Sindbis and Middelburg Old World Alphaviruses Associated with Neurologic Disease in Horses, South Africa

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Old World alphaviruses were identified in 52 of 623 horses with febrile or neurologic disease in South Africa. Five of 8 Sindbis virus infections were mild; 2 of 3 fatal cases involved co-infections. Of 44 Middelburg virus infections, 28 caused neurologic disease; 12 were fatal. Middelburg virus likely has zoonotic potential.

Alphaviruses (Togaviridae) include zoonotic, vectorborne viruses with epidemic potential (1). Phylogenetic analysis defined 2 monophyletic groups: 1) the New World group, consisting of Sindbis virus (SINV), Venezuelan equine encephalitis virus, and Eastern equine encephalitis virus; and 2) the Old World group, consisting of Semliki Forest virus (SFV), Middelburg virus (MIDV), Ndumu virus, Chikungunya virus (CHIKV), and Barmah Forest virus (2). Old World alphaviruses are associated mainly with febrile disease and arthralgia, are often accompanied by a maculopapular rash, and are rarely fatal, although neurologic cases have been reported (3). In contrast, New World alphaviruses are associated with neurologic disease in horses and, potentially, humans (4).

We previously investigated horses as sentinels for detection of neurologic arboviruses and described West Nile virus (WNV) lineage 2 (5) and Shunivirus (SHUV) as previously missed causes of fatal encephalitis in Africa (6), with zoonotic potential (7). Five alphaviruses have been detected in vectors in southern Africa: SINV, CHIKV, MIDV, Ndumu virus, and SFV (8); however, little is known about prevalence, pathogenicity, and host range (9).

The Study

To determine if alphaviruses may contribute to undefined neurologic infections, we investigated specimens (blood, cerebrospinal fluid, or tissue from brain, spinal cord, or visceral organs) from 623 horses with unexplained febrile and acute neurologic infections reported to our surveillance program by veterinarians across South Africa during January 2008–December 2013. Of reported cases, 346 horses had neurologic signs; 277 had mainly febrile illness and other miscellaneous signs, including colic and sudden death (online Technical Appendix Figure 1, panel A, http://wwwnc.cdc.gov/EID/article/21/12/15-0132-Techapp.pdf). Formalin-fixed tissue samples from horses that died were submitted for histopathology. Horses ranged from <1 to 20 years of age and included thoroughbred, Arabian, warmblood, and part-bred horses; most were bred locally.

A generic nested alphavirus nonstructural polyprotein (nsP) region 4 gene reverse transcription PCR (10) was used to screen total nucleic acids. TaqMan probes (Roche, Indianapolis, IN, USA) were developed for rapid differentiation of MIDV and SINV by real-time PCR (online Technical Appendix).

PCR-positive cases were confirmed by sequencing the nsP4 amplicon (200 bp), followed by maximum-likelihood analysis (Figure 1, panel A). Additional amplification, sequencing, and phylogenetic analysis of a 349-bp E1 gene fragment (online Technical Appendix) was attempted (Figure 1, panel B) to investigate recombination events.

Baby hamster kidney cell–culture isolates were obtained for 2 MIDV strains under Biosafety Level 3 conditions. Isolate SAE25/2011 from blood of a horse with neurologic signs was visualized by electron microscopy (Figure 2); the full genome was sequenced as described (11). By using maximum-likelihood and P-distance analysis (online Technical Appendix Figure 2), we compared isolate SAE25/2011 with MIDV-857, which was isolated in 1993 from spleen of a horse with African horse sickness virus (AHSV)–like signs in Zimbabwe (GenBank accession no. EF536323) (12). Differential diagnosis for flaviviruses, WNV, Wesselsbron virus, SHUV, and equine encephalitis viruses was performed on all specimens, which were sent to other laboratories for testing for AHSV, equine herpes viruses, and rabies (5,6).

