In 2013, a 6-week-old female piglet—Sus scrofa—was presented at the University of Veterinary Medicine Hannover in Germany. The piglet was euthanized after necropsy revealed interstitial pneumonia; a mild, multifocal, lymphohistiocytic panencephalitis that affected the cerebrum and cerebellum, including brain stem and medulla oblongata; and a mild, multifocal, lymphohistiocytic pannmyelitis. Results from screening for typical neurotropic viruses (classical swine fever virus, suid herpesvirus 1, rabies virus, teschovirus, porcine enterovirus 8, 9, and 10) were negative; Mycoplasma hyorhinis was detected by multiplex PCR (Institute of Virology, University of Veterinary Medicine Hannover) within the lung and pulmonary lymph nodes. Cerebral tissue from the pig was processed for viral metagenomics by random RNA and DNA virus screening and next-generation sequencing (NGS) with the 454 sequencing platform (GS Junior; Roche, Basel, Switzerland), as described (1), and 21,359 reads were obtained. Analysis by using blastn and blastx (2) showed 10 reads had ≥97% nt identity with porcine bocavirus (PBoV) KU14. No other viral sequences were detected.

By using primers based on sequence data of the PBoV, partially overlapping PCR amplicons were obtained to confirm and extend the NGS data of the isolate, which was named PBoV S1142/13 (1; GenBank accession no. KU311698). A total of 2,176 nt of PBoV S1142/13 were obtained, consisting of the partial nucleoprotein (NP) 1 and the nearly complete viral protein (VP) 1 gene. By using MAFFT version 7 (http://mafft.cbrc.jp/alignment/server/), we aligned the nearly complete VP1 gene of PBoV S1142/13 with various closely related members of the genus Bocaparvovirus and built a maximum-likelihood tree by using the general time reversible plus invariable sites plus gamma distribution method, as determined by jModelTest 2.0 (3) and default parameters in MEGA6.06 (4). Results confirmed that PBoV S1142/13 was most closely related to PBoV KU14 (Figure, panel A). The partial genome of PBoV S1142/13 differed at 8 nt positions from PBoV KU14, resulting in 99.6% nt identity. Of these nucleotide differences, 4 resulted in an amino acid difference, including position 2733 (T→C on the basis of PBoV KU14 as a reference genome), which is part of the NP1 stopcodon of PBoV KU14. These results indicate that the stopcodon was located 39 nt farther downstream than for PBoV KU14. The other 3 aa differences were present in the VP1 protein; each of these differences was within the same group of amino acids as those detected in PBoV KU14.

For further substantiation of a potential cause-effect relationship of histologic (Figure, panel B) and NGS results, we performed fluorescent in situ hybridization (FISH) on formalin-fixed, paraffin-embedded central nervous system (CNS) sections of the diseased animal and of a control pig with no CNS lesions. We used an RNA probe specific for the obtained NP1 and VP1 sequences covering 1,153 nt (Accession no. KU311698) and built a maximum-likelihood tree by using the general time reversible plus invariable sites plus gamma distribution method, as determined by jModelTest 2.0 (3) and default parameters in MEGA6.06 (4). Results confirmed that PBoV S1142/13 was most closely related to PBoV KU14 (Figure, panel A). The partial genome of PBoV S1142/13 differed at 8 nt positions from PBoV KU14, resulting in 99.6% nt identity. Of these nucleotide differences, 4 resulted in an amino acid difference, including position 2733 (T→C on the basis of PBoV KU14 as a reference genome), which is part of the NP1 stopcodon of PBoV KU14. These results indicate that the stopcodon was located 39 nt farther downstream than for PBoV KU14. The other 3 aa differences were present in the VP1 protein; each of these differences was within the same group of amino acids as those detected in PBoV KU14.

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served as a positive control. The spinal cord of the diseased pig showed diffuse intracytoplasmic and intranuclear PBoV-specific signals within scattered neurons adjacent to the histologically detected inflammatory lesions (Figure, panel C). The negative control and the nonprobe incubation lacked PBoV-specific signals. The porcine ubiquitin probe provided a strong intracellular and extracellular staining within the CNS of both pigs.

PBoV (genus Bocaparvovirus, family Parvoviridae) was first described in 2009 as porcine boca-like virus in pigs in Sweden with postweaning multisystemic wasting syndrome (5). PBoV is usually involved in respiratory and intestinal diseases in pigs (5) but has not been detected in the CNS. In the pig in our study, the lack of detection of other viral sequences by using NGS indicates the potential role of PBoV as a pathogen that triggers encephalomylitis. FISH substantiated the NGS results and revealed neuronal intracytoplasmic and intranuclear PBoV-specific signals adjacent to the lesion, indicating intraneuronal transcription and replication (6). Nevertheless, a potential synergistic effect of M. hyorhinis on the PBoV pathogenesis cannot be ruled out. Similarly, co-infection of M. hyorhinis and porcine circovirus type 2 has been associated with enhanced inflammatory lesions in the lungs of pigs (7).

