Viscrotropic Disease Associated with Vaccination against Yellow Fever

Case Studies

GUIDE FOR FACILITATOR
Visceralotropism Disease
Associated with Vaccination
against Yellow Fever
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The case studies presented in this guide should be implemented with the guidance of the Pan American Health Organization and under the supervision of experts in immunization, epidemiological surveillance, and the laboratory and clinical aspects of yellow fever.
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<table>
<thead>
<tr>
<th>Acronym</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALT (SGPT)</td>
<td>alanine transaminase (serum glutamic pyruvic transaminase)</td>
</tr>
<tr>
<td>ARDS</td>
<td>adult respiratory distress syndrome</td>
</tr>
<tr>
<td>AST (SGOT)</td>
<td>aspartate transaminase (serum glutamic-oxaloacetic transaminase)</td>
</tr>
<tr>
<td>CDC</td>
<td>Centers for Disease Control and Prevention (CDC)</td>
</tr>
<tr>
<td>cm</td>
<td>centimeter</td>
</tr>
<tr>
<td>CPK</td>
<td>creatinine phosphokinase</td>
</tr>
<tr>
<td>DGE</td>
<td>General Directorate of Epidemiology (Peru)</td>
</tr>
<tr>
<td>DIRESA</td>
<td>Regional Health Directorate (Peru)</td>
</tr>
<tr>
<td>dL</td>
<td>deciliter</td>
</tr>
<tr>
<td>ESAVI</td>
<td>event supposedly attributable to vaccination or immunization</td>
</tr>
<tr>
<td>IgG</td>
<td>immunoglobulin G</td>
</tr>
<tr>
<td>IgM</td>
<td>immunoglobulin M</td>
</tr>
<tr>
<td>IHR</td>
<td>International Health Regulations</td>
</tr>
<tr>
<td>INR</td>
<td>international normalized ratio (prothrombin time)</td>
</tr>
<tr>
<td>INS</td>
<td>National Institute of Health (Peruvian agency responsible for health research)</td>
</tr>
<tr>
<td>L</td>
<td>liter</td>
</tr>
<tr>
<td>mg</td>
<td>milligram</td>
</tr>
<tr>
<td>mL</td>
<td>milliliter</td>
</tr>
<tr>
<td>mm</td>
<td>millimeter</td>
</tr>
<tr>
<td>MoH</td>
<td>Ministry of Health</td>
</tr>
<tr>
<td>NAMRU-6</td>
<td>Naval Medical Research Unit 6 (formerly NMRC, United States)</td>
</tr>
<tr>
<td>PAHO</td>
<td>Pan American Health Organization</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
</tr>
<tr>
<td>PFU</td>
<td>plaque forming units</td>
</tr>
<tr>
<td>Q-PCR</td>
<td>quantitative polymerase chain reaction</td>
</tr>
<tr>
<td>RNA</td>
<td>ribonucleic acid</td>
</tr>
<tr>
<td>RT-PCR</td>
<td>reverse transcription polymerase chain reaction</td>
</tr>
<tr>
<td>U/L</td>
<td>units per liter</td>
</tr>
<tr>
<td>VTD</td>
<td>viscerotropic disease</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
</tbody>
</table>
GLOSSARY

**Active surveillance of ESAVis:** Surveillance based on the routine and proactive investigation of cases conducted on a case-by-case basis by health workers.

**Brighton Collaboration:** Global network of vaccine-safety experts. The organization’s website can be accessed at http://Brightoncollaboration.org/public.

**Crisis related to vaccination:** A situation caused by a real or perceived adverse event that produces the real or potential loss of confidence in vaccines and immunization services.

**Immunoglobulin M (IgM) and immunoglobulin G (IgG):** Antibodies produced by B-lymphocytes responsible for acute humoral immunity (IgM) and for immunological memory (IgG). A person with a recent infection tends to present IgM in the serum, while IgG appears later.

**National ESAVI Committee:** Committee responsible for assessing serious ESAVI cases in order to guide case investigation and provide final classifications. The committee includes professionals from various fields, including experts in epidemiology, public health, and biostatistics and members of the national regulatory authority (NRA).

**Neurologic disease associated with yellow fever vaccination (YEL-AND):** A neurological syndrome associated with the yellow fever vaccine that presents one or more of the following signs and symptoms: fever, focal neurological deficits, changes in mental status, seizures, or pleocytosis or excessive protein in cerebrospinal fluid. Neurologic disease can be due to direct vaccine viral invasion into the central nervous system or due to autoimmune manifestation triggered by the vaccine.

**Passive surveillance:** Surveillance based on the spontaneous reporting of ESAVI cases conducted on a case-by-case basis by health workers.

**Polymerase chain reaction (PCR):** Test used to detect genetic material (e.g., nucleic acids) in serum or tissue specimens.

**Real-time or quantitative PCR (Q-PCR):** Test that allows for the quantification of genetic material at any time during amplification. For cases of viscerotropic disease associated with yellow fever vaccine, Q-PCR makes it possible to identify viral concentrations in biological samples.

**Reverse transcription polymerase chain reaction (RT-PCR):** Test that detects and amplifies genetic material via retrotranscription of ribonucleic acid (RNA) into deoxyribonucleic acid (DNA).

**Risk communication:** Decision-making process that considers the risk of potential dangers in formulating, studying, and comparing risk control measures that are intended to protect the population in the event of a probable danger.

**Sentinel surveillance:** Surveillance based on reports by a group of sources (sentinel units) within the health system in which a sample of a population group is studied.

**Viscerotropic disease associated with yellow fever vaccination:** A disease associated with vaccination against yellow fever and characterized by systemic multiple organ failure and clinical symptoms similar to those of wild yellow fever.
INTRODUCTION

The purpose of the following case studies is to familiarize participants with case definitions and with the epidemiological investigation of viscerotropic disease (VTD) associated with vaccination against yellow fever. We present here a cluster of events supposedly attributable to vaccination or immunization (ESAVIs) that occurred in Peru in 2007, during a yellow fever vaccination campaign implemented in a non-endemic area.

FOR WHOM IS THIS DOCUMENT INTENDED

1. Health professionals from the national, subnational, and local levels who participate in the investigation of ESAVI cases.

2. Clinicians, laboratorians, and other professionals (academics, investigators, etc.) who participate in vaccination activities or pharmacovigilance.

3. Staff from the national regulatory authority, particularly those responsible for pharmacovigilance and vaccine regulation.

STUDY OBJECTIVES

1. To understand and apply case definitions of VTD.

2. To analyze and implement reporting, investigation, and classification procedures for cases of VTD in a timely manner.

3. To review response measures in the event of a case of VTD associated with yellow fever vaccine.

4. To understand and apply principles of causality assessment for cases of VTD.

5. To understand the importance of planning for establishing strategic alliances with media sources and other important stakeholders.

GENERAL INSTRUCTIONS

1. Participants will work as members of a rapid response team assigned to investigate multiple cases of VTD associated with yellow fever vaccine.

2. Participants will be assigned to teams of approximately 6-8 persons to answer questions related to each case study. Each group will have a facilitator and a rapporteur. After reading the case studies, group members will discuss their responses, sharing their experiences and raising concerns.

3. Group work allows for the exchange of experiences and for participants to learn new information and opinions.

4. In group work, the facilitator should help to stimulate the conversation and assist participants with responses as needed, while maintaining the conceptual framework.
EVALUATION
To measure participant learning during the case studies, evaluations of the participants’ knowledge will be conducted at the beginning and end of the exercises (Annex 1). It is recommended to present a summary of these results and the progress made during the case studies to participants.

PRIOR KNOWLEDGE
In preparation for working with the case studies, participants are encouraged to review the following documents:

ABOUT YELLOW FEVER AND ESAVIS

YELLOW FEVER

Yellow fever is a disease endemic to the tropical areas of Africa and South America. Approximately 200,000 cases and 30,000 deaths from the disease occur each year (1). These cases primarily affect young adult males who, for occupational reasons, must enter enzootic areas. In the Region of the Americas, yellow fever is endemic mainly in Bolivia, Brazil (eastern-central region), Colombia, Ecuador, French Guiana, Guyana, Peru, Suriname, Trinidad and Tobago, and Venezuela. Sporadic cases have also been reported in other countries, such as Argentina, Panama, and Paraguay (2). Yellow fever virus transmission in the Americas predominantly occurs in a jungle cycle. Enzootic areas are rural, isolated from urban areas, and mostly situated in the Amazon jungle (3). Nonetheless, in 2008, yellow fever cases occurred in a peri-urban area in Paraguay (4).

Yellow fever can be prevented with live-attenuated yellow fever vaccine 17D, which is considered safe and effective. Yellow fever vaccination strategies implemented in the Region of the Americas include: 1) introduction of yellow fever vaccine into national immunization programs in all endemic countries for children aged 1 year; 2) vaccination campaigns during inter-epidemic periods; 3) vaccination campaigns in response to outbreaks or epizootics; and 4) vaccination of travelers entering enzootic areas, except when contraindicated.

As of 2011, all countries in the Americas with enzootic areas have included yellow fever vaccine in their national vaccination schedules. In Argentina, Brazil, and Suriname, the vaccine is administered exclusively in areas considered at risk. Vaccination coverage of children aged 1 year was approximately 70% for the period 2007-2011 and has been significantly affected by limited vaccine availability.

The International Health Regulations (IHR) indicate that proof of yellow fever vaccination can be demanded from travelers as a requirement for admission into a country (5).

EVENTS SUPPOSEDLY ATTRIBUTABLE TO VACCINATION OR IMMUNIZATION (ESAVIs)

An ESAVI is defined as a set of clinical symptoms that occur following immunization, which may or may not be attributable to the vaccine, and which causes great concern among the population (6).

The following ESAVIS must be reported, investigated, and classified in a timely manner: 1) serious events that require hospitalization, are life-threatening for the patient, cause disability, or are fatal; 2) events that affect a group of people (disease clusters); 3) events related to the immunization program (programmatic errors); and 4) events that generate rumors or confusion among the population.

It is difficult to determine whether an ESAVI is truly the result of a vaccine. Consequently, the objectives of ESAVI investigation are to confirm or rule out the notified event as vaccine-related, determine if the event might have another cause, determine whether the event is isolated or related to other events, and notify all interested parties of the results of the investigation.

Following a thorough review by vaccine-safety experts, the classification of ESAVIS was recently modified as follows (7):
Vaccine product-related event:
An event that is caused or precipitated by a vaccine due to one or more of the inherent properties of the vaccine product.

Example: Extensive limb inflammation (edema) following diphtheria-tetanus-pertussis (DPT) vaccination.

Vaccine quality defect-related event:
An event that is caused or precipitated by a vaccine that is due to one or more quality defects of the vaccine product including its administration device as provided by the manufacturer.

Example: Failure of the manufacturer to completely inactivate a lot of inactivated polio vaccine (IPV) leads to cases of paralytic polio.

Immunization error-related event:
An event that is caused by inappropriate vaccine handling, prescribing, or administration and thus by nature is preventable.

Example: Sepsis, toxic shock syndrome, infection (e.g., localized abscess at the injection site), or death due to a non-sterile injection.

Immunization anxiety-related event:
An event that arises from anxiety about the vaccine.

Example: Vaso-vagal syncope in adolescents following immunization against the Human Papillomavirus (HPV).

Coincidental events:
An event that is caused by something other than the vaccine product, immunization error, or immunization anxiety.

Example: Fever occurs at the time of vaccination (temporal association) but is caused by another agent such as malaria.

Inconclusive cases:
An event whose cause remains unknown despite a thorough investigation. In such cases, it must be explained why a conclusion was not reached and how far the investigation progressed.

**ESAVIs associated with yellow fever vaccine**

In general, yellow fever vaccine is considered to be among the safest vaccines in existence. More than 500 million people have received the vaccine, with very good results in terms of safety and tolerance. Nevertheless, side effects, such as fever, headache, and back pain, have been observed since the first studies on the 17D vaccine were conducted in the 1930s (8).

Between 1953-1994, 10 clinical trials were conducted to study reactions to the vaccine. A small percentage of those vaccinated experienced mild, self-limited reactions. The most common of these were pain and erythema at the injection site as well as systemic reactions, including fever, chills, headache, myalgia, and malaise. These symptoms typically appeared 3–7 days following vaccination. Reactogenicity was observed to be less in infants than in adults (9).
Serious ESAVIs associated with yellow fever vaccine are rare. According to available data, the incidence of reported ESAVIs is 1.6 cases per 100,000 vaccine doses (10). The most frequently reported serious ESAVIs in the scientific literature are cases of VTD, neurologic disease, and severe hypersensitivity reactions.

