years: initially 30 mg. 6 to 12 years initially 60 mg; dose may be gradually increased by 15-30 mg on daily basis.

CONTRAINDICATIONS
Recent intestinal or bladder surgery; gastrointestinal or urinary tract obstruction; after suxamethonium; pneumonia; peritonitis.

PRECAUTIONS
Asthma; urinary tract infection; cardiovascular disease including arrhythmias (especially bradycardia or atrioventricular block); hyperthyroidism; hypotension; peptic ulcer; epilepsy; parkinsonism; renal impairment; pregnancy (Appendix 7c); lactation.

ADVERSE EFFECTS
Muscarinic effects generally weaker than with neostigmine: increased salivation, nausea, vomiting, abdominal cramps, diarrhoea; signs of overdosage include bronchoconstriction, increased bronchial secretions, lacrimation, excessive sweating, involuntary defecation and micturition, miosis, nystagmus, bradycardia, heart block, arrhythmias, hypotension, agitation, excessive dreaming, weakness eventually leading to fasciculation and paralysis; thrombophlebitis; rash associated with bromide salt; diaphoresis, increased peristalsis.

STORAGE
Store protected from light and moisture.

Succinyl Choline Chloride
EDL-D495 Secondary hospitals

INDICATIONS
Short-term muscle relaxation needed for surgical or diagnostic procedures; adjunct to general anesthesia, facilitate tracheal intubation.

AVAILABILITY
INJECTION 50, 100 and 500 mg Vial (50 mg/ ml).

DOSE
Intravenous injection
Initially 1 mg/kg body weight, maintenance by 0.5 to 1 mg/kg body weight every 5 to 10 min interval (max 500 mg).
Intravenous infusion
1 to 2 mg/ml (0.1 to 0.2%), reduce infusion rate in child.

Contraindications
Low serum levels of serum cholinesterase; myasthenia gravis; hypokalemia; glaucoma.

PRECAUTIONS
Severe burns, pregnancy (Appendix 7c), should not be administered until anaesthetic is fully effective, hepatic and renal failure, reduced plasma cholinesterase activity.

ADVERSE EFFECTS
Cardiac arrest, malignant hyperthermia, arrhythmia, increased intraocular pressure; jaw rigidity; muscle pain.

STORAGE
Store protected from light. Injection should not be allowed to freeze.
**Neostigmine**

**EDL-D 361,362 Secondary hospitals**

**Indications**
Myasthenia gravis; reversal of non-depolarizing neuromuscular block, postoperative urinary retention.

**Availability**
Tablets 15 mg; Injection 1 and 5 ml ampoule (0.5 mg/ml).

**Dose**
Subcutaneous or intramuscular injection
- **Adult** - Myasthenia gravis: as neostigmine metilsulfate: 0.5 to 2.5 mg as required, total daily dose 5 to 20 mg.
- **Child** - 200-500 μg as required. Neonate: 50 to 250 μg 30 min before feeds (not usually required beyond 8 weeks of age).

**Contraindications**
- Recent intestinal or bladder surgery; mechanical intestinal or urinary tract obstruction; after suxamethonium; pneumonia; peritonitis.

**Precautions**
- Asthma; urinary tract infections; cardiovascular disease including arrhythmias (especially bradycardia, vagotonia, recent myocardial infarction or atrioventricular block); hyperthyroidism; hypotension; peptic ulcer; epilepsy; parkinsonism; renal impairment; lactation (Appendix 7b); pregnancy (Appendix 7c); interactions (Appendix 6c).

**Adverse Effects**
- Increased salivation, nausea and vomiting, abdominal cramps, diarrhoea; signs of overdosage include bronchoconstriction, increased bronchial secretions, lacrimation, excessive sweating, involuntary defecation and micturition, miosis, nystagmus, bradycardia, heart block, arrhythmias, hypotension, agitation, excessive dreaming, weakness eventually leading to fasciculation and paralysis; thrombophlebitis reported; rash associated with bromide salt, anaphylaxis; dizziness; rash; frequent urination.

**Storage**
Tablets: Store protected from light and moisture. Injection: Store protected from light.

**Vecuronium**

**EDL-D 519 Tertiary**

**INDICATION**
Non-depolarising, intermediate duration, large doses may have cumulative effect, no histamine release, sympathetic blockade or vagolytic effect and it is ideal for cardiac surgery.

**AVAILABILITY**
Powder for reconstitution - 4 mg/mL - 1 mL. amp.

**DOSE**
By IV injection, initially 80-100 mcg / kg (maximum 250 mcg/kg), then 30- 50 mcg/kg as required; By IV infusion, 50- 80 mcg/kg/h. For children: as adult dose (onset more rapid).

**CONTRAINDICATION**
Same as atracurium.

**ADVERSE EFFECT**
Same as atracurium.

**DRUG INTERACTION**
Same as atracurium.

**PRECAUTION**
- Pregnancy, reduce dose in renal impairment and hepatic impairment.
Anti-infective agents

Blepharitis, conjunctivitis and keratitis are common acute infections of the eye and can be treated topically. However, in some cases, for example, in gonococcal conjunctivitis, both topical and systemic anti-infective treatment may be necessary. Blepharitis and conjunctivitis are often caused by staphylococcus, while keratitis and endophthalmitis may be bacterial, viral or fungal. Bacterial blepharitis is treated with an antibacterial eye ointment or drops. Although most cases of acute bacterial conjunctivitis may resolve spontaneously, anti-infective treatment shortens the infectious process and prevents complications. Acute infective conjunctivitis is treated with antibacterial eye drops by day and eye ointment applied at night. A poor response may indicate viral or allergic conjunctivitis. Keratitis requires immediate specialist treatment.

Gentamicin is a broad-spectrum bactericidal aminoglycoside antibiotic with particular activity against *Pseudomonas aeruginosa*, *Neisseria gonorrhoea* and other bacteria that may be implicated in blepharitis or conjunctivitis. Topical application may lead to systemic absorption and possible adverse effects.

Silver nitrate is a topical anti-infective. Its antibacterial activity is attributed to precipitation of bacterial proteins by silver ions. It is available in 1% ophthalmic solutions and is used for prophylaxis of gonococcal ophthalmia neonatorum.

Tetracycline is a broad spectrum antibiotic with activity against many Gram-positive and Gram-negative bacteria including *N. gonorrhoea*, and most chlamydia, rickettsia, mycoplasma and spirochetes. Ophthalmic tetracycline is used in blepharitis, conjunctivitis, and keratitis produced by susceptible bacteria. Tetracycline is also used in the treatment of trachoma caused by *Chlamydia trachomatis* and in the prophylaxis of neonatal conjunctivitis (ophthalmia neonatorum) caused by *N. gonorrhoea and C. trachomatis*.

**Gentamicin**

**EDL-D 244 PHC**

**INDICATIONS**

Blepharitis; bacterial conjunctivitis; keratitis, corneal ulcers.

**AVAILABILITY**

Ointment (1% w/w); Drops 5 ml (0.3% w/v).

**DOSE**

Instillation into the eye

Adult- Mild to moderate infections: 1 drop every 2 h, reducing frequency as infection is controlled, then continue for 48 h after healing is complete.

**CONTRAINDICATIONS**

Hypersensitivity to aminoglycoside group of antibiotics.

**PRECAUTIONS**

Prolonged use may lead to skin sensitization and emergence of resistant organisms including fungi; discontinue if purulent discharge, inflammation or exacerbation of pain; ophthalmic ointment may retard corneal healing, renal impairment (Appendix 7d), interactions (Appendix 6c), pregnancy (Appendix 7c).
ADVERSE EFFECTS
Burning; stinging; itching; dermatitis; conjunctival epithelial defects; conjunctival hyperemia; thrombocytopenic purpura; hallucination.

Ciprofloxacin
EDL-D 125,126 Secondary hospitals

INDICATIONS
Bacterial infections of eye.

AVAILABILITY
tablets 250, 500 and 750 mg; injection 100 ml infusion (20 mg/10 ml); Ointment 5g (0.3% w/w); Drops 5 and 10 ml (0.3% w/v).

DOSE
Adult and child above 12 years- Instill 2 to 3 drops in affected eye 3 to 4 times daily to start with thereafter reduce slowly as infection subsides. Apply about 0.5 cm ribbon of ointment in lower conjuctival sac for 3 to 4 times daily. Reduce as infection subsides.

CONTRAINDICATIONS
Epilepsy and hypersensitivity to quinolones.

PRECAUTIONS
It should be used with caution in patients with a history of epilepsy or conditions that predispose to seizures; in G-6-PD deficiency; myasthenia gravis (risk of exacerbation); in renal impairment (Appendix 7d); pregnancy (Appendix 7c), during lactation (Appendix 7b), and in children or adolescents. Exposure to excessive sunlight should be avoided (discontinue if photosensitivity occurs). Avoid excessive alkalinity of urine and ensure adequate fluid intake (risk of crystalluria); interactions (Appendix 6c); paediatric use.

ADVERSE EFFECTS
Nausea, vomiting, dyspepsia, abdominal pain, diarrhoea (rarely, antibiotic-associated colitis); headache; dizziness; sleep disorders; rash (rarely, Stevens-Johnson syndrome and toxic epidermal necrolysis) and pruritus.
Less frequent side-effects include anorexia, increase in blood urea and creatinine; drowsiness, restlessness, asthenia, depression, confusion, hallucinations, convulsions, tremor, paraesthesia, hypoaesthesia; photosensitivity, hypersensitivity reactions including fever, urticaria, angioedema, arthralgia, myalgia, and anaphylaxis; blood disorders (including eosinophilia, leucopenia, thrombocytopenia); disturbances in vision, taste, hearing and smell. Other side-effects that have been reported include haemolytic anaemia, renal failure, interstitial nephritis, and hepatic dysfunction (including hepatitis and cholestatic jaundice). The drug should be discontinued if psychiatric, neurological or hypersensitivity reactions (including severe rash) occur. Ophthalmic solution: local burning, discomfort, corneal ulcers, lid oedema, corneal infiltration. Ointment: discomfort, keratopathy, blurred vision, corneal staining, epitheliopathy, photophobia.

STORAGE
Ointment: Store protected from light at a temperature not exceeding 30°C. Drops: Store protected from light.

Povidone Iodine
EDL-D 424 Secondary hospitals

INDICATIONS
Antiseptic; skin disinfection; Mouth wash.

AVAILABILITY
SOLUTIONS 100 and 500 ml (5% w/v), 500 ml (7.5% w/v and 10% w/v); OINTMENT 15g (5% w/w).

DOSE
Adult and Child- Pre- and post-operative skin disinfection: apply undiluted. Antiseptic (minor wounds and burns): apply twice daily.
CONTRAINDICATIONS
Avoid regular or prolonged use in patients with thyroid disorders or those taking lithium; avoid regular use in neonates; avoid in very low birthweight infants; burn covering large surface area; hypersensitivity to iodine.

PRECAUTIONS
Pregnancy (Appendix 7c); lactation (Appendix 7b); broken skin (see below); renal impairment; avoid contact with eyes; neonates. The application of povidone iodine to large wounds or severe burns may produce systemic adverse effects such as metabolic acidosis; hypernatraemia; and impairment of renal function.

ADVERSE EFFECTS
Irritation of skin and mucous membranes; may interfere with thyroid function tests; systemic effects (see under Precautions).

STORAGE
Store protected from light.

Acyclovir
EDL-D 9 Secondary hospitals
INDICATIONS
Treatment of Herpes simplex keratitis; long term suppression of skin infections in Herpes simplex as well as mucous membrane, prophylaxis in immunocompromised patients; Herpes zoster treatment.

AVAILABILITY
Ointment 5g (3% w/w); drops 5 ml (3% w/w).

DOSE
Adult- Herpes simplex keratitis: apply 3% w/w ointment 5 times daily for 3 days.

CONTRAINDICATIONS
Hypersensitivity.

PRECAUTIONS
Maintain adequate hydration (especially with infusion or high doses); monitor neutrophil count at least twice weekly in neonates; renal impairment (Appendix 7d); lactation (Appendix 7b); pregnancy (Appendix 7c); not to be applied on mucous membrane.

ADVERSE EFFECTS
Nausea, vomiting, abdominal pain, diarrhoea, headache, fatigue, rash, urticaria, pruritus, photosensitivity; very rarely, hepatitis, jaundice; dyspnoea; neurological reactions (including dizziness, confusion, hallucinations, convulsions and drowsiness); acute renal failure; anaemia, thrombocytopenia and leucopenia; on intravenous infusion; severe local inflammation (sometimes leading to ulceration), and very rarely, agitation, tremors; psychosis and fever; increase in blood urea and creatinine, encephalopathy; seizures; anorexia, tremors.

Miconazole
EDL-D 348 Universal
AVAILABILITY
Drops 5 and 10 ml (1%w/v).

DOSE
Adult and child- Fungal infection of eye: instill 2 to 3 drops 3 to 4 times a day in infected eye or as required.

INDICATION
Fungal infections of eye.

CONTRAINDICATION
Hypersensitivity.
PRECAUTION
Contact with eyes and mucous membranes should be avoided, pregnancy

ADVERSE EFFECTS
Occasional local irritation and hypersensitivity reactions include mild burning sensation, erythema; pruritus and itching. Treatment should be discontinued if these are severe.

Ofloxacin
EDL-D 551 PHC

INDICATIONS
Acute uncomplicated cystitis, community acquired pneumonia, acute exacerbation of chronic bronchitis.

AVAILABILITY
TABLETS 100, 200 and 400 mg; SYRUP 30 ml (50 mg/5 ml, 100 mg/5 ml); INJECTION 100 ml (2 mg/ml);
EYE DROPS 0.3% w/v.

DOSE
Oral
Community acquired pneumonia:
Adult- 400 mg twice daily for 10 days. Pelvic inflammatory disease:
Adult- 400 mg twice daily for 14 days. Complicated UTI:
Adult- 200 mg twice daily for 10 days.
Parenteral
Complicated UTI:
Adult- 200 mg daily by i.v infusion over atleast 30 minutes, max. 400 mg twice infused over at least 1 h.
Septicaemia, lower respiratory tract infection:
Adult- 200 mg twice daily by i.v infusion over at least 30 minutes, max. 400 mg twice daily infused over at least 1 h.
Bacterial corneal ulcer:
Adult- 0.3%, 1-2 drops every 30 minutes.
Ophthalmic Bacterial conjunctivitis:
Adult- 0.3%, 1-2 drops every 2-4 h.
Child- >1year, 1-2drops every 2-4 h.

CONTRAINDICATIONS
Hypersensitivity.

PRECAUTIONS
Patients with epilepsy, kidney disease, tendon problem, nervous system problem, liver disease (Appendix 7a), limit alcohol intake, pregnancy (Appendix 7c); lactation (Appendix 7b).

ADVERSE EFFECTS
Sinus tachycardia, hallucination, Steven’s Johnson syndrome, seizure; dizziness, headache, nausea, vomiting, diarrhoea; insomnia, pruritus, photosensitivity.

STORAGE
Tablets: Store protected from light and moisture. Eye Drops: Store protected from light.

Tropicamide
EDL-D 512 Secondary hospitals

INDICATIONS
Dilatation of the pupil to examine the fundus; cycloplegia.

AVAILABILITY
EYE DROPS 5 and 10 ml vial (0.08 & 1.0 % w/v).
DOSE  Ocular instillation
Adult and Child- Dilatation of pupil to examine the fundus: 1 drop, 15 to 20 min before examination of eye.

PRECAUTIONS
Patients aged over 60 years and hypermetropic (long-sighted)-may precipitate acute angleclosure glaucoma; darkly pigmented iris; more resistant to pupillary dilatation-exercise caution to avoid overdosage; hyperthyroidism; pregnancy (Appendix 7c); hypertension. Avoid operating machinery or driving for 1-2 h after mydriasis.

ADVERSE EFFECTS
Transient stinging and raised intraocular pressure; on prolonged administrationlocal irritation; hyperaemia; oedema and conjunctivitis; eczematic dermatitis; photophobia; parasympathetic stimulation.

STORAGE
Store protected from light and moisture. For eye drops: store in a refrigerator (8 to 15°C). It should not be allowed to freeze.

Moxifloxacin
EDL-D 695 Secondary Hospitals

INDICATION
A/c bacterial sinusitis, CAP, skin infection, intra abdominal infection

AVAILABILITY
Tablet(5) of 400mg Rs 350/; Injection 400mg - Rs 160/

Dose:
400mg OD

CONTRAINICATION
Hypersensitivity, Age< 18, pregnancy, lactation, bradycardia, heart failure, hypokalemia.

ADVERSE EFFECT
Similar to ciprofloxacin & hematological disturbances, peripheral neuropathy.

Fluconazole
EDL-D 636 Secondary hospitals

INDICATIONS
Systemic mycosis including histoplasmosis, non-meningal coccidioidomycosis, paracoccidioidomycosis and blastomycosis treatment and, in AIDS and other immunosuppressed patients, prophylaxis of cryptococcal meningitis; oesophageal and oropharyngeal candidiasis, vaginal candidiasis and systemic candidiasis

AVAILABILITY
TABLETS/CAPSULES 50, 100, 150 and 200 mg; EYE DROPS 5 ml (0.3% w/v).

DOSE
Adult- Mucosal: 50 to 100 mg daily for 14 to 30 days. Vaginal: 150 mg as a single dose. Oral: systemic loading dose of 400 mg on first day and thereafter 200 to 400 mg once daily for at least 28 days. Prophylaxis of fungal infection: 50 to 100 mg once daily.

CONTRAINDICATIONS
Sensitivity to primaquine; infants below 1 year of age; alcohol; coadministration of cisapride, terfenadine.

PRECAUTIONS
Renal impairment (Appendix 7d); lactation (Appendix 7b); monitor liver functiondiscontinue if signs or symptoms of hepatic disease (risk of hepatic necrosis; Appendix 7a); interactions: (Appendix 6b, 6c); pregnancy (Appendix 7c); immunocompromised patients.
ADVERSE EFFECTS
Nausea, vomiting, abdominal pain; flatulence, diarrhoea; headache, taste disturbance, hepatic disorders, dizziness, seizures, alopecia, pruritus; rash (withdraw treatment); angioedema, anaphylaxis, bullous lesions, toxic epidermal necrolysis and erythema multiforme (Stevens-Johnson syndrome) reported (skin reactions more common in AIDS); hyperlipidaemia, leukopenia, thrombocytopenia, hypokalaemia.

STORAGE
Store in an airtight container.

Fluorescein Sodium
EDL-D 229 Secondary hospitals

AVAILABILITY
EYE DROPS 5 and 10 ml (2% w/v).

DOSE
Ocular instillation Adult and Child- Detection of lesions and foreign bodies in eye: instill sufficient solution dropwise to stain damaged area

INDICATION
Detection of lesions and foreign bodies in the eye

CONTRAINDICATION
Hypersensitivity; avoid use with soft contact lenses.

PRECAUTION
History of allergy or bronchial asthma; lactation; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Gastrointestinal distress; hypotension; syncope; cardiac arrest; thrombophlebitis.

Tobramycin and dexamethasone
EDL-D 750 Tertiary

AVAILABILITY
TABLETS 0.5 mg; INJECTION 2 ml vial (4 mg/ ml); CREAM 5 and 15 g (0.1% w/w).

DOSE
Oral Adult- 0.5 to 10 mg daily in divided doses, repeat if necessary. Child- 0.02 to 0.3 mg/kg in three or four divided doses daily.Intravenous injection 4 to 10 mg every 6 h.

INDICATION
Adjunct in the emergency treatment of anaphylaxis; short-term suppression of inflammation in allergic disorders; adrenocortical insufficiency, ocular inflammation, autoimmune disorders, rheumatic disorder, cerebral oedema, unresponsive shock, bacterial meningitis along with antibiotics.

CONTRAINDICATION
Untreated systemic infection (unless condition life-threatening); administration of live virus vaccines; renal failure, diabetes mellitus, psychosis, osteoporosis, pregnancy, CHF, tuberculosis, fungal infections of the eye.

PRECAUTION
Increased susceptibility to and severity of infection; activation or exacerbation of tuberculosis, amoebiasis, strongyloidiasis; risk of severe chickenpox in non-immune patient (varicella-zoster immunoglobulin required if exposed to chickenpox); avoid exposure to measles (normal immunoglobulin possibly required if exposed); diabetes mellitus; peptic ulcer; hypertension; precautions relating to long-term use of corticosteroids; glaucoma, epilepsy; drug should not be abruptly withdrawn; interactions , lactation
ADVERSE EFFECTS
Nausea, dyspepsia, malaise, hiccups; hypersensitivity reactions including anaphylaxis; perineal irritation after intravenous administration; adverse effects associated with long-term corticosteroid treatment; hyperglycaemia, abdominal distension, angioedema, bradycardia, acne, erythema, Cushing’s syndrome, oropharangeal candidiasis, hypothalamic pituitary adrenal axis suppression

Anti-inflammatory agents
Ophthalamic corticosteroids should only be used under supervision of an ophthalmologist as inappropriate use may lead to blindness. Dangers include the development of open-angle glaucoma (chronic simple glaucoma) and cataracts, and the aggravation of a simple herpes simplex epithelial lesion into an extensive corneal ulcer and subsequent permanent corneal scarring, with possible damage to vision and even loss of the eye. Corticosteroids such as prednisolone are useful in the treatment of inflammatory conditions including uveitis and scleritis. They are also used for reducing postoperative ocular inflammation. Before administration of an ophthalamic corticosteroid, the possibility of bacterial, viral or fungal infection should be excluded. Treatment should be the lowest effective dose for the shortest possible time; if long-term therapy (more than 6 weeks) is unavoidable, withdrawal of an ophthalamic corticosteroid should be gradual to avoid relapse.

Prednisolone
EDL-D 430,719 Secondary hospitals

INDICATIONS
Short-term local treatment of inflammation of the eye; malignant disease; inflammatory and allergic reactions.

AVAILABILITY
tablets 5, 10, 20 and 40 mg; Drop S 5 ml (1% w/v).

DOSE
Doses to be instilled into affected eye 3 to 4 times daily. Ointment at night, preferably at bedtime.

CONTRAINDICATIONS
Undiagnosed ‘red eye’ caused by herpetic keratitis; glaucoma; viral diseases of cornea and conjunctiva.

PRECAUTIONS
Cataract, corneal thinning, corneal or conjunctival infection; discontinue treatment if no improvement within 7 days; risk of adrenal suppression after prolonged use in infants; hepatic impairment (Appendix 7a); lactation (Appendix 7b); interactions (Appendix 6c, 6d); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Secondary ocular infection; impaired corneal healing (due to corneal thinning), optic nerve damage, cataract; glaucoma, mydriasis, ptosis, epithelial punctate keratitis, delayed hypersensitivity reactions including burning, stinging.

Dexamethasone
EDL-D 606 Secondary hospitals

INDICATIONS
Adjunct in the emergency treatment of anaphylaxis; short-term suppression of inflammation in allergic disorders; adrenocortical insufficiency, ocular inflammation, autoimmune disorders,
rheumatic disorder, cerebral oedema, unresponsive shock, bacterial meningitis along with
antibiotics.

AVAILABILITY
TABLETS 0.5 mg; INJECTION 2 ml vial (4 mg/ ml); CREAM 5 and 15 g (0.1% w/w).

DOSE
Oral
Adult- 0.5 to 10 mg daily in divided doses, repeat if necessary.
Child- 0.02 to 0.3 mg/kg in three or four divided doses daily.
Intravenous injection
4 to 10 mg every 6 h.

CONTRAINDICATIONS
Untreated systemic infection (unless condition life-threatening); administration of live virus
vaccines; renal failure, diabetes mellitus, psychosis, osteoporosis, pregnancy (Appendix 7c), CHF,
tuberculosis, fungal infections of the eye.

PRECAUTIONS
Increased susceptibility to and severity of infection; activation or exacerbation of tuberculosis,
amoebiasis, strongyloidiasis; risk of severe chickenpox in non-immune patient (varicella-zoster
immunoglobulin required if exposed to chickenpox); avoid exposure to measles (normal
immunoglobulin possibly required if exposed); diabetes mellitus; peptic ulcer; hypertension;
precautions relating to long-term use of corticosteroids; glaucoma, epilepsy; drug should not be
abruptly withdrawn; interactions (Appendix 6c), lactation (Appendix 7b).

ADVERSE EFFECTS
Nausea, dyspepsia, malaise, hiccups; hypersensitivity reactions including anaphylaxis; perineal
irritation after intravenous administration; adverse effects associated with long-term
corticosteroid treatment; hyperglycaemia, abdominal distension, angioedema, bradycardia,
acline, erythema, Cushing’s syndrome, oropharangeal candidiasis, hypothalamic pituitary adrenal
axis suppression.

STORAGE
Store protected from light at a temperature not exceeding 30°C.

Miotics and anti glaucoma medicines
Glaucoma is one of the leading causes of irreversible blindness world wide. Glaucoma is
normally associated with raised intra-ocular pressure and eventual damage to the optic nerve
which may result in blindness. The rise in pressure is almost always due to reduced outflow of
aqueous humour, the inflow remaining constant. The most common condition is chronicopen-
angle glaucoma (chronic simple glaucoma) in which the intra-ocular pressure increases
gradually and the condition is usually asymptomatic until well advanced. In contrast,
angleclosure glaucoma (closed-angle glaucoma) usually occurs as an acute emergency resulting
from a rapid rise in intra-ocular pressure; if treatment is delayed, chronic angle-closure
glaucoma may develop. Ocular hypertension is a condition in which intra-ocular pressure is
raised without signs of optic nerve damage.

Drugs used in the treatment of glaucoma lower the intraocular pressure by a variety of
mechanisms including reduction in secretion of aqueous humour by the ciliary body, or
increasing the outflow of the aqueous humour by opening of the trabecular network. Antiglaucoma
drugs used include topical application of a beta-blocker (beta-adrenoceptor
antagonist), a miotic, or a sympathomimetic such as epinephrine; systemic administration of a
carbonic anhydrase inhibitor may be used as an adjunct.
Timolol is a non-selective beta-blocker that reduces the secretion of aqueous humour. A beta-blocker is usually the drug of choice for initial and maintenance treatment of chronic open-angle glaucoma. If further reduction in intra-ocular pressure is required a miotic, a sympathomimetic or a systemic carbonic anhydrase inhibitor may be used with timolol. In angle-closure glaucoma, timolol should be used with a miotic and not alone. Since systemic absorption can occur, an ophthalmic betablocker should be used with caution in certain individuals.

A miotic such as pilocarpine, through its parasympathomimetic action, contracts the iris sphincter muscle and the ciliary muscle, and opens the trabecular network. It is used in chronic open-angle glaucoma either alone or, if required, with a beta-blocker, epinephrine or a systemic carbonic anhydrase inhibitor. Pilocarpine is used with systemic acetazolamide in an acute attack of angle-closure glaucoma prior to surgery; however, it is not advisable to use pilocarpine after surgery because of a risk of posterior forming. Systemic absorption of topically applied pilocarpine can occur producing muscarinic adverse effects.

The sympathomimetic drug epinephrine (adrenaline) probably acts by reducing the rate of production of aqueous humour and increasing the outflow through the trabecular network. Epinephrine is usually used with a miotic, a beta-blocker or a systemic carbonic anhydrase inhibitor in the treatment of chronic open-angle glaucoma; however, because epinephrine is also a mydriatic, it is contraindicated for angle-closure glaucoma unless an iridectomy has been carried out.

Acetazolamide, by reducing carbonic anhydrase in the eye, reduces the production of aqueous humour and so reduces intra-ocular pressure. It is used systemically as an adjunct in chronic open-angle glaucoma unresponsive to treatment with topically applied antiglaucoma drugs. Prolonged therapy with acetazolamide is not normally recommended, but if treatment is unavoidable blood count and plasma electrolyte concentration should be monitored. Acetazolamide is also used as part of emergency treatment for an acute attack of angle-closure glaucoma; however it should not be used in chronic angle-closure glaucoma as it may mask deterioration of the condition.

Pilocarpine Hydrochloride or Nitrate
EDL-D 412,413 Secondary hospitals

AVAILABILITY
EYE Drops 5 ml (2% w/v, 4%w/v).

DOSE
Instillation into the eye Adult- Chronic open-angle glaucoma before surgery: 1 drop (2% or 4 %) up to 4 times daily. Acute angle closure glaucoma before surgery: 1 drop (2%) every 10 min for 30 to 60 min, then 1 drop every 1 to 3 h until intra-ocular pressure subsides.

INDICATION
Chronic open-angle glaucoma, ocular hypertension; emergency treatment of acute angle- closure glaucoma; to antagonize effects of mydriasis and cycloplegia following surgery or ophthalmoscopic examination; Accommodative esotropia.
CONTRAINDICATION
Acute iritis, acute uveitis, anterior uveitis, some forms of secondary glaucoma; acute inflammation of anterior segment; not advisable after angle-closure surgery (risk of posterior synechiae).

PRECAUTION
Retinal disease, conjunctival or corneal damage; monitor intra-ocular pressure in chronic open-angle glaucoma and in long-term treatment; cardiac disease, hypertension; asthma; peptic ulceration; urinary-tract obstruction; Parkinson’s disease; stop treatment if symptoms of systemic toxicity develop; ulcer; hyperthyroidism; seizures. Causes difficulty with dark adaptation; may cause accommodation spasm. Do not carry out skilled tasks, for example operating machinery or driving until vision is clear, pregnancy (Appendix 7c).

ADVERSE EFFECTS
Eye pain, blurred vision, ciliary spasm, lacrimation, myopia, browache; conjunctival vascular congestion, superficial keratitis, vitreous haemorrhage and increased pupillary block; lens opacities have occurred following prolonged use; rarely, systemic effects including hypertension, tachycardia; bronchial spasm, pulmonary oedema; salivation; sweating; nausea, vomiting, diarrhoea; flushing, rhinitis, chills, middle ear disturbances.

Acetazolamide
EDL-D1 Secondary hospitals

INDICATIONS
As an adjunct in the treatment of chronic open-angle glaucoma; secondary glaucoma; as part of pre-operative treatment of acute angle-closure glaucoma.

AVAILABILITY
Tablet 250 mg; capsule 250 mg.

DOSE
Oral
Adult- 0.25 to 1g daily in divided doses.

CONTRAINDICATIONS
Hypersensitivity to sulfonamides; chronic angle-closure glaucoma (may mask deterioration); hypokalaemia, hyponatraemia, hyperchloraemic acidosis; renal impairment (Appendix 7d), severe hepatic impairment; renal hyperchloremic acidosis, addison’s disease.

PRECAUTIONS
Elderly; lactation; diabetes mellitus; pulmonary obstruction; monitor blood count and electrolytes if used for long periods; interactions (Appendix 6b, 6c); pregnancy (Appendix 7c); severe respiratory acidosis.
May impair ability to perform skilled tasks, for example operating machinery, driving.

ADVERSE EFFECTS
Nausea, vomiting, diarrhoea, taste disturbance; loss of appetite, paraesthesia, flushing, headache, dizziness, fatigue, irritability, depression; thirst, polyuria; reduced libido; metabolic acidosis and electrolyte disturbances on longterm therapy; occasionally drowsiness, confusion, hearing disturbances, urticaria, melaena, glycosuria, haematuria; abnormal liver function; renal calculi, blood disorders including agranulocytosis and thrombocytopenia; rashes including Stevens- Johnson syndrome and toxic epidermal necrolysis; transient myopia reported; blood dyscrasias; crystalluria.

STORAGE
Store protected from light.
Physostigmine  
Non-EDL Tertiary  
INDICATIONS  
Glaucoma in conjunction with other drugs and not alone (as it is very potent).  
AVAILABILITY  
EYE DROPS 5 ml (0.25% w/v).  
Dose: Instillation into the eye  
Adult- 1 to 2 drops, 4 to 5 times daily.  
PRECAUTIONS  
Care to be taken when administered into eye, pregnancy (Appendix 7c).  
ADVERSE EFFECTS  
Twitching lids, myopia, ocular and periorbital pain, ciliary and conjunctival congestion.  

Betaxolol Hydrochloride  
EDL-D 76,77 Secondary hospitals  
AVAILABILITY  
EYE DROPS 5 ml (0.5% w/v).  
DOSE  
Instillation into the eye  
Adult- 1 to 2 drops, twice daily.  
Child- Not recommended.  
INDICATION  
Glaucoma.  
CONTRAINDICATION  
Systemic absorption may follow topical application to the eyes, therefore they are contraindicated in patients with bradycardia, heart block, or uncontrolled heart failure; hypersensitivity  
PRECAUTION  
Avoid in asthma, poor cardiac reserve, hepatic impairment; not for injection; pregnancy (Appendix 7c)  
ADVERSE EFFECTS  
Ocular stinging, burning, pain, itching, erythema, dry eyes and allergic reactions including anaphylaxis and blepharoconjunctivitis; occasionally corneal disorders have been reported; crusty taste, photophobia, corneal punctuate staining, decreased corneal sensitivity, keratitis, anisocoria; headache; sleep disturbances.  

Timolol Maleate  
EDL-D 505,506 Secondary hospitals  
AVAILABILITY  
Drops 5 ml (0.2% w/v, 0.25% and 0.5% w/v); gel (0.5%/5 ml).  
DOSE  
Instillation into the eye  
Adult- 1 drop (0.25% or 0.5%) twice daily.  
INDICATION  
Ocular hypertension; chronic open-angle glaucoma, aphakic glaucoma, some secondary glaucomas.  
CONTRAINDICATION  
Uncontrolled heart failure, bradycardia, heart block; asthma, obstructive airways disease; hypersensitivity.  
PRECAUTION  
Older people (risk of keratitis); if used in angle-closure glaucoma, use with a miotic, and not alone; interactions (Appendix 6c); pregnancy (Appendix 7c).
ADVERSE EFFECTS
Stinging, burning, pain, itching, erythema, transient dryness, allergic blepharitis, transient conjunctivitis, keratitis, decreased corneal sensitivity, diplopia, ptosis; systemic effects; particularly on the pulmonary, cardiovascular and central nervous systems, may follow absorption; blurred vision; headache.

Mydriatics
Antimuscarinics, by blocking the cholinergic effects of acetylcholine, paralyse the pupillary constrictor muscles causing dilation of the pupil (mydriasis) and paralyse the ciliary muscles resulting in paralysis of accommodation (cycloplegia). Mydriasis may precipitate acute angle-closure glaucoma particularly in elderly or far-sighted patients. In patients with dark iridic pigmentation, higher concentrations of mydriatic drugs are usually required and care should be taken to avoid overdosing.

Atropine is a long-acting antimuscarinic used for cycloplegic refraction procedures, particularly in children. It is also used to immobilize the ciliary muscle and iris and to prevent formation of posterior synechiae in the treatment of inflammatory eye disorders such as iritis and uveitis.

Homatropine
EDL-D 258 Universal

AVAILABILITY
Drop S 5 ml (2% w/v).

DOSE
Adult- 1 to 2 drops in each eye till the desired effect is achieved. Child- 1 to 2 drops in each eye till the desired effect is achieved.

INDICATION
To dilate pupil and paralyze ciliary muscle for fundus examination.

CONTRAINDICATION
Narrow angle glaucoma, tendency for glaucoma.

PRECAUTION
Darkly pigmented iris is more resistant to pupillary dilatation and caution should be exercised to avoid overdosage. Mydriasis can precipitate acute angle-closure glaucoma in a few patients, usually aged over 60 years and hypermetropic (long-sighted), who are predisposed to the condition because of a shallow anterior chamber; glaucoma, check intraocular pressure before use; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Ocular side-effects of mydriatics and cycloplegics include transient stinging and raised intraocular pressure; on prolonged administration, local irritation, hyperaemia, oedema and conjunctivitis can occur. Contact dermatitis can occur with the antimuscarinic mydriatic drugs, especially atropine. Systemic side-effects of atropine and cyclopentolate can occur in the young and the old; posterior synechia, headache, drowsiness, loss of taste, photophobia, brow ache, lacrimation.

Phenylephrine
EDL-D 407 Universal

INDICATIONS
Used in cough syrups, hypotension; mydriatic for eye conditions; uveitis, wide angle glaucoma, refraction, ophthalmoscopic examinations.

AVAILABILITY
Drop S 5 ml (5% w/v).
Dose 1 to 2 drops in affected eye, every 4 to 6 h.
CONTRAINDICATIONS
Hypertension (monitor blood pressure and rate of flow frequently); pregnancy (Appendix 7c); narrow angle glaucoma.

PRECAUTIONS
Coronary, mesenteric, or peripheral vascular thrombosis; following myocardial infarction, Prinzmetal’s variant angina; hyperthyroidism, diabetes mellitus; hypoxia or hypercapnia; uncorrected hypovolaemia; elderly; extravasation at injection site may cause necrosis; infants.

ADVERSE EFFECTS
Headache, hypertension, bradycardia, arrhythmias, peripheral ischaemia.

STORAGE: Store protected from light.

Atropine Sulphate
EDL-D 542 PHC

AVAILABILITY
Drops 5 ml (1% w/v).

DOSE
Instillation into the eye
Adult - Cycloplegic refraction: 1 drop (1%) twice daily for 1 to 2 days before procedure or a single application of 1 drop (1%), 1 h before procedure. Iritis and uveitis: 1 drop (0.5 to 1%) up to 4 times daily. Child - Cycloplegic refraction: 3 months to 1 year: 0.1%; 1 to 5 years: 0.1 to 0.5%; Over 5 years: 0.5 to 1.0%. 1 drop twice daily for 1 to 3 days before procedure with a further dose given 1 h before procedure. Iritis and uveitis: 1 drop (0.5 to 1% w/v) up to 3 times daily.

INDICATION
Iritis, uveitis; cycloplegic refraction procedures; iridocyclitis.

CONTRAINDICATION
Angle-closure glaucoma; pregnancy

PRECAUTION
May precipitate acute attack of angleclosure glaucoma, particularly in the elderly or far-sighted; risk of systemic effects with eye drops in infants under 3 months-eye ointment preferred. May cause sensitivity to light and blurred vision. Do not carry out skilled tasks, for example operating machinery or driving, until vision is clear, lactation,

ADVERSE EFFECTS
Transient stinging and raised intra-ocular pressure; on prolonged administration, local irritation, hyperaemia, oedema and conjunctivitis may occur; contact dermatitis; systemic toxicity may occur in the very young and the elderly; blurred vision, dry mouth, photophobia.

Cyclopentolate
EDL-D 594 Secondary hospitals

INDICATION
Refraction testing ciliary spasm, postoperative state, iridocyclitis.

CONTRAINDICATION
Narrow angle glaucoma.

ADVERSE EFFECT
Visual hallucination, incoherence of speech.

AVAILABILITY: Ointment 1 %, 0.5% eye drops.
Drugs may be used to modify uterine contractions. These include oxytocic drugs used to stimulate uterine contractions both in induction of labour and to control postpartum haemorrhage and β₂-adrenoceptor agonists used to relax the uterus and prevent premature labour.

**Postpartum Haemorrhage:**
Ergometrine and oxytocin differ in their actions on the uterus. In moderate doses oxytocin produces slow generalized contractions with full relaxation in between; ergometrine produces faster contractions superimposed on a tonic contraction. High doses of both substances produce sustained tonic contractions.

Oxytocin is now recommended for routine use in postpartum and post-abortion haemorrhage since it is more stable than ergometrine. However, ergometrine may be used if oxytocin is not available or in emergency situations.

**Premature Labour:**
Salbutamol is a β₂-adrenoceptor agonist which relaxes the uterus and can be used to prevent premature labour in uncomplicated cases between 24 and 33 weeks of gestation. Its main purpose is to permit a delay in delivery of at least 48 h. The greatest benefit is obtained by using this delay to administer corticosteroid therapy or to implement other measures known to improve perinatal health. Prolonged therapy should be avoided since the risk to the mother increases after 48 h and the response of the myometrium is reduced.

**Ergometrine Hydrogen Maleate**

**EDL-D 200,201 PHC**

**AVAILABILITY**
- TABLETS 0.125, 0.25 and 0.5 mg; INJECTION 5 ml ampoule (0.2 mg/ml).

**DOSE**
- Oral Adult and adolescent- Secondary postpartum haemorrhage: 400 μg 3 times daily for 3 days. Intramuscular injection 499 NFI-2011 Hormones, Contraceptives and Related Drugs Adult and adolescent- Prevention and treatment of postpartum haemorrhage: when oxytocin is not available, 200 μg when the anterior shoulder is delivered or immediately after birth. Slow intravenous injection Adult and adolescent- Excessive uterine bleeding: 250 to 500 μg when the anterior shoulder is delivered or immediately after birth.

**INDICATION**
- Prevention and treatment of postpartum and post-abortion haemorrhage in emergency situations and where oxytocin not available

**CONTRAINDICATION**
- Induction of labour, first and second stages of labour; vascular disease, severe cardiac disease especially angina pectoris; severe hypertension; hepatic impairment (Appendix 7a) and renal impairment; sepsis; eclampsia.

**PRECAUTION**
- Cardiac disease, hypertension; multiple pregnancy (Appendix 7c); porphyria.

**ADVERSE EFFECTS**
- Nausea, vomiting; headache; dizziness; tinnitus, abdominal pain; chest pain; palpitations; dyspnoea; bradycardia, transient hypertension, vasoconstriction; stroke, myocardial infarction and pulmonary oedema also reported.
Oxytocin
EDL-D 393 PHC

AVAILABILITY
INJECTION 2 IU/2 ml and 5 IU/ml.

DOSE
Intravenous infusion Adult and adolescent- Induction of labour: initially 0.001 to 0.002 units/min increased in 0.001 to 0.002 units/min increments at intervals of 30 min until a max. of 3 to 4 contractions occur every 10 min; max. recommended rate 0.02 units/min. Slow intravenous injection Adult and adolescent- Prevention of postpartum haemorrhage: 5 units when the anterior shoulder is delivered or immediately after birth. Treatment of postpartum haemorrhage: 5-10 units.Intramuscular injection Adult and adolescent- Prevention of postpartum haemorrhage: 10 units when the anterior shoulder is delivered or immediately after birth. 10 units, followed in severe cases by slow intravenous infusion, a total of 40 units should be infused at a rate of 0.02-0.04 units/min; this should be

INDICATION
Routine prevention and treatment of postpartum and post-abortion haemorrhage; induction of labour.

CONTRAINDICATION
Hypertonic uterine contractions, mechanical obstruction to delivery, fetal distress; any condition where spontaneous labour or vaginal delivery inadvisable; avoid prolonged administration in oxytocin-resistant uterine inertia, in severe pre-eclamptic toxaemia or in severe cardiovascular disease; uterine hyperactivity; major cephalopelvic disproportion, placental previa.

PRECAUTION
Induction or enhancement of labour in presence of borderline cephalopelvic disproportion (avoid if significant); mild to moderate pregnancy (Appendix 7c)-associated hypertension or cardiac disease; age over 5 years; history of low-uterine segment caesarean section; avoid tumultuous labour if fetal death or meconium-stained amniotic fluid (risk of amniotic fluid embolism); water intoxication and hyponatraemia (avoid large volume infusions and restrict fluid intake); caudal block anaesthesia (risk of severe hypertension due to enhanced vasopressor effect of sympathomimetics); interactions

ADVERSE EFFECTS
Uterine spasm, uterine hyperstimulation (usually with excessive doses—may cause fetal distress, asphyxia and death, or may lead to hypertonicity, tetanic contractions,softtissue damage or uterine rupture); water intoxication and hyponatraemia associated with high doses and large-volume infusions; nausea, vomiting, arrhythmias, rashes and anaphylactoid reactions also reported; hypotension; sinus bradycardia; hematoma; fetal asphyxia.

Misoprostol
EDL-D 353,354 Universal

AVAILABILITY
Tablet KIT mifepristone 200 mg, misoprostol 200 μg.

DOSE
Mifepristone 200 mg orally followed 1 to 3 days latter by misoprostol 800 μg vaginally. Patients should return for followup visit after approximately 14 days after administration of mifepristone.

INDICATION
Medical termination of pregnancy of upto 49 days, cervical dilatation prior to surgical termination of pregnancy in the first trimester, therapeutic termination of pregnancy for medical reasons beyond the first trimester, labor induction in case of fetal death in utero.
CONTRAINDICATION
Hypersensitivity to Mifepristone, Misoprostol or other prostaglandin; confirmed or suspected ectopic pregnancy; chronic adrenal failure; haemorrhagic disorders or concurrent anticoagulant therapy; inherited porphyria.

PRECAUTION
Hypersensitivity to Mifepristone, Misoprostol or other prostaglandin; confirmed or suspected ectopic pregnancy (Appendix 7c); chronic adrenal failure; haemorrhagic disorders or concurrent anticoagulant therapy; inherited porphyria

ADVERSE EFFECTS
Abdominal pain, diarrhoea, nausea, vomiting; fever, chills, uterine cramping; vaginal bleeding or spotting; Pelvic inflammatory disease.

Mifepristone
EDL-D 352 Secondary hospitals

AVAILABILITY
Tablet KIT mifopristone 200 mg, misoprostol 200 μg.

DOSE
Mifepristone 200 mg orally followed 1 to 3 days latter by misoprostol 800 μg vaginally. Patients should return for followup visit after approximately 14 days after administration of mifepristone.

