EVALUATION OF A RAPID ANALYTE MEASUREMENT PLATFORM AND
REAL-TIME REVERSE-TRANSCRIPTASE POLYMERASE CHAIN
REACTION ASSAY WEST NILE VIRUS DETECTION SYSTEM IN
MOSQUITO POOLS
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ABSTRACT. We evaluated the commercially available Rapid Analyte Measurement Platform (RAMP®) West Nile virus (WNV) antigen detection test for sensitivity and consistency with real-time reverse transcriptase polymerase chain reaction (RT-PCR) confirmation testing. Panels of samples consisting of WNV-spiked mosquito pools and negative control pools were sent to 20 mosquito abatement districts (MADs) that processed the pools using the RAMP assay. The samples were then sent to the reference laboratories used by the MADs for confirmation by real-time RT-PCR. Positive pools with virus titers of roughly 1–3 log_{10} PFU/ml had RAMP scores above the RAMP test positive cutoff score of 30 RAMP units, but these virus-positive samples could not be reliably confirmed by real-time RT-PCR testing. Pools with virus titers ≥4 log_{10} PFU/ml scored ≥50 RAMP units. Real-time RT-PCR results varied among the confirmation laboratories. With few exceptions, pools returning a RAMP score of ≥100 were confirmed with real-time RT-PCR, while pools returning a RAMP score of 50–99 appeared to be at the limit of real-time RT-PCR detection. Therefore, we recommend using a positive cutoff of 50 RAMP units with no real-time RT-PCR confirmation to maximize speed, efficiency, and economy of the RAMP assay. A more conservative approach would be to implement a “gray zone” range of 50–100 RAMP units. Pools scoring within the gray zone could be submitted for real-time RT-PCR confirmation with the understanding that positive pools may not confirm due to the inhibitory effect of the RAMP buffer on the real-time RT-PCR assay. We also conducted a series of experiments using laboratory-prepared mosquito pools spiked with WNV to compare mosquito homogenization buffers, pool sizes, and grinding methods in order to determine how these variables affect the RAMP and real-time RT-PCR assay results.

KEY WORDS West Nile virus, RAMP® test, real-time reverse transcriptase polymerase chain reaction, rapid detection, mosquito, vector

INTRODUCTION

Since the 1999 introduction of West Nile virus (WNV) into the USA and subsequent expansion of the virus across the country, health departments and mosquito abatement districts (MADs) have established or intensified existing mosquito-based arbovirus surveillance programs. Traditional methods for arbovirus surveillance include reverse-transcriptase polymerase chain reaction (RT-PCR) assays (Kaufman et al. 2003), cell culture plaque assays (Beaty et al. 1995), and enzyme immunoassays (Hunt et al. 2002). Rapid procedures, such as automated high-throughput arboviral RNA extraction protocols and real-time RT-PCR assays (Lanciotti et al. 2000), were developed and utilized to ensure efficient virus detection when testing the large numbers of mosquito pools required to accurately estimate the infection rate in mosquito populations (Gubler 2000). While these assays are excellent tools for arbovirus surveillance, they require significant monetary resources, technical expertise, and appropriate laboratory equipment and space.

Aided by Epidemiology and Laboratory Capacity funding from the federal government, state and local health departments expanded arbovirus surveillance. However, due to budget reductions over the years, many laboratories limited testing capacity or ceased mosquito pool testing altogether. In addition to the budget reductions, the time between submitting samples for testing and receiving the results was viewed as a significant limitation to using central laboratories for mosquito pool testing. Due to sample transport, implementation of weekly batch-processing schedules by the laboratories, and the time required to run the test, the lag from collection to test results can easily be a week or longer.