Viscerotropic disease was first identified as an adverse event of yellow fever vaccination in 2001 (11). The disease occurs by dissemination and widespread replication of the 17D live attenuated virus of the vaccine. Based primarily on a retrospective review of ESAVI reports, 65 cases of VTD following yellow fever vaccination were identified as occurring worldwide between 1973 and March 2011.

Although the incidence of the disease is unknown largely because of underreporting and passive ESAVI surveillance systems, data from travelers in the United States and Europe show a risk of 0.3-0.4 VTD cases per 100,000 yellow fever vaccine doses distributed. Furthermore, based on data from 2008 and 2009 vaccination campaigns in Brazil, the estimated risk of VTD following yellow fever vaccination in that country is similar to those reported in the United States and Europe (12). The mortality rate is estimated to be 60% in notified cases, though this may be an overestimate because fatal cases are more likely to be reported than nonfatal cases (12).

Although it has not been possible to determine the physiopathological mechanism that causes VTD following yellow fever vaccination, some population groups, such as those aged ≥60 years, have demonstrated a greater risk of developing the disease. As compared to the 19-29 year age group, the case ratio of the ≥60-year age group is 5.9, with an estimated relative risk of 4.4-13.4. In addition, a history of thymectomy for benign or malignant thymus disease is considered a risk factor for VTD associated with the yellow fever vaccine (12).

In the United States, 0.4-0.8 cases of neurologic disease following yellow fever vaccination (Yel-aNd) have been reported per 100,000 vaccine doses administered. The highest rate is found among individuals aged ≥60 years (1.4-1.8 cases per 100,000 doses administered). Australia and the United Kingdom reported similar data, suggesting an increased risk for the elderly. Cases of post-vaccination encephalitis have also been attributed to the 17D yellow fever vaccine virus in infants aged <4 months (0.8 cases per 100,000 doses administered). As a result, the vaccine is contraindicated for infants aged <6 months and thereby has a greater safety margin. Neurological sequelae are unusual and deaths are rare. Neurological events reported as related to yellow fever vaccine include acute disseminated encephalomyelitis and Guillain-Barré Syndrome (9).

Lastly, serious hypersensitivity reactions are extremely rare. The yellow fever vaccine is contraindicated for individuals allergic to eggs. The most frequent hypersensitivity reactions are skin rashes, asthma (1 case per 130,000-250,000 doses administered) (9), and anaphylactic shock (0.8 cases per 100,000 doses administered) (13).
BACKGROUND

On 15 August 2007, a severe earthquake struck Peru. The earthquake had a magnitude of 8.0 on the Richter scale and its epicenter was located 40 km from Ica, a city south of the country’s capital, Lima. The Ica Region lies along the coast, and its population at the time was estimated to be 693,411 inhabitants in five provinces: Chincha, Ica, Nazca, Palpa, and Pisco.

Although Ica is a non-endemic area for yellow fever, Peru’s Ministry of Health (MoH) began a vaccination campaign targeting individuals aged 15-59 years on 23 September 2007. The campaign used two vaccine lots, 050VFA121Z (121Z) and 050VFA123Z (123Z), of 17DD vaccine manufactured at Bio-Manguinhos in Brazil. From the first lot, 42,742 doses were administered; from the second, 20,432 were administered. These lots were used solely in the Ica Region.

As part of the campaign, the Ica Regional Health Directorate (DireSa in Spanish) strengthened regional surveillance of adverse events and established daily negative reporting of serious ESAVI cases. A few months before, the MoH’s General Directorate of Epidemiology (DGE in Spanish) had organized a training workshop on ESAVI surveillance for healthcare workers of the Ica DireSa. The training was part of ongoing evaluation and training activities in the country’s surveillance network.

The vaccination campaign in Ica was one of several activities carried out to mitigate the earthquake’s effects and complemented the National Plan for Accelerated Yellow Fever Control started in 2004. The plan included routine vaccination of children aged 1 year throughout the country, and vaccination campaigns for individuals aged 2-59 years in two phases: 1) vaccination of the population living in endemic areas and 2) vaccination of populations in migratory areas, where many people leave non-endemic areas and enter endemic areas during the harvest season. As part of the plan, approximately 12 million doses of yellow fever vaccine were administered between 2004-2007 in endemic and migratory areas. During this period, no cases of VTD associated with yellow fever vaccine were reported. Vaccination campaigns continued throughout 2007 in the regions of Amazonas, Cusco, San Martín, and Piura. Figure 1 shows the assessment of yellow fever risk areas in Peru.
FIGURE 1. ASSESSMENT OF YELLOW FEVER RISK AREAS, PERU, 2012
A. FIRST CASE: MEDICAL STUDENT

On 6 October 2007 at 1:30 a.m., the epidemiologist in charge of the surveillance of vaccine-preventable diseases at the MoH received a call from his counterpart at the Ica Regional Hospital, reporting a serious ESAVI. The Ica epidemiologist indicated that he had already notified officials at the regional level, but given the situation, he also wanted to immediately notify the MoH. The patient was a female medical student aged 23 years and a former student of the Ica epidemiologist. The epidemiologist himself had seen the patient in the hospital’s emergency room on the morning of 5 October. The patient arrived with clinical symptoms of approximately a week’s duration, including fever, nausea, vomiting, and diarrhea. What most alarmed the attending physician was how rapidly the patient’s health had deteriorated. She had arrived at the hospital on her own, lucid, and without any apparent serious clinical symptoms. Twelve hours later, she had to be admitted to the intensive care unit due to liver and renal failure. The only notable precursor to the patient’s illness had been her yellow fever vaccination on Thursday, September 27, a fact confirmed by her vaccination card.

The MoH epidemiologist advised his counterpart at the Ica Regional Hospital to monitor the patient’s progress and to notify him of any changes. At 6:00 a.m., the hospital epidemiologist reported that the patient’s symptoms had worsened and that her prognosis was poor. The diagnosis was multiple organ failure. The patient was provided inotropic support and mechanical ventilation.

The epidemiologist at the Ica Hospital was advised to obtain a blood specimen from the patient to send to the National Institute of Health (INS in Spanish) in order to determine the etiological diagnosis. The patient’s clinical symptoms were similar to those of a rare adverse reaction documented in several medical journal articles: viscerotrophic disease associated with yellow fever vaccine. Based on this information, two epidemiologists were immediately dispatched to investigate the case. Additionally, the National ESAVI Committee was urgently convened and PAHO’s focal point for yellow fever immunization was informed of the case.

1. Did the Ica Hospital epidemiologist notify the case in a timely manner? Explain.

   The notification was made in a timely manner because the case was serious and, as such, should have been reported within 24 hours in accordance with the requirements of Peru’s ESAVI surveillance system.

2. Use IHR regulations to determine the mechanism for official international notification of this case.

   Annex 2 of the IHR requires that answers to the following questions be provided for all ESAVIs detected by the national surveillance system. An adapted version of the algorithm is provided in Figure 2 below.
FIGURE 2. ANNEX 2 OF THE INTERNATIONAL HEALTH REGULATIONS

I. DOES THE EVENT HAVE A SERIOUS IMPACT ON PUBLIC HEALTH?

1. Is the number of cases and/or the number of deaths for this type of event higher than expected for the given time, place, or population?
   Answer: Yes. The risk, 1 case per 42,742 doses of yellow fever vaccine, is higher than the risk of 0.3-0.4 cases per 100,000 doses reported in the literature.

2. Has the event the potential to have a high public health impact?
   Answer: Yes. The yellow fever vaccine was being administered to a large susceptible population. In addition, an earthquake had recently struck the area and the population was still recovering.

3. Is external assistance needed to detect, investigate, respond, and control the current event, or prevent new cases?
   Answer: Yes. The tests required to diagnose the event were not available in the country.

Since questions 1, 2, and 3 were answered “Yes,” THE EVENT HAS A SERIOUS PUBLIC HEALTH IMPACT.

II. IS THE EVENT UNUSUAL OR UNEXPECTED?

4. Is the event unusual?
   Answer: No. The agent, source, and clinical presentation of the event are described in the scientific literature.

5. Is the event unexpected from a public health perspective?
   Answer: No. The event has been notified previously and is not caused by a disease or agent already eliminated or eradicated from the country.

Since questions 4 and 5 were answered “No,” THE EVENT IS NOT UNUSUAL OR UNEXPECTED.

(Continued)
In Ica, the patient continued to evolve unfavorably. Complementary testing showed the following results: metabolic acidosis; moderate increases in transaminases; AST (SGOT): 78 U/L; ALT (SGPT): 65 U/L; bilirubin: 0.85 mg/dL; leukocytes: 66,400 mL; lymphocytes: 8%; monocytes: 2%; neutrophils: 90%; platelets: 54,000 mL; and creatinine: 4.1 mg/dL. Despite the administration of inotropic drugs, the patient had a systolic blood pressure of 60 mm Hg and a central venous pressure of 15 cm H₂O. Chest X-rays showed pulmonary congestion with prominent hilar markings and bilateral pleural effusions, while abdominal ultrasonography showed hepatosplenomegaly and ascites (>500 mL). The patient also presented oliguresis (10 cm³/hour).

Due to the patient’s fragile health, she could not be transferred to a hospital in Lima and remained at the Ica hospital. The patient died on the same day of her admission (October 6) at 10:00 p.m.

---

1 All acronyms related to diagnostic results are explained in the list of abbreviations provided at the beginning of the case studies.
3. What information should the epidemiologists at the Ica Regional Hospital obtain to complete the case investigation?

*Use of Table 1 is recommended for this response.*

**TABLE 1. DATA COLLECTION CHECKLIST FOR THE INVESTIGATION OF VTD**

<table>
<thead>
<tr>
<th>CATEGORY</th>
<th>VARIABLES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographics</td>
<td>• Name</td>
</tr>
<tr>
<td></td>
<td>• Birthday or age</td>
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<tr>
<td></td>
<td>• Sex</td>
</tr>
<tr>
<td></td>
<td>• Place of residence</td>
</tr>
<tr>
<td></td>
<td>• Occupation(s)</td>
</tr>
<tr>
<td></td>
<td>• Infants (&lt;12 months): gestational age at birth and birth weight</td>
</tr>
<tr>
<td></td>
<td>• Address and phone number</td>
</tr>
<tr>
<td>Clinical history</td>
<td>• Medical history of any disease or illness experienced prior to most recent vaccination(s), including hospitalization, surgery (e.g., thymectomy), and any other disease or disorder with underlying pre-immunization signs and symptoms.</td>
</tr>
<tr>
<td></td>
<td>• Genetic diseases.</td>
</tr>
<tr>
<td></td>
<td>• Compromised immune system due to primary or acquired immunodeficiency diseases.</td>
</tr>
<tr>
<td></td>
<td>• Compromised immune system due to cancer, chemotherapy, or immunosuppressive radiation or treatment, among others.</td>
</tr>
<tr>
<td></td>
<td>• Allergies to previous vaccine doses or eggs.</td>
</tr>
<tr>
<td></td>
<td>• For women of childbearing age: date of last menstruation, date of last delivery. For postpartum women: history of breastfeeding.</td>
</tr>
<tr>
<td>History of medications</td>
<td>Medications taken before, during, or after vaccination, including those with and without prescriptions and herbal or homeopathic medicines. These medicines should be different than those used to treat VTD.</td>
</tr>
<tr>
<td></td>
<td>Medications such as immunoglobulins, blood products, immunosuppressants, and immunomodulators should be recorded, even if they had been used several months before vaccination. Recommendations on individual drug pharmacokinetics should be consulted to identify potential interactions.</td>
</tr>
<tr>
<td></td>
<td>Any form of contraception should be registered.</td>
</tr>
<tr>
<td>History of previous vaccinations</td>
<td><strong>Any vaccine administered &gt;30 days before VTD symptom onset,</strong> especially vaccines known to cause VTD, such as yellow fever vaccine and vaccines against varicella, polio, and BCG. Any previous ESAVs should also be noted.</td>
</tr>
<tr>
<td>Other background information</td>
<td>Presence or absence of local outbreaks or recent travel to an area affected by a disease outbreak (e.g., yellow fever).</td>
</tr>
</tbody>
</table>