INDICATION
Medical termination of pregnancy of upto 49 days, cervical dilatation prior to surgical termination of pregnancy in the first trimester, therapeutic termination of pregnancy for medical reasons beyond the first trimester, labor induction in case of fetal death in utero.

CONTRAINDICATION
Hypersensitivity to Mifepristone, Misoprostol or other prostaglandin; confirmed or suspected ectopic pregnancy; chronic adrenal failure; haemorrhagic disorders or concurrent anticoagulant therapy; inherited porphyria.

PRECAUTION
IUD in place; asthma, chronic obstructive pulmonary disease; alcoholism; prosthetic heart valve; infective endocarditis; interactions (Appendix 6c), pregnancy (Appendix 7c)

ADVERSE EFFECTS
Abdominal pain, diarrhoea, nausea, vomiting; fever, chills, uterine cramping; vaginal bleeding or spotting; Pelvic inflammatory disease.

Mifepristone + Misoprostol
Non-EDL Tertiary

INDICATIONS
Medical termination of pregnancy of upto 49 days, cervical dilatation prior to surgical termination of pregnancy in the first trimester, therapeutic termination of pregnancy for medical reasons beyond the first trimester, labor induction in case of fetal death in utero.

AVAILABILITY
Tablet KIT mifopristone 200 mg, misoprostol 200 μg.

DOSE
Mifepristone 200 mg orally followed 1 to 3 days latter by misoprostol 800 μg vaginally. Patients should return for followup visit after approximately 14 days after administration of mifepristone.

CONTRAINDICATIONS
Hypersensitivity to Mifepristone, Misoprostol or other prostaglandin; confirmed or suspected ectopic pregnancy (Appendix 7c); chronic adrenal failure; haemorrhagic disorders or concurrent anticoagulant therapy; inherited porphyria.
PRECAUTIONS
IUD in place; asthma, chronic obstructive pulmonary disease; alcoholism; prosthetic heart valve; infective endocarditis; interactions (Appendix 6c), pregnancy (Appendix 7c).

ADVERSE EFFECTS
Abdominal pain, diarrhoea, nausea, vomiting; fever, chills, uterine cramping; vaginal bleeding or spotting; Pelvic inflammatory disease.

Antioxytocics
Isoxsuprine
EDL-D 293 Secondary hospitals

AVAILABILITY
TABLETS 10 and 20 mg; INJECTION 2 ml ampoule (5 mg/ml).

DOSE
Premature labour and threatened abortion: initially 20 mg 6 hly after food, maintenance dose after improvement 10 mg thrice a day. Intravenous injection/infusion Premature labour and threatened abortion: 0.2 to 0.5 mg/min, adjust according to response, monitor BP and heart rate.

INDICATION
Cerebral and peripheral vascular disorder; threatened abortion and premature labour; night cramps; habitual abortion.

CONTRAINDICATION
Anaemia; heart disease, arterial hemorrhage; postpartum; premature detachment of placenta; hypersensitivity

PRECAUTION
Blood disorders, bleeding episodes or allergies, pregnancy (Appendix 7c), lactation.

ADVERSE EFFECTS
Dizziness, nausea and vomiting; tachycardia, Irregular heart beat, hypotension, chest pain; flushed skin, rashes.

Isoxsuprine Hydrochloride
EDL-D 294,295 Secondary hospitals

AVAILABILITY
TABLETS 10 and 20 mg; INJECTION 2 ml ampoule (5 mg/ml).

DOSE
Premature labour and threatened abortion: initially 20 mg 6 hly after food, maintenance dose after improvement 10 mg thrice a day. Intravenous injection/infusion Premature labour and threatened abortion: 0.2 to 0.5 mg/min, adjust according to response, monitor BP and heart rate.

INDICATION
Cerebral and peripheral vascular disorder; threatened abortion and premature labour; night cramps; habitual abortion.

CONTRAINDICATION
Anaemia; heart disease, arterial hemorrhage; postpartum; premature detachment of placenta; hypersensitivity

PRECAUTION
Blood disorders, bleeding episodes or allergies, pregnancy (Appendix 7c), lactation.

ADVERSE EFFECTS
Dizziness, nausea and vomiting; tachycardia, Irregular heart beat, hypotension, chest pain; flushed skin, rashes.
Medicines used in psychotic disorders

Treatment of psychotic disorders is both pharmacological and psychosocial. Individual and community programmes for relearning old skills and developing new ones and for learning to cope with the illness should be initiated. Classes of antipsychotic drugs include phenothiazines (for example chlorpromazine), butyrophenones (for example haloperidol), thioxanthenes (for example flupentixol) and newer ‘atypical’ neuroleptics including clozapine and risperidone. The various antipsychotic drugs do not, in general, differ in their antipsychotic activity, but differ in range and quality of adverse effects (see below).

**Acute Phase Treatment:**
The administration of chlorpromazine or haloperidol will relieve symptoms such as thought disorder, hallucinations and delusions and prevent relapse. They are usually less effective in apathetic, withdrawn patients. However, haloperidol may restore an acutely ill schizophrenic, who was previously withdrawn, or even mute and akinetic, to normal activity and social behaviour. In the acute phase chlorpromazine may be administered by intramuscular injection in a dose of 25-50 mg which can be repeated every 6-8 h while observing the patient for possible hypotension. In most cases, however, the intramuscular injection is not needed and patients can be treated with an oral dose. Haloperidol may be administered in the acute phase.

**Maintenance Therapy:**
Long-term treatment in patients with a definite diagnosis of schizophrenia may be necessary after the first episode to prevent the manifest illness from becoming chronic. The lowest possible dose of antipsychotic drug that will prevent major exacerbations of florid symptoms is used for long-term management. Too rapid a dose reduction should be avoided. Intramuscular depot preparations such as fluphenazine may be used as an alternative to oral maintenance therapy especially when compliance with oral treatment is unreliable. Exacerbations of illness in patients on maintenance drug therapy can be precipitated by stress. Withdrawal of maintenance drug treatment requires careful surveillance since it is not possible to predict the course of the disease and the patient may suffer a relapse if treatment is withdrawn inappropriately. Further, the need for continuation of treatment may not be evident on withdrawal of treatment because relapse may be delayed for several weeks.

**Adverse Effects**
They are very common with long-term administration of antipsychotic drugs. Hypotension and interference with temperature regulation, neuroleptic malignant syndrome and bone marrow depression are the most life-threatening. Hypotension and interference with temperature regulation are doserelated. They can result in dangerous falls and hypothermia in the elderly and this must be considered before prescribing these drugs for patients over 70 years of age.

Extrapyramidal symptoms are the most troublesome and are caused most frequently by the piperazine phenothiazines such as fluphenazine, the butyrophenones such as haloperidol and the depot preparations. Although easily recognized, they are not so easy to predict because they depend in part on the dose and patient susceptibility as well as the type of drug. However, there is a general tendency for low-potency drugs to have less extrapyramidal adverse effects, while high-potency drugs such as haloperidol have more extrapyramidal effects but less sedation and anticholinergic (more correctly antimuscarinic) effects. Sedation and
anticholinergic effects usually diminish with continued use. Extrapyramidal symptoms consist of parkinsonian-type symptoms including tremor which may occur gradually; dystonia (abnormal face and body movements) and dyskinesia, which may appear after only a few doses; akathisia (restlessness), which may occur after large initial doses and may resemble an exacerbation of the condition being treated; and tardive dyskinesia (an orofacial dyskinesia), which usually takes longer to develop but may develop on short-term treatment with low doses; short-lived tardive dyskinesia may occur after withdrawal of the drug. Parkinsonian symptoms are usually reversible on withdrawal of the drug and may be suppressed by anticholinergic (antimuscarinic) drugs but they may unmask or worsen tardive dyskinesia. Tardive dyskinesia is usually associated with long-term treatment and high dosage of an antipsychotic, particularly in elderly patients. There is no established treatment for tardive dyskinesias, which may be irreversible on withdrawing therapy. However, withdrawal at the earliest signs of tardive dyskinesia may halt its full development. Treatment of all patients on antipsychotics must be carefully and regularly reviewed.

Neuroleptic malignant syndrome (hypothermia, fluctuating levels of consciousness, muscular rigidity, and autonomic dysfunction with pallor, tachycardia, labile blood pressure, sweating and urinary incontinence) is a rare adverse effect of haloperidol and chlorpromazine. It is managed by discontinuing the antipsychotic, correcting fluid and electrolyte defects, and giving bromocriptine and sometimes dantrolene.

**Alprazolam**

**Non-EDL Tertiary**

**INDICATIONS**

Anxiety disorders; panic attacks, insomnia.

**AVAILABILITY**

Tablets 0.25, 0.5 and 1 mg.

**DOSE**

Oral

Adult- 0.25 to 0.5 mg daily 2 to 3 times a day.

Child- Not recommended.

**CONTRAINDICATIONS**

Respiratory depression; marked neuromuscular respiratory weakness including unstable myasthenia gravis; acute pulmonary insufficiency; sleep apnoea syndrome; severe hepatic impairment; not for chronic psychosis; should not be used alone in depression or in anxiety with depression; avoid injections containing benzyl alcohol in neonates; narrow angle glaucoma, hypersensitivity.

**PRECAUTIONS**

Respiratory disease; muscle weakness and myasthenia gravis; history of drug or alcohol abuse; marked personality disorder; pregnancy (Appendix 7c), lactation; reduce dose in elderly and debilitated and in hepatic impairment, renal impairment; avoid prolonged use (and abrupt withdrawal thereafter); interactions (Appendix 6a); periodic blood count; liver function test. Drowsiness may affect performance of skilled tasks (e.g. driving); effects of alcohol enhanced.

**ADVERSE EFFECTS**

Drowsiness and lightheadedness on the next day; confusion and ataxia (especially in the elderly); amnesia; dependence; paradoxical increase in aggression; muscle weakness; occasionally: headache, vertigo, hypotension, salivation changes, gastro-intestinal disturbances,
visual disturbances, dysarthria, tremor, changes in libido, incontinence, urinary retention; blood disorders and jaundice reported; skin reactions; rarely, apnoea and insomnia.

STORAGE
Store protected from light.

Diazepam
Non-EDL Tertiary

INDICATIONS
Short-term treatment of anxiety and insomnia; status epilepticus, recurrent seizures; febrile convulsions, adjunct in acute alcohol withdrawal; premedication; agitation.

AVAILABILITY
Tablets 2.5, 5 and 10 mg; Injection 10mg/2 ml; Capsules 10 and 15 mg.

DOSE
Oral
Adult- Anxiety: 2 mg 3 times daily, increased if necessary to 15 to 30 mg daily in divided doses. Insomnia: 5 to 15 mg at bedtime.
Child- Oral 1-2.5 mg, 3 or 4 times daily (Not for use under 6 months).
Elderly or debilitated- Anxiety: half adult dose.

CONTRAINDICATIONS
Respiratory depression; acute pulmonary insufficiency; sleep apnoea; severe hepatic impairment; myasthenia gravis; hypersensitivity.

PRECAUTIONS
Respiratory disease; muscle weakness; history of alcohol or drug abuse; marked personality disorder; lactation (Appendix 7b); reduce dose in elderly or debilitated and in hepatic impairment (avoid if severe, Appendix 7a); renal impairment; avoid prolonged use and abrupt withdrawal; porphyria; interactions (Appendix 6a, 6c); pregnancy (Appendix 7c); liver function test to be done, least amount of drug should be given in patients in whom depression accompanies anxiety and suicidal tendencies. May impair ability to perform skilled tasks, for example operating machinery, driving.

ADVERSE EFFECTS
Drowsiness and lightheadedness the next day; confusion and ataxia (especially in the elderly); amnesia; dependence; paradoxical increase in aggression; muscle weakness; occasionally headache, vertigo, salivation changes, gastrointestinal disturbances, visual disturbances, dysarthria, tremor, changes in libido, incontinence, urinary retention; blood disorders and jaundice; skin reactions; raised liver enzymes; reduces reflexes; jaundice; psychological dependence; physiological dependence, respiratory arrest.

STORAGE
Tablet: Store protected from light. Injection: Table in single dose or multi dose container protected from light.

Lorazepam
EDL-D318,319 Tertiary

INDICATIONS
Anxiety disorders.

AVAILABILITY
TABLETS 0.5, 1, 2, 2.5 and 3 mg
INJECTIONS 2 ml ampoule (2 mg/ml).

DOSE
2 to 6 mg/day given in divided doses, initial dose of 2 to 3 mg/day given twice or thrice a day. Elderly or debilitated patients: Initial dosage of 1 to 2 mg/day in divided doses.
CONTRAINDICATIONS
Severe hepatic impairment; respiratory depression; acute narrow angle glaucoma; pregnancy (Appendix 7c), lactation.

PRECAUTIONS
Hepatic dysfunction; impaired ability to drive or operate machinery; interactions (Appendix 6a).

ADVERSE EFFECTS
Nausea and vomiting, dizziness; weakness; blurred vision; vertigo.

Chlorpromazine Hydrochloride
EDL-D 118,119,120 Tertiary

AVAILABILITY
Tablets 25, 50 and 100 mg; Syrup 60 ml (25 mg/5 ml); Injection 2 ml ampoule (25 mg/ml).

DOSE
Oral Adult- Schizophrenia and other psychoses, mania, psychomotor agitation, violent behaviour and severe anxiety (adjuvant): initially 25 mg 3 times daily (or 75 mg at night) adjusted to response to usual maintenance dose of 100-300 mg daily (but up to 1.2g daily may be required in psychosis). Elderly or debilitated- Schizophrenia and other psychoses, mania, psychomotor agitation, violent behaviour and severe anxiety (adjuvant): one-third to one-half adult dose. Child- Schizophrenia and other psychoses, mania, psychomotor agitation, violent behaviour and severe anxiety (adjuvant); (for childhood schizophrenia and autism) 1 to 5 years: 500 µg/kg every 4-6 h (max. 40 mg daily). 6 to 12 years: one-third to one-half adult dose (max. 75 mg daily). Deep intramuscular injection Adult- Relief of acute symptoms: 25 to 50 mg every 6 to 8 h. Child- Relief of acute symptoms: 500 µg/kg every 6 to 8 h (1 to 5 years: max. 40 mg daily. 6 to 12 years: max. 75 mg daily).

INDICATION
Schizophrenia and other psychotic disorders, mania, psychomotor agitation and violent behaviour; adjunct in severe anxiety; psychosis, mania, hiccups.

CONTRAINDICATION
Impaired consciousness due to CNS depression; bone-marrow depression; pheochromocytoma; epilepsy, narrow angle glaucoma, Parkinson’s disease; depressed level of consciousness.

PRECAUTION
Cardiovascular and cerebrovascular disorders, respiratory disease, parkinsonism, epilepsy, acute infections, pregnancy (Appendix 7c), lactation (Appendix 7b), renal and hepatic impairment (avoid if severe; Appendixes 7a), history of jaundice, leukopenia (blood counts if unexplained fever or infection); hypothyroidism, myasthenia gravis, prostatic hypertrophy, angle-closure glaucoma; elderly (particularly in very hot or very cold weather); avoid abrupt withdrawal; patients should remain supine and the blood pressure monitored for 30 min after intramuscular injection; interactions (Appendix 6a, 6c); extreme heat, alcohol withdrawal, peptic ulcer. May impair ability to perform skilled tasks, for example operating machinery, driving.

ADVERSE EFFECTS
Extrapyramidal symptoms and on prolonged administration, occasionally potentially irreversible tardive dyskinesias (see notes above); hypothermia (occasionally pyrexia), drowsiness, apathy, pallor, nightmares, dizziness, excitement, insomnia, headache, confusion, depression; more rarely, agitation; EEG changes; convulsions; nasal congestion; anticholinergic symptoms including dry mouth, constipation; blurred vision, difficulty in micturition; hypotension, tachycardia and arrhythmias; ECG changes; respiratory depression; menstrual disturbances, galactorrhoea, gynaecomastia, impotence, weight gain; sensitivity reactions such as agranulocytosis, leukopenia, leukocytosis, haemolytic anaemia, photosensitization, contact sensitization and rashes, jaundice and alterations in liver function; neuroleptic malignant syndrome; lupus erythematosuslike syndrome; with prolonged high dosage, corneal and lens opacities, and purplish pigmentation of the skin, cornea and retina; intramuscular injection may
be painful and cause hypotension and tachycardia (see Precautions) and nodule formation; seizures, temperature disorder, hyperprolactinemia, ocular complication.

**Risperidone**

**EDL-D 459,460 Tertiary**

**INDICATION**
- Acute and chronic psychoses.

**CONTRAINDICATION**
- Same as for chlorpromazine (pg no.284)

**PRECAUTION**
- Same as for chlorpromazine (pg no.284)

**ADVERSE EFFECT**
- Same as for chlorpromazine (pg no.284)

**DRUG INTERACTION**
- Same as for chlorpromazine (pg no.284)

**AVAILABILITY**
- Tablets: 1mg, 2 mg, 3 mg, 4 mg
- Liquid 1 mg / mL

**DOSE**
- Oral: 2 mg in 1-2 divided doses on first day, and increased to 4 mg on second day, 6 mg in 1-2 divided doses on third day upto the usual range of 4-8 mg od Upto 16 mg od may be given exceptionally only if benefit is considered to outweigh the risk. Elderly, 0.5 mg bd, increased in increments of 0.5 mg bd to 1-2 mg bd. For children under 15 years not recommended.

**Fluphenazine Decanoate or Enantate**

**EDL-D 231 Secondary hospitals**

**AVAILABILITY**
- 1 mg; Injection 1 ml ampoule (25 mg/ml).

**DOSE**
- Deep intramuscular injection into gluteal muscle. Adult: Maintenance in schizophrenia and other psychoses: test dose of 12.5 mg, then after 4 to 7 days, 12.5 to 100 mg repeated at intervals of 2 to 5 weeks, adjusted according to the response. Elderly: Maintenance in schizophrenia and other psychoses: test dose of 6.25 mg, then after 4 to 7 days, 12.5 to 100 mg repeated at intervals of 2 to 5 weeks, adjusted according to the response. Child: Maintenance in schizophrenia and other psychoses: not recommended.

**INDICATION**
- Maintenance treatment of schizophrenia and other psychoses; mania, postoperative nausea.

**CONTRAINDICATION**
- As for Chlorpromazine (pg no.284), but less sedating and fewer hypotensive and anticholinergic symptoms; higher incidence of extrapyramidal symptoms (most likely to occur a few hours after injection and continue for about 2 days but may be delayed); systemic lupus erythematosus; pain at injection site, occasionally erythema, swelling, nodules; tardive dyskinesia, neurological disturbances, blood dyscrasias.

**PRECAUTION**
- Treatment requires careful monitoring for optimum effect; initial small test dose as adverse effects are prolonged; extrapyramidal symptoms occur frequently; when transferring from oral to depot therapy, dosage by mouth should be reduced gradually; cardiovascular and cerebrovascular disorders; respiratory disease, epilepsy; acute infections; pregnancy (Appendix 7c), lactation (Appendix 7b); renal and hepatic impairment (avoid if severe; Appendices 7a), history of jaundice; leukopenia (blood counts if unexplained fever or infection); hypothyroidism, myasthenia gravis, prostatic hypertrophy, angle-closure glaucoma; elderly (particularly in very hot or very cold weather); interactions (Appendix 6a, 6c); alcohol withdrawal, extreme heat. May impair ability to perform skilled tasks, for example operating machinery, driving.
ADVERSE EFFECTS

Children; confusional states; impaired consciousness due to CNS depression; parkinsonism; intolerance to antipsychotics; depression; bone-marrow depression; pheochromocytoma; blood dyscrasias, coma, brain damage.

**Haloperidol**

**EDL-D 249,250,251,252 Secondary hospitals**

**AVAILABILITY**

Tablets 1.5, 5, 10 and 20 mg; Liquid 30 ml (25 mg/ml); Injection 5 ml ampoule (5 mg/ml).

**DOSE**

- **Oral Adult**-Schizophrenia and other psychoses, mania, psychomotor agitation and violent behaviour and severe anxiety (adjuvant): initially 1.5 to 3 mg 2 to 3 times daily or 3 to 5 mg 2 to 3 times daily in severely affected or resistant patients (up to 30 mg daily in resistant schizophrenia). Elderly or debilitated-Schizophrenia and other psychoses, mania, psychomotor agitation and violent behaviour and severe anxiety (adjuvant): initially half adult dose. Child-Schizophrenia and other psychoses, mania, psychomotor agitation and violent behaviour and severe anxiety (adjuvant): initially 25 to 50 μg/kg daily in 2 divided doses (max. 10 mg daily).

- **Intramuscular injection**
  - Adult- Acute psychotic conditions: initially 2 to 10 mg, subsequent doses every 4 to 8 h according to response (up to every h if necessary) to max. of 18 mg; severely disturbed patients may require initial dose of up to 18 mg. Elderly or debilitated- Acute psychotic conditions: initially half adult dose. Child- Acute psychotic conditions: not recommended

**INDICATION**

Schizophrenia and other psychotic disorders, mania, psychomotor agitation and violent behaviour; adjunct in severe anxiety; agitation, psychosis, neuroleptanalgesia.

**CONTRAINDICATION**

Impaired consciousness due to CNS depression; bone-marrow depression; pheochromocytoma; porphyria; basal ganglia disease; parkinsonism, thyrotoxicosis, cardiac arrhythmia, depression, close angle glaucoma.

**PRECAUTION**

Cardiovascular and cerebrovascular disorders; respiratory disease; parkinsonism; epilepsy; acute infections; pregnancy (Appendix 7c), lactation (Appendix 7b); renal and hepatic impairment (avoid if severe; Appendices 7a), history of jaundice; leukopenia (blood count required if unexplained fever or infection); hypothyroidism, myasthenia gravis, prostatic hypertrophy, angle-closure glaucoma; also subarachnoid haemorrhage and metabolic disturbances such as hypokalaemia, hypocalcaemia, or hypomagnesaemia; elderly (particularly in very hot or very cold weather); children and adolescents; avoid abrupt withdrawal; patients should remain supine and the blood pressure monitored for 30 min after intramuscular injection; interactions (Appendix 6a, 6c); photosensitisation, peptic ulcers.

May impair ability to perform skilled tasks, for example operating machinery, driving.

**ADVERSE EFFECTS**

As for Chlorpromazine (pg no.284), but less sedating and fewer hypotensive and anticholinergic symptoms; pigmentation and photosensitivity reactions rare; extrapyramidal symptoms are common, particularly acute dystonia and akathisia (especially in thyrotoxic patients); rarely, weight loss, hypoglycaemia, inappropriate antidiuretic hormone secretion.

**Olanzapine**

**EDL-D 380,381 Tertiary**

**AVAILABILITY**

Tablets 2.5, 5, 7.5, 10, 15 and 20 mg.
DOSE
Schizophrenia: initial 5-10 mg, usual dose is 10-20 mg. Acute maniac episodes in bipolar illness: 10-15 mg/day.

INDICATION
Schizophrenia, acute mania episodes in bipolar disorder.

PRECAUTION
Impaired renal, hepatic and cardiovascular function; prostatic hypertrophy; paralytic ileus; parkinsonism; blood dyscrasias; myelosupression; seizures; dementia; pregnancy (Appendix 7c)

ADVERSE EFFECTS
Postural hypotension, dizziness, constipation, weight gain, agitation, insomnia, akathesia, tremors, personality disorder, oedema, increases appetite, antimuscarinic effects, hallucination, bradycardia

Amitriptyline
EDL-D 25 Tertiary

AVAILABILITY
Tablets 10, 25, 50 and 75 mg; injection 10 ml ampoule (10 mg/ml).

DOSE
Oral Adult- Initially 75 mg (adolescents 30 to 75 mg) daily in divided doses or as a single dose at bed time increased gradually as necessary to 150 to 200 mg daily. Prophylaxis of migraine: 10-75 mg at night. Child- Under 16 years; not recommended.

INDICATION
Moderate to severe depression, migraine prophylaxis; tension, headache, enuresis.

CONTRAINDICATION
Recent myocardial infarction, arrhythmias (especially heart block); manic phase in bipolar disorders; severe liver disease; children; porphyria; glaucoma, prostatic hypertrophy

PRECAUTION
Cardiac disease (see Contraindications above); history of epilepsy; lactation (Appendix 7b); elderly; hepatic impairment (Appendix 7a); thyroid disease; pheochromocytoma; history of mania, psychoses (may aggravate psychotic symptoms); angle-closure glaucoma; history of urinary retention; concurrent electroconvulsive therapy; avoid abrupt withdrawal; anaesthesia (increased risk of arrhythmias and hypotension); interactions (Appendix 6a, 6b, 6c); pregnancy (Appendix 7c); pre-existing haematological disorder, abrupt disorientation. May impair ability to perform skilled tasks, for example operating machinery, driving.

ADVERSE EFFECTS
Sedation; dry mouth; blurred vision (disturbance of accommodation, increased intraocular pressure); constipation; nausea; difficulty in micturition; cardiovascular adverse effects particularly with high dosage including ECG changes, arrhythmias, postural hypotension, tachycardia, syncope; sweating, tremor, rash and hypersensitivity reactions (urticaria, photosensitivity); behavioural disturbances; hypomania or mania, confusion (particularly in elderly), interference with sexual function, blood sugar changes; increased appetite and weight gain (occasional weight loss); endocrine adverse effects such as testicular enlargement, gynaecomastia and galactorrhoea; convulsions, movement disorders and dyskinesias, fever, agranulocytosis, leukopenia, eosinophilia, purpura, thrombocytopenia, hyponatraemia (may be due to inappropriate antidiuretic hormone secretion); abnormal liver function test.

Clozapine
EDL-D 137 Tertiary

INDICATION
Schizophrenia in patients unresponsive to, or intolerant of conventional antipsychotic drugs.

CONTRAINDICATION
Severe cardiac disease; history of drug-induced neutropenia or agranulocytosis; bone marrow
disorders; alcoholic and toxic psychoses; history of circulatory collapse or paralytic ileus; drug intoxication, coma or severe CNS depression, uncontrolled epilepsy, pregnancy and breast-feeding.

PRECAUTION
Leucocyte and differential blood counts must be normal before treatment and must be monitored weekly for first 18 weeks, then fortnightly. Avoid drugs which depress leucopoiesis, withdraw treatment if leucocyte count falls below 3000/ mm3 or absolute neutrophil count falls below 1500/ mm3. Patients should report any infections, hepatic or renal impairment, epilepsy, cardiovascular disorders, prostatic enlargement, glaucoma, paralytic ileus. Avoid abrupt withdrawal, avoid in children.

ADVERSE EFFECT
High incidence of antimuscarinic symptoms; extrapyramidal symptoms may occur less frequently, neutropenia and potentially fatal agranulocytosis, fever, headache, dizziness, urinary incontinence, priapism, pericarditis, myocarditis, delirium, hypotension, sialorrhea, skin rashes and convulsions (if dosage is above 800 mg/ day).

AVAILABILITY
Tablets 25 mg, 100 mg

DOSE
Start 12.5 mg od or bd on first day, then 25-50 mg on second day, then increase gradually in steps of 25-50 mg over 7-14 days to 300 mg od in divided doses. Larger dose upto 200 mg od may be taken as a single dose at hs Further increased in steps of 50-100 mg once or twice weekly may be required. Usual antipsychotic dose 200-450 mg od upto a maximum of 900 mg od Subsequent maintenance dose of 150-300 mg. Elderly, 12.5 mg once on first day subsequent adjustments restricted to 25 mg od.

DRUG INTERACTION
Clozapine cause agranulocytosis when used concurrently with drugs associated with a substantia potential for causing agranulocytosis, such as cotrimoxazole, chloramphenicol, sulphonamides, penicillamine, cytotoxics or carbamazepine.

Quetiapine
EDL-D 723,724 PHC

INDICATION
Schizophrenia, acute mania

CONTRAINDICATION
Patients below 18 yrs

PRECAUTION
Renal or hepatic impairment, epilepsy, cardiovascular or cerebro-vascular disease, hypotension

ADVERSE EFFECT
Hyperglycemia, weight gain, sedation, dizziness. Rarely, Neuroleptic Malignant Syndrome ` (NMS),seizures

AVAILABILITY
25mg, 50mg, 100mg, 200mg, 300mg Sustained Release preparation- 50mg, 100mg, 200mg Dose: 50- 800 mg/day. Initially 25 mg twice daily. Increase by 25-50 mg twice daily. Maximum dose-800mg/ day

Fluoxetine
Non-EDL Tertiary

INDICATION
Depression, obsessive compulsive disorder (OCD), panic disorders, anxiety disorders.

CONTRAINDICATION
Pregnancy, lactation.
PRECAUTION
Use with caution in patients with seizures and diabetes.

ADVERSE EFFECT
Insomnia, anorexia, nausea, diarrhoea, headache, nervousness, anxiety, seizures in high doses, sexual dysfunction.

AVAILABILITY
Capsule 20 mg. Suspension 20 mg/5 mL

DOSE
Depression - 20 mg/day. OCD - 60 mg/day.

DRUG INTERACTION
Increased sedation with other drugs having sedative effect on central nervous system. Produces agitation, restlessness and gastric distress with tryptophan. Produces changes in serum lithium level. Produces sedation, dry mouth and constipation with other antidepressants.

Sertraline
EDL-D 470 PHC

INDICATION
Same as for fluoxetine (pg no.288)

PRECAUTION
Same as for fluoxetine (pg no.288).

ADVERSE EFFECT
Same as for fluoxetine (pg no.288).

DRUG INTERACTION
Same as for fluoxetine (pg no.288).

CONTRAINDICATION
Hypersensitivity, pregnancy, lactation, history of drug abuse, hepatic or renal impairment, seizure disorders.

AVAILABILITY
Tablet 50 mg

DOSE
100 - 150 mg/day.

Medicines used in mood disorders
Tricyclic and related antidepressants and the more recently introduced selective serotonin reuptake inhibitors (SSRIs) are the most widely used drugs in the treatment of depressive disorders. The response to antidepressant therapy is usually delayed with a lag-period of up to two weeks and at least six weeks before max. improvement occurs. It is important to use doses that are sufficiently high for effective treatment, but not so high as to cause toxic effects. Low doses should be used for initial treatment in the elderly. The use of more than one antidepressant at a time is not recommended since this does not enhance effectiveness and it may result in enhanced adverse effects or interactions.

Patients should be reviewed every 1-2 weeks at the start of treatment. Treatment should be continued for at least 4 weeks (6 weeks in the elderly) before considering whether to change to another antidepressant due to lack of efficacy. In the case of a partial response, treatment may be continued for a further 2 weeks (elderly patients may take longer to respond). Remission usually occurs after 3-12 months. Treatment at full therapeutic dose should be continued for at least 4-6 months after resolution of symptoms (about 12 months in the elderly). Treatment should not be withdrawn prematurely otherwise symptoms are likely to recur. Patients with a
history of recurrent depression should continue to receive maintenance treatment (for at least 5 years and possibly indefinitely). Lithium may be used as an alternative for maintenance treatment. Reduction in dose should be gradually carried out over a period of about 4 weeks or longer if withdrawal symptoms emerge (6 months in patients who have been on long-term maintenance treatment).

Tricyclic and related antidepressants can be divided into those with more or less sedative effect. Those with sedative properties include amitriptyline and those with less sedative effects include imipramine. These drugs are most effective in the treatment of depression associated with psychomotor and physiological disturbances. Adverse effects include anticholinergic (more correctly antimuscarinic) symptoms of dry mouth, blurred vision, constipation and urinary retention. Arrhythmias and heart block can occur. Minimal quantities of tricyclic antidepressants should be prescribed at any one time because they are dangerous in overdose. The SSRIs characteristically cause gastrointestinal disturbances, sleep disturbances and hypersensitivity reactions including rash (may be a sign of an impending serious systemic reaction and discontinuation should be considered) but they are less sedating and have fewer anticholinergic (antimuscarinic) and cardiotoxic effects than tricyclic antidepressants. The SSRIs are less toxic in overdose than the older tricyclic compounds. They may be preferred in patients in whom the risk of suicide is strong, but there is some concern that SSRIs may increase suicidal ideation.

**Imipramine Hydrochloride**

**EDL-D 275 PHC**

**AVAILABILITY**

- Tablets 5, 25 and 75 mg;
- Capsules 25 and 75 mg.

**DOSE**

- Oral: 75 mg/day initially, usual dose 100 to 200 mg daily. Child:<6 years: not recommended, 6-12 years: 25 mg at bed time, >12 years: 50 mg at bed time.

**INDICATION**

- Panic attacks; chronic pain; nocturnal enuresis; Kleine-Levin syndrome; depression, hyperactivity, attention deficit disorder.

**CONTRAINDICATION**

- Recent myocardial infarction, arrhythmias (particularly heart block), not indicated in manic phase, severe liver disease; epilepsy, mania, narrow angle glaucoma, hypersensitivity.

**PRECAUTION**

- Cardiac disease (particularly with arrhythmias), history of epilepsy, pregnancy, lactation, elderly, hepatic impairment, interactions (Appendix 6a), thyroid disease, pheochromocytoma, history of mania, psychoses (may aggravate psychotic symptoms), susceptibility to angle-closure glaucoma, history of urinary retention, concurrent electroconvulsive therapy; if possible avoid abrupt withdrawal; anaesthesia (increased risk of arrhythmias and hypotension), see surgery; porphyria; for additional nocturnal enuresis warnings; acetylsalicylic acid hypersensitivity. Drowsiness may affect performance of skilled tasks (e.g. driving); effects of alcohol enhanced.

**ADVERSE EFFECTS**

- Dry mouth, sedation, blurred vision (disturbance of accommodation, increased intraocular pressure), constipation, nausea, difficulty with micturition; cardiovascular sideeffects (such as ECG changes, arrhythmias, postural hypotension, tachycardia, syncope, particularly with high doses); sweating, tremor, rashes and hypersensitivity reactions (including urticaria, photosensitivity), behavioural disturbances (particularly children), hypomania or mania, confusion or delirium (particularly elderly), headache, interference with sexual function, blood sugar changes; increased appetite and weight gain (occasionally weight loss); endocrine side-
effects such as testicular enlargement, gynaecomastia, galactorrhoea; also convulsions, movement disorders and dyskinesias, dysarthria, paraesthesia, taste disturbances, tinnitus, fever, agranulocytosis, leucopenia, eosinophilia, purpura, thrombocytopenia, hyponatraemia (see Hyponatraemia and Antidepressant Therapy), abnormal liver function tests (jaundice); impairment of memory, cutaneous vasculitis.

Fluoxetine Hydrochloride (for use above 8 years of age)

EDL-D 230 Tertiary

AVAILABILITY
Tablets 10, 20, 40 and 60 mg; Capsules 10, 20 and 60 mg.

DOSE
Oral 20 mg/day initially (max 60 mg).

INDICATION
Major depression (including pediatric depression); obsessive-compulsive disorder (in both adult and pediatric populations); bulimia nervosa; anorexia nervosa; panic disorder and premenstrual dysphoric disorder; depression illness, Parkinson’s disease.

CONTRAINDICATION
Should not be used if the patient enters a manic phase; renal failure, hypersensitivity.

PRECAUTION
Should be used with caution in patients with epilepsy (avoid if poorly controlled, discontinue if convulsions develop), cardiac disease, diabetes mellitus, susceptibility to angle-closure glaucoma, a history of mania or bleeding disorders (especially gastrointestinal bleeding), and if used with other drugs that increase the risk of bleeding, hepatic impairment (Appendix 7a), renal impairment, pregnancy (Appendix 7c), and lactation. They should also be used with caution in those receiving concurrent electroconvulsive therapy (prolonged seizures reported with fluoxetine). The risk of suicidal behaviour is possibly higher in young adults, calling for close monitoring of those receiving SSRIs. SSRIs may also impair performance of skilled tasks (e.g. driving), interactions

ADVERSE EFFECTS
Gastro-intestinal effects (dose-related and fairly common—include nausea, vomiting, dyspepsia, abdominal pain, diarrhoea, constipation), anorexia with weight loss (increased appetite and weight gain also reported) and hypersensitivity reactions including rash (consider discontinuation—may be sign of impending serious systemic reaction, possibly associated with vasculitis), urticaria, angioedema, anaphylaxis, arthralgia, myalgia and photosensitivity; other side-effects include dry mouth, nervousness, anxiety, headache, insomnia, tremor, asthenia, hallucinations, drowsiness, convulsions, galactorrhoea, sexual dysfunction, urinary retention, sweating, hypomania or mania, movement disorders and dyskinesias, visual disturbances, hyponatraemia; serum sickness, elevation of liver enzymes.

Mirtazapine

EDL-D 691,692 PHC

INDICATION
Depression

AVAILABILITY
Tablets- 15mg and 30mg

DOSE
Tab 15-45 mg/day. Initially 15mg once per day at night. Increase 15 mg/day once in every 5 days to a maximum dose of 60 mg/day

CONTRAINDICATION
Pregnancy, lactation, concomitant use with or within 14 days of use of MAOI

PRECAUTION
Renal or hepatic impairment, epilepsy, organic brain syndrome, cardiac conduction disturbances, angina, glaucoma, mania, psychosis
ADVERSE EFFECT
Increased appetite, weight gain, drowsiness, dizziness, somnolence, nausea, constipation, asthenia, flu syndrome, headache, bone marrow depression, agranulocytosis

DRUG INTERACTION
Potentiates the effect of CNS depressants

Escitalopram
EDL-D 625,626,627 PHC

AVAILABILITY
TABLETS 5, 10 and 20 mg.

DOSE
Initially 10 mg once daily. Maximum- 20 mg daily.

INDICATION
Depression, obsessive compulsive disorder, anxiety disorder, panic disorder.

CONTRAINDICATION
Concomitant use with MAO Inhibitors, thioridazine.

PRECAUTION
History of panic disorder or seizure disorders, renal impairment, hepatic impairment, work requiring mental alertness, concomitant use of escitalopram with other SSRIs, serotonin–norepinephrine reuptake inhibitors (SNRIs) or tryptophan, interactions , pregnancy

ADVERSE EFFECTS
Insomnia, nausea, ejaculation disorder.

Medicines used in bipolar disorders
Treatment of bipolar disorders has to take account of three stages: treatment of the acute episode, continuation phase and prophylaxis to prevent further episodes. Lithium is effective in acute mania but symptomatic control of the florid symptoms with an antipsychotic or benzodiazepine is often necessary whilst waiting for the antimania drug to exert its effect. Benzodiazepines may be given during the initial stages until lithium becomes effective but they should not be used for long periods because of the risk of dependence. Lithium may be given concurrently with antipsychotics and treatment with the antipsychotic should be tailed off as lithium becomes effective. Alternatively, lithium therapy may be delayed until the patient’s mood is stabilized with the antipsychotic. However, there is a risk of neurotoxicity and increased extrapyramidal disorders when lithium and antipsychotics are used concurrently (Appendix 6c). Lithium is the mainstay of treatment but its narrow therapeutic range is a disadvantage. Sodium valproate is effective and carbamazepine may also be used.

Treatment of depressive episodes in bipolar disorders will mostly involve combination treatment using either lithium or Sodium valproate together with a tricyclic antidepressant. Increased adverse effects are a problem which may compromise treatment.

Lithium prophylaxis should usually only be undertaken with specialist advice and the likelihood of recurrence considered. Long-term lithium therapy has been associated with thyroid disorders and mild cognitive and memory impairment. Patients should continue the treatment for longer than 3 to 5 years only if benefit persists.

Withdrawal appears to produce high levels of relapse. If lithium is to be discontinued, the dose should be reduced gradually over a few weeks and patients should be warned of possible relapses if discontinued abruptly.
Lithium salts have a narrow therapeutic/toxic ratio and should only be prescribed if there are facilities for monitoring serum lithium concentrations. Doses are adjusted to achieve serum lithium concentrations of 0.4-1 mmol/litre (lower end of range for maintenance therapy and the elderly) on samples taken 12 h after the preceding dose. The optimum range for each patient should be determined.

Overdosage, usually with serum-lithium concentration of over 1.5 mmol/litre may be fatal and toxic effects include coarse tremor, ataxia, dysarthria, nystagmus, renal impairment and convulsions. If any of these effects occur, treatment should be stopped, serum-lithium concentration determined and in mild overdosage large amounts of sodium and fluid should be given to reverse the toxicity; in severe toxicity, haemodialysis may be required.

For patients who are unresponsive to or intolerant of lithium, carbamazepine may be used in the prophylaxis of bipolar illness particularly in those with rapid cycling affective disorders (more than four affective episodes per year).

**Carbamazepine**

**EDL-D 86,87 Secondary hospitals**

**INDICATIONS**
Prophylaxis of bipolar disorder unresponsive to or intolerant of lithium; epilepsy, trigeminal neuralgia.

**AVAILABILITY**
Tablets 100, 200 and 400 mg Plain; 100 mg (DT) Syrup 100 mg/5 ml.

**DOSE**
Oral
Adult- Initially 400 mg daily in divided doses increased until symptoms are controlled to a max. of 1.6g daily: usual maintenance range 400 to 600 mg daily.
Trigeminal neuralgia: initially 100 mg twice daily, maintenance dose is 400-800 mg/day.

**CONTRAINDICATIONS**
Atrioventricular conduction abnormalities; history of bone-marrow depression; porphyria.

**PRECAUTIONS**
Hepatic impairment (Appendix 7a); renal impairment; cardiac disease (see also Contraindications); skin reactions (see Adverse effects); history of blood disorders (blood counts before and during treatment); glaucoma; (neural tube screening); lactation (Appendix 7b); avoid sudden withdrawal; interactions (Appendix 6b, 6c, 6d); pregnancy (Appendix 7c); patients on anticoagulants. Patients or their caretakers should be told how to recognize signs of blood, liver or skin disorders, and advised to seek immediate medical attention if symptoms such as fever, sore throat, rash, mouth ulcers, bruising or bleeding develop. Leukopenia which is severe, progressive and associated with clinical symptoms requires withdrawal (if necessary under cover of suitable alternative). May impair ability to perform skilled tasks, for example operating machinery, driving.

**ADVERSE EFFECTS**
Dizziness; drowsiness; headache; ataxia; blurred vision; diplopia (may be associated with high plasma concentrations); gastrointestinal intolerance including nausea and vomiting, anorexia, abdominal pain, dry mouth, diarrhoea or constipation; commonly, mild transient generalized erythematous rash (withdraw if worsens or is accompanied by other symptoms); leukopenia and other blood disorders (including thrombocytopenia, agranulocytosis and aplastic anaemia); cholestatic jaundice, hepatitis, acute renal failure, Stevens-Johnson syndrome (erythema
multiforme), toxic epidermal necrolysis, alopecia, thromboembolism, arthralgia, fever, proteinuria, lymph node enlargement, arrhythmias, heart block and heart failure, dyskinesias, paraesthesia, depression, impotence, male infertility, gynaecomastia, galactorrhoea, aggression, activation of psychosis, photosensitivity, pulmonary hypersensitivity, hyponatraemia, oedema, disturbances of bone metabolism with osteomalacia also reported; confusion and agitation in elderly; exfoliative dermatitis, ankle swelling.

**Lithium Carbonate**

**EDL-D 316 Secondary hospitals**

**AVAILABILITY**
Tablets 150, 200, 300 and 400 mg; Capsules 150 and 300 mg.

**DOSE**
Oral Adult-Treatment of mania: initially 0.6 to 1.8g daily. Prophylaxis of mania, bipolar disorder and recurrent depression: initially 0.6 to 1.2g daily. Elderly-Treatment of mania: initially 300 to 900 mg daily. Prophylaxis of mania, bipolar disorder and recurrent depression: initially 300 to 900 mg daily.

**INDICATION**
Treatment and prophylaxis of mania, prophylaxis of bipolar disorder and recurrent depression; ADH secretion syndrome, psychosis.

**CONTRAINDICATION**
Renal impairment; cardiac insufficiency; conditions with sodium imbalance such as Addison’s disease; fetal goiter; heart failure; psoriasis; kidney infection; hypothyroidism

**PRECAUTION**
Measure serum-lithium concentration about 4 days after starting treatment, then weekly until stabilized, then at least every 3 months; monitor thyroid function every 6-12 months on stabilized regimens-risk of hypothyroidism (see below); monitor renal function; maintain adequate fluid and sodium intake; reduce dose or discontinue in diarrhoea, vomiting and intercurrent infection (especially if associated with profuse sweating); lactation; pregnancy; elderly (reduce dose); diuretic treatment, myasthenia gravis; surgery; if possible, avoid abrupt withdrawal (see notes above); interactions (Appendix 6c, 6d); kidney, thyroid and heart function test, children and adolescents. Patients should maintain adequate fluid intake and should avoid dietary changes which may reduce or increase sodium intake. Patients should be advised to seek medical attention if symptoms of hypothyroidism (for example, feeling cold, lethargy) develop (women are at greater risk).

**ADVERSE EFFECTS**
Gastrointestinal disturbances; fine tremor, renal impairment (particularly impaired urinary concentration and polyuria); polydipsia, weight gain and oedema (may respond to dose reduction); hyperparathyroidism and hypercalcaemia reported; signs of intoxication include blurred vision; muscle weakness, increasing gastrointestinal disturbances (anorexia, vomiting, diarrhoea); increased CNS disturbances (mild drowsiness and sluggishness, increasing to giddiness with ataxia, coarse tremor, lack of co-ordination, dysarthria) and require withdrawal of treatment; with severe overdosage (serum concentrations above 2 mmol/litre), hyperreflexia and hyperextension of the limbs; convulsions; toxic psychoses; syncope; renal failure; circulatory failure; coma; occasionally death; goitre, raised antidiuretic hormone concentration, hypothyroidism, hypokalaemia, ECG changes, exacerbation of psoriasis and kidney changes may occur; sinus bradycardia, leukocytosis, glycosuria, weight gain.

