ABBREVIATIONS

Abbreviations used in WHO documentation include the following:

- ACHR – Advisory Committee on Health Research
- ASEAN – Association of Southeast Asian Nations
- CEB – United Nations System Chief Executives Board for Coordination (formerly ACC)
- CIOMS – Council for International Organizations of Medical Sciences
- FAO – Food and Agriculture Organization of the United Nations
- IAEA – International Atomic Energy Agency
- IARC – International Agency for Research on Cancer
- ICAO – International Civil Aviation Organization
- IFAD – International Fund for Agricultural Development
- ILO – International Labour Organization (Office)
- IMF – International Monetary Fund
- IMO – International Maritime Organization
- INCB – International Narcotics Control Board
- ITU – International Telecommunication Union
- OECD – Organisation for Economic Co-operation and Development
- OIE – Office International des Epizooties
- PAHO – Pan American Health Organization
- UNAIDS – Joint United Nations Programme on HIV/AIDS
- UNCTAD – United Nations Conference on Trade and Development
- UNDCP – United Nations International Drug Control Programme
- UNDP – United Nations Development Programme
- UNEP – United Nations Environment Programme
- UNESCO – United Nations Educational, Scientific and Cultural Organization
- UNFPA – United Nations Population Fund
- UNHCR – Office of the United Nations High Commissioner for Refugees
- UNICEF – United Nations Children’s Fund
- UNIDO – United Nations Industrial Development Organization
- UNRWA – United Nations Relief and Works Agency for Palestine Refugees in the Near East
- WFP – World Food Programme
- WIPO – World Intellectual Property Organization
- WMO – World Meteorological Organization
- WTO – World Trade Organization

The designations used and the presentation of the material in this volume do not imply the expression of any opinion whatsoever on the part of the Secretariat of the World Health Organization concerning the legal status of any country, territory, city or area or of its authorities, or concerning the delimitation of its frontiers or boundaries. Where the designation “country or area” appears in the headings of tables, it covers countries, territories, cities or areas.
PREFACE

The 126th session of the Executive Board was held at WHO headquarters, Geneva, from 18 to 23 January 2010. The proceedings are issued in two volumes. The present volume contains the resolutions and decisions, relevant annexes, and details regarding membership of committees. The summary records of the Board’s discussions and list of participants and officers are issued in document EB126/2010/REC/2.
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COMMITTEES AND WORKING GROUPS

1. Programme, Budget and Administration Committee

Dr M. Dahl-Regis (Bahamas, Chairman), Professor A.F.M.R. Haque (Bangladesh), Dr P. Buss (Brazil), Mr D. Houssin (France), Dr M. Kökény (Hungary), Ms K. Sujatha Rao (India), Dr S. Omi (Japan), Dr S. Kabuluzi (Malawi), Mr T. Ryall (New Zealand), Dr A. Djibo (Niger), Dr A.J. Mohamed (Oman), Dr A.A. Bin Shakar (United Arab Emirates), Dr S. Zaramba (Uganda), Chairman of the Executive Board, member ex officio, Professor Sohn Myongsei (Republic of Korea), Vice-Chairman of the Executive Board, member ex officio.

Eleventh meeting, 14 and 15 January 2010: Dr M. Dahl-Regis (Bahamas, Chairman), Mr S.A. Ali (alternate to Professor A.F.M.R. Haque, Bangladesh), Dr L.F. Beskow (alternate to Dr P. Buss, Brazil), Mr A. Allo (alternate to Mr D. Houssin, France), Dr A. Meszaro (alternate to Dr M. Kökény, Hungary), Mr P. Satpathy (alternate to Ms K. Sujatha Rao, India), Dr M. Mugitani (alternate to Dr S. Omi, Japan), Ms D. Roche (alternate to Mr T. Ryall, New Zealand), Dr A. Djibo (Niger), Dr A.J. Mohamed (Oman, Vice-Chair), Dr F. Al Braik (alternate to Dr S. Al Darmaki, United Arab Emirates), Dr S. Zaramba (Uganda), Chairman of the Executive Board, member ex officio, Professor Sohn Myongsei (Republic of Korea), Vice-Chairman of the Executive Board, member ex officio.

2. Standing Committee on Nongovernmental Organizations

Professor A.F.M.R. Haque (Bangladesh), Dr A.J. Mohamed (Oman), Mr C. Vallejos (Peru), Mrs G.A. Gidlow (Samoa), Dr J.A.N. dos Ramos (Sao Tome and Principe).

Meeting of 19 January 2010: Professor A.F.M.R Haque (Bangladesh), Dr A.J. Mohamed (Oman, Chairman), Mr C. Vallejos (Peru), Mr A. Peteru (alternate to Mrs G.A. Gidlow, Samoa), Dr J.A.N. dos Ramos (Sao Tome and Principe).

3. İhsan Doğramacı Family Health Foundation Selection Panel

The Chairman of the Executive Board, a representative of the International Children’s Centre, Ankara, and the President of Bilkent University or his or her appointee.

Meeting of 19 January 2010: Dr S. Zaramba (Uganda, Chairman), Professor P.L. Erdogan (appointee of Professor İ. Doğramacı, President of Bilkent University) and Professor M. Bertan (Representative of the International Children’s Centre, Ankara).

4. Sasakawa Health Prize Selection Panel

The Chairman of the Executive Board, a representative of the founder, and a member of the Executive Board.

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1 Showing their current membership and listing the names of those committee members who attended meetings held since the previous session of the Executive Board.
Meeting of 20 January 2010: Dr S. Zaramba (Uganda, Chairman), Professor K. Kiikuni (representative of the founder), Professor Sohn Myongsei (Republic of Korea).

5. United Arab Emirates Health Foundation Selection Panel

The Chairman of the Executive Board, a representative of the founder, and a member of the Executive Board from a Member State of the WHO Eastern Mediterranean Region.

Meeting of 19 January 2010: Dr S. Zaramba (Uganda, Chairman), Dr S. Al Darmaki (representative of the founder), Dr H. Abdesselem (Tunisia).

6. Dr LEE Jong-wook Memorial Prize Selection Panel

The Chairman of the Executive Board, a representative of the founder, and a member of the Executive Board from a Member State of the WHO Western Pacific Region.

Meeting of 20 January 2010: Dr S. Zaramba (Uganda, Chairman), Ms Chung Kyeong-hee (representative of the founder), Ms P.T. Toelupe (alternate to Mrs G.A. Gidlow, Samoa).
RESOLUTIONS

EB126.R1 Appointment of the Regional Director for Africa

The Executive Board,

Considering the provisions of Article 52 of the Constitution of WHO;

Considering the nomination made by the Regional Committee for Africa at its fifty-ninth session,

1. REAPPOINTS Dr Luis Gomes Sambo as Regional Director for Africa as from 1 February 2010;

2. AUTHORIZES the Director-General to issue to Dr Luis Gomes Sambo a contract for a period of five years from 1 February 2010, subject to the provisions of the Staff Regulations and Staff Rules.

(Third meeting, 19 January 2010)

EB126.R2 Appointment of the Regional Director for Europe

The Executive Board,

Considering the provisions of Article 52 of the Constitution of WHO;

Considering the nomination made by the Regional Committee for Europe at its fifty-ninth session,

1. APPOINTS Ms Zsuzsanna Jakab as Regional Director for Europe as from 1 February 2010;

2. AUTHORIZES the Director-General to issue to Ms Zsuzsanna Jakab a contract for a period of five years from 1 February 2010, subject to the provisions of the Staff Regulations and Staff Rules.

(Third meeting, 19 January 2010)

EB126.R3 Expression of appreciation to Dr Marc Danzon

The Executive Board,

Desiring, on the occasion of the retirement of Dr Marc Danzon as Regional Director for Europe, to express its appreciation of his services to the World Health Organization;

Mindful of his lifelong devotion to the cause of international health, and recalling especially his 10 years as Regional Director for Europe;
1. Expresses its profound gratitude and appreciation to Dr Marc Danzon for his invaluable contribution to the work of WHO;

2. Addresses to him on this occasion its sincere good wishes for many further years of service to humanity.

(Third meeting, 19 January 2010)

**EB126.R4 Monitoring of the achievement of the health-related Millennium Development Goals**

The Executive Board,

Having considered the report on monitoring of the achievement of the health-related Millennium Development Goals,¹

Recommends to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Having considered the report on monitoring of the achievement of the health-related Millennium Development Goals;

Recalling resolution WHA61.18 on monitoring of the achievement of the health-related Millennium Development Goals;

Recalling the outcomes of the major United Nations conferences and summits in the economic, social and related fields, especially those related to global health, in particular the 2005 World Summit Outcome and the commitments made by the international community to attain the Millennium Development Goals and the new commitments made during the United Nations High-level Event on the Millennium Development Goals (New York, 25 September 2008);

Stressing the importance of achieving the health-related Millennium Development Goals, especially with the objective of ensuring socioeconomic development;

Concerned by the fact that achievement of the Millennium Development Goals varies from country to country and from goal to goal;

Welcoming the Ministerial Declaration adopted at the annual ministerial review held by the Economic and Social Council in 2009 on implementing the internationally agreed goals and commitments in regard to global public health;

Recalling United Nations General Assembly resolution 64/108 (10 December 2009) on global health and foreign policy;

¹ Document EB126/7.
Recognizing that the Millennium Development Goals are interlinked, and reiterating the Health Assembly’s commitment to continued reinvigoration and strengthening of the global partnership for development, as a vital element for achieving these Goals, in particular those related to health, inter alia through capacity building, transfer of technology, sharing of best practices and lessons learnt, South–South cooperation, and predictability of resources;

Recalling the Monterrey Consensus of March 2002 to “urge developed countries that have not done so, to make concrete efforts towards the target of 0.7% of the gross national product (GNP) as ODA” and “encourage developing countries to build on progress achieved in ensuring that ODA is used effectively to help achieve development goals and targets”;

Reaffirming the commitments by many developed countries to achieve the target of 0.7% of gross national income on official development assistance by 2015 and to reach 0.56% of gross national income for official development assistance by 2010, as well as the target of 0.15% to 0.20% for least developed countries;

Welcoming increasing efforts to improve the quality of official development assistance and to increase its development impact, such as the Development Cooperation Forum of the Economic and Social Council, the principles contained in the Paris Declaration and the Accra Agenda for Action, and the experience of the International Health Partnership and others, in order to strengthen national ownership, alignment, harmonization and managing for results;

Noting the work of the Leading Group on Innovative Financing for Development and of the High-level Task Force on Innovative International Financing for Health Systems, the additional pledges made by several countries to increase financing for health, and the announcements made by several countries at the United Nations General Assembly High-level Meeting on Health (New York, 23 September 2009) to achieve universal access to affordable basic health care, including provision of free services for women and children at the point of use where countries choose, and financial mechanisms toward social health protection;

Expressing concern at the relatively slow progress in attaining the Millennium Development Goals, particularly in sub-Saharan Africa;

Expressing deep concern that maternal, newborn and child health and universal access to reproductive health services remain constrained by health inequities, and at the slow progress in achieving Millennium Development Goals 4 and 5 on improving child and maternal health;

Welcoming the contribution of all relevant partners and progress achieved towards the goal of universal access to prevention, treatment, care and support related to HIV/AIDS;

Reaffirming WHO’s leading role as the primary United Nations specialized agency for health, including its roles and functions with regard to health policy in accordance with its mandate;

Welcoming WHO’s report on women and health as important in advancing women’s rights and gender equality, underlining the need to address women’s health through comprehensive strategies targeting root causes of discrimination, and stressing the importance of strengthening health systems to better respond to women’s health needs in terms of access and comprehensiveness;

Recognizing that health systems based on the principles of tackling health inequalities through universal access, putting people at the centre of care, integrating health into broader public policy, and providing inclusive leadership for health are essential to achieving sustainable improvements in health;

Recognizing also the growing burden of noncommunicable diseases worldwide, and recalling the importance of preventing infectious diseases that still represent a heavy burden, particularly in developing countries, the adverse impacts of the food, environmental, economic and financial crises on populations, in particular on the poorest and the most vulnerable ones, which may increase the level of malnutrition and reverse the achievement of Millennium Development Goal 1 (Eradicate extreme hunger and poverty) and the health-related Goals and the progress made in the past two decades,

1. **URGES** Member States:

   (1) to strengthen health systems so that they deliver equitable health outcomes as a basis of a comprehensive approach towards achieving Millennium Development Goals 4, 5 and 6, underlining the need to build sustainable national health systems and strengthen national capacities through attention to, inter alia, service delivery, health systems financing, health workforce, health information systems, procurement and distribution of medicines, vaccines and technologies, sexual and reproductive health care and political will in leadership and governance;

   (2) to review policies, including those on recruitment, training and retention, that exacerbate the problem of the lack of health workers, and their imbalanced distribution, within countries and throughout the world, in particular the shortage in sub-Saharan Africa, which undermines the health systems of developing countries;

   (3) to reaffirm the values and principles of primary health care, including equity, solidarity, social justice, universal access to services, multisectoral action, transparency, accountability, decentralization and community participation and empowerment, as the basis for strengthening health systems, through support for health and development;

   (4) to take into account health equity in all national policies that address social determinants of health, and to consider developing and strengthening universal comprehensive social protection policies, including health promotion, infectious and noncommunicable disease prevention and health care, and promoting availability of and access to goods and services essential to health and well-being;

   (5) to renew their commitment to prevent and eliminate maternal, newborn and child mortality and morbidity: through an effective continuum of care, strengthening health systems, and comprehensive and integrated strategies and programmes to address root causes of gender inequalities and lack of access to adequate care and reproductive health, including family planning and sexual health; by promoting respect for women’s rights; and by scaling up efforts to achieve integrated management of newborn and child health care, including actions to address the main causes of child mortality;

   (6) to expand significantly efforts towards meeting the goal of universal access to HIV prevention, treatment, care and support by 2010 and the goal to halt and reverse the spread of HIV/AIDS by 2015;

   (7) to maximize synergies between the HIV/AIDS response and strengthening of health systems and social support;
(8) to enhance policies to address the challenges of malaria including monitoring of drug resistance in artemisinin-based combination therapy;

(9) to sustain and strengthen the gains made in combating tuberculosis, and to develop innovative strategies for tuberculosis prevention, detection and treatment, including means of dealing with new threats such as coinfection with HIV, multidrug-resistant tuberculosis or extensively drug-resistant tuberculosis;

(10) to sustain commitments to support the eradication of poliomyelitis;

(11) to include best practices for strengthening health services in bilateral and multilateral initiatives addressed to the achievement of the Millennium Development Goals, in particular in South–South cooperation initiatives;

(12) to support developing countries in their national endeavours to achieve the Millennium Development Goals, in particular the health-related Millennium Development Goals, inter alia through capacity building, transfer of technology, sharing of lessons learnt and best practices, South–South cooperation, and predictability of resources;

(13) to fulfil their commitments regarding official development assistance by 2015;

2. REQUESTS the Director-General:

(1) to continue to play a leading role in the monitoring of the achievement of the health-related Millennium Development Goals, including progress towards achieving universal coverage of services essential to these Goals;

(2) within the framework of WHO’s Medium-term strategic plan 2008–2013, to continue to cooperate closely with all other United Nations and international organizations involved in the process of achieving the Millennium Development Goals, maintaining a strong focus on efficient use of resources based on the respective mandates and core competencies of each, avoiding duplication of efforts and fragmentation of aid, and promoting the coordination of work among international agencies;

(3) to provide support to Member States in their efforts to strengthen their health systems, address the problem of the lack of health workers, reaffirm the values and principles of primary health care, address the social determinants of health, and strengthen their public policies aimed at fostering full access to health and social protection, including improved access to quality medicines required to support health care for, inter alia, the most vulnerable sectors of society;

(4) to foster alignment and coordination of global interventions for health system strengthening, basing them on the primary health care approach, in collaboration with Member States, relevant international organizations, international health initiatives, and other stakeholders in order to increase synergies between international and national priorities;

(5) to articulate and present to the Health Assembly as part of its action plan for the renewal of primary health care, the actions that the Secretariat envisages will strengthen its support for the realization of Millennium Development Goals 4, 5 and 6;
(6) to work with all relevant partners in order to achieve high immunization coverage rates with affordable vaccines of assured quality;

(7) to lead the work with all relevant partners to help to ensure that action on the health-related Millennium Development Goals is one of the main themes of the United Nations Millennium Development Goals High-level Plenary Meeting, 20–22 September 2010;

(8) to continue to collect and compile scientific evidence needed for achieving health-related Millennium Development Goals and to disseminate it to all Member States;

(9) to continue to submit annually a report on the status of progress made, including on main obstacles and ways to overcome them, in achievement of the health-related Millennium Development Goals, through the Executive Board, to the Health Assembly;

3. INVITES concerned organizations of the United Nations system, international development partners and agencies, international financial institutions, nongovernmental organizations and private sector entities to continue their support and consider further support to countries, particularly in sub-Saharan Africa, for the development and implementation of health policies and national health development plans, consistent with internationally agreed health goals, including the Millennium Development Goals.

(Seventh meeting, 21 January 2010)

**EB126.R5 Infant and young child nutrition**

The Executive Board,

Having considered the quadrennial progress report on infant and young child nutrition,¹

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Having considered the report on infant and young child nutrition;

Recalling resolutions WHA35.26, WHA37.30, WHA39.28, WHA41.11, WHA43.3, WHA45.34, WHA46.7, WHA47.5, WHA49.15 and WHA54.2 on infant and young child nutrition and WHA59.11 on nutrition and HIV/AIDS;

Conscious that achieving the Millennium Development Goals will require the reduction of maternal and child malnutrition;

Aware that worldwide malnutrition accounts for 11% of the global burden of disease, leading to long-term poor health and disability and poor educational and developmental outcomes; that worldwide 178 million children are underweight and 20 million suffer from the

¹Document EB126/9.
most deadly form of severe acute malnutrition each year; and that nutritional risk factors, including underweight, suboptimal breastfeeding and vitamin and mineral deficiencies, particularly of vitamin A, iron, iodine and zinc, are responsible for 3.9 million deaths (35% of total deaths) and 144 million disability-adjusted life years (33% of total disability-adjusted life years) in children less than five years old;

Aware that countries are faced with increasing public health problems posed by the double burden of malnutrition (both undernutrition and overweight), with its negative later-life consequences;

Acknowledging that 90% of stunted children live in 36 countries and that children under two years of age are most affected by undernutrition;

Mindful of the challenges posed by the HIV/AIDS pandemic and the difficulties in formulating appropriate policies for infant and young child feeding, and concerned that food assistance does not meet the nutritional needs of young children infected by HIV;

Aware that inappropriate feeding practices and their consequences are major obstacles to attaining sustainable socioeconomic development and poverty reduction;

Concerned about the vast numbers of infants and young children who are still inappropriately fed and whose nutritional status, growth and development, health and survival are thereby compromised;

Mindful of the fact that implementation of the global strategy for infant and young child feeding and its operational targets requires strong political commitment and a comprehensive approach, including strengthening of health systems and communities and careful monitoring of the effectiveness of the interventions used;

Recognizing that the improvement of breastfeeding practices could save annually the lives of about one million children under five years of age and that each year the deaths of more than half a million such children could be prevented by adequate and timely complementary feeding along with continual breastfeeding for up to two years or beyond;

Aware that multisectoral food and nutrition policies are needed for the successful scaling up of evidence-based safe and effective nutrition interventions;

Recognizing the need for comprehensive national policies on infant and young child feeding that are well integrated within national strategies for nutrition and child survival;

Convinced that it is time for governments, civil society and the international community to renew their commitment to promoting the optimal feeding of infants and young children and to work together closely for this purpose;

Convinced that strengthening of national nutrition surveillance is crucial in implementing effective nutrition policies and scaling up interventions,
1. URGES Member States:

(1) to increase political commitment to reducing malnutrition in all its forms;

(2) to strengthen and expedite the implementation of the global strategy for infant and young child feeding with emphasis on giving effect to the International Code of Marketing of Breast-milk Substitutes, adopted in resolution WHA34.22;

(3) to develop or review current policy frameworks addressing the double burden of malnutrition and allocate adequate human and financial resources to ensure their implementation;

(4) to scale up interventions to improve infant and young child nutrition, including: the protection and promotion of breastfeeding and timely, safe and appropriate complementary feeding; the implementation of supplementary and therapeutic feeding interventions for severe malnutrition; and the control of vitamin and mineral deficiencies;

(5) to include the strategies referred to in subparagraph 1(4) above in comprehensive maternal and child health services and support the aim of universal coverage and principles of primary health care, including strengthening health systems as outlined in resolution WHA62.12;

(6) to strengthen nutrition surveillance systems and improve use and reporting of agreed Millennium Development Goals indicators in order to monitor progress;

(7) to implement the WHO Child Growth Standards by their full integration into child health programmes;

2. CALLS UPON the food industry to observe the International Code of Marketing of Breast-milk Substitutes and enhance its corporate social responsibilities;

3. REQUESTS the Director-General:

(1) to strengthen the evidence base on effective and safe nutrition actions to counteract the public health effects of the double burden of malnutrition, and to describe good practices for successful implementation;

(2) to mainstream nutrition in all WHO’s health policies and strategies and confirm the presence of essential nutrition actions in the context of the reform of primary health care;

(3) to continue and strengthen collaboration with other United Nations agencies and international organizations involved in the process of ensuring improved nutrition including clear identification of leadership, division of labour and outcomes;

(4) to support Member States, on request, in expanding nutritional interventions related to the double burden of malnutrition, monitoring and evaluating impact, strengthening or establishing effective nutrition surveillance systems, and implementing the WHO Child Growth Standards;
(5) to develop a comprehensive implementation plan on infant and young child nutrition as a critical component of a global multisectoral nutrition framework for preliminary discussion at the Sixty-fourth World Health Assembly and for final delivery at the Sixty-fifth World Health Assembly, through the Executive Board and after broad consultation with Member States.

(Seventh meeting, 21 January 2010)

**EB126.R6 Birth defects**

The Executive Board,

Having considered the report on birth defects,

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Concerned by the high number of stillbirths and neonatal deaths occurring worldwide and by the large contribution of neonatal mortality to under-five mortality;

Recognizing the importance of birth defects as a cause of stillbirths and neonatal mortality;

Mindful that effective interventions to prevent birth defects including provision of appropriate community genetic services within the primary health care are available that can be integrated into maternal, reproductive and child health services;

Concerned by the inadequate coverage of maternal, newborn and child health interventions and the barriers to access to health services that still exist in countries with the highest burden of maternal, newborn and child deaths;

Aware that the attainment of Millennium Development Goal 4 (Reduce child mortality) will require accelerated progress in reducing neonatal mortality including prevention and management of birth defects;

Recalling resolution WHA58.31, in which the Health Assembly, calling for universal coverage of maternal, newborn and child health interventions, urged Member States to commit resources and to accelerate national action to build a seamless continuum of care for reproductive, maternal, newborn and child health; and resolution WHA57.13 in which it was recognized that genomics has a significant contribution to make in the area of public health;

Recognizing that the prevalence of birth defects varies between communities, and that insufficient epidemiological data may hamper effective and equitable management;

1. See Annex 1.
Recognizing the diversity of causes and determinants of congenital disorders, including preventable factors such as infections or nutritional factors, vaccine-preventable diseases, consumption of alcohol, tobacco and drugs, and exposure to chemical substances, notably pesticides;

Deeply concerned that birth defects are not still recognized as priorities in public health;

Concerned by the limited resources dedicated to prevention and management of birth defects in particular in middle- and low-income countries;

Welcoming the report on birth defects,

1. URGES Member States:

(1) to raise awareness among all relevant stakeholders, including government officials, health professionals, civil society and the public, about the importance of birth defects as a cause of child morbidity and mortality;

(2) to set priorities, commit resources, and develop plans and activities for integrating effective interventions that include comprehensive guidance, information and awareness raising to prevent birth defects, and care for children with birth defects into existing maternal, reproductive and child health services and social welfare for all individuals who need them;

(3) to promote the application of internationally recognized standards regulating the use of chemical substances in the air, water and soil;

(4) to increase coverage of effective prevention measures, through health education programmes that include ethical, legal and social issues associated with birth defects for the general population and high-risk groups, and by fostering the development of parent-patient organizations and establishing appropriate community genetic services;

(5) to record surveillance data on birth defects as part of national health information systems;

(6) to develop expertise and to build capacity on the prevention of birth defects and care of children with birth defects;

(7) to strengthen research and studies on etiology, diagnosis and prevention of major birth defects and to promote international cooperation in combating with them;

(8) to take all necessary measures to ensure the full enjoyment by children with disabilities of all human rights and fundamental freedoms on an equal basis with other children and give priority to the child’s well-being and support and facilitate families in their child-care and child-raising efforts;

(9) to raise awareness among all relevant stakeholders, including government officials, health professionals, civil society and the public, about the importance of newborn screening programmes and their role in identifying infants born with birth defects;
(10) to support families who have children with birth defects and associated disabilities, and ensure that appropriate habilitation and support is provided to children with disabilities;

2. REQUESTS the Director-General:

(1) to promote the collection of data on the global burden of mortality and morbidity due to birth defects, and to consider broadening the groups of congenital abnormalities included in the classification when the International Statistical Classification of Diseases and Related Health Problems (Tenth Revision) is revised;

(2) to continue to collaborate with the International Clearinghouse for Birth Defects Surveillance and Research in order to improve collection of data on global burden of mortality and morbidity due to birth defects;

(3) to support Member States in developing national plans for implementation of effective interventions to prevent and manage birth defects within their national maternal, newborn and child health plan, strengthening health systems and primary care, including improved coverage of vaccination against diseases such as measles and rubella, and food fortification strategies, for the prevention of birth defects, and promoting equitable access to such services;

(4) to provide support to Member States in developing ethical and legal guidelines in relation to birth defects;

(5) to support Member States in the provision of appropriate community genetic services within the primary health-care system;

(6) to promote technical cooperation among Member States, nongovernmental organizations and other relevant bodies on prevention of birth defects;

(7) to support and facilitate research efforts on prevention and management of birth defects in order to improve the quality of life of those affected by such disorders;

(8) to report on progress in implementing this resolution to the Sixty-seventh World Health Assembly, through the Executive Board, in 2014.