Of the 623 horses, 8 (1.3%) tested positive for SINV and 44 (7.1%) for MIDV (Table 1). Of the 8 horses with SINV, 3 survived febrile illness, 2 survived neurologic disease, and 3 died from neurologic disease. Two of the 3 horses that died with SINV had WNV co-infection and were positive by PCR for both viruses in brain tissue. In those 2 horses, lesions of meningoencephalitis were visible and acute neurologic infections reported to our surveillance program by veterinarians across South Africa during January 2008–December 2013.
Figure 1. Maximum-likelihood trees of strains of Middelburg virus and Sindbis virus identified in horses in South Africa relative to other members of the alphavirus genus. Trees were constructed by using the Tamura-Nei substitution model and midpoint rooted with MEGA5 (http://www.megasoftware.net/). Scale bar indicates 0.1 nt substitutions. Estimates were constructed on the basis of bootstrap resampling performed with 1,000 replicates. Confidence estimates >70 are shown. A) A 200-bp fragment of the nonstructural polyprotein region 4 gene of MDV- and SINV-positive cases. B) A 348-bp fragment of the E1 gene of 7 MIDV cases identified in horses in southern Africa (genome position 10543–10911 corresponding with the MIDV-857 strain in GenBank accession no. EF536323). Reference sequences used in these trees are as previously described (2). Complexes are identified as follows: EEE, Eastern equine encephalitis; MID, Middelburg; SF, Semliki Forest; VEE, Venezuelan equine encephalitis; WEE, Western equine encephalitis. Viruses are identified as follows: AURAV, Aura virus; BEBV, Bebaru virus; BFV, Barmah Forest virus; CABV, Cabassou virus; CHIKV, Chikungunya virus; CEEVV, Eastern equine encephalitis virus; EILT, Ella virus; EEEV, Everglades virus; FMV, Fort Morgan virus; GETV, Getah virus; HJV, Highlands J virus; MAYV, Mayaro virus; MIDV, Middelburg virus; MUCV, Mucambo virus; NDUV, Nduvu virus; ONNV, O’nyong nyong virus; PIXV, Pixuna virus; RNV, Rio Negro virus; RRV, Ross River virus; SAE, South Africa equine virus; SAGV, Sagiyama virus; SESV, Southern elephant seal virus; SFV, Semliki Forest virus; SINV, Sindbis virus; SPDV, Salmon pancreatic disease virus; TONV, Tonate virus; TROV, Trocara virus; UNAV, Una virus; VEEV, Venezuelan equine encephalitis virus; WEEV, Western equine encephalitis virus; WHATV, Whataroaviruses; ZRU, Zoonoses Research Unit virus.
by histopathology but were nonspecific and could not be ascribed to either virus. Immunohistochemistry (IHC) for WNV in equine cases is reportedly a poor diagnostic tool (13), although IHC for SINV and MIDV needs to be established. In the third fatal case, SINV was detected in blood; neurologic samples were unavailable for testing. One horse with SINV had mild colic, dysphagia with tongue paralysis, and pale mucus membranes (Table 2).

Of 44 horses with MIDV, 16 had febrile disease, 28 had neurologic disease, and 12 died. Five dual infections were detected: 1 horse with MIDV and SHUV infections and 2 with MIDV and EEV infections survived; 1 horse with MIDV and AHSV infections and 1 with MIDV and WNV infections died (Table 2). For 6 of 12 horses that died with MIDV, central nervous system samples were available, and MIDV was detected in brain of all 6 horses. AHSV was detected in lung and spleen and MIDV in brain of 1 horse. MIDV and WNV was found in brain of another; only MIDV was detected in brain of the other 4. Altogether, 26 (66.7%) of 39 horses with MIDV single infection showed signs of neurologic disease (Table 2). Clinical signs with less severe illness included fever, stiffness, swollen limbs, hyperreactiveness, and depression; signs of severe neurologic disease included ataxia, paresis, paralysis, recumbency, and seizures. One horse exhibited icterus, 1 had a pregnancy abort, and 1 had laminitic stance. Brain and cord tissues from 4 horses with neurologic disease caused by MIDV single infections were examined microscopically. MIDV was detected in blood of 1 horse but in brain of the others. Lesions of mild to moderate meningoencephalitis were observed in all 4, including perivascular cuffing involving mainly mononuclear cells, glial nodules, and diffuse gliosis.

No clear associations with age, gender, or breed were apparent for either virus. Number of cases peaked during the rainy season (February–May), consistent with vectorborne diseases, although sporadic cases were detected year-round (online Technical Appendix Figure 1, panel B). Both viruses were widely distributed across South Africa (online Technical Appendix Figure 3).

The nsP4 gene fragment for 6 SINV-positive specimens had ≤7% nt differences but 100% aa identity, clustering closely to the Ockelbo strain from Sweden and Babanki strain from Cameroon. One SINV isolate was of unrecorded origin (Figure 1, panel A) (2).

