WHO-recommended standards for surveillance of selected vaccine-preventable diseases

DEPARTMENT OF VACCINES AND BIOLOGICALS

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<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>AFRO</td>
<td>WHO Regional Office for Africa</td>
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<tr>
<td>9GPW</td>
<td>9th Global Programme of Work</td>
<td></td>
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<tr>
<td>AFRO</td>
<td>WHO Regional Office for Africa</td>
<td></td>
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<tr>
<td>AFP</td>
<td>Acute flaccid paralysis</td>
<td></td>
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<tr>
<td>AMRO</td>
<td>WHO Regional Office for the Americas</td>
<td></td>
</tr>
<tr>
<td>CIE</td>
<td>counter immunoelectrophoresis</td>
<td></td>
</tr>
<tr>
<td>CSF</td>
<td>cerebrospinal fluid</td>
<td></td>
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<tr>
<td>DTP3</td>
<td>third dose of diphtheria-tetanus-pertussis vaccine</td>
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<tr>
<td>EMRO</td>
<td>WHO Regional Office for the Eastern Mediterranean</td>
<td></td>
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<tr>
<td>EPI</td>
<td>Expanded Programme on Immunization</td>
<td></td>
</tr>
<tr>
<td>EURO</td>
<td>WHO Regional Office for Europe</td>
<td></td>
</tr>
<tr>
<td>GPV</td>
<td>Global Programme for Vaccines and Immunization</td>
<td></td>
</tr>
<tr>
<td>HAV</td>
<td>hepatitis A virus</td>
<td></td>
</tr>
<tr>
<td>HBc</td>
<td>hepatitis B core</td>
<td></td>
</tr>
<tr>
<td>HBV</td>
<td>hepatitis B virus</td>
<td></td>
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<tr>
<td>HCV</td>
<td>hepatitis C virus</td>
<td></td>
</tr>
<tr>
<td>HDV</td>
<td>hepatitis D virus</td>
<td></td>
</tr>
<tr>
<td>HEV</td>
<td>hepatitis E virus</td>
<td></td>
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<tr>
<td>HBsAg</td>
<td>hepatitis B surface antigen</td>
<td></td>
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<tr>
<td>HepB3</td>
<td>third dose of hepatitis B vaccine</td>
<td></td>
</tr>
<tr>
<td>Hib3</td>
<td>third dose of <em>Haemophilus influenzae</em> type b vaccine</td>
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<tr>
<td>IgA</td>
<td>immunoglobulin A</td>
<td></td>
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<tr>
<td>IgG</td>
<td>immunoglobulin G</td>
<td></td>
</tr>
<tr>
<td>IgM</td>
<td>immunoglobulin M</td>
<td></td>
</tr>
<tr>
<td>ml</td>
<td>millilitre</td>
<td></td>
</tr>
<tr>
<td>NT</td>
<td>neonatal tetanus</td>
<td></td>
</tr>
<tr>
<td>OPV3</td>
<td>third dose of oral polio vaccine</td>
<td></td>
</tr>
<tr>
<td>PAB</td>
<td>protected at birth</td>
<td></td>
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<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
<td></td>
</tr>
<tr>
<td>SEARO</td>
<td>WHO Regional Office for South-East Asia</td>
<td></td>
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<tr>
<td>TT</td>
<td>tetanus toxoid</td>
<td></td>
</tr>
<tr>
<td>TT2+</td>
<td>second and subsequent doses of tetanus toxoid</td>
<td></td>
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<tr>
<td>WHO</td>
<td>World Health Organization</td>
<td></td>
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<tr>
<td>WPRO</td>
<td>WHO Regional Office for the Western Pacific</td>
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</table>
Introduction

The purpose of this document is to provide WHO recommendations on surveillance standards for selected vaccine-preventable diseases. These recommendations should be carefully adapted to meet national needs according to a country’s own disease control priorities, objectives and strategies.

Disease surveillance is defined as the routine ongoing collection, analysis and dissemination of health data. An effective surveillance system has the following functions:

- detection and notification of health events
- collection and consolidation of pertinent data
- investigation and confirmation (epidemiological, clinical and/or laboratory) of cases or outbreaks
- routine analysis and creation of reports
- feedback of information to those providing the data, and
- feed-forward (i.e. the forwarding of data to more central levels).

The rationale for surveillance of a specific health event must be established and based on clear national priorities, disease control objectives and strategies; otherwise, the disease surveillance data collected may be irrelevant. Decisions as to what data to collect should be based on what analyses are needed to guide public health decision-making. To avoid overburdening health staff at peripheral levels, the surveillance system should be as streamlined as possible by collecting the minimum amount of data necessary, and by using the most efficient and appropriate means to collect, consolidate and transfer data. Staff at all levels should be trained and encouraged to analyse and use their data. Data that can be more efficiently collected from other sources (e.g. survey) should not be included in a surveillance system.

An effective surveillance system is:

- useful
- efficient
- flexible
- representative
- simple
These attributes should be assessed when evaluating a surveillance system.

At the national level, clear surveillance “standards” must be established for maximum efficiency and so that data are comparable throughout the country. These standards would include:

- a case definition
- the type of surveillance to be conducted
- the data elements to be collected
- the minimum analyses and routine reports to be created
- the use of data for making decisions

For surveillance to be operational, the following needs to be carefully defined:

- the process of surveillance
- the tasks at each level
- the data/specimen flow
- the logistics, including staff issues
  - designation of staff
  - staff training
  - appropriate tool distribution (e.g. means of communication, transportation, specimen kits)

As a part of supervision, standard performance indicators should be monitored to identify weaknesses in the system so that corrective action can be taken.
# Diphtheria

## Rationale for surveillance

Diphtheria is a widespread severe infectious disease that has the potential for epidemics. The control of diphtheria is based on the following three measures. 1) Primary prevention of disease by ensuring high population immunity through immunization. 2) Secondary prevention of spread by the rapid investigation of close contacts, to ensure their proper treatment. 3) Tertiary prevention of complications and deaths by early diagnosis and proper management. Surveillance data can be used to monitor levels of coverage (target > 90%) and disease as a measure of the impact of control programmes. Recent epidemics have highlighted the need for adequate surveillance and epidemic preparedness.

## Recommended case definition

### Clinical description

An illness characterised by laryngitis or pharyngitis or tonsillitis, and an adherent membrane of the tonsils, pharynx and/or nose.

### Laboratory criteria for diagnosis

Isolation of *Corynebacterium diphtheriae* from a clinical specimen, or fourfold or greater rise in serum antibody (but only if both serum samples were obtained before the administration of diphtheria toxoid or antitoxin).

### Case classification

<table>
<thead>
<tr>
<th>Suspected</th>
<th>Not applicable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Probable</td>
<td>A case that meets the clinical description</td>
</tr>
<tr>
<td>Confirmed</td>
<td>A probable case that is laboratory confirmed or linked epidemiologically to a laboratory confirmed case</td>
</tr>
</tbody>
</table>

**Note:** Persons with positive *C. diphtheriae* cultures and not meeting the clinical description (i.e. asymptomatic carriers) should not be reported as probable or confirmed diphtheria cases.