(Seventh meeting, 21 January 2010)

**EB126.R7 Advancing food safety initiatives**

The Executive Board,

Having considered the report on food safety,¹

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¹ Document EB126/11.
RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Recalling resolution WHA53.15 on food safety, which requested the Director-General to put in place a global strategy for the surveillance of foodborne diseases and for the efficient gathering and exchange of information in and between countries;

Recalling resolution WHA55.16 on the global public health response to natural occurrence, accidental release or deliberate use of biological and chemical agents or radionuclear material that affect health, which noted that such agents can be disseminated through food- and water-supply chains;

Noting the endorsement by the Executive Board in 2002 of WHO’s global strategy for food safety,1 which had as its aim the reduction of the health and social burden of foodborne disease;

Noting also, that other food safety-related activities identified in resolutions WHA53.15 and WHA55.16 have been undertaken, including: the revision of the International Health Regulations in 2005; the establishment of the International Food Safety Authorities Network in 2005; the establishment of WHO’s Foodborne Disease Burden Epidemiology Reference Group in 2006; and increased participation, particularly by developing countries, in the elaboration of international food safety standards by the Codex Alimentarius Commission;

Recognizing that the Codex Alimentarius Commission presents a unique opportunity for all countries to join the international community in formulating and harmonizing food standards and ensuring their global implementation, and in particular the participation of developing countries in this regard should be encouraged;

Further recognizing the important roles of WHO and FAO in support of the Codex Alimentarius Commission as the international reference point for developments associated with food standards;

Confirming that foodborne disease continues to represent a serious threat to the health of millions of people in the world, particularly those in developing countries with poor nutritional status;

Mindful of the inextricable links between food safety, nutrition and food security, and acknowledging the instrumental role of food safety in eradicating hunger and malnutrition, in particular in low-income and food-deficit countries;

Aware of increasing evidence that many communicable diseases, including emerging zoonoses, are transmitted through food, and that exposure to chemicals and pathogens in the food supply is associated with acute and chronic diseases;

Acknowledging that climate change could be a factor in the increasing rates of some foodborne diseases, including those of zoonotic origin, owing to the more rapid growth of

1 Document EB109/2002/REC/2, summary record of the fourth meeting.
microorganisms in food and water with higher temperatures, resulting in the emergence of toxins in new geographical areas and possibly in higher levels of toxins or pathogens in food;

Recognizing that the global trade in food is increasing every year, contributing to the risk of spread of pathogens and contaminants across national borders, thereby creating new challenges for food authorities and necessitating more efficient global sharing of food safety information, taking into account that protection of food safety cannot lead to discrimination or a disguised restriction on international trade;

Acknowledging the continuing need for closer collaboration between the health sector and other sectors, and increased action on food safety at the international and national levels, across the full length of the food-production chain, in order to reduce significantly the incidence of foodborne disease;

Noting the continuing need for updated and comprehensive internationally agreed standards and agreements for risk assessments and scientific advice to support measures and interventions to improve the safety and nutritional quality of food;

Recognizing the importance of international agreement on global management of food safety, the application of scientific principles in finding solutions, the efficient exchange of monitoring and surveillance data, and practical experience,

1. URGES Member States:¹

  (1) to continue to establish and maintain the activities and measures elaborated in resolutions WHA53.15 on food safety and WHA55.16 on the global public health response to natural occurrence, accidental release or deliberate use of biological and chemical agents or radionuclear material that affect health;

  (2) to further develop and implement the core capacities as defined in Annex 1 of the International Health Regulations (2005), as applicable, and those required for participation in the International Food Safety Authorities Network, specifically for food-safety events, including the development of systems for: surveillance for foodborne disease and food contamination; risk assessment, traceability, risk management, including the Hazard Analysis and Critical Control Points system, and risk communication; food safety emergency response; product tracing and recall; and strengthened laboratory capacity;

  (3) to participate fully as members of the International Food Safety Authorities Network in its activities, including supporting the timely transmission of data, information and knowledge about food-safety emergencies through the Network in a transparent manner;

  (4) to enhance the integration of food-safety considerations into food aid, food security and nutrition interventions in order to reduce the occurrence of foodborne diseases and improve the health outcomes of populations in particular the vulnerable groups;

  (5) to establish or improve the evidence base for food safety through systematic efforts on disease-burden estimation and surveillance, and through comprehensive risk and risk-

¹ And, where applicable, regional economic integration organizations.
benefit assessment, and to provide support for international activities in these areas, in particular, WHO’s initiative to estimate the global burden of foodborne diseases from all major causes (microbiological, parasitic and chemical);

(6) to contribute to the timely conduct of international risk assessments through the provision of relevant data and expertise in order to tackle more efficiently and consistently foodborne diseases and food-safety issues that threaten global public health security;

(7) to continue to develop and maintain sustainable preventive measures, including food safety-education programmes, aimed at reducing the burden of foodborne diseases through a systems approach encompassing the complete food-production chain from farm to consumption;

(8) to promote dialogue and collaboration among human health, veterinary and food-related disciplines, within and among Member States, focused on an integrated effort of foodborne risk reduction along the whole food-production chain, including consideration of zoonotic risks;

(9) to participate actively in the Codex Alimentarius Commission’s standard-setting process and to adopt Codex standards whenever appropriate;

2. REQUESTS the Director-General:

(1) to develop the International Food Safety Authorities Network further through the implementation of the WHO’s global strategy for food safety; to encourage communication and technical exchange of risk assessments and best practices among members of the Network; to facilitate Member States’ involvement in the Network’s operation and development; and to encourage additional membership into the International Food Safety Authorities Network;

(2) to strengthen the emergency function of the International Food Safety Authorities Network as a critical component of WHO’s preventive and emergency operations relative to food safety, and linkages to other relevant international organizations and networks in this area;

(3) to continue to provide global leadership in providing technical assistance and tools that meet the needs of Member States and the Secretariat for scientific estimations on foodborne risks and foodborne disease burden from all causes;

(4) to promote the inclusion of food safety into the international debate on food crises and hunger emergencies, and provide technical support to Member States and international agencies for considering food safety, nutrition and food security issues in a comprehensive, integrated manner;

(5) to monitor regularly and report to Member States on the global burden of foodborne and zoonotic diseases from the country, regional and international perspectives;

(6) to promote research, including the safety and quality of traditional foods, and investigation of the association of foodborne hazards with acute and chronic diseases, in order to support evidence-based strategies for the control and prevention of foodborne and zoonotic diseases such as the Hazard Analysis and Critical Control Points system;
(7) to provide support to Member States in building relevant capacity to improve cross-sectoral collaboration and action at international, regional and national levels along the whole food-production chain, including the assessment, management and communication of foodborne and zoonotic risks;

(8) to develop guidance on the public health aspects arising from zoonotic diseases that originate at the human-animal interface, in particular prevention, detection and response;

(9) to provide adequate and sustainable support for the joint expert bodies of FAO and WHO, the Codex Alimentarius Commission and the International Food Safety Authorities Network in order to advance the international development, provision, use, and sharing of scientific risk assessments and advice; to support the development of international food standards that protect the health and nutritional well-being of consumers; and to address and communicate more effectively on food safety issues at the national and international levels;

(10) to establish with the International Food Safety Authorities Network an international initiative for the collaboration of laboratory partners in support of surveillance of foodborne disease, identification of food contamination and emergency response, including outbreak investigation and linking product to illness in order to support recall, with that initiative also including the establishment of mechanisms for data sharing;

(11) to report to the Sixty-fifth World Health Assembly, through the Executive Board, on progress in implementing this resolution.

(Eighth meeting, 21 January 2010)

**EB126.R8 Method of work of the Executive Board**

The Executive Board,

Having considered the report on the method of work of the governing bodies,

DECIDES to amend its Rules of Procedure as proposed in the Annex to the report on the method of work of the governing bodies, with effect from the closure of its 127th session.

(Ninth meeting, 22 January 2010)

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1 See Annex 2.

The Executive Board,

CONFIRMS, in accordance with Staff Regulation 12.2, the amendments to the Staff Rules that have been made by the Director-General with effect from 1 January 2010 concerning the remuneration of staff in the professional and higher categories.

(Ninth meeting, 22 January 2010)

EB126.R10 Salaries of staff in ungraded posts and of the Director-General

The Executive Board,

Having considered the report on confirmation of amendments to the Staff Regulations and Staff Rules,\(^2\)

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Noting the recommendations of the Executive Board with regard to remuneration of staff in ungraded posts and of the Director-General,

1. **ESTABLISHES** the salaries of Assistant Directors-General and Regional Directors at US$ 183 022 gross per annum before staff assessment, resulting in a modified net salary of US$ 131 964 (dependency rate) or US$ 119 499 (single rate);

2. **ESTABLISHES** the salary of the Deputy Director-General at US$ 201 351 gross per annum before staff assessment, resulting in a modified net salary of US$ 143 878 (dependency rate) or US$ 129 483 (single rate);

3. **ESTABLISHES** the salary of the Director-General at US$ 247 523 gross per annum before staff assessment, resulting in a modified net salary of US$ 173 890 (dependency rate) or US$ 154 641 (single rate);

4. **DECIDES** that those adjustments in remuneration shall take effect from 1 January 2010.

(Ninth meeting, 22 January 2010)

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\(^{1}\) See Annex 3.

\(^{2}\) Document EB126/39.
The Executive Board,

Having considered the report on strategies to reduce the harmful use of alcohol,\(^2\)

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Having considered the report on strategies to reduce the harmful use of alcohol and the draft global strategy annexed therein;

Recalling resolutions WHA58.26 on public-health problems caused by harmful use of alcohol and WHA61.4 on strategies to reduce the harmful use of alcohol,

1. ENDORSES the global strategy to reduce the harmful use of alcohol;

2. AFFIRMS that the global strategy aims to give guidance for action at all levels and to set priority areas for global action, and that it is a portfolio of policy options and measures that could be considered for implementation and adjusted as appropriate at the national level, taking into account national circumstances, such as religious and cultural contexts, national public health priorities, as well as resources, capacities and capabilities;

3. URGES Member States:\(^3\)

   (1) to adopt and implement the global strategy to reduce the harmful use of alcohol as appropriate in order to complement and support public health policies in Member States to reduce the harmful use of alcohol, and to mobilize political will and financial resources for that purpose;

   (2) to continue implementation of the resolutions WHA61.4 on the strategies to reduce the harmful use of alcohol and WHA58.26 on public-health problems caused by harmful use of alcohol;

   (3) to ensure that implementation of the global strategy to reduce the harmful use of alcohol strengthens the national efforts to protect at-risk populations, young people and those affected by harmful drinking of others;

   (4) to ensure that implementation of the global strategy to reduce the harmful use of alcohol is reflected in the national monitoring systems and reported regularly to WHO’s information system on alcohol and health;

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\(^1\) See Annex 4.


\(^3\) And regional economic integration organizations, where applicable.
4. REQUESTS the Director-General:

   (1) to give sufficiently high organizational priority to prevention and reduction of harmful use of alcohol and implementation of the global strategy to reduce the harmful use of alcohol;

   (2) to collaborate with and provide support to Member States, as appropriate, in implementing the global strategy to reduce the harmful use of alcohol and strengthening national responses to public health problems caused by the harmful use of alcohol;

   (3) to monitor progress in implementing the global strategy to reduce the harmful use of alcohol and to report progress, through the Executive Board, to the Sixty-sixth World Health Assembly.

(Eleventh meeting, 22 January 2010)

EB126.R12 Improvement of health through safe and environmentally sound waste management

The Executive Board,

Having considered the report on the Strategic Approach to International Chemicals Management;¹

Having also considered the letter of President of the ninth meeting of the Conference of the Parties to the Basel Convention on the Control of Transboundary Movement of Hazardous Wastes and their Disposal to the Director-General of WHO,

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Having considered the report on the Strategic Approach to International Chemicals Management;

Recalling resolution WHA61.19 on climate change and health, and resolutions WHA59.15, WHA50.13, WHA45.32, WHA31.28 and WHA30.47 relating to chemical safety;

Recalling also resolutions of the United Nations General Assembly 44/226 of 22 December 1989 on traffic in and disposal, control and transboundary movements of toxic and dangerous products and wastes and 43/212 of 20 December 1988 on the responsibility of States for the protection of the environment;

Noting the principles set out in Agenda 21, including chapter 20 and chapter 21, as agreed upon at the United Nations Conference on Environment and Development in 1992;

¹ Document EB126/20.
Noting also the Johannesburg Declaration on Sustainable Development and the related Plan of Implementation of the World Summit on Sustainable Development in 2002;


Mindful of the outcomes of the second session of the International Conference on Chemicals Management which relate to human health;

Aware that wastes, if not properly managed in a safe and environmentally sound manner, may have serious consequences for human health and livelihood;

Convinced that the lack of environmentally sound management of waste will harm the environment and be detrimental to human health, through polluted air, water, land and food chains;

Concerned that poor management of health-care waste, including sharps, non-sharp materials, blood, body parts, chemicals, pharmaceuticals and medical devices, puts health-care workers, waste handlers and the community at risk of infections, toxic effects and injuries;

Welcoming the Bali Declaration on Waste Management for Human Health and Livelihood adopted at the ninth meeting of the Conference of the Parties to the Basel Convention on the Control of Transboundary Movement of Hazardous Wastes and their Disposal in 2008,

1. URGES Member States\(^1\) to assess the health aspects of waste management in order to make it safe and environmentally sound and to explore options to work more closely with the United Nations Environment Programme, the Strategic Approach to International Chemicals Management, the Basel Convention on the Control of Transboundary Movement of Hazardous Wastes and their Disposal and the WHO Secretariat towards achieving their shared objectives on the improvement of health through safe and environmentally sound waste management;

2. REQUESTS the Director-General:

   (1) to support the implementation of the actions set out in the Bali Declaration on Waste Management for Human Health and Livelihood, within WHO’s mandate and available resources;

   (2) to work together with the United Nations Environment Programme and the secretariat of the Basel Convention on the Control of Transboundary Movement of Hazardous Wastes and their Disposal on environmentally sound waste management, including collaborating with governments and donor organizations to strengthen the implementation of the Bali Declaration on Waste Management for Human Health and Livelihood, with the aim in particular of:

   (a) promoting the raising of awareness about the link between waste management, health and livelihood, and the environment;

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\(^1\) And regional economic integration organizations, where applicable.
(b) strengthening subregional and regional cooperation on waste and health issues by promoting human and appropriate technical capacities at national, regional and international levels;

(c) improving controls on waste shipment and border procedures in order to prevent illegal movements of hazardous and other wastes, through means that include capacity building, technology transfer and technical assistance;

(d) improving cooperation between national authorities in the waste, chemicals and health sectors and, in collaboration with other relevant authorities and stakeholders, in the development and implementation of effective and sound waste management systems;

(e) increasing capacity building, promoting and, where possible, enhancing public and private investment for the transfer and use of appropriate technology for the safe and environmentally sound waste management;

(3) to continue supporting the prevention of health risks associated with exposure to health-care waste and promoting environmentally sound management of health-care waste in order to support the work of the Basel Convention on the Control of Transboundary Movement of Hazardous Wastes and their Disposal and the Stockholm Convention on Persistent Organic Pollutants;

(4) to explore the development of strategies aimed at minimizing the generation of health-care waste;

(5) to invite governments, relevant intergovernmental and regional economic integration organizations, relevant entities of the industry and business sectors to provide resources and technical assistance to developing countries in developing and implementing instruments to improve health through safe and environmentally sound waste management;

(6) to report to the Sixty-fourth World Health Assembly, through the Executive Board, on implementation of this resolution.

(Eleventh meeting, 22 January 2010)

**EB126.R13 Improvement of health through sound management of obsolete pesticides**\(^1\) and other obsolete chemicals

The Executive Board,

Having considered the report on the Strategic Approach to International Chemicals Management,\(^2\)

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\(^1\) “Obsolete pesticides” are defined as those pesticides that can no longer be used for their intended purpose or whose use is no longer wanted and which therefore must be disposed of.

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Having considered the report on the Strategic Approach to International Chemicals Management;

Recalling resolution WHA59.15 on the Strategic Approach to International Chemicals Management;

Recognizing the outcomes of the second session of the International Conference on Chemicals Management (Geneva, 11–15 May 2009) regarding human health and, in particular, resolution II/8 on health aspects of the sound management of chemicals which drew attention to the need for a greater involvement of health sector, Member States1 and the WHO Secretariat in the implementation of the Global Plan of Action of the Strategic Approach to International Chemicals Management2 because of the adverse effects some chemicals may have on human health, and noting that some of the global priorities for cooperative action identified within the Strategic Approach to International Chemicals Management also have to be dealt with by the health sector;

Recognizing that pesticides are designed to kill or control harmful organisms and pests, and may have adverse acute and chronic effects, and that, although they are regulated in most countries, they may affect populations’ health and the environment, particularly when improperly used and stored, including when they are obsolete;3


Recognizing that all the forums, conventions and instruments mentioned in the preceding paragraph are important global tools for the preservation and protection of human health and the environment that provide measures and guidelines to deal with certain aspects of chemicals

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1 And, where applicable, regional economic integration organizations.

2 Document WHA59/2006/REC/1, Annex 1.

3 The International HCH and Pesticides Association (IHPA) estimates that total amount of obsolete pesticides is about 260 000–265 000 tonnes in central and eastern Europe and the countries of the former Union of Soviet Socialist Republics. Estimated amounts in 25 Member States of the European Union are 22 000–24 000 tonnes, south-east Europe 36 000–41 000 tonnes, the countries of the former Union of Soviet Socialist Republics 199 000 tonnes, Africa 50 000 tonnes (estimated by FAO in its Africa Stockpiles Programme), South-East Asia 6500 tonnes (FAO, first rough indication), Central and South America 30 000 tonnes (FAO, 2005).
life-cycle, and that, in that sense, the closely linked Stockholm Convention on Persistent Organic Pollutants and Basel Convention on the Control of the Transboundary Movements of Hazardous Wastes and Their Disposal\(^1\) foresee the development of appropriate strategies for identification of persistent organic pollutant wastes, stockpiles of persistent organic pollutants and their management;

Recognizing that hazardous waste and highly toxic pesticides fall under the global priority areas identified for cooperative action within the Strategic Approach to International Chemicals Management, and that the Health Assembly in resolution WHA59.15 on Strategic Approach to International Chemicals Management urged Member States to participate in national, regional and international efforts to implement the Strategic Approach;

Mindful of the new challenges and determinants of health and of the need for additional action in order to preserve and protect human health and the environment;

Recognizing the risks to human health and environment from obsolete pesticides and other obsolete chemicals, particularly through local and global chemical accidents and disasters;

Recognizing also the risks to human health and environment from obsolete pesticides and other obsolete chemicals, linked to the creation of stockpiles resulting from their regulation (such as withdrawal from the market without appropriate phase-out period), which might further lead to spreading of improperly stored chemicals worldwide;

Recalling the fact that the exposure of humans and the environment to obsolete pesticides and other obsolete chemicals may also be due to their long-range transport;

Recognizing the threat of unsafe storage of obsolete pesticides and other obsolete chemicals, which, owing to illegal use, package deterioration, or accidents may cause localized or widespread pollution and represent a potential risk to human health and the environment;

Mindful of the clear evidence that, besides environmental benefits, economic benefits can be expected from safe and efficient recovery, reuse, recycling and disposal of obsolete pesticides and other obsolete chemicals;

Acknowledging the progress regarding obsolete pesticides made by African countries through the interagency Africa Stockpiles Programme with the support of FAO, the Global Environment Facility, the World Bank and other partners;

Welcoming the work of the Basel Convention on the Control of the Transboundary Movements of Hazardous Wastes and Their Disposal in developing technical guidelines on the environmentally sound disposal of wastes containing persistent organic pollutants;

Further recognizing that only a comprehensive and long-term strategy of sound management of obsolete pesticides and other obsolete chemicals can be effective,

\(^1\) The fundamental aims of the Basel Convention are the control and reduction of transboundary movement of hazardous and other wastes subject to the Convention, the prevention and minimization of their generation, the environmentally sound management of such wastes and active promotion of the transfer and use of cleaner technologies.
1. **URGES** Member States:

   (1) to adopt, where necessary, or strengthen sound national policies and legislation on safe handling and disposal of obsolete pesticides and other obsolete chemicals;

   (2) to adopt, where this has not already been done in the context of the Stockholm Convention on Persistent Organic Pollutants and other existing instruments, comprehensive national implementation plans or other strategies as the basis for taking action towards the elimination of risks from obsolete pesticides and other obsolete chemicals;

   (3) to enhance social responsibility through awareness-raising in the area of obsolete pesticides and other obsolete chemicals and chemicals with potential transboundary risks to human health;

   (4) to increase support for training and capacity building, and coordinated technical activities for implementing relevant international conventions and instruments;

   (5) to encourage and promote cooperation between Member States in this regard;

   (6) to establish or strengthen capacity for the regulation of the sound management of pesticides and other chemicals throughout their life-cycle, as a preventive measure to avoid accumulation of obsolete chemicals;

2. **INVITES** all relevant stakeholders, including Member States, regional economic integration organizations, bodies in the United Nations system and other intergovernmental organizations including regional, international and national nongovernmental organizations and foundations, waste-management companies, pesticide manufacturers, donors and the remaining international community:

   (1) to promote sound management of obsolete pesticides and other obsolete chemicals in order to minimize and, wherever possible, avoid adverse impacts to human health and the environment;

   (2) to mobilize efforts and cooperate with other stakeholders on the implementation of national implementation plans and strategies, through local, regional and global networks among other means;

   (3) to consider the synergies to be gained from sharing technical experience, expertise and capacity-building efforts among international instruments, conventions, regulations and processes;

3. **REQUESTS** the Director-General:

   (1) to support the development of appropriate and efficient strategies (at national, regional and international levels) for minimizing the risks of obsolete pesticides and other obsolete chemicals and thus promote the relevant WHO policy goals and practices;

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1 And, where applicable, regional economic integration organizations.
(2) to enhance WHO’s capacity to foster the strategies mentioned in subparagraph 3(1) above;

(3) to facilitate implementation of the strategies on sound management of obsolete pesticides and other obsolete chemicals with a view to reducing inequities in health and securing an unpolluted living environment;

(4) to work with UNEP, in connection with the WHO/UNEP Health Environment Linkages Initiative and the Strategic Approach to International Chemicals Management, as well as with UNDP, FAO, the World Bank and other appropriate institutions in assisting Member States to implement their national strategies and existing guidance, for instance under the Basel Convention on the Control of the Transboundary Movements of Hazardous Wastes and Their Disposal and strategies for sound management of obsolete pesticides and other obsolete chemicals at the global level;

(5) to include obsolete pesticides and other obsolete chemicals among WHO’s priorities in order to reduce and prevent risks to human health and the environment from their adverse effects and to support their safe disposal;

(6) to ensure full support of WHO to the activities of the Secretariat of the Strategic Approach to International Chemicals Management;

(7) to support the ongoing joint efforts of FAO and WHO in capacity building of Member States in the sound management of pesticides;

(8) to report to the Sixty-fourth World Health Assembly, through the Executive Board, on progress in implementing this resolution.

(Eleventh meeting, 22 January 2010)

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1 Technical guidelines on the safe disposal of obsolete pesticides (http://www.basel.int/meetings/sbc/workdoc/techdocs.html):

- Updated general technical guidelines for the environmentally sound management of wastes consisting of, containing or contaminated with persistent organic pollutants,
- Technical guidelines for the environmentally sound management of wastes consisting of, containing or contaminated with 1,1,1-trichloro-2,2-bis(4-chlorophenyl)ethane (DDT),
- Technical guidelines on the environmentally sound management of wastes consisting of, containing or contaminated with the pesticides aldrin, chlordane, dieldrin, endrin, heptachlor, hexachlorobenzene (HCB), mirex or toxaphene or with HCB as an industrial chemical.
The Executive Board,

Having considered the report on availability, safety and quality of blood products,

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Recalling resolution WHA58.13 on blood safety: proposal to establish World Blood Donor Day and preceding related resolutions since resolution WHA28.72 on utilization and supply of human blood and blood products, which urged Member States to promote the full implementation of well-organized, nationally coordinated and sustainable blood programmes with appropriate regulatory systems and to enact effective legislation governing the operation of blood services;

Recognizing that achieving self-sufficiency, unless special circumstances preclude it, in the supply of safe blood components based on voluntary, non-remunerated blood donation, and the security of that supply are important national goals to prevent blood shortages and meet the transfusion requirements of the patient population;

Conscious that plasma-derived medicinal products for the treatment of haemophilia and immune diseases are included in the WHO Model List of Essential Medicines and of the need to facilitate access to these products by developing countries;

Concerned by the unequal access globally to blood products, particularly plasma-derived medicinal products, leaving many patients in need of transfusion and with severe congenital and acquired disorders without adequate treatment;

Aware that a major factor limiting the global availability of plasma-derived medicinal products is an inadequate supply of plasma meeting internationally recognized standards for fractionation;

Bearing in mind that treatment using labile blood components is gradually being included in medical practice in developing countries and that thereby increased quantities of recovered plasma should become available for fractionation into plasma-derived medicinal products to meet their needs;

Concerned that in developing countries, blood components separation technology and fractionation capacity are lacking, and that, because of insufficient regulatory controls and

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1 The term “blood products” is defined by the Expert Committee on Biological Standardization as follows: “any therapeutic substances derived from human blood, including whole blood, labile blood components and plasma-derived medicinal products”.

