Senecavirus A (SVA) in finishing swine: diagnosis and viral isolation

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ABSTRACT: Senecavirus A (SVA) has been a problem in Brazil since the end of 2014. The infections caused by SVA have disrupted the productive chain in Brazil, as it can be confused with foot-and-mouth disease. Although, the virus has remained endemic in the country, an increase in the number of cases of the disease was observed in 2018. The aim of the present study was to conduct the differential diagnosis of foot-and-mouth disease in an outbreak of vesicular disease in finishing swine. Animals (160–170 days old) were kept on a farm with 6,000 pigs in Minas Gerais State, Brazil. The morbidity and mortality rates were 20% and 2.2%, respectively. The diagnosis was performed by RT-PCR, using primers that determine the amplification of an internal region of the 3D gene. Furthermore, samples were inoculated into BHK-21 cell culture for viral isolation. In the first passage under cultivation, a cytopathogenic effect compatible with SVA replication (rounding and detachment of the cell monolayer) was observed. The viral identity was confirmed using two additional assays: indirect immunofluorescence assay (IFA) and nucleotide sequencing. Both tests confirmed that the infection was caused by SVA. In summary, we described a method for the diagnosis and viral isolation of SVA, a virus that arrived in Brazil in 2014 and has become endemic in the country.

Key words: emerging diseases, idiopathic vesicular disease, picornavirus, Seneca Valley virus, vesicular disease.

Senecavirus A (SVA) em suínos de terminação: diagnóstico e isolamento viral

RESUMO: Senecavirus A (SVA) é um problema no Brasil desde o final de 2014. As infecções causadas pelo SVA têm causado problemas para a cadeia produtiva no Brasil, pois podem ser confundidas com febre aftosa. Embora o vírus permaneça endêmico no país, foi observado um aumento no número de casos em 2018. O objetivo do presente estudo foi realizar o diagnóstico diferencial de febre aftosa em um surto de doença vesicular em suínos de terminação. Animais (160 a 170 dias de idade) eram mantidos em uma granja com 6.000 suínos no estado de Minas Gerais, Brasil. As taxas de morbidade e mortalidade foram de 20% e 2.2%, respectivamente. O diagnóstico foi realizado por RT-PCR, utilizando primers que determinam a amplificação de um região interna do gene 3D. Além disso, as amostras foram inoculadas na cultura de células BHK-21 para isolamento viral. Na primeira passagem em cultivo, foi observado efeito citopatogênico compatível com SVA (arredondamento e descolamento da monocamada celular). A identidade viral foi confirmada usando duas técnicas adicionais: ensaio de imunofluorescência indireta (IFA) e sequenciamento de nucleotídeos. Ambos os testes confirmaram que a infecção foi causada pelo SVA. Em resumo, descrevemos um método para o diagnóstico e isolamento viral do SVA, um vírus que chegou ao Brasil em 2014 e se tornou endêmico no país.

Palavras-chave: doenças emergentes, doença vesicular, picornavírus, Seneca Valley virus, vesicular disease.
The SVA has created problems for the swine production chain in Brazil, since it is a vesicular disease that can be confused with foot-and-mouth disease (LEME et al., 2019). However, there is little research concerning this virus in Brazil. Thus, the objective of the present study was to conduct the differential diagnosis of foot-and-mouth disease in an outbreak of vesicular disease in finishing pigs.

In May 2019, finishing swine (from 160 to 170 days old) kept on a farm with approximately 6000 pigs in the state of Minas Gerais, Brazil, showed clinical signs compatible with SVA infection. Clinical findings included vesicles, ulcers, and lameness (Figure 1). During the outbreak, the morbidity and mortality rates were 20% and 2.2%, respectively. After the suspicion of foot-and-mouth disease was ruled out by the official veterinary service, vesicle swabs from five animals were placed into 15 mL conical centrifuge tubes, frozen, and sent to the laboratory for investigation into the suspected SVA.