Since MADs often make mosquito control decisions using mosquito-based arbovirus surveillance data, these limitations were troublesome. Commercial companies responded by developing WNV antigen detection assays that are accurate, rapid, and available to entities with limited technical and laboratory resources. One such assay, the Rapid Analyte Measurement Platform (RAMP®) test developed by Response Biomedical (Response Biomedical Corp., Burnaby, British Columbia, Canada), was adopted by many MADs across the USA. Briefly, up to 50 mosquitoes are homogenized in a proprietary
RAMP grinding buffer and centrifuged. An aliquot of the resulting supernatant is mixed with a conjugated-antibody complex and added to the test cartridge that contains an immunochromatographic test strip. During a 90-min incubation period the sample migrates along the strip to the detection and internal control zones. The antigen-bound antibody complex is captured at the detection zone, and excess antibody conjugate is captured at the internal control zone. The cartridge is inserted into the RAMP reader, which reads and calculates the ratio between the fluorescence emitted at each zone to produce the results in RAMP units between 0 and 640. Mosquito pools producing ≥30 RAMP units are considered WNV positive according to the RAMP kit insert. Most MADs then send positive RAMP samples to laboratories for real-time RT-PCR confirmation.

While many MADs incorporated the RAMP assay into their surveillance programs successfully, some programs observed that samples that were RAMP positive in their laboratories produced negative results when tested with real-time RT-PCR in confirmation labs and sought advice from the manufacturer and the Centers for Disease Control and Prevention (CDC). Previously we had suspected that real-time RT-PCR may be inhibited by the grinding solution used for the RAMP assay and suggested a modified extraction protocol incorporating extra washes (Burkhalter et al. 2006), but due to the wide variability in sample handling, storage, and processing protocols among MADs and confirmation laboratories, we were unable to definitively determine the cause of the incongruent RAMP and real-time RT-PCR results. To address these discrepancies, we conducted several experiments using laboratory-prepared mosquito pools. To examine the WNV detection and confirmation system frequently incorporated by MADs and laboratories, and to determine a useful RAMP positive cutoff score, we created proficiency panels consisting of pools of whole mosquitoes spiked with known quantities of WNV and negative control pools to be tested by MADs and their confirmation labs by the RAMP and real-time RT-PCR assays. In addition, we conducted a series of in-house experiments at the CDC on laboratory-prepared mosquito pools spiked with WNV to compare grinding buffers, pool sizes, and grinding methods to determine how these variables affect the RAMP and real-time RT-PCR assay results.

**MATERIALS AND METHODS**

**Proficiency panel preparation**

Each proficiency panel consisted of 10 pools of 50 laboratory-reared female *Culex quinquefasciatus* Say, each containing an aliquot of medium that was either positive or negative for WNV. Tenfold serial dilutions of high-titered stock WNV strain CO08-10517 harvested from a pool of field-collected mosquitoes were prepared in RAMP buffer. These 7 dilutions contained virus titers of approximately 1–7 log<sub>10</sub> PFU/ml. Aliquots of 100 μl of each virus dilution were added to 7 mosquito pools to generate the positive pools for each panel. Aliquots of 100 μl of uninfected RAMP buffer were added to 3 pools of each panel to serve as negative controls. Panels were blind-coded and stored at −70°C until shipped on dry ice to 20 participating MADs in California, Idaho, Illinois, South Carolina, Utah, Virginia, and Washington. An additional 5 panels were tested in the CDC laboratory in Ft. Collins, CO, to provide a standard for comparison with the MAD and reference laboratory results.

**CDC proficiency panel testing methods**

The following procedures were used to test the 5 sample panels evaluated by RAMP and real-time RT-PCR at the CDC laboratory. One ml RAMP buffer and 1 copper-coated BB were added to each pool. Pools were homogenized for 4 min at 25 cycles/sec in a Qiagen Mixer Mill MM 300 (Qiagen, Inc., Valencia, CA) and centrifuged for 3 min at 4,000 rpm in a refrigerated centrifuge. Each pool was tested using the RAMP assay according to the manufacturer’s protocol in triplicate to determine the variation in RAMP scores for a given sample.

After processing the pools for the RAMP assay, RNA was extracted from 2 replicate aliquots from each pool, and each extract was tested for WNV by real-time RT-PCR with 2 primer sets in duplicate (resulting in 8 real-time RT-PCR results for each pool of each panel). RNA was extracted from a 100-μl aliquot of supernatant from each pool using Qiagen’s QIAmp Virus BioRobot 9604 kit on a Qiagen BioRobot 9604 according to the manufacturer’s protocol. Five microliters of each extraction were added to primers and probes specific to the 3’ noncoding (NC) and envelope (ENV) regions of the WNV genome and reagents of Qiagen’s Quantitect Probe RT-PCR kit and subjected to 45 amplification cycles in the Bio-Rad iCycler IQ<sup>TM</sup> Real-time Detection system (Bio-Rad, Hercules, CA) according to previously described cycling conditions (Lanciotti et al. 2000). Results are reported as the cycle threshold (CT) value for each real-time RT-PCR run (poles are considered positive if the CT score is ≤37).

