# Detection of Novel Orthobunyavirus Reassortants in Fatal Neurologic Case in Horse and Culicoides Biting Midges, South Africa

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We detected Shuni virus in horses and ovine fetuses and Shamonda virus in a caprine fetus in South Africa. We identified a Shuni/Shamonda virus reassortant in a horse and Shuni/Caimito, Shamonda/Caimito, and Shamonda/Sango virus reassortants in *Culicoides* midges. Continued genomic surveillance will be needed to detect orthobunyavirus infections in Africa.

The Orthobunyavirus genus, in the family Peribunyaviridae, has 18 serogroups; several of those represent emerging or reemerging viruses of importance to human and animal health transmitted by mosquitoes and *Culicoides* biting midges (1). Orthobunyavirus genomes have 3 negative single-stranded RNA segments, small (S), medium (M), and large (L) (2). Those segments can reassort in nature (3,4), leading to genetically diverse viruses with altered host ranges and pathogenicity (3). We characterized the genomes of orthobunyaviruses detected in animals with neurologic signs and birth defects and in *Culicoides* biting midges isolated in South Africa.

## The Study

During January 2020–February 2023, samples from 280 equids, 35 ruminants, 53 wildlife species, 6 avians species, and 7 cats or dogs displaying unexplained febrile and neurologic signs or sudden unexplained death were submitted from across South Africa to the zoonotic arbovirus surveillance program at the University of Pretoria (Pretoria, South Africa). We extracted RNA from blood, serum, plasma, cerebrospinal

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fluid, or tissue samples in a Biosafety Level 3 laboratory by using the QIAamp Viral RNA Mini Kit or RNeasy Mini Kit (QIAGEN, https://www.qiagen. com). We screened animal samples by using in-house S segment-specific TaqMan real-time reverse transcription PCR (RT-PCR) for *Orthobunyavirus* Simbo serogroup, as previously described (5). We characterized the M and L segments of positive samples by using conventional nested PCR. Primers for the M segment specifically targeted orthobunyaviruses (6), whereas L segment primers targeted the Peribunyaviridae family (7) (Appendix 1 Table 1, https://wwwnc. cdc.gov/EID/article/31/7/24-1800-App1.pdf).

A total of 6/381 (1.6%) animals tested positive for orthobunyaviruses by PCR of the S segment (Table; Appendix 1 Table 2) (5,8). Sanger sequencing identified 5/6 (83.3%) viruses as Shuni virus (SHUV); 4/5 (80.0%) infections were fatal. We found SHUV in samples from 2 aborted ovine fetuses from Mpumalanga and Northwest Provinces and 3 horses from Gauteng and the Northern Cape Provinces. An orthobunyavirus detected in an aborted goat fetus from the Western Cape Province was identified as Shamonda virus (SHAV) through Sanger sequencing, as previously described (6). We attempted next-generation sequencing but were unable to obtain full genomes.

We determined pairwise distances between orthobunyavirus sequences and constructed maximum-likelihood phylogenetic trees by using MEGA X version 10.2.6 (https://www.megasoftware.net) (Appendix 2 Tables 1–4, https://wwwnc.cdc.gov/ EID/article/31/7/24-1800-App2.xlsx). We performed conventional RT-PCR to obtain a larger S segment fragment (291 bp) and successfully amplified 4 (66.7%) of 6 sequences (Figure 1). The ZRU099/22 (ovine fetus) S segment grouped with SHUV strains from Israel (97.0%–98.0% identity) (Figure 1). Two

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	Orthony avirus DT DCD	Conventional nested BCB		
	Onnobunyavirus RT-PCR	Conventional nested PCR		
Sample no.	S segment	S segment	M segment	L segment
Mammals				
ZRU053/20	Shuni virus	IS	IS	IS
ZRU012/21	Shuni virus	Shuni virus	Shamonda virus	Shamonda virus
ZRU027/21	Shuni virus	AF	AF	AF
ZRU093/21	Shamonda virus	Shamonda virus	Shamonda virus	Shamonda virus
ZRU019/22	Shuni virus	Shuni virus	AF	AF
ZRU099/22	Shuni virus	Shuni virus	AF	Shuni virus
Culicoides biting midges				
MAR538 16	Schmallenberg virus	Schmallenberg virus	Schmallenberg virus	Schmallenberg virus
MAR085 13	Sabo virus	AF	Sabo virus	AF
MAR032 12	Sango virus	AF	Shamonda virus	AF
KYA229_16	Shuni virus	Shuni virus	AF	Shuni virus
GAU272 17	Shamonda virus	AF	AF	Caimito virus
GAU110 <sup>-</sup> 14	Shamonda virus	AF	AF	Caimito virus
MAR057_13	Shuni virus	AF	AF	Caimito virus
*AF, amplification failed; IS, insufficient sample; RT-PCR, reverse transcription PCR.				