The CNS tropism of PBoV S1142/13 could result from various factors, including specific amino acid changes that enable the virus to pass the blood–brain barrier and infect neurons. Additional studies are necessary to elucidate a possible role of the amino acid differences between PBoV S1142/13 and PBoV KU14 in the tropism of these viruses.

Human bocavirus has recently been found in the cerebrospinal fluid of patients having encephalitis (8), and related human parvovirus 4 (9) and human parvovirus B19 (10) have been reported in human encephalitis. The correlation of PBoV-specific signals by using FISH for histologic detection of encephalomyelitis assigns PBoV a potential role in provoking CNS lesions. PBoV should be considered as a cause of encephalomylitis but needs further investigation.

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Pegivirus Infection in Domestic Pigs, Germany

Christine Baechlein,¹ Adam Grundhoff,¹ Nicole Fischer, Malik Alawi, Doris Hoeltig, Karl-Heinz Waldmann, Paul Becher

Author affiliations: University of Veterinary Medicine Hannover, Hannover, Germany (C. Baechlein, D. Hoeltig, K.-H. Waldmann, P. Becher); German Center for Infection Research Partner Site Hannover–Braunschweig, Hannover (C. Baechlein, P. Becher); German Center for Infection Research Partner Site Hamburg–Lübeck–Borstel, Hamburg (N. Fischer, A. Grundhoff); Heinrich Pette Institute, Hamburg, Germany (A. Grundhoff, M. Alawi); University Medical Center Hamburg–Eppendorf, Hamburg (N. Fischer, M. Alawi)

Address for correspondence: Wolfgang Baumgärtner, Department of Pathology, University of Veterinary Medicine, Bünteweg 17 D-30559 Hannover, Germany; email: wolfgang.baumgaertner@tiho-hannover.de

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¹These authors contributed equally to this article.

To the Editor: The family Flaviviridae includes many human and animal virus pathogens. Recently, in addition to the genera Flavivirus, Hepacivirus, and Pestivirus, a fourth genus, Pegivirus, has been identified (1). In addition to human pegiviruses, a range of phylogenetic, highly divergent pegiviral sequences have been identified in various animal species, including primates, bats, rodents, and horses (2). We report the detection of a porcine pegivirus (PPgV) in serum samples from pigs.

Initially, we investigated pooled serum samples by using high-throughput sequencing methods and isolated RNA from individual porcine serum samples by using the QiAmp Viral RNA Mini Kit (QIAGEN, Hilden, Germany). We prepared libraries compatible with Illumina (San Diego, CA, USA) sequencing from pooled samples and individual serum samples by using the ScriptSeq version 2 RNA-Seq Library Preparation Kit (Epicenter, Madison, WI, USA) and sequenced them by using a HiSeq 2500 (2 × 150 cycles paired-end; Illumina) for pooled samples and MiSeq (2 × 250 cycles paired-end; Illumina) for individual samples (3).

We conducted quantitative reverse transcription PCR (RT-PCR) by using a Quantitect-SYBR Green Assay (QIAGEN) and primers PPgV_fwd: 5′-CTGTCTCATGCTGGTCAC-3′ and PPgV_rev: 5′-GCCATAGAACGGGAAGTC-3′. By using high-throughput sequencing of the pooled serum sample library (23,167,090 reads), we identified 1 contig (4,582 bp) that had distant nucleotide sequence similarity to bat pegivirus (69% and 4% sequence coverage) and 2 contigs (2,683 bp and 665 bp) that had 73% sequence coverage, thereby covering 8% and 37% of the identified sequence. RT-PCR with primers designed on basis of recovered sequences identified the sample containing pegivirus sequences. Subsequent MiSeq analysis (7,085,595 reads) of an RNA library prepared from a sample from 1 animal identified 1 contig (9,145 nt) with sequence similarity to pegivirus sequences.

We performed 3′ end completion of the viral genome by rapid amplification of cDNA ends and identified the entire open reading frame of PPgV_903 encoding 2,972 aa (GenBank accession no. KU351669). Analysis of the pegivirus 5′ untranslated region identified a highly structured internal ribosome entry site motif (online Technical Appendix, http://wwwnc.cdc.gov/EID/article/22/7/16-0024-Techapp1.pdf), which was similar in structure to previously described 5′ untranslated region structures of other pegiviruses (4,5).

Pegiviruses do not encode a protein homologous to the capsid protein of other viruses of the family Flaviviridae, another common feature of pegiviruses (6). The presence of cleavage sites for cellular signal peptidases and viral proteases indicates that, similar to polyproteins of other pegiviruses and members of the genus Hepacivirus, the pegivirus polyprotein NH2-E1-E2-Px-NS2-NS3-NS4A-NS4B-NS5A-NS5B-COOH (E [envelope], NS [nonstructural],

References

Address for correspondence: Wolfgang Baumgärtner, Department of Pathology, University of Veterinary Medicine, Bünteweg 17 D-30559 Hannover, Germany; email: wolfgang.baumgaertner@tiho-hannover.de