**MAD proficiency panel testing methods**

We requested that each MAD perform the same processing and RAMP assay protocols on
the panel samples as they would on their regular field-collected mosquito pools. The MADs recorded protocol data and test results in a data collection form included with each panel. With 1 exception, all MADs homogenized the pools by vortexing the samples with 2 BBs in 1-ml RAMP buffer (vortexing intervals ranged from 1.5 to 12 minutes) and performed the RAMP assay according to the manufacturer’s protocol. One MAD, according to their state’s established protocol, ground their pools in a virus-transport medium provided by their state’s health department, and added 60 μl of supernatant to 60-μl RAMP buffer, and used this 120-μl aliquot to perform the RAMP assay as described earlier. All MADs forwarded the processed pools to their respective confirmation laboratories; MADs within a state sent their panels to the same confirmation laboratory.

Confirmation laboratory proficiency panel testing methods

The confirmation laboratories performed RNA extractions on each pool using comparable RNA extraction kits by Qiagen®, and Ambion® (Life Technologies, Grand Island, NY). Each laboratory used primers and probes specific to either the 3’ NC or ENV regions of the WNV genome to perform real-time RT-PCR on each pool (Lanciotti et al. 2000); some laboratories tested each pool with both primer sets. The confirmation laboratories used similar real-time RT-PCR kits from Qiagen, Ambion, and Invitrogen (Life Technologies, Grand Island, NY), and the thermocycler protocols used were nearly identical and as described previously (Lanciotti et al. 2000).

Processing variables experiment methods

Samples of varying pool sizes (i.e., number of mosquitoes in the pool) were processed at the CDC by 2 grinding methods (vortex for 1 min, Mixer Mill for 4 min at 25 cycles/sec) in 2 buffers (BA-1 cell culture medium, RAMP buffer) to evaluate the effects of these variables on RAMP and real-time RT-PCR detection of WNV.

Tenfold serial dilutions were made with high-titered stock WNV strain CO08-10142 containing virus titers of approximately 3.5–6.5 log_{10} PFU/ml, and 100 μl of each dilution was added to pools containing 1, 5, 10, 25, and 50 lab-reared Cx. quinquefasciatus. Aliquots of 100 μl of BA-1 cell culture medium without virus were added to additional pools that served as negative controls. One copper-clad BB and 900 μl BA-1 were added to pools of each viral titer and the negative controls. Half the pools of each titer were ground by vortex (1 min), and the other half were ground by the Qiagen Mixer Mill MM 300 (4 min at 25 cycles/sec). Grinding procedures for these pools were the same as for pools ground in RAMP buffer, and aliquots of supernatant were removed for subsequent real-time RT-PCR testing as described earlier in the CDC’s proficiency panel processing methods.

Statistical analysis methods

A linear model was used for the RAMP assay analysis, including estimated titer, grinding method and mosquito pool size as predictors, the logarithm of the RAMP values as the response variable, and the variance was modeled as a power function of the estimated titer for different pool sizes (Pinheiro and Bates 2000). This model was fitted using generalized least squares. A linear model was used for the real-time RT-PCR assay analysis, including the same predictors as in the RAMP assay analysis with the addition of buffer type, and the real-time RT-PCR CT values as a response. The best fitting model was chosen using restricted likelihood ratio tests (RLRT) to compare models. Diagnostic plots were created to ascertain appropriateness of model assumptions and fit. Once a best fitting model was selected, to characterize results using a measure of relevant practical application, we estimated the sensitivities of the assays by estimating the proportion of RAMP scores expected to exceed 50 and real-time RT-PCR CTs expected to be ≤37, for various estimated titer and pool size combinations. Estimates of the sensitivities and 95% confidence intervals (CI) were computed using estimates from the regression models described, as detailed below.

