October 15, 2014 (replaces version dated October 10, 2014)

To: North Carolina Health Care Providers and Laboratories
From: Megan Davies, MD, State Epidemiologist
      Scott Zimmerman, DrPH, MPH, HCLD (ABB), State Laboratory of Public Health
Re: Ebola Hemorrhagic Fever (4 pages)

This memo is intended to provide updated information to all North Carolina health care providers and laboratories regarding Ebola virus disease (EVD) and management of suspected cases.

This version has been updated to include clinical observations shared through the Infectious Disease Society of America by physicians who provided care to Ebola cases at Emory University; updated infection control guidance; and updated testing guidance.

Summary
National and international health authorities are currently working to control a large, ongoing outbreak of Ebola involving areas in West Africa. A map of affected areas is available at http://www.cdc.gov/vhf/ebola/outbreaks/2014uwestuafrica/. The first travel-associated case of Ebola to be diagnosed in the United States was confirmed on September 30, 2014 and the first case acquired in the United States was confirmed on October 12, 2014.

Clinical and Epidemiologic Features
Ebola can spread from person to person by direct contact with a sick person’s blood or body fluids or by contact with contaminated objects (such as needles).

The incubation period for Ebola is usually 8–10 days, but can range from 2–21 days. The risk for person-to-person transmission is greatest during the later stages of illness when viral loads are highest. Ebola is not transmissible during the incubation period (i.e., before onset of fever).

Initial symptoms include fever, headache, joint and muscle aches, sore throat, and weakness, followed by diarrhea, vomiting, and stomach pain. Skin rash, red eyes, and internal and external bleeding may be seen in some patients.

Case Investigation and Risk Assessment

- Any patient with fever or a clinically compatible illness who has been in a country affected by the Ebola outbreak within 3 weeks before illness onset should be placed in appropriate isolation precautions (see below) as soon as possible. Precautions should be maintained while a more thorough risk assessment is completed.
- Clinicians caring for patients meeting these criteria should contact their local health department or the state Communicable Disease Branch (919-733-3419; available 24/7) to discuss laboratory testing and control measures.
• The algorithm and this document are intended as general guidance. Providers are encouraged to use clinical judgment and to contact public health immediately with any questions or concerns.

• Ebola should be suspected and testing is recommended for patients with fever or clinically compatible illness who, within 3 weeks before onset, have had a high-risk exposure, defined as follows:
  o Percutaneous (e.g., needle stick) or mucous membrane exposure to blood or body fluids of an EVD patient;
  o Direct skin contact with, or exposure to blood or body fluids of, an EVD patient without appropriate personal protective equipment (PPE);
  o Processing blood or body fluids of a confirmed EVD patient without appropriate PPE or standard biosafety precautions; or
  o Direct contact with a dead body without appropriate PPE in a country where an EVD outbreak is occurring.

• Ebola and testing should be considered for patients with fever or clinically compatible illness who, within 3 weeks before onset, have had a low-risk exposure, defined as follows:
  o Household contact with an EVD patient; or
  o Other close contact with an EVD patient in health care facilities or community settings. Close contact is defined at http://www.cdc.gov/vhf/ebola/hcp/case-definition.html.

• For persons with fever or clinically compatible symptoms who resided in or traveled to affected areas within 21 days before onset but had no reported high- or low-risk exposures, testing may be indicated depending on other factors, including severity of illness, presence of abnormal laboratory findings (i.e., platelet count <150,000/µL, prolonged PT/PTT or elevated transaminases), and presence or absence of alternative diagnoses.

• Even following travel to areas where Ebola has occurred, persons with fever are more likely to have infectious diseases other than Ebola (e.g., common respiratory viruses, endemic infections such as malaria or typhoid fever). Clinicians should promptly evaluate and treat patients for these more common infections even if Ebola is being considered. Lassa fever should also be considered if Ebola is suspected, since there is overlap in terms of clinical features and geographic areas where exposures could occur.