2 Document EB126/19.

3 The WHO Model List of Essential Medicines identifies individual medicines that together could provide safe and effective treatment for most communicable and noncommunicable diseases. This List includes plasma-derived medicinal products, namely immunoglobulins and coagulation factors, which are needed to prevent and treat a variety of serious conditions that occur worldwide (http://www.who.int/medicines/publications/essentialmedicines/en/index.html).
failure to implement appropriate practices in blood establishments, plasma from developing countries is often unacceptable for contract fractionation, with considerable wastage of plasma as a result;

Convinced that assuring the suitability of plasma for fractionation requires the establishment of a nationally coordinated and sustainable plasma programme within a properly organized, legally established and regulated national blood programme;

Recognizing that, as the capacity to collect plasma is limited and would not suffice to produce enough essential medicines to cover global needs, it is essential that all countries have local capacity to collect plasma of acceptable quality and safety from voluntary and unpaid donations in order to meet their needs;

Convinced that fractionation should be set up as close to the source as possible, and that, where national plasma fractionation capacities are lacking, there should be an option for supply of fractionation capacity in other countries, ensuring that the supply of plasma derived medicinal products can be made available to meet local needs in the country of the plasma supplier;

Recognizing that access to information about strategies to ensure supplies of blood products sufficient to meet demand, effective mechanisms of regulatory oversight, technologies to ensure the quality and safety of blood products, and guidelines on the appropriate clinical use of blood products and the risks of transfusion have become more and more necessary;

Bearing in mind that voluntary and non-remunerated blood donations can contribute to high safety standards for blood and blood components, and being aware that the safety of blood products depends on testing of all donated blood for transfusion-transmissible infections, and correct labelling, storage and transportation of blood products;

Bearing in mind that patient blood management means that before surgery every reasonable measure should be taken to optimize the patient’s own blood volume, to minimize the patient’s blood loss and to harness and optimize the patient-specific physiological tolerance of anaemia following WHO’s guide for optimal clinical use (three pillars of patient blood management);

Recognizing that excessive and unnecessary use of transfusions and of plasma-derived medicinal products, unsafe transfusion practices, and errors (particularly at the patient’s bedside) seriously compromise patient safety;

Concerned that unsafe and/or poor-quality blood products can render patients vulnerable to avoidable risk if the blood programmes are not subject to the level of control now exercised by experienced national or regional regulatory authorities;

Alarmed that patients in developing countries continue to be exposed to the risk of preventable transfusion-transmitted infections by bloodborne pathogens such as hepatitis B virus, hepatitis C virus and HIV;

Noting the increasing movement across boundaries of blood products and blood safety-related in vitro diagnostic devices, together with their rapid development and introduction into health-care systems of both developed and developing countries;
Recognizing the value of international biological reference materials (WHO International Standards) for the quality control of blood products and related in vitro diagnostic devices for detection of known and emerging bloodborne pathogens;

Convinced that traceability at all stages of the preparation of blood products, from the donor to the recipient and vice versa, is essential to identify risks, particularly the transmission of pathogens and transfusion reactions, and to monitor the efficacy of corrective measures aiming to minimize such risks;

Convinced that good practices need to be implemented for recruiting voluntary, non-remunerated healthy blood and plasma donors from low-risk donor populations and testing of all donated blood for transfusion-transmissible pathogens, and that the whole chain of processes in the production of blood products, i.e. correct processing, labelling, storage and transportation, needs to be covered by relevant, reliable quality-assurance systems;

Recognizing that stringent regulatory control is vital in assuring the quality and safety of blood products, as well as of related in vitro diagnostic devices, and that special effort is needed to strengthen globally the technical capacity of regulatory authorities to assure the appropriate control worldwide;

Recalling previous resolutions of the Health Assembly mentioning the vital need to strengthen blood establishments and ensure the quality, safety and efficacy of blood products,

1. **URGES** Member States:¹

(1) to take all the necessary steps to establish, implement and support nationally-coordinated, efficiently-managed and sustainable blood and plasma programmes according to the availability of resources, with the aim of achieving self-sufficiency, unless special circumstances preclude it;

(2) to take all the necessary steps to update their national legislation on donor assessment and deferral, the collection, testing, processing, storage, transportation and use of blood products, and operation of regulatory authorities in order to ensure that regulatory control in the area of quality and safety of blood products across the entire transfusion chain meets internationally recognized standards;

(3) to establish quality systems, for the processing of whole blood and blood components, good manufacturing practices for the production of plasma-derived medicinal products and appropriate regulatory control;

(4) to build human resource capacity through the provision of initial and continuing training of staff to ensure quality of blood services and blood products;

(5) to enhance the quality of evaluation and regulatory actions in the area of blood products and associated medical devices, including in vitro diagnostic devices;

(6) to establish or strengthen systems for the safe and rational use of blood products and to provide training for all staff involved in clinical transfusion, to implement potential solutions in order to minimize transfusion errors and promote patient safety, to promote

¹ And regional economic integration organizations, where applicable.
the availability of transfusion alternatives including, where appropriate, autologous transfusion and patient blood management;

(7) to ensure the reliability of mechanisms for reporting serious or unexpected adverse reactions to blood and plasma donation and to the receipt of blood components and plasma-derived medicinal products, including transmissions of pathogens;

2. REQUESTS the Director-General:

(1) to guide Member States to meet internationally recognized standards in updating their legislation, national standards and regulations for effective control of the quality and safety of blood products and associated medical devices, including in vitro diagnostics;

(2) to advise and build capacity in Member States on leadership and management of blood supply systems in order to strengthen national coordinated and sustainable blood and plasma programmes by sharing best practices about the organizational structure of blood supply systems in order to increase efficiency and minimize error;

(3) to augment the support offered to Member States for developing and strengthening their national regulatory authorities and control laboratories so as to increase their competence in the control of blood products and associated medical devices, including in vitro diagnostic devices, and to foster the creation of regional collaborative and regulatory networks where necessary and appropriate;

(4) to ensure sustainable development and provision of international biological reference materials (WHO International Standards) for use in the quality control and regulation of blood products and related in vitro diagnostic devices;

(5) to improve access by developing countries to international biological reference materials and to the scientific information obtained in their validation in order to assure the appropriate use of these materials;

(6) to develop, provide and disseminate guidance and technical support to strengthen national coordinated blood and plasma programmes and introduction of blood component separation and plasma fractionation technology, to meet local needs, and promote effective regulatory oversight of blood services and implementation of good manufacturing practices in plasma-fractionation programmes, under the responsibility of regulatory authorities;

(7) to provide guidance, training and support to Member States on safe and rational use of blood products and to support the introduction of transfusion alternatives including, where appropriate, autologous transfusion, safe transfusion practices and patient blood management;

(8) to encourage research into new technologies for producing safe and effective blood substitutes;

(9) to inform regularly, at least every four years, the Health Assembly, through the Executive Board, on actions taken by Member States and other partners to implement this resolution.

(Twelfth meeting, 22 January 2010)
EB126.R15 Accelerating progress towards achievement of Millennium Development Goal 4 to reduce child mortality: prevention and treatment of pneumonia

The Executive Board,

Having considered the report on treatment and prevention of pneumonia,\(^1\)

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Aware of the joint WHO/UNICEF report on a global action plan for the prevention and control of pneumonia, presented in November 2009;\(^2\)

Noting the first advance market commitment on the pneumococcal vaccine and the progress made so far in integrating the *Haemophilus influenzae* type b vaccine into routine immunization programmes;

Noting also the introduction of the pneumococcal Accelerated Development and Introduction Plans;

Recalling that resolution WHA58.15 on global immunization strategy requested the Director-General to mobilize resources to promote the availability and affordability in countries of future new vaccines based on evidence of epidemiological profiles;

Concerned at the lack of substantial progress towards reducing morbidity and mortality from pneumonia, despite it being globally the leading cause of mortality of children under the age of five years;

Mindful that decreasing the global burden of pneumonia will be essential for reaching Target 4.A of Millennium Development Goal 4;

Noting that safe and highly effective tools are available for pneumonia control in the form of WHO’s Integrated Management of Childhood Illness approach for case management at all levels, universal childhood immunization against *Haemophilus influenzae* type b and *Streptococcus pneumoniae* infections, improvement of nutrition and low birth weight, control of indoor air pollution arising from household use of solid fuels and second-hand smoking in households, and prevention and management of HIV infection;

Further noting that affordable price of vaccines in preventing pneumonia and significant scaling up of cold-chain capacities determine the adoption and implementation of vaccination programmes particularly in developing countries;

Concerned that pneumonia continues to cause more than 1.8 million preventable deaths in children less than five years of age globally each year;

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\(^1\) Document EB126/40.

Noting that the GAVI Alliance and other donors have made substantial resources available, and that the International Finance Facility for Immunisation and the PAHO revolving fund for immunization provide powerful mechanisms for directing resources to immunization programmes;

Welcoming the contribution to the mobilization of resources for development of voluntary innovative financing initiatives taken by groups of Member States;

Noting in addition that efforts to strengthen the capacity of health systems to detect and manage pneumonia effectively are likely also to contribute positively to efforts to achieve Millennium Development Goal 5 (Improve maternal health);

Aware that pandemic of influenza A (H1N1) 2009 has raised awareness of the need for system-wide strengthening of management of serious acute respiratory infections, and noting that the time is therefore opportune to build upon investments made related to the pandemic and to continue efforts to ensure that patients with acute respiratory infections receive prompt and effective treatment,

1. **URGES** Member States:

   (1) to apply, according to their specific contexts, the policies, strategies and tools recommended by WHO to prevent and treat pneumonia;

   (2) to establish evidence-based national policies and operational plans for strengthening health systems in order to expand coverage of populations at risk with effective preventive and curative interventions;

   (3) to assess programme performance including the coverage and impact of interventions in an effective and timely manner, and use this assessment to inform WHO’s country-profile database;

   (4) to identify national and international resources, both human and financial, for strengthening health systems and for the provision of technical support in order to ensure that the most locally and epidemiologically appropriate strategies are implemented and target populations reached;

   (5) to implement the recommendations in the joint WHO/UNICEF global action plan for the prevention and control of pneumonia, noting the importance of:

      (a) immunization by accelerating the adoption of affordable and cost-effective vaccines based on evidence of national epidemiological profiles;

      (b) case management at community, health-centre and hospital levels;

      (c) exclusive breastfeeding for six months;

      (d) improvement of nutrition and prevention of low birth weight;

      (e) control of indoor air pollution, and;

      (f) prevention and management of HIV infection;
(6) to encourage integrated approaches to pneumonia prevention and treatment through multisectoral collaboration and community responsibility and participation;

2. REQUESTS the Director-General:

(1) to strengthen human resources for prevention and control of pneumonia at all levels, especially the country level, thereby improving the capacity of WHO’s country offices to provide support to national health programmes for coordinating the work of partners on preventing and controlling pneumonia;

(2) to bring together interested Member States, organizations in the United Nations system, the GAVI Alliance, medical research councils, and other interested stakeholders in a forum in order to improve coordination between different stakeholders in the fight against pneumonia and mobilize resources to promote the availability of Haemophilus influenzae type b and pneumococcal vaccines;

(3) to expand the coverage of the report to the Health Assembly through the Executive Board on the status of progress made in achieving the health-related Millennium Development Goals, requested in resolution WHA61.18, to include progress on the implementation of this resolution, starting from the Sixty-fourth World Health Assembly.

(Twelfth meeting, 22 January 2010)

EB126.R16 Viral hepatitis

The Executive Board,

Having considered the report on viral hepatitis,¹

RECOMMENDS to the Sixty-third World Health Assembly the adoption of the following resolution:

The Sixty-third World Health Assembly,

Having considered the report on viral hepatitis;

Taking into account the fact that some 2000 million people have been infected by hepatitis B virus and that about 350 million people live with a chronic form of the disease;

Considering that hepatitis C is still not preventable by vaccination and around 80% of hepatitis C virus infections become a chronic infection;

Considering the seriousness of viral hepatitis as a global public health problem and the need for advocacy to both governments and populations for action on health promotion, disease prevention, diagnosis and treatment;

¹ Document EB126/15.
Expressing concern at the lack of progress in the prevention and control of viral hepatitis in developing countries, in particular in sub-Saharan Africa, due to the lack of access to affordable treatments as well as an integrated approach to the management of the disease;

Considering the need for a global approach to all forms of viral hepatitis – with a special focus on viral hepatitis B and C, which have the higher rates of morbidity;

Recalling that one route of transmission of hepatitis B and C viruses is parenteral and that the Health Assembly in resolution WHA28.72 on utilization and supply of human blood and blood products recommended the development of national public services for blood donation and in resolution WHA58.13 agreed to the establishment of an annual World Blood Donor Day, and that in both resolutions the Health Assembly recognized the need for safe blood to be available to blood recipients;

Reaffirming resolution WHA45.17 on immunization and vaccine quality which urged Member States to include hepatitis B vaccines in national immunization programmes;

Considering the need to reduce liver cancer mortality rates and that viral hepatitides are responsible for 78% of cases of primary liver cancer;

Considering the collaborative linkages between prevention and control measures for viral hepatitis and those for infectious diseases like HIV and other related sexually transmitted and blood-borne infections;

Recognizing the need to reduce incidence to prevent and control viral hepatitis, to increase access to correct diagnosis and to provide appropriate treatment programmes in all regions,

1. RESOLVES that 28 July shall be designated as World Hepatitis Day in order to provide an opportunity for education and greater understanding of viral hepatitis as a global public health problem, and to stimulate the strengthening of preventive and control measures of this disease in Member States;

2. URGES Member States:

   (1) to implement and/or improve epidemiological surveillance systems in order to generate reliable information for guiding prevention and control measures;

   (2) to support or enable an integrated and cost-effective approach to the prevention, control and management of viral hepatitis considering the linkages with associated coinfection such as HIV through multisectoral collaboration among health and educational institutions, nongovernmental organizations and civil society, including measures that strengthen safety and quality and the regulation of blood systems;

   (3) to incorporate in their specific contexts the policies, strategies and tools recommended by WHO in order to define and implement preventive actions, diagnostic measures and the provision of assistance to the population affected by viral hepatitis;

   (4) to strengthen national health systems in order to address prevention and control of viral hepatitis effectively through the provision of health promotion and national surveillance, including tools for prevention, diagnosis and treatment of viral hepatitis, vaccination, information, communication and injection safety;
(5) to provide vaccination strategies, infection-control measures, and means for injection safety for health-care workers;

(6) to use national and international resources, either human or financial, to provide technical support to strengthen health systems in order to provide local populations adequately with the most cost-effective and affordable interventions that suit the needs of local epidemiological situations;

(7) to consider, as necessary, national legislative mechanisms for the use of the flexibilities contained in the Agreement on Trade-Related Aspects of Intellectual Property Rights in order to promote access to specific pharmaceutical products;\(^1\)

(8) to consider, whenever necessary, using existing administrative and legal means in order to promote access to preventive, diagnostic and treatment technologies against viral hepatitis;

(9) to develop and implement monitoring and evaluation tools related to preventive, diagnostic and treatment activities;

(10) to promote the observance of 28 July each year, or on such other day or days as individual Member States may decide, as World Hepatitis Day;

3. REQUESTS the Director-General:

(1) to establish in collaboration with Member States the necessary guidelines, time-bound goals, strategies and tools for the prevention and control of viral hepatitis;

(2) to provide the necessary support to the development of scientific research related to the prevention, diagnosis and treatment of viral hepatitis;

(3) to improve the assessment of the economic impact and estimate the burden of viral hepatitis in the world;

(4) to support, as appropriate, resource-constrained Member States in conducting events to mark World Hepatitis Day;

(5) to invite international organizations and financial institutions to give support to strengthen capacity in developing countries for increasing the use of reliable diagnostic and treatment methods suitable to local epidemiological situations and health systems;

(6) to encourage international organizations and financial institutions to assign resources for the prevention and control of viral hepatitis, providing technical support to countries in an equitable, most efficient and suitable manner;

\(^1\) The WTO General Council in its Decision of 30 August 2003 (i.e. on Implementation of paragraph 6 of the Doha Declaration on the TRIPS Agreement and Public Health) decided that “‘pharmaceutical product’ means any patented product, or product manufactured through a patented process, of the pharmaceutical sector needed to address the public health problems as recognized in paragraph 1 of the Declaration. It is understood that active ingredients necessary for its manufacture and diagnostic kits needed for its use would be included.”
(7) to collaborate with other organizations in the United Nations system, partners, international organizations and other relevant stakeholders in enhancing access to affordable treatments in developing countries;

(8) to report to the Sixty-fifth World Health Assembly, through the Executive Board, on the implementation of this resolution.

(Thirteenth meeting, 23 January 2010)

**EB126.R17 Relations with nongovernmental organizations**

The Executive Board,

Having examined the report of its Standing Committee on Nongovernmental Organizations,

1. DECIDES to admit into official relations with WHO the International Insulin Foundation, International Life Saving Federation, Caritas Internationalis, Stichting Global Network of People Living with HIV/AIDS (GNP+), International Committee for Monitoring Assisted Reproductive Technologies and International Network for Cancer Treatment and Research;

2. DECIDES to defer consideration of an application from the International Union for Physical and Engineering Sciences in Medicine until such time as a new workplan has been agreed, and recommends that working relations be continued;

3. DECIDES to discontinue official relations with Cystic Fibrosis Worldwide, the International Association for Adolescent Health, the International Society of Hematology, and the World Association of Girl Guides and Girl Scouts;

4. DECIDES to suspend official relations with the Association of the Institutes and Schools of Tropical Medicine in Europe and the International Union for Conservation of Nature and Natural Resources until such time as they have submitted a report on their collaboration with WHO or, as the case may be, the status of their relations with WHO from 2006, for review by the Executive Board.

(Thirteenth meeting, 23 January 2010)

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1 See Annex 5, and Annex 7 for the financial and administrative implications for the Secretariat of this resolution.

DECISIONS

EB126(1) Appointment of members of the Independent Expert Oversight Advisory Committee

The Executive Board, having considered the report by the Secretariat on the appointment of members of the Independent Expert Oversight Advisory Committee, and recalling the terms of reference of the Committee, decided to appoint Ms Marion Cowden (Australia/New Zealand), Mr John Fox (United States of America), Mr Graham Miller (United Kingdom of Great Britain and Northern Ireland), Ms Hélène Ploix (France), and Mr Veerathai Santiprabhob (Thailand) as members of the Independent Expert Oversight Advisory Committee.

(Tenth meeting, 22 January 2010)

EB126(2) Appointment of representatives of the Executive Board at the Sixty-third World Health Assembly

Further to decision EB125(4) of 23 May 2009, and in accordance with paragraph 1 of resolution EB59.R7, the Executive Board appointed its Chairman, Dr S. Zaramba (Uganda), ex officio, and its first three Vice-Chairmen, Dr E.R. Sedyaningsih (Indonesia), Dr E. Giménez (Paraguay) and Professor Sohn Myongsei (Republic of Korea) to represent the Board at the Sixty-third World Health Assembly. It was understood that, if any of those members were not available for the Health Assembly, the other Vice-Chairman, Dr A.J. Mohamed (Oman), and the Rapporteur, Professor T. Milosavljević (Serbia), could be asked to represent the Board.

(Twelfth meeting, 22 January 2010)

EB126(3) Provisional agenda for, and duration of, the Sixty-third World Health Assembly

The Executive Board, having considered the report of the Director-General on the provisional agenda for the Sixty-third World Health Assembly, recalling its earlier decision that the Sixty-third World Health Assembly should be held at the Palais des Nations, Geneva, opening on Monday, 17 May 2010, and closing no later than Saturday, 22 May 2010, and recalling also the agreement made during the discussion of item 7.4 of the present session on the provisional draft agenda, approved the provisional agenda of the Sixty-third World Health Assembly, as amended, and decided that the Health Assembly would close no later than Friday, 21 May 2010.

(Twelfth meeting, 22 January 2010)

1 Document EB126/25.
2 Resolution EB125.R1, Annex.
3 See document EB126/27.
4 Decision EB125(6).
EB126(4) Date and place of the 127th session of the Executive Board

The Executive Board decided that its 127th session should be convened on Saturday, 22 May 2010, in Geneva.

(Twelfth meeting, 22 January 2010)

EB126(5) Review of nongovernmental organizations in official relations with WHO

The Executive Board, having considered and noted the report of its Standing Committee on Nongovernmental Organizations concerning the review of one third of the nongovernmental organizations in official relations with WHO,\(^1\) and following up decision EB124(1), reached the decisions set out below.

Noting with appreciation their collaboration with WHO and commending their continuing dedication to the work of WHO, the Board decided to maintain in official relations with WHO the nongovernmental organizations whose names are followed by an asterisk in the Annex to the report.

Noting the reports concerning their relations with WHO, the Board decided to maintain the International Association of Medical Regulatory Authorities, International Federation of Biomedical Laboratory Science, International Federation of Clinical Chemistry and Laboratory Medicine, International Society for Telemedicine & eHealth, International Organization against Trachoma and the International Association of Hydatid Disease in official relations with WHO and to request, as appropriate, reports on their collaboration during the period under review and the outcome of efforts to agree plans for collaboration, the results of which should be reported to the Board at its 128th session.

Noting the successful efforts to agree plans of collaboration the Board decided to maintain in official relations with WHO the International Association of Logopedics and Phoniatrics, Collegium Internationale Neuro-Psychopharmacologicum, and the International Union of Microbiological Societies.

Noting the report concerning the Secretariat’s introduction of an information system for management of the process of application by nongovernmental organizations for admission into official relations with WHO, the Board encouraged nongovernmental organizations in official relations with WHO to use the system and maintain their data therein up-to-date, at least on the occasion of their triennial review.

(Thirteenth meeting, 23 January 2010)

EB126(6) Award of the Dr A.T. Shousha Foundation Prize

The Executive Board, having considered the report of the Dr A.T. Shousha Foundation Prize Committee, awarded the Dr A.T. Shousha Foundation Prize for 2010 to Dr Faïssal A.R.M. Shaheen from Saudi Arabia for his significant contribution to public health in Saudi Arabia, in particular the development of kidney services and organ transplantation. The laureate will receive the equivalent of 2500 Swiss francs in United States dollars.

(Thirteenth meeting, 23 January 2010)

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\(^1\) Document EB126/28.

\(^2\) See Annex 5.
EB126(7) Award of the Sasakawa Health Prize

The Executive Board, having considered the report of the Sasakawa Health Prize Selection Panel, awarded the Sasakawa Health Prize for 2010 to Dr Du Xueping of China for her outstanding innovative work in health development. The laureate will receive US$ 30 000.

(Thirteenth meeting, 23 January 2010)

EB126(8) Award of the United Arab Emirates Health Foundation Prize

The Executive Board, having considered the report of the United Arab Emirates Health Foundation Selection Panel, awarded the United Arab Emirates Health Foundation Prize for 2010 to both the National Center for Diabetes, Endocrinology and Genetics, Jordan, and the Early Childhood Intervention Programme, Regional Administration of Health of Alentejo, Portugal, for their outstanding contribution to health development. The laureates will each receive US$ 20 000.

(Thirteenth meeting, 23 January 2010)

EB126(9) Award of the Dr LEE Jong-wook Memorial Prize for Public Health

The Executive Board, having considered the report of the Dr LEE Jong-wook Memorial Prize Selection Panel, awarded the Dr LEE Jong-wook Memorial Prize for 2010 to Action for AIDS, Singapore, for its outstanding contribution in research into and prevention, treatment and control of HIV/AIDS. The laureate will receive US$ 85 000.

(Thirteenth meeting, 23 January 2010)
ANNEXES
ANNEX 1

Birth defects

[EB126/10 – 3 December 2009]

1. In May 2009 the Executive Board at its 125th session considered an agenda item on birth defects. The Board noted the report on the subject but postponed further discussion to the present session of a draft resolution submitted by China, India and the Republic of Korea. This report is a revised version of the earlier report and reflects comments made by members of the Board. Document EB126/10 Add.1 contains the draft resolution initially considered by the Board and additionally reflects comments and proposals thereon made by members of the Board.

2. This report aims to inform the discussion on birth defects, including definition, epidemiology, burden of disease and interventions for prevention and care, as well as indications of how these interventions might be integrated into existing health services.

DEFINITION

3. The International statistical classification of diseases and related health problems Tenth Revision (ICD10), includes birth defects in Chapter XVII “Congenital malformations, deformations and chromosomal abnormalities”. Birth defects like inborn errors of metabolism and blood disorders of prenatal origin appear in other chapters. Birth defects can be defined as structural or functional abnormalities, including metabolic disorders, which are present from birth. The term congenital disorder is considered to have the same definition; the two terms are used interchangeably. The eleventh revision of the International statistical classification of diseases and related health problems provides an opportunity for a review of the current entry.

4. Irrespective of definition, birth defects can cause spontaneous abortions and stillbirths and are a significant but underrecognized cause of mortality and disability among infants and children under five years of age. They can be life-threatening, result in long-term disability, and negatively impact individuals, families, health-care systems and societies.

BIRTH DEFECTS AND GLOBAL NEWBORN AND CHILD MORTALITY

5. Congenital disorders are a common condition. WHO estimates that some 260 000 deaths worldwide (about 7% of all neonatal deaths) were caused by congenital anomalies in 2004. They are most prominent as a cause of death where overall mortality rates are lower, for example in the WHO European Region, where as many as 25% of neonatal deaths are due to congenital anomalies.

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1 See resolution EB126.R6.
2 See document EB125/2009/REC/1, summary record of the second meeting, section 1.
3 Document EB125/7.
6. Currently, sound estimates do not exist of the number of children who were born with a serious congenital disorder due to genetic or environmental causes. The most common serious congenital disorders are congenital heart defects, neural tube defects and Down syndrome. Haemoglobinopathies (including thalassaemia and sickle cell disease) and glucose-6-phosphate dehydrogenase deficiency, which are not covered by the ICD10 definition of congenital anomalies account for 6% of all congenital disorders.

7. Considerable uncertainties remain as to the incidence and mortality due to congenital disorders, especially in countries which lack adequate registration of deaths. However, these figures indicate that addressing the incidence and mortality associated with congenital anomalies needs to be linked to efforts to achieve the Millennium Development Goal 4 target, of a two thirds reduction in the mortality rate of children under five years of age, between 1990 and 2015.

COMMON CAUSES OF BIRTH DEFECTS

8. Birth defects are a diverse group of disorders of prenatal origin which can be caused by single gene defects, chromosomal disorders, multifactorial inheritance, environmental teratogens and micronutrient deficiencies. Maternal infections such as syphilis and rubella are a significant cause of birth defects in low- and middle-income countries. Maternal illnesses like diabetes mellitus, conditions such as iodine and folic acid deficiency, and exposure to medicinal and recreational drugs including alcohol and tobacco, certain environmental chemicals, and high doses of radiation are other factors that cause birth defects.

PREVENTION

9. The wide range of causes of birth defects means that a portfolio of prevention approaches is needed. Most birth defects of environmental origin can be prevented by public health approaches, including prevention of sexually-transmitted diseases, legislation controlling sound management of toxic chemicals (e.g. certain agricultural chemicals), vaccination against rubella, and fortification of basic foods with micronutrients (iodine and folic acid). Prevention may be considered in terms of life stage (see Appendix).

10. Preconception care aims to ensure the optimal physical and mental well-being of women, and their partners, at the onset of and during early pregnancy, to promote a normal pregnancy and the delivery of a healthy infant. It enables the timely deployment of primary prevention of teratogen-induced birth defects (including those caused by congenital syphilis and rubella), of defects caused by iodine deficiency disorder, of neural tube defects (and possibly other malformations), and of maternal-age-related chromosomal disorders (e.g. Down syndrome). The timely identification of a family risk of inherited disease, and carrier screening with genetic counselling, enable couples to limit family size where there is a known risk.

11. Prevention during pregnancy requires risk identification and management. Some of the interventions and services related to this can raise ethical, legal and social issues and may have cost implications. Such services include prenatal screening and diagnosis for birth defects, selective termination of pregnancy, and the availability of counselling services. Minimally invasive screening methods currently available involve the measurement of several metabolites in the maternal serum. Abnormal levels of biochemical markers are also associated with fetal structural defects such as Down syndrome, neural tube defects and open ventral wall defects. The detection rate of congenital disorders in the first trimester through biochemical screening is improved when it is done in tandem with
ultrasound screening involving nuchal translucency and other ultrasonographical assessments. Ultrasonography in the second trimester is useful to detect major structural defects.