For a random sample \( X_1, X_2, ..., X_n \) from a normal distribution with mean \( \mu \) and variance \( \sigma^2 \), let \( \hat{\mu} \) and \( \hat{\sigma}^2 \) be the usual unbiased estimators of the mean and variance, respectively, and note that \( \hat{\mu} \) is normally distributed with mean \( \mu \) and variance \( \sigma^2/n \). Let \( \theta \) be a fixed value that we use to represent a cutoff value for observations from the common distribution of the \( \{X_i\} \), such that values
for which \( X_i > \theta \) are considered “positive,” and otherwise “negative.” We are interested in computing the probability that future values of \( X \) from this distribution are greater than \( \theta \), and we call this probability the sensitivity. Let \( W = (\theta - \hat{\theta})/\tilde{\sigma} \) and \( P = 1 - \Phi(W) \). The value \( P \) is therefore the sensitivity, and a point estimate for \( P \) is obtained from the observed value for \( W \) via the specification for \( \theta \) and the observed values for \( \hat{\theta} \) and \( \tilde{\sigma}^2 \). CIs for the sensitivity \( P \) can be obtained from its sampling distribution, the CDF for which is given by

\[
P[P \leq p] = P[1 - \Phi(W) \leq p]
\]

\[
= P[\hat{\theta} \leq \theta - \sqrt{\frac{\tilde{\sigma}^2}{n}}\Phi^{-1}(1-p)]
\]

\[
= \int_{-\infty}^{\infty} P[\hat{\theta} \leq \theta - \sqrt{\frac{\tilde{\sigma}^2}{n}}\Phi^{-1}(1-p)]
\]

\[
f(s^2; \sigma^2)ds^2
\]

\[
= \int_{-\infty}^{\infty} \Phi(\theta - \sqrt{\frac{\tilde{\sigma}^2}{n}}\Phi^{-1}(1-p); \mu, \frac{\sigma^2}{n})
\]

\[
f(s^2; \sigma^2)ds^2
\]

where \( f(s^2; \sigma^2) \) is the density of \( \hat{\sigma}^2 \sim \frac{\sigma^2}{n} \chi^2 \), a scaled \( \chi^2 \) distribution with \( n-1 \) degrees of freedom. From this CDF, a 100(1-\( \alpha \))% CI \((p_L, p_U)\) may be found by solving \( P[P \leq p_L] = \frac{\alpha}{2} \) and \( P[P \leq p_U] = 1 - \frac{\alpha}{2} \) for \( p_L \) and \( p_U \), which must be accomplished numerically. Computation of these expressions requires estimates for parameter values from a particular application.

In the regression modeling here, we assumed a normal regression model for the responses (log RAMP scores or real-time RT-PCR CT) as a function of covariates estimated titer, mosquito pool size, and grinding method. We also modeled the variance as a function of the estimated titer. Estimates of the regression parameters from these fits provide estimates for the mean, at specified values of the covariates, and variance function and degrees of freedom, at specific values of estimated titer, for use in the above formula for \( P[P \leq p] \).

Analyses were performed in R (www.r-project.org, ver. 2.15.3) and TIBCO Spotfire S+ (version 8.2.0, TIBCO Software, Inc., Somerville, MA).

RESULTS

Proficiency panel results: RAMP assay

Results of the proficiency panel RAMP assays are presented in Fig. 1. Data from 1 MAD which performed the alternate processing and RAMP protocol described earlier were excluded from the overall RAMP score analysis. All samples containing \( \geq 4 \log_{10} \) PFU/ml WNV produced results >50 RAMP units. Five of the 24 panel samples containing \( \sim 3 \log_{10} \) PFU/ml WNV produced RAMP results >30; the rest scored <30. No samples containing \( \sim 3 \log_{10} \) PFU/ml scored >50. With 2 exceptions, negative pools and pools containing approximately 1–2 \( \log_{10} \) PFU/ml WNV did not produce RAMP scores >30. One negative pool (of 72 total) produced results of 46.5 and 33.0 in 2 of the 3 replicates (the 3rd repeat scored <30). One pool containing \( \sim 2 \log_{10} \) PFU/ml produced a result of 640. None of the negative control pools produced a result \( \geq 50 \).

