Experimental inoculation study indicates swine as a potential host for Hendra virus

Mingyi Li¹*, Carissa EMBURY-HYATT¹, Hana M. WEINGARTL¹,²

¹ National Centre for Foreign Animal Disease, Canadian Science Centre for Human and Animal Health, Canadian Food Inspection Agency, Winnipeg, Manitoba, Canada
² University of Manitoba, Winnipeg, Manitoba, Canada

* Corresponding author: Mingyi.Li@inspection.gc.ca

(Received 12 November 2009; accepted 18 January 2010)

Short title: Hendra virus infection of swine
Abstract
Hendra virus (HeV) is a zoonotic virus from the family Paramyxoviridae causing fatal disease in humans and horses. Five-week-old Landrace pigs and five-month-old Gottingen minipigs were inoculated with approximately $10^7$ plaque forming units per animal. In addition to fever and depression exhibited in all infected pigs, one of the two Landrace pigs developed respiratory signs at 5 dpi and one of the Gottingen minipigs developed respiratory signs at 5 dpi and mild neurological signs at 7 dpi. Virus was detected in all infected pigs at 2-5 dpi from oral, nasal, and rectal swabs and at 3-5 dpi from ocular swabs by real-time RT-PCR targeting the HeV M gene. Virus titers in nasal swab samples were as high as $10^{4.6}$ TCID$_{50}$/mL. The viral RNA was mainly distributed in tissues from respiratory and lymphoid systems at an early stage of infection and the presence of virus was confirmed by virus isolation. Pathological changes and immunohistochemical staining for viral antigen were consistent with the tissue distribution of the virus. This new finding indicates that pigs are susceptible to HeV infections and could potentially play a role as an intermediate host in transmission to humans.

Hendra virus / Nipah virus / infection / swine / real-time RT-PCR
1. INTRODUCTION
Hendra virus (HeV), first described in 1994 as an equine morbillivirus, was identified as the cause of respiratory illnesses involving horses and two humans in the town of Hendra, a suburb of Brisbane, Australia. This outbreak resulted in the death of thirteen horses and one human [2, 7]. Sporadic cases of HeV infections have continued to occur in both horses and humans [2]. The outbreaks reported in 2008 and 2009 resulted in the death of ten horses and two veterinarians [1] (ProMed Digest #379, 390, 395, 398, 401, 409, and 413). Since 1994, 4 of the 8 human cases of HeV infections were fatal [1] (ProMed Digest #413). HeV is a zoonotic and Biosafety Level 4 (BSL4) agent. It is also listed as one of the potential biological agents for Bio-terrorism by the Atlantic Treaty Organization.

HeV belongs to a new genus, Henipavirus, in the Paramyxoviridae family together with Nipah virus (NiV), which was identified as a causative agent of human encephalitis in the 1998 outbreak in Malaysia [6, 10]. Limited in vivo studies have been done with both viruses due to the requirements for BSL4 facilities. Experimental HeV infections were performed in horses, cats, and guinea pigs all of which developed fatal disease during several trials. In contrast, rabbits and fruit bats developed antibodies against HeV without any clinical signs [13].

In nature, HeV infections have been detected in horses, humans and bats, the latter being the natural reservoir host of the virus [5]. In a serological survey of 100 swine herds in Queensland, Australia [4], no anti-HeV antibodies were found in the 500 tested serum samples. Since pigs are susceptible to the closely related NiV and considered to be an intermediate host for this virus, the aim of this study was to determine whether pigs can also be susceptible to HeV infection, shed the virus and develop clinical disease characterized by pathological lesions.

2. MATERIALS AND METHODS

2.1. Viruses and cells
Human isolates of HeV and NiV were kindly provided by Thomas Ksiazek and Pierre Rollin (CDC, Atlanta, GA, USA). The HeV and NiV stocks were produced by infecting Vero-76 cell monolayers (American Type Culture Collection, Manassas, VA, USA) at a multiplicity of infection of 0.1. Inoculated cells were then incubated at 37 °C for 72 h or until 80% of the monolayer exhibited a cytopathic effect. Aliquots of clarified (centrifugation at 2 000 ×g, 4 °C, 20 min) supernatant were stored at −150 °C. Virus titres were determined by virus plaque assay in Vero-76 cells as described previously [11].