DETECTION, TREATMENT AND CARE

12. Screening of newborn infants for congenital disorders facilitates early detection, treatment and care. Neonatal screening programmes (physical examination of all neonates, screening for congenital hypothyroidism, phenylketonuria, sickle cell disease and glucose-6-phosphate dehydrogenase deficiency) as well as training of primary care providers will support the diagnosis and appropriate referral for treatment of infants with congenital disorders. The physical examination of all newborn infants by trained primary care practitioners is practicable in most health systems and will allow the identification and referral of many birth defects, including cardiovascular defects that are associated with a high risk of early mortality.

13. Treatment of birth defects depends on the level of health care available. It comprises medical therapy, surgery, rehabilitation and palliative care when appropriate.

14. Effective life-saving medical treatment is available for several birth defects, including some common functional single gene defects. Examples include treatment of neonatal jaundice in glucose-6-phosphate dehydrogenase deficiency and with Rhesus incompatibility, congenital hypothyroidism, sickle cell disorders, thalassaemia, haemophilia, cystic fibrosis, and other inborn errors of metabolism. Other treatment options include in utero therapy and postnatal surgical corrections. These are now under research and evaluation in a few selected centres for a number of conditions (e.g. congenital diaphragmatic hernia, congenital heart lesions, myelomeningocele, twin-to-twin transfusion syndrome).

15. Surgery is an important but largely unheralded component of the services required to treat children with birth defects. Over 60% of children with a birth defect have a congenital malformation of a single organ, system or limb. Many birth defects are amenable to cost-effective surgery that can be life saving and improve long-term prognosis. Examples are surgery for simple congenital heart defects, cleft lip and palate, clubfoot, congenital cataracts, and gastrointestinal and urogenital abnormalities.

16. Appropriate treatment is also needed for impairments manifesting themselves after the neonatal period. This includes the early detection and treatment of physical, mental, intellectual or sensory impairments. Access to health and rehabilitation services is important to support the participation and inclusion of affected children.

17. With appropriate training, primary health care practitioners can offer basic care for children with birth defects; recognizing birth defects, diagnosing common problems and identifying associated disabilities. This facilitates basic treatment and counselling at the primary care level, taking into account family and community circumstances and available medical services. Referral to specialist advice is considered when diagnosis is not possible at the primary care level.

IMPLICATIONS FOR SERVICES

18. Services and interventions for the prevention and care of birth defects should be part of existing health-care services, in particular those concerned with maternal and child health. They should combine the best possible patient care with a preventive strategy encompassing education, pre-
conception care, population screening, genetic counselling, and the availability of diagnostic services. That strategy must deliver services for the prevention and care of birth defects as part of a continuum of interventions for maternal and child health. Depending on countries’ health-care capacity, the services should go beyond primary health care to include obstetric, paediatric, surgical, laboratory, radiological and, if available, clinical genetic services in secondary and tertiary health care.

19. Effective delivery of services for the prevention and care of birth defects depends on the availability of a range of specialist clinical and diagnostic services, and a primary care system that is able to use them. A nucleus of expertise in medical genetics, paediatric surgery, imaging, and fetal medicine is required, with the potential to expand to meet needs. Conventional laboratory services (haematological, microbiological, biochemical) need to be supplemented with cytogenetic and DNA-based diagnostic services. Introduction may need to be a gradual process. Over time, the new technologies will support more efficient and cost-effective service delivery.

20. The diversity of priority conditions, social structures, cultural conventions and health-care capabilities, means that countries need to be able to consider a range of possible services, assessing costs and relative effectiveness, in order to make a selection and decide the sequence of implementation. However, no organized guidance is yet available on this. The WHO Secretariat has an important potential role to play in identifying successful models, and providing coherent information on community genetics that is accessible to public health policy-makers.

POTENTIAL ACTIONS

21. There are several country-level actions that can support the development of services for the prevention and care of birth defects. Prevention requires basic public health approaches to be integrated into health systems including maternal and child health services. Many of the services and interventions proposed are already within the reach of low- and middle-income countries while others can be added as needs and resources determine.

22. Basic components for a national programme for the prevention and care of birth defects include:

(a) Commitment of policy-makers and provision of adequate managerial support.

(b) A core network of appropriate specialist clinical and laboratory services that can be expanded in response to demand.

(c) Integration of approaches for prevention and care of birth defects into primary health care, with emphasis on maternal and child health.

(d) Education and training for health-care providers, particularly those in primary health care.

(e) Organization of health education programmes for the general population and recognized high-risk groups.

(f) Establishment of effective mechanisms to foster development of patient–parent support organizations, and collaboration with them in caring for people with birth defects and their families.

(g) Definition of the ethical, legal, religious and cultural issues relevant to formulating services appropriate for the local population.
(h) Initiation and monitoring of population-screening programmes such as screening of newborn infants, premarital/pre-pregnancy screening, and screening during pregnancy.

(i) Establishment of appropriate surveillance systems for birth defects.\(^1\)

23. There is a need for technical guidance to establish or to strengthen national programmes for the control of birth defects. The following are priority actions for the international community:

(a) Resolve currently divergent opinions on the health burden of both environmental and constitutional birth defects, using the revision of ICD10 to draw on expert review of available data and to consider broadening the groups of conditions beyond those currently included in the classification of congenital anomalies.

(b) Promote legislation and public health activities to minimize exposure of the population, and particularly of pregnant women, to potentially teratogenic infections, chemicals and other environmental risk factors.

(c) Define effective community services, and support the integration of the prevention and care of birth defects into maternal and child health programmes. Support the provision to ministries of health of an organized assessment of requirements and costs and support in choosing priorities.

(d) Identify successful models that can be applied in low- and middle-income countries.

(e) Facilitate and support international networking on birth defect prevention and care programmes, with emphasis on developing common approaches, and optimizing instruments for information, education, cost analysis and surveillance, among others. Promote informatics approaches in view of their potential to support cost-effectiveness.

**ACTION BY THE EXECUTIVE BOARD**

24. [In this paragraph, the Board was invited to consider a draft resolution that was adopted at the seventh meeting as resolution EB126.R6.]

\(^1\) Support with this may be obtained by collaboration with existing birth defect surveillance systems including the International Clearinghouse for Birth Defects Surveillance and Research (ICBDSR) which includes ECLAMC (Latin American Collaboration Study of Congenital Malformation), the WHO-supported Craniofacial Anomalies database, and EUROCAT (European Registration of Congenital Anomalies).
### Interventions to prevent or treat birth defects

#### Preconception care

**Family planning**
- Introducing women to the concept of reproductive choice
- Reducing total number of children born with a birth defect
- Reducing the proportion of mothers of advanced maternal age, which reduces the birth prevalence of autosomal trisomies, particularly Down syndrome
- Allowing women with affected children the option of not having further children

**Preconception screening and counselling**
- Using family history taken in primary health care to identify individuals at risk of having affected children
- Carrier screening for common recessive disorders (thalassaemia and sickle cell disorder)

**Optimize women’s diet before and throughout pregnancy**
- Promote use of salt fortified with iodine to prevent iodine deficiency disorder
- Promote use of a staple food fortified with folic acid and use of supplementary multivitamins with folic acid to prevent neural tube defects and other malformations
- Avoid alcohol, tobacco and cocaine
- Ensure adequate general diet (protein, calories, iron)

**Prevent and treat teratogen-induced infections before and throughout pregnancy**
- Syphilis
- Rubella (67 countries do not have national rubella immunization programmes)

**Optimize pre-conception maternal health and treatment for**
- insulin-dependent diabetics
- women on treatment for epilepsy
- women on treatment with warfarin

#### Pregnancy care

**Antenatal screening for**
- Rhesus status
- Syphilis
- Individuals at risk of having children with birth defects using a family history
- Down syndrome: advanced maternal age; maternal serum screening; early ultrasound scanning
- Neural tube defects with maternal serum screening
- Major malformations with ultrasound fetal anomaly scanning (18+ weeks gestation)
- Carriers of common recessive disorders (thalassaemia and sickle cell disorder)

**Prenatal diagnosis**
- Ultrasound
- Amniocentesis
- Chorionic villus biopsy

**Fetal treatment**
- For syphilis
- Intrauterine transfusion for fetal anaemia Rhesus negativity

#### Newborn infant and child care

**Newborn infant examination**
- Trained examiner clinically examining all newborn infants for birth defects

**Newborn infant screening**
- Congenital hypothyroidism
- Phenylketonuria
- Cystic fibrosis
- Others, as dictated by each country’s needs and circumstances

**Medical treatment**
- Neonatal jaundice in glucose-6-phosphate dehydrogenase deficiency and Rhesus incompatibility
- Treatment and care for children with blood disorders like sickle cell disorder, thalassaemia, etc.
- Treatment of some inborn errors of metabolism
- Care of children with cystic fibrosis

**Surgery**
- Examples: correction of simple congenital heart defects
- cleft lip and palate
- clubfoot
- congenital cataracts

**Rehabilitation and palliative care**
- As appropriate
ANNEX 2

Text of amended Rules of Procedure of the Executive Board\(^1\)

[EB126/26, Annex – 10 December 2009]

CONDUCT OF BUSINESS

*Rule 30 bis*

The right of reply shall be accorded by the Chairman to any member who requests it. Members should, in exercising this right, attempt to be as brief as possible and preferably deliver their statements at the end of the meeting at which this right is requested.

\[\ldots\ldots\]

VOTING

*Rule 47 bis*

After the voting has been completed, a member may make a brief statement, consisting solely of an explanation of vote. A sponsor of a proposal shall not speak in explanation of vote thereon, except if it has been amended.

\[\ldots\ldots\]

*Rule 51 bis*

In an election each member, unless he abstains, shall vote for that number of candidates equal to the number of elective places to be filled. Any ballot paper on which there are more or fewer names than there are elective places to be filled shall be null and void.

*Rule 51 ter*

If during an election one or more elective places cannot be filled by reason of an equal number of votes having been obtained by two or more candidates, a ballot shall be held among such candidates to determine which of them will be elected. This procedure may be repeated if necessary.

\[\ldots\ldots\]

\(^1\) See resolution EB126.R8.
ANNEX 3

Confirmation of amendments to the Staff Rules

[EB126/39 – 26 November 2009]

1. Amendments to the Staff Rules made by the Director-General are submitted for confirmation by the Executive Board in accordance with Staff Regulation 12.2.1

2. The amendments described in this document reflect decisions expected to be taken by the United Nations General Assembly at its sixty-fourth session on the basis of recommendations made by the International Civil Service Commission in its annual report for 2009.2 Should the United Nations General Assembly not approve the Commission’s recommendations, an addendum to this document will be issued.

3. The financial implications of the amendments for the biennium 2009–2010 include negligible additional costs under the regular budget, which will be met from the appropriate allocations established for each of the WHO regions and for global and interregional activities, and from extrabudgetary sources of funds.

4. The amended Staff Rules are set out [in the Appendix].

AMENDMENTS CONSIDERED NECESSARY IN THE LIGHT OF DECISIONS EXPECTED TO BE TAKEN BY THE UNITED NATIONS GENERAL ASSEMBLY AT ITS SIXTY-FOURTH SESSION ON THE BASIS OF RECOMMENDATIONS OF THE INTERNATIONAL CIVIL SERVICE COMMISSION

Remuneration of staff in the professional and higher categories

5. The Commission recommended to the United Nations General Assembly that the current base/floor salary scale for the professional and higher categories should be increased by 3.04% through the standard consolidation method of increasing base salary and commensurately reducing post adjustment multiplier points (i.e. on a “no loss, no gain” basis) with effect from 1 January 2010.

6. Amendments to Appendix 1 of the Staff Rules have been prepared accordingly and are attached [Appendix].

Salaries of staff in ungraded posts and of the Director-General

7. Subject to the decision of the United Nations General Assembly in respect of the recommendation in paragraph 5 above, the Director-General proposes, in accordance with Staff Regulation 3.1, that the Executive Board should recommend to the Sixty-third World Health Assembly

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1 See resolutions EB126.R9 and EB126.R10.


modifications in the salaries of Assistant Directors-General and Regional Directors. Thus, as from 1 January 2010, the gross salary for Assistant Directors-General and Regional Directors would be US$ 183 022 per annum, and the net salary US$ 131 964 (dependency rate) or US$ 119 499 (single rate).

8. Based on the adjustments to salaries described above, the salary modification to be authorized by the Health Assembly for the Deputy Director-General would entail, as from 1 January 2010, a gross salary of US$ 201 351 per annum with a corresponding net salary of US$ 143 878 (dependency rate) or US$ 129 483 (single rate).

9. The salary adjustments described above would imply similar modifications to the salary of the Director-General. The salary to be authorized by the World Health Assembly, as of 1 January 2010 would therefore be US$ 247 523 per annum gross, US$ 173 890 net (dependency rate) or US$ 154 641 net (single rate).

**ACTION BY THE EXECUTIVE BOARD**

10. [This paragraph contained two draft resolutions, which were adopted at the ninth meeting as resolutions EB126.R9 and EB126.R10, respectively.]
Appendix 1 to the Staff Rules

Salary scale for staff in the professional and higher graded categories: annual gross base salaries and net equivalents after application of staff assessment (in US dollars)\(^1\) (effective 1 January 2010)

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\(D = \) Rate applicable to staff members with a dependent spouse or child; \(S = \) Rate applicable to staff members with no dependent spouse or child.

\(*) = \) The normal qualifying period for a within-grade increase between consecutive steps is one year, except at those steps marked with an asterisk, for which a two-year period at the preceding step is required (Staff Rule 550.2).
ANNEX 4

Draft global strategy to reduce the harmful use of alcohol

[EB126/13, Annex 1 – 3 December 2009]

Setting the scene

1. The harmful use of alcohol has a serious effect on public health and is considered to be one of the main risk factors for poor health globally. In the context of this draft strategy, the concept of the harmful use of alcohol is broad and encompasses the drinking that causes detrimental health and social consequences for the drinker, the people around the drinker and society at large, as well as the patterns of drinking that are associated with increased risk of adverse health outcomes. The harmful use of alcohol compromises both individual and social development. It can ruin the lives of individuals, devastate families, and damage the fabric of communities.

2. The harmful use of alcohol is a significant contributor to the global burden of disease and is listed as the third leading risk factor for premature deaths and disabilities in the world. It is estimated that 2.5 million people worldwide died of alcohol-related causes in 2004, including 320 000 young people between 15 and 29 years of age. Harmful use of alcohol was responsible for 3.8% of all deaths in the world in 2004 and 4.5% of the global burden of disease as measured in disability-adjusted life years lost, even when consideration is given to the modest protective effects, especially on coronary heart disease, of low consumption of alcohol for some people aged 40 years or older.

3. Harmful drinking is a major avoidable risk factor for neuropsychiatric disorders and other noncommunicable diseases such as cardiovascular diseases, cirrhosis of the liver and various cancers. For some diseases there is no evidence of a threshold effect in the relationship between the risk and level of alcohol consumption. The harmful use of alcohol is also associated with several infectious diseases like HIV/AIDS, tuberculosis and pneumonia. A significant proportion of the disease burden attributable to harmful drinking arises from unintentional and intentional injuries, including those due to road traffic crashes and violence, and suicides. Fatal injuries attributable to alcohol consumption tend to occur in relatively young people.

4. The degree of risk for harmful use of alcohol varies with age, sex and other biological characteristics of the consumer as well as with the setting and context in which the drinking takes place. Some vulnerable or at-risk groups and individuals have increased susceptibility to the toxic, psychoactive and dependence-producing properties of ethanol. At the same time low risk patterns of

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1 Revised in the light of comments made by the Executive Board at its 126th session (see document EB126/2010/REC/2, summary record of the eleventh meeting). See resolution EB126.R11.

2 An alcoholic beverage is a liquid that contains ethanol (ethyl alcohol, commonly called “alcohol”) and is intended for drinking. In most countries with a legal definition of “alcoholic beverage” a threshold for content of ethanol by volume in a beverage is set at ≥ 0.5% or 1.0%. The predominant categories of alcoholic beverages are beers, wines and spirits.

3 The word “harmful” in this strategy refers only to public-health effects of alcohol consumption, without prejudice to religious beliefs and cultural norms in any way.

alcohol consumption at the individual level may not be associated with occurrence or significantly increased probability of negative health and social consequences.

5. A substantial scientific knowledge base exists for policy-makers on the effectiveness and cost-effectiveness of strategies and interventions to prevent and reduce alcohol-related harm. Although much of the evidence comes from high-income countries, the results of meta-analyses and reviews of the available evidence provide sufficient knowledge to inform policy recommendations in terms of comparative effectiveness and cost-effectiveness of selected policy measures. With better awareness, there are increased responses at national, regional and global levels. However, these policy responses are often fragmented and do not always correspond to the magnitude of the impact on health and social development.

Challenges and opportunities

6. The present commitment to reducing the harmful use of alcohol provides a great opportunity for improving health and social well-being and for reducing the existing alcohol-attributable disease burden. However, there are considerable challenges that have to be taken into account in global or national initiatives or programmes. These include the following:

(a) **Increasing global action and international cooperation.** The current relevant health, cultural and market trends worldwide mean that harmful use of alcohol will continue to be a global health issue. These trends should be recognized and appropriate responses implemented at all levels. In this respect, there is a need for global guidance and increased international collaboration to support and complement regional and national actions.

(b) **Ensuring intersectoral action.** The diversity of alcohol-related problems and measures necessary to reduce alcohol-related harm points to the need for comprehensive action across numerous sectors. Policies to reduce the harmful use of alcohol must reach beyond the health sector, and appropriately engage such sectors as development, transport, justice, social welfare, fiscal policy, trade, agriculture, consumer policy, education and employment, as well as civil society and economic operators.

(c) **According appropriate attention.** Preventing and reducing harmful use of alcohol is often given a low priority among decision-makers despite compelling evidence of its serious public health effects. In addition, there is a clear discrepancy between the increasing availability and affordability of alcohol beverages in many developing and low- and middle-income countries and those countries’ capability and capacity to meet the additional public health burden that follows. Unless this problem is given the attention it deserves, the spread of harmful drinking practices and norms will continue.

(d) **Balancing different interests.** Production, distribution, marketing and sales of alcohol create employment and generate considerable income for economic operators and tax revenue for governments at different levels. Public health measures to reduce harmful use of alcohol are sometimes judged to be in conflict with other goals like free markets and consumer choice and can be seen as harming economic interests and reducing government revenues. Policy-makers face the challenge of giving an appropriate priority to the promotion and protection of

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1 See document A60/14 for evidence-based strategies and interventions to reduce alcohol-related harm.

population health while taking into account other goals, obligations, including international legal obligations, and interests. It should be noted in this respect that international trade agreements generally recognize the right of countries to take measures to protect human health, provided that these are not applied in a manner which would constitute a means of unjustifiable or arbitrary discrimination or disguised restrictions to trade. In this regard, national, regional and international efforts should take into account the impact of harmful use of alcohol.

(e) **Focusing on equity.** Population-wide rates of drinking of alcoholic beverages are markedly lower in poorer societies than in wealthier ones. However, for a given amount of consumption, poorer populations may experience disproportionately higher levels of alcohol-attributable harm. There is a great need to develop and implement effective policies and programmes that reduce such social disparities both inside a country and between countries. Such policies are also needed in order to generate and disseminate new knowledge about the complex relationship between harmful consumption of alcohol and social and health inequity, particularly among indigenous populations, minority or marginalized groups and in developing countries.

(f) **Considering the “context” in recommending actions.** Much of the published evidence of effectiveness of alcohol-related policy interventions comes from high-income countries, and concerns have been expressed that their effectiveness depends on context and may not be transferrable to other settings. However, many interventions to reduce harmful use of alcohol have been implemented in a wide variety of cultures and settings, and their results are often consistent and in line with the underpinning theories and evidence base accumulated in other similar public health areas. The focus for those developing and implementing policies should be on appropriate tailoring of effective interventions to accommodate local contexts and on appropriate monitoring and evaluation to provide feedback for further action.

(g) **Strengthening information.** Systems for collecting, analysing and disseminating data on alcohol consumption, alcohol-related harm and policy responses have been developed by Member States, the WHO Secretariat, and some other stakeholders. There are still substantial gaps in knowledge and it is important to sharpen the focus on information and knowledge production and dissemination for further developments in this area, especially in developing and low- and middle-income countries. The WHO Global Information System on Alcohol and Health and integrated regional information systems provide the means to monitor better progress made in reducing harmful use of alcohol at the global and regional levels.

### Aims and objectives

7. National and local efforts can produce better results when they are supported by regional and global action within agreed policy frames. Thus the purpose of the global strategy is to support and complement public health policies in Member States.

8. The vision behind the global strategy is improved health and social outcomes for individuals, families and communities, with considerably reduced morbidity and mortality due to harmful use of alcohol and their ensuing social consequences. It is envisaged that the global strategy will promote and support local, regional and global actions to prevent and reduce the harmful use of alcohol.  

9. The global strategy aims to give guidance for action at all levels; to set priority areas for global action; and to recommend a portfolio of policy options and measures that could be considered for implementation and adjusted as appropriate at the national level, taking into account national circumstances, such as religious and cultural contexts, national public health priorities, as well as resources, capacities and capabilities.
10. The strategy has five objectives:

   (a) raised global awareness of the magnitude and nature of the health, social and economic problems caused by harmful use of alcohol, and increased commitment by governments to act to address the harmful use of alcohol;

   (b) strengthened knowledge base on the magnitude and determinants of alcohol-related harm and on effective interventions to reduce and prevent such harm;

   (c) increased technical support to, and enhanced capacity of, Member States for preventing the harmful use of alcohol and managing alcohol-use disorders and associated health conditions;

   (d) strengthened partnerships and better coordination among stakeholders and increased mobilization of resources required for appropriate and concerted action to prevent the harmful use of alcohol;

   (e) improved systems for monitoring and surveillance at different levels, and more effective dissemination and application of information for advocacy, policy development and evaluation purposes.

11. The harmful use of alcohol and its related public health problems are influenced by the general level of alcohol consumption in a population, drinking patterns and local contexts. Achieving the five objectives will require global, regional and national actions on the levels, patterns and contexts of alcohol consumption and the wider social determinants of health. Special attention needs to be given to reducing harm to people other than the drinker and to populations that are at particular risk from harmful use of alcohol, such as children, adolescents, women of child-bearing age, pregnant and breastfeeding women, indigenous peoples and other minority groups or groups with low socioeconomic status.

**Guiding principles**

12. The protection of the health of the population by preventing and reducing the harmful use of alcohol is a public health priority. The following principles will guide the development and implementation of policies at all levels; they reflect the multifaceted determinants of alcohol-related harm and the concerted multisectoral actions required to implement effective interventions.

   (a) Public policies and interventions to prevent and reduce alcohol-related harm should be guided and formulated by public health interests and based on clear public health goals and the best available evidence.

   (b) Policies should be equitable and sensitive to national, religious and cultural contexts.

   (c) All involved parties have the responsibility to act in ways that do not undermine the implementation of public policies and interventions to prevent and reduce harmful use of alcohol.

   (d) Public health should be given proper deference in relation to competing interests and approaches that support that direction should be promoted.

   (e) Protection of populations at high risk of alcohol-attributable harm and those exposed to the effects of harmful drinking by others should be an integral part of policies addressing the harmful use of alcohol.
(f) Individuals and families affected by the harmful use of alcohol should have access to affordable and effective prevention and care services.

(g) Children, teenagers and adults who choose not to drink alcohol beverages have the right to be supported in their non-drinking behaviour and protected from pressures to drink.

(h) Public policies and interventions to prevent and reduce alcohol-related harm should encompass all alcoholic beverages and surrogate alcohol.¹

**National policies and measures**

13. The harmful use of alcohol can be reduced if effective actions are taken by countries to protect their populations. Member States have a primary responsibility for formulating, implementing, monitoring and evaluating public policies to reduce the harmful use of alcohol. Such policies require a wide range of public health-oriented strategies for prevention and treatment. All countries will benefit from having a national strategy and appropriate legal frameworks to reduce harmful use of alcohol, regardless of the level of resources in the country. Depending on the characteristics of policy options and national circumstances, some policy options can be implemented by non-legal frameworks such as guidelines or voluntary restraints. Successful implementation of measures should be assisted by monitoring impact and compliance and establishing and imposing sanctions for non-compliance with adopted laws and regulations.

14. Sustained political commitment, effective coordination, sustainable funding and appropriate engagement of subnational governments as well as from civil society and economic operators are essential for success. Many relevant decision-making authorities should be involved in the formulation and implementation of alcohol policies, such as health ministries, transportation authorities or taxation agencies. Governments need to establish effective and permanent coordination machinery, such as a national alcohol council, comprising senior representatives of many ministries and other partners, in order to ensure a coherent approach to alcohol policies and a proper balance between policy goals in relation to harmful use of alcohol and other public policy goals.

15. Health ministries have a crucial role in bringing together the other ministries and stakeholders needed for effective policy design and implementation. They should also ensure that planning and provision of prevention and treatment strategies and interventions are coordinated with those for other related health conditions with high public health priority such as illicit drug use, mental illness, violence and injuries, cardiovascular diseases, cancer, tuberculosis and HIV/AIDS.

16. The policy options and interventions available for national action can be grouped into 10 recommended target areas, which should be seen as supportive and complementary to each other. These 10 areas are:

   (a) leadership, awareness and commitment

   (b) health services’ response

   (c) community action

   (d) drink-driving policies and countermeasures

¹ In this strategy “surrogate alcohol” refers to liquids usually containing ethanol and not intended for consumption as beverages, that are consumed orally as substitutes for alcoholic beverages with the objective to producing intoxication or other effects associated with alcohol consumption.
(e) availability of alcohol

(f) marketing of alcoholic beverages

(g) pricing policies

(h) reducing the negative consequences of drinking and alcohol intoxication

(i) reducing the public health impact of illicit alcohol and informally produced alcohol

(j) monitoring and surveillance.

17. The policy options and interventions proposed below for consideration by Member States for each of the 10 recommended target areas are based on current scientific knowledge, available evidence on effectiveness and cost-effectiveness, experience and good practices. Not all the policy options and interventions will be applicable or relevant for all Member States and some may be beyond available resources. As such, the measures should be implemented at the discretion of each Member State depending on national, religious and cultural contexts, national public health priorities, and available resources, and in accordance with constitutional principles and international legal obligations. Policy measures and interventions at the national level will be supported and complemented by global and regional efforts to reduce the harmful use of alcohol.

