A large, highly prolific swine farm in Hungary had a 2-year history of neurologic disease among newly weaned (25–35-day-old) pigs, with clinical signs of posterior paraplegia and a high mortality rate. Affected pigs that were necropsied had encephalomyelitis and neural necrosis. Porcine astrovirus type 3 was identified by reverse transcription PCR and in situ hybridization in brain and spinal cord samples in 6 animals from this farm. Among tissues tested by quantitative RT-PCR, the highest viral loads were detected in brain stem and spinal cord. Similar porcine astrovirus type 3 was also detected in archived brain and spinal cord samples from another 2 geographically distant farms. Viral RNA was predominantly restricted to neurons, particularly in the brain stem, cerebellum (Purkinje cells), and cervical spinal cord. Astrovirus was generally undetectable in feces but present in respiratory samples, indicating a possible respiratory infection. Astrovirus could cause common, neuroinvasive epidemic disease.

Astroviruses are small, nonenveloped viruses with single-stranded 6.2–7.8 kb RNA genome of positive polarity (1,2). The family Astroviridae is currently divided into 2 genera: the genus Mamastrovirus of mammal-infecting viruses and the genus Avastrovirus of avian viruses (3,4). The genetically heterogenic astroviruses that are widespread among mammals and birds are generally associated with gastroenteritis, less commonly with respiratory disease, and rarely encephalitis or disseminated infections (2,5–19). Astrovirus infections with central nervous system (CNS) involvement were reported recently in mink, human, bovine, ovine, and swine hosts (the latter in certain cases of AII type congenital tremors) (5,6,12–14). Most neuroinvasive astroviruses belong to the Virginia/Human-Mink-Ovine (VA/HMO) phylogenetic clade and cluster with enteric astroviruses identified from asymptomatic or diarrheic humans and animals (15,16). Recent research shows that pigs harbor one of the highest astrovirus diversities among mammals examined (3,15,20). Porcine astroviruses (PoAstVs) were identified mainly from diarrheic fecal specimens, less commonly from respiratory specimens, although the etiologic role of astrovirus infection in gastroenteritis or in other diseases among swine is not settled (3,9,20–23). We report the detection of neuroinvasive porcine astrovirus type 3 (NiPoAstV-3) by reverse transcription PCR (RT-PCR) and in situ hybridization (ISH) in recent and archived CNS samples of newly weaned paraplegic pigs from 3 highly prolific swine farms in Hungary.

Materials and Methods

Sample Collection and Handling

During November 2015–July 2017, we collected multiple tissue samples from 5 paraplegic and 5 asymptomatic pigs at the index farm located in Hungary (GD; specific location redacted) (Table 1). We also tested nasal and anal swab pairs collected by using polyester-tipped swabs from another 5 paraplegic and 13 healthy animals. We washed tissue samples twice in 10 mmol/L phosphate buffered saline (PBS) to remove excess blood and held them at −80°C until total RNA extraction. For formalin-fixed, paraffin-embedded (FFPE) blocks, we fixed the dissected samples (Table 2) with buffered 8% formaldehyde, dehydrated and embedded into paraffin.

We also analyzed archived FFPE specimens from paraplegic pigs from earlier outbreaks of posterior paraplegia.
in Tázlár in 2011 and in Balmazújváros in 2014 (Table 2). The 3 swine farms are located in the central and eastern parts of Hungary, >100 km from each other, without known connection.

**Previous Laboratory Diagnostics**

CNS homogenates from the index farm tested negative by PCR for the following pathogens (families in parentheses): porcine reproductive and respiratory syndrome virus (Arteriviridae); porcine circovirus 2 (Circoviridae); hemagglutinating encephalitis virus (Coronaviridae); and porcine parvovirus 1, 2, 4, and porcine bocavirus (Paroviridae). Immunohistochemical detection of Toxoplasma gondii and West Nile virus and bacterial cultivation attempts from the CNS samples were also negative. Virus isolation attempts using brain homogenates of affected animals in swine kidney (PK-15) and Caucasian colon adenocarcinoma (Caco-2) cell lines were not successful (no cytopathic effects were visible). We detected no PoAsV type 3 (PoAstV-3) in the cell culture supernatants by nested RT-PCR with RNA-dependent RNA polymerase (RdRp) primer pairs.

**Total RNA Extraction and RT-PCR Screening**

Treatment of FFPE samples included the deparaffinization and rehydration steps, proteinase K digestion, and total RNA extraction. We used the same treatment protocols and the same reaction conditions and reagents used in the RT-PCR and nested RT-PCR reactions as described previously, with minor modifications (24–26) (online Technical Appendix, https://wwwnc.cdc.gov/EID/article/23/13/17-0804-Techapp1.pdf). For the RT-PCR screening of CNS samples for the presence of pestiviruses (Flaviviridae) and swine picornaviruses (Picornaviridae), including teschovirus, enterovirus, sapelovirus, Seneca Valley virus, pasivirus, kobivirus and encephalomyocarditis, we used virus-specific primer pairs as well as the outer and inner primer pairs targeting the RdRp or the capsid regions of PoAstV-3 (Figure 1; online Technical Appendix Tables 1, 2).

