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Evaluation of the Tetracore Orthopox Bio Threat® antigen detection assay using laboratory grown orthopoxviruses and rash illness clinical specimens

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ABSTRACT

The commercially available Orthopox Bio Threat® Alert assay for orthopoxvirus (OPV) detection is piloted. This antibody-based lateral-flow assay labels and captures OPV viral agents to detect their presence. Serial dilutions of cultured Vaccinia virus (VACV) and Monkeypox virus (MPXV) were used to evaluate the sensitivity of the Tetracore assay by visual and quantitative determinations: specificity was assessed using a small but diverse set of diagnostically relevant blinded samples from viral lesions submitted for routine OPV diagnostic testing. The Bio Threat® Alert assay reproducibly detected samples at concentrations of 10² pfu/ml for VACV and MPXV and positively identified samples containing 10⁶ pfu/ml in 4 of 7 independent experiments. The assay correctly identified 9 of 11 OPV clinical samples and had only one false positive when testing 11 non-OPV samples. Results suggest applicability for use of the Bio Threat® Alert assay as a rapid screening assay and point of care diagnosis for suspect human monkeypox cases.

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1. Introduction

More than 30 years after the eradication of smallpox, Orthopoxviruses remain an emerging threat. In the past decade, zoonotic outbreaks of MPXV in central Africa and the US, cowpox virus in Europe, and VACV in Brazil highlight the emerging potential of these viruses (Centers for Disease Control and Prevention, 2003; Learned et al., 2005; Trindade et al., 2006; Vorou et al., 2008). Monkeypox is a zoonotic disease found in Central and West Africa that produces a pustular rash illness in humans. Although human contact with infected animals is the most common route of infection, extended human-to-human transmission chains can occur (Learned et al., 2005). Two genetic clades of MPXV have been characterized (Likos et al., 2005) and include the West African (mortality rate less than 1%) and Central African (up to 10% mortality) clades (Jezeck et al., 1987). MPXV was also the cause of a 2003 outbreak in the United States. Captive black tailed prairie dogs transmitted the virus to humans after being co-housed with MPXV infected African rodents (Reed et al., 2004). Thirty-seven confirmed and ten probable human cases resulted (Reynolds et al., 2006). More recently an outbreak of monkeypox occurred in southern Sudan where MPXV disease had previously never been reported (Damon et al., 2006; Formenty et al., 2010). Historical outbreaks of monkeypox in the Congo basin and West Africa, along with the emergence of this disease in the US and southern Sudan emphasize the importance of MPXV as an emerging infectious agent of global scale.

In addition to a clinical appearance similar to that of smallpox, human monkeypox is often confused with other rash illnesses; up to 50% of suspected monkeypox cases in the Democratic Republic of Congo are actually Varicella-Zoster virus (VZV) infections (Jezeck et al., 1988; Meyer et al., 2002). Early detection of OPV infection is important for making informed decisions regarding patient treatment, defining epidemiology of disease and implementing disease control measures. Current state of the art diagnoses of acute OPV infections largely utilize non-commercial PCR-based methods. Real-time PCR (RT-PCR) assays allow rapid detection and species level identification of multiple different OPVs in high-throughput, high-sensitivity formats (Kulesh et al., 2004; Li et al., 2006; Olson et al., 2004; Shchelkunov et al., 2011). In these assays, viral genomic material is isolated from swabs of lesions or from homogenized tissues prior to diagnostic testing. RT-PCR assays are highly sensitive and can detect as few as 10–100 genomes (Olson et al., 2004; Shchelkunov et al., 2011) and thus are state of the art for virus identification. However, requirements for skilled technicians, expensive instrumentation and strong laboratory controls to prevent cross-contamination limit the use of this technology in resource-poor areas such as rural Africa where rapid diagnosis of monkeypox would be beneficial.
Several alternative in-house assays have also been developed for diagnosis of OPV infection. A shell vial culture assay and direct fluorescent antibody (DFA) assay were tested with patient samples from recent VACV vaccines and detected virus as early as 4 days after vaccination (Fedorko et al., 2005). The DFA assay was the least sensitive (40% positive results OPV samples), but the most rapid and could be completed within an hour. The shell vial culture was sensitive to 89% of the samples tested but required up to 2 days to obtain results. By comparison, the RT-qPCR assay used in that study detected virus in 100% of the samples tested. Both assays require fluorescent probes, expensive instrumentation and extensive laboratory equipment and expertise to perform. An in-house IgM ELISA (Karem et al., 2005) affords detection of antibody markers of recent OPV infection by examining humoral immune response to infection. It uses equipment found in most clinical laboratories, but requires laboratory space, specialized reagents, and technical expertise to perform the assay and interpret the results. It is also necessary to wait at least 5 days after rash onset to obtain predictive results.