POLICY OPTIONS AND INTERVENTIONS

Area 1. Leadership, awareness and commitment

18. Sustainable action requires strong leadership and a solid base of awareness and political will and commitment. The commitments should ideally be expressed through adequately funded comprehensive and intersectoral national policies that clarify the contributions, and division of responsibility, of the different partners involved. The policies must be based on available evidence and tailored to local circumstances, with clear objectives, strategies and targets. The policy should be accompanied by a specific action plan and supported by effective and sustainable implementation and evaluation mechanisms. The appropriate engagement of civil society and economic operators is essential.

19. For this area policy options and interventions include:

(a) developing or strengthening existing, comprehensive national and subnational strategies, plans of action and activities to reduce the harmful use of alcohol;

(b) establishing or appointing a main institution or agency, as appropriate, to be responsible for following up national policies, strategies and plans;

(c) coordinating alcohol strategies with work in other relevant sectors, including cooperation between different levels of governments, and with other relevant health-sector strategies and plans;

1 Informally produced alcohol means alcoholic beverages produced at home or locally by fermentation and distillation of fruits, grains, vegetables and the like, and often within the context of local cultural practices and traditions. Examples of informally produced alcoholic beverages include sorghum beer, palm wine and spirits produced from sugarcane, grains or other commodities.
(d) ensuring broad access to information and effective education and public awareness programmes among all levels of society about the full range of alcohol-related harm experienced in the country and the need for, and existence of, effective preventive measures;

(e) raising awareness of harm to others and among vulnerable groups caused by drinking, avoiding stigmatization and actively discouraging discrimination against affected groups and individuals.

**Area 2. Health services’ response**

20. Health services are central to tackling harm at the individual level among those with alcohol-use disorders and other health conditions caused by harmful use of alcohol. Health services should provide prevention and treatment interventions to individuals and families at risk of, or affected by, alcohol-use disorders and associated conditions. Another important role of health services and health professionals is to inform societies about the public health and social consequences of harmful use of alcohol, support communities in their efforts to reduce the harmful use of alcohol, and to advocate effective societal responses. Health services should reach out to, mobilize and involve a broad range of players outside the health sector. Health services response should be sufficiently strengthened and funded in a way that is commensurate with the magnitude of the public health problems caused by harmful use of alcohol.

21. For this area **policy options and interventions** include:

(a) increasing capacity of health and social welfare systems to deliver prevention, treatment and care for alcohol-use and alcohol-induced disorders and co-morbid conditions, including support and treatment for affected families and support for mutual help or self-help activities and programmes;

(b) supporting initiatives for screening and brief interventions for hazardous and harmful drinking at primary health care and other settings; such initiatives should include early identification and management of harmful drinking among pregnant women and women of child-bearing age;

(c) improving capacity for prevention of, identification of, and interventions for individuals and families living with fetal alcohol syndrome and a spectrum of associated disorders;

(d) development and effective coordination of integrated and/or linked prevention, treatment and care strategies and services for alcohol-use disorders and co-morbid conditions, including drug-use disorders, depression, suicides, HIV/AIDS and tuberculosis;

(e) securing universal access to health including through enhancing availability, accessibility and affordability of treatment services for groups of low socioeconomic status;

(f) establishing and maintaining a system of registration and monitoring of alcohol-attributable morbidity and mortality, with regular reporting mechanisms;

(g) provision of culturally sensitive health and social services as appropriate.

**Area 3. Community action**

22. The impact of harmful use of alcohol on communities can trigger and foster local initiatives and solutions to local problems. Communities can be supported and empowered by governments and other
stakeholders to use their local knowledge and expertise in adopting effective approaches to prevent and reduce the harmful use of alcohol by changing collective rather than individual behaviour while being sensitive to cultural norms, beliefs and value systems.

23. For this area policy options and interventions include:

(a) supporting rapid assessments in order to identify gaps and priority areas for interventions at the community level;

(b) facilitating increased recognition of alcohol-related harm at the local level and promoting appropriate effective and cost-effective responses to the local determinants of harmful use of alcohol and related problems;

(c) strengthening capacity of local authorities to encourage and coordinate concerted community action by supporting and promoting the development of municipal policies to reduce harmful use of alcohol, as well as their capacity to enhance partnerships and networks of community institutions and nongovernmental organizations;

(d) providing information about effective community-based interventions, and building capacity at community level for their implementation;

(e) mobilizing communities to prevent the selling of alcohol to, and consumption of alcohol by, under-age drinkers, and to develop and support alcohol-free environments, especially for youth and other at-risk groups;

(f) providing community care and support for affected individuals and their families;

(g) developing or supporting community programmes and policies for subpopulations at particular risk, such as young people, unemployed persons and indigenous populations, specific issues like the production and distribution of illicit or informal-alcohol beverages and events at community level such as sporting events and town festivals.

Area 4. Drink–driving policies and countermeasures

24. Driving under the influence of alcohol seriously affects a person’s judgment, coordination and other motor functions. Alcohol-impaired driving is a significant public health problem that affects both the drinker and in many cases innocent parties. Strong evidence-based interventions exist for reducing drink–driving. Strategies to reduce harm associated with drink–driving should include deterrent measures that aim to reduce the likelihood that a person will drive under the influence of alcohol, and measures that create a safer driving environment in order to reduce both the likelihood and severity of harm associated with alcohol-influenced crashes.

25. In some countries, the number of traffic-related injuries involving intoxicated pedestrians is substantial and should be a high priority for intervention.

26. For this area policy options and interventions include:

(a) introducing and enforcing an upper limit for blood alcohol concentration, with a reduced limit for professional drivers and young or novice drivers;

(b) promoting sobriety check points and random breath-testing;
(c) administrative suspension of driving licences;

(d) graduated licensing for novice drivers with zero-tolerance for drink–driving;

(e) using an ignition interlock, in specific contexts where affordable, to reduce drink-driving incidents;

(f) mandatory driver-education, counselling and, as appropriate, treatment programmes;

(g) encouraging provision of alternative transportation, including public transport until after the closing time for drinking places;

(h) conducting public awareness and information campaigns in support of policy and in order to increase the general deterrence effect;

(i) running carefully planned, high-intensity, well-executed mass media campaigns targeted at specific situations, such as holiday seasons, or audiences such as young people.

**Area 5. Availability of alcohol**

27. Public health strategies that seek to regulate the commercial or public availability of alcohol through laws, policies, and programmes are important ways to reduce the general level of harmful use of alcohol. Such strategies provide essential measures to prevent easy access to alcohol by vulnerable and high-risk groups. Commercial and public availability of alcohol can have a reciprocal influence on the social availability of alcohol and thus contribute to changing social and cultural norms that promote harmful use of alcohol. The level of regulation on the availability of alcohol will depend on local circumstances, including social, cultural and economic contexts as well as existing binding international obligations. In some developing and low- and middle-income countries, informal markets are the main source of alcohol and formal controls on sale need to be complemented by actions addressing illicit or informally produced alcohol. Furthermore, restrictions on availability that are too strict may promote the development of a parallel illicit market. Secondary supply of alcohol, for example from parents or friends, needs also to be taken into consideration in measures on the availability of alcohol.

28. For this area **policy options and interventions** include:

(a) establishing, operating and enforcing an appropriate system to regulate production, wholesaling and serving of alcoholic beverages that places reasonable limitations on the distribution of alcohol and the operation of alcohol outlets in accordance with cultural norms, by the following possible measures:

   (i) introducing, where appropriate, a licensing system on retail sales, or public health-oriented government monopolies;

   (ii) regulating the number and location of on-premise and off-premise alcohol outlets;

   (iii) regulating days and hours of retail sales;

   (iv) regulating modes of retail sales of alcohol;

   (v) regulating retail sales in certain places or during special events;
(b) establishing an appropriate minimum age for purchase or consumption of alcoholic beverages and other policies in order to raise barriers against sales to, and consumption of alcoholic beverages by, adolescents;

(c) adopting policies to prevent sales to intoxicated persons and those below the legal age and considering the introduction of mechanisms for placing liability on sellers and servers in accordance with national legislations;

(d) setting policies regarding drinking in public places or at official public agencies’ activities and functions;

(e) adopting policies to reduce and eliminate availability of illicit production, sale and distribution of alcoholic beverages as well as to regulate or control informal alcohol.

**Area 6. Marketing\(^1\) of alcoholic beverages**

29. Reducing the impact of marketing, particularly on young people and adolescents, is an important consideration in reducing harmful use of alcohol. Alcohol is marketed through increasingly sophisticated advertising and promotion techniques, including linking alcohol brands to sports and cultural activities, sponsorships and product placements, and new marketing techniques such as e-mails, SMS and podcasting, social media and other communication techniques. The transmission of alcohol marketing messages across national borders and jurisdictions on channels such as satellite television and the Internet, and sponsorship of sports and cultural events is emerging as a serious concern in some countries.

30. It is very difficult to target young adult consumers without exposing cohorts of adolescents under the legal age to the same marketing. The exposure of children and young people to appealing marketing is of particular concern, as is the targeting of new markets in developing and low- and middle-income countries with a current low prevalence of alcohol consumption or high abstinence rates. Both the content of alcohol marketing and the amount of exposure of young people to that marketing are crucial issues. A precautionary approach to protecting young people against these marketing techniques should be considered.

31. For this area **policy options and interventions** include:

   (a) setting up regulatory or co-regulatory frameworks, preferably with a legislative basis, and supported when appropriate by self-regulatory measures, for alcohol marketing by:

      (i) regulating the content and the volume of marketing;

      (ii) regulating direct or indirect marketing in certain or all media;

      (iii) regulating sponsorship activities that promote alcoholic beverages;

      (iv) restricting or banning promotions in connection with activities targeting young people;

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\(^1\) Marketing could refer, as appropriate and in accordance with national legislation, to any form of commercial communication or message that is designed to increase, or has the effect of increasing, the recognition, appeal and/or consumption of particular products and services. It could comprise anything that acts to advertise or otherwise promote a product or service.
(v) regulating new forms of alcohol marketing techniques, for instance social media;

(b) development by public agencies or independent bodies of effective systems of surveillance of marketing of alcohol products;

(c) setting up effective administrative and deterrence systems for infringements on marketing restrictions.

Area 7. Pricing policies

32. Consumers, including heavy drinkers and young people, are sensitive to changes in the price of drinks. Pricing policies can be used to reduce underage drinking, to halt progression towards drinking large volumes of alcohol and/or episodes of heavy drinking, and to influence consumers’ preferences. Increasing the price of alcoholic beverages is one of the most effective interventions to reduce harmful use of alcohol. A key factor for the success of price-related policies in reducing harmful use of alcohol is an effective and efficient system for taxation matched by adequate tax collection and enforcement.

33. Factors such as consumer preferences and choice, changes in income, alternative sources for alcohol in the country or in neighbouring countries, and the presence or absence of other alcohol policy measures may influence the effectiveness of this policy option. Demand for different beverages may be affected differently. Tax increases can have different impacts on sales, depending on how they affect the price to the consumer. The existence of a substantial illicit market for alcohol complicates policy considerations on taxation in many countries. In such circumstances tax changes must be accompanied by efforts to bring the illicit and informal markets under effective government control. Increased taxation can also meet resistance from consumer groups and economic operators, and taxation policy will benefit from the support of information and awareness-building measures to counter such resistance.

34. For this area policy options and interventions include:

(a) establishing a system for specific domestic taxation, on alcohol accompanied by an effective enforcement system, which may take into account, as appropriate, the alcoholic content of the beverage;

(b) regularly reviewing prices in relation to level of inflation and income;

(c) banning or restricting the use of direct and indirect price promotions, discount sales, sales below cost and flat rates for unlimited drinking or other types of volume sales;

(d) establishing minimum prices for alcohol where applicable;

(e) providing price incentives for non-alcoholic beverages;

(f) reducing or stopping subsidies to economic operators in the area of alcohol.

Area 8. Reducing the negative consequences of drinking and alcohol intoxication

35. This target area includes policy options and interventions that focus directly on reducing the harm from alcohol intoxication and drinking without necessarily affecting the underlying alcohol consumption. Current evidence and good practices favour the complementary use of interventions within a broader strategy that prevents or reduces the negative consequences of drinking and alcohol.
intoxication. In implementing these approaches, managing the drinking environment or informing consumers, the perception of endorsing or promoting drinking should be avoided.

36. For this area **policy options and interventions** include:

   (a) regulating the drinking context in order to minimize violence and disruptive behaviour, including serving alcohol in plastic containers or shatter-proof glass and management of alcohol-related issues at large-scale public events;

   (b) enforcing laws against serving to intoxication and legal liability for consequences of harm resulting from intoxication caused by the serving of alcohol;

   (c) enacting management policies relating to responsible serving of beverage on premises and training staff in relevant sectors in how better to prevent, identify and manage intoxicated and aggressive drinkers;

   (d) reducing the alcoholic strength inside different beverage categories;

   (e) providing necessary care or shelter for severely intoxicated people;

   (f) providing consumer information about, and labelling alcoholic beverages to indicate, the harm related to alcohol.

**Area 9. Reducing the public health impact of illicit alcohol and informally produced alcohol**

37. Consumption of illicitly or informally produced alcohol could have additional negative health consequences due to a higher ethanol content and potential contamination with toxic substances, such as methanol. It may also hamper governments’ abilities to tax and control legally produced alcohol. Actions to reduce these additional negative effects should be taken according to the prevalence of illicit and/or informal alcohol consumption and the associated harm. Good scientific, technical and institutional capacity should be in place for the planning and implementation of appropriate national, regional and international measures. Good market knowledge and insight into the composition and production of informal or illicit alcohol are also important, coupled with an appropriate legislative framework and active enforcement. These interventions should complement, not replace, other interventions to reduce harmful use of alcohol.

38. Production and sale of informal alcohol are ingrained in many cultures and are often informally controlled. Thus control measures could be different for illicit alcohol and informally produced alcohol and should be combined with awareness raising and community mobilization. Efforts to stimulate alternative sources of income are also important.

39. For this area **policy options and interventions** include:

   (a) good quality control with regard to production and distribution of alcoholic beverages;

   (b) regulating sales of informally produced alcohol and bringing it into the taxation system;

   (c) an efficient control and enforcement system, including tax stamps;

   (d) developing or strengthening tracking and tracing systems for illicit alcohol;
(e) ensuring necessary cooperation and exchange of relevant information on combating illicit alcohol among authorities at national and international levels;

(f) issuing relevant public warnings about contaminants and other health threats from informal or illicit alcohol.

Area 10. Monitoring and surveillance

40. Data from monitoring and surveillance create the basis for the success and appropriate delivery of the other nine policy options. Local, national and international monitoring and surveillance are needed in order to monitor the magnitude and trends of alcohol-related harms, to strengthen advocacy, to formulate policies and to assess impact of interventions. Monitoring should also capture the profile of people accessing services and the reason why people most affected are not accessing prevention and treatment services. Data may be available in other sectors, and good systems for coordination, information exchange and collaboration are necessary in order to collect the potentially broad range of information needed to have comprehensive monitoring and surveillance.

41. Development of sustainable national information systems using indicators, definitions and data-collection procedures compatible with WHO’s global and regional information systems provides an important basis for effective evaluation of national efforts to reduce harmful use of alcohol and for monitoring trends at subregional, regional and global levels. Systematic continual collection, collation and analysis of data, timely dissemination of information and feedback to policy-makers and other stakeholders should be an integral part of implementation of any policy and intervention to reduce harmful use of alcohol. Collecting, analysing and disseminating information on harmful use of alcohol are resource-intensive activities.

42. For this area policy options and interventions include:

(a) establishing effective frameworks for monitoring and surveillance activities including periodic national surveys on alcohol consumption and alcohol-related harm and a plan for exchange and dissemination of information;

(b) establishing or designating an institution or other organizational entity responsible for collecting, collating, analysing and disseminating available data, including publishing national reports;

(c) defining and tracking a common set of indicators of harmful use of alcohol and of policy responses and interventions to prevent and reduce such use;

(d) creating a repository of data at the country level based on internationally agreed indicators and reporting data in the agreed format to WHO and other relevant international organizations;

(e) developing evaluation mechanisms with the collected data in order to determine the impact of policy measures, interventions and programmes put in place to reduce the harmful use of alcohol.

GLOBAL ACTION: KEY ROLES AND COMPONENTS

43. Given the magnitude and the complexity of the problem, concerted global efforts must be in place to support Member States in the challenges they face at the national level. International
coordination and collaboration create the synergies that are needed and provide increased leverage for Member States to implement evidence-based measures.

44. WHO, in cooperation with other organizations in the United Nations system and other international partners will:

(a) provide leadership;
(b) strengthen advocacy;
(c) formulate, in collaboration with Member States, evidence-based policy options;
(d) promote networking and exchange of experience among countries;
(e) strengthen partnerships and resource mobilization;
(f) coordinate monitoring of alcohol-related harm and the progress countries are making to address it.

45. Action by WHO and other international partners to support the implementation of the global strategy will be taken according to their mandates. International nongovernmental organizations, professional associations, research institutions and economic operators in the area of alcohol, all have important roles in enhancing the global action, as follows.

(a) Major partners within the United Nations system and intergovernmental organizations like ILO, UNICEF, WTO, UNDP, UNFPA, UNAIDS, United Nations Office on Drugs and Crime, and the World Bank group will be urged to increase collaboration and cooperation to prevent and reduce harmful use of alcohol, especially in developing and low- and middle-income countries.

(b) Civil society has an important role in warning about the impact of harmful use of alcohol on individuals, families and communities and in bringing additional commitment and resources for reducing alcohol-related harm. Nongovernmental organizations are especially encouraged to form wide networks and action groups to support the implementation of the global strategy.

(c) Research institutions and professional associations play a pivotal role in generating additional evidence for action and disseminating this to health professionals and the wider community. WHO collaborating centres have an important role in supporting the implementation and evaluation of the global strategy.

(d) Economic operators in alcohol production and trade are important players in their role as developers, producers, distributors, marketers and sellers of alcoholic beverages. They are especially encouraged to consider effective ways to prevent and reduce harmful use of alcohol within their core roles mentioned above, including self-regulatory actions and initiatives. They could also contribute by making available data on sales and consumption of alcohol beverages.

(e) The media play an increasingly important role, not only as a conveyer of news and information but also as a channel for commercial communications, and will be encouraged to support the intentions and activities of the global strategy.
Public health advocacy and partnership

46. International public health advocacy and partnership are needed for strengthened commitment and abilities of the governments and all relevant parties at all levels for reducing harmful use of alcohol worldwide.

47. WHO is committed to raising awareness of the public health problems caused by harmful use of alcohol and of the steps that can be taken to prevent and reduce such use in order to save lives and reduce suffering. WHO will engage with other international intergovernmental organizations and, as appropriate, international bodies representing key stakeholders, to ensure that relevant actors can contribute to reducing the harmful use of alcohol.

48. The Secretariat will provide support to Member States by:

(a) raising the awareness of the magnitude of public health problems caused by harmful use of alcohol and advocating for appropriate action at all levels to prevent and reduce such problems;

(b) advocating that attention is given to addressing the harmful use of alcohol in the agendas of relevant international and intergovernmental organizations in order to support policy coherence between health and other sectors at regional and global levels;

(c) promoting and facilitating international coordination, collaboration, partnerships and information exchange to ensure the needed synergies and concerted actions of all relevant parties;

(d) ensuring consistency, scientific soundness and clarity of key messages about preventing and reducing harmful use of alcohol;

(e) promoting intercountry networking and exchange of experiences;

(f) facilitating international networking in order to tackle specific and similar problems (for example, specific problems among indigenous or other minority groups or changing youth drinking cultures);

(g) advocating appropriate consideration by parties in international, regional and bilateral trade negotiations to the need and the ability of national and subnational governments to regulate alcohol distribution, sales and marketing, and thus to manage alcohol-related health and social costs;

(h) ensuring that the WHO Secretariat has processes in place to work with nongovernmental organizations and other civil society groups, taking into consideration any conflicts of interest that some nongovernmental organizations may have;

(i) continuing its dialogue with the private sector on how they best can contribute to the reduction of alcohol-related harm. Appropriate consideration will be given to the commercial interests involved and their possible conflict with public health objectives.

Technical support and capacity building

49. Many Member States need increased capacity and capability to create, enforce and sustain the necessary policy and legal frames and implementation mechanisms. Global action will support national action through the development of sustainable mechanisms and the provision of the necessary
normative guidance and technical tools for effective technical support and capacity building, with particular focus on developing and low- and middle-income countries. Such actions must be in accordance with the national contexts, needs and priorities. Development of the necessary infrastructure for effective policy responses in countries with higher or increasing alcohol-attributable burden is an important prerequisite for attaining broader public health and developmental objectives.

50. WHO is committed to cooperate with other relevant actors at regional and global levels in order to provide technical guidance and support for strengthening institutional capacity to respond to public health problems caused by harmful use of alcohol. WHO will especially focus on support and building capacity in developing and low- and middle-income countries.

51. The Secretariat will provide support to Member States by:

(a) documenting and disseminating good models of health-service responses to alcohol-related problems;

(b) documenting and disseminating best practices and models of responses to alcohol-related problems in different sectors;

(c) drawing on expertise in other areas like road safety, taxation and justice with public health expertise in order to design effective models to prevent and reduce alcohol-related harm;

(d) providing normative guidance on effective and cost-effective prevention and treatment interventions in different settings;

(e) developing and strengthening global, regional and intercountry networks in order to help in sharing best practices and facilitating capacity building;

(f) responding to Member States’ requests for support of their efforts to build the capacity to understand the implications of international trade and trade agreements for health.

Production and dissemination of knowledge

52. Important areas for global action will be monitoring trends in alcohol consumption, alcohol-attributable harm and the societal responses, analysing this information and facilitating timely dissemination. Available knowledge on the magnitude of harmful use of alcohol, and effectiveness and cost-effectiveness of preventive and treatment interventions should be further consolidated and expanded systematically at the global level, especially information on epidemiology of alcohol use and alcohol-related harm, impact of harmful use of alcohol on economic and social development and the spread of infectious diseases in developing and low- and middle-income countries.

53. The Global Information System on Alcohol and Health and its regional components were developed by WHO for dynamic presentation of the data on levels and patterns of alcohol consumption, alcohol-attributable health and social consequences and policy responses at all levels. Improving the global and regional data on alcohol and health requires development of national monitoring systems, regular reporting of data by designated focal points to WHO and strengthening the relevant surveillance activities.

54. WHO is committed to working with the relevant partners to shape the international research agenda on alcohol and health, build capacity for research and promote and support international research networks and projects to generate and disseminate data to inform policy and programme development.
55. The Secretariat will provide support to Member States by:

(a) providing an international clearinghouse for information on effective and cost-effective interventions to reduce harmful use of alcohol including promoting and facilitating exchange of information about effective treatment services;

(b) strengthening the Global Information System on Alcohol and Health and the comparative risk assessment of the alcohol-attributable disease burden;

(c) developing or refining appropriate data-collection mechanisms, based on comparable data and agreed indicators and definitions, in order to facilitate data collection, collation, analysis and dissemination at the global, regional and national levels;

(d) facilitating regional and global networks to support and complement national efforts, with a focus on knowledge production and information exchange;

(e) continuing its collaboration with international networks of scientists and health experts to promote research on various aspects of harmful use of alcohol;

(f) facilitating comparative effectiveness studies of different policy measures implemented in different cultural and developmental contexts;

(g) facilitating operational research to expand effective interventions and research on the relationship between harmful use of alcohol and social and health inequities.

Resource mobilization

56. The magnitude of alcohol-attributable disease and social burden is in sharp contradiction with the resources available at all levels to reduce harmful use of alcohol. Global development initiatives must take into account that developing and low- and middle-income countries need technical support – through aid and expertise – to establish and strengthen national policies and plans for the prevention of harmful use of alcohol and develop appropriate infrastructures, including those in health-care systems. Development agencies could consider reducing harmful use of alcohol as a priority area in developing and low- and middle-income countries with a high burden of disease attributable to harmful use of alcohol. Official development assistance provides opportunities to build sustainable institutional capacity in this area in developing and low- and middle-income countries, as do mechanisms for collaboration between developing countries. In that regard, Member States are urged to support each other in the implementation of the global strategy through international cooperation and financial assistance including official development assistance for developing countries.

57. WHO is committed to assist countries upon request in resource mobilization and pooling of available resources to support global and national action to reduce harmful use of alcohol in identified priority areas.

58. The Secretariat will provide support to Member States by:

(a) promoting exchange of experience and good practice in financing policies and interventions to reduce harmful use of alcohol;

(b) exploring new or innovative ways and means to secure adequate funding for implementation of the global strategy;
(c) collaborating with international partners, intergovernmental partners and donors to mobilize necessary resources to support developing and low- and middle-income countries in their efforts to reduce harmful use of alcohol.

IMPLEMENTING THE STRATEGY

59. Successful implementation of the strategy will require concerted action by Member States, effective global governance and appropriate engagement of all relevant stakeholders. All actions listed in the strategy are proposed to support the achievement of the five objectives.

60. The Secretariat will report regularly on the global burden of alcohol-related harm, make evidence-based recommendations, and advocate action at all levels to prevent and reduce harmful use of alcohol. It will collaborate with other intergovernmental organizations and, as appropriate, other international bodies representing key stakeholders to ensure that action to reduce harmful use of alcohol receives appropriate priority and resources.

Links and interfaces with other strategies, plans and programmes

61. This global strategy builds upon regional initiatives such as the Framework for alcohol policy in the WHO European Region (resolution EUR/RC55/R1), the Regional strategy to reduce alcohol-related harm in the Western Pacific Region (resolution WPR/RC57.R5), Alcohol consumption control – policy options in the South-East Asia Region (resolution SEA/RC59/R8), Public health problems of alcohol consumption in the Eastern Mediterranean Region (resolution EM/RC53/R.5) and Actions to reduce the harmful use of alcohol in the African Region (document AFR/RC58/3).

62. Harmful use of alcohol is one of the four main risk factors highlighted in the action plan for the global strategy for the prevention and control of noncommunicable diseases (resolution WHA61.14). The strategy to reduce harmful use of alcohol builds on and links to the other risk factors for noncommunicable diseases and the disease-specific programmes, especially through the global strategy on diet, physical activity and health (resolution WHA57.17), tobacco control (resolution WHA56.1), health promotion and healthy lifestyle (resolution WHA57.16) and cancer prevention and control (resolution WHA58.22).