*(Continued)*
**TABLE 1. (Continued).**

<table>
<thead>
<tr>
<th>CATEGORY</th>
<th>VARIABLES</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Vaccination details</strong></td>
<td>Date, time, and geographic location (e.g., city or town, state or province, and country) of all vaccines administered ≤30 days before VTD symptom onset.</td>
</tr>
<tr>
<td></td>
<td>Description of the vaccine(s) administered ≤30 days before VTD symptom onset: name of vaccine manufacturer, lot number, expiration date, multi-dose or single-dose vial (e.g., 0.25 mL, 0.5 mL, etc.), number of doses (if vaccine is part of vaccination series against the same disease), and name of manufacturer, batch number, and expiration date of any diluent used.</td>
</tr>
<tr>
<td></td>
<td>Anatomical sites including the side (left or right) in which vaccines were applied (e.g., vaccine A in superolateral region of the left thigh; vaccine B in left deltoid).</td>
</tr>
<tr>
<td></td>
<td>Means and method of administration: intramuscular, intradermal, subcutaneous, oral, intranasal, with needle (include size and type), needleless, or use of other injection device.</td>
</tr>
<tr>
<td></td>
<td>Context of vaccination: routine childhood vaccination, preventative mass immunization campaign, mass immunization campaign for outbreak response, routine adult immunization, domestic travel from non-endemic to endemic area, international travel, occupational hazards, military requirements, or other requirements.</td>
</tr>
<tr>
<td><strong>Adverse event</strong></td>
<td>Detailed clinical description of the signs and symptoms of VTD, particularly jaundice, hypoxia, hypotension, oliguria, tachypnea, petechiae, purpura, and bleeding.</td>
</tr>
<tr>
<td></td>
<td>• Date and time of onset(^1)</td>
</tr>
<tr>
<td></td>
<td>• First observations(^2)</td>
</tr>
<tr>
<td></td>
<td>• Diagnosis(^3)</td>
</tr>
<tr>
<td></td>
<td>• End of an episode(^4)</td>
</tr>
<tr>
<td></td>
<td>• Last observations(^5)</td>
</tr>
<tr>
<td></td>
<td>Signs, symptoms, and comorbidities that are not the event described.</td>
</tr>
<tr>
<td><strong>Results of laboratory tests, other clinical tests, surgical and/or pathological findings, and diagnoses</strong></td>
<td>If possible, efforts should be made to perform the laboratory tests in reliable and accredited institutions (Annex 2). If more than one measurement of a particular indicator is taken and registered, the expected normal range for that indicator as well as any deviation from it should be recorded (e.g., highest serum creatinine, lowest platelet count). Units for each of the parameters must be specified (e.g., cm, °C, etc.), particularly for those indicating the severity of VTD. Normal ranges observed in specific laboratory or clinical settings should also be reported for each parameter measured, as should the date of specimen collection and/or measurement for each test.</td>
</tr>
<tr>
<td><strong>Treatment for VTD</strong></td>
<td>Especially hospitalization, supplemental oxygen, intravenous fluids, vasopressors, mechanical ventilation, hemodialysis, steroids, antiviral drugs as well as names and addresses of doctors and/or the healthcare facilities responsible for treatment.</td>
</tr>
</tbody>
</table>

(Continued)
TABLE 1. (Continued).

<table>
<thead>
<tr>
<th>CATEGORY</th>
<th>VARIABLES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final health status</td>
<td>- Recovery of health status experienced prior to immunization</td>
</tr>
<tr>
<td></td>
<td>- Spontaneous resolution</td>
</tr>
<tr>
<td></td>
<td>- Ongoing treatment</td>
</tr>
<tr>
<td></td>
<td>- Persistence of the event</td>
</tr>
<tr>
<td></td>
<td>- Sequelae from VTD</td>
</tr>
<tr>
<td></td>
<td>- Significant complications of treatment</td>
</tr>
<tr>
<td></td>
<td>- Death</td>
</tr>
<tr>
<td></td>
<td>Description of any other outcome</td>
</tr>
</tbody>
</table>


1 The date and time of onset are defined as the date post-immunization when the first sign or symptom suggestive of VTD occurred. This may only be possible to determine in retrospect.

2 If the date/time of onset is unknown, the date and/or time of the first observation of the first sign or symptoms suggestive of VTD can be used.

3 The date of diagnosis of an episode is the date following immunization that the event first met the case definition.

4 The end of an episode is the date the event no longer meets the case definition.

5 VTD not resolved at the end of a predefined follow-up period may be followed up as clinically necessary, and additional reporting should be encouraged to describe progress until the clinical status is stable. The date of last observation is the date of the last clinical evaluation of the patient with VTD. The persistence of event refers to events continuing to meet the case definition beyond the follow-up period. Sequelae are long-term clinical consequences resulting from VTD.

B. SUSPENSION OF THE VACCINATION CAMPAIGN

The same day of the patient’s death and in response to increasing media pressure, the director of DIRESA announced at a press conference that vaccination against yellow fever would be suspended throughout the region, until the results of the investigation of the death of the young student who received one dose of vaccine became available (15).

On 8 October, the MoH convened two emergency committee meetings: the National ESAVI Committee and the Crisis Committee, chaired by the vice minister of health. Both committees agreed that, throughout the crisis, the Ministry’s spokespersons would be the president of the ESAVI Committee and the president of the National Immunization Technical Advisory Group (NITAG). They also concurred that these officers would respond to media requests for information on the campaign’s suspension in Ica.

1. What arguments exist in favor of suspending the vaccination campaign?

   - The ESAVI occurred in an area that had recently suffered an earthquake. As a result, all state-run services, including the health system, were impaired.
The ESAVI was a serious event, expected for the vaccine, but unexpected for the number of doses administered.

There was no short- or long-term risk of a yellow fever epidemic.

2. Should yellow fever vaccination have been suspended in other parts of the country? Support your answer.

No, vaccination in other regions should not have been suspended because:

- There was no report of serious ESAVI cases compatible with VTD in those regions.
- Vaccination was part of an established plan launched years ago, and similar cases had not occurred in the population at risk for yellow fever (in Peru, more than 200 cases of jungle yellow fever had been reported in the previous five years).
- The same vaccine lot had not been used in other areas of the country.

3. What is the function of the Crisis Committee, and who should be part of it? Briefly mention the components that should be included in a crisis plan for a situation like the one described above.

The Crisis Committee should organize and implement activities that prevent, control, and mitigate a crisis stemming from vaccination activities. The committee should include officials from the highest levels of government so that required measures can be quickly implemented. Ideally, the committee should be chaired by the area’s highest health authority and be composed of members representing all technical aspects of the immunization program, including experts in epidemiology, laboratories, vaccine regulation, and communication. When possible, representatives from PAHO, professional associations, and scientific societies should be included.

Crisis plans should have the following objectives: 1) organize the response team to address technical issues; 2) confirm the facts of a case and determine the need for further investigation; 3) coordinate response efforts and manage damage control; 4) implement a communication plan for the crisis; 5) revise or develop a plan for legal aspects related to the crisis; 6) designate spokespersons and a leader responsible for managing the crisis; and 7) evaluate the response.

TIME AVAILABLE: 90 MINUTES

A. SECOND CASE: A YOUNG MOTHER

On 8 October 2007, a 24-year-old female sought emergency services at the Chincha provincial hospital, located in the Ica Region. The patient had been referred by her private physician, who had evaluated her hours earlier at her home. Three months before, she had given birth to a healthy child. According to the patient, she had been vaccinated against yellow fever approximately 10 days earlier; however, she did not have her vaccination card on hand. On the day of vaccination, she experienced pain at the injection site. The next day, she experienced headache, malaise, myalgia, and fever. These symptoms lasted 3-4 days. On 30 September, a private physician diagnosed her with a urinary tract infection and prescribed ciprofloxacin, dexamethasone, and amikacin.
By 3 October, in addition to her original symptoms, the patient developed watery stools, nausea, and vomiting. She had as many as 20 bowel movements on 7 October. Following her medical evaluation, the patient was diagnosed with hypovolemic shock, severe dehydration, and renal failure. The epidemiologist at the hospital immediately reported the case to the Ica DIRESA, which in turn notified the DGE. The DGE arranged the patient’s transfer to the Dos de Mayo National Hospital in Lima, where she arrived on the night of 8 October.

Clinical examination upon admission showed: blood pressure: 80/50 mm Hg; heart rate: 100/ min; respiratory rate: 30/min; and mydriatic pupils, 4/4 with response to light. Other signs and symptoms included slow capillary refill time, tachypnea, dyspnea, and vesicular murmur passing properly in both lungs. The patient was moaning, agitated, and able to move her four limbs. On the night of her admission to the hospital, two epidemiologists visited the Dos de Mayo Hospital to obtain the patient’s background and clinical information and to request that a blood specimen be sent to the INS.

1. **What specimens and laboratory tests should be used to classify the case?**

   Since VTD is suspected, the disease’s major diagnostic criteria should be evaluated (Annex 3). Serum specimens should be obtained to determine at minimum: bilirubin level (total, direct, and indirect); transaminases levels (liver criterion); creatinine level (renal criterion); and CPK level (musculoskeletal criterion). Additionally, the following should be assessed: thrombocytopenia (platelet count <100,000/µL); coagulopathies (through determination of the international normalized ratio [INR], prothrombin time, or activated thromboplastin, among others); respiratory disorders (through low oxygen saturation with spontaneous ventilation [88%] or the need for mechanical ventilation); and the need for vasopressors to maintain systolic blood pressure.

   To establish the yellow fever vaccine as the cause of VTD, strain 17D (viral RNA) of the yellow fever virus must be obtained in a blood or tissue specimen ≥10 days following vaccination. Alternatively, the concentration of the viral RNA can be determined in a blood specimen of ≥3 log_{10} plaque forming units (PFU)/mL at any time following vaccination.

   Specimens should remain frozen or in the cold chain (especially when the case has occurred in a distant area), although tissue specimens may be fixed in 10% formalin.

   Tables 2 and 3 summarize the specimens and laboratory tests for both VTD and VTD associated with yellow fever vaccine.
### TABLE 2: LABORATORY TESTS AND SPECIMENS FOR VTD

<table>
<thead>
<tr>
<th>SPECIMEN TYPE</th>
<th>TARGET ORGAN/SYSTEM</th>
<th>LABORATORY TESTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum</td>
<td>Liver</td>
<td>• Bilirubin level</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Transaminases levels</td>
</tr>
<tr>
<td></td>
<td>Kidney</td>
<td>• Creatinine</td>
</tr>
<tr>
<td></td>
<td>Striated muscle</td>
<td>• Creatine phosphokinase (CPK)</td>
</tr>
<tr>
<td></td>
<td>Blood (coagulation)</td>
<td>• Platelet count</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Prothrombin time</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Activated thromboplastin time</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Products of fibrin degradation</td>
</tr>
</tbody>
</table>


### TABLE 3: LABORATORY TESTS AND SPECIMENS FOR VTD ASSOCIATED WITH YELLOW FEVER VACCINATION

<table>
<thead>
<tr>
<th>SPECIMEN TYPE</th>
<th>LABORATORY TESTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum</td>
<td>• Serological tests</td>
</tr>
<tr>
<td></td>
<td>• Viral isolation</td>
</tr>
<tr>
<td></td>
<td>• Viral load</td>
</tr>
<tr>
<td></td>
<td>• Detection of viral RNA</td>
</tr>
<tr>
<td>Tissue (liver, spleen, kidneys, brain, meninges, lungs, thymus, thyroid, pancreas, heart, suprarenal glands, and skin)</td>
<td>• Viral isolation</td>
</tr>
<tr>
<td></td>
<td>• Detection of viral RNA</td>
</tr>
<tr>
<td></td>
<td>• Immunohistochemistry</td>
</tr>
<tr>
<td></td>
<td>• Histopathology</td>
</tr>
</tbody>
</table>


2. **What should the Ica DIRESA epidemiologist investigate while visiting the patient’s home?**

During the home visit, it is necessary to obtain the patient’s vaccination card, corroborate the patient’s case history (onset and evolution of symptoms prior to hospitalization), and obtain the patient’s history of personal and family illnesses as well as evidence of pre- and post-vaccination treatments. The epidemiologist should interview the caregiver—in this case, the husband or another adult family member living with the family.

### B. FIELD INVESTIGATION

The Ica DIRESA and Chincha hospital epidemiologists interviewed the patient’s husband, who indicated that she had given birth to a full-term healthy infant three months earlier, a fact corroborated by her clinical records. The patient’s vaccination history could not be verified, as the husband did not have her vaccination card. Nonetheless, he indicated that she had been vaccinated at the Chincha hospital. The patient had not traveled in previous months. Unsuccessful
attempts were made to locate the patient’s clinical records corresponding to the private care she received, which would have indicated diagnosis of a urinary tract infection days before her hospital admission. However, several empty amikacin containers found at the patient’s home confirmed that she had received treatment for such an infection. As time passed and the case gained notoriety in the media, it became impossible to again speak with the patient’s husband, who refused further contact with investigators.

