Interpretation of Negative Molecular Test Results in Patients with Suspected or Confirmed Ebola Virus Disease: Report of Two Cases

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ABSTRACT

Quantitative reverse-transcription-polymerase-chain reaction (qRT-PCR) is the most sensitive quantitative diagnostic assay for detection of Ebola virus in multiple body fluids. Despite the strengths of this assay, we present two cases of Ebola virus disease (EVD) highlighting potential for false-negative results during early and late stages of EVD. The first case emphasizes the low negative-predictive-value of qRT-PCR during incubation and the early febrile stage of EVD and the second case potential for false-negative results during recovery and late neurologic complications of EVD. Careful interpretation of test results are needed to guide difficult admission and discharge decisions in suspected or confirmed EVD.
INTRODUCTION

Presumptive diagnosis of Ebola virus disease (EVD) is dependent upon a combination of epidemiologic risk factors with a compatible clinical syndrome.(1-4) Confirmation of EVD during the 2014-15 West Africa outbreak has been largely reliant on quantitative reverse transcription polymerase chain reaction (qRT-PCR) testing, the gold-standard diagnostic assay for detecting and quantifying Ebola virus (EBOV).(5) Between August 2014 and March 2015 the United States Food and Drug Administration (FDA) issued Emergency Use Authorizations (EUA) for eight molecular assays for EBOV.(6) At present no FDA approved molecular assay for the detection of EBOV is available for routine use.

Here we report two cases of EVD highlighting challenges in admission and discharge decisions despite ready availability of EBOV molecular diagnostic testing. The first case reveals the low negative-predictive value of EBOV PCR in blood during incubation and the early febrile stage of EVD, the second case highlights the potential for false-negative results during recovery and late complications of EVD. Negative molecular test results cannot be relied upon to rule out infection during the early febrile stage of EVD and in patients with a high-pretest probability, serial testing is recommended.(5,7) False-negative results can occur late in EVD attributable to one or multiple factors including improper specimen collection, transportation, or storage, errors in specimen preparation, or factors related to the assay or equipment performance. Careful clinical interpretation of results is needed to guide admission and discharge decisions.
CASE 1

On September 25, 2014 a six-month-old boy presented with his mother to the Médecins Sans Frontières (MSF) ELWA-3 Ebola management center (EMC) in Monrovia, Liberia. The mother exhibited symptoms of EVD, all primary family members died with syndromes consistent with EVD. The child was asymptomatic on presentation and breastfeeding from his mother, but had no other notable medical history. The mother and child were admitted to the suspect tent in the high-risk zone of the EMC for qRT-PCR EBOV testing from blood, completed by an adjacent National Institutes of Health (NIH) supported laboratory. An effort was made to isolate the child from the mother and other suspect cases during the testing interval. The mother’s test returned positive and the child’s negative with a PCR cycle threshold (Ct) value of 39 (Figure 1). The local cut-off for a “negative” test result was a Ct value $\geq 35$ and during this time in Monrovia there was no indeterminate range utilized. The mother was transferred to a confirmed tent in the high-risk zone and the asymptomatic boy was discharged under observation. It was normal practice to only hold and retest symptomatic patients.

Because the child had no immediate family members he was cared for by EMC staff in an observation tent for individuals who tested negative for EBOV, discharged from either the suspect or confirmed tents. Staff wore “light” personal protective equipment (PPE) consisting of scrubs, boots, surgical gown, single pair of exam gloves, surgical mask, and routine use of hand washing with 0.05% chlorine. On September 27, two days after discharge from the suspect tent, the child developed fever to 38.4°C, without other symptoms of EVD. Repeat EBOV testing was again negative (Ct 40, Figure 1) and he
continued to be observed in the low risk area. On September 29, four days after discharge from the suspect tent in the setting of persistent fever, repeat EBOV testing returned positive (Ct 30, Figure 1). He was admitted to the confirmed tent in the high-risk zone and subsequently died of EVD. No staff members who cared for the child became infected.