**Absolute Quantification Using Quantitative RT-PCR**

For the absolute quantification of viral RNA present in different tissue, urine, and fecal samples, we used the SYBR Green–based quantitative RT-PCR (RT-qPCR) method (Maxima SYBR Green qPCR Master Mix; Thermo Scientific, Waltham, MA, USA). For the generation of standard curve, we used 10-fold dilution series of purified and spectrophotometrically quantified RNA transcripts in the reactions. The RT-qPCR assays contained 3 technical repeats of all samples and standards. The slope of the standard curve was –3.4228 and the calculated PCR efficiency was 99.96%. The detailed protocol is provided in the online Technical Appendix.
Table 2. Results of PoAstV-3 detection, histology, and ISH analyses using formalin-fixed, paraffin-embedded blocks of samples from 3 symptomatic newly weaned pigs from a farm in Hungary and samples from 2 other farms with symptomatic pigs

| Farm ID | Collection year | Animal ID | FFPE block ID | Nested RT-PCR† | Tissue samples | ISH‡ |
|---------|-----------------|-----------|---------------|-----------------|----------------|-----|
| GD      | 2016            | GD-1      | GD-1A         | – (+)           | Spinal cord    | +   |
|         |                 |           |               | – (+)           | Brainstem      | +   |
|         |                 |           |               |                 | Cerebellum     | +   |
|         |                 |           |               |                 | Medulla oblongata | - |
| GD-2    | 2015            | GD-11§    | GD-11A        | – (+)           | Spinal cord    | +   |
|         |                 |           |               | – (+)           | Brainstem      | +   |
| Tázlár  | 2011            | TAZ-1     | TAZ-1A        | – (+)           | Hippocampus    | –   |
|         |                 |           |               | – (+)           | Brainstem      | +   |
|         |                 |           |               |                 | Spinal cord    | +   |
| Balmazújváros | 2014  | BAM-1     | BAM-1A        | – (+)           | Spinal cord    | –   |
|         |                 |           |               | – (+)           | Brainstem      | +   |
|         |                 |           |               |                 | Cerebellum     | +   |

*The screening RT-PCR primers are designed to either the RdRp or the capsid region of PoAstV-3; GD, index farm; ID, identification; ISH, in situ hybridization; PoAstV-3, porcine astrovirus type 3; RdRp: RNA-dependent RNA polymerase; RT-PCR, reverse transcription PCR; +, positive; −, negative.
†Symbols indicate the results of the first screening PCRs; symbols in parentheses indicate the results of the second (nested) RT-PCR. The results of nested RT-PCR refer to a mixture of tissues embedded into the total of 7 paraffin blocks.
‡Indicates results for neuroinvasive PoAstV-3.
§FFPE samples were the only specimens taken from this animal.

Long-range Amplification, 5′/3′ RACE-PCR, and Sanger Sequencing

For the complete genome (or complete 3′ open reading frame [ORF] 1b–ORF2–3′ untranslated region [UTR]) acquisitions of the PoAstVs, we used different long-range and 5′/3′ rapid amplification of cDNA ends RT-PCRs according to previously described protocols (26,27). We designed the sequence-specific primers used for the amplification of overlapping genome fragments based on the genome of PoAstV-3 strain US-MO123 (GenBank accession no. JX556691) and closely related sequences downloaded from the GenBank database (online Technical Appendix Table 3). We sequenced PCR products directly with the BigDye Terminator v1.1 Cycle Sequencing Ready Reaction Kit (Applied Biosystems, Stafford, TX, USA) using the primer-walking method with an automated sequencer (ABI Prism 310 Genetic Analyzer; Applied Biosystems). We have submitted the nucleotide sequences of study astrovirus strains to GenBank under accession nos. KY073229–32.

Sequence and Phylogenetic Analyses

We aligned astrovirus sequences by using the MUSCLE web tool of EMBL-EBI (28) and performed pairwise nucleotide and amino acid identity calculations of the aligned sequences with GeneDoc version 2.7 (http://iubio.bio.indiana.edu/soft/molbio/ibmpc/genedoc-readme.html). We constructed phylogenetic trees of deduced amino acid sequence alignments by using MEGA version 6.06 software (29) and the neighbor-joining method with the Jones–Taylor–Thornton matrix-based model. Bootstrap values were set to 1,000 replicates, and only likelihood percentages of ≥50% were indicated.

Histology and In Situ Hybridization

We performed chromogenic (with 3,3′-diaminobenzidine/DAB) in situ hybridization in FFPE slides (RNAscope 2.0, Brown Kit; Advanced Cell Diagnostics, Newark, CA, USA) according to the manufacturer’s instructions for viral RNA detection of Ni-PoAstV-3. We used 30 probe pairs generated at Advanced Cell Diagnostics designed to hybridize native viral Ni-PoAstV-3 RNA. Negative controls included Dap-B (dihydropicolinate reductase gene from Escherichia coli) probe; an unrelated viral probe; and normal porcine brain region-matched sections.