The epidemiologists, in coordination with the head of the hospital’s immunizations program, verified the patient’s vaccination records at the Chincha hospital. They then interviewed the health care worker who had vaccinated the patient and evaluated injection practices used in the hospital. The patient had been immunized with vaccine from lot 121Z on 27 September—the same lot responsible for the medical student’s death on 6 October.

1. What aspects of vaccination safety should the investigating team have evaluated during its visit to the vaccination center?

- **Safety of person vaccinated:**
  1) quality of vaccine products (laboratory certified by WHO);
  2) safety, transport, and storage of products from time of manufacture until administration;
  3) potency, safety, and proper storage of the vaccine;
  4) handling of multi-dose bottles;
  5) administration technique and proper use of syringes; and
  6) re-evaluation of vaccine quality in light of a suspected operational problem.

- **Safety of vaccinator:**
  1) proper handling (do not cover or remove needles) and
  2) use of safety boxes.

- **Safety of the environment:** Vaccines should be safely and properly discarded. The safest and most reliable means of disposal is by destroying the biohazard container in an incinerator. Discarding or burning vaccine vials in an open field or municipal landfill is not recommended.

2. What hypothesis is suggested by the fact that both patients received vaccine from the same lot?

A programmatic error or a problem with the quality of the vaccine might be suspected, such as contamination or a mutation of the virus used in the lot. If this is the case, the vaccine lot may need to be suspended. In addition, vaccine storage and transportation conditions and procedures would need to be evaluated, as would the quality and safety of the vaccine and its process of production. This would seriously impact other regions of Peru and other countries that have received the lot.

3. What would you do if the patient’s family member refused to cooperate in the investigation?

To prevent refusal, patients and family members should receive whatever care or assistance they need from the health system. In the event that family members refuse to cooperate, other social actors, such as a community leader, may be needed to reestablish communication with the family.
When first contacting the family, investigators should use an extensive list of questions to obtain as much information as possible about the case. Family members should also receive advanced notice of follow-up visits. Lastly, it is preferable that one person serves as the point of contact with the family throughout the investigation.

C. Evaluation of the Vaccination Center and the Cold Chain

Using a structured questionnaire designed specifically to evaluate aspects of vaccination safety, epidemiologists assessed injection practices and the experience and knowledge of the person who administered the vaccine associated with the young mother’s death. Investigators also observed the immunization of other patients at the vaccination center. Based on this evaluation, the nurse’s performance was considered adequate and the health center was determined to be well organized. The cold chain was in good working order and possessed sufficient cooling capacity. Nevertheless, given the rush in implementing the campaign, the nurse had not been specifically trained in yellow fever vaccination. The vaccination center also lacked equipment to properly dispose of vaccine waste products.

The investigating team visited Ica DIRESA to evaluate the quality of the cold chain, which met technical specifications for biological storage and transportation. Around this time, upon learning that the patient had received vaccine from the same lot as the medical student, the MoH evaluated the storage and transportation of vaccines at all levels and requested that samples from the vaccine vials of the lot in question be sent to the INS for safety and potency evaluations. The MoH then requested technical assistance from PAHO, which offered to investigate the lot’s production in its country and laboratory of origin (Bio-Manguinhos, Brazil).

1. Do you agree with the need to perform potency and safety tests? Justify your response.

Although these tests are usually conducted to determine vaccine quality, neither is necessarily useful in investigating ESAVI cases. Potency tests are used when the efficacy of the vaccine is questionable. Safety tests are used to detect product contamination. Nevertheless, other tests may be useful, such as those measuring the effectiveness of the preservative agent, deficiencies in the cold chain, improper use of the vaccine, and storage of the vaccine lot. What is crucial is to inform the national regulatory authority of the situation and provide the agency with samples of vials from the lot and, if possible, from the vial under investigation.

2. What are the recommendations for shipping vaccine samples?

If the vial has not yet been reconstituted, it should be stored at 2°C–8°C and transported in cold packs. If the vial has been reconstituted, there should be at least 1 mL in each vial, which should be kept at -70°C and transported in dry ice (Table 4). Unreconstituted vaccine is preferable to reconstituted vaccine. Several vaccine vials should be reserved for future tests.
TABLE 4: RECOMMENDATIONS FOR SENDING VACCINE SAMPLES

<table>
<thead>
<tr>
<th>VACCINE</th>
<th>VOLUME</th>
<th>STORAGE</th>
<th>TRANSPORT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unreconstituted</td>
<td>Vial</td>
<td>+4°C</td>
<td>Cold packs</td>
</tr>
<tr>
<td>Reconstituted</td>
<td>At least 1 mL per vial</td>
<td>-70°C</td>
<td>Dry ice or cold packs</td>
</tr>
</tbody>
</table>


D. CLINICAL AND LABORATORY ASSESSMENT

While admitted at the Dos de Mayo Hospital, the patient received treatment with intravenous fluids, vasopressors, hydrocortisone, meropenem, vancomycin, and insulin, and was placed on mechanical ventilation. She experienced fever (40.2°C), jaundice, edema, and thrombocytopenia (24,000 blood platelets/mL). Results of liver function tests were as follows: AST (SGOT): 735 U/L; ALT (SGPT): 167 U/L; bilirubin: 6.3 mg/dL; CPK: 3,173 U/L; and INR: 3. The patient also presented renal function deficiency, with serum creatinine of 3.2 mg/dL and urea of 154 mg/dL. The patient’s stool culture tested positive for Escherichia coli O86. Likewise, in her first blood culture, Staphylococcus strains were isolated, although this was attributed to a contaminated specimen. A second blood culture tested positive for Candida spp. The patient died on her third day of hospitalization. The autopsy was conducted at the Dos de Mayo Hospital, and was attended by the epidemiologist of the Lima Health Directorate that has jurisdiction over the hospital.

The serum specimen analyzed by the INS yielded the following results:

- Serum IgM positive for yellow fever.
- Serum IgM negative for hepatitis B, Oropouche virus, Mayaro virus, Hantavirus, Venezuelan equine encephalitis, and rickettsiae.
- ELISA negative for HIV.

As part of the autopsy, a macroscopic anatomopathological analysis was performed and specimens were obtained for the microscopic anatomopathological and toxicological analyses, which were performed by the INS and Central Morgue. Results were as follows:

- Candida in the pharynx, esophagus, and upper portion of the stomach.
- Liver: examination with hematoxylin-eosin showed preserved liver structure, serious liver microvacuolization with midzonal predominance, limited lymphoplasmocytic inflammatory component with predominance of portal spaces, and liver steatosis.
- Kidney: tubular necrosis.
- Brain: cerebral edema.
- Immunohistochemistry: positive for yellow fever antigen in the brain, kidney, liver, and lung.
1. Does the patient meet the case definition for VTD? Justify your response.

The patient meets three or more major criteria of the case definition provided by the Brighton Collaboration. These include hepatic criteria: elevated bilirubin and transaminases; renal criteria: elevated creatinine; respiratory criteria: need for mechanical ventilation; and platelet disorder: thrombocytopenia and coagulopathy (elevated INR). Thus, the case qualifies as a level 1 case of viscerotropic disease (Annex 3).

2. What specimens should be obtained during the autopsy?

Ideally, duplicate tissue specimens should be procured. One set should remain in the country, while the other may be sent to the regional reference laboratory or to an international laboratory to complete tests that cannot be performed in the country. Specimens should be collected from all organs, especially the liver, lungs, brain, kidney, spleen, thymus, lymph nodes, bone marrow, and skin.

Each specimen set should contain four 3-4 cm³ specimens of each organ (Table 5). Brain specimens should contain meninges, pulmonary specimens should include all five lobes of the lung, and kidney specimens should include specimens from the suprarenal glands. In the past, some specimens had to be kept frozen, while others were preserved in formalin. Today, advances in polymerase chain reaction (PCR) processing techniques have enabled testing of formalin fixed tissues. Nevertheless, frozen specimens are required for viral isolation and a full study of viral RNA, even though these results can also be obtained, though less frequently, from serum (2.5 mL) or whole blood specimens in vials with anticoagulant (5 mL). If an autopsy is not performed, a liver tissue specimen should be secured by whatever means necessary. A subcutaneous liver biopsy can be very helpful in this respect.

**TABLE 5: SPECIMENS THAT SHOULD BE OBTAINED DURING AN AUTOPSY**

<table>
<thead>
<tr>
<th>SPECIMEN TYPE</th>
<th>QUANTITY</th>
<th>PRESERVATION</th>
<th>DIAGNOSTIC TEST</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver</td>
<td></td>
<td>Tissue frozen at -80°C</td>
<td>Viral culture and RNA detection.</td>
</tr>
<tr>
<td>Brain with meninges</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kidney</td>
<td></td>
<td>Tissue fixed in 10% formalin.</td>
<td>RNA detection within 24 hours.</td>
</tr>
<tr>
<td>Spleen</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pancreas</td>
<td></td>
<td>Tissue fixed in paraffin.</td>
<td>Histopathology, immunohistochemistry.</td>
</tr>
<tr>
<td>Heart</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lungs</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adrenal glands</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thymus</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
TABLE 5. (Continued).

<table>
<thead>
<tr>
<th>SPECIMEN TYPE</th>
<th>QUANTITY</th>
<th>PRESERVATION</th>
<th>DIAGNOSTIC TEST</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood</td>
<td>One 2 mL specimen of blood.</td>
<td>Frozen specimen of blood and serum at -80°C.</td>
<td>Viral culture, detection of RNA, and viral load.</td>
</tr>
<tr>
<td></td>
<td>Two 5 mL specimens of serum.</td>
<td>A serum specimen, not frozen at +4°C.</td>
<td>Serological tests.</td>
</tr>
<tr>
<td>Lymph nodes</td>
<td>Two nodes near vaccination site.</td>
<td>Frozen at -80°C or fixed in formalin at +4°C.</td>
<td>Detection of viral RNA.</td>
</tr>
</tbody>
</table>


Given the tests results, increasing media pressure, and limited information on viscerotropic disease, the MoH requested support from PAHO, the United States Naval Medical Research Unit 6 (NMRU-6, known as NMRCD at time of the outbreak) in Peru, and the U.S. Centers for Disease Prevention and Control (CDC). Serum and tissue specimens were subsequently provided to NMRU-6 and the CDC. Final results from the CDC became available in February 2008. These included: viral culture: negative; RT-PCR and Q-PCR: positive in the serum, urine, lung, kidney, liver, and brain, with viral concentrations ranging from $2.4 \times 10^2$ to $1.1 \times 10^4$ PFUeq/mL; and immunohistochemistry: positive and limited in the lung, liver, and kidney.

Additionally, anatomo-pathology tests performed by the INS and the Central Morgue yielded the following results:

- **Thyroid**: follicular thyroid neoplasm and chronic thyroiditis.
- **Liver**: hematoxylin-eosin examination showed midzonal necrosis, limited lymphoplasmocytic inflammatory component with predominance of portal spaces, serious liver microvacuolization, and intracellular cholestasis.
- **Kidney**: vascular congestion, hemorrhage foci, and tubular necrosis.
- **Lung**: edema, congestion, and vascular expansion.
- **Brain**: cerebral edema and discreet gliosis.
- **Immunohistochemistry for yellow fever**: positive (polyclonal and monoclonal antibodies) in the lung, liver, kidneys, and brain.
- **Toxicology tests**: negative.
3. Based on this information, how do you believe the ESAVI Committee classified the case? The case was definitively classified as associated with the vaccine (YEL-AVD) because it met the following criteria in the case definition: 1) liver pathology findings compatible with yellow fever; 2) immunohistochemistry positive for yellow fever in organs, including the liver; 3) no precedent of having been in an area endemic for yellow fever; and 4) culture and amplification of the genetic material of the vaccine virus in tissue (Annex 4).

ABOUT THE TERM YEL-AVD

As defined by the Brighton Collaboration, this case would be classified as viscerotropic disease associated with the yellow fever vaccine. In 2012, the group decided to stop using the term YEL-AVD, reasoning that clinical diagnosis of VTD and association with the yellow fever vaccine require separate analyses. As a result, the diagnosis of VTD is first made and the case’s relationship to vaccination is then evaluated in a separate step using a causality algorithm.