CASE 2

On September 20, 2014, a nine-year-old boy was brought with his mother and two siblings to ELWA-3. They met the suspect case definition for EVD and were transferred to the suspect tent in the high-risk zone and tested positive for EBOV by qRT-PCR from blood (Ct 24, Figure 2). The mother died in the suspect tent and the three children were transferred to the confirmed tent. One of the two siblings died and the other recovered and was discharged. The nine-year-old boy complained of fever for five days, and progressive weakness, vomiting, and diarrhea at the time of presentation. No other significant medical history was reported. On admission to the confirmed tent, he was confused and apathetic. He was treated with oral rehydration solution as well as metoclopramide and loperamide for vomiting and diarrhea. Empiric treatment for malaria and bacterial co-infection was provided with artemether-lumefantrine and cefixime per MSF protocol. Fever and gastrointestinal symptoms gradually resolved, he began to walk and communicate. However, he continued to have neurocognitive abnormalities with delayed cognition and difficulty formulating complex sentences, which was felt to represent possible long-standing developmental delay. On September 28, in the absence
of other symptoms for three days and thirteen days post-initial symptom onset, repeat EBOV testing was negative (Ct 40, Figure 2).

The boy was discharged and transported to his community, but was not recognized there. He returned to the EMC, was placed in the observation tent in the low-risk zone and cared for by staff in light PPE, while further disposition options were considered. On September 30, fifteen days post-initial symptom onset, he developed fever to 38.6°C with associated neck rigidity, severe weakness, and diminished consciousness. A malaria rapid diagnostic test was negative and empiric treatment for bacterial meningitis was initiated with ceftriaxone. Repeat EBOV testing was performed which returned positive (Ct 32, Figure 2) and he was readmitted to the confirmed tent in the high-risk zone. Fever and neck rigidity subsided and on October 2, seventeen days post-initial symptom onset repeat EBOV testing was performed. Results showed a rising PCR cycle threshold value consistent with a down trending viral load in blood (Ct 37, Figure 2). The boy survived and was released to a newly opened orphanage. No staff member who cared for the boy subsequently became infected.

**Retrospective Quantification of Viral Load**

We sought to precisely characterize viral expansion in the blood of these two patients by retrospectively quantifying viral copy number in blood specimen. Using an external control specimen with known viral copy number we establish a standard curve for comparison to our patients’ blood specimens by q-PCR. Case 1 had no detectable viral copies in blood from September 25 or 27 during viral incubation and early febrile
disease, but had $6.0 \times 10^5$ viral copies/ml detected from the September 29 specimen, representing greater than a five-log expansion in two days. Case 2 presented to ELWA-3 during the gastrointestinal stage of illness on September 20, five days after initial symptom onset. Retrospective quantification of viral copy number in blood from September 20 showed $3.9 \times 10^6$ viral copies/ml, consistent with high viral load typically observed during the gastrointestinal stage of EVD and was positive for malaria. Retrospective analysis of the September 28 specimen obtained following resolution of gastrointestinal symptoms but in the presence of neurocognitive abnormalities showed that no RNA, neither viral nor host was present, indicating that an error in specimen handling or RNA extraction likely led to a false-negative result. Retrospective analysis of specimens from September 30 and October 2 show down trending viral copy number in blood at $3.1 \times 10^4$ copies/ml and $2.3 \times 10^2$ copies/ml respectively.

**DISCUSSION**

Presumptive diagnosis of EVD is based upon a compatible clinical syndrome and epidemiologic risk factors. Early clinical syndrome of EVD consists of fever, typically associated with lethargy and malaise, which overlaps with many commonly observed infectious syndromes in West Africa, most notably malaria. Presence of epidemiologic risk factors assists in risk-stratifying patients under consideration for EVD but are less helpful when Ebola is widespread in the community. While clinical case-definitions for suspected EVD provide useful guidelines for providers making triage decisions, these definitions are imprecise.
Molecular testing for EBOV may reliably confirm EVD within hours, providing an essential tool for management. However, a recognized limitation of EBOV PCR is the low negative-predictive value during EBOV incubation and early febrile stage of disease. Difficult triage decisions routinely arose at ELWA-3 at the peak of the outbreak. Case 1 presented a distinct dilemma, high-risk exposure but no fever on presentation. Given that he had no alternative caregiver and was at high-risk for becoming EBOV PCR-positive he was admitted to the suspect tent of the EMC. The boy’s EBOV PCR from blood became positive on September 29, 4 days after initial presentation to the EMC and 2 days after fever onset. With an average EBOV incubation period of 6-8 days, his presentation is most consistent with infection prior to EMC admission.

This case reinforces that circulating EBOV remains below the level of detection of sensitive molecular assays during incubation and early febrile illness and also highlights the rapid rate of rise of virus in blood over a short time interval. Patients with high-pretest probability of EVD require serial testing and close clinical observation.