For the laboratory diagnosis, 2 mL of phosphate-buffered saline (PBS) was added to each tube containing a swab. Using a vortex, the tubes were shaken for 20 to 30 s, with 100 μL of the supernatant being used for RNA purification using TRIzol reagent® (Thermo Fisher Scientific™, USA), following the manufacturer’s instructions. Then, the complementary DNA (cDNA) was synthesized using kit SuperScript III reverse transcriptase® (Thermo Fisher Scientific™, USA), according to the manufacturer’s instructions. For the PCR, using software Clone Manager Basic - Sci-Ed, forward and reverse initiators were determined (SVA-3D-F 5’ <TGAACCTGCTGTGTTGTC> 3’ and SVA-3D-R 5’ <CAGAGTAGTCACCGTCTAAG> 3’, respectively) that delimit the amplification of an internal region of the 3D gene of the SVA genome (product size 359 bp). The conditions used in the PCR were as follows: 95 °C for 5 min for denaturation, followed by 35 cycles of 94 °C for 30 s for denaturation, 48 °C for 30 s for primer annealing, 72 °C for 30 s for extension, and a final extension at 72 °C for 10 min. The PCR products were resolved by electrophoresis (80 V, 40 min) on agarose gel 1%, and the DNA was intercalated with Unisafe® (Unisafe Dye, Uniscience do Brasil, Brazil) and visualized in a gel imaging system under UV light (ChemiDoc MP, BioRad, USA).

After adding PBS, the tubes containing the swabs were centrifuged for 10 min at 5000 RPM and 4 °C, and the supernatant was used for inoculation in

![Figure 1 – Lesions suggestive of Senecavirus A (SVA) infection. Finishing swine showing vesicles, ulcers (evidenced by the white arrows), and lameness.](image-url)
cell culture. Baby hamster kidney cells (BHK-21 [C-13 ATCC® CCL-10™]) were cultured in 6-well plates, prepared 24 h in advance, and inoculated with 500 μL of the swab supernatant from the lesions. After 1 h of adsorption, the inoculum was replaced with a culture medium (MEM) supplemented with 5% fetal bovine serum (SFB), L-glutamine (2 mM), penicillin (100 U.mL⁻¹), and streptomycin (100 μg.mL⁻¹). The cultures were kept in an incubator at 37 °C in a 5% CO₂ atmosphere for 48 h.

Cell cultures were subjected to the indirect immunofluorescence assay. Quickly, inoculated and uninoculated cells (negative control) grown on multi-spot slides were fixed and permeabilized with acetone PA for 5 min. An anti-SVA monoclonal antibody (mAb) (dilution 1:1000) (kindly provided by Dr. D. Diel) was incubated for 1 h at 37 °C. Then, an anti-mouse-IgG-FITC conjugated antibody (dilution 1:200) (Sigma-Aldrich, USA) was incubated in similar conditions. The reading was performed using an epifluorescence microscope (Opticam®).

For nucleotide sequencing, the RT-PCR product was purified using a Purification Kit (PureLink™ Quick Gel Extraction and PCR Purification Combo Kit Purification, Invitrogen™, USA), following the manufacturer’s instructions. The positive samples in the cell culture isolation were sequenced in quadruplicate in an automatic sequencer (ABI-PRISM 3100 Genetic Analyzer, Applied Biosystems, Foster City, CA) using the BigDye reagent.

The sequences were analyzed using the Staden Package program to obtain the consensus sequence, which was subsequently submitted to the BLAST sequence analysis tool for comparison with sequences available on GenBank, using the NCBI (National Center for Biotechnology Information) database. Sequence alignment was performed with ClustalW, an internal feature of the BioEdit software. The obtained sequence was aligned with 24 sequences available on GenBank using the BioEdit Sequence Alignment Editor Software, version 7.0.5.3 (http://www.mbio.ncsu.edu/bioedit/bioedit.html). The phylogenetic tree was built using the neighbor-joining method, with 1000 self-initialization replicates, based on the kimura-2 parameter of nucleotide substitution in MEGA6 (TAMURA et al., 2013).