Despite advances in nucleic acid diagnostic assays and serologic assays for post-exposure surveillance, rapid point-of-care OPV diagnostics remain limited. This limitation is particularly acute in rural Africa where the orthopox MPXV is endemic and resources for diagnostics are scarce. A rapid, simple, point-of-care diagnostic assay would eliminate many of these limitations, allowing local health workers to quickly confirm suspected monkeypox cases and enact measures to prevent further spread of the disease. The Tetracore Orthopox BioThreat Alert® assay is the first commercially available lateral-flow based detection assay for OPV. Viral agent detection is accomplished through a combination of antibodies raised against VACV (O’Brien, 2012). One set of antibodies is conjugated with colloidal gold for labeling, while the second set of antibodies is bound at the detection line to capture labeled antibodies for visualization. Prior evaluation was performed at Plum Island with positive visual responses for Orthopoxviruses Horsepox and Camelpox, and negative visual responses for Suidpoxvirus Swinepox, and Capripoxviruses Sheeppox and Goatpox (O’Brien, 2012). Serologic cross-reactivity between various OPV is well known (Mercer et al., 2007), and as such the manufacturer insert suggests the assay should be sensitive to MPXV. However, this has not been verified. The current study addresses sensitivity and specificity of the Tetracore assay for VACV and MPXV, and examines its potential as a method for diagnosing human monkeypox.

2. Materials and methods

2.1. Tetracore assay protocol

The Tetracore assay can be utilized with a variety of sample types including both solid and liquid samples. Manufacturer instructions for dry or solid material include directly resuspending tissue or swab samples in Tetracore sample buffer. Liquid samples are to be diluted in Tetracore assay sample buffer by at least 1:2 ratio. Instructions then detail the application of approximately 150 μl of sample to the strips, which re-hydrates test reagents, and allows binding of OPV agents by labeled antibody. Test results are to be read after 15 min. A positive result is indicated by the formation of a colored line at the sample line window. The control line contains antibodies directed against labeled antibody and verifies the test was run appropriately; this line should appear regardless of the presence or absence of OPV. The test is invalid if no control line appears. A sample test strip image is shown in Fig. 1.

![Sample Line Control Line](image)

Fig. 1. Tetracore Orthopox cassette image and strip layout showing application port, the sample test line for the orthopox detection, and a control line to indicate the test was successfully completed.

2.2. Samples utilized

2.2.1. Laboratory viral isolates

Crude Wyeth Dryvax® VACV and MPXV–ROC Congo Basin strain 2003–385 (Hutson et al., 2009) virus stocks were grown in BSC-40 cells for 2–3 days, purified from cellular debris by centrifugation, and resuspended in 10 mM Tris pH 9.0 (Likos et al., 2005). Purified virus was evaluated in a cell–culture plaque forming assay to determine pfu/ml. Briefly, extracted samples were applied to BSC-40 cell monolayers in 10-fold dilutions, incubated at 36 °C/6% CO₂ for 48–72 h and subsequently fixed and stained with formalin and crystal violet to reveal plaques. Uninfected BSC-40 cells were processed through the virus purification procedure and collected for use as a control (BSC-40 lysate).