Current understanding of EVD pathogenesis suggests that EBOV enters the body through mucosal or skin barrier breaches, encountering the mononuclear phagocyte system. Infected macrophages and dendritic cells migrate to lymph nodes with viral replication, indirectly inducing lymphocyte depletion, and release into blood.(9) Fever, driven by cytokines, typically precedes viral detection in blood. Early after fever onset, viral tropism for multiple cell types and results in exponential viral growth.(10,11) Case 1 had
exponential viral growth occur over a two-day period where viral copy number in blood went from undetectable two days post fever onset to almost six million copies/mL four days post-fever onset. In the absence of novel diagnostic tests that do not rely upon circulating virus, early diagnosis of EVD depends upon careful interpretation of molecular diagnostic test results in the context of clinical and epidemiologic data. In addition, during early epidemic response with limited resources and overwhelming needs, infection control management of those patients at high-risk for EBOV, yet test negative, is especially challenging because of the length of incubation.

The second case highlights that false-negative molecular test results for EBOV may occur in the later stages of EVD reinforcing the need to carefully interpret molecular test results in the context of clinical factors. Per MSF protocol this patient underwent PCR testing three or more days after resolution of symptoms in order to determine eligibility for discharge. Despite resolution of symptoms, however, notable neurocognitive abnormalities persisted. These neurological abnormalities progressed to a clinical syndrome most consistent with meningoencephalitis. In review, the persistent neurological symptoms in our patient might have prompted consideration of a false negative result and the need for repeat testing prior to discharge.

Retrospective analysis of the September 28 blood specimen indicated that likely an error in RNA extraction resulted in a false-negative test result. Late September coincided with the peak of the epidemic in Liberia. At this time staff-patient ratios were low and specimen-testing volume was high. Under these conditions errors can be expected, and
issues with sub-optimal specimen volume, use of incorrect collection tubes, long
specimen transport times, unreliable cold chain and equipment malfunction, all
contributed to the potential for false-negative test results.(12)

Consensus has not been reached on uniform criteria for safe discharge of EVD patients
and some maintain that resolution of clinical symptoms alone without the need for
molecular testing is adequate.(13,14) Others, including the World Health Organization
(WHO), recommend symptom resolution for three days and/or two consecutive negative
PCR tests 48 hours apart.(7,15) At this time, MSF maintains recommending discharging
EVD patients who are asymptomatic for 3 days and have a single negative PCR result.
The likelihood of patients having persisting EBOV infection meeting these discharge
criteria is believed to be extremely low.(15) From a public health perspective during an
escalating epidemic period, withholding discharge would likely be
counterproductive.(16) However, clinicians must be aware of the potential for false-
negative PCR results during resolution of EVD. Therefore it is recommended that if the
clinical context is unclear, two negative EBOV PCR tests be utilized for discharge
eligibility.

In summary these cases highlight the need for careful interpretation of EBOV PCR
results in the context of clinical and epidemiological factors. In the absence of early
diagnostics not reliant on circulating virus, serial testing of patients at high-risk for EVD
and close clinical observation is recommended. Improved understanding of the kinetics of
live virus versus molecularly detectable virus in various body fluids during resolution and
late complications of EVD will guide improved decision making at the time of discharge.

In the meantime, multiple factors will by necessity play into discharge decisions including but not limited to availability of staffing, patient volume, diagnostic capabilities, and bed-space in the isolation ward.

Notes

**Ethics:** This case series report utilized routine program data, which was de-identified; therefore, informed consent was not indicated. The study protocol was evaluated and approved by the ethics committees of Médecins Sans Frontières, Brussels, Belgium and University of Liberia, Monrovia, Liberia.

**Acknowledgement:** The authors would like to thank the staff of ELWA-3 and NIH/CDC laboratory in Monrovia, Liberia for their hard work and dedication during the Ebola epidemic of 2014-2015.

**Financial support:** This research was partly supported by the Intramural Research Program of the National Institute of Allergy and Infectious Diseases (NIAID), National Institutes of Health (NIH). All other expenses came from the routine budgets of each author’s affiliation.

**Conflicts of interest:** The authors have no conflicting interests to report.
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Figure 1. Case #1: initial results of whole blood testing for Ebola virus with threshold cycle (Ct) in Monrovia, Liberia 2014.
Figure 2. Case #2: initial results of whole blood testing for Ebola virus with threshold cycle (Ct) in Monrovia, Liberia 2014.

*Positive for malaria retrospectively using polymerase chain reaction test.
**False-negative quantitative reverse transcription polymerase chain reaction test based upon follow up lab control and retrospective viral load analysis results.