63. The strategy also links and aligns itself with other related activities in WHO, especially the Mental Health Gap Action Programme, including suicide prevention and management of other substance use disorders as well as programmatic activities on violence and health (resolution WHA56.24), road safety and health (resolution WHA57.10), child and adolescent health and development (resolution WHA56.21) and reproductive health (resolution WHA57.12).

64. With emerging evidence, greater attention is being given to the links between harmful use of alcohol and some infectious diseases and between harmful drinking and development. The strategy also links in with WHO’s existing programmes on HIV/AIDS and tuberculosis and its work on reducing health inequities through action on the social determinants of health (resolution WHA62.14) and achieving the health-related development goals including those contained in the United Nations Millennium Declaration (resolution WHA58.30).

65. The implementation of a global strategy to reduce harmful use of alcohol provides a supportive framework for the WHO regional offices to formulate, revisit and implement region-specific policies and, together with the country offices, provide technical support to Member States. Emphasis will also be put on coordination within the Secretariat so that all actions relevant to harmful use of alcohol are in line with this strategy.
Monitoring progress and reporting mechanisms

66. For monitoring progress, the strategy requires appropriate mechanisms at different levels for assessment, reporting and re-programming. A framework with an impact-focused perspective is needed for assessing achievement of the strategy’s objectives.

67. WHO’s Global Survey on Alcohol and Health and the Global Information System on Alcohol and Health will be important parts of the reporting and monitoring mechanisms. The data-collecting tools of the latter will be adjusted to include the relevant reporting on the process and outcomes of implementation of the strategy at the national level.

68. Regular meetings of global and regional networks of national counterparts offer a mechanism for technical discussion of the implementation of the global strategy at different levels. In addition to taking stock of the process, these meetings could include detailed discussions of priority areas and topics relevant to implementation.

69. Reporting on the implementation of the global strategy to Member States will take place through regular reports to WHO regional committees and the Health Assembly. Information about implementation and progress should also be presented at regional or international forums and appropriate intergovernmental meetings.
ANNEX 5

Nongovernmental organizations admitted into, or maintained in, official relations with WHO by virtue of, respectively, resolution EB126.R17 and decision EB126(5)


African Medical and Research Foundation
Aga Khan Foundation
Association of the Institutes and Schools of Tropical Medicine in Europe
CMC – Churches’ Action for Health
Caritas Internationalis
Collegium Internationale Neuro-Psychopharmacologicum
Consumers International
Council for International Organizations of Medical Sciences
Council on Health Research for Development
CropLife International
EuroSafe – European Association for Injury Prevention and Safety Promotion
FDI World Dental Federation
Framework Convention Alliance on Tobacco Control
Global Forum for Health Research
Global Health Council, Inc.
Helen Keller International
International Agency for the Prevention of Blindness
International Alliance of Patients’ Organizations
International Association for Biologics
International Association for Dental Research
International Association of Biologists Technicians
International Association of Cancer Registries
International Association of Hydatid Disease
International Association of Logopedics and Phoniatrics
International Association of Medical Regulatory Authorities
International Catholic Committee of Nurses and Medico-Social Assistants
International College of Surgeons
International Commission on Radiological Protection
International Committee for Monitoring Assisted Reproductive Technologies
International Conference of Deans of French-Language Faculties of Medicines
International Council for Standardization in Haematology
International Council of Nurses
International Diabetes Federation
International Epidemiological Association
International Federation for Medical and Biological Engineering
International Federation of Biomedical Laboratory Science
International Federation of Clinical Chemistry and Laboratory Medicine

1 Activities concern the period 2007–2009.
2 Activities concern the period 2006–2008.
International Federation of Fertility Societies
International Federation of Health Records Organizations
International Federation of Hospital Engineering
International Federation of Medical Students’ Associations
International Federation of Oto-Rhino-Laryngological Societies
International Federation of Pharmaceutical Manufacturers and Associations
International Federation of Surgical Colleges
International Hospital Federation
International Insulin Federation
International League Against Epilepsy
International Leprosy Association
International Life Saving Federation
International Medical Informatics Association
International Medical Parliamentarians Organization
International Network for Cancer Treatment and Research
International Network on Children’s Health Environment and Safety
International Organization against Trachoma
International Organization for Standardization
International Pharmaceutical Federation
International Physicians for the Prevention of Nuclear War
International Society for Burn Injuries
International Society for Telemedicine & eHealth
International Society of Blood Transfusion
International Society of Orthopaedic Surgery and Traumatology
International Society of Radiology
International Society on Thrombosis and Haemostasis
International Solid Waste Association
International Union against Tuberculosis and Lung Disease
International Union for Conservation of Nature and Natural Resources
International Union of Architects
International Union of Basic and Clinical Pharmacology
International Union of Immunological Societies
International Union of Microbiological Societies
International Union of Psychological Science
International Union of Toxicology
Medicus Mundi International – International Organisation for Cooperation in Health Care
OXFAM
Project ORBIS International, Inc. (ORBIS International)
Stichting Global Network of People Living with HIV/AIDS (GNP+)
The Commonwealth Pharmacists Association
The International League of Dermatological Societies
The International Pharmaceutical Students’ Federation
The International Society for Quality in Health Care Incorporated
The International Society of Radiographers and Radiological Technologists

1 Activities concern the period 2007–2009.
2 Activities concern the period 2006–2008.
3 Activities concern the period 2005–2007.
4 Previously known as the Commonwealth Pharmaceutical Association.
The Network: Towards Unity For Health ¹
The Save the Children Fund ¹
The Transplantation Society ¹
The World Federation of Acupuncture-Moxibustion Societies ¹
The World Medical Association, Inc. ¹
World Association of Societies of Pathology and Laboratory Medicine ¹
World Blind Union ²
World Federation for Medical Education ¹
World Federation of Chiropractic ¹
World Federation of Hydrotherapy and Climatotherapy ²
World Federation of Public Health Associations ¹
World Federation of Societies of Anaesthesiologists ¹
World Federation of Ultrasound in Medicine and Biology ¹
World Organization of Family Doctors ¹
World Self-Medication Industry ¹
World Vision International ¹

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¹ Activities concern the period 2007–2009.
² Activities concern the period 2006–2008.
## ANNEX 6

**Guidelines for the WHO review of psychoactive substances for international control**


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### Appendix 1
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1 See document EB126/2010/REC/2, summary record of the twelfth meeting, for approval of these guidelines.
I. MANDATE

1. The World Health Organization (WHO) is the specialized agency of the United Nations that conducts the medical, scientific and public health evaluation of psychoactive substances under the Single Convention on Narcotic Drugs, 1961 (the 1961 Convention), as amended by the 1972 Protocol, and the Convention on Psychotropic Substances, 1971 (the 1971 Convention). The guidance document for this evaluation has been developed pursuant to resolutions of the World Health Assembly and of the United Nations Commission on Narcotic Drugs (CND). This document amends the previous version of these guidelines and sets out guidelines establishing the underlying principles of the review procedure, working arrangements within the Secretariat and with external bodies, and the nature of the documentation to be prepared. The guidelines cover WHO’s responsibilities under Article 3 of the 1961 Convention and Article 2 of the 1971 Convention concerning whether or not to recommend international control of substances, as well as the assessment of exempted preparations under Article 3 of the 1971 Convention. Common terms and abbreviations are listed in Section VII.

2. The Thirty-third World Health Assembly, by resolution WHA33.27 (1980), requested the Director-General “to promote the initiation and strengthening of national and international programmes for the assessment, scheduling, control and appropriate use of narcotic and psychotropic substances including those of plant origin, and to support such programmes by the development of appropriate guidelines”, and further “to strengthen the coordination between the WHO programmes relating to narcotic and psychotropic substances, those dealing with drug policy and management, and other related programmes, and to strengthen collaboration with interested nongovernmental organizations”.

3. In the light of experience gained over later years, and following the guidance of the Executive Board, WHO first developed the guidelines document for the evaluation and assessment of narcotic and psychotropic substances for decisions on international control in consultation with CND in 1986, which document was revised in 1990. Amendments and decisions subsequently adopted by the Executive Board in 1994 and 1999 resulted in a further revision in 2000. Subsequently, at the request of the Expert Committee on Drug Dependence (Expert Committee), supplementary guidelines were submitted to the Executive Board in order to clarify certain issues. The Board considered the proposed supplementary guidelines in May 2004 and January 2005 when it requested the Secretariat and the Expert Committee to continue their work on the issue. This revision of the guidelines has been prepared in response to that request.

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1 See Appendix 1 for the most relevant excerpts from these conventions.
2 Resolution WHA33.27.
3 Resolution EB73.R11.
4 Decision EB77(3).
5 Decision EB85(10).
6 Decision EB93(16).
7 Decision EB103(5).
8 Documents EB114/7 and EB115/12.
9 Document EB114/2004/REC/1, summary record of the third meeting.
10 Document EB115/2005/REC/2, summary record of the sixth meeting.
II. UNDERLYING PRINCIPLES

4. The Preamble of the Single Convention on Narcotic Drugs, 1961 provides:

“The Parties,
Concerned with the health and welfare of mankind,
Recognizing that the medical use of narcotic drugs continues to be indispensable for the relief of pain and suffering and that adequate provision must be made to ensure the availability of narcotic drugs for such purposes,
Recognizing that addiction to narcotic drugs constitutes a serious evil for the individual and is fraught with social and economic danger to mankind,
Conscious of their duty to prevent and combat this evil,
Considering that effective measures against abuse of narcotic drugs require coordinated and universal action,
Understanding that such universal action calls for international cooperation guided by the same principles and aimed at common objectives,
Acknowledging the competence of the United Nations in the field of narcotics control and desirous that the international organs concerned should be within the framework of that Organization,
Desiring to conclude a generally acceptable international convention replacing existing treaties on narcotic drugs, limiting such drugs to medical and scientific use, and providing for continuous international cooperation and control for the achievement of such aims and objectives ...”

The Preamble of the Convention on Psychotropic Substances, 1971 provides:

“The Parties,
Being concerned with the health and welfare of mankind,
Noting with concern the public health and social problems resulting from the abuse of certain psychotropic substances,
Determined to prevent and combat abuse of such substances and the illicit traffic to which it gives rise,
Considering that rigorous measures are necessary to restrict the use of such substances to legitimate purposes,
Recognizing that the use of psychotropic substances for medical and scientific purposes is indispensable and that their availability for such purposes should not be unduly restricted,
Believing that effective measures against abuse of such substances require coordination and universal action,
Acknowledging the competence of the United Nations in the field of control of psychotropic substances and desirous that the international organs concerned should be within the framework of that Organization,
Recognizing that an international convention is necessary to achieve these purposes ...”

The WHO review procedure, grounded in considerations of public health and with an evidence-based approach, will utilize the best available relevant information. Consistent with the requirements of the 1961 and 1971 Conventions, WHO will develop scheduling recommendations guided by the provisions in the Conventions regarding the changes in the scope of control of substances and also taking into account the preambles of the Conventions, the need to reduce the risk to public health, including the risk of abuse and ensuring medical availability, and the relevant resolutions of its governing bodies. The Conventions are legal instruments; the WHO review procedure shall be applied in a manner consistent with the letter and the spirit of the Conventions.
III. PROVISIONS OF THE CONVENTIONS

5. The 1961 and 1971 Conventions entrust WHO with the responsibility of reviewing and assessing substances to determine whether they should be controlled under the Conventions. A request for such a review can be initiated by a notification to the Secretary-General of the United Nations by a Party to the Conventions, or by WHO itself. WHO will forward the results of this review to CND which has the responsibility to decide whether to schedule substances under the provisions of the Conventions.

6. The basis for the scheduling recommendation made by WHO is an evaluation of whether specific criteria set forth in the Conventions have been met. Under the provisions of the 1961 Convention, the CND must accept or refuse the WHO recommendation as a whole, except that it may decide to place a substance only in Schedule I and not in Schedule IV if WHO has recommended simultaneous inclusion in both schedules. The CND should in principle accept the medical, scientific, chemical and pharmacological findings of WHO, and when the CND rejects a recommendation, it should be guided by other considerations such as those of an administrative or social nature. In the case of the 1971 Convention, the CND may accept a WHO proposal, but it may also decide to place a substance in a schedule other than that recommended by WHO. With respect to control under the 1971 Convention, WHO’s assessment is determinative for scientific and medical matters, but CND may also take into account legal, administrative, economic, social and other factors in reaching its decision.

7. Under the provisions of Article 3 of the 1971 Convention, a Party may exempt from specific control measures a preparation containing one or more psychotropic substances if the preparation is compounded in such a way that it presents no, or a negligible, risk of abuse, and the substance cannot be recovered by a readily applicable means in a quantity liable to abuse, so that the preparation does not give rise to a public health and social problem. A party shall notify the Secretary-General of the United Nations who in turn shall transmit the notification to other Parties, to WHO and to the International Narcotics Control Board. If a Party or WHO has information which it believes requires that the exemption of a preparation should be terminated, it should notify the Secretary-General of the United Nations accordingly and submit information in support of that notification. WHO reviews the data submitted by the Parties that wish to avail themselves of this provision for exemption under the 1971 Convention by applying specific guidelines that have been approved by CND.

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1 The Director-General represents WHO for the purpose of receiving notifications under the international drug control conventions and of making recommendations concerning the international control of psychoactive substances under those conventions on the basis of recommendations and advice provided to him or her as described in these guidelines.

2 The scheduling process is defined in Article 3 of the 1961 Convention and Articles 2 and 17 (para. 2) of the 1971 Convention. The scheduling process is described in detail in the commentaries on these Conventions, published by the United Nations.


4 1971 Convention, Art. 2, para. 5; See also, Commentary on the Convention of Psychotropic Substances (1971 Convention), para. 20 (p. 71).

5 The specific WHO procedure for review of exempted preparations was developed in accordance with the Commission’s guidelines for exemption. These guidelines, which were largely based on recommendations made by WHO, were approved by CND at its Eighth special session and are set forth in its resolution 1 (S-VIII). See the report of the Commission in Economic and Social Council, Official Records, 1984, Supplement No. 3 (Document E/CS.7/1984/13).
8. Under the provisions of the 1961 Convention, preparations of narcotic drugs exempted from specific control measures are listed in Schedule II. New exemptions can be made only by including a preparation in Schedule III, and relevant proposals are reviewed by WHO in the same way as those for single substances.

9. The United Nations Convention Against Illicit Traffic in Narcotic Drugs and Psychotropic Substances, 1988 (the 1988 Convention), entered into force in November 1990. Article 12 of the 1988 Convention places under international control substances frequently used in the illicit manufacture of narcotic drugs or psychotropic substances. These substances are listed in Table I and Table II of the 1988 Convention. WHO has no formal role to play in the scheduling of such substances under the 1988 Convention. However, it is possible that the same substance may be considered for control simultaneously under the 1961 Convention, the 1971 Convention, or the 1988 Convention. Guidance on how to address such a situation is provided below under the subsection Assessment for scheduling by the Expert Committee.

IV. WHO REVIEW PROCEDURE

10. The purpose of the WHO review procedure is to evaluate substances for international control. Using data provided by the Secretariat, the Expert Committee conducts pre-reviews and critical reviews in order to provide scheduling advice to the Director-General.

11. The review of exempted preparations notified by a Party involves a preliminary review by the Secretariat and an evaluation by the Expert Committee.

12. The time schedule for the review procedure should be set by the Secretariat bearing in mind the calendar of CND and its procedural requirements.

Information collection

13. The Secretariat should routinely collect relevant data related to psychoactive substances that are being abused or might have abuse potential and substances convertible into such substances from the literature, WHO programmes, WHO collaborating centres, national health and drug control authorities, intergovernmental and nongovernmental organizations, research and academic institutions and other competent sources.

Pre-review

14. The purpose of the pre-review is to determine whether current information justifies an Expert Committee critical review.

15. A pre-review is initiated when a proposal has been submitted to the Expert Committee with supporting information either by (1) the Secretariat, (2) any member of the Expert Committee, or (3) representatives of other organizations invited to participate in the Expert Committee meeting in accordance with paragraph 35. The Secretariat will put the proposed pre-review of a substance on the agenda of the first possible Expert Committee meeting.

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1 See Appendix 3 for a flow chart of the evaluation procedure.
16. The categories of information for evaluating substances in pre-reviews are identical to those used in critical reviews. The Secretariat shall supply the supporting information required for pre-review in the form of a brief summary of relevant information. At this stage the Expert Committee must decide whether the information warrants a critical review. If the Expert Committee determines that a critical review is not warranted then the Expert Committee should recommend no further evaluation of the substance. The pre-review is a preliminary analysis and findings at this stage should not determine whether the control status of a substance should be changed. The confidentiality of information received by WHO for use in the review will be respected if so requested by the provider. Appropriate arrangements to sustain confidentiality will be made when the Expert Committee has access to the information used to prepare the pre-review.

17. The Expert Committee shall recommend a critical review if it finds that information may justify the scheduling or a change in the scheduling of the substance in the 1961 or 1971 Conventions, using the criteria in paragraphs 48 to 59.

**Critical review**

18. The purpose of the critical review is to consider whether the Expert Committee should advise the Director-General to recommend the scheduling of, or amending of the scheduling status of, a substance.

19. A critical review is initiated when:

   (1) there has been notification from a Party to the 1961 or the 1971 Convention concerning the scheduling of a substance;

   (2) there has been an explicit request from CND to review a substance;

   (3) pre-review of a substance has resulted in an Expert Committee recommendation for critical review; or

   (4) information is brought to WHO’s attention that a substance is clandestinely manufactured, of especially serious risk to public health and society, and of no recognized therapeutic use by any Party.

In respect of case (4), if therapeutic use of the substance is confirmed subsequently by any Party the substance shall be subjected to a pre-review.

**Critical review documents**

20. The purpose of the critical review document is to provide comprehensive data for use by the Expert Committee in assessing individual substances.

21. The Secretariat is responsible to collect and assemble the data on substances selected for critical review. The Secretariat will use a questionnaire to request information from ministers of health in the Member States and international drug control bodies and will circulate the agenda of the next meeting.

22. The critical review document should be as thorough as possible, and balanced in its presentation. It should include the adequate and relevant data, including medical literature and abuse studies. In order to accomplish this, the Secretariat may seek assistance from advisers and ad hoc working groups.
23. When preparing the draft critical review document, including a separate report on the questionnaire, the Secretariat should include, where feasible, information under the following headings:

(1) substance identification by International Nonproprietary Name (INN), chemical or other common name and trade names, other identifying characteristics, Chemical Abstracts Service (CAS) registry number;
(2) chemistry, including general information on synthesis, preparation and properties;
(3) ease of convertibility into controlled substances;
(4) general pharmacology, including pharmacokinetics and pharmacodynamics;
(5) toxicology,
(6) adverse reactions in humans;
(7) dependence potential;
(8) abuse potential;
(9) therapeutic applications, extent of therapeutic use and epidemiology of medical use;
(10) listing on the WHO Model List of Essential Medicines;
(11) marketing authorizations (as a medicine);
(12) industrial use;
(13) non-medical use, abuse and dependence;
(14) nature and magnitude of public health problems related to abuse and dependence;
(15) licit production, consumption and international trade;
(16) illicit manufacture and traffic, and related information;
(17) current international controls and their impact;
(18) current and past national controls;
(19) other medical and scientific matters relevant for a recommendation on the scheduling of the substance.

24. The data in the critical review should be presented in a manner that will facilitate an evidence-based assessment by the Expert Committee. The critical review will comprise a summary and a section that compares the data directly against the scheduling criteria.

25. The draft critical review document and the report on the questionnaire are transmitted to all governments, institutions, organizations, or other interested parties that have directly and substantially collaborated in its preparation and have requested it. The recipients may provide comments on the
draft. To help to ensure that all material submitted to the Expert Committee is up to date, the Secretariat will circulate the agenda of the next meeting to those collaborating information sources.

26. For each substance, the draft critical review document and the report on the questionnaire will be peer-reviewed by two experts from WHO’s Expert Advisory Panels, including an evaluation of the strength of evidence they present. If there are data limitations or omissions, they should be identified, discussed and adapted as needed.

27. The critical review document and the report on the questionnaire will be provided to all members of the Expert Committee at least thirty days before its meeting, and posted on the WHO website, according to WHO rules for publication.

28. The confidentiality of information received by WHO for use in the review will be respected if so requested by the provider. Appropriate arrangements to sustain confidentiality will be made when the Expert Committee has access to the information used to prepare the pre-review and the critical review.

Preliminary review of exempted preparations containing psychotropic substances

29. The Secretariat should review the notification of exemption received from a Party to the 1971 Convention in order to ascertain whether the preparation containing a psychotropic substance is for domestic use only, or is being exported outside the exempting country. Where the preparation is for domestic use only, and if the exempting Party gives assurance in its notification that, to the best of its knowledge, there is no significant abuse, the Secretariat will assume that the exemption does not require an evaluation by the Expert Committee. However, if WHO receives evidence of national abuse, or information that the preparation may constitute a public health and social problem to another Party (e.g. illicit trade and/or abuse), the exemption shall be evaluated by the Expert Committee.

Expert Committee on Drug Dependence

30. In accordance with WHO’s regulations, the Expert Committee meets when necessary to discuss the appropriate issues within its responsibility. As a guide, the Expert Committee should meet at least every second year.

31. **Membership.** The Expert Committee members are chosen by the Director-General in accordance with WHO’s Regulations on Expert Advisory Panels and Committees. The Director-General shall establish the number of experts to be invited to a meeting of an Expert Committee on Drug Dependence, determine its date and duration, and convene the Expert Committee meeting.

32. **Functions.** The functions of the Expert Committee are to review information available to it on substances being considered for international control and for exemptions, and to advise the Director-General on such control. The advice of the Expert Committee concerns scientific, medical and public health findings and must comply with the criteria established in the Conventions. Specific responsibilities of the Expert Committee are:

   (1) **pre-review:** to determine whether a substance should be subject to critical review;

   (2) **critical review:** to assess the dependence-producing capability, the likelihood of abuse and of causing public health and social problems, and usefulness in medical therapy of each substance under review; and to advise on the appropriate schedule under one of the Conventions;
(3) exempted preparations: to evaluate the need to terminate notified exemptions of preparations under the 1971 Convention.

33. Procedure. WHO’s Regulations on Expert Advisory Panels and Committees are applicable.

34. Secretariat. The Expert Committee is assisted by a secretariat, in particular by the Expert Committee’s Secretary and furthermore by staff members from appropriate WHO programmes, consultants and temporary advisers, as required. The functions of the Secretary are executed by a technical officer competent in the subject concerned.

35. Other organizations. Representatives of United Nations organizations such as the United Nations Office on Drugs and Crime (UNODC), the International Narcotics Control Board (INCB), and appropriate nongovernmental organizations (NGOs) in official relations with WHO may be invited to attend the meetings of the Expert Committee as observers. In consultation with the members and the Secretariat, the Chair may decide to have a session of the Expert Committee with the members only.

36. The Expert Committee’s recommendations and advice remain confidential until the clearing of their publication according to WHO’s internal rules. All participants are required to respect the confidentiality of all information received as part of the Expert Committee process as well as the confidentiality of the Expert Committee’s deliberations.

Information meeting

37. Interested parties that intend to make submissions of data may request the convening of an information meeting with the Expert Committee for this purpose. Requests for such a meeting should be submitted to the Secretariat at least twenty days before the start of the Expert Committee meeting. The request should state the nature and content of the presentation to be made at the meeting. All participants to the Expert Committee meeting are invited to this information meeting.

38. The purpose of the information meeting is to afford the Expert Committee the opportunity, before the Committee’s meeting, to receive presentations and to question representatives of interested parties concerning data that have been provided about substances under review.

39. The information meeting will be held before the Expert Committee convenes its meeting. The Secretariat at its discretion shall decide the agenda of the information meeting, taking into account the nature of the proposed presentations and the time constraints for the meeting of the Expert Committee. The decisions of the Secretariat concerning the information meeting will be communicated to requesting interested parties at least 10 days before the Expert Committee meeting.

Experts collaborating in the WHO review

40. Experts collaborating in the review should have a well-documented scientific career at a high level and professional background, and should represent relevant behavioural, pharmacological, pharmaceutical, medical, biological, or epidemiological disciplines, as well as public health administration. Scientists representing industry research may be asked to collaborate as advisers in WHO ad hoc working groups, as appropriate, but they are not invited to participate in the Expert Committee meeting.

41. Experts participating in the WHO review should be selected with careful attention given to the avoidance of conflicts of interest. Similar considerations shall apply to all concerned with the process. In this connection, experts invited to participate in the WHO review and, in particular, in the work of the Expert Committee, sign a statement concerning potential conflicts of interest.
Assessment for scheduling by the Expert Committee

42. The Expert Committee bases its deliberations mainly on the documents provided by the Secretariat: these consist of the critical review document, the report on the questionnaire and comments received by the Secretariat concerning the critical review. The Expert Committee may also consider additional information presented in the information meeting. The information on which the critical review is based will be made available to the Expert Committee. The dissemination of this information may otherwise be restricted if needed to protect confidentiality requirements pursuant to paragraph 28.

43. Proposals for the change in control of a substance should be subjected to the same assessment that is given to substances proposed for initial scheduling; the same criteria as mentioned below in paragraphs 46 to 59 should be used in making the assessment.

44. To facilitate efficient administration of the international control system, it is not advisable to place a substance under more than one Convention. This is also true of substances that have been placed in a Table of the 1988 Convention, or have been recommended by INCB for inclusion in a Table.

45. A recommendation to delete a substance from one Convention with a simultaneous recommendation to add the same substance to another Convention may affect administration of the international scheme of regulation. Like all recommendations, consideration of such changes in control may be undertaken in light of new information to justify such a change. Any proposal to move a substance from one convention to another should be made only if specific new control measures are necessary in order to decrease the extent or likelihood of abuse or the use of the substance in illicit drug manufacturing, and will not unduly limit availability for legitimate medical and scientific purposes.

The assessment process

Orientation

46. Both the 1961 and 1971 Conventions provide for control of substances that are liable to “similar abuse and similar ill-effects” as substances already controlled under those Conventions. Many substances exhibit similarity in their “abuse” and “ill-effects” to substances in both the 1961 Convention and the 1971 Convention. Amphetamines, barbiturates and tranquillizers are only subject to the 1971 Convention, by virtue of an understanding of the Parties to the Conventions that the 1961 Convention did not apply to these substances even though the effects of amphetamines, barbiturates and tranquillizers were recognized to be similar to cocaine and morphine in some respects.1 When considering other substances that exhibit abuse characteristics similar to substances regulated under both Conventions, the Expert Committee should follow the sequence for analysis established by the guidelines for all substances; that is, first consider applicability of the 1961 Convention and, if it is found not to apply, then the 1971 Convention. As such, the Committee would first assess whether the substance under review shows similar abuse liability profile (based on animal and human studies) and dependence-producing properties to drugs already controlled under the 1961 Convention. This assessment should not be limited to a narrow consideration of a single pharmacologic property. If the substance under review shows sufficiently similar abuse liability profile and dependence-producing properties to drugs already controlled under the 1961 Convention, then it should be recommended for scheduling under the 1961 Convention; if not, then the analysis should be made using the criteria in the 1971 Convention.