MIDV nsP4 partial sequences of 26 unique strains clustered with MIDV-857 (12), (Figure 1, panel A), with 3.8% nt and 2.8% aa differences from South Africa strains and ≤5% nt and 3.6% aa differences from MIDV-857. No specific relationships to time, geographic origin, or outcome were evident, although the length and conserved nature of the diagnostic amplicon limited the phylogenetic analysis. The E1 gene fragments from 7 MIDV cases (GenBank accession nos. JN226792–JN226795, KF680222–KF680224) clustered with MIDV-857 (Figure 1, panel B), differing by ≤1.2% nt and 0.5% aa levels from each other and by ≤1.4% nt and 0.6% aa levels from MIDV-857.

The genome of MIDV isolate SAE25/2011 (GenBank accession no. KF680222) was 11,674 nt in length, excluding the poly (A) tail, with 98.5% nt and 99.4% aa identity to MIDV-857 (12), clustering similarly as previous alphavirus phylogenetic investigations (2). Seven aa differences exist in the structural polyprotein (98.7% identity) and 17 in the nsP region (99.3% identity). Three aa changes in the nsP and 3 in the structural polyprotein altered hydrophilicity. The nsP1 amino acid sequences differed by 0.8%; the nsP2 and nsP3 were identical; and nsP4 differed by 0.5%. The capsid protein differed by 1.1%; E1 and E2 differed by 0.5%; and E3 and 6K proteins were identical. SAE25/2011, like MIDV-857, also contained the recombinant SFV domains identified previously (12).
Table 1. Prevalence of neurologic and febrile cases of MIDV and SINV infections in horses, South Africa, 2008–2013*

<table>
<thead>
<tr>
<th>Alphavirus results</th>
<th>2008</th>
<th>2009</th>
<th>2010</th>
<th>2011</th>
<th>2012</th>
<th>2013</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIDV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Specimens, no.</td>
<td>76</td>
<td>50</td>
<td>137</td>
<td>166</td>
<td>107</td>
<td>100</td>
<td>636 (100)</td>
</tr>
<tr>
<td>PCR+, no. (% of total specimens)</td>
<td>0 (2)</td>
<td>10 (16)</td>
<td>26 (16)</td>
<td>4 (4)</td>
<td>3 (3)</td>
<td>44 (7)</td>
<td></td>
</tr>
<tr>
<td>Deaths, no. (% of no. PCR+)</td>
<td>NA</td>
<td>0</td>
<td>2 (50)</td>
<td>7 (27)</td>
<td>3 (75)</td>
<td>2 (27)</td>
<td>12 (27)</td>
</tr>
<tr>
<td>Detected in CNS, no. (% of no. PCR+)</td>
<td>NA</td>
<td>0</td>
<td>6 (23)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>6 (14)</td>
<td></td>
</tr>
<tr>
<td>Neurologic disease, no. (% of no. PCR+)</td>
<td>NA</td>
<td>1</td>
<td>7 (70)</td>
<td>16 (62)</td>
<td>4 (100)</td>
<td>2 (50)</td>
<td>30 (68)</td>
</tr>
<tr>
<td>Fever only, no. (% of no. PCR+)</td>
<td>NA</td>
<td>0</td>
<td>6 (60)</td>
<td>5 (19)</td>
<td>2 (50)</td>
<td>0 (0)</td>
<td>13 (30)</td>
</tr>
<tr>
<td>Co-infections, no. and type</td>
<td>NA</td>
<td>0</td>
<td>1 EEV, 1 WNV†</td>
<td>0 (0)</td>
<td>2 EEV, 1 ASHV†</td>
<td>1 EEV, 1 AHSV†</td>
<td>1 SHUV, 1 WNV†</td>
</tr>
</tbody>
</table>

| SINV                |      |      |      |      |      |      |       |
| Specimens, no.      | 76   | 50   | 137  | 166  | 107  | 100  | 636   |
| SINV+, no. (% of total specimens) | 2 (3) | 4 (8) | 0 (0) | 21 (50) | 0 (0) | 1 (100) | 8 (1) |
| Deaths, no. (% of no.) | 0 (0) | 2 (50) | NA | NA | 1 (100) | 0 (0) | 3 (38) |
| Detected in CNS, no. (% of no.) | NA | 0 | 2 (50) | NA | NA | 0 (0) | 4 (50) |
| Neurologic disease, no. (% of no.) | 1 (50) | 2 (50) | NA | NA | 1 (100) | 0 (0) | 4 (50) |
| Fever only, no. (% of no.) | 1 (50) | 2 (50) | NA | NA | 0 (0) | 1 (100) | 4 (50) |
| Co-infections, no. and type | 1 EEV, 1 WNV† | NA | NA | NA | 0 (0) | 1 EEV, 1 AHSV† | 2 WNV† |

*SINV, African horse sickness virus; CNS, central nervous system sample; EEV, equine encephalitis virus; MIDV, Middelburg virus; NA, not applicable; SHUV, Shuvivirus; SINV, Sindbis virus; WNV, West Nile virus; †, undetectable in central nervous system sample; ††, positive.