The suspicion of SVA was confirmed by RT-PCR after the amplification of a product with the predicted molecular weight (359 bp) (Figure 2). The five samples analyzed using RT-PCR tested positive. However, in cell culture

![Figure 2 – Agarose gel after reverse transcription followed by polymerase chain reaction (RT-PCR) for Senecavirus A (SVA). Amplification of an internal region of the gene encoding the 3D protein (359 bp).](image-url)
isolation (CCI), only one of the five samples tested positive. Probably because some of these samples have lost viability (lost infectivity), but the genome has been preserved, making it possible positive result in RT-PCR. After the first pass in cell culture, a cytopathogenic effect compatible with SVA replication was observed (rounding and detachment of the cellular layer) approximately 24 h post-infection (Figure 3). The identity of the isolated virus was confirmed using an indirect immunofluorescence assay (IFA) (Figure 4). The use of an mAb against SVA ensured the specificity of the diagnostic tool. In addition, genetic sequencing and phylogenetic analysis grouped the sample obtained in the present study with Brazilian SVA sequences available on GenBank (Figure 5). Thus, both tests confirmed the identity of the virus causing the vesicular disease during the investigated outbreak. In addition, the isolation of an SVA sample makes it possible to carry out studies in vitro and in vivo viral characterization. Clinically identifying SVA infections remains a challenge. In the first cases in Brazil, piglets were the main category affected and, in addition to the vesicular disease, the animals presented diarrhea, neurological signs (motor incoordination, tremors, and convulsion), and mortality (LEME et al., 2015). In finishing swine, vesicular disease is the main manifestation of the infection and the animals usually recover within two weeks (JOSHI et al., 2016a). From the first detection in 2014 until 2019, SVA has remained endemic in the main swine-producing regions in Brazil. Although, the frequency of the disease has fluctuated and remained low, in late 2018 and early 2019 there was an intense increase in the occurrence of outbreaks (LEME et al., 2019). In addition, the attention of veterinarians and researchers has been drawn, as the outbreaks of SVA started to occur mainly in finishing pigs, which is different from what was described in the first cases in Brazil.

In Brazil, since the first viral detection in 2014, there has been much speculation concerning how the virus has been maintained and circulated among farms. Recently, research has shown the ability of SVA to remain in the infected animals, with carriers of the virus able to spread the agent among herds via viral excretion up to 60 days post-infection (MAGGIOLI et al., 2019). This hypothesis may explain the maintenance of the virus and increase in cases observed in Brazil between late 2018 and early 2019. In addition, a study has detected SVA in environmental samples, as well as the feces and

Figure 3 - Viral isolation of Senecavirus A (SVA) in baby hamster kidney cells (BHK-21). Inoculated = cells inoculated with a swab sample collected from swine with vesicular lesions. Control = cell culture maintained with minimal essential medium (MEM). Images obtained at 0, 24 and 48 hours post-infection (h p.i.) under an optical microscope (100x magnification). Inoculated cells feature cytopathogenic effect compatible with SVA replication (rounding and detachment of the cellular layer).

Figure 4 - Indirect immunofluorescence assay (IFA) using anti-SVA monoclonal antibody and FITC-conjugated mouse anti-IgG antibody after viral isolation of Senecavirus A (SVA) in baby hamster kidney cells (BHK-21). Inoculated = cells inoculated with a swab sample collected from swine with vesicular lesions. Control = cell culture maintained with minimal essential medium (MEM). Images obtained at 24 hours post-infection under UV light microscope.
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In summary, SVA has caused problems for the production chain in Brazil. We performed the differential diagnosis of foot-and-mouth disease in an outbreak of vesicular disease in finishing swine. The SVA diagnosis was confirmed and a contemporary strain from finishing swine was isolated.

Animal sample used in this study were collected as part of routine diagnostics. This article does not contain studies with human or animal participants carried out by any of the authors.

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AUTHORS’ CONTRIBUTIONS

MM, VBF, and MM performed the analyses. All authors critically reviewed the manuscript and approved the final version.

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