CASE III

TIME AVAILABLE: 90 MINUTES

A. STATE OF EMERGENCY SERVICES, THE HEALTH SYSTEM, AND THE PATIENT REFERRAL SYSTEM

The earthquake that struck Peru in August 2007 rendered useless many of the health centers in Ica. To make matters worse, the death of two patients following vaccination against yellow fever generated considerable concern among the local population. As a result, a large number of vaccinated individuals sought care from emergency departments that already faced high demand and logistical constraints. These circumstances, as well as limited knowledge on VTD, impaired the ability of healthcare workers to accurately diagnose and treat the disease.

To improve care, health officials developed a provisional case definition of suspected VTD. Patients meeting the definition were required to be observed for at least 12 hours in the hospital and for an additional 72 hours in their homes under the responsibility of the Ica DIRESA. For all cases, patient vaccination histories had to be obtained. To that end, the immunization program created a database to monitor all individuals vaccinated during the campaign. In a 10-day period, 139 people sought care for fever, diarrhea, nausea, and other symptoms similar to those of the deceased patients.
1. Develop a case definition for suspected VTD associated with the yellow fever vaccine that may be used to rapidly detect cases in a situation similar to the one described above.

The definition of a suspected case should serve as an initial screening of patients and should be based primarily on clinical rather than laboratory criteria. The clinical symptoms of VTD cases already reported provide a starting point for developing this definition. In Peru, a suspected case might have been defined as one presenting fever, jaundice, and hemorrhage or gastrointestinal symptoms, with a history of vaccination in the previous 30 days.

2. What other measures could be taken to decrease demand of health services?

A more efficient referral and counter-referral system could be implemented. It would also be helpful to establish triage areas in health centers, train personnel to better identify cases, and designate some health centers to care exclusively for patients who meet the suspected case definition. In addition to decreasing demand in many healthcare facilities, this approach would make the capture and follow-up of suspected cases more efficient. Lastly, community members should be frequently reminded of where to seek services if they have symptoms suggestive of viscerotropic disease.

B. Third case: A vaccinated 79-year-old man

On 10 October 2007, a new ESAVI case was reported in a 79-year-old man, who had been transferred to the Ica Regional Hospital from a hospital in Nazca (a province south of Ica). The patient presented abdominal pain syndrome, acute diarrheal disease, and hypovolemic shock. He was diagnosed with congestive heart failure, acute dysenteric diarrheal disease, and shock of undetermined etiology. Despite not being part of the campaign’s target age group, the patient had received yellow fever vaccine on 1 October. While hospitalized, the patient deteriorated rapidly. On 11 October, he was placed on mechanical ventilation, but, despite the efforts of health workers, he died the same day. The patient had a history of allergy to sulfa drugs and had undergone prostatectomy in October 2002.

The results of the patient’s tests at the Ica hospital were: AST (SGOT): 416 U/L; ALT (SGPT): 231 U/L; total bilirubin: 2.93 mg/dL; direct bilirubin: 2.25 mg/dL; and prostatic specific antigen: 76 ng/mL. The results of RT-PCR and Q-PCR showed partial viral sequences 100% homologous to the 17DD vaccine virus in the serum, lung, liver, kidney, and brain, with viral concentrations ranging from $1.9 \times 10^2$ to $3.5 \times 10^4$ PFUeq/mL.
1. How would you classify this case? Justify your answer.
   This is unquestionably a case of VTD associated with yellow fever vaccine. Elevated viral concentrations were found in the patient and viral RNA was amplified in tissue specimens from several of his organs (Annexes 3-5).

2. Do aspects of this case suggest a programmatic error? Explain.
   Yes. Vaccination of the patient did not comply with indications of the yellow fever vaccination campaign (target population aged 15-59). To reduce the risk of programmatic errors, some countries use a brief pre-vaccination screening to ask patients questions regarding pregnancy and their history of chronic diseases, drug use, and previous reactions to vaccines.

3. Who else may be at risk for developing VTD associated with yellow fever vaccine? How long should follow-up activities last?
   The risk of additional cases should be evaluated based on the type of vaccine used and other features of vaccination. Everyone vaccinated from the same vial or with the same vaccine batch may be at risk. The at-risk population may also include anyone vaccinated on the same day, at the same health center, or by the same team as the individual who developed VTD.

   It is necessary to determine what risk factors exist for VTD following yellow fever vaccination (contraindications and vaccine precautions) and what clinical symptoms of VTD have already presented. From the standpoint of fieldwork, the follow-up of all individuals who have symptoms suggestive of VTD and of those who received vaccine from the same vial or at the same health center should be prioritized.

   Additionally, due to the possibility of cases of neurologic disease following yellow fever vaccination in infants breastfeeding from vaccinated mothers, these children must be monitored.

   Follow-up should continue throughout the period in which cases of VTD associated with yellow fever vaccine are expected to appear. Symptoms usually manifest 1-8 days following receipt of the vaccine, although in one case symptoms appeared 18 days later. The time leading up to death has ranged from 7 to 30 days (12). Depending on the type of follow-up implemented, follow-up may last 8, 18, or 30 days.

C. IDENTIFICATION AND FOLLOW-UP OF OTHER PERSONS AT RISK

To identify and follow-up with other patients at risk, investigators reviewed lists of patients who had received vaccine from the same vial as the deceased patient in order to determine if they had developed an ESAVI. After a few days, most vaccinated individuals had been located, and it was reported that none had developed important clinical symptoms.
Since the first case of VTD following yellow fever vaccination occurred in a university student who had been vaccinated with many of her classmates, the entire cohort of vaccinated students was evaluated. Although some experienced mild or moderate events (among them the boyfriend of the deceased patient), none presented significant clinical symptoms. Additionally, no one vaccinated at the same facility as the known cases reported significant adverse events.

1. What strategies would you recommend to detect other cases of VTD associated with yellow fever vaccine?

   The definition of a suspected case should be disseminated to all health centers and active institutional case finding must be performed. Due to the disease’s severity, individuals with viscerotropic disease would have presented to a health center, particularly since the population is largely urban. Case finding at the community level would be very difficult due to the number of people vaccinated.

2. What could be done to capture vaccinated individuals in a timely manner? What minimum variables should be collected?

   Ideally, an electronic registry of individual cases would allow investigators to rapidly verify patient’s vaccination histories and the lot of vaccine they had received. Information on affiliation, such as the patient’s name, age, and address, should be included, as should the clinical symptoms (<10 days for VTD and >30 days for neurologic disease) required to identify cases. If an electronic registry is unavailable, a manual registry should be used. In Peru, an electronic registry was created based on existing manual registries in order to facilitate the identification of information related to vaccination history.

3. How would you structure an active case-finding strategy to detect VTD cases, taking into account the largely urban population?

   First, the time period and areas included in the search would need to be determined. In this situation, investigators should review all cases seen in emergency centers within 30 days of the campaign’s end as well as hospital admissions made during this period. Since the number of cases would be large, a sensitive definition of suspected VTD cases would be needed to conduct an initial screening. For cases meeting these criteria, clinical symptoms should then be reviewed in greater detail to determine whether they meet the case definition. For cases meeting the case definition, the patient’s clinical history should be obtained to determine if the illness is associated with yellow fever vaccination. Municipal death registries should also be reviewed.

D. RETROSPECTIVE EPIDEMIOLOGICAL INVESTIGATION

   The surveillance team proposed implementing an active, institutional, and retrospective case-finding system in order to determine whether other cases of VTD associated with yellow fever vaccine had occurred. Since the disease has a high fatality rate and clinical symptoms requiring hospitalization, investigators reviewed emergency service records and discharge databases of all Ica hospitals that cared for patients following the earthquake. Municipal death records were also evaluated.
To identify potential cases, both searches used a time period stretching from the start of the campaign to 30 days after its completion. Reviewing a large number of registries would have required a multi-stage case strategy, and in an initial, rapid search of cases, suspected patients with VTD would have been identified by their clinical symptoms. Due to limited knowledge of the disease, however, it was not clear which cases could be classified as probable or confirmed. For this reason, the country requested support from PAHO and the CDC in implementing a field investigation.

E. METHODS AND RESULTS OF THE INVESTIGATION

The PAHO/CDC team of experts arrived in Lima at the end of October. Shortly thereafter, the team analyzed the cases and reviewed the outbreak investigation. The team decided to use the following definitions:

a. **Suspected case:**
   
   Any patient presenting fever >38°C (or who feels feverish) for >24 hours and one or more of the following signs or symptoms:
   
   - Intense headache.
   - Sensory disturbance.
   - Tonic-clonic seizure.
   - Nausea, vomiting.
   - Watery stools.
   - Myalgia lasting >24 hours.
   - Arthralgia lasting >24 hours.
   - Increased respiratory rate (>20 breaths/min).

b. **Probable case:**
   
   Any suspected case that was vaccinated less than 15 days before the onset of symptoms, with no evidence of other etiologies to explain clinical symptoms, and with fever and one or more of the following signs and symptoms:

   - Nausea, vomiting, malaise, watery stools, myalgia, arthralgia, dyspnea, and one or more of the following:
     
     - Elevated serum transaminases ≥3 times normal level.
     - Elevated total serum bilirubin ≥1.5 times normal level.
     - Serum creatinine ≥1.5 times normal level.
     - Total CPK: ≥5 times normal level.
     - Thrombocytopenia (blood platelets <100,000/mL).
     - Myocarditis (compatible abnormalities detected by electrocardiogram, echocardiogram, or cardiac enzymatic changes, or inflammation confirmed by cardiac tissue biopsy).
     - Elevation of prothrombin time or activated partial thromboplastin time or elevated INR.
     - Histopathology compatible with yellow fever (e.g., midzonal hepatic necrosis or Councilman's bodies).
c. **Confirmed case:**

Any probable case that has one or more of the following:

- Isolation in blood of 17D\(^2\) yellow fever virus >7 days after vaccination, and/or through PCR >11 days following vaccination.
- Specific antigen for yellow fever in visceral tissue demonstrated by immunohistochemistry.
- Isolation of 17D\(^2\) yellow fever virus in visceral tissue.
- Amplification of the 17D\(^2\) yellow fever RNA virus in visceral tissue.

Presently, the case definitions developed by the Brighton Collaboration are recommended for making case classifications (Annexes 3-5).

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1. **Compare the case definition used in Peru with those developed by the Brighton Collaboration Working Group.**

   The Brighton Collaboration’s definition of VTD relies largely on laboratory rather than clinical criteria. Once VTD is diagnosed, the case’s association with yellow fever vaccine is evaluated in a separate step using a causality algorithm. In Peru, the research team used a definition based on the disease’s clinical characteristics and evaluated laboratory evidence at a later date. The team’s definition was based on guidelines employed by the CDC and Brazil, which are oriented more toward surveillance and the epidemiological investigation of VTD following yellow fever vaccination.

2. **Reviewing the causality criteria below, which are consistent with VTD following yellow fever vaccination?**

   The British epidemiologist, Austin Bradford Hill, proposed the causality criteria that are most commonly accepted today in his article about disease and the environment (16). They are:

   Those pertaining to the study itself (internal validity):
   
   a. **Strength of association:** (the stronger the relationship between two variables, the greater the likelihood that a relationship exists). The small number of VTD cases following yellow fever vaccination reported to date makes it impossible to determine the strength of association.
   
   b. **Temporality:** (cause should precede effect). Vaccination against yellow fever should precede the onset of disease symptoms.
   
   c. **Biological gradient:** (the greater the time and/or exposure to the causal factor, the greater the risk of the disease). A biological gradient does not appear to exist because cases only occur following the first dose.

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\(^2\) Presence of virus 17D (and of all 17D vaccine-derived viruses) confirmed by nucleotide sequencing.
Scientific plausibility:

d. Consistency: (the results of a study should be consistent and reproducible by any researcher anywhere). In this case, the use of standardized case definitions by different researchers investigating different areas and populations supports consistency.

e. Specificity of the cause: (one cause leads to one effect). Establishing specificity increases the likelihood of causation; in this case, the yellow fever virus can produce VTD, both in the wild in people with adequate immune systems and through an attenuated vaccine virus in an immunocompromised person.

f. Analogy: (reasoning by analogy applies proven causal associations with causal structures similar to the problem studied; in so doing, analogies may suggest possible causal structures to explain the problem under study). The behavior of the virus in other vaccines might provide such a connection, even though VTD linked to other vaccines has been found in far fewer persons and predominantly in those with immune disorders.

g. Biological plausibility: (the suggested causal relationship must be consistent with the accepted scientific principles of the time—that is, a causal relationship seems more plausible if we understand its pathogenic mechanism). The vaccine virus contains a live attenuated virus and is considered viscerotropic.

h. Consistency: (the interpretation of cause and effect cannot conflict with the behavior of the illness or injury. This criterion combines aspects of consistency and biological plausibility). The principal findings of these case studies are consistent with the disease caused by wild yellow fever virus and do not conflict with current scientific knowledge.

i. Experimental evidence (it is not always possible to perform a study, but such studies are the strongest proof of causality). In this case, experimental evidence does not exist and it would unethical to obtain it. Nevertheless, the virus was identified in the organs of vaccinated individuals.