## Recommended types of surveillance

- Routine monthly reporting of aggregated data of probable or confirmed cases is recommended from peripheral level to intermediate and central levels. Zero reporting should be required at all levels
- All outbreaks should be investigated immediately and case-based data collected
- In countries achieving low incidence (usually where coverage is >85-90%) immediate reporting of case-based data of probable or confirmed cases is recommended from peripheral level to intermediate and central levels
**Diphtheria (continued)**

**Recommended minimum data elements**

**Aggregated data:**
- Number of cases
- Number of third doses of diphtheria toxoid containing vaccine (e.g. DTP3) administered to infants

**Case-based data:**
- Unique identifier
- Geographical area (e.g. district) name
- Date of birth
- Date of onset
- Date of first treatment
- Treatment type:
  - 1=antibiotic & antitoxin; 2=antibiotic only; 3=antitoxin only; 4=no or other treatment; 9=unknown.
- Laboratory result:
  - 1=toxigenic *C. diphtheriae* isolated; 2=non-toxigenic *C. diphtheriae* isolated;
  - 3=*C. diphtheriae* isolated, toxigenicity unknown; 4=*C. diphtheriae* not isolated;
  - 5=no specimen processed; 9=unknown.
- Total diphtheria vaccine (DTP, DT or Td) doses received
- Date of last dose
- Final classification of the case:
  - 1=confirmed; 2=probable; 3=discarded
- Outcome:
  - 1=alive; 2=dead; 3=unknown

**Recommended data analyses, presentation, reports**

**Aggregated data:**
- Incidence rate by month, year, and geographic area
- DTP3 coverage by year and geographic area
- Completeness/timeliness of monthly reporting
- Proportional morbidity (compared to other diseases of public health importance)

**Case-based data:** same as aggregated data plus the following:
- Age-specific incidence rate
- Cases by immunization status, laboratory results, treatment type
- Cases treated "on time" (≤ seven days of onset)
- Case fatality rate
- Proportional mortality (compared to other diseases of public health importance)
## Special aspects

More detailed information available from the Expanded Programme on Immunization (EPI), Department of Vaccines and Biologicals (VAB).

## Principle uses of data for decision-making

- Monitor case fatality rate and, if high, determine cause (e.g. poor case management, lack of antibiotics/anti-toxin, patients not seeking treatment in time) so that corrective action can be taken
- Determine age-specific incidence rate, geographical area, and season of diphtheria cases to know risk groups and risk periods
- Monitor incidence rate to assess impact of control efforts
- Monitor immunization coverage per geographical area to identify areas of poor programme performance
- Detect outbreaks and implement control measures
- Investigate outbreaks to understand epidemiology, determine why the outbreak occurred (e.g. vaccine failure, failure to immunise, accumulation of susceptibles, waning immunity, new toxigenic strain), and ensure proper case management

**Note:** In addition to surveillance, carefully designed serologic studies can be used to monitor the immune status of different age groups.

## Contact information

**Regional Offices and Headquarters (V&B)**

See Annex 1 for addresses and fax numbers.
### Acute viral hepatitis

#### Rationale for surveillance

Several distinct infections are grouped as viral hepatitis. Transmission is mainly through the oral-faecal route for hepatitis A and E, and percutaneous exposure to body fluids, including sexual intercourse, for hepatitis B, C, and D. The course of the disease may be fulminating (e.g. hepatitis E in pregnancy); chronic infection and severe sequel occur mainly for hepatitis B, C, and D.

Control measures for blood-related transmission include ensuring transfusion safety, injection safety, and (for hepatitis A and hepatitis B at least) immunization. Hepatitis B is targeted by WHO (9GPW6.3) for reduced incidence/prevalence.

#### Recommended case definition

##### Clinical description

An acute illness typically including acute jaundice, dark urine, anorexia, malaise, extreme fatigue, and right upper quadrant tenderness. Biologic signs include increased urine urobilinogen and >2.5 times the upper limit of serum alanine aminotransferase.

**Note:** Most infections occur during early childhood. A variable proportions of adult infections are asymptomatic.

##### Laboratory criteria for diagnosis

- **Hepatitis A:** positive for IgM anti-HAV
- **Hepatitis B:** positive for IgM anti-HBc-positive or (less preferably) hepatitis B surface antigen (HBsAg)
- **Non-A, non-B:** negative for IgM anti-HAV and IgM anti-HBc or (less preferably) HBsAg

**Note:** The anti-HBc IgM test, specific for acute infection, is not available in most countries. HBsAg is often available, but is less preferable since it cannot distinguish acute new infections from exacerbation of chronic hepatitis B. Nevertheless, continued HBsAg seropositivity (> six months) is an indicator of chronic infection. For patients with non-A, non-B, the following testing is used for a diagnosis of acute hepatitis C, D, or E:

- **Hepatitis C:** positive for anti-HCV
- **Hepatitis D:** positive for IgM anti-HBc or (less preferably) HBsAg plus anti-HDV positive (N.B. only occurs as co-infection or super-infection of hepatitis B)
- **Hepatitis E:** positive for IgM anti-HEV

##### Case classification

- **Suspected:** A case that is compatible with the clinical description
- **Probable:** Not applicable
- **Confirmed:** A suspected case that is laboratory confirmed or, for hepatitis A only, a case compatible with the clinical description, in a person who has an epidemiological link (i.e. household or sexual contact with an infected person during the 15-50 days before the onset of symptoms) with a laboratory-confirmed case of hepatitis A
Acute viral hepatitis (continued)

Recommended types of surveillance

- Routine monthly reporting of aggregated data of suspected cases, and if available, the number of confirmed cases of each type of hepatitis is recommended from the peripheral level to intermediate and central levels
- Zero reporting should be required at all levels
- All outbreaks should be investigated immediately and confirmed serologically

Recommended minimum data elements

Aggregated data:

- Number of third doses of hepatitis B vaccine (HepB3) administered to infants
- Number of suspect cases
- If available, number of confirmed cases by each type of hepatitis

Recommended data analyses, presentation, reports
(from multiple sources of data, in addition to surveillance data):

- HepB3 coverage in infants by year and geographic area
- Acute viral hepatitis incidence by year, month, geographical area, and (if data exist) age group
- Where data exist on etiologic agent, incidence rate of each type of acute viral hepatitis by geographic area, year, month and age group
- Proportion of all cases of chronic liver disease, cirrhosis, and primary liver cancer that are HBsAg positive or anti-HCV positive. (see special aspects section)

Principle uses of data for decision-making

- Monitor HepB3 coverage by geographic area to measure areas with weak performance and take action
- Investigate all suspected/reported outbreaks
- Determine the specific cause of acute viral hepatitis cases (reported routinely or during outbreaks), so that corrective measures can be taken
- Understand the epidemiology of hepatitis by etiologic agent in terms of distribution over time, by age group, and by geographic area
- Measure the incidence (including age-specific incidence) and prevalence of HBsAg and anti-HCV
- Measure the proportion of acute viral hepatitis, chronic liver disease, cirrhosis, and primary liver cancer that are hepatitis B virus or hepatitis C virus carriers to:
  1) determine the burden of the disease in the population;
  2) prioritize it among other diseases of public health importance; and
  3) choose the proper strategies for its control
Acute viral hepatitis (continued)

Special aspects

Surveillance data of acute viral hepatitis from developing countries should be interpreted with caution. Differentiation of types of viral hepatitis (A to E) based on clinical diagnosis is unreliable and serologic testing is necessary for accurate diagnosis. Unfortunately, many developing countries do not have access to diagnostic reagents. Most infections with hepatitis A, B, C and E virus occur asymptptomatically (in developing countries usually among children) and will not be detected and reported to the surveillance system. Therefore, a low incidence of acute viral hepatitis should not be misinterpreted as a low prevalence of viral hepatitis infection.