Results

Clinical Observations

There are ~2,000 sows and their offspring in the investigated highly prolific index farm (GD). Episodes of neurological disease of unknown etiology have persisted in the past 2 years. The syndrome affects an average of 30–40 weaned pigs monthly (1.5%–2% of total), although the number of monthly cases infrequently rose to ~80 pigs (4%) in the autumn–winter seasons. The clinical signs of posterior leg weakness or paraplegia and pitching (stage 1); later paralysis of both legs and skin pain (stage 2); or loss of consciousness, paresis, and serious flaccid paralysis of muscles (stage 3) typically appear among weaned pigs 25–35 days old, 1 week after the weaning procedure (Video). We did not observe gastroenteric symptoms. All of the affected pigs in stage 3 of the disease were unable to eat or drink; they died due to exsiccosis (dehydration) or were euthanized. Signs persisted typically for 1 week before death or euthanasia. Postmortem examination
results showed no signs of mechanical damage (fractures, abscesses, or hemivertebrae). Pigs are vaccinated against porcine circovirus 2, *Mycoplasma hyopneumoniae*, and *Actinobacillus pleuropneumoniae*. Preventive amoxicillin treatment of the piglets was done routinely at weaning. Due to the preventive measures in effect as of spring 2017, which included extensive decontamination of the piggeries and the physical separation of the newly weaned pigs from different litters, the number of encephalomyelitis cases among weaned pigs decreased with only 1–2 cases/month observed on the index farm.

The 2 additionally examined swine farms located in Tázlár and Balmazújváros each held approximately 500 sows and their offspring. Similar symptoms of staggering and paralysis appeared among pigs 3–5 weeks old in outbreaks in 2011 (Tázlár) and 2014 (Balmazújváros).

**Detection and Analysis of Astroviruses from CNS Samples of Affected Animals**

In March 2016, we collected brain stem, spinal cord, nasal swab, and fecal samples from a newly weaned pig from index farm GD (GD-1, index animal) that showed signs of encephalomyelitis and posterior paraplegia (stage 1). The brain stem and spinal cord samples tested negative by RT-PCR for pestivirus (family *Flaviviridae*) and several swine-infecting picornaviruses (family *Picornaviridae*) (online Technical Appendix Table 1). On the basis of the increasing evidence of the pathogenic role of neurotropic astroviruses among humans and farm animals (5,6,14,17,30) we investigated the presence of astrovirus using panastrovirus PCR primers (online Technical Appendix Table 1) (31). We sequenced 2 samples from the index animal: the full-length genome of the neuroinvasive astrovirus strain NI-Brain/9-2016a/HUN (GenBank accession no. KY073229) from the brain stem sample and the complete capsid-encoding ORF2 from the spinal cord sample NI-SC/9–2016a/HUN (GenBank accession no. KY073230). The 6393-nt (without the poly[A] tail) complete genome showed the typical astrovirus genome organization with 3 putative ORFs, 2529 nt (ORF1a), 1527 nt (ORF1b), and 2265 nt (ORF2), flanked by short 5′ and 3′ UTRs (Figure 1). We identified the conserved proteolytic cleavage site (V\_561HQ\_TNT) of serine protease (ORF1a) and the conserved Y\_358GDD motif of the RdRp (ORF1b) (33). The nonstructural proteins of ORF1a (842 aa) and ORF1b (508 aa) and the capsid protein of ORF2 (754 aa) showed 93%, 95%, and 93% aa identity, respectively, to the corresponding genome parts of the closest known relative PoAstV-3 strain, US-MO123. All of the conserved genomic features of mamastroviruses were present in strain NI-Brain/9-2016a/HUN: the conserved C\_1CAAA pentamer at the 5′ end of the genome; the frame-shift heptamer motif (A\_251AAAAAC) followed by a stem–loop structure at the 3′ end of ORF1a; the conserved sgRNA