The field investigation was conducted on 13-23 November 2007 and included all five provinces of Ica: Chincha, Ica, Nazca, Palpates, and Pisco. Investigators sought medical records for all individuals who received care or died between 23 September and 6 November. A total of 28,788 medical records were reviewed, among which 311 cases met the criteria for suspected cases. Of those, five were classified as probable cases of VTD following yellow fever vaccination. Four of these had been captured by the ESAVI surveillance system and associated with yellow fever vaccination. The fifth was discarded because the patient had not received yellow fever vaccination. No suspected cases of neurologic diseases associated with the yellow fever vaccine were identified.

3. If you find yourself in a similar situation in your country, how would you conduct a prospective search for cases?

The capture of prospective cases requires a slightly different approach. Investigators must first identify locations where cases could be captured and determine the severity
of expected cases (moderate, severe, etc.). To this end, the suspected case definition should take into account the clinical features of cases already reported. Subsequently, all recommended specimens should be obtained from any suspected case. The advantage of this approach is that all cases, including those that are moderate and non-fatal, could be studied. On the other hand, the entire population could not be monitored, which might reduce the representativity of cases.

TIME AVAILABLE: 90 MINUTES

A. FOURTH CASE: A PATIENT WITH AN AUTOIMMUNE DISORDER

On 23 October 2007, a case was reported in a 49-year-old female patient at the emergency room of the Social Security Hospital of Chincha. The patient’s symptoms included intense headache, oliguresis, generalized edema, mild jaundice, and signs of dehydration. She had a history of hypertension, rheumatoid arthritis, systemic lupus erythematosus, and chronic renal failure. She also had difficulty moving due to stroke sequelae.

The patient had been vaccinated on 24 September at the insistence of a nurse in the Ica region, despite the patient’s and her family members’ initial reluctance. An analysis of the patient’s vaccination history revealed she had received a vaccine from lot 121Z. Three days prior to vaccination, the patient had developed pain in her hip that hindered her ability to walk. When the pain increased, she went to the Chincha hospital on 28 September, where the attending physician prescribed treatment with methotrexate and tenoxicam, both in tablet form, as well as diclofenac and dexamethasone in ampoules. Days later, the patient experienced various episodes of melena and three episodes of vaginal bleeding. On 12 October, she was again seen at the Social Security Hospital for intense headache and malaise. This was the patient’s last contact with the health care system prior to 23 October, and she was prescribed analgesics and prednisone.

On 24 October, the patient was transferred to the Rebagliati National Hospital in Lima. She arrived in poor health and was diagnosed with metabolic encephalopathy, metabolic acidosis, decompensated chronic renal insufficiency, electrolyte disorder, icteric syndrome of unknown etiology, systemic lupus erythematosus, and exacerbation of lupus activity. Despite receiving treatment in the emergency intensive care unit, the patient died on the afternoon of 24 October.

Laboratory tests performed at the Chincha hospital showed AST (SGOT): 91 U/L; ALT (SGPT): 128 U/L; total bilirubin: 5.2 mg/dL; direct bilirubin: 4.2 mg/dL; and indirect bilirubin: 1.0 mg/dL. Yellow fever immunohistochemistry (monoclonal and polyclonal antibodies) was slightly positive in the liver and kidney. Additionally, results from the RT-PCR and Q-PCR were positive in the kidney and slightly positive in the liver, with a viral concentration of $1 \times 10^4$ PFUeq/mL in the kidney.
1. **Would you have vaccinated this patient? Explain.**

   A patient with an autoimmune disorder should not be vaccinated. Prior to vaccination, individuals should be asked about their health and allergies. If a person has a chronic disease, a compromised immune system, or is undergoing potentially immunosuppressive treatment, he or she should not be vaccinated without first being evaluated by a physician. Other situations requiring evaluation include pregnancy, allergies to any vaccine component, and previous allergic reactions to immunizations. In this patient’s case, the correct assessment of her history would have led healthcare professionals to postpone vaccination until an adequate medical assessment was performed (to evaluate potential relapse of previous diseases).

2. **What are the indications, contraindications, and precautions of yellow fever vaccine (2,17)?**

   **Indications:**
   - Yellow fever vaccine should be administered to children aged 12 months as part of the routine vaccination program.
   - In the case of outbreaks, the vaccine can be administered as early as age 6 months.
   - Per International Health Regulations, travelers to enzootic areas should be vaccinated every 10 years for the purpose of validating the International Yellow Fever Vaccination Certificate. However, routine revaccination of residents in enzootic areas is not necessary.
   - Laboratory staff that may be exposed to yellow fever virus should be vaccinated.

   **Contraindications:**
   - People with acute febrile diseases, whose general health status is compromised.
   - People with a history of hypersensitivity to chicken eggs and their derivatives.
   - Pregnant women, except in an epidemiological emergency and following recommendations of health authorities.
   - People with severely compromised immune systems, including those with:
     - Acquired immunodeficiency syndrome (AIDS) or C4+ cell counts <200 cell/mm³.
     - Diseases involving the thymus.
     - Malignant neoplasms treated with chemotherapy.
     - Recent transplants of stem cells.
     - Intake or consumption of medications with immunosuppressive or immuno-compromising properties (high doses of cortical steroids, diluents, anti-metabolites, interferon-alpha inhibitors).
     - Recent or ongoing radiotherapy.
   - Infants aged <6 months.
Precautions:

- The yellow fever vaccine can be administered to patients infected with human immunodeficiency virus (HIV), but only if they are asymptomatic or as per medical guidelines.
- For people aged <60 years, it is recommended that the epidemiological risk of contracting the disease be evaluated individually and be weighed against the probability of an adverse event. A similar risk assessment should be conducted in deciding to vaccinate breastfeeding women who are traveling to enzootic areas and have not been previously vaccinated.

3. Could this case be considered the result of a programmatic error? Support your answer.

Yes. A programmatic error occurred because a person with an immunosuppressive disease and with a chronic disease and symptoms suggestive of a relapse should not be vaccinated until evaluated by a specialist. The health worker should have inquired about the patient’s medical history prior to vaccination in order to become aware of the use of any immunosuppressive drugs and thereby of any contraindications for attenuated vaccines.

B. Risk Communication

Shortly before the international team’s arrival, the parents of a 1-year-old girl who died in Lima following yellow fever vaccination confronted ministry spokespersons. While it was later determined that the cause of the child’s death was not VTD, the time required to complete the investigation and the MoH’s poor relationship with the press meant that the true cause of the child’s death was not widely reported. The child’s case was the last reported serious ESAVI following yellow fever vaccination, and it was officially classified as a coincidental event after the patient’s death from sepsis and hemolytic anemia.

The PAHO/CDC team of technical experts arrived in the country at the end of October and remained until mid-November (Annex 6). The team supported the MoH in two fundamental ways: 1) by participating in field investigations and helping to classify suspected cases of VTD associated with yellow fever vaccine and 2) by responding to media inquiries, reinforcing the messages of ministry spokespersons, and thereby improving communication with the public.

1. What traits should a spokesperson possess?

Spokespersons should:

- Be credible, tell the truth, not deny everything that is asked, and express themselves well using clear language.
- Clearly articulate messages pertaining to immunization.
• Maintain a close relationship with the public, be empathetic, understand the public’s concerns in order to achieve greater understanding, and be capable of handling problems and difficult questions.

• Be capable of interacting with diverse groups and media sources and be flexible and able to anticipate problems.

• Be well informed of EAVIs in general and specific cases in particular.

2. How should a spokesperson prepare for an interview with mass media outlets?

A spokesperson’s preparation should include an understanding of the media—its dynamics, work processes, and editorial approaches—as well as interview techniques. Interviews are conversations and can therefore be guided to the extent that spokespersons are familiar with key messages and the types of questions that will be asked. This skill is the most important aspect of the spokesperson’s preparation and should be oriented to putting forth concepts and messages that are part of the MoH’s communication strategy.

C. EVALUATION OF THE VACCINE LOT

The vaccine administered in Peru was donated by the government of Bolivia, which, in turn, had received it as a donation from the Bolivarian Republic of Venezuela. In Venezuela, 73,000 doses of lot 121Z had been administered, with no reported serious EAVI cases. In Bolivia, vaccines from lot 121Z had not been administered. However, 23,000 doses were stored in the country’s central warehouse.

Test results on the potency and quality of the vaccine performed by the INS of Peru showed no abnormalities. Production procedures were evaluated at the manufacturing laboratory, even though the WHO had prequalified the vaccine. Using a validated assay, a laboratory contracted by WHO analyzed retention samples from the manufacturer and samples of vaccine lot 121Z used in Ica at the end of its storage period. No differences in vaccine potency were observed between distributed samples and those obtained at the end of the storage period. These results were consistent with those from the manufacturer, INS, and CDC, which showed no reductions in the dose or potency of the vaccine lot.

The manufacturer also performed tests to detect bacteria, fungi, mycoplasma, and several known polluting avian viral agents (although not mammalian viruses) in the vaccine’s preparation products. Results were negative.

All other evaluations also yielded satisfactory results, and no defects in product quality were found to explain the four cases in Ica.

D. VIROLOGICAL EVALUATION

Studies revealed high viremia, elevated viral load, and broad viral tissue distribution (including in many vital organs) in those who had died following vaccination. These results are consistent with previous reports of fatal cases of VTD following yellow fever vaccination.
Stability studies of lot 121Z indicated there had not been a significant loss of viral viability, despite the fact that the vaccine was nearing its expiration date (October 2007) and had been transported on multiple occasions among countries. Nor was a variant of the virus found in tissue specimens of those who had died following receipt of lot 121Z. The consensus nucleotide sequence of the 17DD vaccine virus isolated was essentially identical to the sequence that the laboratory has used for vaccine production since 1984. In short, laboratory studies did not reveal changes in the vaccine virus that could explain the ESAVIS.

1. Do you believe the following hypothesis can be rejected: lot 121Z used in Ica was inherently “more dangerous than other lots?” Please provide arguments supporting and refuting the hypothesis.

Contrary to the hypothesis:

- All analyses related to aspects of vaccine production showed appropriate practices.
- The vaccine virus was genetically indistinguishable from the original virus, suggesting that there were no differences between the vaccine virus and those used in other lots (e.g., 123Z).
- Studies showed that the vaccine was safe and had normal potency.
- The incidence among individuals who received the vaccine from lot 121Z was not statistically greater \((p<0.05)\) than the incidence observed among people who received a vaccine from a different lot (123Z) during the mass campaign.

Consistent with the hypothesis:

- The tests did not include animal models that would have more accurately mimicked the infection process in those affected.
CONCLUSIONS

In conjunction with national authorities, a panel of international experts (Annex 6) was formed to analyze the reported cases and determine what causal relationship existed between the cases and the yellow fever vaccine. The panel reached the following conclusions (18):

- The occurrence of cases of viscerotropic disease associated with yellow fever vaccine clustered in time and space with a single lot of yellow fever vaccine (050VFA-12Z, Bio-Manguinhos, Brazil) had no precedent.

- The incidence of VTD following yellow fever vaccination was 7.9-11.7 cases per 100,000 doses administered—over 20 times the incidence previously notified. The incidence among individuals who received the vaccine from lot 121Z was not statistically greater (p<0.05) than the incidence observed among people who received a vaccine from a different lot (123Z) during the mass campaign.3

- Clinical, virological, and pathological evidence was sufficient to classify four cases as VTD following yellow fever vaccination (all fatal) and one case as probable (survived). The cause of death was a generalized infection with the 17DD vaccine virus, probably associated with severe immunity response syndrome.

- The consensus sequence of the genome of vaccine lot 121Z, in both the parental seed lot and in viral RNA collected in patient specimens, indicated there were no genetic changes in the vaccine virus that could have caused the cases of VTD associated with yellow fever vaccine.

- No sign of virological or other factors related to the vaccine production process were found that pointed to anomalies inherent to vaccine lot 121Z that could explain the higher frequency of VTD following yellow fever vaccination among individuals vaccinated from the lot.

- Peru’s surveillance system for detecting adverse events proved sensitive. No other fatal cases of VTD following yellow fever vaccination in Ica were detected after the retrospective review of hospital and death records.