Understanding the epidemiology and burden of disease of viral hepatitis requires an understanding of the sequelae of hepatitis B, C and D infection. These includes asymptomatic chronic infection, chronic hepatitis, cirrhosis, and primary liver cancer. Measuring the burden of these conditions requires data collection from sources not traditionally used by infectious disease epidemiologists, including data on hospital discharge and mortality data (for chronic hepatitis, cirrhosis, and liver cancer), and cancer registers. Special sero-prevalence surveys may be needed to measure the prevalence of hepatitis B and hepatitis C infection in the general population and in special groups such as blood donors, pregnant women, military recruits, health care workers, certain patient groups (e.g. patients with liver disease, people on dialysis, haemophiliacs), and ethnic sub-populations.

Assessing coverage of hepatitis B vaccine is similar to that for other EPI vaccines. Vaccine is given to infants (and in some industrial countries to adolescents) primarily to prevent the development of chronic liver disease and liver cancer; serological testing to document sero-conversion in children is usually not necessary because numerous studies have shown that the vaccine is 85% to 100% effective in preventing chronic infection.

Contact information

Regional Offices and Headquarters (VAB and CDS)
See Annex 1 for addresses and fax numbers.
### Rationale for surveillance

Hib is the most common cause of bacterial meningitis in children, and one of the two most common causes of severe bacterial pneumonia. Pneumonia is the largest single remaining infectious disease killer of young children in the developing world. Hib may also cause other diseases, including arthritis, skin infection, and epiglottitis. Surveillance data are critical for clarifying the burden of disease and evaluating the impact of immunization programmes. Although in many countries Hib pneumonia is more common than the other types of infection, diagnosis of Hib pneumonia is extremely difficult. Routine surveillance should focus on meningitis and other Hib infections, diagnosed with microbiologic tests on blood, cerebrospinal fluid (CSF), and other body fluids (such as pleural fluid) that usually do not contain bacteria. Such infections are often called “invasive Hib disease”. Countries may also wish to report potential cases of bacterial meningitis, both as a performance indicator for Hib detection, and to clarify the burden of meningitis attributable to all bacteria.

### Recommended case definition

#### Clinical description

Bacterial meningitis is characterized by acute onset of fever, headache and stiff neck. Meningitis is not specific for Hib disease, and Hib disease cannot be diagnosed on clinical grounds.

#### Laboratory criteria for diagnosis

Culture: isolation of Hib from a normally sterile clinical specimen, such as cerebrospinal fluid (CSF) or blood (i.e. culture of Hib from a non-sterile site, such as the throat, does not define Hib disease, since the bacteria can grow in these other areas and not cause disease). Antigen detection methods: identification of Hib antigen in normally sterile fluids (i.e. CSF or blood) by antigen detection methods such as latex agglutination or counter immunoelectrophoresis (CIE).

#### Case classification

**Potential:** Bacterial meningitis case: a child with a clinical syndrome consistent with bacterial meningitis

**Probable:** Not applicable

**Confirmed:** A case that is laboratory confirmed by growing or identifying Hib in the CSF or blood

**Note:** Any person with Hib isolated from CSF or blood may be reported as a confirmed case, regardless of whether their clinical syndrome was meningitis.
Haemophilus influenzae type B (Hib) disease (continued)

### Recommended types of surveillance
- Routine monthly reporting of aggregate data of confirmed cases is recommended from peripheral level to intermediate and central levels
- Zero reporting should be required at all levels
- All potential cases should also be reported if laboratory performance indicators are to be monitored (see Note)

**Note:** Since laboratory confirmation is required for all cases, the extent of surveillance will of necessity vary depending on the capabilities of individual countries. Surveillance does not need to be national in scope to fulfil goals as noted in “Rationale” section above. It is more important to have a well-functioning system in some areas than to have a national system that functions poorly.

### Recommended minimum data elements

**Aggregated data for reporting**
- Number of cases
- Number of 3rd doses of Hib vaccine (Hib3) administered to infants

**Case-based data for reporting and investigation**
- Unique identifier
- Geographical area (e.g. district and province) names
- Date of birth
- Date of onset
- Specimen type, if specimen collected:
  1=blood; 2=CSF; 3=both; 4=other
- Culture result, if done:
  1=positive; 2=negative; 3=pending; 4=not done
- Antigen detection result, if done:
  1=positive; 2=negative; 3=pending; 4=not done
- CSF white cell count/ml, if done
- Outcome:
  1=alive; 2=dead; 9=unknown
- Number of Hib doses received:
  9=unknown
- Final classification:
  1=potential; 2=confirmed

### Recommended data analyses, presentation, reports

**Aggregated data**
- Incidence rate by year and geographic area
- Hib3 coverage by year and geographic area
- Completeness and timeliness of reporting
Haemophilus influenzae type B (Hib) disease (continued)

Recommended data analyses, presentation, reports (continued)

Case-based data: Same as aggregate plus:
- Age-specific incidence rate
- Case fatality rate
- Cases by immunization status

- Performance indicators of surveillance quality
  - Percent of all potential bacterial meningitis cases for which CSF/blood was obtained for evaluation ≥ 90%
  - Percent of potential bacterial meningitis cases in which a bacterial pathogen was identified from CSF or blood:
    - Among CSF with 10 or more white blood cells/ml ≥ 20%
    - Among CSF with 100 or more white blood cells/ml ≥ 50%

Note: Although persons with bacterial meningitis have a wide range of CSF white blood cell counts, the proportion of potential bacterial meningitis cases with identifiable bacterial causes increases with increasing CSF cell counts. For evaluation of performance, programme personnel may wish to determine proportion of potential bacterial meningitis cases in which bacterial causes have been identified in one or both of the above categories. Results below the target levels suggest some cases of bacterial meningitis are not being identified, and that review of laboratory and clinical practices should be performed.

Principle uses of data for decision-making
- To determine incidence of Hib meningitis and invasive disease for estimation of Hib disease burden
- To measure impact of immunization program and identify areas needing additional input
- To monitor coverage and take action to correct low coverage areas

Special aspects

Since Hib surveillance requires laboratory confirmation, nation-wide surveillance may not be practical in many countries. However, most surveillance goals may be approached with a less comprehensive plan. Surveillance in areas with appropriate clinical and laboratory capacity can provide necessary information on burden and immunization impact. Coverage data should be obtained nation-wide. Evaluating the combination of nation-wide coverage data, and area-specific disease data can provide necessary information for making immunization programme decisions. Additional guidance on surveillance methodology can be obtained in WHO publication WHO/VRD/GEN/95.05.

Contact information

Regional Offices and Headquarters (VAB)
See Annex 1 for addresses and fax numbers.
Measles

Rationale for surveillance

Measles is targeted for a reduction by 90% in incidence and by 95% in mortality (9GPW 6.2). Surveillance for measles is important to monitor/adjust strategies and should evolve based on level of control. Countries in the “control” phase are endemic and should concentrate on raising routine measles immunization coverage and focusing extra immunization efforts in areas with high measles mortality. Countries in the “measles elimination phase” are achieving high routine measles coverage and low incidence with periodic outbreaks. Surveillance in these countries should be used to identify high risk populations and to predict and prevent potential outbreaks. Those countries in which the objective is to completely interrupt measles transmission require very intensive case-based surveillance to detect, investigate, and confirm every suspect measles case in the community.