Conclusions

Besides CHIKV, SINV is the most widely distributed Old World alphavirus. It is associated with fever, rash, and arthralgia in humans in Europe, Asia, Africa, and Australia (8) and has been isolated from mosquitoes, birds, and humans in South Africa. Although antibodies described in livestock and horses, it has been unrecognized as a potential pathogen of horses. Less is known about MIDV. Isolated in 1957 from mosquitoes in South Africa, subsequent surveys identified antibodies in humans, horses, and livestock, and the virus has been isolated in mosquitoes and 2 humans elsewhere in Africa (14). In 1974, MIDV was isolated from blood of a horse during an outbreak of fever and icterus and was implicated in a horse with anorexia and muscular stiffness in South Africa (14) and from spleen of a horse in Zimbabwe in 1993 (11). Additional investigations have been limited, and the virus has not previously been associated with neurologic disease.

This study was not structured to provide definitive information on the prevalence and incidence of disease and likely underestimates the situation. Development of serologic assays for diagnosis of cases past the viremic stage are needed to establish the true prevalence and effects of disease, and IHC is needed for investigating the pathogenesis in horses and other species. Our findings strongly suggest that Old World alphaviruses, particularly MIDV, may constitute an overlooked cause of febrile and neurologic disease in horses and, like New World alphaviruses, may pose threat to horses, livestock, and humans.

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References


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Technical Appendix

Detailed Methods and Results

1. Generic Real-Time Reverse Transcription PCR for Diagnosis of Middelburg and Sindbis Virus Infections

Viral RNA was extracted from plasma or serum samples derived from EDTA-clotted blood and from cerebrospinal fluid samples with the High Pure Viral Total Nucleic Acid Extraction Kit (Roche, Indianapolis, IN, USA). Nervous tissue or visceral organ samples were homogenized by using a mortar and pestle, and RNA was extracted by using the RNeasy Plus Mini Kit (QIAGEN, Valencia, CA, USA).

The first round and nested PCR primers that were based on a highly conserved region of the nsP4 gene were prepared as described previously (1). First-round PCR was performed by using the Titan one tube reverse transcription (RT)-PCR system (Roche). Each reaction contained 10 μL of RNA, 10 μL 10× reaction buffer, 1 μL of dNTP mix (10 mmol/L), 2 μL of 10 pmol of each primer (Alpha1+; Alpha1–), 2.5 μL Dithiothreitol solution (100 mmol/L), 0.25 μL RNase inhibitor (40 U/μL), and 1 μL of the Titan enzyme mix. The final volume was made up to 50 μL with distilled water. Cycling conditions were 50°C for 30 min, 94°C for 2 min, (94°C for 10 s, 52°C for 30 s, 68°C for 1 min) × 35 cycles, and 68°C for 7 min.

By using Primer 3 (2), two probes were designed for rapid differentiation of Sindbis virus (SINV, a New World virus) and Middelburg virus (MIDV, an Old World virus) as follows: SINV: 5’ ATGACGAGTATTGGGAGGAGTTG 3’-FAM; MIDV: 5’ GCTTTAAGAAGTACGCATGCAACA 3’ –VIC. The probe position for MIDV is 6,132 by using MIDV EF536323 as reference and 6,478 for SINV (AF103734).
Differentiation of MIDV and SINV was performed on first-round RT-PCR products by real-time PCR by using the Lightcycler TaqMan Master kit (Roche): 2 μL of RT-PCR product was used with 1 μL of 20 pmol of each primer (Alpha2+; Alpha2–), 0.2 μL of 10 pmol probe each for SINV and MIDV, 4 μL of FastStart enzyme mixture to a final volume of 20 μL. The following program was followed on a Lightcycler 2 machine (Roche): preincubation at 95°C for 10 min (95°C for 10 s, 52°C for 1 min, 72°C for 1 s) for 45 cycles. Qualitative analysis that used different channels could distinguish between SINV and MIDV.