Crude VZV (Webster strain) was grown in human lung fibroblast (HLF) cells, lysed with radio immunoprecipitation assay (RIPA) buffer (50 mM Tris–HCl pH 8.0, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS), centrifuged to remove cellular debris, and the supernatant containing the virus was collected (Behrman et al., 2003). The virus was used without further characterization. Uninfected HLF cells were processed with this same protocol and used as a control (HLF lysate). Purified VZV (vaccine strain v0ka) was also grown in HLF cells. Infected cells were rinsed with PBS, removed by agitation with glass beads in DMEM media supplemented with 10% sorbitol, and vortexed after transfer to a 50 ml conical tube. A low speed spin pelletted the glass beads and cellular debris. Supernatant was then further centrifuged to enhance purity (Rahaus et al., 2003; Schmidt and Lennette, 1975), then aliquoted and frozen at −70 °C. Virus titer was determined with an in-house colorimetric assay. Briefly, 48-well plates of HLF cells were infected with serial dilutions of the virus stock and fixed in 2% paraformaldehyde after 48 h. Wells were blocked with PBS containing 10% BSA, 0.5% FBS and 0.1% tween–20. Primary antibody (MAb to VZV gp1, Biodesign, Saco, ME) was diluted 1:1000 in blocking buffer and applied to each well for 1 h, followed by a 1:1000 diluted goat anti–mouse HRP (Invitrogen) for 30 min. BM Blue POD (Roche, Indianapolis, IN) was applied until color developed. Wells were washed with PBS to stop the reaction and color stabilization buffer (50 mM Tris pH 6, 100 mM NaCl, 1 mM EDTA) was added (Chen et al., 2007).

2.2.2. Clinical specimens

Samples submitted to the CDC poxvirus program for OPV testing are held in a clinical samples database after diagnostics are complete and include swabs of lesions or vesicle fluid, or scab material. Swabs are resuspended in 400 μl 1 × PBS (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na₂HPO₄ and 1.47 mM KH₂PO₄), extracted using SETS (Swab Extraction Tube System, Roche Applied Science, Indianapolis, IN) then sonicated. Scab material is resuspended with 300 μl of 1 × PBS, then frozen in a dry-ice/ethanol bath, ground using a disposable pestle, vortexed, and sonicated. After diagnostic testing is complete, remaining sample is stored frozen at −80 °C. In this study, rash lesion clinical samples were used to test the
efficacy of the BioThreat Alert strips for detection of viral antigen in clinical samples. Samples were blinded and included human specimens from monkeypox, vaccinia, herpesvirus (Varicella and HSV-1) and parapoxvirus infections. Specimen remainders from 22 human cases were chosen; 11 were from OPV infections (monkeypox and vaccinia) and 11 were non-OPV (VZV, pseudocowpox, Orf and HSV1). By design, OPV clinical samples with early RT-PCR cycle threshold (Ct) values were selected to ensure antigen presence; earlier Ct values indicate higher concentration of genomic DNA. Genome equivalents were extrapolated by comparison with standard curves of Ct values generated from 10-fold dilutions of purified viral DNA (17.5 fg to 1.75 ng) using an E9L RT-PCR assay (Li et al., 2006). A generic genome size of 200,000 base pairs was used to calculate individual genome mass, and thus genome equivalents. All orthopox samples utilized had Ct values less than 24, or approximately 8 × 10^4 genome equivalents. Infectious virus was then evaluated by plaque assay (described in Section 2.2.1). The CDC’s Human Research Protection Office determined that use of de-identified diagnostic remainder for this activity was exempt from consideration under 45 CFR 46.101(4)(b) (3/11/2009).

2.3. Tetracore BioThreat testing procedures

2.3.1. Sensitivity to viral isolates

Tetracore Orthopox BioThreat Alert® assay (catalog number TC-8001-025) was provided by Tetracore (Rockville, MD). The BioThreat Alert assay was tested for sensitivity to lab grown OPV strains on duplicate strips by two users. Viruses were diluted in Tetracore sample buffer in 10-fold dilution series to final concentrations ranging from 10^4 to 10^6 pfu/ml of VACV, or 10^4 to 10^6 pfu/ml of MPXV, and 150 μl aliquots applied to the test strips (1.5 × 10^2–1.5 × 10^3 pfu/strip). Initial dilutions of VACV were 1:70 (from a stock concentration of 6.8 × 10^9 to 10^6 pfu/ml), so BSC-40 lysate was also diluted at 1:70, and Crude VZV and HLF cell lysates were diluted at 1:100. The 150 μl samples were applied to each test strip and results were visually scored as positive or negative after 15 min. For strips used to test VACV, detection was also performed using the ESEQuant Lateral Flow Reader (Qiagen, Valencia, CA) and analyzed with Lateral Flow Studio (Qiagen) to quantify line intensity. The reader measures reflectance, where increased line intensity reduces reflectance, and thus reduced signal (mV) received at the detector; the resulting line measurement is displayed as an inverted peak (i.e., a trough). Lateral Flow Studio software (Qiagen) calculates peak height, and along with the number of data point measurements quantifies the peak area. For MPXV, restrictions in moving select agent materials from our laboratory prevented ESEQuant analysis of these samples (reader is in another laboratory), thus MPXV isolates were only analyzed visually. A second study assessed improved sensitivity of later measurement time points and utilized the field portable BioThreat Alert Reader. This reader provides quantitative signal value (SV) measurements of sample line intensity. Purified VZV and DMEM media used for VZV resuspension were diluted at 1:2 in Tetracore buffer prior to application. Strips were analyzed between 15 and 37 min after sample application and those with an SV above the instrument default threshold value (TV) of 0.01 were scored positive.