2.2. Animals and animal care
Hartley guinea pigs (GP) with a body weight of approximately 500 g were obtained from Charles River Laboratories International Inc (St. Constant, QC, Canada). Four-week-old Landrace pigs (P) were obtained from a high health status herd (Sunnyside Colony LTD, Sunnyside, MB, Canada). Five-month-old female Gottingen minipigs (MP) were obtained from Marshall BioResources (North Rose, NY, USA). All animals were acclimatized for a minimum of one week prior to inoculation. All procedures involving live virus were performed under BSL4 conditions. Animal housing and manipulations met the Canadian Council on Animal Care guidelines and were approved by the Animal Care Committee of the Canadian Science Centre for Human and Animal Health.
2.3. Animal experimental design and sample collection

2.3.1. Guinea pigs
Six guinea pigs were inoculated intra-nasally with $3.4 \times 10^6$ plaque forming units (PFU) of HeV per guinea pig. Oral swabs and serum samples were collected on 0, 2, 6, 8 days post-inoculation (dpi). Swab samples were collected using polyester fiber swabs (BD, Sparks, MD, USA) emerged into 2 mL of Dulbecco’s phosphate buffered saline (DPBS) (Sigma, St. Louis, MO, USA), supplemented with 100 IU/mL of Penicillin and 100 µg/mL of Streptomycin. Surviving animals were re-inoculated at 16 dpi with the same dose as above and bled out by cardiac puncture on 25 dpi for serum collection.

2.3.2. Landrace pigs
Two Landrace pigs (P1-2) were inoculated oro-nasally with $6.6 \times 10^7$ PFU of HeV per pig. Two control pigs were inoculated with DPBS and kept separately in a different cubicle.
Nasal, oral, ocular and rectal swabs as well as blood samples were collected on 0, 2, 4, and 5 dpi. The following tissues were collected at the time of necropsy: nasal turbinates, trachea, lung, submandibular, bronchial, inguinal, mesenteric lymph nodes, tonsil, spleen, thymus, frontal lobe of the brain (forebrain), cerebellum, pons, thalamus, basal ganglia, medulla, olfactory bulb, meninges, optic nerve, tongue, heart, liver, kidney, bladder, colon, and duodenum. Bronchial-alveolar lavage fluid (BALF), cerebral spinal fluid (CSF) and urine were also collected. Blood, swabs and selected tissue samples were collected from control pigs on 0 and 6 dpi for analysis along with those from infected animals.

2.3.3. Minipigs
Five Gottingen minipigs (MP2-6) were inoculated intra-nasally only with $2.0 \times 10^7$ PFU of HeV per pig, due to the tight jaw in this breed during anesthesia. MP6, MP5 and MP4 were sacrificed on 3, 5, and 7 dpi, respectively. MP3 and MP2 were re-inoculated on 13 dpi intra-nasally with the same dose of HeV for antiserum production and kept until 20 and 21 dpi, respectively. Four control minipigs were kept separately in a different cubicle. Blood and swab samples were collected at 0, 3, 5, 7, 13, 20 or 21 dpi. A set of tissue and fluid samples corresponding to the set collected from the Landrace pigs was collected at necropsy. For these animals, individual lung lobes were harvested instead of the BALF.

2.4. Histology and immunohistochemistry (IHC)
Tissues collected from pigs and minipigs were fixed in 10% formalin for a minimum of five days and trimmed inside a BSL4 lab. These tissues were left in 10% formalin for an additional 24 hours before removal from BSL4. Paraffin embedded tissue sections were stained with hematoxylin and eosin (HE) according to standard protocols.
Immunohistochemistry was conducted using the protocols described elsewhere [11] with monoclonal antibodies against NiV-N (mixture of F45G2, F45G3 and F45G6) [3].

2.5. Homogenization of tissue samples
The 10% (w/v) tissue homogenates were prepared in cold, Ca\(^{2+}\) and Mg\(^{2+}\) free DPBS supplemented with 1% Pen/Strep (Wisent Inc, St-Bruno, QC, Canada) by homogenization in a closed plastic bag using a MiniMix blender (Interscience Lab. Inc. Rockland, MA, USA) for 30 s at a setting of 9 strokes per second. The homogenized tissues were clarified by centrifugation (2 000 × g, 20 min) and the supernatants were collected for analyses.

2.6. RNA extraction and real-time RT-PCR

2.6.1 RNA extraction
Total RNA was extracted from 100 µL of the serum, swab, and tissue homogenates using TriPure Reagent (Roche Diagnostic Corporation, Indianapolis, IN, USA) as per the manufacturer's instructions. All samples were tested in duplicate. The final RNA pellets were re-suspended in 40 µL of Nuclease-Free water (Ambion, Austin, TX, USA).

2.6.2. Real-time RT-PCR
Viral RNA was detected by semi-quantitative real-time RT-PCR (HeV-M rRT-PCR) using primers (Invitrogen Canada, Burlington, ON, Canada) and probe (Applied Biosystems Ltd., Foster City, CA, USA) targeting a coding region of the HeV matrix protein (M) with an amplification of a 69-bp product [9]. Real-time RT-PCR (rRT-PCR) assays were performed using the ABI Sequence Detection System model 7900HT (Taqman; Applied Biosystems Ltd.) with reagents from the QuantiTec Probe RT-PCR kit (Qiagen, Mississauga, ON, Canada). Each reaction contained 5 µL of template RNA in a final volume of 25 µL. The conditions for real-time RT-PCR were 30 min at 50 °C and 15 min at 95 °C, followed by 45 cycles of 95 °C for 15 s, and 60 °C for 60 s.