1 Commentary on the 1961 Convention, Art.3, para.3, subpara (iii), Comment 6 (p.87).
47. The 1961 Convention provides for the control of substances convertible to narcotic drugs. The 1971 Convention provides for no such control of precursors. The 1988 Convention fills the void that existed for controlling precursors of psychotropic substances and the control of other chemicals frequently used in the illicit production of all controlled substances. INC has responsibility for reviewing precursors of both narcotic and psychotropic substances for potential control. The Expert Committee might be asked to assess a substance to determine if it is convertible to a substance controlled under the 1961 Convention. If so, the Committee should determine if the substance is “convertible” as defined in paragraph 49 of these Guidelines, and then determine whether it is convertible to a substance controlled by the 1961 Convention.

**Step 1: 1961 Convention**

48. The Expert Committee, when deciding whether to recommend international control, or a change in international control, after completion of its discussions, first decides, with regard to the 1961 Convention, whether the substance in accordance with Article 3, paragraph 3 (iii) of that Convention: (1) is liable to similar abuse and productive of similar ill-effects as the substances in Schedule I or Schedule II; or (2) is convertible into a substance already in Schedule I or Schedule II.

49. In addition to the principle of “Similarity”, laid down in Article 3, paragraph 3 (iii) of that Convention and mentioned in paragraph 48, the Convention also contains the principle of “Convertibility”. A substance is convertible if it is of such a kind as to make it, by the ease of the process and by the yield, practicable and profitable for a clandestine manufacturer to transform the substance in question into controlled drugs.1

50. The Secretariat will promptly advise the INCB Secretariat of all Expert Committee assessments relating to substances that might be convertible into a narcotic drug. If the advice of the Expert Committee is to schedule a substance, whether psychoactive or convertible into a psychoactive substance, that is already in Table I or Table II of the 1988 Convention, the Secretariat will take steps to coordinate its proceedings with the INCB Secretariat. Such steps will enable INC to review the possibility of recommending deletion of the substance from the Table of the 1988 Convention before WHO communicates its recommendation to the United Nations. If both WHO and INC make such recommendations, CND could consider the two proposals simultaneously.

51. If a substance meets the criteria for inclusion in Schedule I of the 1961 Convention, the Expert Committee should further consider whether the drug meets the requirements for inclusion in Schedule IV in accordance with Article 3, paragraph 5 of that Convention, with regard to substances being particularly liable to abuse and to produce ill-effects and if such liability is not offset by substantial therapeutic advantages not possessed by substances other than drugs in Schedule IV.

52. If the Expert Committee finds that a substance does not meet the criteria for control under the 1961 Convention, then it makes an assessment in accordance with the 1971 Convention.

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1 Commentary on the 1961 Convention, para. 13 (p.89).
Step 2: 1971 Convention

53. In considering the scheduling under the 1971 Convention, the Expert Committee determines whether, in accordance with Article 2, paragraph 4:

(a) The substance has the capacity to produce:

(i) (1) A state of dependence, and

(2) Central nervous system stimulation or depression, resulting in hallucinations or disturbances in motor function or thinking or behaviour or perception or mood, or

(ii) Similar abuse and similar ill-effects as a substance in Schedule I, II, III or IV, and

(b) There is sufficient evidence that the substance is being or is likely to be abused so as to constitute a public health and social problem warranting the placing of the substance under international control.

54. In applying paragraph 53 of the Guidelines, the principle of similarity described in Article 2, paragraph 4(a)(ii) of the 1971 Convention applies only in situations when the substance does not produce a state of dependence. In the absence of a finding that a substance produces dependence, similarity takes on importance; otherwise it is secondary.

55. The Commentary on the 1971 Convention provides the following considerations to be taken into account in such an evaluation:

(i) “The assessment of the substance (…) should not only comprise the factual results of [WHO’s] examination (…) but also an evaluation of the data which it may have found in the light of such considerations of public health as it may consider appropriate (…)” Commentary on the 1971 Convention, para. 41 (p. 58);

(ii) “WHO must also establish the extent of abuse or the degree of likelihood of abuse (…) in order to be able to determine whether [this] constitutes a public health and social problem warranting the placing of the substance under international control.” Id., para. 42 (p. 58);

(iii) WHO must “assess the degree of seriousness of the public health and social problem (…). Since in arriving at its decision the [Commission on Narcotics Drugs will] weigh the dangerous properties of the substance against the non-medical considerations … it would find it useful to have the views of WHO on the degree of seriousness of the health and social problem which it has to take into account.” Article 2, paragraph 5 … Id., para. 43 (p. 59);

(iv) WHO is required to include an assessment of “the degree of usefulness of the substance in medical therapy based on two considerations: (a) the degree of risk to public health and

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1 Dependence was defined by the 28th Expert Committee on Drug Dependence as: “A cluster of physiological, behavioural and cognitive phenomena of variable intensity, in which the use of a psychoactive drug (or drugs) takes on a high priority. The necessary descriptive characteristics are preoccupation with a desire to obtain and take the drug and persistent drug-seeking behaviour. Determinants and the problematic consequences of drug dependence may be biological, psychological or social, and usually interact.” The Committee also mentioned that in its opinion this definition is compatible with the ICD-10 diagnostic guidelines. (WHO Expert Committee on Drug Dependence. Twenty-eighth report. Geneva, World Health Organization, 1993 (WHO Technical Report Series, No. 836)).
(b) the usefulness of the drug in medical therapy… [which means] not only its potential beneficial effects, its value in the case of grave medical indications and the extent and frequency of its employment, but also the intensity of its dangerous properties (…) and other harmful side effects may have to be taken into account.” Id., para. 44 (pp. 59–60);

(v) It is safe to state that WHO in recommending a particular Schedule for a substance, “will be guided by its views of the degree of risk to public health which the substance presents and its usefulness in medical therapy.” Id., para. 49 (p. 61).

56. On the basis of the above considerations, more specific criteria for proposing to include a substance for control in a particular schedule were developed by the Expert Committee at its seventeenth meeting.¹ They are as follows:

For inclusion in Schedule I:

Substances whose liability to abuse constitutes an especially serious risk to public health and which have very limited, if any, therapeutic usefulness.

For inclusion in Schedule II:

Substances whose liability to abuse constitutes a substantial risk to public health and which have little to moderate therapeutic usefulness.

For inclusion in Schedule III:

Substances whose liability to abuse constitutes a substantial risk to public health and which have moderate to great therapeutic usefulness.

For inclusion in Schedule IV:

Substances whose liability to abuse constitutes a smaller but still significant risk to public health and which have a therapeutic usefulness from little to great.

In cases where the above criteria apply only in part, the scheduling recommendation should be made with a higher regard to the risk to that dimension of public health specific to abuse liability.

Notwithstanding the above, recommendations for inclusion in Schedule I should be made only when the above criteria are fully met, with respect to both therapeutic usefulness and the risk to public health.

57. The criteria given in the foregoing paragraph do not specifically address the dimension of social problems, although the Commentary on the 1971 Convention does. It is also noted that the above criteria do not cover all cases. The “risk to public health” in the above criteria should be interpreted to mean both social and public health problems. Note that under Article 2, paragraph 4(b) there must be a finding of an “international” need for control, meaning that controls of the Convention are suitable to solve or alleviate the problem and that lack of those controls in one country, no matter whether it has itself the public health and social problem caused by the substance under examination, weakens the control in other countries which have such a problem. International control is also warranted if the

public health and social problem exists only in a single country if the efforts of control by that country are weakened by the lack of control in other countries.

58. If the advice of the Expert Committee is to include a substance that is already in Table I or Table II of the 1988 Convention, the WHO Secretariat will take steps to coordinate its proceedings with the INCB Secretariat. Such steps will enable INCB to review the possibility of recommending deletion of the substance from the Table of the 1988 Convention before WHO communicates its recommendation to the United Nations. If both WHO and INCB make such recommendations, CND could consider the two proposals simultaneously.

59. The Expert Committee shall provide its recommendation on the scheduling status on all drugs or substances under review as described in paragraph 60. Should the Expert Committee be unable to make a recommendation concerning substances under review, then it should request another critical review in order to refer the matter to a subsequent Expert Committee.¹

**Step 3: The report**

60. The Expert Committee prepares a summary assessment of each substance reviewed. This assessment should include the Expert Committee’s findings regarding pharmacological similarity, similar abuse, and similar ill-effects of the substance to substances in Schedules I and II of the 1961 Convention and, in the case of a “convertible” substance, an assessment of the convertibility of the substance into a substance already controlled as a narcotic drug. If the substance is recommended for control under the 1971 Convention, the assessment should also indicate whether the substance is being recommended for such control as a dependence-producing substance or on the basis of similarity. For all substances reviewed, the summary assessment should give a description of the Expert Committee’s findings on the extent or likelihood of abuse, the degree of seriousness of the public health and social problem, and the degree of usefulness of the substance in medical therapy, together with the advice on the control measures, if any, that would be appropriate in the light of its assessment. The Expert Committee will advise the Director-General on its assessment and recommendation. The Expert Committee’s report will be published and made available on the WHO web site in conformity with WHO rules for publication of Expert Committee reports.

**Assessment of exempted preparations by the Expert Committee**

61. The assessment of exempted preparations by the Expert Committee should evaluate the following elements: (1) conformity with the requirements of Article 3, paragraph 2, of the 1971 Convention concerning abuse liability and recoverability of the psychotropic substances as well as with CND resolution 1 (S-VIII); and (2) the evidence available to WHO that the preparation may constitute a public health and social problem to an importing country or to a country where it is illicitly traded. On conclusion of the assessment, the Expert Committee advises the Director-General accordingly.²

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² See also Appendix 2.
V. COMMUNICATION OF WHO RECOMMENDATIONS

62. After receiving the advice of the Expert Committee to schedule or to amend the scheduling status of a substance, the Director-General will, as appropriate, communicate the recommendation on behalf of WHO to the United Nations. Copies of the recommendation are made available on the WHO web site concurrently.

63. Any recommendation to terminate an exemption in whole or in part will be communicated by the Director-General to the exempting Party if the abuse problem is limited to the country of origin of the preparation, or to the United Nations if the problems are widespread.

VI. PUBLICATION OF DOCUMENTS RELATED TO THE WHO REVIEW

64. The Director-General will submit to the Executive Board a report of the meetings of the ECDD in accordance with paragraph 4.23 of the Regulations on Expert Advisory Panels and Committees, and the report of the Expert Committee is published according to the WHO rules, both in the WHO Technical Report Series and on the WHO web site. The publication of any other document prepared for the Expert Committee is subject to Rule 4.15 of the Regulations for Expert Advisory Panels and Committees, which states that the Director-General may publish or authorize the publication of any document prepared for an expert committee, with due recognition of authorship if applicable.

VII. ABBREVIATIONS AND DEFINITIONS

CND The Commission on Narcotic Drugs of the Economic and Social Council of the United Nations.

Expert Committee In this document, “the Expert Committee” refers to the WHO Expert Committee on Drug Dependence. The First World Health Assembly decided in 1948, by resolution WHA1.25, to establish the Expert Committee on Habit-Forming Drugs which, since its sixteenth meeting (1968), is named Expert Committee on Drug Dependence.


Member State A State which is a Member of WHO.


Notification A formal communication addressed to the Secretary-General of the United Nations by a Party to an international drug control convention or by WHO, or by the Secretary-General of the United Nations to a Party to an international drug control convention or to WHO. In the context of the present guidelines, reference to a notification means a notification relating to the scheduling of a substance under the
provisions of either Article 3 of the Single Convention or Articles 2 and 3 of the Convention on Psychotropic Substances.

**Party**

A State which has become a Party to an international drug control convention, through signature, ratification, accession, or succession.

**Psychoactive substance**

Any substance, natural or synthetic, or any natural substance material, which has psychoactive properties.

**Psychotropic substance**

Any substance, natural or synthetic, or any natural material in Schedule I, II, III or IV of the Convention on Psychotropic Substances, 1971.

**Secretariat**

The Secretariat of WHO.

**The 1961 Convention**


**The 1971 Convention**


**The 1988 Convention**


**UNODC**

United Nations Office on Drugs and Crime.
APPENDIX 1

EXCERPTS FROM THE UNITED NATIONS DRUG CONTROL CONVENTIONS

Single Convention on Narcotic Drugs, 1961, as amended by the 1972 Protocol (Extract)¹

Article 3

CHANGES IN THE SCOPE OF CONTROL

1. Where a Party or the World Health Organization has information which in its opinion may require an amendment to any of the Schedules, it shall notify the Secretary-General and furnish him with the information in support of the notification.

2. The Secretary-General shall transmit such notification, and any information which he considers relevant, to the Parties, to the Commission, and, where the notification is made by a Party, to the World Health Organization.

3. Where a notification relates to a substance not already in Schedule I or in Schedule II,

   i) The Parties shall examine in the light of the available information the possibility of the provisional application to the substance of all measures of control applicable to drugs in Schedule I;

   ii) Pending its decision as provided in subparagraph iii) of this paragraph, the Commission may decide that the Parties apply provisionally to that substance all measures of control applicable to drugs in Schedule I. The Parties shall apply such measures provisionally to the substance in question;

   iii) If the World Health Organization finds that the substance is liable to similar abuse and productive of similar ill-effects as the drugs in Schedule I or Schedule II or is convertible into a drug, it shall communicate that finding to the Commission which may, in accordance with the recommendation of the World Health Organization, decide that the substance shall be added to Schedule I or Schedule II.

4. If the World Health Organization finds that a preparation because of the substances which it contains is not liable to abuse and cannot produce ill-effects (paragraph 3) and that the drug therein is not readily recoverable, the Commission may, in accordance with the recommendation of the World Health Organization, add that preparation to Schedule III.

5. If the World Health Organization finds that a drug in Schedule I is particularly liable to abuse and to produce ill-effects (paragraph 3) and that such liability is not offset by substantial therapeutic advantages not possessed by substances other than drugs in Schedule IV, the Commission may, in accordance with the recommendation of the World Health Organization, place that drug in Schedule IV.

6. Where a notification relates to a drug already in Schedule I or Schedule II or to a preparation in Schedule III, the Commission, apart from the measure provided for in paragraph 5, may, in accordance with the recommendation of the World Health Organization, amend any of the Schedules by:

   a) Transferring a drug from Schedule I to Schedule II or from Schedule II to Schedule I;

   or

   b) Deleting a drug or a preparation as the case may be, from a Schedule.

7. Any decision of the Commission taken pursuant to this article shall be communicated by the Secretary-General to all States Members of the United Nations, to non-member States Parties to this Convention, to the World Health Organization and to the Board. Such decision shall become effective with respect to each Party on the date of its receipt of such communication, and the Parties shall thereupon take such action as may be required under this Convention.

8. a) The decisions of the Commission amending any of the Schedules shall be subject to review by the Council upon the request of any Party filed within ninety days from receipt of notification of the decision. The request for review shall be sent to the Secretary-General together with all relevant information upon which the request for review is based;

   b) The Secretary-General shall transmit copies of the request for review and relevant information to the Commission, the World Health Organization and to all the Parties inviting them to submit comments within ninety days. All comments received shall be submitted to the Council for consideration;

   c) The Council may confirm, alter or reverse the decision of the Commission, and the decision of the Council shall be final. Notification of the Council’s decision shall be transmitted to all States Members of the United Nations, to non-member States Parties to this Convention, to the Commission, to the World Health Organization, and to the Board;

   d) During pendency of the review the original decision of the Commission shall remain in effect.

9. Decisions of the Commission taken in accordance with this article shall not be subject to the review procedure provided for in article 7.
Convention on Psychotropic Substances, 1971 (Extracts)\(^1\)

**Article 2**

SCOPE OF CONTROL OF SUBSTANCES

... 

4. If the World Health Organization finds:

   a) That the substance has the capacity to produce

      i) 1) A state of dependence, and

      2) Central nervous system stimulation or depression, resulting in hallucinations or disturbances in motor function or thinking or behaviour or perception or mood, or

      ii) Similar abuse and similar ill-effects as a substance in Schedule I, II, III or IV, and

   b) That there is sufficient evidence that the substance is being or is likely to be abused so as to constitute a public health and social problem warranting the placing of the substance under international control, the World Health Organization shall communicate to the Commission an assessment of the substance, including the extent or likelihood of abuse, the degree of seriousness of the public health and social problem and the degree of usefulness of the substance in medical therapy, together with recommendations on control measures, if any, that would be appropriate in the light of its assessment.

5. The Commission, taking into account the communication from the World Health Organization, whose assessments shall be determinative as to medical and scientific matters, and bearing in mind the economic, social, legal, administrative and other factors it may consider relevant, may add the substance to Schedule I, II, III or IV. The Commission may seek further information from the World Health Organization or from other appropriate sources.

... 

**Article 3**

SPECIAL PROVISIONS REGARDING THE CONTROL OF PREPARATIONS

1. Except as provided in the following paragraphs of this article, a preparation is subject to the same measures of control as the psychotropic substance which it contains, and, if it contains more than one such substance, to the measures applicable to the most strictly controlled of those substances.

2. If a preparation containing a psychotropic substance other than a substance in Schedule I is compounded in such a way that it presents no, or a negligible, risk of abuse and the substance cannot be recovered by readily applicable means in a quantity liable to abuse, so that the preparation does not give rise to a public health and social problem, the preparation may be exempted from certain of the measures of control provided in this Convention in accordance with paragraph 3.

3. If a Party makes a finding under the preceding paragraph regarding a preparation, it may decide to exempt the preparation, in its country or in one of its regions, from any or all of the measures of control provided in this Convention except the requirements of:

   a) article 8 (licences), as it applies to manufacture;
   b) article 11 (records), as it applies to exempt preparations;
   c) article 13 (prohibition of and restrictions on export and import);
   d) article 15 (inspection), as it applies to manufacture;
   e) article 16 (reports to be furnished by the Parties), as it applies to exempt preparations; and
   f) article 22 (penal provisions), to the extent necessary for the repression of acts contrary to laws or regulations adopted pursuant to the foregoing obligations.

A Party shall notify the Secretary-General of any such decision, of the name and composition of the exempt preparation, and of the measures of control from which it is exempted. The Secretary-General shall transmit the notification to the other Parties, to the World Health Organization and to the Board.

4. If a Party or the World Health Organization has information regarding a preparation exempted pursuant to paragraph 3 which in its opinion may require the termination, in whole or in part, of the exemption, it shall notify the Secretary-General and furnish him with the information in support of the notification. The Secretary-General shall transmit such notification, and any information which he considers relevant, to the Parties, to the Commission and, when the notification is made by a Party, to the World Health Organization. The World Health Organization shall communicate to the Commission an assessment of the preparation in relation to the matters specified in paragraph 2, together with a recommendation of the control measures, if any, from which the preparation should cease to be exempted. The Commission, taking into account the communication from the World Health Organization, whose assessment shall be determinative as to medical and scientific matters, and bearing in mind the economic, social, legal, administrative and other factors it may consider relevant, may decide to terminate the exemption of the preparation from any or all control measures. Any decision of the Commission taken pursuant to this paragraph shall be communicated by the Secretary-General to all States Members of the United Nations, to non-member States Parties to this Convention, to the World Health Organization and to the Board. All Parties shall take measures to terminate the exemption from the control measure or measures in question within 180 days of the date of the Secretary-General’s communication.
United Nations Convention against Illicit Traffic in Narcotic Drugs and Psychotropic Substances, 1988 (Extracts)\(^1\)

**Article 12**

**SUBSTANCES FREQUENTLY USED IN THE ILLICIT MANUFACTURE OF NARCOTIC DRUGS OR PSYCHOTROPIC SUBSTANCES**

1. The Parties shall take the measures they deem appropriate to prevent diversion of substances in Table I and Table II used for the purpose of illicit manufacture of narcotic drugs or psychotropic substances, and shall co-operate with one another to this end.

2. If a Party or the Board has information which in its opinion may require the inclusion of a substance in Table I or Table II, it shall notify the Secretary-General and furnish him with the information in support of that notification. The procedure described in paragraphs 2 to 7 of this article shall also apply when a Party or the Board has information justifying the deletion of a substance from Table I or Table II, or the transfer of a substance from one Table to the other.

3. The Secretary-General shall transmit such notification, and any information which he considers relevant, to the Parties, to the Commission, and, where notification is made by a Party, to the Board. The Parties shall communicate their comments concerning the notification to the Secretary-General, together with all supplementary information which may assist the Board in establishing an assessment and the Commission in reaching a decision.

4. If the Board, taking into account the extent, importance and diversity of the licit use of the substance, and the possibility and ease of using alternate substances both for licit purposes and for the illicit manufacture of narcotic drugs or psychotropic substances, finds:

   a) That the substance is frequently used in the illicit manufacture of a narcotic drug or psychotropic substance;

   b) That the volume and extent of the illicit manufacture of a narcotic drug or psychotropic substance creates serious public health or social problems, so as to warrant international action, it shall communicate to the Commission an assessment of the substance, including the likely effect of adding the substance to either Table I or Table II on both licit use and illicit manufacture, together with recommendations of monitoring measures, if any, that would be appropriate in the light of its assessment.

5. The Commission, taking into account the comments submitted by the Parties and the comments and recommendations of the Board, whose assessment shall be determinative as to scientific matters, and also taking into due consideration any other relevant factors, may decide by a two-thirds majority of its members to place a substance in Table I or Table II.

APPENDIX 2

RESOLUTION 1 (S-VIII) OF THE UNITED NATIONS COMMISSION
ON NARCOTIC DRUGS

Guidelines for the exemption of preparations from certain control measures under the
provisions of Article 3 of the 1971 Convention on Psychotropic Substances

The Commission on Narcotic Drugs,

Having taken note of documents MNH/78.1 and MNH/82.51 containing proposals by World
Health Organization consultative groups concerning guidelines for granting exemptions under the
provisions of article 3 of the 1971 Convention on Psychotropic Substances,

Having considered the report by the Secretary-General of 16 December 1983 entitled Review of
establishment of guidelines for the exemption of preparations under the provisions of article 3 of the
1971 Convention on Psychotropic Substances (E/CN.7/1984/4),

Recalling its resolutions 2 (S-VI) of 19 February 1980 and 5 (XXX) of 16 February 1983,

Bearing in mind that decisions taken by it in respect of the termination of an exemption must
consider the social and economic conditions pertaining in the country granting the exemption,
including the level of development of its national medical services and national drug distribution system,

Convinced of the need for Governments to contribute to the development of further guidelines,
in light of the experience gained during the application of the guidelines currently in force,

Approves the following guidelines for use by national authorities, the World Health
Organization and the Commission on Narcotic Drugs:

Guidelines proposed for use by national authorities

(a) A preparation containing a psychotropic substance in association with (i) another
psychotropic substance, (ii) a narcotic drug or (iii) a psychoactive substance not under
international control with known abuse potential, should not be exempted; nevertheless,
exemption of a preparation in any of the three above categories which is compounded in such a
manner that it presents a negligible risk of abuse may be envisaged;

(b) A preparation containing a psychotropic substance in association with a narcotic drug
listed in Schedule I or II of the Single Convention on Narcotic Drugs, 1961, should not be
exempted; exemption can only be authorized if the preparation has been listed in Schedule III of
that Convention by the Commission, in accordance with the amendment procedure established
by the provisions of article 3, paragraph 4, of the Convention;

c) A preparation containing a psychotropic substance in injectable dosage form should not
be exempted;

(d) A preparation containing a psychotropic substance should not be exempted from the provisions of article 10, paragraph 1, of the 1971 Convention on Psychotropic Substances;

e) A preparation containing a psychotropic substance should not be exempted from the provisions of article 10, paragraph 2, of the 1971 Convention on Psychotropic Substances, unless such exemption would be in keeping with national statutory requirements;

f) A preparation containing a psychotropic substance should not be exempted from the requirements of article 12 of the 1971 Convention on Psychotropic Substances;

g) Guidelines (d), (e), and (f) notwithstanding, in vitro diagnostic reagents, buffers and analytical standards containing psychotropic substances may be exempted from the provisions of articles 10 and 12 of the 1971 Convention.

Guidelines proposed for use by the World Health Organization

h) The World Health Organization should not routinely review Parties’ notifications of exemptions intended only for domestic use; however, where there is evidence that a specific exemption granted by a competent national authority does not comply with guidelines (a)–(e) above, and might constitute a danger to the public health of the country concerned, the World Health Organization should immediately draw the attention of the competent national authority to the possible public health hazard and advise the Commission on Narcotic Drugs of its action in this regard. If however, there is evidence that such exemption constitutes a danger to another country, the World Health Organization should proceed to examine the exemption as a matter of urgency.
APPENDIX 3

FLOW CHART OF THE EVALUATION PROCEDURE

WHO review of psychoactive substances for international control

Pre-review

- Proposal for pre-review by secretary
- Proposal for pre-review by expert
- Proposal for pre-review by observer

Expert Committee on Drug Dependence Decision does current information justify a critical review?

Critical review

- Positive decision on pre-review in previous meeting of the Expert Committee on Drug Dependence
- Notification by Treaty Party
- Explicit request by Commission on Narcotic Drugs
- Information on clandestine manufacturing of substance with no recognized therapeutic use

Circulation of agenda and questionnaires to WHO Member States

Report on questionnaires

Scientific part of critical review report

Circulation of combined reports among substantial contributors of information

Peer review by two experts

Report adaptation by WHO Secretariat

Comments

Final critical review report

Meeting documents including critical review reports published on the web

Report distributed among members of the Expert Committee on Drug Dependence

Information meeting requested?

Yes

No

Information meeting (preceding Expert Committee meeting)

Expert Committee meeting

No

Yes

Expert Committee on Drug Dependence proposed change in scheduling status?