2.3.2. Specificity with clinical samples

The specificity experiment was performed in duplicate on 22 unique clinical samples; 44 BioThreat Alert strips were utilized. Samples were supplied in 75 μl of PBS and diluted at 1:2 with Tetracore sample buffer (final 150 μl volume) and run on the Orthopox BioThreat Alert strips according to manufacturer instructions. The blind study samples were examined 15 min after sample application and visually scored, including qualitative notations of intensities for positive samples of strong, medium, weak or faint. “Strong” line intensity was defined as intensity equivalent to the control line. Signal for “medium” and “weak” notations had reduced intensity and a score of “faint” was given to indicate the weakest detectable line by naked eye. Although the control line can vary in intensity due to improper storage of the testing reagents or test age, no qualitative differences in intensity were noted throughout the testing period.

3. Results

3.1. Detection of orthopox viral isolates using the Tetracore assay

Visual analysis of laboratory preparations of VACV resulted in positive identifications at concentrations of 10^8 and 10^7 pfu/ml, but had varying success at 10^6 pfu/ml concentration with two samples identified as positive and three as negative (Table 1). Dilutions below 10^7 were negative. Crude VZV and uninfected HLF and BSC-40 cell lysates controls were negative. Quantitative peak area measurements using the ESEQuant® Lateral Flow Reader show the 10^8–10^7 pfu/ml peak area 100-fold above control sample peak areas and were 2248 ± 446 or 2346 ± 85, respectively. The 10^6 pfu/ml peak area average was almost 10-fold higher than lysate controls, and averaged 165 ± 69 units. Control values (e.g., VZV and cell lysates) were between 20 and 26 units based on peak area measurement. MPXV preparations were only analyzed visually (Table 1), but yielded similar results with 10^6–10^5 pfu/ml positive, and the 10^6 pfu/ml scoring weakly positive by comparison to the higher concentrations. Overall, a strong correlation was noted between visual analysis and quantitative measurements.

To assess the utility of the Tetracore BioThreat field portable reader and potential higher signal at later time points, VACV dilutions were run in duplicate and scanned at approximately 7.5 min
intervals, beginning at 15 min after sample application (Fig. 2a). Variability in processing time resulted in some adjustments in measurement time resulting in three measurement windows from 15 to 19 min, 21 to 27 min and 29 to 37 min. Measurements for all concentrations show increases in signal with additional incubation time. Response between 15 and 19 min revealed 10^6, 10^7, and 10^8 pfu/ml virus samples positive, although one of the two 10^6 pfu/ml sample was nominally above the 0.01 cutoff value. At later time points the 10^8, 10^7 and 10^6 pfu/ml virus samples produced greater signal. One of the 10^5 pfu/ml samples had increasing signal throughout the time course but remained negative, however, the second 10^5 sample became positive after 33 min. Lysates from BSC-40 cells were negative (Fig. 2b). Unlike the crude VZV preparations tested with the ESE Quant Reader, purified VZV was above the positive cutoff; however, the DMEM media used to resuspend VZV after purification was also positive. When diluted an additional 10-fold, both samples were negative throughout the measurement timeframe (Fig. 2b). Additionally, intensities for the purified VZV and DMEM media samples did not change over time indicating that the reaction observed is non-specific (i.e. does not titrate in this specific assay).