**Valproic acid**

**EDL-D 514,515 Secondary hospitals**

**INDICATION**
acute mania; epilepsy
DOSE
Acute mania, by mouth, ADULT initially 750 mg daily in divided doses, increased as quickly as possible to achieve the optimal response (maximum 60 mg/kg daily).

CONTRAINDICATIONS
Active liver disease, family history of severe hepatic dysfunction; pancreatitis; porphyria

PRECAUTIONS
Monitor liver function before and during therapy, especially in patients at most risk (those with metabolic disorders, degenerative disorders, organic brain disease or severe seizure disorders associated with mental retardation); ensure no undue potential for bleeding before starting and before major surgery or anticoagulant therapy; renal impairment; pregnancy (Appendix 2 (neural tube screening)); breastfeeding (Appendix 3); systemic lupus erythematosus; false-positive urine tests for ketones; avoid sudden withdrawal; Interactions: Appendix 1

BLOOD OR HEPATIC DISORDERS. Patients or their carers should be told how to recognize signs of blood or liver disorders, and advised to seek immediate medical attention if symptoms including malaise, weakness, anorexia, lethargy, oedema, vomiting, abdominal pain, drowsiness, jaundice, or spontaneous bruising or bleeding develop.

ADVERSE EFFECTS
Gastrointestinal irritation, nausea, increased appetite and weight gain, hyperammonaemia; ataxia, tremor; transient hair loss (regrowth may be curly); oedema, thrombocytopenia, inhibition of platelet aggregation; impaired hepatic function and rarely fatal hepatic failure (see Precautions—withdraw treatment immediately if malaise, weakness, lethargy, oedema, abdominal pain, vomiting, anorexia, jaundice, drowsiness); sedation reported and also increased alertness; behavioural disturbances; rarely pancreatitis (measure plasma amylase if acute abdominal pain), leukopenia, pancytopenia, red cell hypoplasia, fibrinogen reduction; irregular periods, amenorrhoea, gynaecomastia, hearing loss, Fanconi syndrome, dementia, toxic epidermal necrolysis Stevens-Johnson syndrome (erythema multiforme) and vasculitis reported.

Medicines used in generalized anxiety and sleep disorders
A sedative drug decreases activity, moderates excitement and calms the recipient, whereas, a hypnotic drug produces drowsiness and facilitates the onset and maintenance of a sleep state that resembles natural sleep. The most widely used anxiolytics and hypnotics are the benzodiazepines. Treatment of anxiety should be limited to the lowest effective dose for the shortest possible time. The cause of insomnia should be established and appropriate treatment for underlying factors instituted before hypnotics are considered. Hypnotics may be of value for a few days but rarely, longer than a week.

Tolerance and dependence (both physical and psychological) and subsequent difficulty in withdrawing the drug may occur after regular use for more than a few weeks. Patients with chronic anxiety, alcohol or drug dependence or those with personality disorders are more likely to become dependent. Anxiolytics and hypnotics should be prescribed in carefully individualized dosage and use should be limited to control of acute conditions such as panic attacks and acute anxiety and severe, incapacitating insomnia. There is usually no justification for prolonging treatment with anxiolytics and hypnotics for more than one to two weeks.
If used for longer periods, withdrawal should be gradual by reduction of the dose over a period of weeks or months, as abrupt discontinuation may produce confusion, toxic psychosis, convulsions or a condition resembling delirium tremens. The benzodiazepine withdrawal syndrome may develop at any time up to 3 weeks after stopping a long-acting benzodiazepine but may occur within a few hour in the case of a short-acting one. The syndrome is characterized by insomnia, anxiety, loss of appetite and body-weight, tremor, perspiration, tinnitus and perceptual disturbances. These symptoms may be similar to the original complaint and encourage further prescribing. Some symptoms may continue for weeks or months after stopping benzodiazepines.

Patients should be warned that their ability to drive or operate machinery may be impaired and that the effects of alcohol may be enhanced.

**Diazepam**

**EDL-D 165 PHC**

**INDICATIONS**
Short-term treatment of anxiety and insomnia; status epilepticus, recurrent seizures; febrile convulsions, adjunct in acute alcohol withdrawal; premedication; agitation.

**AVAILABILITY**
Tablets 2.5, 5 and 10 mg; Injection 10mg/2 ml; Capsules 10 and 15 mg.

**DOSE**
Oral
Adult- Anxiety: 2 mg 3 times daily, increased if necessary to 15 to 30 mg daily in divided doses. Insomnia: 5 to 15 mg at bedtime.
Child- Oral 1-2.5 mg, 3 or 4 times daily (Not for use under 6 months).
Elderly or debilitated- Anxiety: half adult dose.

**CONTRAINDICATIONS**
Respiratory depression; acute pulmonary insufficiency; sleep apnoea; severe hepatic impairment; myasthenia gravis; hypersensitivity.

**PRECAUTIONS**
Respiratory disease; muscle weakness; history of alcohol or drug abuse; marked personality disorder; lactation (Appendix 7b); reduce dose in elderly or debilitated and in hepatic impairment (avoid if severe, Appendix 7a); renal impairment; avoid prolonged use and abrupt withdrawal; porphyria; interactions (Appendix 6a, 6c); pregnancy (Appendix 7c); liver function test to be done, least amount of drug should be given in patients in whom depression accompanies anxiety and suicidal tendencies.
May impair ability to perform skilled tasks, for example operating machinery, driving.

**ADVERSE EFFECTS**
Drowsiness and lightheadedness the next day; confusion and ataxia (especially in the elderly); amnesia; dependence; paradoxical increase in aggression; muscle weakness; occasionally headache, vertigo, salivation changes, gastrointestinal disturbances, visual disturbances, dysarthria, tremor, changes in libido, incontinence, urinary retention; blood disorders and jaundice; skin reactions; raised liver enzymes; reduces reflexes; jaundice; psychological dependence; physiological dependence, respiratory arrest.

**STORAGE**
Tablet: Store protected from light. Injection:
Store in single dose or multi dose container protected from light.
Chlordiazepoxide
EDL-D 585 Secondary hospitals

**INDICATION**
For short term use in anxiety. Adjunct in acute alcohol withdrawal symptoms.

**AVAILABILITY**
Tablets 10 mg and 25 mg

**CONTRAINDICATION**
Same as diazepam.

**PRECAUTION**
Same as diazepam.

**ADVERSE EFFECT**
Same as diazepam.

**DRUG INTERACTION**
Same as diazepam.

**DOSE**
Anxiety 10 mg tds increased if necessary to 60-100 mg od in divided doses; For elderly half adult dose.
Adjunct in acute alcohol withdrawal symptom: 10-50 mg qds., gradually reducing over 7-14 days.

Clonazepam
EDL-D 588 PHC

**AVAILABILITY**
Tablets 0.25, 0.5, 1 and 2 mg.

**DOSE**
Adult: 0.5 - 5 mg thrice daily, initial dose should not exceed 1.5 mg/day, slow titration is recommended Maintenance dose 4-8 mg daily, Maximum dose 20 mg daily. Infants and child: Initial dose 0.01-0.03 mg/kg/day (not to exceed 0.05 mg/kg/day) given in 2-3 divided doses. Maintenance dose 0.1-0.2 mg/kg/day in 3 divided doses. Panic disorder: Adult- Initial dose 0.25 mg twice daily, usual maintenance dose 1 mg/day, maximum dose 4 mg/day.

**INDICATION**
Absence seizures, myoclonic seizures, akinetic seizures, panic disorder, subcortical myoclonus, adjuvant treatment of refractory epilepsy.

**CONTRAINDICATION**
Hypersensitivity to benzodiazepines, acute pulmonary insufficiency, acute narrow angle glaucoma.

**PRECAUTION**
Neonates, chronic pulmonary insufficiency, hepatic and renal dysfunction, porphyria, elderly, pregnancy, lactation, avoid sudden withdrawal.

**ADVERSE EFFECTS**
Sedation, dullness, CNS depression, ataxia, bronchial hypersecretion, abnormal eye movement, blood dyscrasias
Antiasthmatic medicines

Asthma:
Asthma is a chronic inflammatory disease characterized by episodes of reversible airways obstruction due to bronchial hyperresponsiveness; inflammation may lead to irreversible obstruction in few patients. A classification based on severity before the start of treatment and disease progression is of importance when decisions have to be made about management. It can be divided by severity into intermittent, mild persistent, moderate persistent and severe persistent. Antiasthmatics are useful in the management of the disease since therapy has a stepwise approach which must be discussed with the patient before commencing therapy. The level of therapy is increased as the severity of the asthma increases with stepping-down if control is sustained (see tables on treatment below).

Inhalation:
Medications for asthma can be administered in several different ways, including inhalation, oral and parenteral (subcutaneous, intramuscular or intravenous routes). The main advantage of delivering drugs directly into the airways via inhalation is that high concentrations can be delivered more effectively and rapidly to the airways, and systemic adverse effects avoided or minimized.

It is important that patients receive careful instruction in the use of pressurized (aerosol) inhalation (using a metereddose inhaler) to obtain optimum results. Before use, the inhaler should be shaken well. After exhaling as completely as possible, the mouthpiece of the inhaler should be placed well into the mouth and the lips firmly closed around it. The patient should inhale deeply through the mouth while actuating the inhaler. After holding the breath for 10 seconds or as long as is comfortable, the mouthpiece should be removed and the patient should exhale slowly.

It is important to check that patients continue to use their inhalers correctly as inadequate technique may be mistaken for drug failure. Spacing devices provide a space between the inhaler and the mouth. They may be of benefit for patients such as the elderly, small children and the asthmatic who find inhalers difficult to use or for those who have difficulty synchronizing their breathing with administration of the aerosol. A large volume spacing device is also recommended for inhalation of high doses of corticosteroids to reduce oropharyngeal deposition which can cause candidosis. The use of metered-dose inhalers with spacers is less expensive and may be as effective as use of nebulizers, although drug delivery may be affected by choice of spacing device. Breath-actuated devices including dry powder inhalers are also available. Solutions for nebulization are available for use in acute severe asthma. They are administered over a period of 5-10 min from a nebulizer, usually driven by oxygen in hospital.

Oral:
The oral route is used when administration by inhalation is not possible. Systemic adverse effects occur more frequently when a drug is given orally rather than by inhalation. Drugs given by mouth for the treatment of asthma include β2-agonists, corticosteroids and theophylline.
Parenteral:
Drugs such as corticosteroids, aminophylline etc. may be given by injection in acute severe asthma when administration by nebulization is inadequate or inappropriate. If the patient is being treated in the community, urgent transfer to hospital should be arranged.

Pregnancy:
Poorly controlled asthma in pregnant women can have an adverse effect on the fetus, resulting in perinatal mortality, increased prematurity and low birth-weight. For this reason using medications to obtain optimal control of asthma is justified. Administration of drugs by inhalation during pregnancy has the advantage that plasma drug concentrations are not likely to be high enough to have an effect on the fetus. Acute exacerbations should be treated aggressively in order to avoid fetal hypoxia.

Acute Exacerbation of Asthma:
Severe asthma can be fatal and must be treated promptly and energetically. Acute severe asthma attacks require hospital admission where resuscitation facilities are immediately available.
Severe asthma is characterized by persistent dyspnoea poorly relieved by bronchodilators, exhaustion, a high pulse rate (usually more than 110/min) and a very low peak expiratory flow.

As asthma becomes more severe, wheezing may be absent. Patients should be given oxygen 40-60% (if available). Patients should also be given salbutamol or terbutaline via a nebulizer. In emergencies where a nebulizer is not available, salbutamol 100 μg by aerosol inhalation can be repeated 10-20 times preferably using a large-volume spacing device. Patients should also be given a corticosteroid; for adults, prednisolone 30-60 mg by mouth or hydrocortisone 200 mg intravenously; for children, prednisolone 1-2 mg/kg by mouth (1-4 years, max. 20 mg, 5-15 years, max. 40 mg) or hydrocortisone 100 mg intravenously; if the patient experiences vomiting the parenteral route may be preferred for the first dose.

If response is inadequate, ipratropium by nebulizer should be considered. Most patients do not benefit from the addition of intravenous aminophylline or a parenteral β2-agonist; both cause more adverse effects than nebulized β2-agonists. Nevertheless, an occasional patient who has not been taking theophylline, may benefit from a slow intravenous infusion of aminophylline.

The use of epinephrine (adrenaline) in asthma has generally been superseded by β2-selective adrenoceptor agonists.

Treatment should never be delayed for investigations, patients should never be sedated and the possibility of pneumothorax should be considered. Patients who deteriorate further despite treatment may need intermittent positive pressure ventilation.
Aminophylline
EDL-D 25 PHC

AVAILABILITY
TABLETS 100, 200, 225, and 350 mg; INJECTION 10 ml (250 mg/2 ml, 25 mg/ml); ORAL LIQUID 105 mg/5 ml; SUPPOSITORY 250 mg, 500 mg.

DOSE
Parenteral/Oral Adult- 250-500 mg orally or by slow i.v injection. Loading dose- 5 mg/kg. Maintenance dose- 0.5 mg/kg/h. Child- (6 months – 9 years) 1 mg/kg/h. (10 – 16 years) 800 μg/kg/h

INDICATION
Status asthmaticus, chronic obstructive pulmonary disease (COPD), reversible airway obstruction, chronic bronchitis, pulmonary edema, adjunct in treating CHF, apnoea in premature infants.

CONTRAINDICATION
Hypersensitivity to theophyllines

PRECAUTION
Alcohol dependence; hyperthyroidism; peptic ulcer; febrile illness; patients with severe heart, liver or kidney disease; lactation (Appendix 7b); renal impairment (Appendix 7d); interactions (Appendix 6c); congestive heart failure; neonates and elderly patients; epilepsy; high blood pressure; glaucoma; diabetes; allergies, pregnancy (Appendix 7c).

ADVERSE EFFECTS
Convulsions; hypokalemia; dizziness, headache; palpitation, tachycardia, diarrhoea; anxiety; urinary retention; restlessness; tremors; abdominal pain; exfoliative dermatitis; erythema.

Adrenaline (Epinephrine)
EDL-D 199 PHC

INDICATIONS
Severe anaphylactic reaction; severe angioedema; cardiac arrest; hemostatic agent.

AVAILABILITY
INJECTION 1 ml ampoule (1 mg/ml).

DOSE
Intramuscular injection Anaphylaxis: preferable site is the midpoint in anterior thigh [1:1000 solution]. This route should be used by specialists only with extreme care.
Slow intravenous injection When there is doubt regarding adequacy of circulation and absorption from the intramuscular site; slow intravenous injection of 1:10000 (10 mg/ml) solution be injected in severely ill patients only.

CONTRAINDICATIONS
Narrow angle glaucoma, organic brain damage, cardiac dilation, coronary insufficiency.

PRECAUTIONS
Hyperthyroidism, hypertension, diabetes mellitus, heart disease, arrhythmias, cerebrovascular disease; second stage of labour; elderly; interactions (Appendix 6c); pregnancy (Appendix 7c); lactation (Appendix 7b).

ADVERSE EFFECTS
“Epinephrine fastness”, tachycardia and arrhythmias, hypertension, tremor, anxiety, sweating, nausea, vomiting, weakness, hyperglycaemia, dizziness, pulmonary oedema have all been reported; headache common.

STORAGE
Store protected from light preferably in containers filled with nitrogen.
Salbutamol Sulphate

EDL-D 461,462,463,464,465,466,467 PHC

AVAILABILITY
TABLETS 2 and 4 mg; SYRUP 2 mg/5 ml (100 ml); CAPSULES 4 mg; INHALER 100, 200 doses (100 μg per actuation).

DOSE

Oral Adult- Chronic asthma (when inhalation is ineffective): 2 to 4 mg, 3 or 4 times daily; in some patients up to max. of 8 mg, 3 or 4 times daily. Child- Chronic asthma (when inhalation is ineffective): under 2 years; 100 μg/kg, 4 times daily. 2 to 6 years; 1 to 2 mg, 3 to 4 times daily. Slow intravenous injection Adult- Severe acute bronchospasm: 250 μg, repeated if necessary. Aerosol inhalation and intramuscular or subcutaneous injection Adult- Relief of acute bronchospasm: 100 to 200 μg (1 to 2 puffs) by aerosol inhalation and 500 μg by intramuscular or subcutaneous injection; repeated every 4 h if necessary. Child- Relief of acute bronchospasm: 100 μg (1 puff) increased to 200 μg (2 puffs); if necessary. Aerosol inhalation Adult- Prophylaxis of exercise-induced bronchospasm: 200 μg (2 puffs). Chronic asthma (as adjunct in stepped treatment): 100 to 200 μg (1 to 2 puffs), up to 3 to 4 times daily. Child- Prophylaxis of exercise-induced bronchospasm: 100 μg (1 puff) increased to 200 μg (2 puffs); if required. Chronic asthma (as adjunct in stepped treatment): 100 μg (1 puff) 3 to 4 times daily, increased to 200 μg (2 puffs) 3 to 4 times daily; if necessary. Inhalation of nebulized solution Adult- Severe acute asthma or chronic bronchospasm unresponsive to conventional treatment: 2.5 mg repeated up to 4 times daily; may be increased to 5 mg, if necessary medical assessment should be considered since alternative therapy may be indicated. Child- Severe acute asthma or chronic bronchospasm unresponsive to conventional treatment, over 18 months: 2.5 mg repeated up to 4 times daily; may be increased to 5 mg, if necessary medical assessment should be considered since alternative therapy may be indicated. Under 18 months: clinical efficacy uncertain (transient hypoxaemia may occur consider oxygen supplementation).

INDICATION
Prophylaxis and treatment of asthma; premature labour; reversible airway obstruction

CONTRAINDICATION
β2agonists are contraindicated in cardiac disease; antepartum haemorrhage; intrauterine infection; intrauterine fetal death; placenta praevia; abruptio placenta; threatened miscarriage; cord compression; eclampsia or severe pre-eclampsia; diabetes mellitus; thyrotoxicosis.

PRECAUTION
Hyperthyroidism; myocardial insufficiency; arrhythmias; susceptibility to QT-interval prolongation; hypertension; pregnancy (Appendix 7c) but appropriate to use; lactation diabetes mellitus-especially intravenous administration (monitor blood glucose; ketoacidosis reported); interactions (Appendix 6c).

ADVERSE EFFECTS
Hypokalaemia after high doses; arrhythmias; tachycardia; palpitations; peripheral vasodilation; fine tremor (usually hands); muscle cramps; headache; insomnia; behavioural disturbances in children; hypersensitivity reactions including paradoxical bronchospasm; urticaria and angioedema; slight pain on intramuscular injection.

Terbutaline

EDL-D 497,498 Secondary hospitals

AVAILABILITY
TABLETS 2.5 and 5 mg; INJECTION 1 ml ampoule (0.5 mg/ml), NEBULISING SOLUTION 10 mg/ml, METERED DOSE INHALER (MDI) 250 μg/puff.
DOSE

Oral: Premature abortion: 2.5 to 5 mg thrice daily. Acute bronchospasm: Adult- 2.5 to 5 mg thrice daily. Subcutaneous, intramuscular or intravenous injection Uncomplicated premature labour: Adult- 5 μg/min for 20 min, increased every 20 min in steps of 2.5 μg/min until contractions have ceased; continue for 1 h then decreased every 20 min in steps of 205 μg/min to lowest dose that maintain suppression, max. dose 20 μg/min. Severe bronchospasm: Adult- 250-500 μg, 4 times daily. Child: >2 years- 10 μg/kg, max. dose- 300 μg. Inhalation: Acute bronchospasm: Adult/Child-MDI- 250 or 500 μg every 4-6 h, max. dose- 2000 μg/24 h; As nebuliser- 5-10 mg inhaled 2-4 times. Child- As nebuliser- 2-5 mg inhaled 2-4 times.

INDICATION

Bronchial spasm in bronchial asthma and chronic bronchitis; emphysema; premature labour; lymphoma.

CONTRAINDICATION

Cardiac disease; antepartum haemorrhage; intrauterine infection; intrauterine fetal death; placenta praevia; abruptio placenta; threatened miscarriage; cord compression; and eclampsia or severe pre-eclampsia; thyrotoxicosis; toxaeemia.

PRECAUTION

Suspected cardiovascular disease (such patients should be assessed by a cardiologist before initiating therapy), hypertension, mild to moderate pre-eclampsia, hyperthyroidism, and hypokalaemia (particular risk with potassium-depleting diuretics). It is important to monitor pulse rate (should not exceed 140 beats per min) and the patient’s fluid and electrolyte status (avoid overhydration- discontinue drug immediately and initiate diuretic therapy if pulmonary oedema occurs). It should also be used with caution in diabetes-monitor blood glucose (risk of hyperglycaemia and ketoacidosis, especially with intravenous β2 agonist); pregnancy (Appendix 7c).

ADVERSE EFFECTS

Nausea, vomiting; pulmonary oedema; palpitation; tachycardia, arrhythmias, peripheral vasodilation; headache, tremor, hyperglycaemia, hypokalaemia, muscle cramps and tension and hypersensitivity reactions (including angioedema, urticaria, rash, bronchospasm, hypotension, and collapse).

Theophylline

**EDL-D 212 PHC**

**INDICATION**

Acute asthma, long term control and prevention of symptoms, COPD.

**AVAILABILITY**

Tablets 200mg, 300mg, 400mg, 600mg Capsule 100mg, 200mg, 250mg Injection 2mL ampoule Syrup 20mg/mL

**DOSE**

Etohylline 169.4 mg/mL IV dose 2 mL 8hrly.

Oral dose: 80 — 240 mg tid.

Children: 24 mg/kg/bw in divided doses.

Controlled release preparation: 400 — 600 mg o.d.

**CONTRAINICATION**

Hypersensitivity, neonates, lactation.

**PRECAUTION**

Hypertension, myocardial infarction, hyperthyroidism, pregnancy, lactation, hepatic disease and acid peptic disease,
ADVERSE EFFECT
Nausea, vomiting, gastric disturbances, headache, gastric reflux, diuresis, cardiac arrhythmias, epilepsy.

DRUG INTERACTION
Metabolism is enhanced by rifampicin, phenobarbitone and alcohol, while it is reduced by ciprofloxacin, cimetidine, erythromycin and allopurinol.

Ipratropium bromide
EDL-D 670 Secondary hospitals

AVAILABILITY
METERED DOSE INHALER 200 doses (200 μg per actuation); CAPSULE 40 mg.

DOSE
Aerosol inhalation Adult- Metered dose inhaler; 20 to 40 μg, in early treatment up to 80 μg at a time, 3 to 4 times daily. Child- Metered dose inhaler; up to 6 years; 20 μg 3 times daily. 6 to 12 years; 20 to 40 μg 3 times daily.

INDICATION
Chronic asthma; chronic obstructive pulmonary disease; bronchospasm; rhinorrhoea, rapid reversal of sinus rhythm.

CONTRAINDICATION
Glaucoma; hypersensitivity; bladder obstruction; urinary retention.

PRECAUTION
Prostatic hypertrophy; pregnancy; glaucoma (standard doses unlikely to be harmful; reported with nebulized drug; particularly in association with nebulized salbutamol); lactation; allergy to atropine or Atropa belladona leaves.

ADVERSE EFFECTS
Occasionally dry mouth; constipation; angina; tremors; palpitation; nasal congestion.

Formoterol + Fluticasone propionate
EDL-D 643 Secondary hospitals

INDICATIONS
Asthma, severe chronic obstructive pulmonary disease (COPD).

AVAILABILITY
Inhalation Aerosol-
Formoterol + Fluticasone Propionate
6 μg + 125 μg
6 μg + 250 μg

DOSE
Inhalation
Asthma: Adults- 1-2 inhalations twice daily.
Child- 1 rotacap twice daily.
(Rotacaps to be used with a rotahaler device only. Do not swallow the capsules).
COPD: Adults- 2 inhalations twice daily.
Not recommended for children below 4 years of age.

CONTRAINDICATIONS
Hypersensitivity, acute asthma symptoms.

PRECAUTIONS
Severe cardiovascular disorders, cardiac rhythm abnormalities, seizure disorder, diabetes, thyrotoxicosis, hypokalemia, pulmonary tuberculosis, pregnancy (Appendix 7c), lactation, interactions (Appendix 6c).
ADVERSE EFFECTS
Headache, pharyngitis, throat irritation, upper respiratory tract infections, pneumonia, bronchitis, oral candidiasis, nausea, vomiting, diarrhea, chest pain, musculoskeletal pain, back pain, allergic reactions, wheezing, cough, skin rash, tremors, paradoxical bronchospasm, insomnia, adrenal suppression.

STORAGE
Store protected from light and moisture at a temperature not exceeding 30°C.

Montelukast
EDL-D 693,694 Secondary hospitals

INDICATIONS
Prophylaxis of mild to moderate asthma.

AVAILABILITY
TABLETS 5 and 10 mg.

DOSE
Oral
Adult- 10 mg once a day.
Child- 2-5yrs: 4 mg once daily; 6-14 yrs: 5 mg once daily; ≥ 15 yrs: 10 mg once daily.

CONTRAINDICATIONS
Hypersensitivity.

PRECAUTIONS
History of liver disease, pregnancy (Appendix 7c).

ADVERSE EFFECTS
Headache; rashes; eosinophilia; neuropathy; Churg-strauss syndrome.

STORAGE
Store protected from light and moisture.

Anti tussive and expectorant
Cough acts as protective reflex. It is helpful in the expulsion of respiratory secretion and other foreign particles from respiratory tract. Cough is of non-productive and productive type. Non-productive cough should be suppressed, whereas productive cough should not be suppressed. Cough suppressants are used only for the control of non-productive cough.

Bromhexine
EDL-D 543 PHC

INDICATION
Conditions where the sputum is viscid and tenaceous.

CONTRAINDICATION
1 Hypersensitivity.

PRECAUTIONS
Use with caution in patients with gastric ulceration.

ADVERSE EFFECT
Gastric irritation, allergic reactions, rhinorrhea, lacrimation.

AVAILABILITY
Tablet 8 mg Syrup 4 mg / 5 mL

DOSE
8-16 mg, tds-qds.
SECTION - 24
SOLUTIONS CORRECTING WATER, ELECTROLYTE AND ACID-BASE DISTURBANCES

Oral
Oral Rehydration Salts
EDL-D 386,387 Universal

INDICATIONS
Dehydration from acute diarrhoea.

AVAILABILITY
GLUCOSE SALT SOLUTION 5 and 37.5g.
Sodium chloride 2.6 g/litre of water
Sodium citrate 2.9 g/litre of water
Potassium chloride 1.5 g/litre of water
Glucose (anhydrous) 13.5 g/litre of water
When glucose and sodium citrate are not available, they may be replaced by
Sucrose (common sugar) 27 g/litre of water
Sodium bicarbonate 2.5 g/litre of water
In cases of cholera, oral rehydration salts containing a higher concentration of sodium may be required to prevent hyponatraemia.

Note: The solution may be prepared either from prepackaged sugar/salt mixtures or from bulk substances and water. Solutions must be freshly prepared, preferably with recently boiled and cooled water. Accurate weighing and thorough mixing and dissolution of ingredients in the correct volume of clean water is important. Administration of more concentrated solutions can result in hypernatraemia.

DOSE
Oral
5g (single use): dissolve in water and drink; 37.5g: to reconstitute it with 1 litre of clean water.
Adult- Fluid and electrolyte loss in acute diarrhoea; 200 to 400 ml solution after every loose motion.

PRECAUTIONS
Renal impairment.

ADVERSE EFFECTS
Vomiting- may indicate too rapid administration; hypernatraemia and hyperkalaemia may result from overdose in renal impairment or administration of too concentrated a solution.

STORAGE
Store protected from moisture in a sachet preferably made of aluminium foil containing sufficient powder for single dose or for a day treatment or for use in hospital.
Parentral
Solutions of electrolytes are given intravenously, to meet normal fluid and electrolyte requirements or to replenish substantial deficits or continuing losses, when the patient is nauseating or vomiting and is unable to take adequate amounts by mouth.

The nature and severity of the electrolyte imbalance must be assessed from the history and clinical and biochemical examination of each individual. Sodium, potassium, chloride, magnesium, phosphate, and water depletion can occur singly and in combination with or without disturbances of acid-base balance.

Isotonic solutions may be infused safely into a peripheral vein. More concentrated solutions, for example 20% glucose, are best given through an indwelling catheter positioned in a large vein.

Sodium chloride in isotonic solution provides the most important extracellular ions in near physiological concentrations and is indicated in sodium depletion which may arise from conditions such as gastroenteritis, diabetic ketoacidosis, ileus and ascites. In a severe deficit of from 4 to 8 litres, 2 to 3 litres of isotonic sodium chloride may be given over 2 to 3 h; thereafter infusion can usually be at a slower rate.

Excessive administration should be avoided; the jugular venous pressure should be assessed; the bases of the lungs should be examined for crepitations, and in elderly or seriously ill patients it is often helpful to monitor the right atrial (central) venous pressure.

Chronic hyponatraemia should ideally be managed by fluid restriction. However, if sodium chloride is required, the deficit should be corrected slowly to avoid risk of osmotic demyelination syndrome; the rise in plasma-sodium concentration should be limited to no more than 10 mmol/litre in 24 h.

The more physiologically appropriate compound solution of sodium lactate can be used instead of isotonic sodium chloride solution during surgery or in the initial management of the injured or wounded.

Sodium chloride and glucose solutions are indicated when there is combined water and sodium depletion. A 1:1 mixture of isotonic sodium chloride and 5% glucose allows some of the water (free of sodium) to enter body cells which suffer most from dehydration while the sodium salt with a volume of water determined by the normal plasma Na+ remains extracellular. Combined sodium, potassium, chloride, and water depletion may occur, for example, with severe diarrhoea or persistent vomiting; replacement is carried out with sodium chloride intravenous infusion 0.9% and glucose intravenous infusion 5% with potassium as appropriate.

Glucose solutions (5%) are mainly used to replace water deficits and should be given alone when there is no significant loss of electrolytes. Average water requirement in a healthy adult are 1.5 to 2.5 litres daily and this is needed to balance unavoidable losses of water through the skin and lungs and to provide sufficient for urinary excretion. Water depletion (dehydration) tends to occur when these losses are not matched by a comparable intake, as for example may
occur in coma or dysphagia or in the aged or apathetic who may not drink water in sufficient amount on their own initiative.

Excessive loss of water without loss of electrolytes is uncommon, occurring in fevers, hyperthyroidism, and in uncommon waterlosing renal states such as diabetes insipidus or hypercalcaemia. The volume of glucose solution needed to replace deficits varies with the severity of the disorder, but usually lies within the range of 2 to 6 litres.

Glucose solutions are also given in regimens with calcium, bicarbonate, and insulin for the emergency treatment of hyperkalaemia. They are also given, after correction of hyperglycaemia, during treatment of diabetic ketoacidosis, when they must be accompanied by continuing insulin infusion.

If glucose or sugar cannot be given orally to treat hypoglycaemia, glucose 50% may be given intravenously into a large vein through a large-gauge needle; this concentration is very irritant on extravasation and it is also viscous and difficult to administer. Larger volumes of less concentrated glucose solutions (10% or 20%) can be used as alternatives and are less irritant.

Sodium hydrogen carbonate (sodium bicarbonate) is used to control severe metabolic acidosis (as in renal failure). Since this condition is usually attended by sodium depletion, it is reasonable to correct this first by the administration of isotonic sodium chloride intravenous infusion, provided the kidneys are not primarily affected and the degree of acidosis is not so severe as to impair renal function. In these circumstances, isotonic sodium chloride alone is usually effective as it restores the ability of the kidneys to generate bicarbonate.

In renal acidosis or in severe metabolic acidosis of any origin, for example blood pH < 7.1, sodium hydrogen carbonate (1.4%) may be infused with isotonic sodium chloride when the acidosis remains unresponsive to correction of anoxia or fluid depletion; a total volume of up to 6 litres (4 litres of sodium chloride and 2 litres of sodium hydrogen carbonate) may be necessary in the adult. In severe shock due for example to cardiac arrest, metabolic acidosis may develop without sodium depletion; in these circumstances sodium hydrogen carbonate is best given in a small volume of hypertonic solution (for example 50 ml of 8.4% solution intravenously); plasma pH should be monitored. Sodium hydrogen carbonate is also used in the emergency management of hyperkalaemia.

Intravenous potassium chloride in sodium chloride infusion is the initial treatment for the correction of severe hypokalaemia when sufficient potassium cannot be taken by mouth. Potassium chloride concentrate may be added to sodium chloride 0.9% infusion, thoroughly mixed and given slowly over 2 to 3 h with specialist advice and ECG monitoring in difficult cases. Repeated measurements of plasma potassium are necessary to determine whether further infusions are required and to avoid the development of hyperkalaemia which is especially likely to occur in renal impairment.

Initial potassium replacement therapy should not involve glucose infusions because glucose may cause a further decrease in the plasma-potassium concentration.
**Glucose Non-EDL Universal**

**INDICATIONS**
Fluid replacement without significant electrolyte deficit; treatment of hypoglycaemia; varicose veins.

**AVAILABILITY**
Injection Infusion 250 ml, 500 ml and 1L (5% w/v); i.v.solution 10 and 20 ml (5% w/v).

**DOSE**
Intravenous infusion
Fluid replacement
Adult and Child- Determined on the basis of clinical and wherever possible, electrolyte monitoring.
Treatment of hypoglycaemia
Infusion of 50% glucose solution into a large vein.
Adult-25 ml.

**CONTRAINDICATIONS**
Anuria; thiamine deficiency; trauma; intracranial haemorrhage; haemodilution; acute ischaemic shock; hypophosphatemia; sepsis.

**PRECAUTIONS**
Diabetes mellitus (may require additional insulin); mannitol fluid balance.

**ADVERSE EFFECTS**
Glucose injections, especially if hypertonic, may have a low pH and cause venous irritation and thrombophlebitis; fluid and electrolyte disturbances; oedema or water intoxication (on prolonged administration or rapid infusion of large volumes of isotonic solutions); hyperglycaemia (on prolonged administration of hypertonic solutions); anaphylactoid reaction.

**STORAGE**
Store in a single dose container at a temperature not exceeding 30°C.

**Glucose + Sodium Chloride Non-EDL Secondary hospitals**

**INDICATIONS**
Fluid and extracellular volume depletion with excess diuresis; gastroenteritis.

**AVAILABILITY**
Injection 250, 450, 500 ml and 1 L.
(Dextrose 5% and sodium chloride 0.9%).

**DOSE**
Intravenous infusion
Adult and Child- Fluid replacement: determined on the basis of clinical and wherever possible, electrolyte monitoring.

**PRECAUTIONS**
Restrict intake in impaired renal function; cardiac failure, hypertension, peripheral and pulmonary oedema; toxaemia of pregnancy.

**ADVERSE EFFECTS**
Administration of large doses may give rise to oedema.

**STORAGE**
Store in a single dose container at a temperature not exceeding 30°C.
**Dextrose**

**EDL-D 158,159,160 PH**

**AVAILABILITY**
- INFUSION 10% dextran 40 + 5% dextrose or 0.9% sodium chloride

**DOSE**
- Intravenous To improve local circulation in peripheral vascular occlusion: Adult- 500-1000 ml (10-20 ml/kg) in first 24 hours; thereafter 500 ml every 1-2 days for up to 2 weeks. Thromboembolism prophylaxis: Adult- 500-1000 ml (10-20 ml/kg) on day of surgery, then 500 ml daily for 2-3 days, then 500 ml every second or third day, for up to 2 weeks. Shock: Adult- initially 500-1000 ml (10-20 ml/kg) infused as rapidly as needed; may follow with 500 ml (10 ml/kg) during the same 24 hour period; thereafter 500 ml (10 ml/kg) may be repeated daily for up to 5 days.

**INDICATION**
- Plasma volume expansion during hypovolemic shock when blood not available, Prophylaxis of thromboembolic disorders to improve local circulation in peripheral vascular occlusion

**CONTRAINDICATION**
- Hypersensitivity, cardiac decompensation, oliguria or anuria, hemostatic defects, thrombocytopenia, blood coagulation disorder, pulmonary oedema, neonates.

**PRECAUTION**
- Renal and hepatic impairment, pregnancy, lactaion, diabetes, cardiac patients, elderly, monitor urine output, monitor for signs of circulatory overload, interactions

**ADVERSE EFFECTS**
- Nausea, vomiting, local injection site reaction, hypersensitivity and anaphylactoid reactions, increased serum SGOT and SGPT concentrations, osmotic nephrosis.

**Dextrose with Sodium Chloride**

**EDL-D 161,162 Secondary hospital**

**AVAILABILITY**
- INFUSION 10% dextran 40 + 5% dextrose or 0.9% sodium chloride

**DOSE**
- Intravenous To improve local circulation in peripheral vascular occlusion: Adult- 500-1000 ml (10-20 ml/kg) in first 24 hours; thereafter 500 ml every 1-2 days for up to 2 weeks. Thromboembolism prophylaxis: Adult- 500-1000 ml (10-20 ml/kg) on day of surgery, then 500 ml daily for 2-3 days, then 500 ml every second or third day, for up to 2 weeks. Shock: Adult- initially 500-1000 ml (10-20 ml/kg) infused as rapidly as needed; may follow with 500 ml (10 ml/kg) during the same 24 hour period; thereafter 500 ml (10 ml/kg) may be repeated daily for up to 5 days.

**INDICATION**
- Plasma volume expansion during hypovolemic shock when blood not available, Prophylaxis of thromboembolic disorders to improve local circulation in peripheral vascular occlusion

**CONTRAINDICATION**
- Hypersensitivity, cardiac decompensation, oliguria or anuria, hemostatic defects, thrombocytopenia, blood coagulation disorder, pulmonary oedema, neonates.

**PRECAUTION**
- Renal and hepatic impairment, pregnancy, lactaion, diabetes, cardiac patients, elderly, monitor urine output, monitor for signs of circulatory overload, interactions
ADVERSE EFFECTS
Nausea, vomiting, local injection site reaction, hypersensitivity and anaphylactoid reactions, increased serum SGOT and SGPT concentrations, osmotic nephrosis.

**Sodium Chloride**
EDL-D 478,479 PHC

**AVAILABILITY**
Injection 250 and 500 ml (0.9% Solution); NASAL drop 5 5 ml (0.9% w/v).

**DOSE**
Intravenous infusion Adult and Child- Fluid and electrolyte replacement: determined on the basis of clinical and wherever possible, electrolyte monitoring.

**INDICATION**
Electrolyte and fluid replacement; hyponatremia; diabetic ketoacidosis; leg cramps; poisoning.

**CONTRAINDICATION**
Hypertension; liver cirrhosis; ischaemic heart disease; nephrotic syndrome; congestive heart failure.

**PRECAUTION**
Restrict intake in impaired renal function; cardiac failure, hypertension; peripheral and pulmonary oedema; toxemia of pregnancy; interactions (Appendix 6d).

**ADVERSE EFFECTS**
Administration of large doses may give rise to sodium accumulation and oedema; vomiting; intraocular coagulopathy.

**Potassium Chloride**
EDL-D 419 PHC

**INDICATIONS**
Electrolyte imbalance; hypokalaemia.

**AVAILABILITY**
INJECTION 10 ml ampoule (11.2%w/v).

**DOSE**
Slow Intravenous infusion
Adult and Child- Electrolyte imbalance; depending on the deficit or the daily maintenance requirements.

**CONTRAINDICATIONS**
Plasma-potassium concentrations above 5mmol/litre; chronic renal failure; systemic acidosis; acute dehydration; adrenal insufficiency.

**PRECAUTIONS**
For intravenous infusion the concentration of solution should not usually exceed 3.2g (43 mmol)/litre; specialist advice and ECG monitoring (see notes above); renal impairment; interactions (Appendix 6d); pregnancy (Appendix 7c); acute alkalosis, paediatric use.

**ADVERSE EFFECTS**
Cardiac toxicity on rapid infusion; nausea, vomiting, flatulence, diarrhoea.

**STORAGE**
Store in a single dose container at a temperature not exceeding 30°C.

**Sodium Bicarbonate**
EDL-D 475 Secondary Hospitals

**INDICATIONS**
Metabolic acidosis; cardiopulmonary resuscitation; hyperkalaemia; muscle spasm.

**AVAILABILITY**
INJECTION 10 ml ampoule (1.4%w/v),(8.4%w/v).
DOSE
Slow intravenous infusion
Adult and Child-Metabolic acidosis: a strong solution (up to 8.4%), an amount appropriate to the body base deficit.
Continuous intravenous infusion
Adult and Child- a weaker solution (up to 1.4%), an amount appropriate to the body base deficit.

CONTRAINDICATIONS
Metabolic or respiratory alkalosis, hypocalcaemia, hypochlorhydria; hypoventilation; hypoosmolarity.

PRECAUTIONS
Restrict intake in impaired renal function, cardiac failure, hypertension, peripheral and pulmonary oedema, toxaemia of pregnancy (Appendix 7c); monitor electrolytes and acidbase status; stomach disorder; allergies.

ADVERSE EFFECTS
Excessive administration may cause hypokalaemia and metabolic alkalosis, especially in renal impairment; large doses may give rise to sodium accumulation and oedema seizures; lactic acidosis; pulmonary oedema; hyperventilation.

STORAGE
Store in a single dose container at a temperature not exceeding 30°C.

Sodium Lactate, Compound Solution (Ringer Lactate Solution)
EDL-D 481 Universal

AVAILABILITY
INJECTIONS 250, 500 ml and 1L (1.87% w/v).

DOSE
Intravenous infusion Adult and Child-Fluid and electrolyte replacement or hypovolaemic shock: determined on the basis of clinical and wherever possible, electrolyte monitoring. Common adult dose is 1 to 3 litre/day.

INDICATION
Perioperative fluid and electrolyte replacement; hypovolaemic shock; metabolic acidosis; peritoneal dialysis.

CONTRAINDICATION
Metabolic or respiratory alkalosis; hypocalcaemia or hypochlorhydria; hypernatremia

ADVERSE EFFECTS
Excessive administration may cause metabolic alkalosis; administration of large doses may give rise to oedema; tissue necrosis; hypernatremia; hypervolemia; reaction at injection site.

Calcium gluconate
EDL-D 84 Secondary hospitals

AVAILABILITY
Tablets 250 and 500 mg; Injection 10 ml (1g/10 ml).

DOSE
Slow intravenous injection and continuous intravenous infusion Adult- Hypocalcaemic tetany: 1g (2.2 mmol) by slow intravenous injection, followed by continuous intravenous infusion of about 4g (8.8 mmol) daily.

INDICATION
Hypocalcaemic tetany; cardiopulmonary bypass.

CONTRAINDICATION
Conditions associated with hypercalcaemia and hypercalciuria (for example some forms of malignant disease).
PRECAUTION
Monitor plasma calcium concentration; renal impairment; interactions; diarrhoea, parathyroid disease; stomach trouble. Adverse Effects Mild gas

ADVERSE EFFECTS
Conditions associated with hypercalcaemia and hypercalciuria (for example some forms of malignant disease).

Water for Injection
EDL-D 532,533,534 PHC

INDICATIONS
In preparations intended for parenteral administration and in other sterile preparations.

AVAILABILITY
Ampoule 1, 5 and 10 ml.

STORAGE
Store in a single dose container at a temperature not exceeding 30°C.
Vitamins:
Vitamins are used for the prevention and treatment of specific deficiency states or when the diet is known to be inadequate. It has often been suggested but never convincingly proved, that subclinical vitamin deficiencies cause much chronic ill-health and liability to infections. This has led to enormous consumption of vitamin preparations, which have no more than placebo value. Most vitamins are comparatively non-toxic but prolonged administration of high doses of retinol (vitamin A), ergocalciferol (vitamin D$_2$) and pyridoxine (vitamin B$_6$) may have severe adverse effects.

**Retinol** (vitamin A) is a fat-soluble substance stored in body organs, principally the liver. Periodic high-dose supplementation is intended to protect against vitamin A deficiency which is associated with ocular defects particularly xerophthalmia (including night blindness which may progress to severe eye lesions and blindness), and an increased susceptibility to infections, particularly measles and diarrhea. Universal vitamin A distribution involves the periodic administration of supplemental doses to all preschool-age children with priority given to age groups, 6 months to 3 years, or regions at greatest risk. All mothers in high-risk regions should also receive a high dose of vitamin A within 8 weeks of delivery. Since vitamin A is associated with a teratogenic effect it should be given in smaller doses (no more than 10,000 units/day) to women of child-bearing age. It is also used in the treatment of active xerophthalmia. Doses of vitamin A should be administered orally immediately upon diagnosis of xerophthalmia and thereafter patients with acute corneal lesions should be referred to a hospital on an emergency basis. In women of child-bearing age there is a need to balance the possible teratogenic effects of vitamin A should they be pregnant with the serious consequences of xerophthalmia. Where there are severe signs of xerophthalmia high dose treatment as for patients over 1 year should be given. When less severe symptoms are present (for example night blindness) a much lower dose is recommended. Vitamin A therapy should also be given during epidemics of measles to reduce complications.