Recommended case definition

Clinical case definition
Any person in whom a clinician suspects measles infection or

Any person with fever, and maculopapular rash (i.e. non-vesicular), and cough, coryza (i.e. runny nose) or conjunctivitis (i.e. red eyes)

Laboratory criteria for diagnosis
Presence of measles-specific IgM antibodies

Case classification

<table>
<thead>
<tr>
<th>Clinically confirmed*</th>
<th>A case that meets the clinical case definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory confirmed**</td>
<td>A case that meets the clinical case definition and that is laboratory confirmed***</td>
</tr>
<tr>
<td>Epidemiologically confirmed**</td>
<td>A case that meets the clinical case definition and is linked epidemiologically to a laboratory confirmed case</td>
</tr>
<tr>
<td>Discarded</td>
<td>A suspect case that does not meet the clinical or laboratory definition</td>
</tr>
</tbody>
</table>

* During the “control” phase, cases are confirmed (and classified) on clinical grounds.
** Laboratory confirmation is used during the “elimination phase” and to confirm an outbreak.
*** Lab confirmation is used during the “elimination phase” and to confirm an outbreak. If the case has been vaccinated within 6 weeks prior to serum collection refer to the Manual for the Laboratory Diagnosis of Measles Virus Infection for classification.
**Recommended types of surveillance**

- **Control phase**: When measles is endemic, routine monthly reporting of aggregated data of clinical measles cases is recommended by district, age group and immunization status. In order not to overburden the system, it is recommended that only outbreaks (not each case) be investigated to determine their causes.*

- **Elimination phase**: Case-based surveillance should be conducted and every case reported and investigated immediately (and also included in the weekly reporting system). Laboratory specimens should be collected from every suspect case. Suspected measles outbreaks should be confirmed by conducting serology on the first 5-10 cases only. Urine specimens (for virus isolation and genetic characterisation) should be collected from sporadic/outbreak cases (10 cases from each chain of transmission) to characterize viral circulation and importation patterns. All outbreaks should be thoroughly investigated.

- **During both phases**: zero reporting should be required at all levels.
  - Laboratory confirmation may be attempted in control phase with sampling ~10 cases per outbreak

**Recommended minimum data elements**

**Control phase (aggregated data):**

- Number of cases by age group and immunization status, month and geographic area
- Number of measles vaccine doses administered to infants (< 12 months) and children 12-23 months
- Target population

**Elimination phase (case-based data):** in addition to data from the control phase

- Unique identifier
- Geographical area (e.g. district and province)
- Date of birth
- Date of rash onset
- Number of prior measles vaccine doses received:
  - 9=unknown
- Date of receipt of last dose
- Date of notification
- Date of case investigation
- Date of blood specimen collection
- Date blood specimen sent to laboratory
- Date blood specimen received by the laboratory
- Condition of blood specimen on receipt
  - 1=adequate; 2=inadequate; 9=unknown
- Date measles serology results reported
- Results of measles serology:
  - 1=positive; 2=negative; 3=indeterminate; 4=no specimens processed; 9=unknown
Measles (continued)

- Results of differential serology (make separate variable for each disease):
  1=positive; 2=negative; 3=indeterminate; 9=unknown
- Collection of specimen for viral culture/identification
  1=yes; 2=no
- Specimen type
  1=urine; 2=respiratory; 3=lymphocytes
- Date specimen received for viral culture/identification
- Results of measles viral culture/identification
  1=positive; 2=negative; 9=unknown
- Final classification:
  1=clinically confirmed; 2=laboratory confirmed; 3=epidemiologically-linked to laboratory confirmed case; 9=discarded
- Source of infection identified:
  1=yes; 2=no; 9=unknown and nature of source
- Outcome

Note: In every phase, completeness and timeliness of monthly (control phase) or weekly (elimination phase) measles reporting should be monitored. To avoid successive changes in forms and other data collection instruments, countries likely to move soon to the elimination phase may wish to move to case-based data while still in the control phase is this is not too burdensome.

Recommended data analyses, presentation, reports

Control phase
- Incidence rate by month, year, and geographic area
- Measles vaccine coverage by year and geographic area
- DTP1-measles or BCG-measles drop out rate
- Completeness/timeliness of monthly reporting
- Proportion of known outbreaks confirmed by the laboratory
- Age-specific incidence rate
- Proportion of cases by age group and immunization status: core age groups suggested 0-8 months, 9-11 months, 2-4 years, 5-9 years, 10-14 years, 15-19 years, 20-24 years, 25 years and over.
Measles (continued)

Measles elimination phase: same as control phase plus the following

Performance indicators                               Target
- % of weekly reports received                        ≥ 80%
- % of cases* notified ≤ 48 hours of rash onset       ≥ 80%
- % of cases* investigated ≤ 48 hours of notification ≥ 80%
- % of cases* with adequate specimen** and lab results within 7 days ≥ 80%
- % of confirmed cases with source of infection identified ≥ 80%
- % of outbreaks investigated                        ≥ 80%

* All cases that meet the clinical case definition
** Adequate specimen is one blood specimen collected within 28 days of rash onset

Principal uses of data for decision-making

- Control phase: Monitor incidence and coverage to determine progress (i.e. decreasing incidence and increasing coverage), and to identify areas with poor programme performance. Describe the changing epidemiology of measles in terms of age, immunization status and inter-epidemic period.
- Elimination phase: Identify chains of transmission. Monitor the epidemiology (age groups at risk, inter-epidemic period, immunization status) of measles and determine when the next outbreak may occur due to a build up of susceptibles and accelerate immunization activities accordingly to avert a potential outbreak. Determine where measles virus is circulating or may circulate.
- Use epidemiologic data to classify cases (See special aspects section).
- During all phases: Detect and investigate outbreaks to ensure proper case management, and determine why the outbreak occurred (e.g. failure to vaccinate, vaccine failure or inadequate strategies)

Special aspects

Use additional methods such as epidemiologic modelling and sero-epidemiology studies to monitor the build-up of susceptibles (guidelines on how to do this are under development)

Also see "manual for laboratory diagnosis of measles virus infection" and "guidelines on measles outbreaks"
Measles (continued)

Final classification of measles cases

- Adequate blood specimen
  - IgM negative → Discard
  - IgM positive → Laboratory confirmed
- Suspect measles cases
- No adequate blood specimen
- Epidemiologic link to laboratory confirmed case
  - Epidemiologically confirmed
- No epidemiologic link to laboratory confirmed case
  - Clinically confirmed

Opportunities should be seized to integrate measles and rubella surveillance depending on control strategies and immunization programmes in place

Contact information

Regional Offices
See Annex 1

Headquarters
WHO Department of Vaccines and Biologicals
(See Annex 1 for address and fax number)
E-mail: EPIdata@who.int
Tel: +41 22 791 4359
Neonatal tetanus