![Figure 1. Genome map of the neuroinvasive PoAstV-3 strain NI-Brain/9-2016a/HUN (GenBank accession no. KY073229) from a symptomatic newly weaned pig from a farm in Hungary together with the location of RT-PCR products used for different astrovirus screening reactions and quantitative RT-PCR analyses. The black arrow indicates the possible localization of a ribosomal frame-shift during the synthesis of ORF1ab peptide. The first and last nucleotide positions of the ORFs are marked with numbers at the top and bottom of each box. ORF, open reading frame; PanAstV, panastrovirus; PoAstV-3, porcine astrovirus type 3; RT-PCR, reverse transcription PCR; UTR, untranslated region.](https://www.cdc.gov/eid/content/23/12/1518_f1.jpg)
promoter sequence motif of U_{\text{508}}UGGAGgGGaGGAC-CaAA_{\text{345}}UUGG (variable nts are in lowercase, start codon of ORF2 is underlined) at the junction of ORF1b/ORF2; and the stem loop II–like motif (s2m) in the 3’ end of the genome between nt position=s 6322 and 6353. The 3’ UTR of Ni-Brain/9-2016a/HUN is 27 nt shorter and did not contain the short sequence repeat found at the 3’ end of strain US-MO123 (G_{\text{501}}AUUUCUUUNA). Based on the high sequence identity and the similar genomic features, the Ni-Brain/9-2016a/HUN strain most likely belongs to the US-MO123 (G_{\text{501}}AUUUCUUUNA). Most of the aa differences between the Ni-PoAstV-3 study strains and the other enteric PoAstV-3 strains are located in the N-terminal part of ORF1a and in the C-terminal part of ORF2 (Table 3). Phylogenetic analysis showed a close relationship between the identified Ni-PoAstV-3 sequences and the known PoAstV-3 strains located within the same larger clade containing most other mammastroviruses with known neurotropic potential (Figure 2).

Detection of Ni-PoAstV-3 in Non-CNS Samples

We detected Ni-PoAstV-3 in multiple non-CNS samples from the respiratory system, lymphoid system, circulatory system, and salivary glands of affected animals (Table 1). We detected virus only in the second PCR round in 1 ileum sample and in 2 of the 3 analyzed fecal samples using nested RT-PCR (Table 1). Samples from internal organs (spleen and kidney) and urine samples tested negative by nested RT-PCR (Table 1).

We determined the copy number of Ni-PoAstV-3 using SYBR Green–based –qPCR. All of the samples that showed nested RT-PCR positivity only in the second (nested) PCR round had negative test results by RT-qPCR, indicating low copy number (<100 copies/μg total RNA) of the virus in that tissue sample. The highest copy number was detected in the brain stem, followed by the spinal cord (Figure 3). Of note, we detected relatively high copy numbers in the tonsil and nasal mucosa samples (Figure 3). The serum of animal GD-3 contained 2.07 ×

### Table 3. Amino acid differences between neuroinvasive PoAstV-3 strains from 3 symptomatic newly weaned pigs from a farm in Hungary and reference enteric PoAstV-3 strains detected from fecal samples*

| Category | Genomic region | ORF1a | ORF1a | ORF1a | ORF1a | ORF1b | ORF1b | ORF2 | ORF2 | ORF2 | ORF2 | ORF2 |
|----------|----------------|-------|-------|-------|-------|-------|-------|------|------|------|------|------|
| PoAstV-3 type | 1–400 | 1–400 | 401–844 | 401–844 | 1–508 | 1–508 | 1–415 | 1–415 | 416–754 | 416–754 |
| Amino acid position | Amino acid changes | Ni | Ent | Ni | Ent | Ni | Ent | Ni | Ent | Ni | Ent |
| M25 | S/L | F408 | L | N54 | D | R29N | KT/I/A/V | L439 | H/P/V |
| Y41 | F | I434 | V | D106G | [A/E]D | S434 | R | S453 |
| R17 | K | S481 | P | A181 | S | R38 | Y | F457 |
| T120 | S/L | S576 | T/V | I206 | V | V55 | T | Y589 |
| T122 | S/L | G608 | N | R213 | K | T57R | SK | A570[P] |
| K151G | RC | N646 | H | Y293 | H/N | T81 | A | N572[Y] |
| L170 | M | E679 | D | E343 | D | D581 | N |
| L179 | M | K375 | R | I601 | V |
| M185 | L | I378 | T | S617 | N |
| D208 | E/N | N382 | D | T628 | S |
| D202S[P/Q] | NPTDG | i415 | A/T | S678 | T |
| P217A | TT | | | |
| T220[V/A] | IS | | | |
| P224 | H/R | | | |
| I299 | V | | | |
| E332 | D | | | |
| V338 | L/I | | | |
| L346 | F | | | |
| I349 | V | | | |

*We identified 3 PoAstV-3 isolates: Ni-Brain/9-2016a/HUN (GenBank accession no. KY073230); Ni-Brain/173-2016a/HUN (accession no. KY073231); and Ni-Brain/386-2015/HUN (accession no. KY073232). We compared these with enteric strains from GenBank (accession nos. JX556691, LC201595-7, and LC201599). AD, presumed particle assembly domain; Ent, enteric; Ni, neuroinvasive; ORF, open reading frame; PoAstV-3, porcine astrovirus type 3; RID, presumed receptor-interaction domain.
10^6 virus copies/mL and of animal GD-4. 1.64 × 10^3 virus copies/mL.

To validate the general presence of Ni-PoAstV-3 in the respiratory system and the absence of the virus in the feces during the acute phase of the illness, we collected additional nasal and anal swab pairs from 5 affected pigs and 13 clinically healthy pigs of the same age (±25–35 days) from the index farm. Four (80%) of the 5 nasal swab samples from affected animals tested positive but all of the anal swab samples tested negative using nested RT-PCR with primers...
targeting the RdRp region of Ni-PoAstV-3. The nasal and anal swab samples of the asymptomatic animals were all negative by nested RT-PCR. Because we collected varying amounts of samples by polyester-tipped swabs, we did not perform absolute quantification of Ni-PoAstV-3 by RT-qPCR.