**Vitamin B** is composed of widely differing substances which are, for convenience, classed as ‘vitamin B complex’. Thiamine (vitamin B$_1$) is used orally for deficiency due to inadequate dietary intake. Severe deficiency may result in ‘beri-beri’. Chronic dry ‘beri-beri’ is characterized by peripheral neuropathy, muscle wasting and weakness, and paralysis; wet ‘beriberi’ is characterized by cardiac failure and oedema. Wernicke- Korsakoff syndrome (demyelination of the CNS) may develop in severe deficiency. Thiamine is given by intravenous injection in doses of up to 300 mg daily (parenteral preparations may contain several B group vitamins) as initial treatment in severe deficiency states. Potentially severe allergic reactions may occur after parenteral administration. Facilities for resuscitation should be immediately available.

**Riboflavin** (vitamin B$_2$) deficiency may result from reduced dietary intake or reduced absorption due to liver disease, alcoholism, chronic infectio or probenecid therapy. It may also occur in association with other deficiency states such as pellagra. **Pyridoxine** (vitamin B$_6$) deficiency is rare as the vitamin is widely distributed in foods, but deficiency may occur during isoniazid therapy and is characterized by peripheral neuritis. High doses are given in some metabolic disorders, such as hyperoxaluria and it is also used in sideroblastic anaemia. **Nicotinic acid** inhibits the synthesis of cholesterol and triglyceride and is used in some
hyperlipidaemias. Nicotinic acid and **nicotinamide** are used to prevent and treat nicotinic acid deficiency (pellagra). Nicotinamide is generally preferred as it does not cause vasodilation. **Hydroxocobalamin** is the form of vitamin B₁₂ used to treat vitamin B₁₂ deficiency due to dietary deficiency or malabsorption. **Folic acid** is essential for the synthesis of DNA and certain proteins. Deficiency of folic acid or vitamin B₁₂ is associated with megaloblastic anaemia. Folic acid should not be used in undiagnosed megaloblastic anaemia unless vitamin B₁₂ is administered concurrently, otherwise neuropathy may be precipitated (see chapter 13.1). Supplementation with folic acid 500 μg daily is recommended for women of child-bearing potential in order to reduce the risk of serious neural tube defects in their offspring. **Ascorbic acid** (vitamin C) is used for the prevention and treatment of scurvy. Claims that ascorbic acid is of value in the treatment of common colds are unsubstantiated. The term **vitamin D** covers a range of compounds including ergocalciferol (vitamin D₂) and cholecalciferol (vitamin D₃). These two compounds are equipotent and either can be used to prevent and treat rickets. Simple deficiency of vitamin D occurs in those who have an inadequate dietary intake or who fail to produce enough cholecalciferol (vitamin D₃) in their skin from the precursor 7-dehydrocholesterol in response to ultraviolet light. **Vitamin K** is necessary for the production of blood clotting factors.

**Minerals:**

**Calcium gluconate:** Calcium supplements are usually only required where dietary calcium intake is deficient. This dietary requirement varies with age and is relatively greater in childhood, pregnancy and lactation due to an increased demand, and in old age, due to impaired absorption. In osteoporosis, a calcium intake which is double the recommended daily amount reduces the rate of bone loss. In hypocalcaemic tetany calcium gluconate must be given parenterally but plasma calcium must be monitored. Calcium gluconate is also used in cardiac resuscitation.

**Iodine** is among the body’s essential trace elements. The recommended intake of iodine is 150 μg daily (200 μg daily in pregnant and lactation women); in children the recommended intake of iodine is 50 μg daily for infants under 1 year, 90 μg daily for children aged 2-6 years, and 120 μg daily for children aged 7-12 years. Deficiency causes endemic goitre and results in endemic cretinism (characterized by deaf-mutism, intellectual deficit, spasticity and sometimes hypothyroidism), impaired mental function in children and adults and an increased incidence of still-births and perinatal and infant mortality. Iodine and iodides may suppress neonatal thyroid function and in general iodine compounds should be avoided in pregnancy. Where it is essential to prevent neonatal goitre and cretinism, iodine should not be withheld from pregnant women. Control of iodine deficiency largely depends upon salt iodization with potassium iodide or potassium iodate and through dietary diversification. In areas where iodine deficiency disorders are moderate to severe, iodized oil given either before or at any stage of pregnancy is found to be beneficial.

**Sodium fluoride:** Availability of adequate fluoride confers significant resistance to dental caries. It is now considered that the topical action of fluoride on enamel and plaque is more important than the systemic effect. Where the natural fluoride content of the drinking water is significantly less than 1 mg per litre, artificial fluoridation is the most economical method of supplementing fluoride intake. Daily administration of fluoride tablets or drops is a suitable alternative, but systemic fluoride supplements should not be prescribed without reference to
the fluoride content of the local water supply; they are not advisable when the water contains more than 700 μg per litre. In addition, infants need not receive fluoride supplements until the age of 6 months. Dentifrices which incorporate Sodium fluoride are a convenient source of fluoride. Individuals who are either particularly caries prone or medically compromised may be given additional protection by the use of fluoride rinses or by application of fluoride gels. Rinses may be used daily or weekly; daily use of a less concentrated rinse is more effective than weekly use of a more concentrated one. High-strength gels must be applied on a regular basis under professional supervision; extreme caution is necessary to prevent the child from swallowing any excess.

**Ascorbic Acid (Vit C)**

**EDL-D 51 PHC**

**AVAILABILITY**
Tablets 100 and 500 mg; Drop 100 mg/ml; INJECTION 5 ml ampoule (100 mg/ml)

**DOSE**
Oral Adult and child- Prophylaxis of scurvy: 25 to 75 mg daily. Treatment of scurvy: 0.5 to 1.5g/day.

**INDICATION**
Prevention and treatment of scurvy.

**CONTRAINDICATION**
Hyperoxaluria.

**PRECAUTION**
Acetylsalicylic acid hypersensitivity; G-6-PD deficiency; large doses may cause renal calcium oxalate calculi; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
Gastrointestinal disturbances reported with large doses; failure of conception; kidney oxalate stones.

**Folic Acid**

**EDL-D 232,233 Secondary hospitals**

**AVAILABILITY**
TABLETS 1, 5 and 10 mg.

**DOSE**
Oral Adult- Treatment of folate-deficiency, megaloblastic anaemia: 5 mg daily for 4 months (up to 15 mg daily may be necessary in malabsorption states). Prevention of first occurrence of neural tube defect: 400 to 500 μg daily before conception and during the first twelve weeks of pregnancy. Prevention of recurrence of neural tube defect: 5 mg daily (reduced to 4 mg daily, if suitable preparation available) from at least 4 weeks before conception until twelfth week of pregnancy.

**INDICATION**
Treatment of folate-deficiency megaloblastic anaemia; prevention of neural tube defect in pregnancy.

**CONTRAINDICATION**
Should never be given without vitamin B12 in undiagnosed megaloblastic anaemia or other vitamin B12 deficiency states because risk of precipitating subacute combined degeneration of the spinal cord; folate-dependent malignant disease.

**PRECAUTION**
Women receiving antiepileptic therapy need counselling before starting folic acid; pernicious anaemia; folate dependent tumor; interactions (Appendix 6c); pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
Neuropathy; bronchospasm; skin eruption; anorexia; skin rash; status epilepticus.
Vitamin A Palmitate
EDL-D 524, 525 PHC

AVAILABILITY
Tablets 5000 and 10,000 Iu; Injection 50,000 Iu/ml.

DOSE
Oral
Adult- Prevention of vitamin A deficiency: 2,00,000 units every 6 months; pregnant woman, max. of 10,000 units daily or max. 25,000 units weekly; mothers, 200,000 units at delivery or within 6 weeks. Treatment of xerophthalmia; (except woman of childbearing age) 2,00,000 units on diagnosis, repeated next day and then after 2 weeks; (woman of child-bearing age), 5000 to 10,000 units daily for at least 4 weeks or up to 25000 units weekly. Child- Prevention of vitamin A deficiency: infant under 6 months, 50,000 units; 6 to 12 months, 100,000 units every 4 to 6 months, preferably at measles vaccination; over 1 year, 200,000 units every 4 to 6 months. Treatment of xerophthalmia; infant under 6 months, 50,000 units on diagnosis, repeated next day and then after 2 weeks; 6 to 12 months, 1,00,000 units immediately on diagnosis, repeated next day and then after 2 weeks; over 1 year, same as adults.

INDICATION
Prevention and treatment of vitamin A deficiency; prevention of complications of measles.

CONTRAINDICATION
Hypervitaminosis

PRECAUTION
Pregnancy (Appendix 7c), lactation.

ADVERSE EFFECTS
No serious or irreversible adverse effects in recommended doses; high intake may cause birth defects; transient increased intracranial pressure in adults or a tense and bulging fontanelle in infants (with high dosage); massive overdose can cause rough skin, dry hair, enlarged liver, raised erythrocyte sedimentation rate, raised serum calcium and raised serum alkaline phosphatase concentrations; hair loss; redness of skin; anorexia; weight loss.

Riboflavin (Vit B2)
EDL-D 455 Universal

AVAILABILITY
Tablets 5 mg

DOSE
Oral Adult and child- Treatment of vitamin B2 deficiency: up to 30 mg daily in divided doses. Prophylaxis of vitamin B2 deficiency: 1 to 2 mg daily.

INDICATION
Vitamin B2 deficiency; arabinoflavinosis.

CONTRAINDICATION
Cataract; hypersensitivity.

PRECAUTION
Large doses result in dark yellow discolouration of urine; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Swelling of lips, face and tongue and difficulty in breathing.

Vitamin B12 (Cyanocobalamin)
EDL-D 526 Secondary hospitals

AVAILABILITY
TABLETS 50, 500 and 1500 μg; CAPSULES 50 μg; LIQUID 35 μg/5 ml; INJECTION vial 500 μg/30 ml.
DOSE
Oral Adult- Vitamin-B12 deficiency of dietary origin: 50 to 150 μg daily between meals. Child- 50 to 105 μg daily in 1 to 3 divided doses. Intramuscular injection Initially 1 mg repeated 10 times at intervals of 2 to 3 days, maintenance 1 mg every month.

INDICATION
Cyanocobalamin deficiency; peripheral neuropathy; diabetic neuropathy; medicine related or alcoholic neuropathy

CONTRAINDICATION
Hypersensitivity, tobacco amblyopia.

PRECAUTION
Cobalt hypersensitivity, pregnancy (Appendix 7c).

ADVERSE EFFECTS
Asthenia; dyspepsia; pulmonary edema; shivering; bronchospasm.

Iodine
EDL-D 280 Secondary hospitals

INDICATIONS
Prevention and treatment of iodine deficiency; thyrotoxicosis; hyperthyroidism.

AVAILABILITY
Crystals Bulk.

DOSE
Oral
Adult- Endemic moderate to severe iodine deficiency: during pregnancy and one year postpartum, 300 to 480 mg once a year or 100 to 300 mg every 6 months; women of child-bearing age, 400 to 960 mg once a year or 200 to 480 mg every 6 months. Iodine deficiency; 400 mg, during pregnancy, single dose of 200 mg.
Child- Iodine deficiency: infant under 1 year, single dose 100 mg; 1 to 5 years, 200 mg once a year; above 6 years 400 mg once a year.

Intramuscular injection
Endemic moderate to severe iodine deficiency: women of child-bearing age, including any stage of pregnancy, 480 mg once each year; Iodine deficiency: 380 mg (if aged over 45 or with nodular goiter then 76 mg).
Child- Iodine deficiency; 380 mg but for infant up to 1 year, 190 mg.

CONTRAINDICATIONS
Lactation (Appendix 7b); bronchitis; goitre; hyperkalaemia; asthma; acne vulgaris; tuberculosis.

PRECAUTIONS
Over 45 years old or with nodular goiter (especially susceptible to hyperthyroidism when given iodine supplements-iodized oil may not be appropriate); may interfere with thyroid-function tests; pregnancy (see notes above and Appendix 7c); acute iodide toxicity; cardiac toxicity; interactions (Appendix 6c).

ADVERSE EFFECTS
Hypersensitivity reactions; goitre and hypothyroidism; hyperthyroidism; bronchitis; eosinophilia; rashes; headache; salivation.

Thiamine
EDL-D 502 Secondary hospitals

INDICATIONS
Prevention and treatment of vitamin B1 deficiency, acute alcohol intoxication.

AVAILABILITY
Tablets 25, 50 and 100 mg.
DOSE

Oral
Adult- Mild chronic thiamine deficiency: 10 to 25 mg daily.
Acute alcohol intoxication: 50-100 mg daily. Wernicke-Korsakoff syndrome: 50-100 mg daily.

PRECAUTIONS
Parenteral administration (see notes above); lactation (Appendix 7b); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Nausea; urticaria; gastrointestinal bleeding; oedema; pruritus; dizziness; anorexia.

STORAGE
Store protected from light and moisture in a non-metallic container.

Pyridoxine

EDL-D 722 PHC

INDICATIONS
Treatment of pyridoxine deficiency due to metabolic disorders; isoniazid neuropathy; sideroblastic anaemia.

AVAILABILITY
Tablets 10, 25, 40, 50 and 100 mg.

DOSE
Oral
Adult- Deficiency states: 25 to 50 mg up to 3 times daily. Isoniazid neuropathy, prophylaxis: 10 mg daily. Isoniazid neuropathy, treatment: 50 mg, 3 times daily. Sideroblastic anaemia: 100 to 400 mg daily in divided doses.

PRECAUTIONS
Interactions (Appendix 6c), pregnancy (Appendix 7c).

ADVERSE EFFECTS
Generally well tolerated, but chronic administration of high doses may cause peripheral neuropathies; paresthesia; neurotoxicity; muscular weakness.

Ergocalciferol (Vitamin D2)
Non-EDL Tertiary

INDICATIONS
Prevention of vitamin D deficiency; vitamin D deficiency caused by malabsorption or chronic liver disease; hypocalcaemia of hypoparathyroidism; osteomalacia; osteoporosis.

AVAILABILITY
Capsules 0.25 and 1 mg (50,000 IU).

DOSE
Oral
Adult and child- Prevention of vitamin D deficiency: 10 μg (400 units) daily.

CONTRAINDICATIONS
Hypercalcaemia; metastatic calcification.

PRECAUTIONS
Ensure correct dose in infants; monitor plasma calcium at weekly intervals in patients receiving high doses or those with renal impairment; nausea and vomiting may indicate overdose and hypercalcaemia; lactation (Appendix 7b); interactions (Appendix 6a); pregnancy (Appendix 7c).

ADVERSE EFFECTS
Symptoms of overdosage include anorexia; lasitude; nausea and vomiting, diarrhea; weight loss; polyuria; sweating; headache; thirst, vertigo and raised concentrations of calcium and phosphate in plasma and urine; tissue calcification may occur if dose of 1.25 mg continued for several months; cardiac arrhythmia; hypervitaminosis D; over psychosis; paralytic ileus.

STORAGE
Store protected from light in a hermetically sealed container.
SECTION - 26
EAR, NOSE AND THROAT PREPARATIONS

Xylometazoline

**EDL-D 535 Tertiary restricted**

**AVAILABILITY**
- Drops 10 and 15 ml (0.05 to 0.1 %w/v).

**DOSE**
- Instill 3 to 4 drops every 3 to 4 h or as required.

**INDICATION**
- Nasal congestion, conjunctival decongestant

**CONTRAINDICATION**
- Narrow angle glaucoma, atrophic rhinitis, vasomotor rhinitis.

**PRECAUTION**
- Avoid excessive or prolonged use; caution in infants under 3 months (no good evidence of value-if irritation occurs, might narrow nasal passage); infants and cardiac patients; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
- Local irritation, nausea, headache; after excessive use tolerance with diminished effect, rebound congestion; cardiovascular effects also reported; dryness of eye and nose, rhinitis medicamentose.

Tetracaine Hydrochloride

**EDL-D 500 Tertiary**

**AVAILABILITY**
- DROPS (0.5 and 1% w/v).

**DOSE**
- Instillation into the eye Instill 2 to 3 drops every 15 to 20 min till the desired effect is achieved.

**INDICATION**
- Short-acting local anaesthesia of cornea and conjunctiva.

**CONTRAINDICATION**
- Hypersensitivity to ester-type local anaesthetics; eye inflammation or infection

**PRECAUTION**
- Avoid prolonged use (cause of severe keratitis, permanent corneal opacification, scarring, delayed corneal healing); protect eye from dust and bacterial contamination until sensation fully restored; not to be applied on highly vascular surface; pregnancy (Appendix 7c).

**ADVERSE EFFECTS**
- Burning, stinging, redness; rarely, allergic reactions may occur; twitching; nystagmus; numbness of tongue; convulsions.

Cinnarizine

**EDL-D 121 Tertiary**

**AVAILABILITY**
- TABLETS 25 & 75 mg Plain and 75 mg SR.

**DOSE**
- Oral Motion sickness Adult: 30 mg 2 hr before travel and 15 mg every 8 hr during travel if needed. Vertigo Adult: 30 mg thrice daily. Child: 5-12 year: half of adult dose. Peripheral circulatory disorders Adult: 75 mg tablets three times daily.
INDICATION
Motion sickness, nausea, vomiting, vertigo and tinnitus associated with Meniere disease and other middle ear disorders, as a nootropic drug, adjunct therapy for symptoms of peripheral arterial disease.

CONTRAINDICATION
Hypersensitivity, Parkinson’s disease, children below 5 years.

PRECAUTION
Hypotension, patients should not drive or operate machinery, pregnancy (Appendix 7c), lactation, elderly, children and neonates, interactions (Appendix 6c).

ADVERSE EFFECTS
Drowsiness, rarely skin and hypersensitivity reactions, dry mouth, extrapyramidal symptoms sometimes associated with severe depression, muscular weakness, headache, euphoria, GI upsets, blurred vision, urinary difficulty or retention, constipation and increased gastric reflux, fatigue, hypolipidaemic effect.

Ciprofloxacin
EDL-D 125,126 Secondary hospitals

INDICATIONS
Bacterial infections of eye.

AVAILABILITY
tablets 250, 500 and 750 mg; injection 100 ml infusion (20 mg/10 ml); Ointment 5g (0.3% w/w);
Drops 5 and 10 ml (0.3% w/v).

DOSE
Adult and child above 12 years- Instill 2 to 3 drops in affected eye 3 to 4 times daily to start with thereafter reduce slowly as infection subsides. Apply about 0.5 cm ribbon of ointment in lower conjunctival sac for 3 to 4 times daily. Reduce as infection subsides.

CONTRAINDICATIONS
Epilepsy and hypersensitivity to quinolones.

PRECAUTIONS
It should be used with caution in patients with a history of epilepsy or conditions that predispose to seizures; in G-6-PD deficiency; myasthenia gravis (risk of exacerbation); in renal impairment (Appendix 7d); pregnancy (Appendix 7c), during lactation (Appendix 7b), and in children or adolescents. Exposure to excessive sunlight should be avoided (discontinue if photosensitivity occurs). Avoid excessive alkalinity of urine and ensure adequate fluid intake (risk of crystalluria); interactions (Appendix 6c); paediatric use.

ADVERSE EFFECTS
Nausea, vomiting, dyspepsia, abdominal pain, diarrhoea (rarely, antibiotic-associated colitis); headache; dizziness; sleep disorders; rash (rarely, Stevens-Johnson syndrome and toxic epidermal necrolysis) and pruritus. Less frequent side-effects include anorexia, increase in blood urea and creatinine; drowsiness, restlessness, asthenia, depression, confusion, hallucinations, convulsions, tremor, paraesthesia, hypoaesthesia; photosensitivity, hypersensitivity reactions including fever, urticaria, angioedema, arthralgia, myalgia, and anaphylaxis; blood disorders (including eosinophilia, leucopenia, thrombocytopenia); disturbances in vision, taste, hearing and smell. Other side-effects that have been reported include haemolytic anaemia, renal failure, interstitial nephritis, and hepatic dysfunction (including hepatitis and cholestatic jaundice). The drug should be discontinued if psychiatric, neurological or hypersensitivity reactions (including severe rash) occur. Ophthalmic solution:local burning, discomfort, corneal ulcers, lid oedema, corneal infiltration. Ointment: discomfort, keratopathy, blurred vision, corneal staining, epitheliopathy, photophobia.
STORAGE
Ointment: Store protected from light at a temperature not exceeding 30°C. Drops: Store protected from light.

Gentamicin
EDL-D 244 PHC

INDICATIONS
Blepharitis; bacterial conjunctivitis; keratitis, corneal ulcers.

AVAILABILITY
Ointment (1% w/w); Drops 5 ml (0.3% w/v).

DOSE
Instillation into the eye
Adult- Mild to moderate infections: 1 drop every 2 h, reducing frequency as infection is controlled, then continue for 48 h after healing is complete.

CONTRAINDICATIONS
Hypersensitivity to aminoglycoside group of antibiotics.

PRECAUTIONS
Prolonged use may lead to skin sensitization and emergence of resistant organisms including fungi; discontinue if purulent discharge, inflammation or exacerbation of pain; ophthalmic ointment may retard corneal healing, renal impairment (Appendix 7d), interactions (Appendix 6c), pregnancy (Appendix 7c).

ADVERSE EFFECTS
Burning; stinging; itching; dermatitis; conjunctival epithelial defects; conjunctival hyperemia; thrombocytopenic purpura; hallucination.

Neomycin with hydrocortisone
EDL-D 700 PHC

AVAILABILITY
CREAM 5, 10 and 15g (Aluminium tubes).

DOSE
Adult and child- Bacterial skin infections over 2 years: apply as a thin layer 3 times daily.

INDICATION
Superficial bacterial infections of the skin due to staphylococci and streptococci

CONTRAINDICATION
Avoid occlusive dressings; interactions

PRECAUTION
Avoid application to substantial areas of skin or to broken skin (risk of significant systemic absorption); overgrowth of resistant organisms on prolonged use; pregnancy (Appendix 7c).

ADVERSE EFFECTS
Local irritation; skin and fabrics stained brown
Appendix 1
Antimicrobial Resistance

Development and spread of antimicrobial resistance (AMR) is commonly due to overuse, misuse, and indiscriminate use of antimicrobials by doctors, nurses and pharmacists, noncompliance and self medication by patients and use in animal husbandry and agriculture. It is estimated that 70-80% of prescriptions for antimicrobials are probably advised unnecessarily by the health professionals. In spite of the fact that most common colds and diarrhoeal episodes are viral in origin, yet, antimicrobials are used indiscriminately. Reasons for over prescribing are often lack of confidence, peer pressure, patient pressure and pharmaceutical company pressure. Antimicrobial use is a key driver of the resistance. Poverty and inadequate access to antibiotics constitute a major factor in the development of resistance. Another common cause of developing resistance is improper diagnosis. In many instances death of an adequately equipped diagnostic laboratory in the vicinity compels the physician to prescribe antibiotics empirically, thus, increasing the likelihood of the patient receiving a wrong antibiotic. Furthermore, ready availability of antibiotics over-the-counter and sales promotion schemes by the pharmaceutical manufacturers also leads to the promotion of indiscriminate use, thus, increasing the likelihood of development of resistance. Counterfeit drugs are also a problem contributing to development of resistance. These contain either the wrong ingredient, or lesser amount of the active ingredient. In some instances, the medication poisons are capable of causing disability or even death. The impact of the media has also contributed to the development of resistance. Patients often demand antibiotics for their ailment on the basis of advertisements read or seen. Unwitting use of more active drugs at sub therapeutic doses leads directly to the development of multi drug resistance. Irrational use of antimicrobials is widespread throughout the world. This is harmful in terms of increased cost of therapy, unnecessary adverse drug reactions, therapeutic failure, reduced quality of care and worst of it is AMR.

The bacterial infections which contribute most to human mortality and morbidity are also those in which emerging antimicrobial resistance is most obvious: diarrhoeal diseases, respiratory infections, meningitis, sexually transmitted diseases, and hospital-acquired infections. Some important NFI-2011 626 examples include penicillin-resistant Streptococcus pneumoniae, vancomycin-resistant enterococci, methicillin-resistant Staphylococcus aureus, multi-resistant Salmonella typhi, Shigella dysenteriae, Neisseria gonorrhoea, Pseudomonas aeruginosa and multi-resistant Mycobacterium tuberculosis. The development of resistance to drugs commonly used to treat P. falciparum malaria is of particular concern, as is the emerging resistance to antiretroviral drugs.

Established mechanisms of AMR

For an antibiotic to be effective, it must reach the target site in an active form, bind to the target, and interfere with its function. Thus, bacterial resistance to an antimicrobial agent can occur due to three general mechanisms:
The drug does not reach its target
In Gram negative bacteria, many antibiotics enter the cell through protein channels called porins. Mutations or loss of these channels can prevent/slow the rate of antibiotic entry into a cell, effectively reducing drug concentration at the target site. If the drug target is intracellular and the drug requires active transport across the cell membrane, a mutation that interferes with the transport mechanism can confer resistance e.g. aminoglycosides. Bacteria can also transport antimicrobial drugs out of the cell through efflux pumps. Resistance to numerous drugs, including fluoroquinolones, macrolides, tetracyclines and beta lactam antibiotics, is mediated by this mechanism.

The drug is inactivated
Bacterial resistance to aminoglycosides can be due to a plasmid encoded aminoglycoside-modifying enzymes. Similarly, β-lactamase production is the most common mechanism of resistance to penicillins and other β-lactam drugs. Many hundreds of different β-lactamases have now been identified. A variation of this mechanism is failure of the bacterial cell to activate a prodrug e.g. loss of ability of M. tuberculosis to activate isoniazid (INH).

The target site is altered
This may be due to mutations in drug binding region of target enzyme e.g. fluoroquinolones, target modification e.g. ribosomal protection type of resistance to macrolides and acquisition of a resistant form of the susceptible target e.g., methicillin resistance in Staphylococcus Spp. due to production of a low-affinity penicillin-binding protein (PBP).

Strategies to prevent AMR in healthcare settings
Prudent antibiotic use: Antibiotics should be used only when they improve patient outcome. Not all infections need antibiotic treatment e.g. in patients with sore throat, benefit from antimicrobial therapy is small and is counterbalanced by the risk of adverse events like rash. Narrow spectrum agents should be used whenever possible. Broad spectrum agents should not be used as a cover for lack of diagnostic precision. Antibiotics should be prescribed in optimal doses, regimens, and should be stopped when the infection is treated. Restrict the use of last line antibiotics for serious infections and only when simpler agents are likely to be ineffective. Whenever used for prophylaxis, antibiotics should be used for short courses and at appropriate times (e.g. during surgical prophylaxis, antibiotics should be given within an hour prior to incision). Prevention of infection: Use of antimicrobials can also be reduced if infections are prevented in the first place. This can be achieved by improved use of vaccines and improved hygiene and infection control practices like compliance with hand washing protocols and aseptic techniques for catheterization. Catheters and drains should be removed when no longer needed. Clinicians should be familiar with local antibiotic sensitivity profiles and should comply with the local antibiotic guidelines. A hospital antibiotic policy should be formulated based on local antimicrobial resistance data. Prescribers should be educated about the use of antibiotics, when not to use them and also the infection control strategies. Hospitals should carry out surveillance of resistance patterns show much, where, in which organisms and to what antibiotics. Similarly antibiotic use pattern can be studied and these data can be used to devise targeted interventions to minimize antimicrobial use. The intent of giving this write up in the formulary is to encourage rational prescribing of antimicrobials and minimize the development of resistance to antimicrobials.
Milliequivalents
The milliequivalent is related to the total number of ionic charges in solution and it takes note of the valency of the ions. In other words, it is a unit of measurement of the amount of chemical activity of an electrolyte. The total concentration of cations always equals the total concentration of anions. A milliequivalent (mEq) weight is 1/1000 of the equivalent weight. An equivalent weight of an element is the atomic weight expressed in grams, divided by its valency

\[
1 \text{mEq} = \frac{\text{atomic weight in mg}}{\text{valency}}
\]

\[
e.g. 1 \text{mEq} \text{Na}^+ = \frac{23}{1} = 23 \text{ mg}
\]

\[
1 \text{mEq} \text{Cl}^- = \frac{35.5}{1} = 35.5 \text{ mg}
\]

Thus, a solution containing 1 mEq of Sodium per litre contains 23 mg Sodium. A solution containing 1 mEq of Sodium Chloride contains 23 mg Na+ and 35.5 mg Cl−, i.e. 58.5 mg Sodium Chloride per litre.

In a salt containing ions of different valencies,

\[
\text{Weight of a salt} = \frac{\text{Sum of the atomic weights (valency containing 1 mEq of specified ion x no. of specified ions in molecule)}}{\text{of specified ion}}
\]

e.g. Weight of magnesium chloride (mgCl2.6H2O) required to prepare a solution containing 1 mEq of magnesium per litre.

\[
24.3 + (2 \times 35.5) + 6[(2 \times 1) + 16] = 101.7 \text{ mg mgCl2.6H2O}
\]

For the conversion of grams per 100 ml (percentage) of a solution to mEq/litre, the following formula may be used.
percentage strength x 10000

\[
\text{mEq per litre} = \frac{\text{mg of salt containing 1 mEq of specified ions}}{\text{mEq weight (g)}}
\]

e.g. Number of mEq of Na\(^+\) per litre contained in Sodium Chloride injection 0.9\% (1 mEq Sodium is contained in 58.5 mg Sodium Chloride)

\[
\text{mEq per litre of Na}^+ \text{ in Sodium Chloride 0.9\%} = \frac{0.9 \times 10,000}{58.5} = 154 \text{ approximately}
\]

\[
\text{mEq} = \frac{\text{Wt (g)}}{\text{mEq weight (g)}}
\]

e.g. mEq in 5g of Potassium Chloride (atomic weight of Potassium Chloride = 74.6, valency = 1)

\[
\text{mEq Wt} = \frac{74.6}{1000} = \frac{0.0746}{1} = 0.0746
\]

\[
\text{mEq} = \frac{5}{0.0746} = 67 \text{ approximately}
\]

To convert mg per 100 ml (mg\%) to mEq per litre, the following formula is used:

\[
\frac{\text{mg. per 100 ml.} \times 10 \times \text{valency}}{\text{atomic weight}} = \text{mEq. per litre}
\]

In the case of gases (i.e. carbon dioxide) the volume per cent of the gas is multiplied by 10 and divided by 22.4, the later figure being the volume in litres occupied by a molecular weight of a gas. The results are expressed in millimoles per litre.

\[
\frac{\text{vol. percent} \times 10}{22.4} = \text{mM. per litre}
\]

To convert mg per 100 ml (mg\%) to milliosmoles:

\[
\frac{\text{mg. per 100 ml} \times 10}{\text{molecular weight}} = \text{mOsm per litre}
\]
Table 1: Atomic weight, Valency, and Normal Ranges of the Common Electrolytes

<table>
<thead>
<tr>
<th>Electrolyte</th>
<th>Atomic Weight</th>
<th>Valency</th>
<th>mg. per 100 ml</th>
<th>mEq. per litre</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>23</td>
<td>1</td>
<td>310-34</td>
<td>135-147</td>
</tr>
<tr>
<td>NaCl</td>
<td>58.5</td>
<td>1</td>
<td>570-620</td>
<td>98-106</td>
</tr>
<tr>
<td>K</td>
<td>39</td>
<td>1</td>
<td>16-22</td>
<td>4.1-5.7</td>
</tr>
<tr>
<td>Ca (total)</td>
<td>40</td>
<td>1</td>
<td>9.1-11.5</td>
<td>2.1-2.6</td>
</tr>
<tr>
<td>Ca(^{2+}) (ionized)</td>
<td>40</td>
<td>1</td>
<td>4.25-5.25</td>
<td>2.1-2.6</td>
</tr>
<tr>
<td>mg</td>
<td>24</td>
<td>2</td>
<td>1.8-3.6</td>
<td>1.5-3.1</td>
</tr>
<tr>
<td>(HCO(_3))^−</td>
<td>Use formula for gases.</td>
<td>55-70</td>
<td>25-31</td>
<td></td>
</tr>
<tr>
<td>(mM. CO(_2)) Infants</td>
<td>(vol. percent) (mM/L)</td>
<td>45-60</td>
<td>20-2</td>
<td></td>
</tr>
<tr>
<td>Cl</td>
<td>35.5</td>
<td>1</td>
<td>350-375</td>
<td>98-106</td>
</tr>
<tr>
<td>NH(_4)Cl</td>
<td>Contains 66.28 per-cent of Cl.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SO(_4)^{2−}</td>
<td>96</td>
<td>2</td>
<td>0.5-2.5</td>
<td>0.3-1.5</td>
</tr>
<tr>
<td>Ion</td>
<td>weight of mEq in mg</td>
<td>Salt</td>
<td>mg of salt containing 1 mEq of the specified ion</td>
<td></td>
</tr>
<tr>
<td>-----------</td>
<td>---------------------</td>
<td>----------------------------------------------------------------------</td>
<td>-----------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Ca^{2+}</td>
<td>20.0</td>
<td>Calcium Chloride, CaCl\textsubscript{2},2H\textsubscript{2}O</td>
<td>147</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Calcium Gluconate, C\textsubscript{12}H\textsubscript{22}CaO\textsubscript{14},H\textsubscript{2}O</td>
<td>224</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Calcium Lactate, C\textsubscript{6}H\textsubscript{10}CaO\textsubscript{6},5H\textsubscript{2}O</td>
<td>154</td>
<td></td>
</tr>
<tr>
<td>K^{+}</td>
<td>39.1</td>
<td>Potassium Chloride, KCl</td>
<td>74.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Potassium Citrate, C\textsubscript{6}H\textsubscript{12}K\textsubscript{2}O\textsubscript{7},H\textsubscript{2}O</td>
<td>108</td>
<td></td>
</tr>
<tr>
<td>mg^{2+}</td>
<td>12.5</td>
<td>Magnesium Sulphate, mgSO\textsubscript{4},7H\textsubscript{2}O</td>
<td>123</td>
<td></td>
</tr>
<tr>
<td>Na^{+}</td>
<td>23.0</td>
<td>Sodium Acetate, C\textsubscript{2}H\textsubscript{3}NaO\textsubscript{2},3H\textsubscript{2}O</td>
<td>136</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium Acid Citrate, C\textsubscript{2}H\textsubscript{3}NaO\textsubscript{2},1\frac{1}{2}H\textsubscript{2}O</td>
<td>131</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium Bicarbonate, NaHCO\textsubscript{3}</td>
<td>84</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium Chloride, NaCl</td>
<td>58.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium Citrate, C\textsubscript{6}H\textsubscript{5}NaO\textsubscript{2},2H\textsubscript{2}O</td>
<td>98</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium Lactate*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cl</td>
<td>35.5</td>
<td>Calcium Chloride, CaCl\textsubscript{2},2H\textsubscript{2}O</td>
<td>73.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Potassium Chloride, KCl</td>
<td>74.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium Chloride, NaCl</td>
<td>58.5</td>
<td></td>
</tr>
<tr>
<td>C\textsubscript{2}H\textsubscript{3}O\textsubscript{2} (Acetate)</td>
<td>59.0</td>
<td>Sodium Acetate, C\textsubscript{2}H\textsubscript{3}NaO\textsubscript{2},3H\textsubscript{2}O</td>
<td>136</td>
<td></td>
</tr>
<tr>
<td>C\textsubscript{6}H\textsubscript{10}O\textsubscript{3} (Lactate)</td>
<td>89.0</td>
<td>Calcium Lactate, C\textsubscript{6}H\textsubscript{10}CaO\textsubscript{6},5H\textsubscript{2}O</td>
<td>308.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium Lactate*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HCO\textsubscript{3}^{-}</td>
<td>61.0</td>
<td>Sodium Bicarbonate, NaHCO\textsubscript{3}</td>
<td>84</td>
<td></td>
</tr>
</tbody>
</table>

*Prepared in solution by neutralising lactic acid with sodium hydroxide: 1.0 ml of 1 M sodium contains the equivalent of 112 mg.
### Appendix 3
Common Laboratory Parameters

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Biochemical Parameter</th>
<th>Reference Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Liver Function Tests</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Aspartate Aminotransferase (AST/SGOT)</td>
<td>12-38 Units/L</td>
</tr>
<tr>
<td></td>
<td>Alanine Aminotransferase (ALT/SGPT)</td>
<td>7-41 Units/L</td>
</tr>
<tr>
<td></td>
<td>Bilirubin</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>0.3-1.3 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Direct</td>
<td>0.1-0.4 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Indirect</td>
<td>0.2-0.9 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Protein</td>
<td>6.7-8.6 g/dL</td>
</tr>
<tr>
<td></td>
<td>Albumin</td>
<td>3.5-5.5 mg/dL</td>
</tr>
<tr>
<td></td>
<td>y Glutamyl Transpeptidase (GGT)</td>
<td>9-58 Units/L</td>
</tr>
<tr>
<td>2.</td>
<td>Kidney Function Tests</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Blood Urea Nitrogen (BUN)</td>
<td>7-20 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Creatinine (Cr) Uric Acid</td>
<td>0.6-1.2 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Male</td>
<td>3.1-7.0 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>2.5-5.6 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Glomerular Filtration Rate</td>
<td>&gt;60 ml/min/1.73 m2</td>
</tr>
<tr>
<td>3.</td>
<td>Lipid Profile</td>
<td></td>
</tr>
<tr>
<td></td>
<td>HDL Cholesterol</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Male</td>
<td>27-67 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>34-88 mg/dL</td>
</tr>
<tr>
<td></td>
<td>LDL Cholesterol</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Desirable</td>
<td>&lt;130 mg/dL</td>
</tr>
<tr>
<td></td>
<td>High Risk</td>
<td>160 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Triglycerides</td>
<td>&lt;165 mg/dL</td>
</tr>
<tr>
<td>4.</td>
<td>Pancreatic Function Tests</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Amylase</td>
<td>20-96 Units/L</td>
</tr>
<tr>
<td></td>
<td>Lipase</td>
<td>3-43 Units/L</td>
</tr>
<tr>
<td>5.</td>
<td>Thyroid Function Tests</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Thyroxine</td>
<td></td>
</tr>
</tbody>
</table>
Total (T<sub>T</sub>) 5.4-11.7 μg/dL
Triiodothyronine (T<sub>3</sub>) Total 77-135 ng/dL
Thyroid Stimulating Hormone (TSH) 0.34-4.25 μIU/ml

6. Parathyroid Function Tests
Calcium 8.5-10.5 mg/dL
Calcium (Ionized) 4.6-5.3 mg/dL

7. Reproductive Function
Tests
Follicle Stimulating Hormone (FSH)
Male 1.0-12.0 m IU/ml
Female
Menstruating
Follicular phase 3.0-20.0 m IU/ml
Ovulatatory phase 9.0-26.0 m IU/ml
Luteal phase 1.0-12.0 m IU/ml
Post menopausal 18.0-153.0 m IU/ml
Leuteinsing Hormone (LH)
Male 2.0-12.0 U/L
Female
Menstruating
Follicular phase 2.0-15.0 U/L
Ovulatatory phase 22.0-105.0 U/L
Luteal phase 0.6-19.0 U/L
Postmenopausal 16.0-64.0 U/L
Testosterone toal, morning sample
Male 270.0-1070.0 ng/dL
Female 6.0-86 ng/ml

8. Electrolytes
Sodium (Na<sub>+</sub>) 136-146 mEq/L
Potassium (K<sub>+</sub>) 3.5-5.0 mEq/L
Magnesium (mg<sub>2+</sub>) 1.5-2.3 mg/dL
9. **Arterial Blood Gases**

- Carbondioxide, Partial Pressure (PCO$_2$) 32-45 mm Hg
- Oxygen, Partial Pressure (PO$_2$) 72-104 mm Hg
- Osmolality Serum 275-295 mOsm/kg serum water
- pH blood (arterial) 7.35-7.45
- Anion Gap 7-16 mmol/L
- Bicarbonate 22-30 mEq/L

10. **Normal Haematological Parameters**

- Total Leucocyte Count (TLC) 4.8-10.8 x 10$^3$/mm$^3$
- Basophil Count 0-2%
- Eosinophil Count 0-6%
- Lymphocyte Count 20-50%
- Monocyte Count 4-8%
- Platelet Count 165-415 x 10$^3$/mm$^3$
- Reticulocyte Count Males 0.8-2.3% red cells
- Females 0.8-2.0% red cells
- Neutrophil Count 0.40-0.7 (40-70%)/mm$^3$
- Red Blood Cell Count Male 4.3-5.6 x 10$^6$/mm$^3$
- Female 4.0-5.2 x 10$^6$/mm$^3$
- Hemoglobin Male 13.3-16.2 g/dL
- Female 12.0-15.8 g/dL
- Erythrocyte Sedimentation Rate (ESR) Male 0-15 mm/h
- Female 0-20 mm/h

11. **Corpuscular Values of Erythrocytes**

- Mean Corpuscular Hemoglobin (MCH) 26.7-31.9 pg/cell
Mean Corpuscular Hemoglobin Concentration (MCHC) 32.3-35.9 g/dL
Mean Corpuscular Volume (MCV) 79-93.3 fL

12. Coagulogram
Fibrinogen 233-496 mg/dL
Partial Thromboplastin Time, Activated (aPTT) 26.3-39.4 seconds
Prothrombin Time (PT) 12.7-15.4 seconds
Thrombin Time 15.3-18.5 seconds
Bleeding Time (Adult) < 7.1 min
Coagulation factors 50-150% of normal

13. Cerebrospinal Fluid
Leukocytes - Total less than 5 cells/cu mm (all mononuclear)
Differential Lymphocytes 60–70%
Monocytes 30–50%
Neutrophils None
Chloride 116 – 122 mEq/L
Glucose 40-70 mg/dL
Pressure 50-180 mm of water
Total protein 15-50 mg/dL (lumbar)
Albumin 6.6-44.2 mg/dL

14. Miscellaneous
Glucose 75-110 mg/dL
Glycosylated Hemoglobin (HbA1C) 4.0-6.0%
Iron 41-141 μg/dL
Total Iron Binding Capacity (TIBC) 251-406 μg/dL
Methemoglobin (Met Hb) Ferritin <1% of total Hb
Male 29-248 ng/ml
Female 10-150 ng/ml
Transferrin 190-375 mg/dL
Lactate Dehydrogenase (LDH) 115-221 Units/L
Folic Acid (Red Cells) 165-760 ng/ml
Glucose-6-Phosphate Dehydrogenase (G-6-PD) 5-14 Units/g Hb
Vitamin B12 140-820 pg/ml
Appendix 4
Disposal of Unused/Expired Pharmaceutical Products

Shelf life of a drug is defined as the time interval within which it remains physically, chemically and/or biologically stable as well as safe and effective for human consumption if stored under the label specified conditions and it is in the original container closure system. Shelf life can be determined by the accelerated stability testing method. Expiry/expiration date is the actual date placed on the label/container indicating the time during which a batch of drug product is expected to remain with the approved shelf life specifications if stored under defined conditions and after which it should not be used. Expired medicines lose their potency and are capable of producing toxins, causing serious reaction or failure of therapy.

Thus disposal of unused/expired pharmaceutical products is required for every pharmacy - retail and wholesale, clinic, dispensary, hospital, manufacturing unit and testing laboratory. Indiscriminate disposal of drugs is likely to pollute the environment resulting in contamination of vegetables, fruits, fish and other aquatic life and even drinking water. Pharmaceuticals and Personal Care Products (PPCP) have been found present as pollutant in water and environment and this poses a serious issue of ecological imbalance due to indiscriminate disposal of expired pharmaceutical products.

With increasing awareness of pollution and their effect on human beings, animals and environment, it is imperative to assure Regulatory compliance by Individuals, Retail and Wholesale Chemists, Clinics, Hospitals, Manufacturers of PPCP, Clinical Research Organizations conducting Biostudies, Analytical Testing Laboratories and other organizations involved in drug distribution (in the event of any disaster) to augment Government efforts.

Managing Disposal of PPCP:

Mass awareness, at every level, of impact of casual approach in such disposal of unwanted and expired PPCP is very much desirable. Following steps are suggested for safe disposal of unused/not required/expired PPCP:

1. Expired or near expiry or unused/not required PPCP in large quantity should be returned by Retail Chemists and Druggists/Pharmacies/Clinics/Hospitals/NGO involved in drug distribution to Wholesalers or stockists of manufacturer(s) who in turn will return the same to the location of the manufacturer where the products were manufactured, for proper disposal.

2. Expired/short expiry PPCP are received at the segregated area for Expired Goods/Market Returns in the Finished Goods Warehouse of the manufacturing location with proper documentation in compliance with Regulatory requirements under i) Drugs and Cosmetics Rules 1945 and amendments therein, ii) Central Excise/State Excise Laws- as applicable and iii) Local authority of Pollution Control Board of the State before under taking disposal of PPCP. Short expiry PPCP should be analyzed and if found satisfactory, could be used for treatment ensuring that it is consumed before expiration.
3. In the event of expired goods returned from overseas customers, appropriate documentation for disposal of PPCP should be provided to the customs authority at the importing port for Bill of Entry.