2. Alphavirus E-Protein Amplification

SINV E-Protein RT-PCR

Titan one tube RT-PCR system (Roche) was used for first-round PCR by using primer SIN8136EF (5’ TCGTCAGCATACGACATGGAG 3’) and A2 (3). PCR amplification began at 94°C for 2 min and 40 cycles of 94°C for 10 s, 52°C for 30 s and 72°C for 1 min with a final elongation step of 72°C for 7 min.

Expand High FidelityPLUS PCR system (Roche) was used for the semi-nested reaction. A total of 5μL of first-round product was used with primers SIN8136EF and SIN8787ER (5’GTATCCAAACTGGCGGGAAGT 3’). The following cycling program was used: 94°C for 2 min and 10 cycles of 94°C for 30 s, 52°C for 30 s, and 72°C for 1 min; 25 cycles of 94°C for 30 s, 52°C for 30 s, and 72°C for 1 min + 10 s/cycle with a final elongation step of 72°C for 7 min.

MIDV E-Protein RT-PCR

Titan one tube RT-PCR system (Roche) was used for first-round PCR with primer MID10475E+ (5’ GGTGCACGTCCATATACCC 3’) and MID11045E (5’ TCCCAATAGCATACTACAC 3’). Cycling began at 94°C for 2 min and 40 cycles of 94°C for 30 s, 48°C for 45 s, and 72°C for 1 min with a final extension of 72°C for 7 min.

Nested PCR was performed with Expand High FidelityPLUS PCR system (Roche) by using 2μL of first round template, 5μL Q-solution (QIAGEN, Valencia, CA, USA), and primers MID10543EN+ (5’ TGAACCACAAGGCCCTCCTTC 3’) and MID10911EN- (5’ CACTTTGCTGTGCAAGTGGT 3’). Cycling commenced at 94°C for 2 min and 40 cycles of 94°C for 30 s, 50°C for 45 s and 72°C for 1 min, with a final elongation step of 72°C for 7 min.
References


Technical Appendix Figure 1. Seasonal distribution of febrile and neurologic alphavirus–positive cases in South Africa. A) Bar chart showing monthly distribution of 623 cases of febrile and neurologic disease investigated in horses in South Africa during 2008–2013. Chart shows proportion of cases resulting in fatalities. B) Seasonal distribution of Middelburg virus (MIDV) cases investigated in horses in South Africa during 2008–2013. C) Seasonal distribution of Sindbis virus (SINV) cases investigated in horses and identified by reverse transcription PCR in South Africa during 2008–2013.
Technical Appendix Figure 2. A maximum-likelihood tree of the complete genome of Middelburg virus (MIDV SAE25/2011; KF680222) isolated in 2010 from a horse with neurologic disease, relative to strain MID-857 (identified as MIDV in the tree) isolated in 1993 from a horse that died with signs similar to those of African horse sickness virus. The tree was constructed with the Tamura-Nei substitution model by using the Mega 5 program (http://www.megasoftware.net/). Scale bar indicates 0.1 nt substitutions. Estimates were constructed on the basis of bootstrap resampling performed with 1,000 replicates. Complexes are identified as follows: EEE, Eastern equine encephalitis; SF, Semliki Forest; VEE, Venezuelan equine encephalitis; WEE, Western equine encephalitis. Viruses are identified as follows: BEBV, Bebaru virus; BFV, Barmah Forest virus; CHIKV, Chikungunya virus; EEEV, Eastern equine encephalitis virus; EVEV, Everglades virus; GETV, Getah virus; MAYV, Mayaro virus; MIDV, Middelburg virus; NDUV, Ndumu virus; ONNV, O’nyong nyong virus; RRV, Ross River virus; SAE, South Africa equine virus; SAGV, Sagiyama virus; SESV, Southern elephant seal virus; SFV, Semliki Forest virus; SINV, Sindbis virus; SPDV, Salmon pancreatic disease virus; TONV, Tonate virus; UNAV, Una virus; VEEV, Venezuelan equine encephalitis virus.
Technical Appendix Figure 3. Geographic distribution of alphavirus cases investigated in horses in South Africa, 2008–2013. Map shows numbers of cases of Sindbis virus (SINV) and Middelburg virus (MIDV) for specific locations. White area inside map indicates the independent nation of Lesotho.