Detection of Ni-PoAstV-3 in Archived FFPE Samples
All but 1 archived FFPE samples from Tázlár and Balmazújváros were positive by nested RT-PCR for Ni-PoAstV-3 using 2 sets of primer pairs targeting the RdRp and capsid genes of Ni-PoAstV-3 (Table 2; Figure 1; online Technical Appendix Table 2). The spinal cord FFPE sample from Balmazújváros had a negative result using both nested RT-PCR primer sets. The nested RT-PCR positive samples had positive results by ISH (data not shown).

Histology and ISH
Histologically, shared CNS lesions among the animals examined were moderate to marked lymphohistiocytic cell perivascular cuffing with marked vasculitis and neuronal degeneration, necrosis, and neurophagia with multifocal microgliosis and satellitosis (Figure 4). The neuronal necrosis was especially evident in the dorsal and ventral horns of the cervical spinal cord gray matter, although it was also detected in neurons of the Purkinje layer (cerebellum), the medulla oblongata, cerebellar peduncles, and midbrain (Figure 5). Necrotic neurons were variously swollen and hypereosinophilic or shrunken with tinctorial changes including faded, amphophilic, or eosinophilic cytoplasm (Figure 5). Nuclei of affected neurons are pyknotic, karyorrhectic, or losing border definition within the cytoplasm. We performed ISH on 5 affected animals (Table 2). Ni-PoAstV-3 hybridization was predominantly restricted to neurons, including those with visible necrosis and, in the cerebellum in particular, some that were histologically unaffected, although some regions of gliosis (presumed inflammation after neuronal necrosis) also contained viral RNA (Figure 5, panel M). Hybridization was distinct, with punctate to diffuse cytoplasmic staining throughout the cytoplasm. The unique microarchitecture of the Purkinje layer of the cerebellum offered the clear demonstration that viral nucleic acid was present within dendritic processes coursing through the molecular layer (Figure 5, panels G, J). We found no pathologic lesions in other samples from kidneys, liver, gastrointestinal tract, or immune system (data not shown). The samples from the immune system were also negative by Ni-PoAstV-3 ISH (Table 2).
Discussion

We detected astrovirus RNA in multiple tissues collected during 2015–2017 from newly weaned pigs with encephalomyelitis and posterior paraplegia of unknown origin, with the highest viral load detected in brain stem and spinal cord samples. We detected the same virus in archived brain and spinal cord FFPE samples from similarly affected animals from 2 additional swine herds collected in 2011 and 2014. These data indicate that a genetically similar, neurovirulent astrovirus is circulating in multiple swine farms since 2011 or earlier in Hungary.

According to the refined classification for the assessment of causation (34), the Ni-PoAstV-3 and the observed encephalitis and paraplegia are in a probable causal relationship (Level 2). Paraplegia associated with astrovirus neuroinfection is not unprecedented; minks had astrovirus-induced “shaking mink syndrome” and were reported paraplegic at the final stage of the disease (12,35).

Neurologic signs were observable mainly among newly weaned pigs (Video, https://wwwnc.cdc.gov/EID/article/23/12/17-0804-V1.htm). The time of weaning, which involves nutritional (from milk to solid feed), social (mixing with different litters without the sow), and environmental (moving to a new pen) changes, is known to be the most stressful period in a pig’s lifetime and is associated with dysfunction of the immune system (36). Furthermore, the inadequate quantity and quality of colostrum intake of sucking piglets, and therefore the presumably low level of specific maternal antibodies due to highly prolific sows with large litters in the index farm, might also contribute to the emergence of the clinical disease. Decreased immune status was frequently present with extraintestinal dissemination of astroviruses in humans and in mice (5,37–41).

Our sequence analyses indicate that the identified astrovirus strains belong to the PoAstV-3 genotype, which clusters within the VA/HMO phylogenetic clade (Figure 2), as do most mammalian strains with known neurotropic potential (6,14,19). However, other canonical human astroviruses outside of the VA/HMO clade could also be associated with CNS disease (41). At the molecular level, the most conspicuous difference between the genomes of neuroinvasive virus and the enteric PoAstV-3 strain U.S.-MO123 is the 27 nt deletions found in the 3′ UTR of the CNS-associated astroviruses. The possible impact of this 3′ UTR deletion on viral tropism is unknown, although neuroinvasive bovine astroviruses also possess 3′ UTR architecture that differs from the diarrhea-associated astroviruses (42).

At the amino acid level, one of the most divergent regions between the neuroinvasive and other PoAstV-3 strains was found at the receptor-interaction domain of ORF2 (Table 3), which contains potential receptor binding sites (43,44). This finding could indicate an altered receptor spectrum and therefore altered tissue tropism of neuroinvasive and enteric PoAstV-3 strains.