- Factors related to the host may have increased the risk of severe infection with viral strain 17DD. These include old age in one case and autoimmune disease (lupus) in another. In two cases, potentially immunosuppressive medication was administered following vaccination. Prior to these cases, autoimmune disease was not recognized as a risk factor for VTD following yellow fever vaccination. However, greater attention should be paid to autoimmune diseases in the future. The apparent association of diarrhea in cases of VTD suggests the direct participation of the yellow fever virus in gastrointestinal tract infection, or as an aggregate cofactor with another infectious agent (Annex 7).

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3 Given an expected rate of 0.4 cases of viscerotropic disease associated with yellow fever vaccine per 100,000 vaccine doses, it is extremely unlikely that four cases would occur after 42,742 doses of lot 121Z by chance alone (p<0.001) (15).
In this outbreak, factors related to the population were probably significant. Among these are the wide use of yellow fever vaccine among adults (many with risk factors, such as old age) in a non-endemic area and a population that had not been previously vaccinated against yellow fever and thus lacked protective immunity against the disease.

Today, the search continues for cases of viscerotropic disease following yellow fever vaccination or factors that increase predisposition to this adverse event. International research protocols have been developed, as have case definitions to identify and investigate suspected cases. The comprehensive study of all cases has the potential to make a great difference in our understanding of this adverse event and the strategies needed to prevent it.
REFERENCES


ANNEX 1: PRE- AND POST-TESTS ON YELLOW FEVER CASE STUDIES

Pseudonym: ________________________________

1. What countries are endemic for yellow fever in the Region of the Americas?
   a. Argentina, Chile, and Paraguay.
   b. Brazil, Bolivia, and Venezuela.
   d. Ecuador, French Guiana, and Peru.
   e. b and d are correct.

2. Which of the following is a **contraindication** against yellow fever vaccination?
   a. Laboratory personnel exposed to the yellow fever virus.
   b. Pregnant women, except in epidemiological emergencies when the risk of yellow fever is very high.
   c. Women who are breastfeeding and will be traveling to endemic areas.
   d. b and c are correct.

3. Which of the following is a **precaution** against yellow fever vaccination?
   a. People of any age with a thymus disorder.
   b. Asymptomatic patients infected with HIV.
   c. People with severe febrile diseases compromising their general health status.
   d. Women who are breastfeeding and will be traveling to endemic areas.
   e. b and d are correct.

4. Which of the following is/are not a serious ESAVI associated with the yellow fever vaccine?
   a. Viscerotropic disease.
   b. Guillain-Barré Syndrome (GBS).
   c. Fever, chills, headache, myalgia.
   d. Hypersensitivity reactions.
   e. a and d are correct.

5. Why should this cluster of cases of viscerotropic disease associated with vaccination against yellow fever be reported immediately to the World Health Organization (WHO), as stipulated by the International Health Regulations (IHR)?
   a. The event is unusual.
   b. The number of cases is higher than expected, since the incidence was 1 case per 42,742 doses.
   c. The vaccine is being massively administered to a susceptible population.
   d. b and c are correct.
   e. a and b are correct.
6. Which of the following criteria would you consider in evaluating a case of viscerotropic disease?
   b. Clinical: jaundice and hemorrhage.
   c. Timing post-vaccination: the patient was vaccinated 7-30 days after the onset of symptoms.
   d. b and c are correct.
   e. All of the above are correct.

7. Which of the following is a major criterion used in the definition of viscerotropic disease?
   a. Jaundice.
   b. Petechiae or purpura.
   c. Melena.
   d. Need for mechanical ventilation.
   e. c and d are correct.

8. Which of the following criteria must be met in order to establish definitive causality between viscerotropic disease and the yellow fever vaccine?
   a. Yellow fever 17D virus concentration in blood ≥2 log_{10} PFU/mL, but <3 log_{10} PFU/mL on any day 1-10 days post-vaccination.
   b. Isolation of yellow fever 17D virus from blood >18 days post-vaccination.
   c. Isolation of yellow fever 17D virus from blood >10 days post-vaccination.
   d. a and b are correct.
   e. All of the above are correct.

9. During an ESAsVi investigation, it is recommended to request the reference laboratory to conduct potency tests of the vaccine in question. What is your opinion of this recommendation?
   a. It is always correct.
   b. It is never correct.
   c. It depends on the results of the investigation.
   d. It depends on the results of the sterility, toxicity, and identity tests.

10. During the investigation of a serious ESAsVi that was widely publicized in the media, it was not possible to determine the cause of the event. What actions should health workers take?
    a. Report the findings of the investigation to interested parties.
    b. Clearly communicate the results and include mass media outlets.
    c. Do not make any statement, since no conclusion was reached.
    d. Avoid mentioning the issue with the media in order to maintain the public’s trust in the immunization program.
    e. a and b are correct.
Answer key

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<th>QUESTION</th>
<th>CORRECT ANSWER</th>
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ANNEX 2: RECOMMENDED LABORATORY TESTS FOR SERIOUS ESAVIS FOLLOWING YELLOW FEVER VACCINATION

A. All serious ESAVs
First set of laboratory tests

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<th>SPECIMEN</th>
<th>LABORATORY TEST</th>
<th>CLINICAL REASONING</th>
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<td>Blood</td>
<td>Complete blood count and platelet count</td>
<td>Establish clinical baseline, rule out bacterial infection</td>
</tr>
<tr>
<td>Blood</td>
<td>Thick blood film</td>
<td>Rule out malaria, infection by <em>Borrelia</em></td>
</tr>
<tr>
<td>Urine</td>
<td>Urinalysis</td>
<td>Evaluate proteinuria, bleeding</td>
</tr>
<tr>
<td>Cerebrospinal fluid</td>
<td>Turbidity</td>
<td>Establish clinical baseline, rule out bacterial infection</td>
</tr>
<tr>
<td></td>
<td>Cell count</td>
<td>Establish clinical baseline, rule out bacterial infection</td>
</tr>
<tr>
<td></td>
<td>Protein</td>
<td>Rule out meningitis</td>
</tr>
</tbody>
</table>

B1. Suspected cases of serious viscerotropic disease
Second set of laboratory tests for clinical evaluation and differential diagnosis

<table>
<thead>
<tr>
<th>SPECIMEN</th>
<th>LABORATORY TEST</th>
<th>CLINICAL REASONING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood</td>
<td>Complete blood count and platelet count</td>
<td>Rule out other etiologies</td>
</tr>
<tr>
<td></td>
<td>Blood culture</td>
<td>Rule out bacteremia</td>
</tr>
<tr>
<td>Serum</td>
<td>Transaminases</td>
<td>Evaluate liver enzymes and function</td>
</tr>
<tr>
<td></td>
<td>Direct and indirect bilirubin</td>
<td>Evaluate liver enzymes and function</td>
</tr>
<tr>
<td></td>
<td>Alkaline phosphatase</td>
<td>Evaluate liver enzymes and function</td>
</tr>
<tr>
<td></td>
<td>Tests for Hepatitis B and C</td>
<td>Rule out other hepatitis viruses</td>
</tr>
<tr>
<td></td>
<td>Urea nitrogen</td>
<td>Evaluate renal function</td>
</tr>
<tr>
<td></td>
<td>Creatinine</td>
<td>Evaluate renal function</td>
</tr>
<tr>
<td></td>
<td>Amylase</td>
<td>Evaluate pancreas function</td>
</tr>
<tr>
<td></td>
<td>Creatine phosphokinase</td>
<td>Rhabdomyolysis evaluation</td>
</tr>
<tr>
<td></td>
<td>Partial prothrombin time</td>
<td>Coagulation panel</td>
</tr>
<tr>
<td></td>
<td>Partial thromboplastin time</td>
<td></td>
</tr>
<tr>
<td>Urine</td>
<td>Urinalysis</td>
<td>Rhabdomyolysis evaluation</td>
</tr>
<tr>
<td></td>
<td>Urea antigen</td>
<td>Rule out leptospirosis</td>
</tr>
<tr>
<td>Saliva</td>
<td>PCR</td>
<td>Detect yellow fever virus</td>
</tr>
<tr>
<td>Feces</td>
<td>PCR</td>
<td>Detect yellow fever virus</td>
</tr>
<tr>
<td>Other fluids</td>
<td>PCR</td>
<td>Detect yellow fever virus</td>
</tr>
<tr>
<td>Serum</td>
<td>IgM, IgG antigens against yellow fever (acute and convalescent)</td>
<td>Confirm infection by or vaccination against yellow fever</td>
</tr>
<tr>
<td></td>
<td>PCR/viral culture</td>
<td>Rule out wild-type yellow fever</td>
</tr>
</tbody>
</table>
### B2. Suspected cases of neurologic disease

#### Second set of laboratory tests

<table>
<thead>
<tr>
<th>SPECIMEN</th>
<th>LABORATORY TEST</th>
<th>CLINICAL REASONING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cerebrospinal fluid</td>
<td>Gram stain and culture</td>
<td>Rule out bacterial infection</td>
</tr>
<tr>
<td></td>
<td>Blood cell count: red blood cells, white</td>
<td>Rule out bacterial infection</td>
</tr>
<tr>
<td></td>
<td>blood cells, and differential</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Glucose</td>
<td>Rule out bacterial infection</td>
</tr>
<tr>
<td></td>
<td>Protein</td>
<td>Rule out bacterial infection</td>
</tr>
<tr>
<td>Blood</td>
<td>Thick blood film</td>
<td>Rule out malaria</td>
</tr>
<tr>
<td>Cerebrospinal fluid (paired serum specimen)</td>
<td>PCR/culture within first 7 days</td>
<td>Confirm presence of yellow</td>
</tr>
<tr>
<td></td>
<td>IgM against yellow fever</td>
<td>fever virus</td>
</tr>
<tr>
<td>Serum</td>
<td>PCR/culture</td>
<td>Confirm presence of yellow</td>
</tr>
<tr>
<td></td>
<td>Antibody tests</td>
<td>fever virus</td>
</tr>
<tr>
<td>Feces</td>
<td>Viral culture</td>
<td>Rule out polio, other</td>
</tr>
<tr>
<td></td>
<td></td>
<td>enteroviruses</td>
</tr>
</tbody>
</table>

ANNEX 3: CASE DEFINITION OF VISCEROTROPIC DISEASE,¹ ACCORDING TO THE BRIGHTON COLLABORATION²

Level 1 of diagnostic certainty
≥ 3 major criteria³

Level 2 of diagnostic certainty
2 major criteria³
or
1 major criterion and ≥2 minor criteria³

Level 3 of diagnostic certainty
≥ 3 minor criteria
or
1 major criterion and 1 minor criterion³

Major and minor criteria used in the case definition of viscerotropic disease

<table>
<thead>
<tr>
<th>MAJOR CRITERIA</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepatic:</td>
<td>Total bilirubin &gt;1.5X ULN⁴</td>
</tr>
<tr>
<td></td>
<td>[≥1.5X patient’s baseline value if known]</td>
</tr>
<tr>
<td></td>
<td>OR</td>
</tr>
<tr>
<td></td>
<td>ALT or AST ≥3X ULN⁴</td>
</tr>
<tr>
<td></td>
<td>[≥3X patient’s baseline value if known]</td>
</tr>
<tr>
<td>Renal:</td>
<td>Creatinine ≥1.5X ULN⁴</td>
</tr>
<tr>
<td></td>
<td>[≥1.5X patient’s baseline value if known]</td>
</tr>
<tr>
<td>Musculoskeletal:</td>
<td>CPK ≥5X ULN⁴</td>
</tr>
<tr>
<td>Respiratory:</td>
<td>Oxygen saturation ≤88% on room air</td>
</tr>
<tr>
<td></td>
<td>OR</td>
</tr>
<tr>
<td></td>
<td>Requirement for mechanical ventilation</td>
</tr>
<tr>
<td>Platelet disorder:</td>
<td>Platelets &lt;100,000/µl</td>
</tr>
</tbody>
</table>

¹ The case definition should be applied when there is temporal association with vaccination and no clear alternative diagnosis to account for the symptoms.
² Previously published WHO guidelines regarding the case definition and data collection for VTD associated with yellow fever vaccine preceded the development of these guidelines. However, the more recent and detailed Brighton Collaboration case definitions and guidelines are preferred.
³ Whenever ≥1 major criteria or both a major and minor criteria are used to meet the case definition, they must each represent different organ systems (e.g., hepatic or renal).
⁴ ULN = Upper limit of normal for the reference range of normal values reported by the clinical laboratory performing the indicated test.
Annex 3 Table. (Continued).