<table>
<thead>
<tr>
<th>Rationale for surveillance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonatal tetanus is targeted by WHO for elimination as a major public health burden (9GPW 6.2). High tetanus toxoid (TT) coverage of pregnant women, clean delivery and the identification of, and implementation of corrective action in high risk areas (i.e. TT immunization of childbearing-aged women) are the three primary strategies towards this goal. Epidemiological surveillance is useful in the identification of areas at high risk for neonatal tetanus (NT) and for monitoring impact of interventions.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Recommended case definition</th>
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</thead>
<tbody>
<tr>
<td><strong>Clinical case definition and case classification</strong></td>
</tr>
<tr>
<td><strong>Suspected case:</strong> Any neonatal death between 3-28 days of age in which the cause of death is unknown; or any neonate reported as having suffered from neonatal tetanus between 3-28 days of age and not investigated</td>
</tr>
<tr>
<td><strong>Confirmed case:</strong> Any neonate with a normal ability to suck and cry during the first two days of life, and who between three and 28 days of age cannot suck normally, and becomes stiff or has convulsions (i.e. jerking of the muscles) or both</td>
</tr>
<tr>
<td><strong>Note:</strong> The basis for case classification is entirely clinical and does not depend upon laboratory confirmation. NT cases reported from hospitals are considered confirmed.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Recommended types of surveillance</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Number of confirmed NT cases should be included in routine monthly surveillance reports of all countries and should be reported as a separate item from other (non-neonatal) tetanus. Zero reporting should be required at all levels</td>
</tr>
<tr>
<td>- Active surveillance for NT should be conducted in major health facilities on a regular basis</td>
</tr>
<tr>
<td>- A retrospective record review for NT cases should be conducted at least once annually in major hospitals</td>
</tr>
<tr>
<td>- In “low risk” geographical areas where NT incidence&lt;1/1000 live births and surveillance is performing well (i.e. surveillance data are reasonably representative of the population and there is good reporting completeness), all suspect cases should be investigated to confirm and identify the cause</td>
</tr>
<tr>
<td>- Community surveillance is recommended in “silent” areas (i.e. where routine reporting is not functional but, based on other indicators, where neonatal tetanus could be a problem)</td>
</tr>
</tbody>
</table>
**Recommended minimum data elements**

**Aggregated data:**
- Number of cases
- Doses of TT administered to pregnant or child-bearing aged women (depending on national policy) or % of newborns protected at birth (PAB) [see special aspects section]
- Completeness/timeliness of monthly reports

**Case-based data:**
- Unique identifier
- Geographical area (e.g. district and province) names
- Date of birth of baby
- Age (in days) of baby at onset
- Sex of baby
- Parity (number of deliveries including this most recent one) of mother
- Date of case investigation
- Location/type of birth:
  - 1= institution; 2= home with trained attendent; 3= home with untrained attendent; 4= home without attendent; 5= other; 9= unknown
- Tetanus immunization status of mother when she gave birth:
  - 1= up-to-date; 2= not up-to-date; 3= unimmunised; 9= unknown
- Final classification:
  - 1= confirmed; 2= suspected; 3= discarded
- Mother given protective TT dose within three months of report:
  - 1= yes; 2= no; 9= unknown
- Supplemental immunization conducted within same locality as the case:
  - 1= yes; 2= no; 9= unknown

**Recommended data analyses, presentation, reports**

**Aggregated data (i.e. routine monthly reporting)**
- Incidence rate per 1000 live births by geographic area, month, and year
- TT2+ (or PAB) by year and geographic area
- Completeness/timeliness of monthly reporting
- Geographic areas considered at high risk for NT compared to those where corrective actions were taken

**Case-based data (i.e. from case investigations only)** same as for aggregated data plus the following:
- Confirmed NT cases by delivery type, sex, TT2+ status of the mother
- % of confirmed cases for which the mother subsequently received a protective TT dose
Neonatal tetanus (continued)

Principle uses of data for decision-making/action

- Monitor progress towards achieving and sustaining high routine TT2+ (or PAB) coverage in all geographical areas
- Monitor progress towards eliminating NT in every geographical area
- Investigate suspect NT cases in areas not considered at risk for NT to confirm and determine cause
- Identify high risk geographical areas
- Monitor whether corrective actions were taken in those areas considered at high risk
- Periodically validate sensitivity of NT reporting by comparing number of reported cases with cases identified through active surveillance

Special aspects

"% protected at birth" (PAB) is an alternative method of determining coverage (particularly where TT2+ is unreliable). To monitor PAB, health workers record during DTP1 visits whether the infant was protected at birth by the mother’s TT status and/or delivery status (clean/unclean). % PAB is then estimated as: number of infants protected/number of live births. If the child was un protected, the mother should receive a dose of tetanus toxoid during the same visit and should be followed up with a subsequent TT dose if needed for protection.

Contact information

Regional Offices and Headquarters (VAB)
See Annex 1 for addresses and fax numbers.
## Pertussis (whooping cough)

### Rationale for surveillance

Pertussis is a major cause of childhood morbidity and mortality. An estimated 45 million cases and 400,000 deaths occur annually. Case fatality rates in developing countries can reach 15%. High routine coverage with effective vaccine is the mainstay of prevention. Surveillance data on the disease can monitor the impact of immunization programmes as well as identify high-risk areas and outbreaks.

### Recommended case definition:

**Clinical description**

A person with a cough lasting at least two weeks with at least one of the following:

1. Paroxysms (i.e. fits) of coughing
2. Inspiratory “whooping”
3. Post-tussive vomiting (i.e. vomiting immediately after coughing) and without other apparent cause

**Laboratory criteria for diagnosis**

Isolation of *Bordetella pertussis* or detection of genomic sequences by polymerase chain reaction (PCR)

**Case classification**

- **Suspected**: A case that meets the clinical description
- **Confirmed**: A person with a cough that is laboratory confirmed

### Recommended types of surveillance

- Routine monthly reporting of aggregated data of suspected and confirmed cases from peripheral level to intermediate and central levels. Zero reporting should be required at all levels
- All outbreaks should be investigated immediately and laboratory confirmed. During an outbreak, case-based data should be collected
- To describe the changing pertussis epidemiology in countries with low pertussis incidence (usually where coverage is >80%), additional information on age group and immunization status should be collected; or, as an alternative case-based surveillance, sentinel surveillance, active surveillance, and/or occasional surveys and laboratory confirmation of suspect cases should be considered

### Recommended minimum data elements

**Aggregated data:**

- Number of cases
- Number of third doses of diphtheria-pertussis-tetanus vaccine (DTP3) administered to infants
- Completeness/timeliness of monthly reports
**Pertussis (whooping cough) - (continued)**

### Recommended minimum data elements (continued)

**Case-based data:**
- Unique identifier
- Geographical area (e.g. district and province) names
- Date of birth
- Date of onset
- Total pertussis vaccine doses:
  - 99=unknown
- Date of last pertussis vaccine dose:
- Outcome:
  - 1=alive; 2=dead; 9=unknown
- Classification:
  - 1=confirmed; 2=suspect; 3=discarded

### Recommended data analyses, presentation, reports

**Aggregated data:**
- Incidence rate by month, year, and geographic area
- DTP3 coverage by year and geographic area
- Completeness/timeliness of monthly reporting
- Proportional morbidity (compared to other diseases of public health importance)

**Case-based data:** same as aggregated data plus the following:
- Age-specific incidence rate
- Immunization status of cases
- Case fatality rate
- Proportional mortality (compared to other diseases of public health importance)

### Principle uses of data for decision-making

- Investigate outbreaks to understand epidemiology of pertussis in the country, why the outbreak occurred (e.g. failure to immunize, vaccine failure, accumulation of susceptibles/waning immunity), and to ensure proper case management
- Monitor case fatality rate. If high, determine cause (e.g. poor case management, lack of antibiotics/supportive care, patients not seeking treatment in time)
- Determine age-specific incidence rate, and incidence rate by geographical area to know risk groups/areas
- Monitor incidence rate to assess impact of control efforts

### Contact information

**Regional Offices and Headquarters (VAB)**
See Annex 1 for addresses and fax numbers.
## Poliomyelitis

### Rationale for surveillance

Poliomyelitis is targeted for eradication (9GPW 6.1). Highly sensitive surveillance for acute flaccid paralysis (AFP), including immediate case investigation, and specimen collection is critical to detect wild poliovirus circulation with the ultimate objective of polio eradication. AFP surveillance is also critical for documenting the absence of poliovirus circulation for polio-free certification.