PoAstV-3 strains were previously detected only from fecal samples of healthy or diarrheic piglets worldwide (20,22,45). We found that Ni-PoAstV-3 was either undetectable or detected only at low viral loads in the
analyzed fecal samples, whereas the virus was generally detectable in the respiratory system of paraplegic pigs. This finding may indicate that CNS infection and replication occur later than enteric replication or that initial
replication (e.g., in the respiratory tract). Multiple types of astroviruses were recently identified from nasopharyngeal swabs or lung tissue samples from swine, bovines, and humans with respiratory symptoms including the neurotropic human VA1 strain from a patient with febrile acute respiratory disease (9–11,23), although neither the respiratory tropism nor the airborne transmission of astroviruses has been experimentally confirmed. Therefore, testing of only fecal samples from sick animals may result in underestimation of the incidence of astrovirus in pigs.

We measured the highest viral loads of Ni-PoAstV-3 in brain and spinal cord samples, similar to those found in diseased ovine and human patients with astrovirus-associated encephalitis (5,14). Ni-PoAstV-3 was also detectable in serum specimens and multiple organs of the respiratory, lymphoid, and cardiovascular systems of diseased swine. These results indicate that Ni-PoAstV-3 can result in viremia and disseminated infection involving the brain, spinal cord, and multiple organs during the acute phase of encephalomyelitis and posterior paraplegia. Astroviruses seem to play a role in a common and severe disease (encephalomyelitis and paralysis) in pigs.

The observable histopathologic changes, as well as the neuronal localizations of Ni-PoAstV-3 RNA in CNS samples of paraplegic pigs, are comparable to astrovirus-associated encephalitic cases of minks, humans, and cattle. Similar neuronal degeneration or necrosis with microgliosis in the brain or cerebellum, as well as inflammation of gray matter of the spinal cord, were previously described in cattle with astrovirus-associated nonsuppurative encephalitis (6,35,46,47), which suggests the general course of an astrovirus neuroinfection.

While some astroviruses are known to cause outbreaks of gastroenteritis, astrovirus-associated encephalitis cases have been reported only sporadically among humans, cattle, and sheep (6,14–16,47). The constant presence with recurrent increases of neurologic disease cases in swine farms indicates that natural neuroinvasive astrovirus infections may cause common, severe, persistent epidemics among domestic pigs and constitute an economically important agent threatening livestock and even humans, considering the possible zoonotic and recombinant potential of astroviruses (48).

Our results must be interpreted in the light of some potential limitations, which are currently true for other astrovirus-associated encephalitis studies: the absence of experimental evidence such as in vivo inoculation experiments, which could clarify the true causality between the astrovirus neuroinfection and the manifested CNS symptoms; and the roles of presumed respiratory replication and decreased immune state. Therefore, despite a growing body of scientific data regarding the presence of astroviruses in CNS in different animals, the direct association of astrovirus neuro-infection and encephalomyelitis should be treated with caution. Newly weaned pigs could potentially provide an in vivo animal model to study and clarify this association.

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Outbreaks of Neuroinvasive Astrovirus Associated with Encephalomyelitis, Weakness, and Paralysis among Weaned Pigs, Hungary

Technical Appendix

Total RNA Extraction and RT-PCR Screening

We conducted pretreatment, including deparaffination and rehydration steps and Proteinase K digestion of FFPE samples, as described previously. (1,2).

For total RNA extraction, we used 50–100 mg of tissue samples, 400µL of Proteinase K–digested FFPE samples, and 150µL of urine, serum specimens, nasal swab samples, and fecal suspensions (35%–40% v/v, diluted in 0.1M PBS) prepared with TRI Reagent (Molecular Research Center, Cincinnati, OH, USA) according to the manufacturer’s instructions. We ground the tissue samples manually to homogenize them using a Potter-Elvehjem pestle (Sigma-Aldrich Co., Munich, Germany).

We used the same reaction conditions and reagents in the RT and nested PCR reactions as were described previously, with minor modifications (3). Briefly, 1µg of total RNA was transcribed in a final volume of 25µL. We used the total amount of the RT product in the first PCR round in a final volume of 50µL. On RT-PCR negative samples, we performed a second PCR round with Ni-PoAstV-3 specific inner primer pairs for the specific and sensitive detection of small quantities of viral RNA. In the second (nested) PCR round, we used 1µL of the first PCR product as a template. The thermal program of the first PCR round contained 39 cycles, and the second, 35 cycles.