**Ebola Virus Testing:**

• **Testing Employed at the North Carolina State Laboratory of Public Health (NCSLPH):** Specimens will not be accepted without prior consultation. The NCSLPH utilizes the CDC/USAMRID Ebola Zaire rRT-PCR assay that has been granted FDA Emergency Use Authorization. The estimated turn-round-time for presumptive results is 6 hours for a single specimen and up to 24 hours for multiple specimens. It is important to note that it may take up to 3 days after symptoms appear for the virus to reach detectable levels. Therefore, if specimens are collected <3 days after onset, a later specimen may be needed to completely rule-out Ebola. Since all specimens tested at the NCSLPH will be forwarded to the CDC for more extensive laboratory testing, we are requiring 2 purple top tubes for submission – see table on the following page. CDC testing will include rRT-PCR with multiple primer probe sets for Ebola to confirm the initial results, tests for other hemorrhagic fever viruses, virus isolation, and serology when indicated by the clinical or epidemiological presentation.

**USE APPROPRIATE PRECAUTIONS WHEN COLLECTING SPECIMENS FOR EBOLA TESTING.**


• All specimen submissions must be accompanied by a completed BTEP Specimen Submission Form (http://slph.ncpublichealth.com/Forms/DHHS-5010-BTEmergingPathogens-0313.pdf), a CDC 50.34 DASH Form (http://slph.state.nc.us/Forms/CDC-Dash-NCSLPH-013114.pdf) and a Viral Special Pathogens Branch Diagnostic Specimen Submission Form (http://www.cdc.gov/ncezid/dhcpp/vspb/pdf/specimen-submission.pdf).

• The NCSLPH highly recommends that individuals packaging and shipping these diagnostic specimens use their professional judgment and consider packing instruction 620, IATA guidelines for Category A, which utilizes a triple packaging system. *We anticipate active discussion with all entities requesting diagnostic testing for Ebola and we will provide more specific guidance on a case-by-case basis.*
### Appropriate Specimens for Ebola rRT-PCR Testing at NCSLPH

<table>
<thead>
<tr>
<th>Specimen Type</th>
<th>Quantity</th>
<th>Testing</th>
<th>Transport</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whole blood with EDTA anticoagulant (purple top tube) in non-glass collection tube</td>
<td>≥ 4ml X 2</td>
<td>rRT-PCR</td>
<td>Refrigerated (4°C), placed on cold packs. Package specimens using Category A guidelines.</td>
</tr>
</tbody>
</table>

### Appropriate Specimens for Ebola Testing Conducted at the CDC

<table>
<thead>
<tr>
<th>Specimen Type</th>
<th>Quantity</th>
<th>Testing</th>
<th>Transport</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncoagulated whole blood (purple, yellow, or blue top) in non-glass collection tube</td>
<td>≥ 4ml</td>
<td>Culture, PCR</td>
<td>Refrigerated (4°C), placed on cold packs if shipment is to be received within 72 hrs. For delays exceeding 72 hrs freeze serum at -70°C &amp; ship on dry ice.</td>
</tr>
<tr>
<td>Serum (red top, collected in non-glass tube)</td>
<td>≥ 4ml</td>
<td>Culture, PCR, Serology</td>
<td>Ship at room temperature. Note: An autopsy or surgical report must accompany the specimen.</td>
</tr>
<tr>
<td>Formalin-fixed or paraffin-embedded tissues</td>
<td>As Appropriate</td>
<td>Immuno-histochemistry</td>
<td>Ship specimen frozen on dry ice in a plastic container.</td>
</tr>
<tr>
<td>Fresh frozen tissue</td>
<td>1 cm³ (except for biopsies)</td>
<td>Culture, PCR</td>
<td></td>
</tr>
</tbody>
</table>

### CONTACT THE BTEP UNIT (919-807-8600) PRIOR TO ANY SHIPMENT OR IF YOU HAVE QUESTIONS.

Address all specimen shipments as follows:

-Attention: Bioterrorism & Emerging Pathogens Unit
-North Carolina State Laboratory of Public Health
-4312 District Drive
-Raleigh, NC 27607-5490

**Routine Laboratory Testing on Suspect EVD Cases**

- Clinicians should ensure that laboratory staff are aware if a diagnosis of EVD is being considered so that appropriate precautions can be taken in the laboratory when handling routine or diagnostic specimens.