### Recommended case definition

#### Clinical case definition

Any child under fifteen years of age with acute, flaccid paralysis\(^1\) or any person with paralytic illness at any age when polio is suspected.

#### Case classification

- **Suspected case:** A case that meets the clinical case definition
- **Confirmed case:** See diagram in special aspects section

\(^1\) Including Guillain Barré syndrome

### Recommended types of surveillance

- Aggregated data of AFP cases should be included in routine monthly surveillance reports
- Zero reporting should be required at all levels
- All outbreaks should be investigated immediately
- All AFP cases under 15 years of age or with paralytic illness at an age where polio is suspected, should be reported immediately, investigated within 48 hours and two stool specimens collected 24-48 hours apart and within 14 days of paralysis onset
- Active surveillance should be implemented in selected hospitals
**Poliomyelitis (continued)**

### Recommended minimum data elements

**Aggregated data:**
- Number of third doses of oral polio vaccine (OPV3) administered to infants
- Number of AFP cases

**Case-based data** (to be linked to specimen-based data for analysis):
- Unique identifier
- Geographical area (e.g. district & province) names
- Date of birth
- Date of paralysis
- Date of notification
- Date of case investigation
- Total polio vaccine doses received:
  - 99=unknown
- Fever at onset of paralysis:
  - 1=yes; 2=no; 9=unknown
- Progression of paralysis within four days:
  - 1=yes; 2=no; 9=unknown
- Asymmetric paralysis:
  - 1=yes; 2=no; 9=unknown
- Date of 60-day follow-up exam
- Findings at 60-day follow-up:
  - 1=residual weakness; 2=no residual weakness; 3=lost to follow-up; 4=death before follow-up
- Final classification:
  - 1=confirmed; 2=compatible; 3=discarded

**Specimen-based data** (to be linked to case-based data for analysis):
- Unique identifier
- Specimen number:
  - 1=first specimen; 2=second specimen; 3=other; 9=unknown
- Date of paralysis onset
- Date of last OPV
- Date of stool specimen collection
- Date stool specimen sent to lab
- Date stool specimen received in lab
- Condition of stool:
  - 1=good; 2=poor; 9=unknown
- Date final culture results sent from lab to EPI
- Date intratypic differentiation results sent from lab to EPI
### Poliomyelitis (continued)

#### Recommended minimum data elements (continued)

- **Results**
  - Polio type 1 isolated?
    - 1=yes, wild; 2=yes, Sabin; 3=yes, pending intratypic differentiation; 4=yes, mixture of wild & Sabin; 5=no P1 isolated; 6=specimen not processed
  - Polio type 2 isolated?
    - 1=yes, wild; 2=yes, Sabin; 3=yes, pending intratypic differentiation; 4=yes, mixture of wild & Sabin; 5=no P2 isolated; 6=specimen not processed
  - Polio type 3 isolated?
    - 1=yes, wild; 2=yes, Sabin; 3=yes, pending intratypic differentiation; 4=yes, mixture of wild & Sabin; 5=no P3 isolated; 6=specimen not processed
  - Non-polio enterovirus (NPEV) isolated?
    - 1=yes; 2=no NPEV isolated; 3=specimen not processed

#### Recommended data analyses, presentation, reports

**Aggregated data:**
- Cases by month, year, and geographic area
- OPV3 coverage by year and geographic area
- Completeness/timeliness of monthly reporting

**Case-based data:** same as aggregated data plus the following:
- Confirmed cases by age group, immunization status, geographic area, month and year
- Confirmed cases from which wild poliovirus was isolated by geographic area, month, year
- Compatible cases by geographic area and month
- All suspect cases by final classification
- Non-polio enterovirus isolation rate

**Indicators of surveillance performance**

- % of all expected monthly reports that were received
- Annualized non-polio AFP rate per 100 000 children under 15 years of age
- % of AFP cases investigated within 48 hours
- % of AFP cases with two adequate stool specimens collected 24-48 hours apart and ≤ 14 days of onset
- % of specimens arriving at the laboratory in "good" condition
- % of specimens arriving at a WHO-accredited laboratory within 3 days of being sent
- % of specimens with laboratory results sent within 28 days of specimen receipt

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>% of all expected monthly reports that were received</td>
<td>≥ 90%</td>
</tr>
<tr>
<td>Annualized non-polio AFP rate per 100 000 children under 15 years of age</td>
<td>≥ 1/100 000</td>
</tr>
<tr>
<td>% of AFP cases investigated within 48 hours</td>
<td>≥ 80%</td>
</tr>
<tr>
<td>% of AFP cases with two adequate stool specimens collected 24-48 hours</td>
<td>≥ 80%</td>
</tr>
<tr>
<td>% of specimens arriving at the laboratory in &quot;good&quot; condition</td>
<td>≥ 80%</td>
</tr>
<tr>
<td>% of specimens arriving at a WHO-accredited laboratory within 3 days</td>
<td>≥ 80%</td>
</tr>
<tr>
<td>% of specimens with laboratory results sent within 28 days of specimen</td>
<td>≥ 80%</td>
</tr>
</tbody>
</table>
Poliomyelitis (continued)

Principle uses of data for decision-making

- Track wild poliovirus circulation
- Use data for classifying cases as confirmed, polio compatible or discarded (see special aspects section)
- Monitor routine coverage in all geographical areas and focus efforts in low performing geographical areas
- Monitor seasonality to determine low season of poliovirus transmission for National Immunization Day (NIDs) planning
- Identify high risk areas for planning mopping up immunization
- Monitor performance of surveillance using standard indicators listed above and focus efforts in low performing geographical areas
- Provide evidence to Certification Commissions of the interruption of wild poliovirus circulation
Poliomyelitis (continued)

Special aspects

The scheme in the following illustration (Figure 1) should be used to classify AFP cases. A country should use the clinical classification until their surveillance performance meets the following three criteria: 1) a non-polio AFP rate of at least 1/100,000 children under 15 years of age; 2) two adequate specimens\(^1\) collected from at least 60% of detected AFP cases; 3) all specimens processed in a WHO-accredited laboratory.

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**Figure 2a: Clinical criteria**
(early stages of polio eradication)

- Wild poliovirus → confirm
- residual weakness, died or lost to follow-up → confirm
- inadequate specimens → confirm
- No wild poliovirus
  - inadequate specimens → discard
  - two adequate specimens → discard

**Figure 2b: Virologic criteria**
(advanced stages)

- confirm
- compatible\(^2\)
- National Expert Committee Review
- discard

---

\(^1\) "Adequate specimens" means two specimens collected 24-48 hours apart and within 14 days of onset of paralysis. The specimen arriving at the laboratory must be of adequate volume (approximately 8-10 grams), have appropriate documentation (i.e. laboratory request form) and be in "good condition". "Good condition" = no leakage, no desiccation, and evidence that the reverse cold chain was maintained (based on presence of ice or temperature indicator).