Proficiency panel results: real-time RT-PCR assay

Real-time RT-PCR results varied between laboratories, as seen in Fig. 2. Using a CT value of >37 as the negative real-time RT-PCR cutoff (Lanciotti et al. 2000), CDC and 4 confirmation laboratories were unable to detect WNV in negative pools and pools containing \( \leq 3 \log_{10} \) PFU/ml. The other 3 confirmation labs reported equivocal or positive results for some of the negative pools; the same confirmation laboratories also reported equivocal or positive results in some pools containing \( \leq 3 \log_{10} \) PFU/ml. The CDC and 4 confirmation laboratories were able to detect WNV in pools containing \( \geq 4 \log_{10} \) PFU/ml. The other 3 confirmation laboratories returned equivocal or negative results for pools containing \( \sim 4 \log_{10} \) PFU/ml, and 2 of these laboratories returned equivocal or negative results for pools containing \( \sim 5 \log_{10} \) PFU/ml. All laboratories were able to detect WNV in pools containing \( \sim 6 \log_{10} \) PFU/ml.

The comparison of RAMP results and CT values is also depicted in Fig. 2. With few exceptions, pools scoring \( \geq 100 \) RAMP units contained \( \geq 4 \log_{10} \) PFU/ml and were confirmed with real-time RT-PCR. Three pools scoring between 50 and 100 RAMP units were confirmed by real-time RT-PCR; however, the RAMP results were closer to 100 than 50 (81.3, 86.6, and 93.7) and pools scoring less than these were not confirmed by real-time RT-PCR in the proficiency panel evaluation.

Processing variables experiment results: RAMP assay

The RAMP results for the evaluation comparing grinding methods and pool sizes conducted at CDC are shown in Fig. 3. For samples containing 3.8 \log_{10} \) PFU/ml, pool size must be reduced to \( \leq 20 \) to affect qualitative RAMP interpretation (pos/neg); pools containing \( \geq 25 \) mosquitoes are qualitatively similar. Samples with titers \( >4 \log_{10} \)
PFU/ml were within the positive range (>50 RAMP units) even though the RAMP scores were lower as pool size increased; all samples with titers <3.6 log_{10} PFU/ml were negative (<50 RAMP units).

Results of the linear model including mosquito pool size and estimated titer indicated statistically significant interaction effects among these predictors and grinding method (RLRT \( P < 0.001 \)), complicating interpretation. We describe the comparisons relevant to the utility of the RAMP assay.

While mean RAMP score did significantly depend on the grinding method’s interaction with mosquito pool size and estimated titer, this dependence was not qualitatively important when evaluating RAMP assay sensitivity (i.e., the probability that a sample of a particular mosquito pool size and estimated titer will produce \( \geq 50 \) RAMP units). We therefore used a model excluding grinding method but retaining the mosquito pool size and estimated titer main effects and interaction for estimation of RAMP assay sensitivity; this in effect estimates the average sensitivity over the grinding method effect at each combination of mosquito pool size and estimated titer. As inspection of the RAMP scores in Fig. 3 suggests, reactivity of the RAMP assay was dynamic over the range of estimated titers used, with the estimated titer 3.8 log_{10} PFU/ml yielding the range of qualitative differences in sensitivity of interest when using a positivity cutoff of 50 for the RAMP score. Indeed, sensitivities using this cutoff for estimated titers below 2.8 log_{10} PFU/ml were nearly 0, for those at 3.6 log_{10} PFU/ml were all well below 0.5, and for those at least 4.5 log_{10} PFU/ml were nearly 1. In our evaluation, the sensitivities and 95% CIs for estimated titer 3.8 log_{10} PFU/ml for the different pool sizes evaluated are given in Table 1. For estimated titer 3.8 log_{10} PFU/ml and all mosquito pool sizes tested, sensitivities are at least 50%, while for samples with 15 or fewer individuals in the pools the sensitivities are generally above 95%.

**Processing variables experiment results: real-time RT-PCR assay**

All pools processed in RAMP buffer produced significantly higher real-time RT-PCR CT values when compared to pools of the same titer, size, and grinding protocols that were processed in BA-1 (Fig 4).