4. Finished Goods Warehouse receiving the PPCP shall inform Quality System/Quality Assurance authority of the organization for verification and appropriate documentation before undertaking disposal.

5. For small quantity of expired products in Clinics, Hospitals, Health Care Centers and Dispensaries, Pharmacist at the location should be assigned responsibility for disposal of PPCP. He/she should be trained for proper documentation and disposals as indicated below.

6. Disposal of expired retention samples of API, Excipients and that of Pharmaceutical dosage forms in the manufacturing unit or Testing laboratories should be assigned to a responsible Quality control person under supervision of Quality System/Quality Assurance ensuring appropriate documentation.

7. After compliance to administrative control procedure, the expired PPCP should be transferred to a segregated area under the control of Safety, Health and Environment department in the manufacturing location for undertaking disposal.

Disposal Methods of Pharmaceutical and Personal Care Products

Sorting of Materials:

Materials to be disposed off should be segregated. Different methods are employed depending on

i) Type of dosage forms - Tablets, Capsules, Powders, Injectables, Creams, Ointments, Liquids, Ampoules, Vials, Intravenous Infusions etc.

ii) Chemical nature of drugs e.g. Antineoplastics/Anticancer, β-Lactams, Hormones, Steroids, Anti-infective, Narcotics, Antiseptics and Psychotropic substances etc. Tertiary (Printed/Labelled Corrugated Boxes) and Secondary (Printed Cartons/Paper box) packaging materials are removed and destroyed with the help of heavy duty paper shredder. The methods of disposal of various pharmaceutical dosage forms and that of specific category medicines are mentioned below.

Pharmaceutical Dosage Forms

1. Tablets/Capsules:

   Up to 50 tablets or capsules soak in about 100 ml of water and collect the same in a polyethylene bag containing used Tea/Coffee grind. Seal the bag and put in trash. Big quantity - Pulverize using heavy duty crusher. Collect in a poly bag and seal. Dispose it in high temperature incinerator (Temp. 850°C to 1200°C)/approved site for solid waste disposal by the Pollution Control Board of the State.

2. Injectables - ampoules/vials:

   Up to 50 Ampoules/Vials (up to 10 ml)-break ampoules/open vials and collect liquid in a polyethylene bag containing used Tea/Coffee grind. Seal the bag and put in trash. For bigger
3. In the event of expired goods returned from overseas customers, appropriate documentation for disposal of PPCP should be provided to the custom authority at the importing port for Bill of Entry.

4. Finished Goods Warehouse receiving the PPCP shall inform Quality System/Quality Assurance authority of the organization for verification and appropriate documentation before undertaking disposal.

5. For small quantity of expired products in Clinics, Hospitals, Health Care Centers and Dispensaries, Pharmacist at the location should be assigned responsibility for disposal of PPCP. He/she should be trained for proper documentation and disposals as indicated below.

6. Disposal of expired retention samples of API, Excipients and that of Pharmaceutical dosage forms in the manufacturing unit or Testing laboratories should be assigned to a responsible Quality control person under supervision of Quality System/Quality Assurance ensuring appropriate documentation.

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i) Type of dosage forms-
- Tablets, Capsules, Powders, Injectables, Creams, Ointments, Liquids, Ampoules, Vials, Intravenous Infusions etc.

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1. Tablets/Capsules:
   Up to 50 tablets or capsules soak in about 100 ml of water and collect the same in a polyethylene bag containing used Tea/Coffee grind. Seal the bag and put in trash. Big quantity - Pulverize using heavy duty crusher. Collect in a poly bag and seal. Dispose it in high temperature incinerator (Temp. 850⁰C to 1200⁰C)/approved site for solid waste disposal by the Pollution Control Board of the State.

2. Injectables - ampoules/vials:
   Up to 50 Ampoules/Vials (up to 10 ml)-break ampoules/open vials and collect liquid in a polyethylene bag containing used Tea/Coffee grind. Seal the bag and put in trash. For bigger quantity, use heavy duty crusher to separate liquid and dilute it with water and transfer it to Effluent Treatment Plant (ETP) of the manufacturing unit. Broken glass/vials (after removal of label), rubber stoppers and seals should be disposed off as scrap. Powder Injectables (in Vials/Ampoules) to be disposed off in an incinerator as indicated above.

3. Oral liquids and Intravenous fluids:
   Small quantity – Dilute the liquid with water and drain it. For bigger quantity, dilute collected liquid with water and transfer it to ETP of the manufacturing unit. Liquids with high solid contents to be disposed off in an incinerator as indicated above.

4. Semi solids:
   Small quantity, mix it with used Tea/Coffee grind in a polyethylene bag. Seal the bag and put in a trash. Deshape the containers/remove the label and discard the containers. Semisolids in bigger quantity to be disposed off in an incinerator mentioned earlier.

Containers - Tubes to be deshaped and remove the label from glass/plastic container before disposal as a scrap.

**Specific categories:**

1. **Anti-infectives-β-lactams:**
   Small quantity of all β-lactam antibiotics to be destroyed by soaking in 1N Sodium Hydroxide for 30 mins or 1% Hydroxylamine in Water for 10 mins and trash. Bigger quantity to be disposed off in an incinerator (Temp. 850⁰C to 1200⁰C) indicated above.

2. **Anti-infectives - others:**
   Tetracyclines- Small quantity to be soaked in 10% of Calcium Hydroxide/any other Calcium salt in Water for 30 mins and trash. Macrolides- (Erythromycin, Clarithromycin etc.)-Small quantity, soak in 1N Hydrochloric Acid and trash. Amino glycosides ( Gentamycin, Amikacin etc.) - Small quantity dilute with large volume of water and drain it. Bigger quantity of all the above anti-infective should be disposed of in an incinerator as mentioned above.

3. **Steroids:**
   Small quantity- Soak in 1N Sodium Hydroxide for 30 mins and trash. Bigger quantity- all dosage forms (taken out from the primary packing materials) to be incinerated at the temperature range indicated above.

4. **Hormones:**
   Small quantity- Aqueous solution to be exposed to UV for 20 minutes and trash. Estrogens- small quantity in aqueous solution should be exposed to ultrasound at 0.6 and 2 kw in a sonicator for 60 mins. and trash. Bigger quantity- all solid dosage forms (taken out from primary packaging materials) to be incinerated as indicated above.
5. **Disinfectants:**

   Small quantity- use it. Bigger quantity- Not more than 50L. Dilute with enough quantity of water to ensure dilution with loss of activity and drain it in ETP.

6. **Controlled substances:**

   Small quantity- Flush down the toilet to avoid misuse. Bigger quantity- All dosage forms (take out from primary packaging material) to be incinerated as mentioned above.

Disposal by incineration is preferred over chemical inactivation for all dosage forms/APIs.

**Cost of Disposal of Pharmaceutical Products:**

In India Solid Waste Disposal of PPCP at an approved solid waste disposal site by the local Pollution Control Board works out to Rs. 15 to 25 per kg. Disposal of PPCP by incineration requires about 5% of the fuel feed to that of the total quantity to be disposed of. Total cost of such disposal works out to about 0.5 % to 2% of the total sales.

There is an urgent need to minimize this wastage on account of disposal of Unused/Expired Pharmaceutical products. This not only involves huge cost but also danger of contaminating water bodies, rivers, sea, air, land and aquatic lives, which ultimately get recycled and comes back to human beings, animals and vegetation in one or the other form.

**Reuse of Expired Pharmaceutical Products:**

In the event of Emergency situation, Regulatory Agencies do allow Revalidation, after Analysis of Expired Pharmaceutical Drug Products, if found satisfactory, for Human/Animal consumption.

**Reduction in Level of Disposal:**

Reduced level of disposal of PPCP can help significantly. If recovery of api (Active Pharmaceutical Ingredient) and purification there of, out of near expiry/expired PPCP be attempted and regulated, it will reduce the level of PPCP disposal. This will be a challenging task for pharmaceutical professionals, which will help industry in reducing quantum of disposal.

**Education and Training:**

Continuing education and training at every level is desirable to generate awareness of hazards associated with indiscriminate disposal of unused/expired pharmaceutical products - an emerging environmental issue. It is expected that awareness of the stakeholders with systematic preventive and corrective actions in time will ensure the safe disposal of unused/expired pharmaceutical products.
### Appendix 5
#### Drugs and Poisons Information Centres in India

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Address</th>
<th>Contact No.</th>
<th>E-mail</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>National Poisons Information Centre, Department of Pharmacology, All India Institute of Medical Sciences (AllIMS) New Delhi-110029</td>
<td>011-26589391 011-26593677</td>
<td><a href="mailto:npicaiims@hotmail.com">npicaiims@hotmail.com</a>, <a href="mailto:npicaiims2010@gmail.com">npicaiims2010@gmail.com</a></td>
</tr>
<tr>
<td>2.</td>
<td>Medicine and Poison- Antidote Information Centre (MAPIC) Delhi Institute of Pharmaceutical Sciences and Research (DIPSAR) Pushp Vihar, Sector-III, M.B. Road, New Delhi- 110017</td>
<td>011-29553173 011-29554649</td>
<td><a href="mailto:mapicdipsar@indiatimes.com">mapicdipsar@indiatimes.com</a>, <a href="mailto:dipsarmapic@gmail.com">dipsarmapic@gmail.com</a></td>
</tr>
<tr>
<td>3.</td>
<td>Poison Information Centre, Toxicology and IMCU Unit, Govt. General Hospital, Chennai-600003</td>
<td>044-536 3208 044-536 3131 ext. 108 Fax: 044-538 8521</td>
<td><a href="mailto:thiruma@satyam.net.in">thiruma@satyam.net.in</a> <a href="mailto:ghpictn@vsnl.net">ghpictn@vsnl.net</a> <a href="http://www.chennaipic.com">www.chennaipic.com</a></td>
</tr>
<tr>
<td>4.</td>
<td>Sri Ramachandra Hospital, Porur, Chennai No.1, Ramachandra Nagar Porur, Chennai-600 116</td>
<td>044-24768403 ext. 8927 Fax: 044-2476 7008</td>
<td><a href="mailto:dicsrmc@yahoo.co.in">dicsrmc@yahoo.co.in</a></td>
</tr>
<tr>
<td>5.</td>
<td>Poisons Information Centre National Institute of Occupational Health, Meghani Nagar Ahmedabad-380 016</td>
<td>079 286 7351 079 562 1400 Fax: 079 286 6630</td>
<td><a href="mailto:dewan4@satyam.net.in">dewan4@satyam.net.in</a></td>
</tr>
<tr>
<td>6.</td>
<td>Poison Information and Laboratory Services (Department of Toxicology) Amrita Institute of Medical Sciences and Research Cochin-682 026</td>
<td>0484 400 8056 0484 400 1234 9895282388 Fax: 0484 280 2051 toxicology@</td>
<td>medical.amrita.edu; <a href="mailto:poisonunit@aimshospital.org">poisonunit@aimshospital.org</a></td>
</tr>
</tbody>
</table>
Mixing alcohol with medications can cause a variety of symptoms like nausea, vomiting, headache, drowsiness, fainting, or loss of coordination. By virtue of its effects on the CNS consumption of alcohol, even in small quantities, puts the patient at a high risk. There are medicines which should never be taken with alcohol (Table 1). However, there are many other medicines which should be used with high level of caution while the patient is on alcohol (Table 2).

### Table 1: Drugs not to be used with alcohol

<table>
<thead>
<tr>
<th>Drug</th>
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</tr>
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<tbody>
<tr>
<td>Acetylsalicylic Acid</td>
<td>Lovastatin</td>
</tr>
<tr>
<td>Alprazolam</td>
<td>Lovastatin + Niacin</td>
</tr>
<tr>
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</tr>
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<td>Metronidazole</td>
</tr>
<tr>
<td>Benazepril</td>
<td>Naproxen</td>
</tr>
<tr>
<td>Butalbital + Codeine</td>
<td>Nicotinic acid</td>
</tr>
<tr>
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<td>Nitrazepam</td>
</tr>
<tr>
<td>Cefoperazone</td>
<td>Nitrofurantoin</td>
</tr>
<tr>
<td>Chlordiazepoxide</td>
<td>Nitroglycerin</td>
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<td>Clomipramine</td>
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<td>Phenobarbital</td>
</tr>
<tr>
<td>Cyclobenzaprine</td>
<td>Phenytoin</td>
</tr>
<tr>
<td>Desipramine</td>
<td>Pravastatin</td>
</tr>
<tr>
<td>Diazepam</td>
<td>Pravastatin + Acetylsalicylic acid</td>
</tr>
<tr>
<td>Diphenhydramine</td>
<td>Prazosin</td>
</tr>
<tr>
<td>Doxazosin</td>
<td>Propoxyphene</td>
</tr>
<tr>
<td>Doxylamine</td>
<td>Quinapril</td>
</tr>
<tr>
<td>Enalapril</td>
<td>Ramipril</td>
</tr>
<tr>
<td>Eszopiclone</td>
<td>Rosuvastatin</td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>Simvastatin</td>
</tr>
<tr>
<td>Griseofulvin</td>
<td>Simvastatin + Ezetimibe</td>
</tr>
<tr>
<td>Herbal Preparations</td>
<td>Temazepam</td>
</tr>
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</table>

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<table>
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</tr>
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<td>Insulin</td>
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<tr>
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<td>Simvastatin + Ezetimibe</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Temazepam</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>Tinidazole</td>
</tr>
<tr>
<td>Isoniazid</td>
<td>Vitamin D</td>
</tr>
<tr>
<td>Isosorbide</td>
<td>Warfarin</td>
</tr>
<tr>
<td>Ketoconazole</td>
<td>Zolpidem</td>
</tr>
<tr>
<td>Leflunomide</td>
<td>Lorazepam</td>
</tr>
</tbody>
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Table 2: Drugs to be avoided with alcohol

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<tr>
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<tr>
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<tr>
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</tr>
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<tr>
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<tr>
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</tr>
<tr>
<td>Glyburide</td>
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</tr>
<tr>
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<tr>
<td>Haloperidol</td>
<td>Trihexyphenidyl</td>
</tr>
<tr>
<td>Imipramine</td>
<td></td>
</tr>
<tr>
<td>Insulin</td>
<td></td>
</tr>
<tr>
<td>Loratadine</td>
<td></td>
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</tbody>
</table>
## Appendix 6b
**Drug–Contraceptive Interactions**

### CONTRACEPTIVES, ORAL

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetazolamide</td>
<td>Antagonism of diuretic effect</td>
</tr>
<tr>
<td>Amiloride</td>
<td>Antagonism of diuretic effect</td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Antagonism of antidepressant effect but adverse effects increased due to increased plasma concentration of amitriptyline</td>
</tr>
<tr>
<td>Amoxycillin</td>
<td>Reduced contraceptive effect of estrogen-containing preparations</td>
</tr>
<tr>
<td>Ampicillin</td>
<td>Reduced contraceptive effect of estrogen-containing preparations</td>
</tr>
<tr>
<td>Atenolol</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Accelerated metabolism of medroxyprogesterone (does not apply to injectable medroxyprogesterone acetate for contraception); reduced contraceptive effect (does not apply to injectable norethisterone enantate for contraception)</td>
</tr>
<tr>
<td>Ceftazidime</td>
<td>Reduced contraceptive effect of estrogen-containing preparations</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>Reduced contraceptive effect of estrogen-containing preparations</td>
</tr>
<tr>
<td>Cefuroxime</td>
<td>Reduced effect of contraceptives</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Oral contraceptives increase plasma concentration of corticosteroids</td>
</tr>
<tr>
<td>Clomipramine</td>
<td>Antagonism of antidepressant effect but adverse effects increased due to increased plasma concentration of clomipramine</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Inhibition of cyclosporine metabolism (increased plasma–cyclosporine concentration)</td>
</tr>
<tr>
<td>Doxycycline</td>
<td>Reduced contraceptive effect of estrogen-containing preparations</td>
</tr>
<tr>
<td>Efavirenz</td>
<td>Efficacy of oral contraceptives reduced</td>
</tr>
<tr>
<td>Enalapril</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Anecdotal reports of contraceptive failure</td>
</tr>
<tr>
<td>Fosphenytoin</td>
<td>Reduced contraceptive effect</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Antagonism of diuretic effect</td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Glimepiride</td>
<td>Reduced hypoglycaemic action</td>
</tr>
<tr>
<td>Glucagon</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Glyceryl trinitrate</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Griseofulvin</td>
<td>Reduced contraceptive effect of levonorgestrel, accelerated metabolism of medroxyprogesterone (does not apply to injectable</td>
</tr>
</tbody>
</table>
CONTRACEPTIVES, ORAL

Acetazolamide Antagonism of diuretic effect
Amiloride Antagonism of diuretic effect
Amitriptyline Antagonism of antidepressant effect but adverse effects increased due to increased plasma concentration of amitriptyline
Amoxycillin Reduced contraceptive effect of estrogen-containing preparations
Ampicillin Reduced contraceptive effect of estrogen-containing preparations
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Ceftazidime Reduced contraceptive effect of estrogen-containing preparations
Ceftriaxone Reduced contraceptive effect of estrogen-containing preparations
Cefuroxime Reduced effect of contraceptives
Corticosteroids Oral contraceptives increase plasma concentration of corticosteroids
Clomipramine Antagonism of antidepressant effect but adverse effects increased due to increased plasma concentration of clomipramine
Cyclosporine Inhibition of cyclosporine metabolism (increased plasma–cyclosporine concentration)
Doxycycline Reduced contraceptive effect of estrogen-containing preparations
Efavirenz Efficacy of oral contraceptives reduced
Enalapril Antagonism of hypotensive effect
Fluconazole Anecdotal reports of contraceptive failure
Fosphenytoin Reduced contraceptive effect
Furosemide Antagonism of diuretic effect
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Glimepiride Reduced hypoglycaemic action
Glucagon Antagonism of hypotensive effect
Glyceryl trinitrate Antagonism of hypotensive effect
Griseofulvin Reduced contraceptive effect of levonorgestrel, accelerated metabolism of medroxyprogesterone (does not apply to injectable medroxyprogestrone acetate for contraception); does not apply to injectable norethisterone enantate for contraception
Hydralazine Antagonism of hypotensive effect
Hydrochlorothiazide Antagonism of diuretic effect
Insulins Antagonism of hypoglycaemic effect
Isosorbide dinitrate Antagonism of hypotensive effect
Metformin Antagonism of hypoglycaemic effect
Methyldopa Antagonism of hypotensive effect
Nelfinavir Accelerated metabolism of levonorgestrel, medroxyprogesterone and norethisterone (reduced contraceptive effect); does not apply to injectable medroxyprogestrone acetate for contraception
Nevirapine Accelerated metabolism of levonorgestrel, medroxyprogesterone and norethisterone (reduced contraceptive effect); does not apply to injectable medroxyprogestrone acetate for contraception
Nifedipine Antagonism of hypotensive effect
Phenobarbital Metabolism accelerated (reduced contraceptive effect); does not apply to injectable medroxyprogesterone acetate for contraception; does not apply to injectable norethisterone enantate for contraception
Phenytoin Accelerated metabolism of levonorgestrel, norethisterone (reduced contraceptive effect); does not apply to injectable medroxyprogestrone acetate for contraception; does not apply to injectable norethisterone enantate for contraception
Propranolol Antagonism of hypotensive effect
Rifampicin Accelerated metabolism of levonorgestrol and medroxyprogesterone (reduced contraceptive effect); does not apply to injectable medroxyprogestrone acetate for contraception; does not apply to injectable norethisterone enantate for contraception
Ritonavir Accelerated metabolism of levonorgestrol and norethisterone (reduced contraceptive effect); does not apply to injectable medroxyprogestrone acetate for contraception; does not apply to injectable norethisterone enantate for contraception
Sodium nitroprusside Antagonism of hypotensive effect
Spironolactone Antagonism of diuretic effect
Topiramate Failure of contraceptive effect
Theophylline Delayed excretion of theophylline; increased plasma concentration
Verapamil Antagonism of hypotensive effect
Warfarin Antagonism of anticoagulant effect
Appendix 6c
Drug–Drug Interactions

Two or more drugs administered at the same time may interact with each other. The interactions may be potentiation or antagonism of one drug by another or occasionally some other effect. Drug interactions may be of pharmacokinetic or pharmacodynamic type. The pharmacokinetic interactions can be because of absorption mechanism, competition of two drugs at the protein binding sites, metabolizing enzyme system or excretion. When two or more drugs are concomitantly administered there is always a possibility of pharmacokinetic or pharmacodynamic interaction. The pharmacodynamic interactions can be at the receptor level for competition at same drug target (enzyme/receptor) acting synergistically or antagonizing the effect of each other. The drugs which have narrow therapeutic window have greater potential to cause unexpected adverse effect when their pharmacokinetics or pharmacodynamics is altered. In such situation, the following precautions are advisable:

1. Concomitant administration of drugs should possibly be avoided.
2. When unavoidable, care should be taken and TDM is recommended.
3. When TDM is not possible logistically, clinical symptomatology be done.
4. Careful dose titration (upward/downward) be done to get optimum dose modification.

The following drug categories are considered as drugs of narrow therapeutic window: Antiepileptics, anticoagulants, anticancers, xanthenes, antidepressants, antiarrhythmics etc.

Some representative clinically relevant drug–drug interactions are listed below:

**ABICIXIMAB**
- Anticoagulants: Increased risk of bleeding
- Antiplatelet agents: Increased risk of bleeding

**ACETAZOLAMIDE**
- Carbamazepine: Increased risk of hyponatraemia; acetazolamide increases plasma–carbamazepine concentration
- Digoxin: Cardiac toxicity of digoxin increased if hypokalaemia occurs
- Furosemide: Increased risk of hypokalaemia
- Nifedipine: Enhanced hypotensive effect
- Phenytoin: Increased risk of osteomalacia

**ACETYLSALICYLIC ACID**
- Corticosteroids: Increased risk of gastrointestinal bleeding and ulceration; corticosteroids reduce plasma–salicylate concentration
- Heparin: Enhanced anticoagulant effect
- Methotrexate: Reduced excretion of methotrexate (increased toxicity)
- Warfarin: Increased risk of bleeding due to antiplatelet effect
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1. Concomitant administration of drugs should possibly be avoided.
2. When unavoidable, care should be taken and TDM is recommended.
3. When TDM is not possible logistically, clinical symptomatology be done.
4. Careful dose titration (upward/downward) be done to get optimum dose modification.

The following drug categories are considered as drugs of narrow therapeutic window: Antiepileptics, anticoagulants, anticancer, xanthenes, antidepressants, antiarrhythmics etc.

Some representative clinically relevant drug–drug interactions are listed below:

**ALENDRONATE**

- Calcium supplements
  Reduced absorption of alendronate
- Antacids
  Reduced absorption of alendronate

**ALLOPURINOL**

- Azathioprine
  Effects of azathioprine enhanced with increased toxicity; reduce dose when given with allopurinol
- Mercaptopurine
  Effects of 6-mercaptopurine enhanced with increased toxicity; reduce dose when given with allopurinol

**ALTEPLASE**

- Prostacyclin, nitrates
  Increased plasma–alteplase clearance
- Abciximab
  Additive effect
- Nitroglycerin
  Decreased thrombolytic effect of alteplase
- Warfarin, Antiplatelet agents
  Increased risk of bleeding
- NSAIDs
  Increased risk of GI bleeding

**AMILORIDE**

- Artemether + Lumefantrine
  Increased risk of ventricular arrhythmias if electrolyte disturbance occurs
- Cisplatin
  Increased risk of nephrotoxicity and ototoxicity
- Cyclosporine
  Increased risk of hyperkalaemia
- Enalapril
  Enhanced hypotensive effect; risk of severe hyperkalaemia

**AMINOPHYLLINE**

- Febuxostat
  Increased effect of aminophylline
- Rifamycin
  Decreased effect of aminophylline

**AMITRIPTYLINE**

- Artemether + Lumefantrine
  Increased risk of ventricular arrhythmias
- Carbamazepine
  Antagonism of anticonvulsant effect
- Haloperidol
  Increased plasma–amitriptyline concentration; increased risk of ventricular arrhythmias
- Phenobarbital
  Antagonism of anticonvulsant effect
- Phenytoin
  Antagonism of anticonvulsant effect
- Valproic acid
  Antagonism of anticonvulsant effect

**AMOXICILIN**

- Methotrexate
  Reduced excretion of methotrexate; increased risk of toxicity

**AMOXICILIN + CLAVULANIC ACID**

- Probenecid
  Increased concentrations of amoxycillin in serum and bile
- Allopurinol
  Occurrence of allergic cutaneous reactions
<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Digoxin</td>
<td>Increased absorption</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Increased incidence of bleeding</td>
</tr>
<tr>
<td><strong>AMPHOTERICIN B</strong></td>
<td></td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Increased risk of nephrotoxicity</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Increased digoxin toxicity if hypokalaemia occurs</td>
</tr>
<tr>
<td>Tacrolimus</td>
<td>Synergistic effect of amphotericin</td>
</tr>
<tr>
<td><strong>AMPICILLIN</strong></td>
<td></td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Reduced excretion of methotrexate; increased risk of toxicity</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Studies have failed to demonstrate an interaction, but common experience in anticoagulant clinics is that INR can be altered by a course of ampicillin</td>
</tr>
<tr>
<td><strong>ANTACIDS (Aluminium Hydroxide; Magnesium Hydroxide)</strong></td>
<td>Note: Antacids should preferably not be taken at the same time as other drugs since they may impair absorption</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>Reduced absorption of ciprofloxacin</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Reduced absorption of digoxin</td>
</tr>
<tr>
<td>Isoniazid</td>
<td>Reduced absorption of isoniazid</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Reduced absorption of phenytoin</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Reduced absorption of rifampicin</td>
</tr>
<tr>
<td><strong>ARTEMETHER + LUMEFANTRINE</strong></td>
<td></td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Increased risk of ventricular arrhythmias</td>
</tr>
<tr>
<td>Azithromycin</td>
<td>Avoid concomitant use</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>Increased risk of ventricular arrhythmias</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>Avoid concomitant use</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Avoid concomitant use</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Increased risk of ventricular arrhythmias if electrolyte disturbance occurs</td>
</tr>
<tr>
<td>Mefloquine</td>
<td>Increased risk of ventricular arrhythmias</td>
</tr>
<tr>
<td>Ofloxacin</td>
<td>Avoid concomitant use</td>
</tr>
<tr>
<td>Pyrimethamine</td>
<td>Avoid concomitant use</td>
</tr>
<tr>
<td>Quinine</td>
<td>Increased risk of ventricular arrhythmias</td>
</tr>
<tr>
<td>Sulfadoxine + Pyrimethamine</td>
<td>Avoid concomitant use</td>
</tr>
<tr>
<td><strong>ATENOLOL</strong></td>
<td></td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Masking of warning signs of hypoglycaemia such as tremor</td>
</tr>
<tr>
<td>Insulins</td>
<td>Enhanced hypoglycaemic effect; masking of warning signs of hypoglycaemia such as tremor</td>
</tr>
<tr>
<td>Drug / Combination</td>
<td>Interaction</td>
</tr>
<tr>
<td>----------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Lidocaine</td>
<td>Increased risk of myocardial depression</td>
</tr>
<tr>
<td>Nifedipine</td>
<td>Severe hypotension and heart failure occasionally</td>
</tr>
<tr>
<td>Verapamil</td>
<td>Asystole, severe hypotension and heart failure</td>
</tr>
<tr>
<td><strong>ATORVASTATIN</strong></td>
<td></td>
</tr>
<tr>
<td>Ketoconazole</td>
<td>Increased plasma concentration of atorvastatin and risk of myotoxicity in frequent</td>
</tr>
<tr>
<td>Itraconazole</td>
<td>Increased plasma concentration of atorvastatin and risk of myotoxicity in frequent</td>
</tr>
<tr>
<td>Ritonavir</td>
<td>Increased plasma concentration of atorvastatin and risk of myotoxicity in frequent</td>
</tr>
<tr>
<td>Erythromycin</td>
<td>Increased plasma concentration of atorvastatin and risk of myotoxicity in frequent</td>
</tr>
<tr>
<td>Fibrates</td>
<td>Increased risk of rhabdomyolysis</td>
</tr>
<tr>
<td><strong>AZATHIOPRINE</strong></td>
<td></td>
</tr>
<tr>
<td>Allopurinol</td>
<td>Effects of azathioprine enhanced</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Reduced absorption of phenytoin</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Transplants rejected</td>
</tr>
<tr>
<td>Sulfamethoxazole + Trimethoprim</td>
<td>Increased risk of haematological toxicity</td>
</tr>
<tr>
<td>Vaccines, Live</td>
<td>Avoid use of live vaccines with azathioprine (impairment of immune response)</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Reduced effect of anticoagulant</td>
</tr>
<tr>
<td><strong>AZITHROMYCIN</strong></td>
<td></td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Plasma concentration of cyclosporine increased</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Effect of digoxin enhanced</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Enhanced anticoagulant effect of warfarin</td>
</tr>
<tr>
<td><strong>BACLOFEN</strong></td>
<td></td>
</tr>
<tr>
<td>Tricyclic antidepressents</td>
<td>Risk of muscle weakness</td>
</tr>
<tr>
<td>MAO inhibitors</td>
<td>Depression of brain function as well as low blood pressure</td>
</tr>
<tr>
<td>Antidiabetic drugs</td>
<td>Increased blood sugar level</td>
</tr>
<tr>
<td><strong>BENZATHINE BENZYLPPENICILLIN</strong></td>
<td></td>
</tr>
<tr>
<td>Aminoglycosides</td>
<td>Reduced effect of aminoglycosides in patient with renal impairment</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Reduced excretion of methotrexate</td>
</tr>
<tr>
<td><strong>BLEOMYCIN</strong></td>
<td></td>
</tr>
<tr>
<td>Vaccines, Live</td>
<td>Avoid use of live vaccines with bleomycin (impairment of immune response)</td>
</tr>
</tbody>
</table>
Vinblastine Increased risk of cardiovascular toxicity

BROMOCRIPTINE
Ergot derivatives Additive dopamine agonistic activity

BUDESONIDE
Ketoconazole Plasma concentration of orally administered budesonide increased
Itraconazole Metabolism of budesonide inhibited
Clarithromycin Metabolism of budesonide inhibited
Erythromycin Metabolism of budesonide inhibited

BUPIVACAINE
Lidocaine Increased myocardial depression
Procainamide Increased myocardial depression
Quinidine Increased myocardial depression

BUSULPHAN
Itraconazole Increased level of busulphan
Metronidazole Increased level of busulphan
Nalidixic acid Risk of gastrointestinal toxicity
Thioguanine Risk of portal hypertension and esophageal varices

CALCIUM CARBONATE + VITAMIN D₃
Quinolones Risk of decreased absorption into the body
Tetracycline Risk of decreased absorption into the body
Mycophenolate mofetil Decreased effectiveness of mycophenolate mofetil

CALCIUM SALTS
Digoxin Large intravenous doses of calcium can precipitate arrhythmias
Tetracyclines Reduced absorption of tetracyclines

CAPREOMYCIN
BCG vaccine May make the vaccine ineffective
Neuromuscular blocking agents Increase in neuromuscular blocking effects
Typhoid vaccine May make the vaccine ineffective

CARBAMAZEPINE
Acetazolamide Increased risk of hyponatraemia; acetazolamide increases plasma–carbamazepine concentration
Amitriptyline Antagonism (convulsive threshold lowered); accelerated metabolism of amitriptyline; reduced plasma concentration; reduced effect antidepressant
Chloroquine  Convulsive threshold occasionally lowered
Chlorpromazine  Antagonism of anticonvulsant effect (convulsive threshold lowered)
Corticosteroids  Accelerated metabolism of corticosteroids
Cyclosporine  Accelerated metabolism (reduced plasma–cyclosporine concentration)
Diltiazem  Increased carbamazepine level
Erythromycin  Increased plasma–carbamazepine concentration
Fluphenazine  Antagonism of anticonvulsant effect (convulsive threshold lowered)
Haloperidol  Antagonism of anticonvulsant effect
Isoniazid  Increased plasma–carbamazepine concentration (also isoniazid hepatotoxicity increased)
Lopinavir  Reduced plasma–lopinavir concentration
Progestins  Accelerated metabolism of progestins
Sulfamethoxazole + Trimethoprim  May be enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of carbamazepine often lowered
Phenytoin  May be enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of phenytoin often lowered
Ritonavir  Plasma concentration increased by ritonavir
Valproic acid  Plasma concentration of valproic acid often lowered; plasma concentration of active metabolite of carbamazepine often raised
Verapamil  Enhanced effect of carbamazepine
Warfarin  Accelerated metabolism of warfarin (reduced anticoagulant effect)

CEFAZOLIN
Oral anticoagulants  Increased hypoprothrombinemic effect of anticoagulant.

CEFIXIME
Carbamazepine  Elevated carbamazepine levels
Anticoagulants  Increased prothrombin time

CEFTAZIDIME
Furosemide  Nephrotoxicity of ceftazidime increased
Warfarin  Enhanced anticoagulant effect
### CEFTRIAXONE

- **Warfarin**
  - Enhanced anticoagulant effect

### CHLORAMPHENICOL

- **Cyclosporine**
  - Plasma concentration of cyclosporine increased
- **Iron**
  - Avoid as can cause bone marrow depression which appears treatment of anaemia
- **Phenobarbital**
  - Metabolism of chloramphenicol accelerated (reduced chloramphenicol concentration)
- **Phenytoin**
  - Plasma–phenytoin concentration increased (risk of toxicity)
- **Vitamin B\textsubscript{12}**
  - Avoid concomitant use, can cause bone marrow depression

### CHLOROQUINE

- **Artemether + Lumefantrine**
  - Increased risk of ventricular arrhythmias
- **Carbamazepine**
  - Convulsive threshold occasionally lowered
- **Cyclosporine**
  - Increased plasma–cyclosporine concentration (increased risk of toxicity)
- **Digoxin**
  - Plasma–digoxin concentration increased
- **Mefloquine**
  - Increased risk of convulsions
- **Phenytoin**
  - Convulsive threshold occasionally lowered
- **Valproic acid**
  - Convulsive threshold occasionally lowered

### CHLORPROMAZINE

- **Amitriptyline**
  - Increased antimuscarinic adverse effects; increased plasma–amitriptyline concentration; increased risk of ventricular arrhythmias
- **Artemether + Lumefantrine**
  - Increased risk of ventricular arrhythmias
- **Clomipramine**
  - Increased antimuscarinic adverse effects; increased plasma–clomipramine concentration; increased risk of ventricular arrhythmias
- **Ether, Anaesthetic**
  - Enhanced hypotensive effect
- **Halothane**
  - Enhanced hypotensive effect
- **Ketamine**
  - Enhanced hypotensive effect
- **Nitrous oxide**
  - Enhanced hypotensive effect
- **Phenobarbital**
  - Antagonism of anticonvulsant effect (convulsive threshold lowered)
- **Phenytoin**
  - Antagonism of anticonvulsant effect (convulsive threshold lowered)
- **Procainamide**
  - Increased risk of ventricular arrhythmias

### CINNARIZINE

- CNS depressants (alcohol, barbiturates, hypnotics, narcotic analgesics, tricyclic antidepressants, sedatives and tranquillizers)
  - Additive sedation
- **Zolpidem**
  - Additive toxicity

### CIPROFLOXACIN

- **Artemether + Lumefantrine**
  - Avoid concomitant use
- **Cyclosporine**
  - Increased risk of nephrotoxicity
- **Glibenclamide**
  - Enhanced effect of glibenclamide
- **Ibuprofen**
  - Increased risk of convulsions
- **Warfarin**
  - Enhanced anticoagulant effect

### CISPLATIN

- **Aminoglycoside antibiotics**
  - Increased risk of nephrotoxicity and ototoxicity
- **Furosemide**
  - Increased risk of nephrotoxicity and ototoxicity
- **Hydrochlorothiazide**
  - Increased risk of nephrotoxicity and ototoxicity
- **Vancomycin**
  - Increased risk of nephrotoxicity and ototoxicity

### CLARITHROMYCIN

- **Oral anticoagulants**
  - Increased anticoagulant effect.
- **Carbamazepine**
  - Increased serum concentration of carbamazepine.
- **Digoxin**
  - Increased concentration of digoxin.
- **Lovastatin**
  - Avoid concomitant use
- **Sildenafil**
  - Dose reduction of sildenafil may be required.
- **Simvastatin**
  - Avoid concomitant use
- **Sirolimus**
  - Elevation in serum sirolimus level
- **Tacrolimus**
  - Elevation in serum sirolimus level
- **Tadalafil**
  - Dose reduction of tadalafil may be required.

### CLINDAMYCIN

- **Erythromycin**
  - Antagonist activity
- **Pancuronium**
  - Neuromuscular blockade exaggerated
<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Propranolol</td>
<td>Concomitant administration may increase plasma concentration of both drugs; enhanced hypotensive effect</td>
</tr>
<tr>
<td>Quinidine</td>
<td>Increased risk of ventricular arrhythmias</td>
</tr>
<tr>
<td>Ritonavir</td>
<td>Plasma concentration increased by ritonavir</td>
</tr>
<tr>
<td>Thiopental</td>
<td>Enhanced hypotensive effect</td>
</tr>
<tr>
<td>Valproic acid</td>
<td>Antagonism of anticonvulsant effect (convulsive threshold lowered)</td>
</tr>
</tbody>
</table>

**CINNARIZINE**

CNS depressants (alcohol, barbiturates, hypnotics, narcotic analgesics, tricyclic antidepressants, sedatives and tranquillizers)

Zolpidem Additive toxicity

**CIPROFLOXACIN**

Artemether + Lumefantrine Avoid concomitant use

Cyclosporine Increased risk of nephrotoxicity

Glibenclamide Enhanced effect of glibenclamide

Ibuprofen Increased risk of convulsions

Warfarin Enhanced anticoagulant effect

**CISPLATIN**

Aminoglycoside antibiotics Increased risk of nephrotoxicity and ototoxicity

Furosemide Increased risk of nephrotoxicity and ototoxicity

Hydrochlorothiazide Increased risk of nephrotoxicity and ototoxicity

Vancomycin Increased risk of nephrotoxicity and ototoxicity

**CLARITHROMYCIN**

Oral anticoagulants Increased anticoagulant effect.

Carbamazepine Increased serum concentration of carbamazepine.

Digoxin Increased concentration of digoxin.

Lovastatin Avoid concomitant use

Sildenafil Dose reduction of sildenafil may be required.

Simvastatin Avoid concomitant use

Sirolimus Elevation in serum sirolimus level

Tacrolimus Elevation in serum sirolimus level

Tadalafil Dose reduction of tadalafil may be required.