3.2. Blind study clinical sample screening with the Tetracore assay

Using visual analysis only, 9 of 11 clinical OPV samples were identified positively (Table 2). All samples were tested in duplicate resulting in four false negative tests (4 of 22 strip tests, or 18%). Among OPV positive samples some correlation between real time PCR Ct value and correct visual identification of OPV was seen as samples 3 and 8 had among the later Ct values (21 and 22, respectively) and were associated with the false negatives. There was however no correlation between the Ct value and line intensity. For example, sample 1 had a late Ct value (22) but produced the strongest intensity line while samples 10 and 11 had very early Ct values (17 and 15, respectively) but produced the weakest detectable lines with the assay. Modest correlation was seen between line intensity and titer; samples 1, 2, and 4 had titers above 4.6 x 10^3 pfu/ml and were positive with medium or weak line intensities, and samples 8, 9 and 11 had titers below 2.8 x 10^1 pfu/ml and were only weakly positive or negative. Sample 3, which contained 4.9 x 10^3 pfu/ml, was the exception as one strip became positive after the defined 15 min measurement time and the replicate was scored negative. VACV samples were identified as positive as few as 3 days post rash onset; days post rash were unavailable for MPXV samples. Non-OPV samples including VZV, pseudocowpox, Orf and HSV were also tested, with 10 of 11 samples correctly identified as negative (Supplementary Table 1). One positive was noted for one of the duplicate strip test for a single VZV sample; that sample was unusual visually, but marked positive as it had a weak intensity line at the sample line location.

4. Discussion

The Tetracore Orthopox BioThreat Alert assay is marketed as a rapid, point-of-care diagnostic for first responders in the event of a bioterrorism attack. Its ease of use and stability at ambient temperature suggest that the test would be applicable for use under field conditions which might be encountered by first-responders, or by rural health workers in geographic areas such as Central Africa where MPXV is endemic. In these situations, specimen transport and on-site laboratory capabilities would likely be limited. This is the first study to assess the BioThreat Alert assay utility for identifying laboratory OPV isolates and clinical samples. Visual analysis was preferred for its simplicity and used to evaluate initially assay performance. In all cases, the assay reproducibly allowed detection of 10^7 pfu/ml of VACV or MPXV, with no disagreement between replicates and independent users. Although only two OPVs were tested herein, they are known to possess broad cross-reactivity (Mercer et al., 2007) and results should be
similar for all OPV. Assay sensitivity, as defined for non-quantitative measurements as the threshold level, below which there is no detectable response and above which a response occurs (Delfert et al., 1987), would be 10^2 pfu/ml; all laboratory isolates tested at this concentration produced a positive response. Quantitative analysis with the ESEQunt® reader confirmed these results. Peak areas, as reported by the ESEQunt® were nearly equivalent for the 10^6 and 10^7 pfu/ml samples, and were 50–100-fold higher than negative control samples; the 10^6 pfu/ml VACV sample was only 5–10-fold above negatives. Thus, reproducible positive results can be anticipated with 10^7 pfu/ml samples, with some positives observed at the 10^6 pfu/ml concentration.

All visual experiments were read at 15 min after sample application according to the manufacturer instructions, which included a notation that samples that become positive after 20 min may not be valid. Extended run times are known to increase assay sensitivity but also can increase false positives (Wong and Tse, 2009). Observations suggested that a later measurement time point could increase sensitivity without sacrificing specificity. To test this assumption a dilution series of VACV was utilized and read with the field portable BioThreat reader. This reader provides quantitative measurements in fewer than 45 s and outputs positive or negative identification based on a default cutoff value of 0.01. Extending the assay incubation time from 15 to over 30 min improved VACV reader output as much as 4-fold, but did not increase background signals from control samples (Fig. 2a and b). Thus, sensitivity could be augmented by allowing longer incubation times as increases in signal were specific for VACV. Specifically, samples in the 10^5–10^6 range would likely benefit from additional incubation time. The 10^6 pfu/ml sample with measurements at 19 min (Fig. 2a) was visually negative at 15 min (Table 1, VACV Test 3), but was positive quantitatively by 19 min (SV = 0.0101), and was then easily identified as positive by 27 min (SV = 0.0289) post sample application. This correlates with visual analysis results where after 15 min many strips were determined to be negative, but would have been positive if read at a later time point. Furthermore, one of the two 10^5 samples became positive at 33 min. Additionally, while false positives were noted with purified lab grown VZV and the DMEM media used to resuspend the VZV virus, use of a longer incubation prior to test measurement would not have affected test outcome in these samples. However, a thorough testing of extended run times with clinical specimens is necessary to determine the maximum time allowed without significant increases in false positive results.