Table. Orthobunyaviruses identified by PCR and sequencing of S, M, and L segments in samples from mammals and *Culicoides* biting midges, South Africa\*

horse samples (ZRU12/22 and ZRU19/22) grouped with SHUV strains from South Africa (97.0%–100.0% identity). The S segment of the aborted goat sample (ZRU93/21) clustered with SHAV (6). We amplified

M segments for 2 (33.3%) of 6 animal samples. Both M segments of ZRU093/21 (6) and ZRU012/21 (detected in horse brain) clustered with SHAV (Figure 2); the S segment of ZRU012/21 also clustered with







SHUV. The M segment of ZRU012/21 had 63.0%–65.0% identity with SHAV. We amplified L segments of 3/6 (50.0%) animal samples. Of those, 2 samples grouped with SHAV, including the aborted goat fetus (ZRU093/21) (6) and horse sample (ZRU012/21) (Figure 3), suggesting ZRU012/21 is a reassortment of the SHUV S segment and SHAV M and L segments; the L segment had 96.0%–99.0% identity with SHAV. The third sample (ZRU099/22) from the ovine fetus clustered with SHUV and had 96% identity with strains from Israel and 92% identity with strains from South Africa.

To determine if reassortants could be identified in *Culicoides* midges previously identified as orthobunyavirus positive, we screened pools of midges collected from 6 surveillance sites in Mpumalanga, Limpopo, and Gauteng Provinces during 2013–2017 (9). We performed M and L segment-specific RT-PCR on RNA from 7 orthobunyavirus S segment-positive *Culicoides* pools previously characterized by Sanger sequencing (Table; Figure 1). We successfully amplified the M segment from 3 pooled samples, all from Marakele National Park in Limpopo Province. MAR538\_16 grouped with the Schmallenberg virus (SBV) M segment (97.0%–98.0% identity) (Figure 2), as well as the SBV S segment (9) (Table). MAR085\_13,

previously positive for the Sabo virus (SABOV) S segment, also grouped with the SABOV M segment (95.0% identity). MAR032\_12 grouped with the SHAV M segment (97.0% identity) but also grouped with the Sango virus (SANV) S segment, suggesting reassortment. We amplified L segments for 5 pooled Culicoides midge samples: 2 from Marakele National Park in Limpopo Province, 2 from Boschkop in Pretoria Province, and 1 from Kyalami in Midrand, Gauteng Province. KYA299\_16 grouped with L segments of SHUV strains from South Africa (97.0% identity) (Figure 3) and SHUV S segments. MAR538\_16 grouped with SBV L segments (97.0%–98.0% identity) and both S and M segments. GAU272\_17 and GAU110\_14 grouped with the Caimito virus (CAIV) L segment (61.0% identity); however, both grouped with SHAV S segments, suggesting a possible reassortment. MAR057\_13 also grouped with the CAIV L segment (59.0% identity) (Figure 3) but grouped with SHUV S segments, suggesting another possible reassortment. We summarized the orthobunyavirus-positive cases tested in this study (Table). We deposited sequences in GenBank (accession numbers in Figures 1–3).

SHUV has been circulating in South Africa in both *Culicoides* midge and mosquito vectors and has been detected in wildlife, ruminants, horses, birds,

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and, more recently, in humans (5,8-12). SHAV was first detected in Culicoides biting midges in Gauteng Province of South Africa in 2017 and in an aborted goat fetus from the Western Cape in 2021 (6,9). SHUV was detected in 3 horses with neurologic signs, 2 from Gauteng and 2 from the Western Cape during 2020-2022. One of the 2 fatal cases was caused by a novel SHUV/SHAV reassortment. SHUV was also detected in 2 aborted ovine fetuses. Abortions associated with SHUV have been described in ruminants in Israel. We previously reported SHUV in cerebrospinal fluid from a 13-day-old, hospitalized infant with hydrops fetalis in Gauteng Province (12), suggesting SHUV might be a risk factor for birth defects in humans. SHUV, SHAV, and SBV, as well as reassortants, were detected in animals and *Culicoides* midges in this study. Therefore, orthobunyavirus infections should be considered a possible causes of birth defects in humans and animals in Africa.

Reassortments might alter pathogenesis, which has been observed for Ngari virus (13,14). Reassortants were detected in *Culicoides* midges collected in Gauteng and Limpopo Provinces; 2 were SHAV/ CAIV, 1 SHUV/CAIV, and 1 SANV/SHAV reassortants. SANV was previously detected in a springbok that displayed neurologic signs in Mpumalanga Province in South Africa (5). Little is known about CAIV, which belongs to the genus *Pacuvirus* isolated in Brazil from rodents and sandflies (15). The M segment of orthobunyaviruses detected in South Africa had only 61% identity to CAIV viruses from South America, suggesting different virus species within that genus.

The first limitation of our study is that full genome sequencing was not successful for mammal and *Culicoides* samples, likely because of low RNA levels. We were also unable to reamplify all samples, suggesting RNA degradation in older samples.

## Conclusions

Co-circulation of several orthobunyaviruses in Africa might give rise to novel reassortants with altered host range, tissue tropism, and pathogenesis. Reassortants might be missed without characterization of all 3 virus segments. A SHUV/SHAV reassortant described in this study was associated with severe neurologic infection and death in a horse, and SHUV and SHAV were both associated with abortions in ruminants. Several orthobunyaviruses were detected in Culicoides midges, including variants clustering with SBV and reassortants involving SHUV, SHAV, SANV, and a CAIV-like virus. Continued One Health genomic surveillance will be needed to detect those and other orthobunyaviruses to determine risks for infections in animals and humans in Africa and elsewhere.

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The study was approved by the University of Pretoria animal ethics committee (155/19), and Section 20 approval was obtained from the Department of Agriculture of South Africa (reference no. 12/11/1/1 [1552SS]).

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