**CLINDAMYCIN**

Erythromycin Antagonist activity

Pancuronium Neuromuscular blockade exaggerated
<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kaoli-pectin</td>
<td>Reduced absorption rate</td>
</tr>
<tr>
<td>Gentamycin</td>
<td>Synergistic effect</td>
</tr>
<tr>
<td><strong>CLOBAZAM</strong></td>
<td></td>
</tr>
<tr>
<td>Cimetidine</td>
<td>Increased effect of clobazam</td>
</tr>
<tr>
<td>Barbiturates</td>
<td>Decreased serum level of clobazam</td>
</tr>
<tr>
<td><strong>CLONAZEPAM</strong></td>
<td></td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Decreased level of carbamazepine</td>
</tr>
<tr>
<td>Ketoconazole</td>
<td>Inhibition of metabolism of clonazepam</td>
</tr>
<tr>
<td><strong>CLOPIDOGREL</strong></td>
<td></td>
</tr>
<tr>
<td>Omeprazole</td>
<td>Plasma concentration of active metabolite of clopidogrel is decreased</td>
</tr>
<tr>
<td><strong>NSAIDs</strong></td>
<td></td>
</tr>
<tr>
<td>Beta blockers</td>
<td>Sinus bradycardia, monitor heart rate</td>
</tr>
<tr>
<td>Clomipramine</td>
<td>Risk of hypertensive crisis</td>
</tr>
<tr>
<td><strong>CODEINE</strong></td>
<td></td>
</tr>
<tr>
<td>Diazepam</td>
<td>Enhanced sedative effect</td>
</tr>
<tr>
<td>Ritonavir</td>
<td>Ritonavir increases plasma concentration of codeine</td>
</tr>
<tr>
<td><strong>CORTICOSTEROIDS</strong></td>
<td></td>
</tr>
<tr>
<td>Acetylsalicylic acid</td>
<td>Increased risk of gastrointestinal bleeding and ulceration; hydrocortisone reduces plasma salicylate concentration</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Atenolol</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Accelerated metabolism of hydrocortisone (reduced effect)</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Enalapril</td>
<td>Antagonism of diuretic effect; increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Antagonism of diuretic effect; increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Insulins</td>
<td>Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Levonorgestrel</td>
<td>Levonorgestrel increases plasma concentration of corticosteroids</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Increased risk of haematological toxicity</td>
</tr>
<tr>
<td>Nifedipine</td>
<td>Antagonism of hypotensive effect</td>
</tr>
</tbody>
</table>
Phenobarbital  Metabolism of hydrocortisone accelerated (reduced effect)
Phenytoin  Metabolism of hydrocortisone accelerated (reduced effect)
Rifampicin  Accelerated metabolism of corticosteroids (reduced effect)
Salbutamol  Increased risk of hypokalaemia if high doses of corticosteroids given with high doses of salbutamol
Warfarin  Anticoagulant effect altered

CYCLOPHOSPHAMIDE
Vaccines, Live  Avoid use of live vaccines with cyclophosphamide (impairment of immune response)

CYCLOSPORINE
Amphotericin B  Increased risk of nephrotoxicity
Ciprofloxacin  Increased risk of nephrotoxicity
Digoxin  Reduced clearance of digoxin (risk of toxicity)
Enalapril  Increased risk of hyperkalaemia
Erythromycin  Increased plasma-cyclosporine concentration
Methotrexate  Increased toxicity
Metoclopramide  Plasma-cyclosporine concentration increased
Ofloxacin  Increased risk of nephrotoxicity
Phenobarbital  Metabolism of cyclosporine accelerated
Phenytoin  Accelerated metabolism
Rifampicin  Accelerated metabolism (reduced plasma-cyclosporine concentration)
Ritonavir  Plasma concentration increased by ritonavir
Rosuvastatin  Marked rise in serum rosuvastatin level
Sulfonamides and Trimethoprim  Increased toxicity
Vaccines, Live  Avoid use of live vaccines with cyclosporine
Vancomycin  Increased risk of nephrotoxicity

DANAZOL
Anticoagulants (warfarin)  Danazol inhibits metabolism of

DANAZOL
Anticoagulants (warfarin)  Danazol inhibits metabolism of coumarins
Cyclosporine  Danazol inhibits metabolism of cyclosporine
Lovastatin  Increased risk of myopathy
Simvastatin  Increased risk of myopathy
Tacrolimus  Danazol increases plasma concentration of tacrolimus
### Dapsone

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rifampicin</td>
<td>Reduced plasma-dapsone concentration</td>
</tr>
<tr>
<td>Sulfamethoxazole + Trimethoprim</td>
<td>Plasma concentration of both dapsone and trimethoprim increased with concomitant use</td>
</tr>
</tbody>
</table>

### Desferrioxamine Mesylate

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ascorbic acid</td>
<td>May worsen iron toxicity</td>
</tr>
</tbody>
</table>

### Dexamethasone

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetazolamide</td>
<td>Increased risk of hypokalaemia; antagonism of diuretic effect</td>
</tr>
<tr>
<td>Acetylsalicylic acid</td>
<td>Increased risk of gastrointestinal bleeding and ulceration; dexamethasone reduces plasma-salicylate concentration</td>
</tr>
<tr>
<td>Albendazole</td>
<td>Plasma-albendazole concentration increased</td>
</tr>
<tr>
<td>Amiloride</td>
<td>Antagonism of diuretic effect</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Increased risk of hypokalaemia (avoid concomitant use unless dexamethasone needed to control reactions)</td>
</tr>
<tr>
<td>Atenolol</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Accelerated metabolism of dexamethasone (reduced effect)</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Enalapril</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Ephedrine</td>
<td>Metabolism of dexamethasone accelerated</td>
</tr>
<tr>
<td>Erythromycin</td>
<td>Erythromycin inhibits metabolism of dexamethasone</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Antagonism of diuretic effect; increased risk of hypokalaemia</td>
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<td>Glibenclamide</td>
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**DICYCLOMINE**

Antidepressants: Increased risk of antimuscarinic side effects

Antipsychotics: Antimuscarinics reduce effects of haloperidol; increased risk of antimuscarinic side effects when antimuscarinics given with clozapine; antimuscarinics reduce plasma concentration of phenothiazines, but risk of antimuscarinic side effects increased

**DIDANOSINE**

Divalproex: Risk of additive toxicity

Ganciclovir: Increased didanosine concentration

Metronidazole: Risk of additive toxicity

Pentamidine: Risk of additive toxicity

Stavudine: Risk of additive toxicity

Vinblastine: Risk of additive toxicity

**DIGOXIN**

Acetazolamide: Cardiac toxicity of digoxin increased if hypokalaemia occurs

Amphotericin B: Increased digoxin toxicity if hypokalaemia occurs

Atenolol: Increased AV block and bradycardia

Corticosteroids: Increased risk of hypokalemia

Cyclosporine: Reduced clearance of digoxin (risk of toxicity)

Furosemide: Cardiac toxicity of digoxin increased if hypokalaemia occurs

Hydrochlorothiazide: Cardiac toxicity of digoxin increased if hypokalaemia occurs

Nifedipine: Increased plasma concentration of digoxin

Timolol: Increased AV block and bradycardia

Verapamil: Increased plasma concentration of digoxin; increased AV block and bradycardia

**DIHYDROERGOTAMINE**

Amiodarone: Increased cardiac depressant effects

Azoles antifungal: Increased level of alkoloid Buspirone; Increased serum level of buspirone

Macrolide antibiotics: Increased plasma level of unchanged alkaloid and peripheral vasoconstriction

Protease inhibitors: Elevated levels of ergot alkaloids

Sumatriptan: Additive effect with dihydroergotamine

**DILTIAZEM**

Carbamazepine: Increased serum level of carbamazepine

Rifampin: Decreased diltiazem plasma concentration
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<tr>
<td>Spironolactone</td>
<td>Enhanced hypotensive effect, risk of severe hyperkalaemia</td>
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EPINEPHRINE (ADRENALINE)
Halothane Risk of arrhythmias
ERYTHROMYCIN
Artemether + Lumefantrine Avoid concomitant use
Carbamazepine Increased plasma-carbamazepine concentration
Corticosteroids Inhibits metabolism of corticosteroids
Cyclosporine Increased plasma-cyclosporine concentration
Digoxin Enhanced effect of digoxin
Warfarin Enhanced anticoagulant effect
ERYTHROPOIETIN
Haematinics Enhanced efficiency of erythropoietin.
ESCITALOPRAM
Carbamazepine Carbamazepine toxicity may be precipitated
ESMOLOL
Verapamil Chances of cardiac arrest
ETHINYL ESTRADIOL
Hydantoin Decreased effect of estrogen
ETOPOSIDE
Vaccines, Live Avoid use of live vaccines with etoposide
EZETIMIBE
Bile Acid Sequestrants Decreased levels and clinical effectiveness of ezetimibe
Fibrates Elevated levels of ezetimibe leading to toxicity.
Cyclosporine Increased ezetimibe levels in patients with severe renal insufficiency.
FACTOR IX
Acetylsalicylic acid Risk of bleeding
FAMOTIDINE
Antacids Reduced absorption of famotidine
Ketoconazole, itraconazole Reduced absorption of ketoconazole and itraconazole
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Ethanol Gastric mucosal irritation may occur.
FENOFIBRATE
Anticoagulants Increased effect of anticoagulants
Statins Increased risk of kidney and muscle problems
Cyclosporine Increased risk of nephrotoxicity
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<td>Absorption of ciprofloxacin reduced by oral ferrous salts</td>
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<td>Reduced hypotensive effect of methylidopa</td>
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<td>Ketoconazole</td>
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<td>Artemether + Lumefantrine</td>
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<td>Cyclosporine</td>
<td>Metabolism of cyclosporine inhibited</td>
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<td>Renal excretion of flucytosine decreased and cellular uptake increased (flucytosine toxicity increased)</td>
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<td><strong>GEMCITABINE</strong></td>
<td></td>
</tr>
<tr>
<td>Live vaccines</td>
<td>Serum antibody response may not be obtained</td>
</tr>
<tr>
<td>Zidovudine</td>
<td>Additive toxicity</td>
</tr>
<tr>
<td><strong>GENTAMICIN</strong></td>
<td></td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Increased risk of nephrotoxicity</td>
</tr>
<tr>
<td>Cisplatin</td>
<td>Increased risk of nephrotoxicity and ototoxicity</td>
</tr>
<tr>
<td>Suxamethonium</td>
<td>Enhanced muscle relaxant effect</td>
</tr>
<tr>
<td>Vancomycin</td>
<td>Increased risk of nephrotoxicity and ototoxicity</td>
</tr>
<tr>
<td>Vecuronium</td>
<td>Enhanced muscle relaxant effect</td>
</tr>
<tr>
<td><strong>GLIBENCLAMIDE</strong></td>
<td></td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>Enhanced effect of glibenclamide</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Enalapril</td>
<td>Hypoglycaemic effect enhanced</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Plasma concentration of glibenclamide increased</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Levonorgestrel</td>
<td>Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Sulfadoxine + Pyrimethamine</td>
<td>Effect of glibenclamide may be enhanced</td>
</tr>
<tr>
<td>Sulfamethoxazole + Trimethoprim</td>
<td>Effect of glibenclamide may be enhanced</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Enhanced hypoglycaemic effects and changes to anticoagulant effect</td>
</tr>
<tr>
<td><strong>GLICLAZIDE</strong></td>
<td></td>
</tr>
<tr>
<td>Acetylsalicylic acid</td>
<td>Effect of gliclazide is potentiated</td>
</tr>
<tr>
<td>Clofibrate</td>
<td>Effect of gliclazide is potentiated</td>
</tr>
<tr>
<td>Sulphonamides</td>
<td>Effect of gliclazide is potentiated</td>
</tr>
<tr>
<td>Oral anticoagulants</td>
<td>Effect of gliclazide is potentiated</td>
</tr>
<tr>
<td>MAO inhibitors</td>
<td>Effect of gliclazide is potentiated</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Effect of gliclazide is antagonized</td>
</tr>
<tr>
<td>Medication</td>
<td>Action</td>
</tr>
<tr>
<td>-------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Barbiturates</td>
<td>Effect of gliclazide is antagonized</td>
</tr>
<tr>
<td>Diuretics</td>
<td>Effect of gliclazide is antagonized</td>
</tr>
<tr>
<td>Diazoxide</td>
<td>Effect of gliclazide is antagonized</td>
</tr>
<tr>
<td>Glucocorticoids</td>
<td>Effect of gliclazide is antagonized</td>
</tr>
<tr>
<td>Sympathomimetics</td>
<td>Effect of gliclazide is antagonized</td>
</tr>
<tr>
<td><strong>GLIMEPIRIDE</strong></td>
<td></td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Reduced hypoglycaemic action</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Reduced hypoglycaemic action</td>
</tr>
<tr>
<td>Thiazides</td>
<td>Reduced hypoglycaemic action</td>
</tr>
<tr>
<td><strong>GLUCAGON</strong></td>
<td></td>
</tr>
<tr>
<td>Anticoagulants</td>
<td>Excess hypoprothrombinemia and bleeding complications</td>
</tr>
<tr>
<td><strong>GLYCERYL TRINITRATE</strong></td>
<td></td>
</tr>
<tr>
<td>Atenolol</td>
<td>Enhanced hypotensive effect</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td><strong>GRISEOFULVIN</strong></td>
<td></td>
</tr>
<tr>
<td>Levonorgestrel</td>
<td>Accelerated metabolism of levonorgestrel (reduced contraceptive effect)</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Metabolism of warfarin accelerated (reduced anticoagulant effect)</td>
</tr>
<tr>
<td><strong>HALOPERIDOL</strong></td>
<td></td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Increased plasma-amitriptyline concentration; increased risk of ventricular arrhythmias</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Antagonism of anticonvulsant effect, metabolism of haloperidol accelerated</td>
</tr>
<tr>
<td>Lithium</td>
<td>Increased risk of extrapyramidal effects and neurotoxicity</td>
</tr>
<tr>
<td>Metoclopramide</td>
<td>Increased risk of extrapyramidal effects</td>
</tr>
<tr>
<td>Phenobarbital</td>
<td>Antagonism of anticonvulsant effect, metabolism of haloperidol accelerated</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Antagonism of anticonvulsant effect (convulsive threshold lowered)</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Accelerated metabolism of haloperidol (reduced plasma-haloperidol concentration)</td>
</tr>
<tr>
<td>Valproic acid</td>
<td>Antagonism of anticonvulsant effect (convulsive threshold lowered)</td>
</tr>
<tr>
<td><strong>HALOTHANE</strong></td>
<td></td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Increased risk of arrhythmias and hypotension</td>
</tr>
<tr>
<td>Drug</td>
<td>Effect</td>
</tr>
<tr>
<td>----------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Atenolol</td>
<td>Enhanced hypotensive effect</td>
</tr>
<tr>
<td>Diazepam</td>
<td>Enhanced sedative effect</td>
</tr>
<tr>
<td>Levodopa</td>
<td>Risk of arrhythmias</td>
</tr>
<tr>
<td>Vancomycin</td>
<td>Hypersensitivity-like reactions can occur with concomitant intravenous vancomycin</td>
</tr>
<tr>
<td>Verapamil</td>
<td>Enhanced hypotensive effect and AV delay</td>
</tr>
<tr>
<td><strong>HEPARIN</strong></td>
<td></td>
</tr>
<tr>
<td>Acetylsalicylic acid</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Enalapril</td>
<td>Increased risk of hyperkalaemia</td>
</tr>
<tr>
<td><strong>HYDRAZINE</strong></td>
<td></td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td><strong>HYDROCHLOROTHIAZIDE</strong></td>
<td></td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Increased risk of postural hypotension</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Artemether + Lumefantrine</td>
<td>Increased risk of ventricular arrhythmias if electrolyte disturbance occurs</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Increased risk of hyponatraemia</td>
</tr>
<tr>
<td>Cisplatin</td>
<td>Increased risk of nephrotoxicity and ototoxicity</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Cardiac toxicity of digoxin increased if hypokalaemia occurs</td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>Risk of nephrotoxicity of ibuprofen increased; antagonism of diuretic effect Insulins Antagonism of hypoglycaemic effect</td>
</tr>
<tr>
<td>Lithium</td>
<td>Reduced lithium excretion (increased plasma-lithium concentration and risk of toxicity); furosemide safer than hydrochlorothiazide</td>
</tr>
<tr>
<td>Salbutamol</td>
<td>Increased risk of hypokalaemia with high doses of salbutamol</td>
</tr>
<tr>
<td><strong>IBUPROFEN</strong></td>
<td></td>
</tr>
<tr>
<td>Acetylsalicylic acid</td>
<td>Avoid concurrent administration (increased adverse effects, including gastrointestinal damage); antiplatelet effect of acetylsalicylic acid reduced</td>
</tr>
<tr>
<td>Atenolol</td>
<td>Antagonism of hypotensive effect</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Increased risk of nephrotoxicity</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>Increased risk of convulsions</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Increased risk of gastrointestinal bleeding and ulceration</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Exacerbation of heart failure, reduced GFR, and increased plasma-digoxin concentration</td>
</tr>
</tbody>
</table>
Enalapril Antagonism of hypotensive effect, increased risk of renal impairment
Glibenclamide Enhanced effect of glibenclamide
Hydrochlorothiazide Risk of nephrotoxicity of ibuprofen increased; antagonism of diuretic effect
Lithium Reduced excretion of lithium
Methotrexate Excretion of methotrexate reduced
Nifedipine Antagonism of hypotensive effect
Warfarin Anticoagulant effect enhanced
Zidovudine Increased risk of haematological toxicity
IMATINIB
Rifampin Increased clearance of imatinib
Warfarin Imatinib may inhibit metabolism of warfarin
IMIPENEM + CILASTATIN
Ganciclovir May result in generalised seizures
INDINAVIR
Carbamazepine Reduced plasma concentration of indinavir
Efavirenz Reduced plasma concentration of indinavir
Ergotamine Increased risk of ergotism (avoid concomitant use)
Nelfinavir Combination may lead to increased plasma concentration of either drug (or both)
Nevirapine Reduced plasma concentration of indinavir
Phenobarbital Reduced plasma concentration of indinavir
Rifampicin Metabolism enhanced by rifampicin
INSULINS
Atenolol Enhanced hypoglycaemic effect; masking of warning signs of hypoglycaemia such as tremor
Corticosteroids Antagonism of hypoglycaemic effect
Enalapril Hypoglycaemic effect enhanced
Furosemide Antagonism of hypoglycaemic effect
Hydrochlorothiazide Antagonism of hypoglycaemic effect
Levonorgestrel Antagonism of hypoglycaemic effect
Nifedipine Occasionally impaired glucose tolerance
IODINE
Lithium Synergistic toxicity
**IOPANOIC ACID**

- Atenolol: Iopanoic acid toxicity may occur
- Methotrexate: Methotrexate toxicity may occur

**ISONIAZID**

- Carbamazepine: Increased plasmacarbamazepine concentration
- Diazepam: Metabolism of diazepam inhibited
- Phenytoin: Metabolism of phenytoin inhibited

**ISOSORBIDE DINITRATE**

- Sildenafil: Serious hypotension, MI may be precipitated

**ISOTRETINOIN**

- Vitamin A: Additive toxicity
- Progesterone: Decreased efficacy of microdosed progesterone
- Corticosteroids, phenytoin: Increased risk of osteoporosis
- Carbamazepine: Decreased plasma levels of carbamazepine
- Tetracyclines: Increased risk of pseudotumor cerebri

**ISPAGHULA**

- Lithium: Decreased effect of lithium

**IVERMECTIN**

- Vitamin K Antagonists (eg, warfarin): Enhanced anticoagulant effect

**KETOCONAZOLE**

- Amphotericin B: Increased adverse effect
- Cyclosporine: Increased level of cyclosporine
- Tolbutamide: Reduces blood glucose level

**LAMIVUDINE**

- Foscarnet: Concurrent use not recommended

**LATANOPROST**

- Thiomersal: Risk of precipitate formation leflunomide
- Acenocoumarol: Increased anticoagulant effect
- Warfarin: Increased anticoagulant effect
- Methotrexate: Increased risk of hepatotoxicity
- Cholestyramine: Enhanced leflunomide excretion and increased total clearance by approximately 50%

**LEVOCETIRIZINE**

- Alcohol or CNS depressants: Additive sedation
- Theophylline: Increases the levels of levocetirizine in blood
### LEVODOPA

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metoclopramide</td>
<td>Antagonism of effects of levodopa</td>
</tr>
<tr>
<td>Ether, Anaesthetic</td>
<td>Risk of arrhythmias</td>
</tr>
<tr>
<td>Ferrous salts</td>
<td>Absorption of levodopa may be reduced</td>
</tr>
<tr>
<td>Halothane</td>
<td>Risk of arrhythmias</td>
</tr>
<tr>
<td>Methyldopa</td>
<td>Enhanced hypotensive effect; antagonism of antiparkinson-sonian effect</td>
</tr>
<tr>
<td>Nifedipine</td>
<td>Enhanced hypotensive effect</td>
</tr>
<tr>
<td>Propranolol</td>
<td>Enhanced hypotensive effect</td>
</tr>
<tr>
<td>Pyridoxine</td>
<td>Antagonism of levodopa unless carbidopa also given</td>
</tr>
</tbody>
</table>

### LEVOTHYROXINE

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenobarbital</td>
<td>Metabolism of levothyroxine accelerated (may increase levothyroxine requirements in hypothyroidism)</td>
</tr>
<tr>
<td>Theophylline</td>
<td>Metabolism of theophylline is increased; larger doses are required</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Enhanced anticoagulant effect</td>
</tr>
</tbody>
</table>

### LIDOCAINE

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetazolamide</td>
<td>Action of lidocaine antagonised by hypokalaemia</td>
</tr>
<tr>
<td>Atenolol</td>
<td>Increased risk of myocardial depression</td>
</tr>
<tr>
<td>Bupivacaine</td>
<td>Increased myocardial depression</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Action of lidocaine antagonised by hypokalaemia</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Action of lidocaine antagonised by hypokalaemia</td>
</tr>
<tr>
<td>Procaainamide</td>
<td>Increased myocardial depression</td>
</tr>
<tr>
<td>Propranolol</td>
<td>Increased risk of myocardial depression; increased risk of lidocaine toxicity</td>
</tr>
<tr>
<td>Quinidine</td>
<td>Increased myocardial depression</td>
</tr>
<tr>
<td>Timololol</td>
<td>Increased risk of myocardial depression</td>
</tr>
<tr>
<td>Verapamil</td>
<td>Increased risk of myocardial depression</td>
</tr>
</tbody>
</table>

### LITHIUM

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetazolamide</td>
<td>Excretion of lithium increased</td>
</tr>
<tr>
<td>Amiloride</td>
<td>Reduced lithium excretion (increased plasma-lithium concentration and risk of toxicity)</td>
</tr>
<tr>
<td>Enalapril</td>
<td>Enalapril reduces excretion of lithium (increased plasma-lithium concentration)</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Reduced lithium excretion (increased plasma-lithium concentration and risk of toxicity); furosemide safer than hydrochlorothiazide</td>
</tr>
</tbody>
</table>
Haloperidol  Increased risk of extrapyramidal effects and possibility of neurotoxicity
Hydrochlorothiazide Reduced lithium excretion (increased plasma-lithium concentration and risk of toxicity); furosemide safer than hydrochlorothiazide
Ibuprofen Reduced excretion of lithium (risk of toxicity)
Methyldopa Neurotoxicity may occur without increased plasma-lithium concentration
Spironolactone Reduced lithium excretion (increased plasma-lithium concentration and risk of toxicity)
Suxamethonium Enhanced muscle relaxant effect
LOPERAMIDE
Quinidine Increased CNS level of loperamide
MEBENDAZOLE
Carbamazepine Reduced plasma-mebendazole concentration (increase mebendazole dose for tissue infection)
Phenytoin Reduced plasma-mebendazole concentration (increase mebendazole dose for tissue infection)
MEFENAMIC ACID
Warfarin Risk of serious GI bleeding higher than users of either drug alone.
Lithium Reduced renal clearance and increased risk of lithium toxicity.
Methotrexate Reduced excretion of methotrexate and possible increased risk of toxicity
Phenobarbital Reduced plasma-mebendazole concentration (increase mebendazole dose for tissue infection)
6-MERCAPTOPURINE
Allopurinol Effects of 6-mercaptopurine enhanced with increased toxicity, reduce dose when given with allopurinol
Phenytoin Reduced absorption of phenytoin
Sulfamethoxazole + Trimethoprim Increased risk of haematological toxicity
Sulfasalazine Increased risk of leukopenia
Trimethoprim Increased risk of haematological toxicity
Vaccines, Live Avoid use of live vaccines with 6-mercaptopurine (impairment of immune response)
Warfarin Anticoagulant effect reduced
**MEROPENEM**

Probenecid  
Renal excretion of meropenem is inhibited

Valproic acid  
Serum valproic acid concentration is decreased

**METFORMIN**

Atenolol  
Masking of warning signs of hypoglycaemia such as tremor

Corticosteroids  
Antagonism of hypoglycaemic effect

Enalapril  
Hypoglycaemic effect enhanced

Levonorgestrel  
Antagonism of hypoglycaemic effect

Lithium  
May occasionally impair glucose tolerance

Medroxyprogesterone  
Antagonism of hypoglycaemic effect

Norethisterone  
Antagonism of hypoglycaemic effect

**METHADONE**

Cimetidine  
Effect of methadone may be increased

MAO Inhibitors  
Risk of hypotension, hyperexia etc.

**METHOTREXATE**

Acetylsalicylic acid  
Reduced excretion of methotrexate (increased toxicity)

Amoxycillin  
Reduced excretion of methotrexate (increased risk of toxicity)

Cyclosporine  
Increased toxicity

Ibuprofen  
Excretion of methotrexate reduced (increased risk of toxicity)

Nitrous oxide  
Increased antifolate effect (avoid concomitant use)

Phenytoin  
Reduced absorption of phenytoin; antifolate effect of methotrexate increased

Pyrimethamine  
Antifolate effect of methotrexate increased

Sulfadoxine + Pyrimethamine  
Antifolate effect of methotrexate increased; risk of methotrexate toxicity increased

Sulfamethoxazole + Trimethoprim  
Antifolate effect of methotrexate increased (avoid concomitant use); risk of methotrexate toxicity increased

Trimethoprim  
Antifolate effect of methotrexate increased (avoid concomitant use)

Vaccines, Live  
Avoid use of live vaccines with methotrexate (impairment of immune response)

**METHYLDOPA**

Ferrous salts  
Reduced hypotensive effect of methyldopa

Propranolol  
Enhanced hypotensive effect
MEROPENEM
Probenecid Renal excretion of meropenem is inhibited
Valproic acid Serum valproic acid concentration is decreased

METFORMIN
Atenolol Masking of warning signs of hypoglycaemia such as tremor
Corticosteroids Antagonism of hypoglycaemic effect
Enalapril Hypoglycaemic effect enhanced
Levonorgestrel Antagonism of hypoglycaemic effect
Lithium May occasionally impair glucose tolerance
Medroxyprogesterone Antagonism of hypoglycaemic effect
Norethisterone Antagonism of hypoglycaemic effect

METHADONE
Cimetidine Effect of methadone may be increased
MAO Inhibitors Risk of hypotension, hyperexia etc.

METHOTREXATE
Acetylsalicylic acid Reduced excretion of methotrexate (increased toxicity)
Amoxycillin Reduced excretion of methotrexate (increased risk of toxicity)
Cyclosporine Increased toxicity
Ibuprofen Excretion of methotrexate reduced (increased risk of toxicity)
Nitrous oxide Increased antifolate effect (avoid concomitant use)
Phenytoin Reduced absorption of phenytoin; antifolate effect of methotrexate increased
Pyrimethamine Antifolate effect of methotrexate increased
Sulfadoxine + Pyrimethamine Antifolate effect of methotrexate increased; risk of methotrexate toxicity increased
Sulfamethoxazole + Trimethoprim Antifolate effect of methotrexate increased (avoid concomitant use); risk of methotrexate toxicity increased
Trimethoprim Antifolate effect of methotrexate increased (avoid concomitant use)
Vaccines, Live Avoid use of live vaccines with methotrexate (impairment of immune response)

METHYLDOPA
Ferrous salts Reduced hypotensive effect of methyldopa
Propranolol Enhanced hypotensive effect

METHYL PREDNISOLONE
Amphotericin B Chances of potentiation of K+ concentration
Cyclosporine Levels increased up to 2 fold

METRONIDAZOLE
Phenytoin Metabolism of phenytoin inhibited (increased plasma phenytoin concentration)
Warfarin Enhanced anticoagulant effect MMR vaccine See vaccines, live

MIDAZOLAM
Ketoconazole Increased levels of midazolam
Verapamil Increased levels of midazolam

MIFEPRISTONE
Dexamethasone Decreased serum levels of mifepristone

MOMETASONE
Anticoagulants Increased or decreased effects of anticoagulants
Bupropion Increased risk of seizures
Quinolones Increased risk of tendonitis and/or tendon rupture
Quetiapine Decreased levels of quetiapine

MORPHINE
Ciprofloxacin Avoid premedication with morphine (reduced plasma ciprofloxacin concentration)
Quinidine Decreased analgesic effect
Ritonavir Ritonavir increases plasma concentration of morphine

MYCOPHENOLATE
Bile acid sequestrants Decreased level and clinical effect of mycophenolate
Antacids Decreased effect

NALIDIXIC ACID
Cyclosporine Increased risk of nephrotoxicity
Ibuprofen Increased risk of convulsions
Theophylline Increased risk of convulsions
Warfarin Enhanced anticoagulant effect

NELFINAVIR
Ergotamine Increased risk of ergotism (avoid concomitant use)
Phenobarbital Plasma concentration of nelfinavir reduced
Quinidine Increased risk of ventricular arrhythmias (avoid concomitant use)
Rifampicin          Plasma concentration of nelfinavir significantly reduced
                     (avoid concomitant use)

NEOSTIGMINE
Gentamicin          Antagonism of effect of neostigmine
Streptomycin        Antagonism of effect of neostigmine

NEVIRAPINE
Lopinavir           Plasma concentration of lopinavir reduced
Rifampicin          Reduced plasma concentration of nevirapine (avoid
                     concomitant use)
Saquinavir          Plasma concentration of saquinavir reduced (avoid
                     concomitant use)

NICOTINIC ACID
Ganglionic blocking agents and      Potentiates the effects of ganglionic blocking agents and
vasoactive drugs                    vasoactive drugs resulting in postural hypotension
Bile acid sequestrants (for example,
cholesterylamine)                   Bind and prevent absorption of niacin, should be separated by 4-6 hours.

NIFEDIPINE
Atenolol             Severe hypotension and heart failure occasionally
Cyclosporine         Increased plasma-nifedipine concentration (increased risk of adverse effects such as gingival hyperplasia)
Digoxin              Increased plasma concentration of digoxin
Magnesium (parenteral) Profound hypotension reported with nifedipine and intravenous magnesium sulphate in pre-eclampsia
Phenobarbital        Effect of nifedipine reduced
Phenytoin            Reduced effect of nifedipine
Propranolol          Severe hypotension and heart failure occasionally
Ritonavir            Plasma concentration increased by ritonavir
Rifampicin           Accelerated metabolism of nifedipine (plasma concentration significantly reduced)
Theophylline          Enhanced theophylline effect (increased plasma-theophylline concentration)
Timolol              Severe hypotension and heart failure occasionally

NITROUS OXIDE
Chlorpromazine       Enhanced hypotensive effect
Fluphenazine         Enhanced hypotensive effect
Haloperidol          Enhanced hypotensive effect
Methotrexate
Verapamil

**NORADRENALINE**
Guanethidine + methyldopa + reserpine + tricyclic antidepressants
Cocaine
MAOIs
Nonselective β-blockers

**OMEPRAZOLE**
Cilostazol
Nelfinavir
Raltegravir

**ONDANSETRON**
Tramadol

**OXCARBAZEPINE**
Lamotrigine

**OXYTETRACYCLINE**
Calcium and Iron dextran
Penicillins
Etritrenate and isotretinoin
Oral contraceptives

**PHENOBARBITAL**
Amitriptyline
Carbamazepine
Cyclosporine
Haloperidol
Nifedipine
Phenytoin
<table>
<thead>
<tr>
<th>Compound</th>
<th>Interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valproic acid</td>
<td>Enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of valproic acid often lowered; phenobarbital concentration often raised</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Metabolism of warfarin accelerated (reduced anticoagulant effect)</td>
</tr>
<tr>
<td>PHENOXYMETHYL PENICILLIN</td>
<td></td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Reduced excretion of methotrexate (increased risk of toxicity)</td>
</tr>
<tr>
<td>PHENTTOIN</td>
<td></td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Antagonism (convulsive threshold lowered); reduced plasma-amitriptyline concentration</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of phenytoin often lowered but may be raised; plasma concentration of carbamazepine often lowered</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>Plasma-phenytoin concentration increased (risk of toxicity)</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>Convulsive threshold occasionally lowered</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Accelerated metabolism (reduced plasma-cyclosporine concentration)</td>
</tr>
<tr>
<td>Clonazepam</td>
<td>Enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of clonazepam often lowered</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Effect of phenytoin enhanced; plasma concentration increased</td>
</tr>
<tr>
<td>Haloperidol</td>
<td>Antagonism of anticonvulsant effect (convulsive threshold lowered)</td>
</tr>
<tr>
<td>Isoniazid</td>
<td>Metabolism of phenytoin inhibited (enhanced effect)</td>
</tr>
<tr>
<td>Mefloquine</td>
<td>Antagonism of anticonvulsant effect</td>
</tr>
<tr>
<td>Metronidazole</td>
<td>Metabolism of phenytoin inhibited (increased plasma-phenytoin concentration)</td>
</tr>
<tr>
<td>Nifedipine</td>
<td>Reduced effect of nifedipine</td>
</tr>
<tr>
<td>Pyrimethamine</td>
<td>Antagonism of anticonvulsant effect; increased antifolate effect</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Accelerated metabolism of phenytoin (reduced plasma concentration)</td>
</tr>
</tbody>
</table>
Sulfadoxine + Pyrimethamine
- Plasma-phenytoin concentration increased; increased antifolate effect

Sulfamethoxazole + Trimethoprim
- Antifolate effect and plasma-phenytoin concentration increased

Valproic acid
- Enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of valproic acid often lowered; plasma concentration of phenytoin often raised (but may also be lowered)

Warfarin
- Accelerated metabolism of warfarin (Reduced anticoagulant effect, but enhancement also reported)

PIOGLITAZONE
- NSAID: Increased risk of fluid retention
- Rifampicin: Decreased plasma concentration.
- Ketoconazole: Increased plasma concentration.

PIPERACILLIN + TAZOBACTAM
- Aminoglycosides: Inactivation of aminoglycosides
- Methotrexate: Reduced clearance of methotrexate

PREDNISOLONE
- Amphotericin B: Increased risk of hypokalaemia (avoid concomitant use unless prednisolone needed to control reactions)
- Carbamazepine: Accelerated metabolism of prednisolone (reduced effect)
- Phenobarbital: Metabolism of prednisolone accelerated (reduced effect)
- Phenytoin: Metabolism of prednisolone accelerated (reduced effect)
- Rifampicin: Accelerated metabolism of prednisolone (reduced effect)
- Vaccines, Live: High doses of prednisolone impair immune response; avoid use of live vaccines

Warfarin: Anticoagulant effect altered

PROPOFOL
- Fentanyl: Concomitant use in pediatric patients may result in serious bradycardia
- CNS depressants: Increased sedative, anaesthetic and cardiorespiratory effects

PYRIDOXINE
- Levodopa: Antagonism of levodopa unless carbidopa also given

PYRIMETHAMINE
- Artemether + Lumefantrine: Avoid concomitant use
- Methotrexate: Antifolate effect of methotrexate increased
Phenytoin  Antagonism of anticonvulsant effect; increased antifolate effect
Sulfonamides + Trimethoprim  Increased antifolate effect
RALOXIFENE
Estrogen  Increased risk of adverse effects.
RAMIPRIL
Diuretics  Excessive reduction of blood pressure
Potassium supplements/Potassium sparing diuretics  Increased risk of hyperkalemia
Lithium  Increased serum lithium levels and lithium toxicity
RIFAMPICIN
Azathioprine  Transplants rejected
Cyclosporine  Accelerated metabolism (reduced plasma-cyclosporine concentration)
Dapsone  Reduced plasma-dapsone concentration
Fluconazole  Accelerated metabolism of fluconazole (reduced plasma concentration)
Glibenclamide  Accelerated metabolism (reduced effect) of glibenclamide
Haloperidol  Accelerated metabolism of haloperidol (reduced plasma-haloperidol concentration)
Nifedipine  Accelerated metabolism of nifedipine (plasma concentration significantly reduced)
Phenytoin  Accelerated metabolism of phenytoin (reduced plasma concentration)
Corticosteroids  Accelerated metabolism of corticosteroids
Verapamil  Accelerated metabolism of verapamil (plasma concentration significantly reduced)
Warfarin  Accelerated metabolism of warfarin (reduced anticoagulant effect)
RITONAVIR
Carbamazepine  Plasma concentration increased by ritonavir
Cyclosporine  Plasma concentration increased by ritonavir
Diazepam  Plasma concentration increased by ritonavir (risk of extreme sedation and respiratory depression-avoid concomitant use)
Fluconazole by ritonavir  Plasma concentration increased
<table>
<thead>
<tr>
<th>Drug / Combination</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Verapamil</td>
<td>Plasma concentration increased by ritonavir</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Plasma concentration increased by ritonavir</td>
</tr>
<tr>
<td><strong>SALBUTAMOL</strong></td>
<td></td>
</tr>
<tr>
<td>Methyldopa</td>
<td>Acute hypotension reported with salbutamol infusion</td>
</tr>
<tr>
<td><strong>SILDENAFIL</strong></td>
<td></td>
</tr>
<tr>
<td>Protease inhibitors</td>
<td>Sildenafil metabolism is inhibited</td>
</tr>
<tr>
<td>Alpha blockers</td>
<td>Avoid concomitant use (may lead to low blood pressure)</td>
</tr>
<tr>
<td>Ketoconazole</td>
<td>Increased action of sildenafil Erythromycin Increased action of sildenafil</td>
</tr>
<tr>
<td>Verapamil</td>
<td>Increased action of sildenafil</td>
</tr>
<tr>
<td>Nitrates</td>
<td>Vasoconstrictor activity of nitrates is potentiated</td>
</tr>
<tr>
<td><strong>STREPTOMYCIN</strong></td>
<td></td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Increased risk of nephrotoxicity</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Increased risk of nephrotoxicity</td>
</tr>
<tr>
<td>Cisplatin</td>
<td>Increased risk of nephrotoxicity and ototoxicity</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Increased risk of ototoxicity</td>
</tr>
<tr>
<td>Neostigmine</td>
<td>Antagonism of effect of neostigmine</td>
</tr>
<tr>
<td>Suxamethonium</td>
<td>Enhanced muscle relaxant effect</td>
</tr>
<tr>
<td><strong>STRONTIUM RANELATE</strong></td>
<td></td>
</tr>
<tr>
<td>Calcium products</td>
<td>Reduced bioavailability of strontium ranelate.</td>
</tr>
<tr>
<td>Tetracycline</td>
<td>Reduced absorption of oral tetracycline</td>
</tr>
<tr>
<td>Quinolone antibiotics</td>
<td>Reduced absorption of quinolone antibiotics</td>
</tr>
<tr>
<td>Alminium and Magnesium Hydroxides</td>
<td>Decreased absorption of strontium ranelate.</td>
</tr>
<tr>
<td><strong>SULFADOXINE + PYRIMETHAMINE</strong></td>
<td></td>
</tr>
<tr>
<td>Artemether + Lumefantrine</td>
<td>Avoid concomitant use</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Increased risk of nephrotoxicity</td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Effect of glibenclamide rarely, enhanced</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Antifolate effect of methotrexate increased; risk of methotrexate toxicity increased</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Plasma-phenytoin concentration increased; increased antifolate effect</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td><strong>SULFASALAZINE</strong></td>
<td></td>
</tr>
<tr>
<td>Azathioprine</td>
<td>Increased risk of leukopenia</td>
</tr>
<tr>
<td>Mercaptopurine</td>
<td>Increased risk of leukopenia</td>
</tr>
</tbody>
</table>
**TACROLIMUS**

- Aminoglycosides: Increased risk of renal dysfunction
- Carbamazepine: Decreased tacrolimus blood concentration
- Cisplatin: Increased risk of renal dysfunction
- Clarithromycin: Increased tacrolimus blood concentration
- Chloramphenicol: Increased tacrolimus blood concentration
- Clotrimazole: Increased tacrolimus blood concentration
- Phenytoin: Decreased tacrolimus blood concentration
- Rifampin: Decreased tacrolimus blood concentration
- Diltiazem: Increased tacrolimus blood concentration
- Nifedipine: Increased tacrolimus blood concentration
- Verapamil: Increased tacrolimus blood concentration

**TELMISARTAN**

- Lithium: Increased in serum lithium concentration and toxicity

**THALIDOMIDE**

- Barbiturates: Enhanced sedative activity
- Alcohol: Enhanced sedative activity
- Chlorpromazine: Enhanced sedative activity
- Reserpine: Enhanced sedative activity
- Vincristine: Potential to cause peripheral neuropathy
- Bortezomib: Potential to cause peripheral neuropathy

**THEOPHYLLINE**

- Ciprofloxacin: Increased plasma-theophylline concentration; increased risk of convulsions
- Erythromycin: Inhibition of theophylline metabolism (increased plasmatheophylline concentration resulting in theophylline toxicity)
- Fluconazole: Plasma-theophylline concentration increased

**TIMOLOL**

*Note: Systemic absorption may follow topical application of timolol to the eye*

- Epinephrine: Severe hypertension
- Verapamil: Asystole, severe hypotension and heart failure

**TOPIRAMATE**

- Carbamazepine: Reduced plasma level of topiramate
- Phenytoin: Reduced plasma level of topiramate
- Rifampin: Reduced plasma level of topiramate
**TRANEXAMIC ACID**
- Clotting factor complexes: Increased risk of thrombotic complications
- Hormonal contraception: Exacerbate the increased thrombotic risk associated with combination hormonal contraceptives
- all-trans Retinoic acid: Concomitant use in women with acute promyelocytic leukemia taking all-trans retinoic acid for remission induction may cause exacerbation of the procoagulant effect of all-trans retinoic acid

**TRIMETHOPRIM**
- Mercaptopurine: Increased risk of haematological toxicity
- Methotrexate: Antifolate effect of methotrexate increased (avoid concomitant use)
- Phenytoin: Antifolate effect and plasmaphenytoin concentration increased
- Pyrimethamine: Increased antifolate effect
- Sulfadoxine + Pyrimethamine: Increased antifolate effect

**VALPROIC ACID**
- Carbamazepine: Enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of valproic acid often lowered; plasma concentration of active metabolite of carbamazepine often raised
- Chloroquine: Convulsive threshold occasionally lowered
- Mefloquine: Antagonism of anticonvulsant effect
- Phenobarbital: Enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of valproic acid often lowered; phenobarbital concentration often raised
- Phenytoin: Enhanced toxicity without corresponding increase in antiepileptic effect; plasma concentration of valproic acid often lowered; plasma concentration of phenytoin often raised (but may also be lowered)

**VANCOMYCIN**
- Cyclosporine: Increased risk of nephrotoxicity
- Furosemide: Increased risk of ototoxicity

**VARICELLA VACCINE**
- Salicylates: Increased risk of Reye’s syndrome

**VERAPAMIL**
- Atenolol: Asystole, severe hypotension and heart failure
<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbamazepine</td>
<td>Enhanced effect of carbamazepine</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Increased plasma concentration of digoxin; increased AV block and bradycardia</td>
</tr>
<tr>
<td>Halothane</td>
<td>Enhanced hypotensive effect and AV delay</td>
</tr>
<tr>
<td>Ketamine</td>
<td>Enhanced hypotensive effect and AV delay</td>
</tr>
<tr>
<td>Lidocaine</td>
<td>Increased risk of myocardial depression</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Accelerated metabolism of verapamil (plasma concentration significantly reduced)</td>
</tr>
<tr>
<td>Vinblastine</td>
<td>v</td>
</tr>
<tr>
<td>Bleomycin</td>
<td>Increased risk of cardiovascular toxicity</td>
</tr>
<tr>
<td>Warfarin</td>
<td>v</td>
</tr>
<tr>
<td>Acetylsalicylic acid</td>
<td>Increased risk of bleeding due to antiplatelet effect</td>
</tr>
<tr>
<td>Azathioprine</td>
<td>Anticoagulant effect reduced</td>
</tr>
<tr>
<td>Azithromycin</td>
<td>Enhanced anticoagulant effect of warfarin</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Accelerated metabolism of warfarin (reduced anticoagulant effect)</td>
</tr>
<tr>
<td>Ceftazidime</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Anticoagulant effect altered</td>
</tr>
<tr>
<td>Doxycycline</td>
<td>Anticoagulant effect enhanced</td>
</tr>
<tr>
<td>Erythromycin</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>5-Fluorouracil</td>
<td>Anticoagulant effect enhanced</td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Enhanced hypoglycaemic effects and changes to anticoagulant effect</td>
</tr>
<tr>
<td>Griseofulvin</td>
<td>Metabolism of warfarin accelerated (reduced anticoagulant effect)</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>Anticoagulant effect enhanced</td>
</tr>
<tr>
<td>Levamisole</td>
<td>Anticoagulant effect enhanced</td>
</tr>
<tr>
<td>Levonorgestrel</td>
<td>Antagonism of anticoagulant effect</td>
</tr>
<tr>
<td>Levothyroxine</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Medroxyprogesterone</td>
<td>Antagonism of anticoagulant effect</td>
</tr>
<tr>
<td>Mercaptopurine</td>
<td>Anticoagulant effect reduced</td>
</tr>
<tr>
<td>Metronidazole</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Drug</td>
<td>Effect</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Nalidixic acid</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Norethisterone</td>
<td>Antagonism of anticoagulant effect</td>
</tr>
<tr>
<td>Ofloxacin</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Phenobarbital</td>
<td>Metabolism of warfarin accelerated (reduced anticoagulant effect)</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Accelerated metabolism of warfarin (reduced anticoagulant effect, but enhancement also reported)</td>
</tr>
<tr>
<td>Phytomenadione</td>
<td>Antagonism of anticoagulant effect by phytomenadione</td>
</tr>
<tr>
<td>Proguanil</td>
<td>Isolated reports of enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Quinidine</td>
<td>Anticoagulant effect may be enhanced</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Accelerated metabolism of warfarin (reduced anticoagulant effect)</td>
</tr>
<tr>
<td>Ritonavir</td>
<td>Plasma concentration increased by ritonavir</td>
</tr>
<tr>
<td>Sulfadiazine</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Sulfadoxine + Pyrimethamine</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Sulfamethoxazole + Trimethoprim</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td>Tamoxifen</td>
<td>Enhanced anticoagulant effect</td>
</tr>
<tr>
<td><strong>ZIDOVUDINE</strong></td>
<td></td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Increased plasma concentration of zidovudine (increased risk of toxicity)</td>
</tr>
<tr>
<td>Stavudine</td>
<td>May inhibit effect of stavudine (avoid concomitant use)</td>
</tr>
<tr>
<td><strong>ZOLPIDEM</strong></td>
<td></td>
</tr>
<tr>
<td>Rifampin</td>
<td>Pharmacodynamic effects of zolpidem are decreased</td>
</tr>
<tr>
<td>Ketoconazole</td>
<td>Pharmacodynamic effects of zolpidem are increased</td>
</tr>
</tbody>
</table>
## Appendix 6d
### Drug – Food Interactions

Several drugs when given orally can interact with food consumed by the patients. Table 1 shows the medications which should be taken on an empty stomach.

**Table 1: Medications which should be taken on an EMPTY stomach**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Food interactions and effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ampicillin</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Alendronate</td>
<td>Decreased bioavailability</td>
</tr>
<tr>
<td>Azithromycin</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Bisacodyl</td>
<td>Dissolves enteric coating</td>
</tr>
<tr>
<td>Didanosine</td>
<td>Decreased absorption</td>
</tr>
<tr>
<td>Indinavir</td>
<td>Reduced absorption with fat, proteins</td>
</tr>
<tr>
<td>Isoniazid</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Isosorbide dinitrate</td>
<td>Delayed absorption</td>
</tr>
<tr>
<td>Levothyroxine</td>
<td>Reduced absorption; anionic exchange resins reduce absorption</td>
</tr>
<tr>
<td>Melphalan</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Mycophenolate</td>
<td>Enhanced absorption</td>
</tr>
<tr>
<td>Omeprazole</td>
<td>Delayed absorption</td>
</tr>
<tr>
<td>Oxytetracycline</td>
<td>Reduced absorption when taken with dairy products.</td>
</tr>
<tr>
<td>Rifampin</td>
<td>Delayed absorption</td>
</tr>
<tr>
<td>Roxithromycin</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Sulfadiazine</td>
<td>Formation of crystalluria on consumption with vitamin C or acidifying agents</td>
</tr>
<tr>
<td>Tacrolimus</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Tetracycline</td>
<td>Reduced absorption, especially when taken with antacids or dairy products</td>
</tr>
<tr>
<td>Thyroid</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Typhoid vaccine (oral)</td>
<td>Reduced absorption</td>
</tr>
<tr>
<td>Zidovudine</td>
<td>Enhanced absorption</td>
</tr>
</tbody>
</table>
Food can also impact the effectiveness of a drug due to the way it is consumed. Generally, medicine is to be taken almost at the same time the food is eaten. This is because the medicine may upset the stomach if the stomach is empty. Certain medications are recommended to be taken with food (Table 2).