Linear models fit to real-time RT-PCR CT showed statistically significant differences in average CT between the 2 buffers, the magnitude
of which depended on the numbers of mosquitoes \((P < 0.01)\), while adjusting for estimated titer. Given that the RAMP assay and subsequent real-time RT-PCR confirmation are performed on samples processed in RAMP buffer and not BA-1, we consider only the RAMP buffer variable as we further analyze CT. Regression of CT on mosquito pool size, grinding method, and estimated titer resulted in the best-fitting model retaining number of mosquitoes and estimated titer (with both linear and squared terms) in the model \((P < 0.05\) for each); differences in average CT did not differ by grinding method when accounting for these variables \((P = 0.11)\). Dependence of CT on pool size and estimated titer resulted in different sensitivity estimates (and 95% CIs) by these values, and we report in Table 1 the estimated sensitivities for the observed mosquito pool sizes 5, 25, and 50, at estimated titer 3.8 log\(_{10}\) PFU/ml, values for which were estimated using the model.

As expected, sensitivity increases with increasing estimated titer, and sensitivities were estimated at essentially 100% for estimated titers at least 4.5 log\(_{10}\) PFU/ml.

**DISCUSSION**

The development of commercial WNV detection assays has allowed MADs across the USA to expand WNV surveillance in mosquitoes without the need for developing a costly virus isolation or PCR laboratory, enabling MADs to use surveillance data to guide abatement decisions and determine the effectiveness of mosquito control strategies. However, conflicting results from RAMP and real-time RT-PCR assays have been frustrating for MADs who have encountered them. One study that systematically examined RAMP and real-time RT-PCR data generated from field-collected mosquito pools described incongruent results between the 2 assays, the most troubling being cases where...
samples were RAMP-positive but real-time RT-PCR negative (Kesavaraju et al. 2012). These samples were described as “false-positive,” because detection of WNV by real-time RT-PCR is considered more sensitive than antigen detection (Kesavaraju et al. 2012). We have shown in this evaluation that real-time RT-PCR performed on samples processed in RAMP buffer is inhibited by the buffer and does not detect RNA in lower-titered pools. Our experiments used laboratory-generated WNV positive and negative mosquito pools to serve as controls so we could accurately gauge the performance of the RAMP and real-time RT-PCR assays together as part of a WNV detection system utilized by many states, and to subject RAMP and real-time RT-PCR to varying conditions to determine their effects on each assay.

Almost all of the RAMP results generated from the proficiency panels fell within an expected range of RAMP units for each titer (Fig. 1). Using the manufacturer’s suggested RAMP score of ≥30 RAMP units as the positive cutoff for mosquito pools is represented by the dashed line; our proposed cutoff of ≥50 RAMP units is represented by the solid line. Samples that were vortexed are represented by black diamonds; samples processed by Mixer Mill are shown as open diamonds.

Table 1. Sensitivities (i.e., the probability that a positive sample will produce a positive result in the RAMP or real-time reverse-transcriptase polymerase chain reaction [RT-PCR] assays) and 95% confidence intervals (CI) of samples containing 3.8 log10 PFU/ml West Nile virus (WNV) using a positivity cutoff of ≥50 for RAMP units and ≤37 for real-time RT-PCR cycle threshold (CT) values.

<table>
<thead>
<tr>
<th>Pool size</th>
<th>No. of pools</th>
<th>RAMP scores, ≥50 RAMP units</th>
<th>Real-time RT-PCR, CT values ≤37</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Sensitivity (%)</td>
<td>95% CI (%)</td>
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<tr>
<td>50</td>
<td>10</td>
<td>56.5</td>
<td>51.3, 61.8</td>
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<td>25</td>
<td>10</td>
<td>67.8</td>
<td>62.7, 72.9</td>
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<tr>
<td>20</td>
<td>10</td>
<td>87.6</td>
<td>83.9, 90.9</td>
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<tr>
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<td>10</td>
<td>99.9</td>
<td>99.9, 1.00</td>
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<tr>
<td>10</td>
<td>9</td>
<td>88.4</td>
<td>84.9, 91.7</td>
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<tr>
<td>5</td>
<td>9</td>
<td>99.8</td>
<td>99.6, 99.9</td>
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<tr>
<td>3</td>
<td>10</td>
<td>96.3</td>
<td>94.4, 97.8</td>
</tr>
<tr>
<td>1</td>
<td>10</td>
<td>97.6</td>
<td>96.2, 98.7</td>
</tr>
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NT, not tested.
produced false negatives for all but 4 samples with titers of \( \leq 3 \log_{10} \text{PFU/ml} \); however, this was expected based on previous sensitivity evaluations (Burkhalter et al. 2006, Sutherland and Nasci 2007). One MAD’s sample containing \( 2 \log_{10} \text{PFU/ml} \) WNV produced an unexpected RAMP result of 640, while all other samples with that titer produced negative RAMP results. The reason for this discrepancy is unknown. All but 1 negative control produced a result \( \geq 30 \) RAMP units. Changing the RAMP positive cutoff from \( < 30 \) RAMP units would not significantly impact the sensitivity limits of the RAMP assay as determined previously (Burkhalter et al. 2006) because most of the pools in the panel that scored less than 50 also scored less than 30 and thus would be interpreted as negative using either cutoff. However, raising the cutoff to 50 eliminates the chance that false positives will be produced by negative pools. No negative control pools in this evaluation or any we have conducted thus far have produced RAMP results \( \geq 50 \).