\(^2\) "Compatible" cases indicate surveillance failures and should be monitored for clustering in space and time.

Contact information

Regional Offices and Headquarters (VAB)
See Annex 1 for addresses and fax numbers.
# Rubella and Congenital Rubella Syndrome

## Rationale for surveillance

Rubella is a mild illness, but rubella in pregnancy can lead to congenital rubella syndrome (CRS) in the infant. The birth defects associated with CRS include heart disease, blindness, deafness, and mental retardation. By 1999, more than 100 countries had introduced rubella vaccine into their national immunization programme. Additional countries, especially those in the elimination phase of measles control, are considering the introduction of rubella vaccine. All countries that include rubella vaccine in their immunization programme should conduct surveillance for CRS and rubella. In the CRS prevention stage, disease surveillance should focus on detecting cases of congenital rubella syndrome (CRS). In the CRS/rubella elimination phase (usually conducted in conjunction with measles elimination), case-based surveillance of febrile rash illness is necessary.

## Recommended case definitions

Case definitions for CRS and rubella are provisional, pending field-testing.

### Congenital rubella syndrome (CRS)

**Suspected CRS case:** Any infant less than one year of age in whom a health worker suspects CRS. A health worker should suspect CRS when an infant (0-11 months of age) presents with heart disease and/or suspicion of deafness and/or one or more of the following eye signs: cataract, diminished vision, nystagmus, squint, microphthalmus, or congenital glaucoma. A health worker should also suspect CRS when an infant’s mother has a history of suspected or confirmed rubella during pregnancy, even when the infant shows no signs of CRS.

**Clinically-confirmed CRS case:** An infant in whom a qualified physician detects two of the complications listed in (a) below or one in (a) and one in (b):

(a) Cataract(s), congenital glaucoma, congenital heart disease, loss of hearing, pigmented retinopathy

(b) Purpura, splenomegaly, microcephaly, mental retardation, meningoencephalitis, radiolucent bone disease, jaundice the onset of which is within 24 hours after birth

**Laboratory-confirmed CRS case:** An infant with clinically-confirmed CRS who has a positive blood test for rubella-specific IgM; (100% of such infants will be positive at age 0-5 months; 60% at 6-11 months). Where special laboratory resources are available, detection of rubella virus in specimens from the pharynx or urine of an infant with suspected CRS provides laboratory confirmation of CRS (60% of such infants shed rubella virus at age 1-4 months; 30% at 5-8 months; 10% at 9-11 months).

**Congenital rubella infection (CRI):** When a mother has suspected or confirmed rubella in pregnancy, her infant should have a rubella-specific IgM blood test. An infant who does not have clinical signs of CRS but who has a positive rubella-specific IgM test is classi-
**Rubella and Congenital Rubella Syndrome (continued)**

**Recommended case definitions (continued)**

**Rubella**

*Suspected rubella case*: Any patient of any age in whom a health worker suspects rubella. A health worker should suspect rubella when a patient presents with: fever, maculopapular rash; and cervical, sub-occipital, or post-auricular adenopathy or arthralgia/arthritis.

*Clinical confirmation*: Rubella cannot be confirmed clinically: laboratory confirmation is required.

*Laboratory-confirmed rubella case*: Because of the difficulty of clinical diagnosis of rubella, laboratory confirmation is required. A laboratory-confirmed case is a suspected case with a positive blood test for rubella-specific IgM. The blood specimen should be obtained within 28 days of rash onset.

*Epidemiologically-confirmed rubella case*: A patient with a febrile rash illness that is linked epidemiologically to a laboratory-confirmed rubella case.

**Recommended types of surveillance**

**CRS prevention stage** – minimum requirements
- Routine monthly reporting of number of suspected CRS cases; zero reporting should be required. All suspected CRS cases in infants under one year of age should be investigated, including clinical and laboratory analysis.
- Routine monthly reporting of number of suspected rubella cases; zero reporting should be required.
- All febrile rash illnesses in pregnant women should be investigated.
- When a rubella outbreak is detected, a limited number of suspected rubella cases should be investigated with laboratory tests periodically during the outbreak (perhaps 5 to 10 cases investigated per month). Active surveillance should be initiated to detect suspected CRS in infants under one year of age and continued for nine months after the last reported case of rubella.

**CRS/rubella elimination stage** – minimum requirements
- Same as CRS prevention stage, plus
- Routine monthly reporting of number of confirmed rubella cases; zero reporting should be required.
- All febrile rash cases regardless of age must be investigated, including laboratory analysis of each case for measles and, if the result is negative, for rubella. Febrile rash illnesses in pregnant women should be given priority in investigation.
### Rubella and Congenital Rubella Syndrome (continued)

**Recommended minimum data elements**

**Routine surveillance data**
- Number of suspected CRS cases in each health facility, by month.
- For countries with febrile rash illness surveillance, number of febrile rash illnesses reported in each health facility, by month.
- For other countries, number of suspected rubella cases in each health facility, by month.

**Routine immunization coverage data**
- Rubella vaccine coverage (%) for each target group (women of childbearing age, schoolgirls, infants, pre-school children) in each health facility, by month.

**CRS case investigation data**
- Clinical signs and symptoms
- Patient's date of birth
- Date of notification
- Date of investigation
- Date of blood specimen collection
- Results of rubella-specific IgM test
- Mother's history of febrile rash illness or exposure to febrile rash illness during this pregnancy
- Mother's address(es) during this pregnancy
- Mother's history of rubella immunization
- Number of mother's previous pregnancies

**Rubella case investigation data (elimination phase)**
- Clinical signs and symptoms
- Patient's date of birth or age
- Date of rash onset
- Date of investigation
- Date of blood specimen collection
- Results of rubella-specific IgM test
- Address or addresses during the 3 weeks before onset febrile rash illness
- History of rubella and measles immunizations
Rubella and Congenital Rubella Syndrome (continued)

Recommended data analyses, presentation, reports

Aggregated data (i.e. from routine reporting)
- CRS incidence - Number of CRS cases per month and number of CRS cases per 1000 live births per year, by geographic area
- Rubella incidence - Number of rubella cases per month, by geographic area
- Rubella vaccine coverage by target group and geographic area per year

Case-based data (i.e. from case investigations only)
- Final classification of all suspected cases of CRS and rubella
- Immunization status of mothers of cases of CRS
- Proportion of all cases of febrile rash illness with laboratory investigation that are rubella-specific IgM positive

Surveillance quality
- Completeness and timeliness of routine reporting, notification, and clinical and laboratory investigation

Principle uses of data for decision making/action
- Understand the epidemiology of CRS and its burden in the population to guide rubella immunization strategies. Because children with CRS may be blind, deaf, retarded, or have major heart disease, CRS creates a long term burden on health care, social, and education systems.
- Use rubella outbreak investigation as a tool for activating CRS surveillance
- Investigate rash illness in pregnancy and provide culturally appropriate follow-up to women who have rubella, including follow-up of their infants
- Investigation of febrile rash illnesses in the measles/rubella elimination phase to determine the proportion of such illnesses due to rubella to identify high risk areas, age groups and/or populations.