**Table 2: Medications which should be taken WITH FOOD**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Food interactions and effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetylsalicylic</td>
<td>Acid Reduced side effects.</td>
</tr>
<tr>
<td>Allopurinol</td>
<td>Reduced side effects; reduced clearance of active metabolite with protein-poor diet</td>
</tr>
<tr>
<td>Amiodarone</td>
<td>Enhances both the rate and extent of absorption.</td>
</tr>
<tr>
<td>Amoxicillin/clavulanic acid</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Azathioprine</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Baclofen</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Bromocriptine</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Increased absorption</td>
</tr>
<tr>
<td>Cefuroxime</td>
<td>Increased absorption</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Clofazimine</td>
<td>Increased drug absorption</td>
</tr>
<tr>
<td>Conjugated estrogens</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Diclofenac</td>
<td>Reduced peak concentration but not extent of absorption; reduced side effects</td>
</tr>
<tr>
<td>Doxycycline</td>
<td>Reduced side effects; reduced absorption with milk</td>
</tr>
<tr>
<td>Ethinyl estradiol</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Ferrous salts</td>
<td>Take between meals, if gastrointestinal upset occurs take with food</td>
</tr>
<tr>
<td>Griseofulvin</td>
<td>Increased rate or extent of absorption with fats; reduced side effects</td>
</tr>
<tr>
<td>Hydroxychloroquine</td>
<td>Reduced bowel side effects; masks the bitter taste of drug</td>
</tr>
<tr>
<td>Hydrocortisone</td>
<td>Slows rate of absorption; reduced peak levels; reduced side effects</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Iron preparations</td>
<td>See ferrous salts</td>
</tr>
<tr>
<td>Levocetirizine</td>
<td>May be taken with or without food</td>
</tr>
<tr>
<td>Lithium</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Mebendazole</td>
<td>Increased absorption</td>
</tr>
<tr>
<td>Methadone</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Drug</td>
<td>Effect</td>
</tr>
<tr>
<td>-----------------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>Methylprednisolone</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Metronidazole</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Mexiletine</td>
<td>Reduced side effects; slows rate of absorption; reduces rate of caffeine clearance</td>
</tr>
<tr>
<td>Morphine</td>
<td>Increased absorption</td>
</tr>
<tr>
<td>Nelfinavir</td>
<td>Greatly increases absorption and AUC</td>
</tr>
<tr>
<td>Niacin</td>
<td>Reduced absorption; decreases side effects</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>Increased absorption</td>
</tr>
<tr>
<td>Pioglitazone</td>
<td>Food slightly delays absorption rate but extent of absorption is not affected.</td>
</tr>
<tr>
<td>Potassium salts</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Prednisolone</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Prednisone</td>
<td>Reduced stomach irritation</td>
</tr>
<tr>
<td>Procainamide</td>
<td>Reduced side effects; increased absorption with fat</td>
</tr>
<tr>
<td>Propranolol</td>
<td>Slows rate but increases extent of absorption</td>
</tr>
<tr>
<td>Quinine</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Ritonavir</td>
<td>Increased absorption</td>
</tr>
<tr>
<td>Salsalate</td>
<td>Reduced stomach irritation.</td>
</tr>
<tr>
<td>Saquinavir</td>
<td>Increased absorption.</td>
</tr>
<tr>
<td>Sodium chloride</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Spironolactone</td>
<td>Increased absorption; reduced side effects</td>
</tr>
<tr>
<td>Sulfasalazine</td>
<td>Reduced side effects</td>
</tr>
<tr>
<td>Sodium valproate</td>
<td>Reduced side effects</td>
</tr>
</tbody>
</table>
Table 3: Selected herbal or food products resulting in adverse effects

<table>
<thead>
<tr>
<th>Herb/Food</th>
<th>Drug</th>
<th>Adverse Effects/Reported Drug Interactions/Remark</th>
</tr>
</thead>
<tbody>
<tr>
<td>Licorice</td>
<td>Digoxin</td>
<td>Elevates serum digoxin levels 4-fold, arrhythmias Hypokalemia and muscle weakness</td>
</tr>
<tr>
<td></td>
<td>Spironolactone</td>
<td></td>
</tr>
<tr>
<td>Foods high in vitamin K (broccoli, sprouts,</td>
<td>Anticoagulants (warfarin)</td>
<td>Such foods may reduce the effectiveness of anticoagulants, increasing the risk of clotting. Intake of such</td>
</tr>
<tr>
<td>turnip greens, spinach, cauliflower, legumes,</td>
<td></td>
<td>foods should be limited, and the amount consumed daily should remain constant.</td>
</tr>
<tr>
<td>mayonnaise, soybean oils and fish</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Foods high in sodium (like licorice, processed</td>
<td>Amlodipine</td>
<td>Such foods decrease the effectiveness of the drug</td>
</tr>
<tr>
<td>meats, canned foods)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium or foods containing calcium (milk and</td>
<td>Tetracycline</td>
<td>These foods can reduce the absorption of tetracycline, which should be taken 1 hr before or 2 hr after</td>
</tr>
<tr>
<td>other dairy products)</td>
<td></td>
<td>eating</td>
</tr>
<tr>
<td>Foods high in tyramine, (includes cheese,</td>
<td>MAO - inhibitors (such as phenelzine and</td>
<td>Severe headache and a potentially fatal increase in BP (hypertensive crisis) can occur if people taking</td>
</tr>
<tr>
<td>yoghurt, sour cream, cured meats, liver,</td>
<td>tranylcypromine)</td>
<td>MAO - inhibitors consume these foods. These foods must be avoided.</td>
</tr>
<tr>
<td>dried fish, bananas, yeast extracts, raisins,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>soya sauce, red wine, certain beers)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Appendix 7a
Hepatic Impairment

Dosing considerations in hepatic impairment

Hepatobiliary system plays an important role in the interactions between drugs and the body. Liver diseases can affect pharmacokinetics and pharmacodynamics of various drugs. However there has to be moderate to severe hepatic impairment to significantly alter the response to drugs as liver has a large reserve capacity. Hepatic impairment may alter response to drugs not only because of its role in metabolism of drugs but it also affects their absorption and distribution. Looking at the importance of liver in dealing with the drug, knowledge of a patient's hepatic function is required for the safe prescribing of many drugs. Unlike renal disease, where estimation of renal function based on creatinine clearance can fairly help in knowing the drug elimination and hence dose adjustment, there is no endogenous marker for hepatic clearance that can be used as a guide for drug dosing.

Hepatic impairment can lead to altered response to drugs due to all or some of the following reasons:

- Metabolism of many drugs depend on adequate liver function. Generally, metabolism result in the loss of pharmacological activity and therefore reduced metabolism in case of impaired liver function can lead to the accumulation of drug in the body to the toxic level at the normal dose. However in some cases drugs are metabolised to the active form and in these drugs normal dose may not be able to achieve desired response.

- For drugs with low bioavailability (high hepatic extraction), bioavailability increases and hepatic clearance decreases in cirrhotic patients. If such drug is to be administered orally to cirrhotic patients, their initial dose has to be reduced according to their hepatic extraction. For drugs with low bioavailability (low hepatic extraction), hepatic clearance may be affected due to impaired metabolism. For such drugs only the maintenance dose has to be adjusted according to estimated decrease in their hepatic metabolism.

- Portal hypertensive gastropathy and ulcers of upper gastrointestinal tract, frequently seen in cirrhotic patients may alter the absorption of orally administered drugs. Absorption of drugs may be increased because of high intestinal permeability in patients with portal hypertension. Impaired gastrointestinal motility seen in cirrhotic patients can lead to delayed drug absorption.

- Volume of distribution of hydrophilic drugs is increased due to presence of oedema and/or ascitis. Hence, loading dose of these drugs may have to be increased if a rapid action is required. On the other hand increase in volume of distribution is associated with an increase in the elimination half life of such drugs.
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- Impaired elimination of drugs which are excreted in the bile can lead to their accumulation in the body.

- Impaired albumin production can lead to decreased protein binding and increased toxicity of highly plasma protein bound drugs.

- High percentage of drugs may reach systemic circulation without passing through liver due to development of portosystemic shunts in cirrhotic patients.

- Cirrhotic patients can often have impaired renal function and in these cases dosage of the drugs have to be carefully adjusted.

The use of certain drugs in patients with cirrhosis may increase the risk of hepatic decompensation. In patients with impaired liver function dose related hepatotoxic reaction may occur at lower doses. Drugs that cause fluid retention (for example, prednisolone, ibuprofen, dexamethasone etc.) may exacerbate oedema and ascites in chronic liver disease. Sensitivity of brain to depressant action of some drugs (for example, morphine and barbiturates) is markedly increased in cirrhotic patients and can precipitate hepatic encephalopathy at normal doses.

As evident from above, there is a complex interactions between the drugs and liver function. Absence of any endogenous marker for hepatic clearance makes it highly difficult to accurately adjust the dose of various drugs in hepatic impairment. Therefore, if no immediate pharmacological effect is needed, drug therapy should be started cautiously in these patients and titrated individually until desired effect is achieved or toxicity appears. Drugs with a narrow therapeutic range and low hepatic extraction for e.g. theophylline are the most dangerous drugs. If such drugs are administered orally, both loading dose and maintenance doses have to be reduced by 50% of the normal dose, depending on the severity of hepatic impairment.

The following table contains information to help prescribing common drugs in hepatic impairment. The table provided is not exhaustive and absence from this table does not imply safety of drug, it is therefore important to refer to the individual drug entries.
<table>
<thead>
<tr>
<th>Drug</th>
<th>Status</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetylsalicylic acid</td>
<td>Avoid in severe hepatic impairment</td>
<td>Increased risk of Gastrointestinal bleeding</td>
</tr>
<tr>
<td>Allopurinol</td>
<td>Reduce the dose</td>
<td></td>
</tr>
<tr>
<td>Aluminium hydroxide</td>
<td>Avoid in severe hepatic impairment</td>
<td>Can precipitate hepatic encephalopathy by causing constipation. Antacids containing high amount of sodium to be avoided in patients with fluid retention.</td>
</tr>
<tr>
<td>Amidotrizoate</td>
<td>Use with caution</td>
<td></td>
</tr>
<tr>
<td>Amitryptiline</td>
<td>Avoid in severe hepatic impairment</td>
<td>Increased sedation</td>
</tr>
<tr>
<td>Amlodipine</td>
<td>Reduce dose</td>
<td>Half life of a mldipine is prolonged</td>
</tr>
<tr>
<td>Amodiaquine</td>
<td>Avoid in hepatic Impairment</td>
<td></td>
</tr>
<tr>
<td>Amoxycillin + Clavulanic acid</td>
<td>Use with caution</td>
<td>Monitor liver function, cholestatic jaundice reported either during or shortly after therapy (more common in males and patients over 65 years), duration of treatment should not exceed 2 weeks.</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Avoid in severe moderate to severe hepatic impairment</td>
<td>Cautiously given in mild hepatic impairment</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>Reduce dose and monitor plasma concentration if there is associated renal impairment</td>
<td></td>
</tr>
<tr>
<td>Chlorambucil</td>
<td>Reduce dose and use cautiously in hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Drug</td>
<td>Dose/Usage</td>
<td>Adverse Effect</td>
</tr>
<tr>
<td>-------------------</td>
<td>-----------------------------</td>
<td>-----------------------------------------------------</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>Avoid if possible, reduce dose and monitor plasma concentration</td>
<td>Increased risk of bone marrow depression</td>
</tr>
<tr>
<td>Chlorpheniramine</td>
<td>Avoid</td>
<td>May cause inappropriate sedation</td>
</tr>
<tr>
<td>Chlorpromazine</td>
<td>Use with caution</td>
<td>May precipitate coma</td>
</tr>
<tr>
<td>Clindamycin</td>
<td>Reduce dose</td>
<td></td>
</tr>
<tr>
<td>Clomifene</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Clomipramine</td>
<td>Avoid in severe hepatic impairment</td>
<td>Increased sedation</td>
</tr>
<tr>
<td>Cloxacillin</td>
<td>Use with caution</td>
<td>Cholestatic jaundice may occur up to several weeks after treatment has stopped. Risk increases with increasing age and if given for more than 2 weeks.</td>
</tr>
<tr>
<td>Codeine</td>
<td>Avoid or reduce dose</td>
<td>May precipitate coma. Causes constipation</td>
</tr>
<tr>
<td>Contraceptive, oral</td>
<td>Avoid in case of active liver disease</td>
<td>Avoid if history of cholestasis and pruritus during pregnancy.</td>
</tr>
<tr>
<td>Cyclophosphamide</td>
<td>Reduce dose</td>
<td>Monitor plasma level</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Reduce dose and use with caution</td>
<td>Hepatotoxic</td>
</tr>
<tr>
<td>Cytarabine</td>
<td>Reduce dose</td>
<td></td>
</tr>
<tr>
<td>Dacarbazine</td>
<td>Avoid in severe hepatic impairment</td>
<td>Dose reduction in mild to moderate hepatic impairment.</td>
</tr>
<tr>
<td>Daunorubicin</td>
<td>Reduce dose</td>
<td>Use with caution as toxicity increases in hepatic impairment.</td>
</tr>
<tr>
<td>Diazepam</td>
<td>Avoid in severe hepatic impairment.</td>
<td>Can precipitate coma</td>
</tr>
<tr>
<td>Didanosine</td>
<td>Monitor for toxicity</td>
<td></td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>Reduce dose according to bilirubin concentration</td>
<td></td>
</tr>
<tr>
<td>Doxycycline</td>
<td>Avoid or use with caution</td>
<td></td>
</tr>
<tr>
<td>Drug</td>
<td>Precaution Information</td>
<td></td>
</tr>
<tr>
<td>--------------------</td>
<td>----------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Efavirenz</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dose reduction and/or use with caution in mild to moderate hepatic impairment.</td>
<td></td>
</tr>
<tr>
<td>Enalapril</td>
<td>Use with caution</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Closely monitor liver function in patients with hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Ergometrine</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Erythromycin</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td></td>
<td>May cause idiosyncratic hepatotoxicity</td>
<td></td>
</tr>
<tr>
<td>Ethinylestradiol</td>
<td>Avoid</td>
<td></td>
</tr>
<tr>
<td></td>
<td>See also Contraceptives, Oral</td>
<td></td>
</tr>
<tr>
<td>Etoposide</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increased risk of toxicity in case of hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Use with caution</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hepatotoxicity</td>
<td></td>
</tr>
<tr>
<td>5-Fluorouracil</td>
<td>Use with caution; dose reduction may be required</td>
<td></td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>Reduce dose or administer on alternate days</td>
<td></td>
</tr>
<tr>
<td>Fluphenazine</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hepatotoxic, can precipitate coma</td>
<td></td>
</tr>
<tr>
<td>Furosemide</td>
<td>Avoid or use with caution in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hypokalaemia may precipitate coma (use potassium sparing diuretic to prevent this); Increased risk of hypomagnesaemia in alcoholic cirrhosis</td>
<td></td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Avoid or reduce the dose</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increased risk of hypoglycaemia. Can produce jaundice</td>
<td></td>
</tr>
<tr>
<td>Griseofulvin</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Haloperidol</td>
<td>Use with caution</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Can precipitate coma</td>
<td></td>
</tr>
<tr>
<td>Heparin</td>
<td>Reduce dose in severe liver disease</td>
<td></td>
</tr>
<tr>
<td>Hydralazine</td>
<td>Reduce dose</td>
<td></td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hypokalaemia may precipitate coma (use potassium sparing diuretic to prevent this); Increased risk of hypomagnesaemia in alcoholic cirrhosis</td>
<td></td>
</tr>
<tr>
<td>Drug</td>
<td>Advice to Patients</td>
<td>Considerations</td>
</tr>
<tr>
<td>-----------------------</td>
<td>-----------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>Avoid in severe hepatic impairment</td>
<td>Increased risk of gastrointestinal bleeding and can also cause fluid retention</td>
</tr>
<tr>
<td>Indinavir</td>
<td>Reduce dose to 600 mg 8th hourly in mild to moderate hepatic impairment, not studied in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Isoniazid</td>
<td>Use with caution</td>
<td>Regularly monitor liver function and particularly frequently in first 2 months.</td>
</tr>
<tr>
<td>Levonorgestrel</td>
<td>Use with caution in active liver disease and recurrent cholestatic jaundice</td>
<td></td>
</tr>
<tr>
<td>Lidocaine</td>
<td>Avoid or reduce the dose in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Magnesium hydroxide/sulphate</td>
<td>Avoid in hepatic coma if risk of renal failure</td>
<td></td>
</tr>
<tr>
<td>Medroxyprogesterone</td>
<td>Avoid in active liver disease.</td>
<td>Avoid if history of pruritus and cholestasis during pregnancy</td>
</tr>
<tr>
<td>Mefloquine</td>
<td>Avoid for prophylaxis in severe liver disease</td>
<td></td>
</tr>
<tr>
<td>6-Mercaptopurine</td>
<td>May need dose reduction</td>
<td></td>
</tr>
<tr>
<td>Metformin</td>
<td>Avoid</td>
<td>Withdraw if tissue hypoxia likely</td>
</tr>
<tr>
<td>Methadone</td>
<td>Avoid or reduce the dose</td>
<td>May precipitate coma</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Avoid in severe hepatic impairment</td>
<td>Hepatotoxic, monitor hepatic impairment liver functions</td>
</tr>
<tr>
<td>Methyldopa</td>
<td>Avoid in active liver disease</td>
<td></td>
</tr>
<tr>
<td>Metoclopramide</td>
<td>Reduce dose</td>
<td></td>
</tr>
<tr>
<td>Metronidazole</td>
<td>Reduce total daily dose to one third and give once daily in case of severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Drug</td>
<td>Adverse Effect(s)</td>
<td>Precaution(s)</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Morphine</td>
<td>May precipitate coma</td>
<td>Avoid or reduce the dose</td>
</tr>
<tr>
<td>Nevirapine</td>
<td>Use with caution in moderate hepatic impairment.</td>
<td>Avoid in severe hepatic impairment.</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>Cholestatic jaundice and chronic active hepatitis reported</td>
<td>Use with caution</td>
</tr>
<tr>
<td>Norethisterone</td>
<td>Avoid if history of pruritus and cholestasis during pregnancy</td>
<td>Avoid in active liver disease.</td>
</tr>
<tr>
<td>Ofloxacin</td>
<td>Hepatic dysfunction reported</td>
<td>Reduce dose in severe hepatic impairment</td>
</tr>
<tr>
<td>Paracetamol</td>
<td>Adverse effects more common</td>
<td>Avoid large doses-related toxicity.</td>
</tr>
<tr>
<td>Phenobarbital</td>
<td></td>
<td>Avoid in severe hepatic impairment.</td>
</tr>
<tr>
<td>Phenytoin</td>
<td></td>
<td>Reduce dose to avoid toxicity</td>
</tr>
<tr>
<td>Prednisolone</td>
<td></td>
<td>Use with caution</td>
</tr>
<tr>
<td>Procainamide</td>
<td></td>
<td>Avoid or reduce the dose</td>
</tr>
<tr>
<td>Procarbazine</td>
<td></td>
<td>Avoid in severe hepatic impairment.</td>
</tr>
<tr>
<td>Promethazine</td>
<td></td>
<td>Avoid in severe hepatic impairment. May precipitate coma,</td>
</tr>
<tr>
<td>Propylthiouracil</td>
<td></td>
<td>Hepatotoxic</td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td></td>
<td>Avoid in severe hepatic impairment. Monitor hepatic function - idiosyncratic</td>
</tr>
<tr>
<td>Pyrimethamine</td>
<td></td>
<td>Hepatotoxicity more common</td>
</tr>
<tr>
<td>Ranitidine</td>
<td></td>
<td>Use with caution</td>
</tr>
<tr>
<td>Ribavirin</td>
<td></td>
<td>Reduce dose</td>
</tr>
<tr>
<td>Rifampicin</td>
<td></td>
<td>Avoid in severe hepatic impairment.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Avoid or do not exceed 8 mg/kg daily.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Monitor liver function</td>
</tr>
<tr>
<td>Substance</td>
<td>Considerations</td>
<td></td>
</tr>
<tr>
<td>---------------------------------</td>
<td>--------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Saquinavir</td>
<td>Avoid in severe hepatic impairment. Caution in moderate hepatic impairment.</td>
<td></td>
</tr>
<tr>
<td>Simvastatin</td>
<td>Avoid in active liver disease or unexplained persistent elevation in serum transaminases</td>
<td></td>
</tr>
<tr>
<td>Sodium nitroprusside</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Sulfadiazine</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Sulfamethoxazole + trimethoprim</td>
<td>Avoid in severe hepatic impairment</td>
<td></td>
</tr>
<tr>
<td>Suxamethonium</td>
<td>Prolonged apnoea may occur in severe liver disease due to reduced hepatic synthesis of plasma cholinesterase</td>
<td></td>
</tr>
<tr>
<td>Testosterone</td>
<td>Preferably avoid. Possibility of dose related toxicity and fluid retention.</td>
<td></td>
</tr>
<tr>
<td>Thiopental</td>
<td>Reduce dose in severe liver disease</td>
<td></td>
</tr>
<tr>
<td>Valproic acid</td>
<td>Avoid if possible. Hepatotoxicity and hepatic failure may occasionally occur (usually in first 6 months)</td>
<td></td>
</tr>
<tr>
<td>Verapamil</td>
<td>Reduce oral dose</td>
<td></td>
</tr>
<tr>
<td>Vinblastine</td>
<td>Reduction of dose may require</td>
<td></td>
</tr>
<tr>
<td>Vincristine</td>
<td>Reduction of dose may require</td>
<td></td>
</tr>
<tr>
<td>Warfarin</td>
<td>Avoid in severe liver disease. Reduced production of clotting factors in hepatic impairment, may increase risk of bleeding</td>
<td></td>
</tr>
<tr>
<td>Zidovudine</td>
<td>Reduction of dose as accumulation may occur</td>
<td></td>
</tr>
</tbody>
</table>
Administration of some drugs (for example, ergotamine) to nursing mothers may harm the infant, whereas administration of others (for example, digoxin) has little effect. Some drugs inhibit lactation (for example, estrogens).

Toxicity to the infant can occur if the drug enters the milk in pharmacologically significant quantities. The concentration in milk of some drugs (for example, iodides) may exceed that in the maternal plasma so that therapeutic doses in the mother may cause toxicity to the infant. Some drugs inhibit the infant’s sucking reflex (for example, phenobarbital). Drugs in breast milk may, at least theoretically, cause hypersensitivity in the infant even when the concentration is too low for a pharmacological effect.

The following table lists drugs:

- which should be used with caution or which are contraindicated in lactation for the reasons given above;
- which are not known to be harmful to the infant although they are present in milk in significant amounts.

For many drugs insufficient evidence is available to provide guidance and it is advisable to administer only drugs essential to a mother during lactation. Because of the inadequacy of information on drugs in breast milk the following table should be used only as a guide; absence from the table does not imply safety.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Comment</th>
</tr>
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<tbody>
<tr>
<td>Abacavir</td>
<td>Lactation recommended during first 6 months if no safe alternative to breast milk</td>
</tr>
<tr>
<td>Acetylsalicylic acid</td>
<td>Short course safe in usual dosage; monitor infant; regular use of high doses could impair platelet function and produce hypoprophospholipinemia in infant if neonatal vitamin K stores low; possible risk of Reye syndrome</td>
</tr>
<tr>
<td>Acyclovir</td>
<td>Significant amount in milk after systemic administration, but considered safe to use</td>
</tr>
<tr>
<td>Alcohol</td>
<td>Large amounts may affect infant and reduce milk consumption</td>
</tr>
<tr>
<td>Aminophylline</td>
<td>Present in milk-irritability in infant reported</td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Detectable in breast milk; continue lactation; adverse effects possible, monitor infant for drowsiness</td>
</tr>
<tr>
<td>Amoxycillin</td>
<td>Trace amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Amoxycillin + Clavulanic acid</td>
<td>Trace amounts in milk</td>
</tr>
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</table>
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Toxicity to the infant can occur if the drug enters the milk in pharmacologically significant quantities. The concentration in milk of some drugs (for example, iodides) may exceed that in the maternal plasma so that therapeutic doses in the mother may cause toxicity to the infant. Some drugs inhibit the infant’s sucking reflex (for example, phenobarbital). Drugs in breast milk may, at least theoretically, cause hypersensitivity in the infant even when the concentration is too low for a pharmacological effect.

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<tr>
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<tr>
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<tr>
<td>Amitriptyline</td>
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</tr>
<tr>
<td>Amoxycillin</td>
<td>Trace amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Amoxycillin + Clavulanic acid</td>
<td>Trace amounts in milk</td>
</tr>
<tr>
<td>Ampicillin</td>
<td>Trace amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Artemether + Lumefantrine</td>
<td>Discontinue lactation during and for 1 week after stopping treatment; present in milk in animal studies</td>
</tr>
<tr>
<td>Asparaginase</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Atenolol</td>
<td>Significant amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Atropine</td>
<td>Small amount present in milk; monitor infant</td>
</tr>
<tr>
<td>Azathioprine</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Beclomethasone</td>
<td>Systemic effects in infant unlikely with maternal dose of less than equivalent of prednisolone 40 mg daily; monitor infant’s adrenal function with higher doses</td>
</tr>
<tr>
<td>Benzathine benzylpenicillin</td>
<td>Trace amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Benzylpenicillin</td>
<td>Trace amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Betamethasone</td>
<td>Systemic effects in infant unlikely with maternal dose of less than equivalent of prednisolone 40 mg daily; monitor infant’s adrenal function with higher doses</td>
</tr>
<tr>
<td>Bleomycin</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Continue lactation; adverse effects possible (severe skin reaction reported in 1 infant); monitor infant for drowsiness;</td>
</tr>
<tr>
<td>Ceftazidime</td>
<td>Excreted in low concentrations; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>Excreted in low concentrations; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Chlorambucil</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>Continue lactation; use alternative drug if possible; may cause bone-marrow toxicity in infant; concentration in milk usually insufficient to cause ‘grey syndrome’</td>
</tr>
<tr>
<td>Chlormethine</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>For malaria prophylaxis, amount probably too small to be harmful; inadequate for reliable protection against malaria, avoid lactation when used for rheumatic disease</td>
</tr>
<tr>
<td>Chlorpheniramime</td>
<td>Safe in usual dosage; monitor infant for drowsiness</td>
</tr>
<tr>
<td>Chlorpromazine</td>
<td>Continue lactation; adverse effects possible; monitor infant for drowsiness</td>
</tr>
<tr>
<td>Drug</td>
<td>Lactation status</td>
</tr>
<tr>
<td>-----------------------</td>
<td>--------------------------------------------</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>Continue lactation; use alternative drug if possible; high concentrations in breast milk</td>
</tr>
<tr>
<td>Cisplatin</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Clindamycin</td>
<td>Amount probably too small to be harmful but bloody diarrhoea reported in 1 infant</td>
</tr>
<tr>
<td>Clomifene</td>
<td>May inhibit lactation</td>
</tr>
<tr>
<td>Clomipramine</td>
<td>Small amount present in milk; continue lactation; adverse effects possible; monitor infant for drowsiness</td>
</tr>
<tr>
<td>Clonazepam</td>
<td>Continue lactation; adverse effects possible; monitor infant for drowsiness;</td>
</tr>
<tr>
<td>Cloxacillin</td>
<td>Trace amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Colchicine</td>
<td>Present in milk but no adverse effects reported; caution because of risk of cytotoxicity</td>
</tr>
<tr>
<td>Contraceptives, oral</td>
<td>Combined oral contraceptives may inhibit lactation-use alternative method of contraception until weaning or for 6 months after birth; progestogen-only contraceptives do not affect lactation (start 3 weeks after birth or later)</td>
</tr>
<tr>
<td>Cyclophosphamide</td>
<td>Lactation contraindicated during and for 36 h after stopping treatment</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Present in milk-avoid</td>
</tr>
<tr>
<td>Cytarabine</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Dacarbazine</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Dactinomycin</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Dapsone</td>
<td>Although significant amount in milk risk to infant very small; continue lactation; monitor infant for jaundice</td>
</tr>
<tr>
<td>Daunorubicin</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Dexamethasone</td>
<td>Systemic effects in infant unlikely with maternal dose of less than equivalent of prednisolone 40 mg daily; monitor infant’s adrenal function with higher doses</td>
</tr>
<tr>
<td>Diazepam</td>
<td>Continue lactation; adverse effects possible; monitor infant for drowsiness;</td>
</tr>
<tr>
<td>Didanosine</td>
<td>Lactation recommended during first 6 months if no safe alternative to breast milk</td>
</tr>
<tr>
<td>Difloxanide Avoid</td>
<td>Doxorubicin Lactation contraindicated</td>
</tr>
<tr>
<td>Drug</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Doxycycline</td>
<td>Continue lactation; use alternative drug if possible (absorption and therefore discolouration of teeth in infant probably usually prevented by chelation with calcium in milk)</td>
</tr>
<tr>
<td>Efavirenz</td>
<td>Lactation recommended during first 6 months if no safe alternative to breast milk</td>
</tr>
<tr>
<td>Eflornithine Avoid</td>
<td>Ephedrine Irritability and disturbed sleep reported</td>
</tr>
<tr>
<td>Ergocalciferol</td>
<td>Caution with high doses; may cause hypercalcaemia in infant</td>
</tr>
<tr>
<td>Ergotamine</td>
<td>Use alternative drug; ergotism may occur in infant; repeated doses may inhibit lactation</td>
</tr>
<tr>
<td>Erythromycin</td>
<td>Only small amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Ethinylestradiol</td>
<td>Use alternative method of contraception; may inhibit lactation; see also Contraceptives, Oral</td>
</tr>
<tr>
<td>Etoposide</td>
<td>Lactation contraindicated</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Present in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Flucytosine</td>
<td>Avoid</td>
</tr>
<tr>
<td>5-Fluorouracil</td>
<td>Discontinue lactation</td>
</tr>
<tr>
<td>Fluphenazine</td>
<td>Amount excreted in milk probably too small to be harmful; continue lactation; adverse effects possible; monitor infant for drowsiness</td>
</tr>
<tr>
<td>Glibenclamide</td>
<td>Hypoglycaemia in infant</td>
</tr>
<tr>
<td>Haloperidol</td>
<td>Amount excreted in milk probably too small to be harmful; continue lactation; adverse effects possible; monitor infant for drowsiness</td>
</tr>
<tr>
<td>Halothane</td>
<td>Excreted in milk</td>
</tr>
<tr>
<td>Hydralazine</td>
<td>Present in milk but not known to be harmful; monitor infant</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Use alternative drug; may inhibit lactation</td>
</tr>
<tr>
<td>Hydrocortisone</td>
<td>Systemic effects in infant unlikely with maternal dose of less than equivalent of prednisolone 40 mg daily; monitor infant’s adrenal function with higher doses</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>Amount too small to be harmful; short courses safe in usual doses Imipenem + Cilastatin Present in milk-avoid</td>
</tr>
<tr>
<td>Indinavir</td>
<td>Lactation recommended during first 6 months if no safe alternative to breast milk</td>
</tr>
</tbody>
</table>
Iodine
Stop lactation; danger of neonatal hypothyroidism or goitre; appears to be concentrated in milk

Isoniazid
Monitor infant for possible toxicity; theoretical risk of convulsions and neuropathy; prophylactic pyridoxine advisable in mother and infant

Ivermectin
Avoid treating mother until infant is 1 week old

Lamivudine
Present in milk; lactation recommended during first 6 months if no safe alternative to breast milk

Levamisole
Lactation contraindicated

Levonorgestrel
Combined oral contraceptives may inhibit lactation-use alternative method of contraception until weaning or for 6 months after birth; progestogen-only contraceptives do not affect lactation (preferably start 6 weeks after birth or later)

Lithium
Present in milk and risk of toxicity in infant; continue lactation; monitor infant carefully, particularly if risk of dehydration

Lopinavir + Ritonavir
Lactation recommended during first 6 months if no safe alternative to breast milk

Lumefantrine
See Artemether + Lumefantrine

Medroxyprogesterone
Present in milk-no adverse effects reported (preferably start injectable contraceptive 6 weeks after birth or later)

Mefloquine
Present in milk but risk to infant minimal 6-Mercaptopurine Lactation contraindicated

Metformin
Present in milk but safe in usual doses; monitor infant

Methotrexate
Lactation contraindicated

Metoclopramide
Present in milk; adverse effects possible; monitor infant for adverse effects

Metronidazole
Significant amount in milk; continue lactation; avoid large doses; use alternative drug if possible

Morphine
Short courses safe in usual doses; monitor infant

Nalidixic acid
Continue lactation; use alternative drug if possible; one case of haemolytic anaemia reported

Nelfinavir
Lactation recommended during first 6 months if no safe alternative to breast milk
<table>
<thead>
<tr>
<th>Drug</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neostigmine</td>
<td>Amount probably too small to be harmful; monitor infant</td>
</tr>
<tr>
<td>Nevirapine</td>
<td>Present in milk; lactation recommended during first 6 months if no safe alternative to breast milk</td>
</tr>
<tr>
<td>Nifedipine</td>
<td>Small amount in milk; continue lactation; monitor infant</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>Only small amounts in milk but could be enough to produce haemolysis in G-6-PD-deficient infants</td>
</tr>
<tr>
<td>Norethisterone</td>
<td>Combined oral contraceptives may inhibit lactation-use alternative method of contraception until weaning or for 6 months after birth; progestogen-only contraceptives do not affect lactation (preferably start injectable contraceptive 6 weeks after birth or later)</td>
</tr>
<tr>
<td>Ofloxacin</td>
<td>Continue lactation; use alternative drug if possible</td>
</tr>
<tr>
<td>Paracetamol</td>
<td>Small amount present in milk: short courses safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Pentamididine</td>
<td>Avoid unless essential</td>
</tr>
<tr>
<td>Pentavalent antimony compounds</td>
<td>Avoid</td>
</tr>
<tr>
<td>Phenobarbital</td>
<td>Continue lactation; adverse effects possible; monitor infant for drowsiness;</td>
</tr>
<tr>
<td>Phenoxy methylpenicillin</td>
<td>Trace amounts in milk; safe in usual dosage; monitor infant</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Small amount present in milk; continue lactation; adverse effects possible; monitor infant for drowsiness;</td>
</tr>
<tr>
<td>Potassium iodide</td>
<td>Stop lactation; danger of neonatal hypothyroidism or goitre; appears to be concentrated in milk</td>
</tr>
<tr>
<td>Povidone–iodine</td>
<td>Avoid; iodine absorbed from vaginal preparations is concentrated in milk</td>
</tr>
<tr>
<td>Praziquantel</td>
<td>Avoid lactation during and for 72 h after treatment; considered safe to continue lactation in treatment of schistosomiasis</td>
</tr>
<tr>
<td>Prednisolone</td>
<td>Systemic effects in infant unlikely with maternal dose of less than prednisolone 40 mg daily; monitor infant’s adrenal function with higher doses</td>
</tr>
<tr>
<td>Primaquine</td>
<td>Avoid; risk of haemolysis in G-6-PD-deficient infants</td>
</tr>
<tr>
<td>Procainamide</td>
<td>Present in milk; continue lactation; monitor infant</td>
</tr>
<tr>
<td>Procarbazine</td>
<td>Lactation contraindicated</td>
</tr>
</tbody>
</table>
Promethazine Safe in usual dosage; monitor infant for drowsiness
Propranolol Present in milk; safe in usual dosage; monitor infant
Propylthiouracil Monitor infant’s thyroid status but amounts in milk probably too small to affect infant; high doses might affect neonatal thyroid function
Pyrimethamine Significant amount-avoid administration of other folate antagonists to infant
Quinidine Significant amount but not known to be harmful
Ranitidine Significant amount present in milk, but not known to be harmful
Ritonavir See Lopinavir with Ritonavir
Salbutamol Safe in usual dosage; monitor infant
Saquinavir Lactation recommended during first 6 months if no safe alternative to breast milk
Senna Avoid; large doses may cause increased gastric motility and diarrhoea
Silver sulfadiazine Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants particularly with longacting sulphonamides, and of haemolysis in G-6-PD-deficient infants
Sodium valproate see Valproic acid
Stavudine Lactation recommended during first 6 months if no safe alternative to breast milk
Sulfadiazine Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants particularly with longacting sulphonamides, and of haemolysis in G-6-PD-deficient infants
Sulfadoxine + Pyrimethamine Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants and of haemolysis in G-6-PD-deficient infants (due to sulfadoxine)
Sulfamethoxazole + Trimethoprim Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants and of haemolysis in G-6-PD-deficient infants (due to sulfamethoxazole)
Sulfasalazine Continue lactation; monitor infant for jaundice-small amounts in milk (1 report of bloody diarrhoea and rashes);
The following table summarizes the safety of various medications during lactation:

<table>
<thead>
<tr>
<th>Medication</th>
<th>Lactation Status</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Promethazine</td>
<td>Safe in usual dosage; monitor infant for drowsiness</td>
<td>theoretical risk of neonatal haemolysis especially in G-6-PD-deficient infants</td>
</tr>
<tr>
<td>Propranolol</td>
<td>Present in milk; safe in usual dosage; monitor infant</td>
<td></td>
</tr>
<tr>
<td>Propylthiouracil</td>
<td>Monitor infant’s thyroid status but amounts in milk probably too small to affect infant; high doses might affect neonatal thyroid function</td>
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</tr>
<tr>
<td>Pyrimethamine</td>
<td>Significant amount-avoid administration of other folate antagonists to infant</td>
<td></td>
</tr>
<tr>
<td>Quinidine</td>
<td>Significant amount but not known to be harmful</td>
<td></td>
</tr>
<tr>
<td>Ranitidine</td>
<td>Significant amount present in milk, but not known to be harmful</td>
<td></td>
</tr>
<tr>
<td>Ritonavir</td>
<td>See Lopinavir with Ritonavir</td>
<td></td>
</tr>
<tr>
<td>Salbutamol</td>
<td>Safe in usual dosage; monitor infant</td>
<td></td>
</tr>
<tr>
<td>Saquinavir</td>
<td>Lactation recommended during first 6 months if no safe alternative to breast milk</td>
<td></td>
</tr>
<tr>
<td>Senna</td>
<td>Avoid; large doses may cause increased gastric motility and diarrhoea</td>
<td></td>
</tr>
<tr>
<td>Silver sulfadiazine</td>
<td>Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants particularly with longacting sulphonamides, and of haemolysis in G-6-PD-deficient infants</td>
<td></td>
</tr>
<tr>
<td>Sodium valproate</td>
<td>see Valproic acid</td>
<td></td>
</tr>
<tr>
<td>Stavudine</td>
<td>Lactation recommended during first 6 months if no safe alternative to breast milk</td>
<td></td>
</tr>
<tr>
<td>Sulfadiazine</td>
<td>Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants and of haemolysis in G-6-PD-deficient infants (due to sulfadoxine)</td>
<td></td>
</tr>
<tr>
<td>Sulfadoxine + Pyrimethamine</td>
<td>Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants and of haemolysis in G-6-PD-deficient infants (due to sulfamethoxazole)</td>
<td></td>
</tr>
<tr>
<td>Sulfamethoxazole + Trimethoprim</td>
<td>Continue lactation; monitor infant for jaundice-small risk of kernicterus in jaundiced infants and of haemolysis in G-6-PD-deficient infants (due to sulfamethoxazole)</td>
<td></td>
</tr>
<tr>
<td>Sulfasalazine</td>
<td>Continue lactation; monitor infant for jaundice-small amounts in milk (1 report of bloody diarrhoea and rashes); theoretical risk of neonatal haemolysis especially in G-6-PD-deficient infants</td>
<td></td>
</tr>
<tr>
<td>Tamoxifen</td>
<td>Suppresses lactation; avoid unless potential benefit outweighs risk</td>
<td></td>
</tr>
<tr>
<td>Testosterone</td>
<td>Avoid; may cause masculinization in the female infant or precocious development in the male infant; high doses suppress lactation</td>
<td></td>
</tr>
<tr>
<td>Tetracycline</td>
<td>Continue lactation; use alternative drug if possible (absorption and therefore discolouration of teeth in infant probably usually prevented by chelation with calcium in milk)</td>
<td></td>
</tr>
<tr>
<td>Theophylline</td>
<td>Present in milk-irritability in infant reported; modified-release preparations preferable</td>
<td></td>
</tr>
<tr>
<td>Thiamine</td>
<td>Severely thiamine-deficient mothers should avoid lactation as toxic methylglyoxal excreted in milk</td>
<td></td>
</tr>
<tr>
<td>Trimethoprim</td>
<td>Present in milk; safe in usual dosage; monitor infant</td>
<td></td>
</tr>
<tr>
<td>Valproic acid</td>
<td>Small amount present in milk; continue lactation; adverse effects possible; monitor infant for drowsiness; (Sodium valproate)</td>
<td></td>
</tr>
<tr>
<td>Vancomycin</td>
<td>Present in milk-significant absorption following oral administration unlikely</td>
<td></td>
</tr>
<tr>
<td>Vinblastine</td>
<td>Lactation contraindicated</td>
<td></td>
</tr>
<tr>
<td>Vincristine</td>
<td>Lactation contraindicated</td>
<td></td>
</tr>
<tr>
<td>Warfarin</td>
<td>Risk of haemorrhage; increased by vitamin-K deficiency; warfarin appears safe</td>
<td></td>
</tr>
<tr>
<td>Zidovudine</td>
<td>Lactation recommended during first 6 months if no safe alternative to breast milk</td>
<td></td>
</tr>
</tbody>
</table>
Appendix 7c
Pregnancy

Drugs can have harmful effects on the fetus at any time during pregnancy. It is important to remember this when prescribing for a woman of childbearing age. However, irrational fear of using drugs during pregnancy can also result in harm. This includes untreated illness, impaired maternal compliance, suboptimal treatment and treatment failures. Major congenital malformations occur in 2–4% of all live births, 15% of all diagnosed pregnancies will result in fetal loss. During the first trimester drugs may produce congenital malformations (teratogenesis), and the greater risk is from third to the eleventh week of pregnancy. During the second and third trimester, drugs may affect the growth and functional development of the fetus or have toxic effects on fetal tissues. Drugs given shortly before term or during labor may have adverse effects on labor or on the neonate after delivery. Few drugs have been shown conclusively to be teratogenic in man but no drug is safe beyond all doubt in early pregnancy. Screening procedures are available where there is a known risk of certain defects.

Prescribing in Pregnancy

Since, approximately 50% of pregnancies are unplanned and rest 50% are planned, if possible, counseling of women before a planned pregnancy should be carried out including discussion of risks associated with specific therapeutic agents, traditional drugs (alternative medicines), over the counter drugs and substances of abuse such as opioids, smoking, alcohol etc. Drugs should be prescribed in pregnancy only if the expected benefits to the mother are thought to be greater than the risk to the fetus. All drugs should be avoided if possible during the first trimester. Drugs which have been used extensively in pregnancy and appear to be usually safe should be prescribed in preference to new or untried drugs and the smallest effective dose should be used. Keeping in view the prevalence of irrational polypharmacy, emphasis should be laid on promoting the use of well known single component drugs to multicomponent drugs. Since, there does appear to be an association of very potent topical corticosteroids with low birth weight, even the dermatological drug products being used should be cautiously selected and used.

The pronounced and progressive change in drug disposition that occurs during pregnancy is another major reason which calls for attention. Major physiological changes which influence drug disposition in mother and fetus are:

<table>
<thead>
<tr>
<th>S. No</th>
<th>Physiologic changes</th>
<th>Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Plasma albumin concentration of mother is reduced</td>
<td>Drug protein binding alteration</td>
</tr>
<tr>
<td>2.</td>
<td>Increased body fat in mother</td>
<td>Distribution of drug is effected</td>
</tr>
<tr>
<td>3.</td>
<td>Increased hepatic metabolism in mother</td>
<td>Faster hepatic clearance</td>
</tr>
</tbody>
</table>
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The pronounced and progressive change in drug disposition that occurs during pregnancy is another major reason which calls for attention. Major physiological changes which influence drug disposition in mother and fetus are:

| 4. | Increased cardiac output in mother | Increased renal blood flow and glomerular filtration and hence, increased elimination of drug |
| 5. | Presence of placental barrier | Selectivity of drug permeation based on its hydrophobicity or molecular weight of drug |
| 6. | Drug metabolizing enzymes activity in fetal liver is very low | Slow elimination of drugs by fetus |

Though maternal medication carry the risk of increase in the incidence of abortion, stillbirths, fetal death, premature or delayed labor or create perinatal problems; but certain medications like folic acid are recommended for all pregnant women to reduce the rate of congenital anomalies specifically, the neural tube defect. The Food and Drug Administration has categorized the drug risks to the fetus that runs from: “Category A” (safest) to “Category X” (known danger--do not use!)

**Category A**

Controlled studies in women fail to demonstrate a risk to the fetus in the first trimester (and there is no evidence of a risk in later trimesters), and the possibility of fetal harm appears remote.

**Category B**

Either animal-reproduction studies have not demonstrated a fetal risk but there are no controlled studies in pregnant women, or animalreproduction studies have shown an adverse effect (other than a decrease in fertility) that was not confirmed in controlled studies in women in the first trimester (and there is no evidence of a risk in later trimesters).

**Category C**

Either studies in animals have revealed adverse effects on the fetus (teratogenic or embryocidal or other) and there are no controlled studies in women, or studies in women and animals are not available. Drugs should be given only if the potential benefit justifies the potential risk to the fetus.

**Category D**

There is positive evidence of human fetal risk, but the benefits from use in pregnant women may be acceptable despite the risk (e.g., if the drug is needed in a life-threatening situation or for a serious disease for which safer drugs cannot be used or are ineffective).

**Category X**

Studies in animals or human beings have demonstrated fetal abnormalities, or there is evidence of fetal risk based on human experience or both, and the risk of the use of the drug in pregnant women clearly outweighs any possible benefit. The drug is contraindicated in women who are or may become pregnant.
Dosing considerations in renal impairment

The number of patients with chronic kidney disease (CKD) and reduced renal function have been inexorably increasing. Reduced renal function may need adjustment in drug therapy as kidney plays a major role in the pharmacokinetics of a large number of drugs.

- Renal insufficiency frequently alters drug distribution volume. Edema and ascites increase the apparent volume of distribution of highly water-soluble or protein-bound drugs. Usual doses of such drugs given to edematous patients result in inadequate, low plasma levels.
- The alteration of plasma protein binding in patients with renal insufficiency is an important factor affecting both efficacy and toxicity. In patients with uremia the unbound fraction of several acidic drugs is substantially increased which may lead to serious toxicity.
- Although renal insufficiency is thought to affect primarily the renal elimination of drugs or metabolites, renal failure substantially affects drug biotransformation. Uremia slows the rate of reduction and hydrolysis reactions.
- Many active or toxic metabolites are produced during drug metabolism. Many of these metabolites depend on the kidneys for their removal from the body. The accumulation of active metabolites can explain in part the high incidence of ADRs seen in renal failure.

A few points should be kept in mind while prescribing:

- Renal function declines with age so that by the age of 80 it is half that in healthy young subjects.
- It is advisable to determine renal function not only before but also during the period of treatment and adjust the maintenance dose as necessary.
- One should try to keep drug prescription to minimum.
- Nephrotoxic drugs should, if possible, be avoided in all patients with renal disease because the nephrotoxicity is more likely to be serious.
- One should stay alert for unexpected ADRs.