Factors that may account for the inability to obtain RT-PCR confirmation of RAMP positive samples in field-collected mosquito pools may be the improper utilization of the RAMP, mishandling of samples before or after RAMP testing and during transit to confirmation laboratories, or the effect of the RAMP buffer on the RT-PCR reaction. In our assessment, it appears unlikely that the RAMP assay was improperly conducted because we were able to ascertain from protocol data reported as part of the proficiency panel evaluation by the MADs that the RAMP assay protocol was followed satisfactorily. To promote proper use of the RAMP kit, Response Biomedical (2010) has published technical documents providing guidelines for handling, processing, and testing samples beyond what is described in the RAMP assay kit insert. It is possible that sample degradation due to high temperatures or repeated freeze-thaws in shipping and handling may render WNV RNA in the samples undetectable in real-time RT-PCR. In our proficiency panel evaluation frozen panels were sent to MADs on dry ice and samples were kept cold before being ground and tested with the RAMP assay. In the processing variables evaluation mosquito pool samples were kept frozen at \(-70^\circ C\) until ground and tested with the RAMP and real-time RT-PCR assays.

A more likely cause for incongruent RAMP and real-time RT-PCR results appears to be the effect of the RAMP buffer on the ability of real-time RT-PCR to detect WNV RNA. In both the proficiency panel and processing variables comparison evaluations, RAMP samples containing \( \leq 3 \log_{10} \text{PFU/ml} \) were undetectable by real-time RT-PCR. By comparing real-time RT-PCR results of WNV positive samples ground in RAMP buffer and BA-1, we were able to show a significant increase in real-time RT-PCR CT values of the samples ground in RAMP buffer. Beyond general reduction in sensitivity due to inhibition by RAMP buffer, however, we found that real-time RT-PCR results and sensitivity limits varied among labs that participated in the proficiency panel evaluation; in some cases even samples with WNV titers of \( \geq 4 \log_{10} \text{PFU/ml} \) were not confirmed. Determining the causes of the lower sensitivity in these labs was beyond the scope of this evaluation and not investigated further. Response Biomedical (2008) recommends freezing an aliquot of supernatant from samples to be tested with real-time RT-PCR at \(-70^\circ C\) as soon as possible to reduce RNA degradation. However, the best results from real-time RT-PCR testing of pools spiked with WNV were produced when the supernatant was extracted for RNA before the sample was frozen and little time had passed between processing and RNA extraction (Response Biomedical 2008).
We also tested pools of various sizes ground in RAMP buffer using 2 different grinding methods to determine the effects of these variables on RAMP and real-time RT-PCR results. Even though RAMP scores decreased as pool size increased the results remained in the positive RAMP unit range for pools containing $\geq 4 \log_{10}$ PFU/ml. Decreasing the pool size did not affect the qualitative negative results of samples containing $\leq 3 \log_{10}$ PFU/ml when using a RAMP cutoff of $\leq 50$. Operationally, testing the maximum pool size of 50 individuals does not diminish the RAMP’s ability to detect WNV in samples containing $\geq 4 \log_{10}$ PFU/ml, nor does decreasing the pool size to $< 50$ enhance RAMP’s ability to detect WNV in samples containing $\leq 3 \log_{10}$ PFU/ml, regardless of grinding method used. For pools containing $3.8 \log_{10}$ PFU/ml, reducing pool size to $\leq 20$ individuals increases the number of pools that would be expected to produce $\geq 50$ RAMP units, and vortexed pools containing $\leq 20$ mosquitoes produced higher RAMP results than pools of the same size ground by Mixer Mill (Table 1). However, we suspect that the few numbers of previously undetected samples in this narrow titer range that would be detected by reducing pool sizes does not warrant incurring the additional cost of testing more pools.