VZV has been difficult historically to grow to high titers, particularly when used in cell free preparations (Harper et al., 1998). In this case, the stock used was only 10^3 pfu/ml. Accounting for infective/inactive particles (Carpenter et al., 2009), the likely total virions in this cell-free purification were less than 10^2 pfu/ml. To address lower number of virions present relative to OPV samples, purified VZV and resuspension buffer were added to the strips after only a 1:2 dilution. Both resulted in false positives. Dilution of VZV virus or DMEM an additional 10-fold (to a 1:20 final dilution) reduced signals below the positive cutoff and eliminated these false positives. Thus, a component of the virus resuspension media and application of samples at such a minimal dilution was likely the reason for the false positives. Examination of signals from these false positives indicated no increase in signal over time. As a precaution for field screening assays, a minimum sample dilution should be established or limits placed on acceptable sample types. Any concerns of false positives can be minimized by adding multiple measurement times after sample application to observe specificity of the reaction indicated by increasing signal over time; field portable instruments such as the BioThreat Reader would allow quantization of multiple time point measurements.

Prior to testing clinical samples, the relationship between virus found in laboratory isolates and a real-time PCR E9L assay (Li et al., 2006) was assessed. The E9L assay is the current standard for identification of OPV in clinical specimens. A correlation between the real-time PCR extrapolated genome equivalents with the assay sensitivity to laboratory virus was established. Estimates for total particles to infectious ones range from 2.4 to 50:1 (Contreras and Ohlbaum, 1968; Overman and Sharp, 1959). Based on 10^7 pfu/ml sensitivity to laboratory grown virus, and using an intermediate value of 10:1 virus particles to infectious units suggests a detection limit of approximately 10^6 particles/ml, or assuming one particle contains one genome, 10^6 genomes/ml. In the 150 μl sample applied to each strip, this equates to 1.5 × 10^7 genomes. Vesiculopustular samples obtained between 5 and 13 days post rash onset during the 2003 monkeypox outbreak had genome copies ranging from 6 × 10^4 to 3 × 10^7. Therefore, for higher concentration clinical samples sensitivity would be adequate.

Within the context of the specificity determination for the Orthopox BioThreat Alert® assay, OPV samples with Ct values ranging from 15 to 22, or 1.5 × 10^7 to 2.3 × 10^5 genome equivalents were selected for testing. Orthopox BioThreat Alert false negative results with samples 3 and 8 had later Ct values of 21 and 22. Additionally, sample 8 had been re-extracted during initial diagnostic testing. Although this re-extracted sample Ct value was still strongly positive, it is possible that the integrity of some antigens was altered during re-processing which reduced binding of antigen to the Orthopox BioThreat Alert assay, but viral genomes were left intact. Variability of sample quality may have also affected the results and subsequent correlation to PCR Ct values. It is also of use to note that samples 1 and 5 had Ct values of 22, or approximately 2.3 × 10^5 genome equivalents. Based on estimations of non-infectious to infectious virions (10:1), this correlates with
2.3 \times 10^4 \text{ pfu/ml}. This concentration was negative with all tested laboratory viruses. This further suggests that sample integrity or quality may directly impact test results. As such, assay sensitivity to clinical specimens may prove to be better with freshly prepared field samples, although this remains to be proven. Additionally, while the number of clinical samples tested was limited, the diversity of samples as well as their well-characterized nature provides a good starting point for determining assay specificity.

5. Conclusion

In conclusion, VACV and MPXV samples with 10^6 pfu/ml were identified reproducibly by visual and quantitative methods when applied to the Bio Threat strips, and after extending the manufacturers recommended incubation time, concentrations as low as 10^5 pfu/ml produced positive results. While visual analysis was sufficient to identify positive readings, the availability of a field portable reader such as the Bio Threat Reader would allow documentation of intensity and standardize readings. When testing clinical samples there was variability in correlation with other methods to identify viruses such as RT-PCR or virus culture, but indications suggest the assay works well for higher concentration samples and discriminates well against other cutaneous rash causing viruses. These studies suggest that the Orthopox Biothreat Alert assay may be a valuable method for screening and identification of human monkeypox. Rapid diagnosis would allow prioritization of samples for further testing and would be crucial for implementation of infection control, outbreak detection, and clinical care.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jviromet.2012.08.023.

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