<table>
<thead>
<tr>
<th>MAJOR CRITERIA</th>
<th>MINOR CRITERIA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertension: Requirement for vasopressor drugs to maintain systolic blood pressure.</td>
<td>Hepatic: Jaundice</td>
</tr>
</tbody>
</table>
| Coagulopathy: INR ≥ 1.5 or prothrombin time ≥ 1.5X ULN⁴ or activated partial thromboplastin ≥ 1.5X ULN⁴ or elevated FDP⁵ or hemorrhage from more than one site.⁶ | Renal: Urine output <500 mL urine/24 h for adults  
Urine output <0.5 mL/kg/h for children⁷ |
|                                 | Musculoskeletal: Positive urine dipstick test for blood with a negative urine microscopy exam for red blood cells | |
| Respiratory: Increased respiratory rate for age⁸ | Platelet disorder: Petechiae or purpura present |
| Hypertension: Systolic BP <90 mm Hg for adults  
Systolic BP <5th percentile for children aged <16 years | Coagulopathy: Clinically evident hemorrhage (one of the following):⁹  
• Epistaxis  
• Hematemesis  
• Melena  
• Hematochezia  
• Hemoptyisis  
• Metrorrhagia or menorrhagia  
• Gingival hemorrhage  
• Persistent bleeding from needle puncture sites |

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⁵ FDP = Fibrin degradation products.  
⁶ See coagulopathy criteria below for list of included hemorrhage sites.  
⁷ Applies to children aged <13 years.  
⁹ Due to the relatively high baseline of hematuria due to various causes, hematuria is too nonspecific to be used as an indicator of coagulopathy for the VTD case definition.
ANNEX 4: VISCEROTROPIC DISEASE (VTD)-CAUSALITY CRITERIA FOR YELLOW FEVER VACCINE, ACCORDING TO THE BRIGHTON COLLABORATION

I. Definite yellow fever vaccine-associated causality
   One or more of the following are present:
   1. Yellow fever 17D virus isolation from blood >10 days post-vaccination.
   2. Yellow fever 17D virus concentration in blood ≥3 log_{10} PFU/mL on any day.
   3. Yellow fever 17D viral amplification from blood ≥14 days post-vaccination.
   4. Isolation of yellow fever 17D virus OR amplification of yellow fever 17D viral RNA from tissue and histopathology compatible with yellow fever (e.g., liver midzonal necrosis, Councilman’s bodies).
   5. Yellow fever virus-specific antigen in tissue with characteristic vaccine-associated distribution (extrahepatic or mesenchymal cell involvement) demonstrated by immunohistochemistry AND histopathology consistent with yellow fever (e.g., liver midzonal necrosis, Councilman bodies) AND NO history of being in a yellow fever-endemic or -epidemic area within 10 days of symptom onset.

II. Probable yellow fever vaccine-associated causality
   One or more of the following are present:
   1. Yellow fever 17D virus isolation from blood 8-10 days post-vaccination.
   2. Yellow fever 17D virus concentration in blood ≥2 log_{10} PFU/mL, but <3 log_{10} PFU/mL on any day 1-10 days post-vaccination.
   3. Yellow fever 17D viral RNA amplification from blood ≥14 days post-vaccination.
   4. Isolation of yellow fever 17D virus OR amplification of yellow fever-endemic or 17D viral RNA from tissues.
   5. Histopathology consistent with yellow fever (e.g., liver midzonal necrosis, Councilman bodies) AND no history of being in a yellow fever-endemic or -epidemic area within 10 days of symptom onset.

III. Suspect yellow-fever vaccine-associated causality
   One or more of the following are present:
   1. Histopathology consistent with yellow fever (e.g., liver midzonal necrosis, Councilman bodies) AND history of being in a yellow fever-endemic or -epidemic area within 10 days of symptom onset.
   2. Yellow fever virus-specific antigen in tissue demonstrated by immunohistochemistry AND history of being in a yellow fever-endemic or -epidemic area within 10 days of symptom onset.

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1 Confirmed as 17D virus by nucleotide sequencing.
2 Immunohistochemistry (IHC) performed by using polyclonal antibody to yellow fever viral antigen that reacts to both 17D yellow fever virus and wild type yellow fever virus.
3 If both the fourth and fifth criteria for probable yellow fever vaccine-associated causality are present, this is equivalent to the fourth criteria for definite yellow fever vaccine-associated causality. In this case, the condition of “no history of recently being in a yellow-fever endemic/epidemic area within 10 days of symptom onset” is unnecessary, because the possibility of wild-type yellow fever virus infection has been eliminated by the identification of the 17D yellow fever vaccine virus in the tissues.
IV. Insufficient data to determine yellow fever vaccine-associated causality

One or more of the following:

1. No yellow fever testing done.

   OR

2. Yellow fever testing done and results do not meet any of the criteria for causality levels 1, 2, or 3 as previously indicated.  

\[4\] The presence or absence of serum yellow fever virus-specific antibodies (IgM or IgG) has not been demonstrated to correlate with or be predictive of yellow fever vaccine causality in cases of VTD.
ANNEX 5. EVENT CLASSIFICATION ACCORDING TO THE BRIGHTON COLLABORATION

A. The case definition applies when there is temporal association and there is no clear alternative diagnosis to explain the symptoms.

B. Once this is determined, **it should be determined if the clinical picture meets the case definition:**

1. **The event meets the case definition** (Annex 3): that is, the case meets one of the three levels of diagnostic certainty:
   a. Level 1 of diagnostic certainty: ≥3 major criteria.
   b. Level 2 of diagnostic certainty: 2 major criteria or 1 major criterion and 2 or more minor criteria.
   c. Level 3 of diagnostic certainty: ≥3 minor criteria or 1 major criterion and 1 minor criterion.

   The case definition levels differ in diagnostic certainty, not in clinical severity of VTD. Similarly, levels of diagnostic certainty do not reflect causal association with a given vaccine.

2. **If an adverse event reported as VTD does not meet levels 1, 2, or 3 of the VTD case definition,** then additional diagnostic studies should be done to further search for diagnoses not previously considered that could explain the clinical picture of the ESAVI.

3. **Reported VTD with insufficient evidence to meet the case definition.** If the evidence available for the event is insufficient to permit classification by any level of diagnostic certainty (e.g., because of missing information), such an event should be categorized as “reported VTD with insufficient evidence to meet the case definition.” Notations should be made as to what evidence is missing.

4. **If there is adequate evidence that an event does not meet the case definition,** the event should be rejected and should be reported as “not a case of VTD.” Such evidence is considered adequate if the investigation reveals negative findings for all necessary criteria (necessary conditions) for diagnosis.

C. **Once it is determined that an event meets the case definition** (levels 1-3) or is classified as reported VTD with insufficient evidence to meet the case definition, the event should be classified according to its causality with the yellow fever vaccine in one of the categories of Annex 4.
ANNEX 6: INTERNATIONAL TECHNICAL COOPERATION

Following the report to PAHO/WHO of four fatal adverse events associated with yellow fever vaccination (sub-strain 17DD), technical collaboration activities began with Peru’s MoH, which had already started investigating the cases. A panel of experts on hemorrhagic yellow fever, virology, epidemiology, yellow fever vaccination, and vaccine quality was established. In conjunction with national authorities, the panel aimed to analyze the reported cases and determine if a causal relationship with the vaccine existed. The panel included experts from WHO, PAHO, CDC, the University of Texas in Galveston, the Butantan Institute of Brazil, and Kleiner, Perkins, Caufield, & Byers (KPCB) Pandemic and Biodefense Fund of the United States.

As a part of its work, the panel issued alerts to countries in the Americas endemic for yellow fever. The first message, sent on 2 November 2007, provided an update of the outbreak in Peru and requested that countries:

- Immediately suspend use of yellow fever vaccine lot 05OVFA121Z from Bio-Manguinhos, and lots related to it in production, specifically lots 05OVFA118Z, 05OVFA119Z, 05OVFA120Z, 05OVFA122Z, 05OVFA123Z, 05OVFA124Z, 05OVFA125Z, and 05OVFA126.
- Intensify surveillance to detect serious adverse events following vaccination.

The message was followed by conference calls held with all target countries, with the aim of reviewing and classifying serious ESAMI and VTD cases. Other measures taken by the international team included:

- Sending a team of experts from PAHO/WHO and the CDC to the field to continue gathering the epidemiological, virological, molecular, and pathological data needed to classify reported cases. The regional network of reference laboratories facilitated technical cooperation between countries in the use of technologies not available in many national reference laboratories.
- Performing independent analyses of manufacturing processes, vaccine quality, and the distribution and use of the aforementioned lots.

Additionally, laboratory tests on vaccine samples were conducted to determine whether the notified events had an association with the lots in question. One set of vaccine vials remains stored in the CDC and in Peru, so that they may be reexamined when more advanced technology becomes available.
### ANNEX 7: CLINICAL CHARACTERISTICS OF CONFIRMED CASES OF VTD ASSOCIATED WITH YELLOW FEVER VACCINE IN PERU

<table>
<thead>
<tr>
<th>CASE</th>
<th>AGE/SEX</th>
<th>PRE-EXISTING CONDITIONS</th>
<th>DAYS BETWEEN VACCINATION AND ONSET</th>
<th>SIGNS OR SYMPTOMS</th>
<th>LABORATORY ABNORMALITIES</th>
<th>RESULTS AND FINDINGS OF THE AUTOPSY</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>23 / F</td>
<td>Acne rosacea</td>
<td>1</td>
<td>9</td>
<td>Fever, headache, myalgia, malaise, nausea, vomiting, diarrhea; 8 days after vaccination, developed shock, adult respiratory distress syndrome (ARDS), encephalopathy, multi-organ failure. Death.</td>
<td>Leukocytes: 66,400 mm$^3$ Platelets: 54,000/mm$^3$ AST: 78 U/L ALT: 65 U/L Creatinine: 4.1 mg/dL CPK: 4,055 U/L Midzonal necrosis, steatosis (liver), acute tubular necrosis, thyroid neoplasia, and chronic thyroiditis.</td>
</tr>
<tr>
<td>2</td>
<td>24 / F</td>
<td>None known</td>
<td>&lt;1</td>
<td>14</td>
<td>Fever, headache, malaise, myalgia, nausea, vomiting, diarrhea. Eleven days after vaccination, presented shock, encephalopathy, acidosis, gastrointestinal bleeding, jaundice, ARDS, multi-organ failure. Death.</td>
<td>Hematocrit 15.5% Leukocytes: 16,300/mm$^3$ Platelets: 15,000/mm$^3$ AST: 735 U/L ALT: 167 U/L Bilirubin: 6.2 mg/dL Blood urea nitrogen: 112 mg/dL CPK: 3173 U/L Steatosis, focal necrosis (liver), cerebral edema, pulmonary edema, and severe <em>Candida</em> infection (larynx, trachea, stomach).</td>
</tr>
<tr>
<td>3</td>
<td>79 / M</td>
<td>Cardiac disease, prostate cancer</td>
<td>3</td>
<td>11</td>
<td>Fever, malaise, dyspnea, abdominal pain, vomiting, diarrhea. Nine days after vaccination, presented progressive shocks, ARDS, acidosis, renal failure. Death.</td>
<td>Leukocytes: 17,400/mm$^3$ Platelets: 249,000/mm$^3$ AST: 416 U/L ALT: 231 U/L Creatinine: 2.8 mg/dL Midzonal necrosis, steatosis (liver), acute tubular necrosis, and depletion of white pulp (spleen).</td>
</tr>
</tbody>
</table>

(Continued)
Annex 7 (Continued).

<table>
<thead>
<tr>
<th>CASE</th>
<th>AGE/SEX</th>
<th>PRE-EXISTING CONDITIONS</th>
<th>DAYS BETWEEN VACCINATION AND ONSET</th>
<th>SIGNS OR SYMPTOMS</th>
<th>LABORATORY ABNORMALITIES</th>
<th>RESULTS AND FINDINGS OF THE AUTOPSY</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>49 / F</td>
<td>Systemic lupus erythematosus, rheumatoid arthritis</td>
<td>Unclear (7-18 days)</td>
<td>Headache, malaise, arthralgia; 29 days after vaccination, hospitalized with generalized edema, jaundice, altered mental status, acidosis bleeding, difficulty breathing. Death.</td>
<td>Hematocrit: 31%</td>
<td>Leukocytes: 5,530/mm³ Platelets: 57,000/mm³ AST: 91 U/L ALT: 128 U/L Bilirubin: 5.2 mg/dL Creatinine: 3.3 mg/dL</td>
</tr>
</tbody>
</table>