Based on the data generated by our evaluations, we offer 2 recommendations for interpreting RAMP assay data. To maximize the speed, efficiency, and economy of testing mosquito pools with the RAMP assay, the RAMP positive threshold could be set at 50 RAMP units, with no real-time RT-PCR confirmation testing required. In this evaluation samples producing RAMP scores $\geq 100$ contained approximately $\geq 4 \log_{10}$ PFU/ml and were within the sensitivity limits of real-time RT-PCR confirmation of samples processed in RAMP buffer. A more conservative approach would be to set the RAMP positive threshold at 100 above which samples do not require real-time RT-PCR to be considered positive and to implement a “gray zone” of 50–99 RAMP units. Samples that produce scores within the established “gray zone” would be confirmed by real-time RT-PCR to be considered positive. In the CDC processing variables evaluation, pools of 25 and 50 individuals containing between 3.6–4 $\log_{10}$ PFU/ml produced RAMP results between 50 and 100, and most were confirmed by real-time RT-PCR. In the proficiency panel evaluation, a few pools of 50 mosquitoes containing approximately $4 \log_{10}$ PFU/ml produced RAMP scores between 80 and 100, and all were confirmed by real-time RT-PCR. These data support the cautionary warning that mosquito pools processed in RAMP buffer producing scores within the range of 50–100 RAMP units are at the limit of real-time RT-PCR detection and may or may not confirm even if positive. Implementing either recommendation would reduce the number of RAMP positive samples needing to be confirmed using real-time RT-PCR.

In the proficiency panel evaluation we were limited to evaluating pools of 1 mosquito species and pool size, and all mosquitoes were deplete (not blood fed). These constants, as well as the standard processing procedures and assay protocols used in this study do not account for the wide variability in sample collecting, handling, processing and testing that field-collected mosquitoes may be subjected to across the country by MADs and confirmation laboratories. We recommend that samples are handled carefully and maintained in a cold chain, and that the guidelines from the manufacturer are heeded to ensure the best results from the RAMP assay and greatest likelihood of correlating real-time RT-PCR results. When the RAMP assay is being performed correctly by the MAD, careful considerations should be made to determine whether sending pools for additional confirmation by real-time RT-PCR is necessary when pools score above a RAMP cutoff of 50 or 100. Results from this evaluation suggest that from an operational standpoint, the RAMP assay is a moderately sensitive and accurate tool for MADs to incorporate in their surveillance programs in lieu of molecular detection capabilities.

ACKNOWLEDGMENTS

We are extremely grateful to all study participants at agencies in Idaho (Gem County MAD, Canyon County MAD, Ada County MAD, Idaho Bureau of Laboratories), California (Consolidated MAD, Contra Costa MVCD, Turlock MAD, Coachella Valley MVCD, University of California–Davis), South Carolina (Beaufort County Mosquito Control, South Carolina Department of Health and Environmental Control), Illinois (Des Plaines Valley MAD, North Shore MAD, Northwest MAD, University of Illinois), Utah (Southwest MAD, Utah County MAD, Salt Lake City MAD, Utah Department of Health), Washington (Benton County MCD, Cowlitz County MCD, Clark County MCD), Oregon (Oregon State University), Virginia (Henrico County Department of Public Works, Fairfax County Health Department, Alexandria Health Department), and Pennsylvania (Pennsylvania WNV Control Program).

REFERENCES CITED


Response Biomedical Corporation. 2010. Storage and handling of WNV samples for testing by RAMP® and PCR. Technical guide 51173. Burnaby, British Columbia, Canada: Response Biomedical Corp.