Further handling by the United Nations Office on Drugs and Crime on behalf of the United Nations Secretary-General

Advice to the Director-General to make a recommendation to the United Nations

Note Verbale from the United Nations Secretary-General

Publication of Note Verbale on the web

WHO 09.28
ANNEX 7

Financial and administrative implications for the Secretariat of resolutions adopted by the Executive Board

1. Resolution EB126.R4  Monitoring of the achievement of the health-related Millennium Development Goals

2. Linkage to programme budget

<table>
<thead>
<tr>
<th>Strategic objective</th>
<th>Organization-wide expected results:</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. To combat HIV/AIDS, tuberculosis and malaria.</td>
<td>All expected results for this strategic objective.</td>
</tr>
<tr>
<td>4. To reduce morbidity and mortality and improve health during key stages of life, including pregnancy, childbirth, the neonatal period, childhood and adolescence, and improve sexual and reproductive health and promote active and healthy ageing for all individuals.</td>
<td>All expected results except 4.8.</td>
</tr>
<tr>
<td>10. To improve health services through better governance, financing, staffing and management, informed by reliable and accessible evidence and research.</td>
<td>All expected results except 10.6, 10.7 and 10.9.</td>
</tr>
</tbody>
</table>

(Briefly indicate the linkage with expected results, indicators, targets, baseline)

The resolution is particularly relevant to strategic objective 10, which concerns both improved management and organization of health service delivery through a primary health-care approach, and enhanced monitoring and evaluation of progress. The resolution is also highly relevant to all strategic objectives concerned with the achievement of specific health outcomes, especially strategic objectives 1–4. Of the latter group, it particularly concerns strategic objective 2 and strategic objective 4. Other strategic objectives with relevance to the resolution include strategic objective 5 on emergencies and crises (expected results 5.1–5.3), strategic objective 6 on reducing risk factors for health conditions linked to unhealthy lifestyles (expected result 6.6), strategic objective 7 on tackling social and economic determinants of health and enhancing health equity (expected result 7.3), strategic objective 8 on promoting healthy environments (expected results 8.1–8.2), strategic objective 9 on nutrition and food safety and security (expected results 9.1–9.4), strategic objective 11 on ensuring access to medical products and technologies (expected results 11.1–11.3) and strategic objective 12 on leadership, partnerships and collaboration with countries (expected results 12.1–12.3).

3. Budgetary implications

(a) Total estimated cost for implementation over the life-cycle of the Secretariat’s activities requested in the resolution (estimated to the nearest US$ 10,000, including staff and activities)

The two major streams of work and associated costs relate to: (i) the production of the annual report on the health-related Millennium Development Goals; and (ii) the provision of technical support to countries for enhanced programme implementation, monitoring and evaluation. The former is primarily carried out at headquarters, the latter through the regional offices.

Total production costs for the annual report (as part of the publication, World health statistics): US$ 350,000.

Staff costs: 33% full-time equivalent, grade P6; 40% full-time equivalent, grade P5; 50% full-time equivalent, grade P4; and 50% full-time equivalent, grade P3. Total: US$ 1.9 million.
Total costs for regional office inputs: US$ 2 million.

Total costs: US$ 4 250 000.

(b) Estimated cost for the biennium 2010–2011 (estimated to the nearest US$ 10 000 including staff and activities, and indicating at which levels of the Organization the costs will be incurred, identifying specific regions where relevant)

Total costs for implementation: US$ 1.7 million (US$ 900 000 at headquarters level and US$ 800 000 at regional office level).

(c) Is the estimated cost noted in (b), included within the existing approved Programme budget for the biennium 2010–2011?

Yes.

4. Financial implications

How will the estimated cost noted in 3(b) be financed (indicate potential sources of funds)?

Headquarters: 50% from assessed contributions, 50% from voluntary contributions.

Regional offices: 100% from voluntary contributions.

5. Administrative implications

(a) Implementation locales (indicate the levels of the Organization at which the work will be undertaken, identifying specific regions where relevant)

Implementation activities in respect of the annual report on the health-related Millennium Development Goals will take place at headquarters. Collaboration with all regional offices to enable improved availability of up-to-date information and support to countries for data analysis, reconciliation and estimation will be organized as needed.

(b) Can the resolution be implemented by existing staff? If not, please specify in (c) below

Headquarters: yes, providing that funding continues to be available.

Regional offices: no for the Regional Office for Africa and the Regional Office for South-East Asia.

(c) Additional staffing requirements (indicate additional required staff – full-time equivalents – by levels of the Organization, identifying specific regions where relevant and noting necessary skills profile)

The Regional Office for Africa and the Regional Office for South-East Asia will need one staff member (full-time equivalent) to work with countries.

(d) Time frames (indicate broad time frames for implementation of activities)

From 2010 to 2015.
4. To reduce morbidity and mortality and improve health during key stages of life, including pregnancy, childbirth, the neonatal period, childhood and adolescence, and improve sexual and reproductive health and promote active and healthy ageing for all individuals.

4.4 Guidelines, approaches and tools for improving neonatal survival and health applied at country level, with technical support provided to Member States for intensified action towards universal coverage, effective interventions and monitoring of progress.

4.5 Guidelines, approaches and tools for improving child health and development applied at the country level, with technical support provided to Member States for intensified action towards universal coverage of the population with effective interventions and for monitoring progress, taking into consideration international and human-rights norms and standards, notably those stipulated in the Convention on the Rights of the Child.

7. To address the underlying social and economic determinants of health through policies and programmes that enhance health equity and integrate pro-poor, gender-responsive, and human rights-based approaches.

7.2 Initiative taken by WHO in providing opportunities and means for intersectoral collaboration at national and international levels to address social and economic determinants of health, including understanding and acting upon the public health implications of trade and trade agreements, and to encourage poverty reduction and sustainable development.

9. To improve nutrition, food safety and food security, throughout the life-course, and in support of public health and sustainable development.

9.1 Partnerships and alliances formed, leadership built and coordination and networking developed with all stakeholders at country, regional and global levels, in order to promote advocacy and communication, stimulate intersectoral actions, increase investment in nutrition, food-safety and food-security interventions, and develop and support a research agenda.

9.2 Norms, including references, requirements, research priorities, guidelines, training manuals and standards, produced and disseminated to Member States in order to increase their capacity to assess and respond to all forms of malnutrition, and zoonotic and non-zoonotic foodborne diseases, and to promote healthy dietary practices.

9.3 Monitoring and surveillance of needs and assessment and evaluation of responses in the area of nutrition and diet-related chronic diseases strengthened, and ability to identify best policy options improved, in stable and emergency situations.
9.4 Capacity built and support provided to target Member States for the development, strengthening and implementation of nutrition plans, policies and programmes aimed at improving nutrition throughout the life-course, in stable and emergency situations.

(Briefly indicate the linkage with expected results, indicators, targets, baseline)

- Guidelines on nutrition interventions to be reviewed and developed, scaling up from 10 to 40 guidelines.
- Monitoring child growth, infant and young child feeding practices, micronutrient status and implementation of nutrition interventions performed, scaling up from 50 to 70 countries implementing growth standards; 5 micronutrient surveys implemented; and implementation data collected in 150 countries.
- Interagency collaborative initiatives developed at global and country levels in order to scale up nutrition programmes: 5 initiatives.
- Technical support for scaling up nutrition interventions provided to 20 countries and capacity building provided to an additional 15 countries.
- Technical support provided to Member States in strengthening and implementing national nutrition policies and strategies to scale up action in tackling the double burden of malnutrition: an additional 15 countries provided with support.
- Technical guidance provided on priority nutrition actions aimed at the prevention of tuberculosis and support against HIV: an additional 15 countries provided with support.

3. Budgetary implications

(a) Total estimated cost for implementation over the life-cycle of the Secretariat’s activities requested in the resolution (estimated to the nearest US$ 10 000, including staff and activities)

US$ 12 million (staff: US$ 5 million, activities: US$ 7 million) for three years.

(b) Estimated cost for the biennium 2010–2011 (estimated to the nearest US$ 10 000 including staff and activities, and indicating at which levels of the Organization the costs will be incurred, identifying specific regions where relevant)


(c) Is the estimated cost noted in (b), included within the existing approved Programme budget for the biennium 2010–2011?

Yes.

4. Financial implications

How will the estimated cost noted in 3(b) be financed (indicate potential sources of funds)?

A global resource mobilization plan is being prepared to approach potential donors. Partial funding has been provided by the Governments of Italy, Japan, Luxembourg, Spain and United States of America and philanthropic bodies.
5. Administrative implications

(a) Implementation locales (indicate the levels of the Organization at which the work will be undertaken, identifying specific regions where relevant)

Although the normative work (including guidelines development and scientific reviews) will be carried out at headquarters, the majority of activities will be undertaken at country and regional levels. Priority will be given to the 36 countries identified by WHO upon which malnutrition places the highest burden of mortality and morbidity.

(b) Can the resolution be implemented by existing staff? If not, please specify in (c) below

As already recommended in the review conducted of WHO’s nutrition work, staffing needs to be strengthened. This is particularly the case at country level and in some of the regional offices.

(c) Additional staffing requirements (indicate additional required staff – full-time equivalents – by levels of the Organization, identifying specific regions where relevant and noting necessary skills profile)

Staffing needs to be strengthened in all regions, particularly in African, South-East Asia and Eastern Mediterranean Regions. There is a particular need for technical nutrition staff at the P.2 and P.3 levels.

(d) Time frames (indicate broad time frames for implementation of activities)

Implementation of some activities under this resolution has already started in the current biennium. Most activities will be implemented by the first quarter of 2011.

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1. Resolution EB126.R7 Advancing food safety initiatives

2. Linkage to programme budget

   Strategic objective:
   1. To reduce the health, social and economic burden of communicable diseases.
   5. To reduce the health consequences of emergencies, disasters, crises and conflicts, and minimize their social and economic impact.
   9. To improve nutrition, food safety and food security, throughout the life-course, and in support of public health and sustainable development.

   Organization-wide expected result:
   1.3. Effective coordination and support provided to Member States in order to provide access for all populations to interventions for the prevention, control, elimination and eradication of neglected tropical diseases, including zoonotic diseases.
   5.5. Support provided to Member States for strengthening national preparedness and for establishing alert and response mechanisms for food-safety and environmental health emergencies.
   9.1. Partnerships and alliances formed, leadership built and coordination and networking developed with all stakeholders at country, regional and global levels, in order to promote advocacy and communication, stimulate intersectoral actions, increase investment in nutrition, food-safety and food-security interventions, and develop and support a research agenda.
   9.2. Norms, including references, requirements, research priorities, guidelines, training manuals and standards, produced and disseminated to Member States in order to increase their capacity to assess and respond to all forms of malnutrition, and zoonotic and non-zoonotic foodborne diseases, and to promote healthy dietary practices.
9.5. Systems for surveillance, prevention and control of zoonotic and non-zoonotic foodborne diseases strengthened; food-hazard monitoring and evaluation programmes established and integrated into existing national surveillance systems, and results disseminated to all key players.

9.6. Capacity built and support provided to Member States, including their participation in international standard-setting in order to increase their ability to assess risk in the areas of zoonotic and non-zoonotic foodborne diseases and food safety, and to develop and implement national food-control systems, with links to international emergency systems.

(Briefly indicate the linkage with expected results, indicators, targets, baseline)

The resolution will provide an updated framework for WHO’s normative work on food safety within the existing expected results, with an additional emphasis on strengthening scientific advice, the estimation of the health burden imposed by foodborne disease, support for the International Food Safety Authorities Network (INFOSAN) and cross-sectoral prevention of zoonotic diseases.

3. Budgetary implications
   (a) Total estimated cost for implementation over the life-cycle of the Secretariat’s activities requested in the resolution (estimated to the nearest US$ 10 000, including staff and activities):

   (b) Estimated cost for the biennium 2010–2011 (estimated to the nearest US$ 10 000 including staff and activities, and indicating at which levels of the Organization the costs will be incurred, identifying specific regions where relevant):
   US$ 9.7 million at headquarters.

   (c) Is the estimated cost noted in (b) included within the existing approved Programme budget for the biennium 2010–2011?
   US$ 8 million are included in the Programme budget.

4. Financial implications
   How will the estimated cost noted in 3(b) be financed (indicate potential sources of funds)?
   Through the extrabudgetary funds of interested Member States and relevant nongovernmental organizations.

5. Administrative implications
   (a) Implementation locales (indicate the levels of the Organization at which the work will be undertaken, identifying specific regions where relevant)
   Implementation will be organized at headquarters, in coordination with all six regional offices and selected countries of each region.

   (b) Can the resolution be implemented by existing staff? If not, please specify in (c) below
   No.
(c) **Additional staffing requirements (indicate additional required staff – full-time equivalents – by levels of the Organization, identifying specific regions where relevant and noting necessary skills profile)**

Two staff members in the professional category at headquarters.

**(d) Time frames (indicate broad time frames for implementation of activities)**

Food safety initiatives are continuing activities. The next progress report will be made to the Sixty-fifth World Health Assembly; thus the major new initiatives should be evaluated by the end of the period 2010–2011.

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1. **Resolution EB126.R12** The improvement of health through safe and environmentally sound waste management

2. **Linkage to programme budget**

   **Strategic objective:**

   8. To promote a healthier environment, intensify primary prevention and influence public policies in all sectors so as to address the root causes of environmental threats to health.

   **Organization-wide expected result:**

   8.2 Technical support and guidance provided to Member States for the implementation of primary prevention interventions that reduce environmental hazards to health, enhance safety and promote public health, including in specific settings (e.g. workplaces, homes or urban settings) and among vulnerable population groups (e.g. children).

   *(Briefly indicate the linkage with expected results, indicators, targets, baseline)*

   The resolution is consistent with the expected result and implementation would facilitate achievement of the target for 2011 of 12 Member States implementing primary prevention interventions in order to reduce environmental risks to health, with WHO technical support, in at least one of the following settings: workplaces, homes or urban settings. The baseline figure for 2010 will remain the same as for 2009 (namely 8).

3. **Budgetary implications**

   (a) **Total estimated cost for implementation over the life-cycle of the Secretariat’s activities requested in the resolution (estimated to the nearest US$ 10,000, including staff and activities)**

   No additional costs will be incurred regarding activities. The resolution requests the Secretariat to support the implementation of the actions set out in the Bali Declaration, within the Organization’s mandate and available resources. The workplan for the biennium 2010–2011 already includes activities aimed at responding to the problem of hazardous waste, in particular health care waste, in conjunction with other relevant bodies including UNEP and the Basel Convention secretariat.

   (b) **Estimated cost for the biennium 2010–2011 (estimated to the nearest US$ 10,000 including staff and activities, and indicating at which levels of the Organization the costs will be incurred, identifying specific regions where relevant)**

   No additional costs are envisaged specifically under the resolution.

   (c) **Is the estimated cost noted in (b), included within the existing approved Programme budget for the biennium 2010–2011?**

   Not applicable.
4. Financial implications

How will the estimated cost noted in 3(b) be financed (indicate potential sources of funds)?

Not applicable.

5. Administrative implications

(a) Implementation locales (indicate the levels of the Organization at which the work will be undertaken, identifying specific regions where relevant)

Primary prevention activities will mainly be conducted at country level; multicountry projects will be undertaken through headquarters and the regional offices; and liaison with UNEP and the Basel Convention will mainly be managed through headquarters and the regional offices.

(b) Can the resolution be implemented by existing staff? If not, please specify in (c) below

Yes.

(c) Additional staffing requirements (indicate additional required staff – full-time equivalents – by levels of the Organization, identifying specific regions where relevant and noting necessary skills profile)

Not applicable.

(d) Time frames (indicate broad time frames for implementation of activities)

There will be a continuing need to tackle the problem of health-care and other hazardous waste to ensure that waste generation is minimized on a lasting basis. Therefore, the initial time frame under this resolution will follow the Medium-term strategic plan until 2013. Thereafter, it will be reviewed as necessary.

1. Resolution EB126.R13 Improvement of health through sound management of obsolete pesticides and other obsolete chemicals

2. Linkage to programme budget

Strategic objective:

8. To promote a healthier environment, intensify primary prevention and influence public policies in all sectors so as to address the root causes of environmental threats to health.

Organization-wide expected result:

8.1 Evidence-based assessments made, and norms and standards formulated and updated on major environmental hazards to health (e.g. poor air quality, chemical substances, electromagnetic fields, radon, poor-quality drinking-water and waste-water reuse).

8.2 Technical support and guidance provided to Member States for the implementation of primary prevention interventions that reduce environmental hazards to health, enhance safety and promote public health, including in specific settings (e.g. workplaces, homes or urban settings) and among vulnerable population group (e.g. children).

8.3 Technical assistance and support provided to Member States for strengthening national occupational and environmental health risk management systems, functions and services.

8.5 Health-sector leadership enhanced for creating a healthier environment and changing policies in all sectors so as to tackle the root causes of environmental threats to health, through means such as responding to emerging and re-emerging consequences of
development on environmental health and altered patterns of consumption and production and to the damaging effect of evolving technologies.

(Briefly indicate the linkage with expected results, indicators, targets, baseline)

The resolution is consistent with the listed expected results and supports indicators relating to: Member States assessing environmental threats to health, implementing primary prevention of environmental risks to health and implementing national action plans or policies for the management of occupational health. The baselines remain the same.

<table>
<thead>
<tr>
<th>3. Budgetary implications</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Total estimated cost for implementation over the life-cycle of the Secretariat’s activities requested in the resolution (estimated to the nearest US$ 10 000, including staff and activities)</td>
</tr>
<tr>
<td>The timeline is not specified and implementation of the resolution is expected to continue over time since additional pesticides and chemicals are likely to be rendered obsolete by risk management actions. The Secretariat’s planned activities are largely directed towards specific substances of concern, such as phasing out the use of lead, mercury and asbestos. The current level of activity at headquarters on such substances entails a cost of some US$ 500 000 per biennium. Implementation of comprehensive activities on obsolete pesticides and obsolete chemicals in general is estimated to require an additional cost of approximately the same amount.</td>
</tr>
<tr>
<td>(b) Estimated cost for the biennium 2010–2011 (estimated to the nearest US$ 10 000 including staff and activities, and indicating at which levels of the Organization the costs will be incurred, identifying specific regions where relevant)</td>
</tr>
<tr>
<td>See section 3(a) above.</td>
</tr>
<tr>
<td>(c) Is the estimated cost noted in (b) included within the existing approved Programme budget for the biennium 2010–2011?</td>
</tr>
<tr>
<td>The current level of activity on specific substances of concern, estimated at US$ 500 000, is included within the existing Programme budget.</td>
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<tr>
<th>4. Financial implications</th>
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<tbody>
<tr>
<td>How will the estimated cost noted in 3(b) be financed (indicate potential sources of funds)?</td>
</tr>
<tr>
<td>Voluntary contributions from interested Member States and relevant nongovernmental resources.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>5. Administrative implications</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Implementation locales (indicate the levels of the Organization at which the work will be undertaken, identifying specific regions where relevant)</td>
</tr>
<tr>
<td>Headquarters (coordination of implementation and provision of general guidance to regional and country offices) and regional offices in consultation with countries.</td>
</tr>
<tr>
<td>(b) Can the resolution be implemented by existing staff? If not, please specify in (c) below</td>
</tr>
<tr>
<td>The current level of activity can be sustained by existing staff. Implementation of comprehensive activities on obsolete pesticides and obsolete chemicals in general will require additional staff (see (c) below).</td>
</tr>
<tr>
<td>(c) Additional staffing requirements (indicate additional required staff – full-time equivalents – by levels of the Organization, identifying specific regions where relevant and noting necessary skills profile)</td>
</tr>
<tr>
<td>Comprehensive activities will require an additional staff member in the professional category (25% full-time equivalent).</td>
</tr>
<tr>
<td>(d) Time frames (indicate broad time frames for implementation of activities)</td>
</tr>
<tr>
<td>The timeline is not specified and implementation is expected to continue over time (see 3(a) above).</td>
</tr>
</tbody>
</table>

2. Linkage to programme budget

   Strategic objective:
   1. To reduce the health, social and economic burden of communicable diseases.
   4. To reduce morbidity and mortality and improve health during key stages of life, including pregnancy, childbirth, the neonatal period, childhood and adolescence, and improve sexual and reproductive health and promote active and healthy ageing for all individuals.

   Organization-wide expected result:
   1.1 Policy and technical support provided to Member States in order to maximize equitable access of all people to vaccines of assured quality, including new immunization products and technologies, and to integrate other essential child-health interventions with immunization.
   4.5 Guidelines, approaches and tools for improving child health and development applied at the country level, with technical support provided to Member States for intensified action towards universal coverage of the population with effective interventions and for monitoring progress, taking into consideration international and human-rights norms and standards, notably those stipulated in the Convention on the Rights of the Child.

   (Briefly indicate the linkage with expected results, indicators, targets, baseline)
   The resolution is consistent with the listed expected results and supports the following indicators: for strategic objective 1 – (i) number of Member States with at least 90% immunization coverage (DPT 3); and (ii) number of Member States that have introduced Haemophilus influenzae type b vaccine into their national immunization schedule; and for strategic objective 4 – (i) number of targeted Member States that have an integrated policy on universal access to effective interventions for improving maternal, newborn and child health and (ii) number of Member States implementing strategies for increasing coverage with child health and development interventions.

3. Budgetary implications

   (a) Total estimated cost for implementation over the life-cycle of the Secretariat’s activities requested in the resolution (estimated to the nearest US$ 10 000, including staff and activities)
   US$ 110 million are required for the period 2010–2015 in respect of costs at headquarters and at regional and country office levels.

   (b) Estimated cost for the biennium 2010–2011 (estimated to the nearest US$ 10 000 including staff and activities, and indicating at which levels of the Organization the costs will be incurred, identifying specific regions where relevant)
   In respect of strategic objective 1: headquarters – total US$ 5.9 million (US$ 900 000 for staff, US$ 5 million for activities); regional and country offices – total US$ 27.4 million (US$ 5 million for staff, US$ 22.4 million for activities).
   In respect of strategic objective 4: headquarters – total US$ 600 000 (US$ 450 000 for staff, US$ 150 000 for activities); regional and country offices – total US$ 8 million (US$ 2.4 million for staff, US$ 5.6 million for activities).

   (c) Is the estimated cost noted in (b), included within the existing approved Programme budget for the biennium 2010–2011?
   Yes.
4. Financial implications

How will the estimated cost noted in 3(b) be financed (indicate potential sources of funds)?

In respect of strategic objective 1: funds for 2010 are available through the GAVI Alliance and the Gates Foundation. Some of the funding required for 2011 may be available through the same sources, but a funding gap is likely to appear in 2011.

In respect of strategic objective 4: voluntary contributions will be sought to fund these activities.

5. Administrative implications

(a) Implementation locales (indicate the levels of the Organization at which the work will be undertaken, identifying specific regions where relevant)

All levels of WHO, with a specific focus on the 68 priority countries that are the focus of the “Countdown to 2015” initiative, on which the disease places a high burden. Most of the countries concerned are also eligible for funding from the GAVI Alliance.

(b) Can the resolution be implemented by existing staff? If not, please specify in (c) below

No. Additional staff will be needed to implement the resolution, especially in countries with a high burden of the disease.

(c) Additional staffing requirements (indicate additional required staff – full-time equivalents – by levels of the Organization, identifying specific regions where relevant and noting necessary skills profile)

Three staff members (full-time equivalent) at the P4 grade: 1 at headquarters, 1 at the Regional Office for South-East Asia and 1 at country level.

(d) Time frames (indicate broad time frames for implementation of activities)

The period 2010–2015.

1. Resolution EB126.16 Viral hepatitis

2. Linkage to programme budget

<table>
<thead>
<tr>
<th>Strategic objective:</th>
<th>Organization-wide expected result:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. To reduce the health, social and economic burden of communicable diseases.</td>
<td>1.1 Policy and technical support provided to Member States in order to maximize equitable access of all people to vaccines of assured quality, including new immunization products and technologies, and to integrate other essential child-health interventions with immunization.</td>
</tr>
<tr>
<td></td>
<td>1.4 Policy and technical support provided to Member States in order to enhance their capacity to carry out surveillance and monitoring of all communicable diseases of public health importance.</td>
</tr>
<tr>
<td></td>
<td>1.5 New knowledge, intervention tools and strategies that meet priority needs for the prevention and control of communicable diseases developed and validated, with scientists from developing countries increasingly taking the lead in this research.</td>
</tr>
</tbody>
</table>

(Briefly indicate the linkage with expected results, indicators, targets, baseline)

The resolution is consistent with the expected result. Indicators specific to prevention of viral hepatitis will be designed as needed.
### 3. Budgetary implications

(a) **Total estimated cost for implementation over the life-cycle of the Secretariat’s activities requested in the resolution (estimated to the nearest US$ 10 000, including staff and activities)**

US$ 30 million are needed for the next five years. Of this amount, one third (US$ 10 million) is needed at headquarters for global planning and coordination between stakeholders, global policy guidance, and the provision of support to regional and country offices; two thirds (US$ 20 million) are needed for support activities at regional and country levels.

(b) **Estimated cost for the biennium 2010–2011 (estimated to the nearest US$ 10 000 including staff and activities, and indicating at which levels of the Organization the costs will be incurred, identifying specific regions where relevant)**

Total costs are estimated at US$ 6 million per year.

(c) **Is the estimated cost noted in (b) included within the existing approved Programme budget for the biennium 2010–2011?**

The Organization’s hepatitis prevention activities involve a number of technical units. It is difficult to know the true amount of resources available for these activities as they may not be directly identified in the Programme budget and may, for example, be covered under references to blood safety, injection safety, food safety, cancer prevention, child immunization or treatment of opportunistic infections in HIV/AIDS.

### 4. Financial implications

**How will the estimated cost noted in 3(b) be financed (indicate potential sources of funds)?**

Additional funding from voluntary contributions is expected through active resource mobilization.

### 5. Administrative implications

(a) **Implementation locales (indicate the levels of the Organization at which the work will be undertaken, identifying specific regions where relevant)**

Currently, most activities are performed at headquarters (policy and technical guidance, global advocacy and stakeholder coordination, and fund-raising) and in two WHO regions (Eastern Mediterranean and Western Pacific).

(b) **Can the resolution be implemented by existing staff? If not, please specify in (c) below**

No.

(c) **Additional staffing requirements (indicate additional required staff – full-time equivalents – by levels of the Organization, identifying specific regions where relevant and noting necessary skills profile)**

At headquarters, at least two additional staff (full-time equivalents) will be required in the professional category, together with one staff member (full-time equivalent) in the general service category. During the biennium 2010–2011, one additional staff member (full-time equivalent) will be needed in the professional category in each of three regional offices (plus administrative support); during the biennium 2012–2013, three more staff (full-time equivalents) will be needed for the other regional offices (plus administrative support). A total of eight staff (full-time equivalents) will therefore be required in the professional category, together with three or four staff (full-time equivalents) in the general service category. In at least 10 countries, a dedicated national programme officer will be needed.

(d) **Time frames (indicate broad time frames for implementation of activities)**

The global programme will be expanded into the African, European, and Eastern Mediterranean regions in 2010, and into all regions during the biennium 2010–2011.