Special aspects
- Febrile rash illness surveillance should link surveillance for rubella, measles, and dengue (where applicable).
- Outbreaks of rubella and measles have occurred simultaneously.
- CRS cases are likely to be under-reported in areas and among populations where a high proportion of births occur at home and where infant deaths may not be reported.
- Infants with CRS are likely to be seen at specialty facilities that do not normally participate in the immunization programme or the routine communicable disease surveillance system, for example, eye hospitals and hospitals specializing in cardiac surgery. For comprehensive CRS surveillance, these facilities should be included in CRS detection, investigation, and reporting activities.
- Infants with CRS and CRI shed rubella virus for long periods (60% for the first 4 months of life) and appropriate infection control measures should be applied. It is particularly important that pregnant women who are not rubella-immune are not exposed to infants with CRS or CRI.
Special aspects (continued)

- Serological monitoring of rubella susceptibility of women attending selected antenatal clinics can be used to monitor rubella immunization programme performance. However, serological monitoring requires a different laboratory test, e.g. rubella-specific IgG. If serological screening is conducted, arrangements should be made to provide postpartum rubella vaccination to women found to be seronegative.

Contact information

Regional Offices and Headquarters (VAB and CDS)
See Annex 1 for addresses and fax numbers.
## Yellow fever

### Rationale for surveillance
This mosquito-borne virus disease occurs in tropical regions of Africa and South America and is maintained by sylvatic transmission of virus involving forest-dwelling mosquitoes and monkeys. Transmission to humans may occur in forest transition zones and subsequently may enter an urban cycle through the *Aedes aegypti* mosquito. Many cities are now threatened with major epidemics as yellow fever is undergoing a major resurgence especially in the African region. The strategies for yellow fever control are: control of *Aedes aegypti* in urban centres, infant immunization, immunization campaigns to prevent epidemics, epidemic detection and emergency immunization when an epidemic is confirmed. Surveillance data allows for monitoring disease incidence, the prediction and early detection of outbreaks and the monitoring of control measures. Case reporting of yellow fever is universally required by International Health Regulations.

### Recommended case definition
**Clinical description**
An illness characterised by acute onset of fever followed by jaundice within two weeks of onset of first symptoms **AND** one of the following: 1) bleeding from nose, gum, skin, or GI tract; or 2) death within 3 weeks of illness onset.

**Laboratory criteria for diagnosis**
Isolation of yellow fever virus, or presence of yellow fever specific IgM or a four-fold or greater rise in serum IgG levels (acute or convalescent) or positive post-mortem liver histopathology or detection of yellow fever antigen in tissues by immunohistochemistry or detection of yellow fever virus genomic sequences in blood or organs by PCR.

**Case classification**
- **Suspected:** A case that is compatible with the clinical description
- **Probable:** Not applicable
- **Confirmed:** A suspected case that is laboratory confirmed or epidemiologically linked to a laboratory-confirmed case or outbreak

### Recommended types of surveillance
- Routine weekly/monthly reporting of aggregated data on suspected and confirmed cases from peripheral level to intermediate and central levels. Zero reporting should be required at all levels.
- Immediate reporting of suspected cases from peripheral level to intermediate and central levels.
- All suspected cases and outbreaks should be investigated immediately and laboratory confirmed.
- Case-based surveillance should be implemented in countries identified by WHO as high risk for yellow fever. Specimens should be collected to confirm an epidemic as rapidly as possible. Then priority should be placed on collecting specimens from new or neighbouring areas (other than the area where the epidemic is already confirmed).
**Yellow fever (continued)**

**Recommended types of surveillance (continued)**

**Note:** There is mandatory reporting to WHO of all suspected and confirmed yellow fever cases within 24 hours of detection.

**Recommended minimum data elements**

**Aggregated data for reporting**
- Number of cases
- Doses of yellow fever vaccine administered to infants by geographical area
- Completeness / timeliness of monthly reports

**Case-based data for reporting and investigation**
- Unique identifier
- Geographical area (e.g. district and province) names
- Date of birth
- Date of onset
- Date of notification
- Date of investigation
- Ever received a dose of yellow fever vaccine:
  - 1=yes; 2=no; 9=unknown
- Date acute blood specimen received in laboratory
- Date convalescent blood specimen received in laboratory (if applicable)
- Date histopathology specimen collected (if applicable)
- Depending on which laboratory tests used:
  - IgM results:
    - 1=positive; 2=negative; 3=no specimen processed; 9=unknown
  - Virus isolation results:
    - 1=positive; 2=negative; 3=no specimen processed; 9=unknown
  - IgG (4-fold rise) results:
    - 1=positive; 2=negative; 3=no specimen processed; 9=unknown
  - Liver histopathology:
    - 1=positive; 2=negative; 3=no specimen processed; 9=unknown
- Date IgM results first sent
- Date virus isolation results first sent
- Date IgG results first sent
- Date histopathology report first sent
- Final classification:
  - 1=confirmed; 2=suspected; 9=discarded
- Final outcome:
  - 1=alive; 2=dead; 9=unknown
Yellow fever (continued)

Recommended data analyses, presentation, reports

Aggregated data
- Incidence rate by month, year, and geographic area
- Yellow fever vaccine coverage by year and geographic area
- Completeness/timeliness of monthly reporting

Case-based data same as aggregated data plus the following:
- Confirmed cases by age group, immunization status, geographic area, month and year
- Case fatality rate
- Final classification of all suspect cases

- Performance indicators of surveillance quality
  - Completeness of monthly reporting. > 90%
  - Percent of all suspect cases for which specimens were collected. > 50%
  - For IgM test: Laboratory results sent < three days of receipt of acute blood specimen. > 80%
  - For virus isolation: Laboratory results sent < 21 days of receipt of acute blood specimen. > 80%
  - For IgG test: Lab results sent < three days of receipt of convalescent blood specimen. > 80%

  ¹ This is the target during non-outbreak periods. Once an outbreak is confirmed, the priority is to detect and laboratory-confirm outbreaks in neighbouring areas.

Principle uses of data for decision-making
- Investigate suspect cases and collect laboratory specimens to confirm an outbreak and mobilise emergency immunization activities
- Monitor yellow fever coverage by geographic region to assess progress towards outbreak prevention and identify areas of poor performance so that corrective actions can be taken
- Monitor incidence rate to assess impact of control efforts

Special aspects

The following 34 countries are at risk for yellow fever epidemics in Africa: Angola, Benin, Burkina Faso, Burundi, Cameroon, Cape Verde, Central African Republic, Chad, Congo, Côte d'Ivoire, Democratic Republic of Congo, Equatorial Guinea, Eritrea, Ethiopia, Gabon, Gambia, Ghana, Guinea, Guinea-Bissau, Liberia, Kenya, Mali, Mauritania, Niger, Nigeria, Rwanda, Sao Tomé and Principe, Senegal, Sierra Leone, Somalia, Sudan, Tanzania, Togo, Uganda. The following countries are at risk for yellow fever in South America: Bolivia, Brazil, Colombia, Ecuador, Guyana, French Guyana, Panama, Peru, Suriname, Trinidad and Tobago, Venezuela.
Yellow fever (continued)

Contact information

Regional Offices and Headquarters (VAB and CDS)
See Annex 1 for addresses and fax numbers.
Annex 1

WHO Headquarters and Regional Office contacts

WHO Headquarters
WHO Department of Vaccines and Biologicals
Vaccine Assessment and Monitoring (VAM)
20 Avenue Appia, CH-1211 Geneva 27, Switzerland
Fax: 41 22 791 4193; Email: vaccines@who.int
EPIdata@who.int

WHO Department of Communicable Disease
Surveillance and Response
Epidemic Disease Control (EDC)
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WHO AFRO
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