Absolute Quantification Using RT-qPCR

For the absolute quantification of viral RNA, we transcribed 1µg of total RNA, and for serum, urine, and fecal samples 2µL of total RNA using random hexamer (500ng/reaction) and
oligo dT (250ng/reaction) primers. For astrovirus quantification, we relied on primers of PoAsV3-qPCR-R/PoAsV3-qPCR-F designed for the highly conserved ORF1b using real-time PCR assay (Maxima SYBR Green qPCR Master Mix, Thermo Scientific, Waltham, MA, USA) (Figure 1 in main text; Technical Appendix Table 1). We followed the quantification steps of the qPCR reactions with a dissociation assay and ran both in a plate format on an ABI 7500 qPCR thermal cycler. For absolute quantification, a single PCR product generated by the PoAstV3-Screen-R and PoAstV3-Screen-F primers was cloned into a pTZ57R/T vector which contains a T7 promoter (InsTAclone PCR Cloning Kit, Thermo Scientific Waltham, MA, USA). We determined the orientation of the insert by sequencing. We linearized and transcribed the purified vectors with TranscriptAid T7 High Yield Transcription Kit (Thermo Scientific Waltham, MA, USA). After DNase-I treatment, we purified the transcribed RNA by phenol-chloroform extraction and ethanol precipitation. We used 10-fold dilution series of purified and spectrophotometrically quantified RNA transcripts in the RT reactions to generate standard curve.

Histology and in situ Hybridization

We designed 30 probe pairs generated at Advanced Cell Diagnostics (Newark, CA, USA) to hybridize to bps 4304–5246 of JX556692.1 within the PoAstV genome (ACD catalog No., V-PoAstV4-ORF2-C2) and to hybridize native viral Ni-PoAstV-3 RNA. As a positive control for nucleic acid integrity in the tissue sections, we used a probe that hybridizes to a conserved region of the highly expressed PPIB of *M. musculus* (Cat. #313919). Negative controls included Dap-B (dihydridopicolinate reductase gene from *Escherichia coli* probe), an unrelated viral probe, and normal porcine brain region-matched sections. We conducted chromogenic in situ hybridization according to the manufacturer’s instructions for nucleic acid detection (RNAscope 2.0 Brown Kit; Advanced Cell Diagnostics, Newark, CA, USA). We baked FFPE slides at 60°C for 1 hour, then deparaffinized with xylene twice, 10 min each time, followed by immersion in 100% ethanol twice before air-drying, then applied a hydrogen peroxide–blocking step, pretreatment 1, to the slides for 10 min at room temperature. We then boiled the slides in pretreatment solution 2 at 100°C for 15 min, followed by protease digestion for 30 min at 40°C to enhance target accessibility. We applied specific or control probes and then incubated the slides at 40°C for 2 h. We washed the slides twice with 1X wash buffer for 2 min at room temperature. We performed a series of 6 signal amplification steps according to manufacturer’s instructions. We incubated
slides with 3,3'-Diaminobenzidine solution at room temperature for 8 min and quenched the reaction with dH2O. We applied Gill’s Hematoxylin for 2 min, rinsed slides in water and “bluing solution,” and dehydrated them in 100% ethanol, 70% ethanol, and xylene before adding a coverslip.

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| Target viruses                        | Oligonucleotide primer ID | Primer sequence (5’→3’) | Size of the PCR product (bp) | Reference                                               |
|--------------------------------------|---------------------------|--------------------------|-----------------------------|---------------------------------------------------------|
| Pestivirus (Flaviviridae)            | PoPestiV-Screen-R         | TTT CCT CTG GCC CTG TTC TT | 323                         | This study                                              |
| Pestivirus (Flaviviridae)            | PoPestiV-Screen-F         | ACA GGR AAG AAY TGC CTG GT |                             | This study                                              |
| Teschovirus (Picornaviridae)         | Teschovirus-5UTR-R        | CCA GCC GCC ACC CTG TCA GGC A | 321                         | Zell et al., 2000†                                      |
| Teschovirus (Picornaviridae)         | Teschovirus-5UTR-F        | AGT TTT GGA TTA TCT TGT GCC C |                             | Zell et al., 2009                                      |
| Senecavirus and Cardiovirus (Picornaviridae) | SVV-Cardio-3D-screen-R   | GTT GGC AGG MGT MAT CTT ATA | 280                         | This study                                              |
| Senecavirus and Cardiovirus (Picornaviridae) | SVV-Cardio-3D-screen-F   | TCA GAT CYY TGG CTG TST CG |                             | This study                                              |
| Sapelovirus (Picornaviridae)         | PSV-5UTR-F                | CCC TGG GAC GAA AGA GCC TG | 383                         | Zell et al., 2000                                      |
| Sapelovirus (Picornaviridae)         | PSV-5UTR-R                | CTT TTA AGT AAG TAG TAA AGG G |                             | Zell et al., 2000                                      |
| Pasaivirus (Picornaviridae)          | SPaV-3Dscreen-R           | CCA TGC ARA GCA AGC TCT AT | 540                         | This study                                              |
| Pasaivirus (Picornaviridae)          | SPaV-3Dscreen-F           | GGT TAT GAT GGT TCT ATA CCA CG |                             | This study                                              |
| Enterovirus (Picornaviridae)         | UnivEnt-5UTR-F            | GTA CCY TTG TRC GCC TGT T | 536                         | Boros et al., 2011‡                                     |
| Enterovirus (Picornaviridae)         | UnivEnt-5UTR-R            | ATT GTC ACC ATA AGC AGC CA |                             | Boros et al., 2011†                                    |
| Kobuvirus (Picornaviridae)           | Univ-Kobo-F               | TGG AYT ACA AGR TGT TTT GAT GC | 216                         | Reuter et al., 2009§                                   |
| Kobuvirus (Picornaviridae)           | Univ-Kobo-R               | ATG TTG TTR ATG ATG GTG TGG A |                             | Reuter et al., 2009                                   |
| Astrovirus (Astroviridae)            | PanAstV-Screen-R          | GGY TTK ACC CAC ATN CCR AA | 443                         | Chu et al., 2008‡                                     |
| Astrovirus (Astroviridae)            | PanAstV-Screen-F1         | GAR TTY GAT TGG RCK CGK TAY |                             | Chu et al., 2008‡                                     |
| Astrovirus (Astroviridae)            | PanAstV-Screen-F2         | GAR TTY GAT TGG RCK AGG TAY |                             | Chu et al., 2008‡                                     |