- The NCSLPH encourages institutions to conduct an internal risk assessment to review all handling and testing procedures that are associated with specimens from a suspect Ebola case. The NCSLPH highly recommends the use of professional judgment to determine the need for enhanced safety precautions.

- The NCSLPH strongly recommends that laboratories consider the following guidelines for handling of routine laboratory specimens from persons under investigation for Ebola:

**Infection Control**

- Every North Carolina hospital must be prepared to assess and provide initial management to a patient with suspected or confirmed EVD.

- Infection prevention procedures must be adhered to strictly.


- The Division of Public Health recommends having a dedicated monitor observe donning and doffing of PPE by all healthcare personnel caring for patients with suspected or confirmed EVD.

- The following are minimum infection prevention measures recommended when caring for persons with known or suspected EVD - SEE LINK BELOW FOR ADDITIONAL INFECTION CONTROL RECOMMENDATIONS.
  - Patient placement: Patients should be placed in a single patient room (containing a private bathroom) with the door closed.
  - Personal Protective Equipment (PPE): All persons entering the patient room should wear at least gloves, gown (fluid resistant or impermeable), eye protection (goggles or face shield), and facemask.
  - Patient care equipment: Dedicated medical equipment (preferably disposable, when possible) should be used for the provision of patient care.
Aerosol-generating procedures: Aerosol-generating procedures should be avoided. If such procedures are necessary, Airborne Precautions (use of N95 respirator or higher and airborne isolation room) should be implemented for the duration of the procedure.

Environmental infection control: Diligent environmental cleaning and disinfection and safe handling of potentially contaminated materials is paramount, as blood, sweat, emesis, feces and other body secretions are potentially infectious.


- If a patient with suspected EVD is going to be transported to another facility, the transport staff and the receiving facility should be notified before transport so that appropriate precautions can be taken.

### Assessment and Monitoring of Asymptomatic Persons with Ebola Exposure

- All persons arriving in North Carolina who travelled to an affected region within 21 days should contact their local health department or the Communicable Disease Branch epidemiologist on call to undergo a thorough risk assessment.

- Control measures may be recommended by the local health department based on findings of the risk assessment.


### Treatment

- Supportive care only; no antivirals are currently available for treatment of Ebola.

- Key interventions include:
  - Providing intravenous fluids and balancing electrolytes (body salts)
  - Maintaining oxygen status and blood pressure
  - Treating other infections if they occur

- Clinical observations shared through the Infectious Disease Society of America by physicians who provided care to Ebola cases at Emory University include the following:
  - Despite weight gains of 15–20 kg, the patients were profoundly hypovolemic due to their low serum albumin and vascular leak with third spacing. Fluid losses in their patients were 5–10 L/day.
  - Electrolyte losses were significant and included profound hyponatremia, hypokalemia and hypocalcemia. Arrhythmias were noted, and both intravenous and oral electrolyte repletion was necessary.
  - Nutritional depletion was evident as well.
  - Ebola virus RNA was detected in blood, urine, vomitus, stool, endotracheal suctioning, and semen and on skin. It was not detected in dialysate. Environmental testing in the patient rooms had no detection of viral RNA and included many high touch surfaces such as bed rails and surfaces in the bathroom.
  - Intensive 1:1 nursing care was necessary around the clock. Patients were monitored continuously and this level of nursing care allowed for rapid responses to clinical changes. Nursing and other team members provided emotional support, and as the patients improved, help with self-care and physical therapy.

### Reporting

- Physicians are required to contact their local health department or the state Communicable Disease Branch (919-733-3419) as soon as Ebola or any other hemorrhagic fever virus infection is reasonably suspected.

This is an evolving situation and recommendations are likely to change as new information becomes available. Updated information and guidance are available from the CDC at [http://www.cdc.gov/vhf/ebola](http://www.cdc.gov/vhf/ebola) and from North Carolina Public Health at [http://epi.publichealth.nc.gov/cd/diseases/hemorrhagic.html](http://epi.publichealth.nc.gov/cd/diseases/hemorrhagic.html).