The recommendations in the table below are meant only as a guide and do not imply efficacy or safety of a recommended dose in an individual patient.

A loading dose equivalent to the usual dose in patients with normal renal function should be considered for drugs with a particularly long half-life.

The table below gives the common drugs where in renal impairment dose adjustment is required.

When the dose method (D) is suggested, the percentage of the dose for normal renal function is given and when the interval method (I) is suggested, the actual dose interval is provided.
Dosing considerations in renal impairment

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The recommendations in the table below are meant only as a guide and do not imply efficacy or safety of a
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<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose Method</th>
<th>GFR &gt;50 (ml/min)</th>
<th>GFR 10-50 (ml/min)</th>
<th>GFR &lt;10 (ml/min)</th>
<th>CAPD</th>
<th>HD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetaminophen</td>
<td>I</td>
<td>q4h</td>
<td>q6h</td>
<td>q8h</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
</tr>
<tr>
<td>Acetazolamide</td>
<td>I</td>
<td>q6h</td>
<td>q12h</td>
<td>Avoid</td>
<td>No data</td>
<td>No data</td>
</tr>
<tr>
<td>Acetylsalicyclic</td>
<td>I</td>
<td>Q4h</td>
<td>Q4-6h</td>
<td>Avoid</td>
<td>As normal</td>
<td>As normal</td>
</tr>
<tr>
<td>Acid</td>
<td>D, I</td>
<td>5 mg/kg</td>
<td>5 mg/kg</td>
<td>2.5 mg/kg</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
</tr>
<tr>
<td>Acyclovir</td>
<td>D, I</td>
<td>5 mg/kg</td>
<td>q12-24h</td>
<td>q24h</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
</tr>
<tr>
<td>Allopurinol</td>
<td>D</td>
<td>75%</td>
<td>50%</td>
<td>33%</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
</tr>
<tr>
<td>Amikacin</td>
<td>D, I</td>
<td>60–90%</td>
<td>30–70%</td>
<td>20–30%</td>
<td>15–20 mg 5 mg/kg</td>
<td>5 mg/kg post HD</td>
</tr>
<tr>
<td>Amiloride</td>
<td>D</td>
<td>100%</td>
<td>50%</td>
<td>Avoid</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Aminophylline</td>
<td>D</td>
<td>100%</td>
<td>200–400</td>
<td>200–300</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
</tr>
<tr>
<td>Amphotericin</td>
<td>B I</td>
<td>q24h</td>
<td>q24h</td>
<td>q24-36h</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
</tr>
<tr>
<td>Ampicillin</td>
<td>I</td>
<td>q6h</td>
<td>q6–12h</td>
<td>q12-24h</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
</tr>
<tr>
<td>Cefazolin I</td>
<td>q8h</td>
<td>q12h</td>
<td>q24–48h</td>
<td>0.5 g</td>
<td>q12h 0.5–1.0 g</td>
<td>0.5–1.0 g post HD</td>
</tr>
<tr>
<td>Cefixime</td>
<td>D</td>
<td>00%</td>
<td>75%</td>
<td>50%</td>
<td>200 mg 200 mg</td>
<td>200 mg q24h dose post HD</td>
</tr>
<tr>
<td>Cefotaxime</td>
<td>I</td>
<td>100%</td>
<td>100%</td>
<td>50%</td>
<td>1 g q24h GFR &lt; 10</td>
<td>Dose as dose post HD</td>
</tr>
<tr>
<td>Drug</td>
<td>Route</td>
<td>Dose as GFR &lt; 10</td>
<td>Dose as GFR &lt; 10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------------</td>
<td>-------</td>
<td>------------------</td>
<td>------------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chloroquine</td>
<td>D</td>
<td>100%</td>
<td>100%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>D</td>
<td>100% 50-75% 50%</td>
<td>250 mg 250 mg q8h q12h</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cisplatin</td>
<td>D</td>
<td>100% 75% 50%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyclophosphamide</td>
<td>D</td>
<td>100% 75-100% 50-75%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dapsone</td>
<td></td>
<td>100% 100% 50%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Didanosine</td>
<td>I</td>
<td>100% 50% 25%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Digoxin</td>
<td>D, I</td>
<td>100% 25–75% 10–25%</td>
<td>Dose as Dose as q24h q36h q48h q12h</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Enalapril</td>
<td>D</td>
<td>100% 75-100% 50-75%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Erythromycin</td>
<td>D</td>
<td>100% 100% 50-75%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethambutol</td>
<td>I</td>
<td>q24h q24-36h q48h</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Etoposide</td>
<td>D</td>
<td>100% 75% 50%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fentanyl</td>
<td>D</td>
<td>100% 75% 50%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluconazole</td>
<td>D</td>
<td>100% 100% 50%</td>
<td>Dose as Dose as</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gentamicin</td>
<td>D, I</td>
<td>60–90% 30–70% 20–30%</td>
<td>3–4 mg/L/day q8–12h q12h q24–72h</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

HD: Hemodialysis; CAPD: Chronic Ambulatory Peritoneal Dialysis.
<table>
<thead>
<tr>
<th>Drug</th>
<th>Form</th>
<th>100%</th>
<th>50–150 mg qd</th>
<th>25 mg qd</th>
<th>Dose as</th>
<th>GFR &lt; 10</th>
<th>Dose as</th>
<th>GFR &lt; 10</th>
<th>dose post HD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chloroquine</td>
<td>D</td>
<td>100%</td>
<td>75%</td>
<td></td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>D</td>
<td>50%</td>
<td>100%</td>
<td>75%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Enalapril</td>
<td>D</td>
<td>75%</td>
<td>25–75%</td>
<td>10–25%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>D</td>
<td>100%</td>
<td>75%</td>
<td>50%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>D, I</td>
<td>50%</td>
<td>100%</td>
<td>75%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Lamivudine</td>
<td>D, I</td>
<td>100%</td>
<td>50–150 mg qd</td>
<td>25 mg qd</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Metformin</td>
<td>D</td>
<td>75%</td>
<td>50%</td>
<td>25%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Metoclopramide</td>
<td>D</td>
<td>100%</td>
<td>50–75%</td>
<td>25–75%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Penicillin</td>
<td>G, D</td>
<td>100%</td>
<td>75%</td>
<td>50%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td>D</td>
<td>100%</td>
<td>50%</td>
<td>25%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Quinine</td>
<td>I</td>
<td>50%</td>
<td>100%</td>
<td>75%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Streptomycin</td>
<td>I</td>
<td>75%</td>
<td>50%</td>
<td>25%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Triamterene</td>
<td>I</td>
<td>75%</td>
<td>50%</td>
<td>25%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Tubocurarine</td>
<td>D</td>
<td>75%</td>
<td>50%</td>
<td>25%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Vancomycin</td>
<td>D, I</td>
<td>75%</td>
<td>50–150 mg qd</td>
<td>25 mg qd</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
<tr>
<td>Zidovudine (AZT)</td>
<td>D, I</td>
<td>100%</td>
<td>50%</td>
<td>25%</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>Dose as</td>
<td>GFR &lt; 10</td>
<td>dose post HD</td>
</tr>
</tbody>
</table>

**HD:** Hemodialysis; **CAPD:** Chronic Ambulatory Peritoneal Dialysis.
Appendix 8

National Health Programmes (NHPs)

The area of government healthcare professionals covers Rural Medical Dispensary (RMD); Primary Health Centre (PHC); Community Health Centre (CHC); Urban Health Centre (UHC); district, Sub-divisional and Rural hospitals; large tertiary care and teaching hospitals and other public hospitals including Railways, ESIS, Coal India, CGHS, MPT, Airlines, Armed Forces, Jail hospitals, etc.

The government has been bringing health care to the public in special areas of concern through a top down approach through its various NHPs.

A list of NHPs is mentioned below:

1. National Vector Borne Disease Control Programme
2. National Filaria Control Programme
3. National Leprosy Eradication Programme
4. Revised National TB Control Programme
5. National Programme for Control of Blindness
6. National Iodine Deficiency Disorders Control Programme
7. National Mental Health Programme
8. National Aids Control Programme
9. National Programme on Prevention and Control of Cancer, Diabetes, Cardiovascular Disease and Stroke
10. Universal Immunization Programme
11. National Programme for Prevention and Control of Deafness
12. Pilot Programme on Prevention and Control of Diabetes, Cardiovascular Disease and Stroke
13. National Tobacco Control Programme
14. School Health Programme
15. Prevention and Control of Non-Communicable Diseases
## Appendix 9
### National Immunization Schedule

<table>
<thead>
<tr>
<th>Beneficiary</th>
<th>Age</th>
<th>Vaccine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonates</td>
<td>Birth</td>
<td>BCG* and OPV**</td>
</tr>
<tr>
<td>Infants</td>
<td>6 weeks</td>
<td>DPT, OPV and Hepatitis B</td>
</tr>
<tr>
<td></td>
<td>10 weeks</td>
<td>DPT, OPV and Hepatitis B</td>
</tr>
<tr>
<td></td>
<td>14 Weeks</td>
<td>DPT, OPV and Hepatitis B</td>
</tr>
<tr>
<td></td>
<td>9 months</td>
<td>Measles</td>
</tr>
<tr>
<td></td>
<td>9, 18, 24, 30, 36, 42, 48, 54 and 60 months</td>
<td>Vitamin A</td>
</tr>
<tr>
<td></td>
<td>18 months</td>
<td>DPT &amp; OPV (Booster dose)</td>
</tr>
<tr>
<td>Children</td>
<td>5 years</td>
<td>DT</td>
</tr>
<tr>
<td></td>
<td>10 years Tetanus toxoid</td>
<td></td>
</tr>
<tr>
<td>Adolescents</td>
<td>16 years</td>
<td>Tetanus toxoid</td>
</tr>
<tr>
<td>Pregnant women</td>
<td></td>
<td>Tetanus toxoid is given during pregnancy (0.5 ml intramuscularly). If there is no prior history of vaccination, 2 doses are administered, the first in 2nd trimester and the second dose one month later. If there is confirmed documentary evidence of proper and complete immunization during childhood, then a single booster dose is administered in the 2nd trimester.</td>
</tr>
</tbody>
</table>

* At birth or at the time of DPT/OPV;

** Dose called as Zero dose and can be given till 14 days of age, if missed early.
Pharmacogenetics refers to the genetic variation in drug response. This could be due to:
(a) Single mutant gene or genetic polymorphism.
(b) Polygenic influence.

However the later is not of much significance in Clinical Practice. Variations in drug responses amongst fraternal twins (dizygotic) may be relatively wide when compared to identical (monozygotic) twins. Importance of Pharmacogenetics lies in the development of new drugs from information available from human genome project. It aims at individualizing and improving precision of pharmacotherapy.

When polymorphic DNA sequence variation occurs in the coding region or regulatory regions of genes, it causes variation in gene product through alteration of activity, function or level of expression. The variation to drug response can also be brought about by:

- Metabolic variation
- Idiosyncratic reaction

As an example metabolic variation can be best explained by the varied metabolic response exhibited to the antitubercular drug Isoniazid due to the presence of two different phenotypes in a population. These varied phenotypes are expressed as larger or smaller amount of enzyme N-acetylase in liver, and the population being termed as rapid acetylators and slow acetylators respectively. Some of the pharmacogenetic conditions and the drugs involved are summarized below.

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Pharmacogenetic variation</th>
<th>Frequency of occurrence</th>
<th>Drugs involved</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Acetylator with race</td>
<td>Varies with race</td>
<td>Isoniazid - slow acetylator- rapid acetylator Procainamide Hydralazine Sulphas, Sulphones Phenezine</td>
<td>Neuropathy Hepatotoxicity SLE SLE ADR ADR Slower recovery from surgical paralysis Postural hypotension, Diplopia, blurred vision</td>
</tr>
<tr>
<td>2.</td>
<td>Butyrylcholinesterase enzymes</td>
<td>1:3000 caucasian</td>
<td>Suxamethonium</td>
<td></td>
</tr>
<tr>
<td>3.</td>
<td>Aromatic hydroxylase N-oxidation enzyme (aminoxydase)</td>
<td>1.5-9.0%</td>
<td>Debrisoquine Sparteine</td>
<td></td>
</tr>
</tbody>
</table>
Pharmacogenetics refers to the genetic variation in drug response. This could be due to:

(a) Single mutant gene or genetic polymorphism.
(b) Polygenic influence.

However the later is not of much significance in Clinical Practice. Variations in drug responses amongst fraternal twins (dizygotic) may be relatively wide when compared to identical (monozygotic) twins. Importance of Pharmacogenetics lies in the development of new drugs from information available from human genome project. It aims at individualizing and improving precision of pharmacotherapy.

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Some of the pharmacogenetic conditions and the drugs involved are summarized below.

<table>
<thead>
<tr>
<th>Variation</th>
<th>Frequency</th>
<th>Outcome</th>
<th>Drugs Involved</th>
</tr>
</thead>
<tbody>
<tr>
<td>G-6-PD deficiency</td>
<td>Varies</td>
<td>Haemolysis</td>
<td>Antimalarials</td>
</tr>
<tr>
<td></td>
<td>with race</td>
<td></td>
<td>Primaquine</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>Mepacrine</td>
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<td></td>
<td></td>
<td></td>
<td>Pamaquine</td>
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<td></td>
<td></td>
<td></td>
<td>Pentaquine</td>
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<td></td>
<td></td>
<td></td>
<td>Chloroquine</td>
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<td></td>
<td></td>
<td></td>
<td>Quinine</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Proguanil</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Pyrimethamine</td>
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<tr>
<td>Cardiovascular drugs</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Procainamide</td>
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<td></td>
<td></td>
<td></td>
<td>Quinidine</td>
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<td></td>
<td></td>
<td></td>
<td>Hydralazine</td>
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<td></td>
<td></td>
<td></td>
<td>Thiazide diuretics</td>
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<tr>
<td>Central Nervous System Drugs</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Methylldopa</td>
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<td></td>
<td></td>
<td></td>
<td>Benzhexol</td>
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<td></td>
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<td></td>
<td>Phenytoin</td>
</tr>
<tr>
<td>Anti-infectives</td>
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<td></td>
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<td></td>
<td>Dapsone</td>
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<td></td>
<td></td>
<td></td>
<td>Sulfacetamide</td>
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<td></td>
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<td></td>
<td>Sulfamethoxypyrimidine</td>
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<td></td>
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<td></td>
<td>Sulfanilamide</td>
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<td></td>
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<td></td>
<td>Sulfapyridine</td>
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<td></td>
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<td></td>
<td>Sulfasalazine</td>
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<td></td>
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<td></td>
<td>Sulfisoxazole</td>
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<td></td>
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<td></td>
<td>Sulfadiazine</td>
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<td></td>
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<td></td>
<td>Cotrimoxazole</td>
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<td></td>
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<td></td>
<td>Trimethoprim</td>
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<td></td>
<td></td>
<td></td>
<td>Chloramphenicol</td>
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<td></td>
<td></td>
<td></td>
<td>Ciprofloxacin</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Moxifloxacin</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Nalidixic acid</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>Ofloxacin</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Norfloxacin</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Nitrofurantoin</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Nitrofurazone</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Isoniazid</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Furazolidone</td>
</tr>
</tbody>
</table>

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Streptomycin
Antineoplastics
Doxorubicin
Rasburicase
Anthelmintics
Niridazole
Stibophen
Analgesics
Acetylsalicylic acid
Antipyrine
Antigout drugs
Probenecid
Colchicine
Antidote
Dimercaprol
Phenylhydrazine
Antimethemoglobinemic Agent
Methylene Blue
Antidiabetics
Glibenclamide
Antihistamines
Diphenhydramine
Tripelennamine
Antazoline
Hormonal contraceptives
Mestranol
Vitamins
Ascorbic acid
Menadione
Diagnostic agent for cancer
Toluidine blue

5. Calcium release channel (ryanodine receptor) 1:20,000
   Halothane  Malignant hyperthermia,
   Calcium release
6. Narrow iridocorneal angle 5% US population
   Corticosteroids  Attack of angle closure glaucoma
   narrow iridocorneal angle
7. Hb variants Rare
   Oxidizing agents like quinolones
   Hb variants

8. Hepatic porphyrias
   Rare Haem-containing Acute porphyria
   hepatic oxidizing (GIT, CNS, CVS enzyme inducers symptoms)
   like barbiturates, Sulphonamides Sulphanylureas etc.

9. Altered receptor 2 large Warfarin Warfarin resistance
   or enzyme pedigrees in liver with increased affinity
   Warfarin resistance
10. Mixed function Only 1 Dicoumarol Dicoumarol oxidase in liver small Phenacetin sensitivity
    Mixed function Dicoumarol oxidase induced cardiovascular death
11. N-oxidation 5% Sparteine Sparteine -induced enzyme diplopia, blurred vision, overstimulated
    N-oxidation Sparteine
12. Mixed function 25% Tolbutamide Tolbutamide oxidase induced cardiovascular death
    Mixed function Tolbutamide oxidase induced cardiovascular death
13. Cytochrome Ondansetron Ondansetron P450 2D6 - lesser efficacy in ultrarapid metabolisers
    Cytochrome Ondansetron P450 2D6 - lesser efficacy in ultrarapid metabolisers
14. Tramadol Lesser efficacy of tramadol Codeine Codeine - poor analgesia
    Tramadol Therapeutic failure of Tamoxifen in poor metabolisers ~ 7% caucasians Debrisoquine Poor metabolism
    Tramadol Therapeutic failure of Tamoxifen in poor metabolisers ~ 7% caucasians Debrisoquine Poor metabolism

407
8. Hepatic porphyrias  
   Rare  
   Haem-containing hepatic oxidizing enzyme inducers like barbiturates, Sulphonamides Sulphany lureas etc.  
   Acute porphyria (GIT, CNS, CVS symptoms)

9. Altered receptor or enzyme in liver with increased affinity for vitamin K  
   2 large pedigrees  
   Warfarin  
   Warfarin resistance

10. Mixed function oxidase in liver microsomes  
    Only 1 small pedigree  
    Dicoumarol  
    Warfarinin sensitivity  
    Methemoglobinemia

11. N-oxidation enzyme (aminoxydase)  
    5%  
    Sparteine  
    Sparteine-induced diplopia, blurred vision, overstimulated uterus

12. Mixed function oxidase  
    25%  
    Tolbutamide  
    Tolbutamide induced cardiovascular death

13. Cytochrome P450 2D6  
    Ondansetron  
    Ondansetron - lesser efficacy in ultrarapid metabolisers

    Tramadol  
    Lesser efficacy of tramadol

    Codeine  
    Codeine - poor analgesia

    Tamoxifen  
    Therapeutic failure of Tamoxifen in poor metabolisers

    ~7% caucasians  
    Debrisoquine  
    Poor metabolism of Debrisoquine
<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>14.</td>
<td>Cytochrome P450 2C9 –</td>
<td>Phenytoin*</td>
<td>Decreased hydroxylation of Phenytoin</td>
</tr>
<tr>
<td>15.</td>
<td>Cytochrome Warfarin and Vitamin K epoxide reductase complex subunit 1</td>
<td>Warfarin</td>
<td>Longer times to dose stabilisation and higher risk of serious and life threatening bleeding</td>
</tr>
<tr>
<td>16.</td>
<td>Increased expression of p-glycoprotein</td>
<td>Chloroquine, anticancer drugs</td>
<td>Development of resistance</td>
</tr>
<tr>
<td>17.</td>
<td>An enzyme or receptor site with altered affinity for vitamin K</td>
<td>Rare</td>
<td>Simultaneous administration of inducing agents with warfarin</td>
</tr>
<tr>
<td>18.</td>
<td>Thiopurine methyl transferase enzyme</td>
<td>3%</td>
<td>Azathioprine 6-mercaptopurine</td>
</tr>
<tr>
<td>19.</td>
<td>Uridine-5-diphosphoglucuronosyltransferase 1A1</td>
<td>61% Caucasians 84% Asians 47% African americans</td>
<td>Irinotecanq</td>
</tr>
<tr>
<td>20.</td>
<td>Dihydropyrimidine dehydrogenase</td>
<td>5-fluorouracil</td>
<td>Risk of severe toxicity</td>
</tr>
<tr>
<td>21.</td>
<td>α-Thalessemia</td>
<td>Artesunate</td>
<td>Rise in plasma drug concentration</td>
</tr>
<tr>
<td>22.</td>
<td>β-Thalessemia</td>
<td>Somatomedin</td>
<td>May depress somatomedin activity</td>
</tr>
</tbody>
</table>

Appendix 11
Pharmacovigilance Programme of India

To provide safe and effective health care system in India and promote rational use of medicines, the Pharmacovigilance Programme of India (PvPI) has been established by the Ministry of Health and Family Welfare, Government of India. The programme is being coordinated by the Indian Pharmacopoeia Commission, Ghaziabad as a National Coordination Centre (NCC). The mission of the programme is to ensure that the benefits of use of medicine outweigh the risk and thus safeguard the health of the Indian population.

The PvPI has the following objectives:

- Monitoring Adverse Drug Reactions (ADRs) in Indian population
- Creating awareness amongst health care professionals about the importance of ADR reporting in India
- Monitoring benefit-risk profile of medicines
- To generate independent, evidence-based recommendations on the safety of medicines
- Support the Central Drugs Standard Control Organization (CDSCO) for formulating safety related regulatory decisions for medicines.

The PvPI is being supported by ADRs Monitoring Centres from across the country. This programme is on expansion mode by enabling more centres to join the reporting of ADRs to NCC and in turn to be linked up to Uppsala Monitoring Centre, WHO Collaborating Centre for International Drug Monitoring, Uppsala, Sweden. All stakeholders using this formulary are encouraged to report ADRs by downloading the Form (Suspected Adverse Drug Reaction Reporting Form) either from the website of the Central Drugs Standard Control Organization (www.cdsco. nic.in) or the Indian Pharmacopoeia Commission (www.ipc. gov.in).

Integrated efforts from the regulatory authorities, pharmaceutical industry and healthcare workers are necessary for the success and effectiveness of this programme. Therefore the active participation of the all concerned in this programme will not only be useful for Indian health care system but also facilitate rational prescribing globally.

A specimen form is attached at the end of the book.
Appendix 12
Pictograms

Pictorial Labeling

In spite of the best efforts there are many patients who are not so literate and may forget the instructions given and are not able to read the prescriptions even if given. They may need to repeatedly refer to the instructions and they may still need help. For taking care of such patients, use of pictorial labeling is recommended. While there has not been any national standard of pictorial label adopted so far, based on a project by Delhi Pharmaceutical Trust, in collaboration with Apothecaries Foundation the following pictorial labels that were developed and used and found appropriate, are recommended for use. Such labels can be pre-printed using self-adhesive sticker label and made available in Pharmacies. Pharmacists should paste relevant pictorial label either on the bottle/pack of the drugs dispensed or on the prescription sheet against each drugs prescribed. Based on the most commonly needed instructions, 11 types of messages were identified and pictorial labels have been developed. More instructions if required can be added to this list and additional labels need to be developed.

Pictorial Labels
Appendix 13
Principles of Dose Calculation in Special Conditions

A. Dosing considerations for the pediatric patient

Determination of a safe and effective drug dose for the pediatric patient is essential for the treating physician. Doses and dosing intervals in children differ from that of an adult because of age-related variations in drug absorption, distribution, metabolism, and elimination. Oral drug absorption matures by four to five months of age. Drugs like phenytoin and chloramphenicol are absorbed slowly and erratically whereas penicillin and ampicillin are absorbed more efficiently than in the adults because of a higher gastric pH in the neonates. Most drug metabolizing enzymes are expressed at low levels at birth followed by postnatal induction of specific isoenzymes. For most drugs including phenytoin, barbiturates, digoxin and analgesics the plasma half lives are 2-3 times longer in neonates as compared to adults. Renal elimination of drugs is also reduced in the neonates. As a result, neonatal dosing regimens for a number of drugs must be reduced to avoid toxicity. Drug pharmacodynamics may also be different in children, for e.g. antihistamines and barbiturates that are generally sedative in adults may be excitatory in pediatric age group. Similarly, specific drug toxicities may be unique to this age group as evident in case of tetracyclines affecting teeth and glucocorticoids reducing linear growth of bones.

Because of these maturational differences in infants and children, simple proportionate reduction in the adult dose may not be adequate to determine an optimal pediatric dose. The most reliable dose information is usually the one provided by the drug manufacturer in the package insert or pediatric doses listed in the formulary. However, such information is not available for the majority of drugs since proper dose optimization studies are often not performed in the pediatric age range. Consequently, initial doses are derived by scaling down the dosages used in adults and then titrating according to clinical response.

In the absence of specific pediatric dose recommendations, an estimate can be made by any of several methods based on age, weight, or surface area.

Age-based rules

Various rules of dosage in which the pediatric dose is a fraction of adult dose based on relative age have been used. Two of these are mentioned below.

Young’s rule (for children 2 years and older)

\[
\text{Child’s dose (approx.)} = \frac{\text{Age (years)}}{\text{Age(years) + 12}} \times \text{Adult dose}
\]
Fried’s rule (for children up to 2 years old)

\[
\text{Child's dose (approx.)} = \frac{\text{Age (months)}}{150} \times \text{Adult dose}
\]

**Weight based rule**

Because of large variability in weight among children of same age group, estimation of drug dosage for children on the basis of body weight is considered more reliable than that based solely on age. A rule proposed by Professor A. J. Clark (known as the Clark’s rule) introduced weight proportional regimen for drug therapy.

\[
\text{Child's dose (approx.)} = \frac{\text{Weight (kg)}}{70} \times \text{Adult dose}
\]

**Body surface area based rule**

The most dependable methods for calculation of pediatric drug doses are those based on body surface area (BSA). Rate of metabolism and redistribution of drug, organ size, blood volume, extracellular fluid volume, renal blood flow and assays of blood concentration of drugs correlate closely with the BSA.

\[
\text{Child's dose (approx.)} = \frac{\text{Weight (kg)}}{1.73 \text{ m}^2} \times \text{Adult dose}
\]

For calculation of doses based on BSA, standard nomogram which includes both body weight and height as factors determining BSA should be used. To calculate a child’s BSA, draw a straight line from the height column to the weight column. The point at which the line intersects the surface area (SA) column is the BSA (m²). If the child is of roughly normal proportion, BSA can be calculated from the weight alone (in the enclosed area).
Fried's rule (for children up to 2 years old)

\[ \text{Child's dose (approx.)} = x \times \text{Adult dose} \]

Because of large variability in weight among children of same age group, estimation of drug dosage for children on the basis of body weight is considered more reliable than that based solely on age. A rule proposed by Professor A. J. Clark (known as the Clark’s rule) introduced weight proportional regimen for drug therapy.

\[ \text{Weight (kg)} \]

\[ \text{Child's dose (approx.)} = x \times \text{Adult dose} \]

The most dependable methods for calculation of pediatric drug doses are those based on body surface area (BSA). Rate of metabolism and redistribution of drug, organ size, blood volume, extracellular fluid volume, renal blood flow and assays of blood concentration of drugs correlate closely with the BSA.

\[ \text{Weight (kg)} \]

\[ \text{Child's dose (approx.)} = x \times \text{Adult dose} \]

\[ 1.73 \text{ m}^2 \]

For calculation of doses based on BSA, standard nomogram which includes both body weight and height as factors determining BSA should be used. To calculate a child's BSA, draw a straight line from the height column to the weight column. The point at which the line intersects the surface area (SA) column is the BSA (m²). If the child is of roughly normal proportion, BSA can be calculated from the weight alone (in the enclosed area).

Note: This nomogram was published in Nelson Textbook of Pediatrics, 18th Edition, Richard E. Behrman, Robert M. Kliegman, MD, Hal B. Jenson, MD and Bonita F. Stanton, MD, Nomogram for the estimation of surface area, page no. 2951, fig no. 715-1, W. B. Saunders Company, 2007 and has been reproduced with permission.
The above mentioned rules are helpful in situations requiring the use of a drug that is unlicensed in children and for which no pediatric prescribing information is available. However, these rules are not precise and doses should not be calculated if it is possible to obtain the actual pediatric dose. Whatever be the method chosen to calculate the child’s dose, it should never exceed that of the adult.

B. Dosing considerations for the geriatric patient

Aging is a natural process of human development and is characterized by a progressive loss of physiologic and reproductive functions. Altered response to drugs with aging occurs at both pharmacokinetic and pharmacodynamic levels.

Pharmacokinetic changes occur with the age as a result of the inevitable anatomical and physiological changes which occur with time, such as loss of an organ’s functional units (nephrons, neurons) and disruption of some regulatory processes between cells and organs, resulting in decrease in function of body systems. For example, first pass metabolism decreases due to decrease in liver mass and blood flow, resulting in an increase in bioavailability of drugs which undergo extensive first pass metabolism, for example, propranolol. Another example of a pharmacokinetic change is the reduced clearance of renally-cleared drugs due to reduced renal plasma flow and glomerular filtration. This increases the potential for toxic effects particularly with those drugs where even marginal accumulation can have toxic effects, for example digoxin and lithium. Changes in body composition such as increase in body fat proportion and decrease in total body water result in a decreased volume of distribution for water soluble drugs such as digoxin, which increases their serum concentrations and potential for adverse effects.

Geriatric patients are much more “sensitive” to the action of many drugs, implying a change in the pharmacodynamic interactions of the drugs with their receptors. Elderly are more sensitive to some sedative-hypnotics and analgesics. Certain homeostatic control mechanisms appear to be blunted in elderly. Since homeostatic responses are often important components of the total response to a drug, these physiological alterations may change the pattern or intensity of drug response.

The age-related changes in the functions and composition of the human body require adjustments of drug selection and dosage for old individuals. Drug excretion via the kidneys declines with age, the elderly should therefore be treated as renally insufficient patients. A rough estimate of creatinine clearance can be obtained from the Cockcroft-Gault formula:

\[
\text{Creatinine clearance} = \frac{\text{(140 - Age)} \times \text{(Weight in kg)}}{72 \times \text{serum creatinine in mg/dL}} \quad \text{(for males)}
\]

For females, the result is multiplied by 0.85. The formula is applicable to patients between the age of 40 and 80.

The metabolic clearance is primarily reduced with drugs that display high hepatic extraction (‘blood flow-limited metabolism’), whereas the metabolism of drugs with low hepatic extraction (‘capacity-limited metabolism’) usually is not diminished. Reduction of metabolic drug elimination is more pronounced in malnourished or frail subjects. The water content of the aging body decreases, the fat content rises, hence
the distribution volume of hydrophilic compounds is reduced in the elderly, whereas that of lipophilic drugs is increased. Intestinal absorption of most drugs is not altered in the elderly. Aside of these pharmacokinetic changes, one of the characteristics of old age is a progressive decline in counterregulatory (homeostatic) mechanisms. Therefore drug effects are mitigated less, the reactions are usually stronger than in younger subjects, the rate and intensity of adverse effects are higher. Examples of drug effects augmented in this manner are, postural hypotension with agents that lower blood pressure, dehydration, hypovolemia, and electrolyte disturbances in response to diuretics, bleeding complications with oral anticoagulants, hypoglycemia with antidiabetics, and gastrointestinal irritation with non-steroidal anti-inflammatory drugs. The brain is an especially sensitive drug target in old age. Psychotropic drugs but also anticonvulsants and centrally acting antihypertensives may impede intellectual functions and motor coordination. The antimuscarinic effects of some antidepressants and neuroleptic drugs may be responsible for agitation, confusion, and delirium in elderly. Hence drugs should be used very restrictively in geriatric patients. If drug therapy is absolutely necessary, the dosage should be titrated to a clearly defined clinical or biochemical therapeutic goal starting from a low initial dose.
1. Definition and Concept

1.1. Storage
The term used to describe the safe keeping of all finished drugs and pharmaceuticals awaiting dispatch. The term is also applied for safe stores in hospitals and dispensaries under the specified conditions, so as to maintain their quality and potency.

1.2. Storage Conditions
The condition specified for storing the product e.g. temperature, humidity, container etc.

1.3. Quality
The ability of drug product to satisfy the users need.

1.4. Dosage Form
Refers to the gross physical form in which a drug is administered to or used by a patient.

1.5. Drug Product
A dosage form containing one or more active therapeutic ingredients along with other substance included during manufacturing process.

1.6. Finished Product
A medicinal product which has completed all stages of manufacture including packaging.

1.7. Strength
The concentration of the drug substance (for example weight/weight, weight/volume or unit dose/volume basis) and the potency i.e. the therapeutic activity of the drug product as indicated by appropriate laboratory tests or by adequately developed and controlled clinical data (expressed, for example in terms of units by reference to a standard).

1.8. Stability
Degree of resistance to chemical and physical changes, the efficacy of the preparation must remain constant or change only within the limit specified by official compendia.

1.9. Expiration Date
The date placed on the immediate container label of a drug product that designates the date through which the product is expected to remain within specifications. Kinetically it is the time required for 10% of the material to disappear.
2. Storage Procedure and Instructions

Drugs must be stored under conditions which minimize deterioration, contamination or damage. They must be stored under conditions compatible with their recommended storage requirements of temperature and humidity and where necessary to comply with legal requirements, under secured or segregated conditions.

**Appropriate storage conditions are:**

Temperature or humidity controlled environment must be equipped with suitable indicators, recorders and/or failure warning devices which must be checked at appropriate intervals and the results are coded. Recording thermometers should be used. Temperature in uncontrolled storage products should also be monitored.

Temperature should be measured at different levels in the warehouse and if necessary storage of sensitive drugs should be restricted to locations in the warehouse where they will be protected from extreme conditions. Temperatures of the refrigerators, deep freezers, and Relative Humidity in humidity control area as well as general areas of storage at room temperature should be recorded on a daily basis.

**Storage conditions not related to temperature are indicated in following terms:**

Drug storage should be regularly checked for cleanliness and good order and for misplaced/deteriorated/ out dated stock. All stocks should be checked regularly for obsolescence and degradation. Drugs with expired shelf life should be destroyed unless an extension of shelf life is granted following the satisfactory results or re-analysis. All due precautions should be observed to preclude issues of outdated Drugs.

Some categories of supplies require special storage conditions which include vaccines, narcotics, and combustibles e.g. vaccines require both refrigerator and freezers. Narcotics and other controlled substances should be kept in secure locking rooms with only one entrance. The keys should be kept in a secure place, preferably a safe. Only the warehouse director and one another person should have access to them.

3. Inspection for Deterioration

Pharmacists should be aware that deterioration of drug product may happen even before their expiration. This may occur perhaps due to improper storage or the fact that the product may require critical storage conditions not stated on

**Appendix 14**

the label. Hence inspection should include frequent product examination to detect signs of product deterioration which differ according to dosage form. Some examples, where deterioration may be physically detected are given here. The Pharmacists in the Stores should prepare an exhaustive list of following deterioration/spoilage indicators and keep them.
3.1. Liquid Dosage Forms
Slight gradual discolouration, Swirly precipitation, Whickering: pin hole at ampoule tip that leaks solution which precipitate or crystalline solid matter, clouding, fading of colour, Cake sedimentation (suspension), Creaming and cracking (emulsion), Discolouration.

3.2. Semisolid Dosage Forms
Ointments creams, gels and suppositories - Change in consistency and feel to touch, Phase separation, Discolouration, Surface crystal growth

3.3. Solid Dosage Forms
Surface chipping or pitting (plain tablets), Deformation (capsules), Increased hardness, Discolouration, Colour fading (coloured tablets), Chipping of coat (coated tablets).

Most vitamins, hormones enzymes are highly sensitive to oxidation and photo decomposition.
The integrity of packaging of dosage form is one of the important tasks of inspection for pharmacist as these protect the drug in a tailored fashion.

After each inspection, products showing any signs of instability should be subjected to sample analysis to ensure quality.

4. Drug Products Requiring Special Storage Conditions

4.1. Aerosols
Aerosols should be stored in a clean separate area away from heat and sunlight because the container contents are under pressure, filled containers must be checked for weight loss over the expiration dating period, for contents under pressure. The label should display “Do not expose to heat or store at a temperature above 40⁰C, keep out of reach of children”.

4.2. Creams
Creams can be destroyed under extreme temperature fluctuations hence they should be stored at temperature above 10oC and not exceeding 30⁰C. If the creams are opened and diluted they should not be kept for more than 14 days to avoid microbial contamination.

4.3. Ophthalmic Solutions and Drops
They should be stored according to the conditions specified on the label. After opening they should not be used for more than one month at home and not more than 15 days in hospitals.

4.4. Capsules
Extremes of humidity and temperature should be avoided. High humidity (> 60% RH) at 21ºC to 24ºC produce more lasting effects. Capsules become softer, tackier and blotted. If temperature is increased the capsule shells may melt and fuse together. High temperature (>40ºC) in dry place may cause cracking of capsule shell therefore capsules should be stored in air-conditioned area in which the humidity does not exceed 45% RH at 21 to 24ºC.
4.5. Suppositories

Suppositories should be protected from heat and preferably stored in the refrigerator. Polyethylene glycol suppositories and suppositories enclosed in solid shell are less prone to distortion at temperature slightly above body temperature. Glycerinated gelatin suppositories should be protected from heat, moisture and dry air by packaging in well sealed containers and storing in a cold place.

4.6. Vaccines

Liquid vaccines are to be stored between 2⁰ - 8⁰C and should not be frozen. All lyophilized vaccines should be stored between 2⁰ - 8⁰C and for long term storage can be kept at or below -20⁰C or otherwise as specified in the individual monographs. Oral polio should be stored in a freezer -2⁰ to -18⁰C.

5. Communicating the Prescription to the Patient

It is important that the drugs reach the patient in good and potent conditions and the patient should know and understand fully how to keep them till they are consumed. It is equally important that the patient should know the way each medicine is used. This will improve compliance and health outcome desired by the physician.

Communicating how and where to store the drugs to the Patient:

The following table may be used to guide and provide information on the way to store the drugs when they are dispensed to the patients. This is based on the recommended storage conditions as given on the labels of the drug products and Indian Pharmacopoeial notes in the General Chapters.

<table>
<thead>
<tr>
<th>On the label</th>
<th>Meaning</th>
<th>Tell the Patient/ Representative of the Patient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do not store over 8⁰C</td>
<td>To be stored in refrigerator (from +2⁰C to +8⁰C)</td>
<td>Keep in the General Compartment of the refrigerator and do not keep in the place where you make Ice.</td>
</tr>
<tr>
<td>Do not store over 30 ⁰C</td>
<td>To be stored at room temperature (from +2⁰C to +30⁰C)</td>
<td>Keep in any part of the house, except in Bath room/Kitchen. Do not keep near or in the window area.</td>
</tr>
<tr>
<td>Do not freeze</td>
<td>To be kept in refrigerator (from +2⁰C to +8⁰C but not in the freezer chamber)</td>
<td>Keep in the General Compartment of the refrigerator and do not keep in the place where you make Ice.</td>
</tr>
</tbody>
</table>
Transit period care and Use of Cool Packs:

It is equally important to ensure that patients who carry drugs requiring special storage conditions like anti-cancer drugs, several types of insulins, vaccines, sera, toxoids, would need to carry them in cold conditions till they reach the place where they will keep for some time before usage or to another hospital/nursing home till it is administered. In such cases during transit they need to be packed in “Thermo cool boxes with lid”, (#) with the drug product packs kept surrounded by adequate number of “Cool Packs”. (#) “Cool Packs are available which come ready filled with such special liquid in sealed bags or plastic packs, which on keeping overnight in freezer compartment of a refrigerator becomes solid ice. Such packs help in keeping the drug products in the box retain temperatures below 8°C for as much as 8 to 10 hours, which is generally adequate for transit protection. In case such cool packs are not available, it is recommended to use normal “Hot cases” (#) that people use to carry food, but stuffing the inside of the hot case boxes with sufficient ice cubes surrounding the drug packs kept inside, and the hot case suitably closed and sealed with sealing tapes. Cool packs can also be made by packing sufficient ice cubes into suitable sized self sealing polybags. (#) Several Pharmacists are known to innovate this way and they do serve for short transit times of up to one to two hours.
Appendix 15
Therapeutic Drug Monitoring

Therapeutic drug monitoring (TDM) is defined as measurement of drug levels in the biological fluids usually blood (serum or plasma). It has been carried out in saliva, urine, sweat, tear fluids etc also. It is carried out for specific drugs at various time intervals in order to maintain a relatively constant concentration of the particular drug in the bloodstream and to optimize drug therapy. The main focus of TDM is on drugs with narrow therapeutic range. Apart from this, it also plays a significant role for drugs having large inter-individual variations; relatively toxic drugs used in concomitant disease conditions, for escalation of dose, drugs showing wide variation in their metabolism, major organ failure, poisoning cases, failure of therapeutic response, to enhance patient compliance, etc. It is very important in such situations in which the drugs are to be taken on chronic or life long basis (chronic disease conditions such as bipolar disorder, organ transplant rejection, neurological disorders etc.). The timing and frequency of blood collection after the medication and correct interpretation of results of analysis and their correlation with clinical features ensures the best therapeutic outcome.

Indications for drug monitoring:

- Drugs whose efficacy is difficult to establish clinically, like Phenytoin.
- Drugs with a narrow therapeutic index. Examples: Lithium, phenytoin, digoxin.
- Patients who have impaired clearance of a drug with a narrow therapeutic index. Example: Patients with renal failure have decreased clearance of digoxin and therefore are at a higher risk of toxicity.
- Drugs whose toxicity is difficult to distinguish from a patient’s underlying disease. Example: Patients with chronic obstructive pulmonary disease treated with theophylline.

When not to do TDM

1. Drugs whose pharmacological effects can easily be used to dose titration, like oral hypoglycemic agents, anti-hypertensive drugs.
2. When easier and/or cheaper methods/alternatives to TDM are available to titrate the drug like International normalized ratio(INR) for warfarin.

Time of sample collection

1. Sample should be collected after steady state has been reached (5 half lives), unless TDM is intended to predict toxicity after single dose.
2. Usually “trough” concentrations are measured by taking the sample just before the subsequent dose.
3. Drugs whose half-lives are much shorter than the dosing interval, the peak and trough levels may be indicated to evaluate the dosage of drugs. Example: Gentamicin

TDM could be affected because of one or more of the factors relating to pharmacokinetics of the drug, or drug administration, or sample collection. Renal and hepatic alterations to half-life must also be considered. Laboratory variations also affect the TDM.

The following table summarizes the therapeutic concentration range of various drugs

**Table: Important drugs requiring therapeutic monitoring**

<table>
<thead>
<tr>
<th>S. No</th>
<th>Pharmacological category</th>
<th>Drugs</th>
<th>Therapeutic drug conc. range</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Drugs acting on cardiovascular system</td>
<td>Amiodarone, Digoxin, Procainamide</td>
<td>1.0 - 2.5 µg/ml, 0.8-2.0 ng/ml, 4.0-10.0 µg/ml</td>
</tr>
<tr>
<td>2.</td>
<td>Antibiotics</td>
<td>Gentamycin, Amikacin, Vancomycin, Tobramycin</td>
<td>5.0-10.0 µg/ml, 15.0-25.0 µg/ml, 15.0-25.0 µg/ml, 5.0-10.0 µg/ml</td>
</tr>
<tr>
<td>3.</td>
<td>Antiepileptics</td>
<td>Phenobarbital, Phenytoin, Valproic acid, Carbamazepine, Ethosuximide, Gabapentin, Lamotrigine</td>
<td>15.0-40.0 µg/ml, 10.0-20.0 µg/ml, 50.0-100.0 µg/ml, 5.0-12.0 µg/ml, 40.0-100.0 µg/ml, 2.0-20.0 µg/ml, 4.0-18.0 µg/ml</td>
</tr>
<tr>
<td>4.</td>
<td>Immunosuppressants</td>
<td>Cyclosporine, Tacrolimus, Sirolimus, Mycophenolate, Mofetil</td>
<td>50.0-300.0 µg/ml, 5.0-20.0 µg/l, 5.0 - 15.0 µg/l, 1.0 - 60.0 mg/l</td>
</tr>
<tr>
<td>5.</td>
<td>Psychopharmacological agents</td>
<td>Lithium, Imipramine, Amitriptyline, Nortriptyline, Desipramine, Clozapine</td>
<td>0.8-1.2 mEq/l, 0.15-0.3 µg/ml, 0.12-0.15 µg/ml, 0.05-0.15 µg/ml, 0.15-0.3 µg/ml, 0.35 to 0.6 mg/l</td>
</tr>
<tr>
<td>6.</td>
<td>Anti-infective</td>
<td>Cycloserine, Ethambutol, Pyrazinamide, Streptomycin</td>
<td>20-35 µg/ml, 2.0-6.0 µg/ml, 20.0-50.0 µg/ml, 35.0-45.0 µg/ml</td>
</tr>
</tbody>
</table>
Drugs whose half-lives are much shorter than the dosing interval, the peak and trough levels may be indicated to evaluate the dosage of drugs. Example: Gentamicin TDM could be affected because of one or more of the factors relating to pharmacokinetics of the drug, or drug administration, or sample collection. Renal and hepatic alterations to half-life must also be considered. Laboratory variations also affect the TDM.

The following table summarizes the therapeutic concentration range of various drugs:

<table>
<thead>
<tr>
<th>Important drugs requiring therapeutic monitoring</th>
<th>Table</th>
</tr>
</thead>
</table>

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