*The numbers in the oligonucleotide primer IDs are the first (forward) or last (reverse) nucleotide positions of the target sites in the genome of Ni-PoAstV-3 NI-Brain/9-2016a/HUN. The PCR products were sequenced in both directions by primer-walking method using sequence-specific primers (data not shown).
†Zell R, Krumpholz A, Henke A, Birch-Hirschfeld E, Stelzner A, Doherty M, et al. Detection of porcine enteroviruses by nRT-PCR: differentiation of CPE groups I-III with specific primer sets. J Virol Methods. 2000;86:205–18. PubMed http://dx.doi.org/10.1016/S0166-0934(00)00189-0
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### Technical Appendix Table 2. Porcine astrovirus 3 primers used in screening and quantification reactions*

| Reaction type                  | Target genome region | Oligonucleotide primer ID | Primer sequence (5′ → 3′) | Size of the PCR product (bp) |
|--------------------------------|----------------------|---------------------------|---------------------------|------------------------------|
| Nested RT-PCR, first round    | ORF1b - RdRP         | PoAsV3-Screen-R           | AAC Cyt CTC CAC ATA ATT GGC | 356                          |
|                                | ORF1b - RdRP         | PoAsV3-Screen-F           | GAA GTG TTT ATG CAC ATC AAG A | 190                          |
| Nested RT-PCR, second round   | ORF1b - RdRP         | PoAsV3-Screen-F-in        | GCG AAG GAG TTT AGG AGG ACG AA | 256                          |
| Nested RT-PCR, first round    | ORF2 - capsid        | PoAsV3-CAP-R-out          | GGC CCA GTR TTA GGB GCA TT | 250                          |
| Nested RT-PCR, second round   | ORF2 - capsid        | PoAsV3-CAP-R-in           | ACC ATY TGG CAG AT A GTK GA | 100                          |

*The numbers in the oligonucleotide primer IDs are the first (forward) or last (reverse) nucleotide positions of the target sites in the genome of NiPoAstV-3 Ni-Brain/9–2016a/HUN. The PCR products were sequenced in both directions by primer-walking method using sequence-specific primers (data not shown). 

### Technical Appendix Table 3. Oligonucleotide primers used in genome acquisition reactions*

| Reaction type                  | Oligonucleotide primer ID | Primer sequence (5′ → 3′) | Size of the PCR product (bp) |
|--------------------------------|---------------------------|---------------------------|------------------------------|
| long range RT-PCR (3′ RACE)   | PoAsV3–3403-F #          | AAG CCT ACC AGT GGT ACT GCG A | 2991                         |
| RT-PCR                         | PoAsV3–3420-R            | AGT ACC ACT GGT AGG CTT CT | 1755                         |
| RT-PCR                         | PoAsV3–1666-Fgen         | ATG AGT GGK TCA CCW GTT AC |                              |
| RT-PCR                         | PoAsV3–1864-R            | TGT CCT CAT GGC CAG AAG ACT | 1579                         |
| RT-PCR                         | PoAsV3–286-Fgen          | GGC GTC AAY GAR TGG GTT GA |                              |
| RT-PCR (5′RACE)                | PoAsV3–5RACE-R2          | GTT GCT GAG GAC GTA CAC GTT | 469                          |
| RT-PCR (5′RACE)                | PoAsV3–5RACE-R1          | CCA ACA GAG AGC CAA TAA GTA A |                              |

**The numbers in the oligonucleotide primer IDs are the first (forward) or last (reverse) nucleotide positions of the target sites in the genome of NiPoAstV-3 Ni-Brain/9–2016a/HUN. # is used for the 3′ RACE RT-PCR reactions of Ni-Brain/9–2016a/HUN, Ni-SC/9–2016a/HUN, Ni-Brain/173–2016a/HUN and Ni-Brain/386–2015/HUN. The PCR products were sequenced in both directions by primer-walking method using sequence-specific primers (data not shown)**