2.6.3. Plasmid DNA standard for quantification
A pair of complemented forward (5’-
TCGACTCTTCGACAAAGACGGAACCAATCTGGCATCTTTATGCTCATCTCGGCAGAAGACTGGATGCGCCCCTCTCAATTGGTCCGACGAGCTGGG-3’) and reverse (5’-
AATTCCAGCTCGTCGGACAAATTGAGGAGGGCCATCCACAGTCTTCTGCGGAGATGGGACATGACATCCAGATTGGTTTCCGTCGTTTGTGGG-3’) oligonucleotides with binding sequences of HeV-M primers and probe, were synthesized in order to generate a standard for real-time RT-PCR. The two oligonucleotides were denatured and re-annealed. The product with overhanging SalI and EcoRI ends was subsequently cloned into the respective sites in pGEM-Teasy plasmid (Promega Corporation, Madison, WI, USA) and transformed into Escherichia coli (DH5α) for the standard plasmid stock preparation. The correct nucleotide sequence of the cloned product was confirmed by sequencing and restriction enzyme digestion. The cloned plasmid DNA was prepared using the Qiagen plasmid purification kit and quantified using a NanoDrop ND1000 spectrophotometer (Thermo Scientific, Wilmington, DE, USA). A series of 10-fold dilutions of the plasmid DNA were prepared and dilutions with copy numbers ranging from 10\(^1\) to 10\(^8\) per reaction were used for quantification in HeV-M rRT-PCR. Samples with the number of copies (copy number) per reaction lower than 10\(^2\) (Cycle threshold (Ct) value of 36.0) were considered negative.
2.7. Virus isolation
Virus isolations were performed on Vero-76 cell monolayers seeded in 96-well plates (Corning Costar Corp) by end-point titration, from the same preparations used for HeV-M rRT-PCR. Ten-fold serial dilutions of the tissue homogenate supernatants, sera, or swab samples were made in Dulbecco’s modified Eagles’s medium (DMEM) (Sigma) and incubated on cells (50 µL/well) for 1 h at 37 °C, 5% CO2. Following the incubation, an equal volume of DMEM with 4% fetal bovine serum was added to each well. Plates were incubated for 3 days at 37 °C, 5% CO2. The virus titer was determined in 50% tissue culture infective dose (TCID50) calculated by the method of Reed & Muench [8].

2.8. Indirect enzyme-linked immunosorbent assay (ELISA) for the detection of antibodies against HeV
Binary ethylenimine (BEI)-inactivated HeV was subjected to ultracentrifugation through a 30% sucrose cushion following procedures described elsewhere [3]. The resulting virus pellet was resuspended in ice-cold DPBS and sonicated with a Microson ultrasonic cell disruptor (Misonix Incorporated, Famingdale, NY, USA). Aliquots of semi-purified virus (BEI-HeV) were stored at −70 °C until further use. Nunc ELISA plates were coated with BEI-HeV at 1.3 µg/well in carbonate buffer, pH 9.6 (100 µL/well) and incubated overnight at 4 °C. The plates were then blocked with 5% skim milk in phosphate buffered saline (PBS) for 1 h at 37 °C. Pig serum samples, diluted 1:100 in 5% skim milk in PBS with 0.05% Tween-20 (Sigma) (PBST), were then added (100 µL/well) and incubated for 1 h at 37 °C. The ELISA plates were washed 5 times with PBST. Horse radish peroxidase (HRP)-conjugated goat anti-porcine IgG antibody (Kirkegaard and Perry Laboratories Inc., Gaithersburg, MD, USA), diluted 1:1000 in 5% skim milk in PBST, was added (100 µL/well) and incubated for 1 h at 37 °C. The plates were washed as described above and 100 µL of 2, 2-azino-bis (3-ethylbenzothiazoline-6-sulphonic acid) (ABTS) solution (Roche Diagnostic Corporation), prepared according to the manufacturer’s instructions, was added to the individual wells. The color development was measured at 405 nm (OD405) after 10 min of incubation.

2.9. Plaque reduction neutralization test (PRNT)
The level of HeV-specific neutralizing antibody in heat-inactivated sera (1 h, 56 °C) was determined using the plaque reduction neutralization test with Vero V-76 cells seeded into 24-well plates. Briefly, working virus solutions were made in cell culture medium with a target of 100 PFU/well. Mixtures of virus (200 µL) and serial dilutions of swine serum (200 µL) were incubated at 37 °C for 1 h and then added to the cell monolayers in triplicate wells (100 µL/well). After a 1 h incubation, the virus-serum mixture in each well was replaced with 1 mL of 1% carboxymethylcellulose (CMC) / 2% fetal bovine serum. The plates were incubated at 37 °C, 5% CO2 for 2-3 days followed by fixation with 4% formaldehyde and staining with 0.5% crystal violet in 80% methanol in PBS. Wells with 70% plaque reduction were considered positive for the presence of HeV neutralizing antibodies. The sera were also tested in parallel for cross-neutralization with NiV using the same procedure.

3. RESULTS
3.1. Guinea pigs
Starting at 4 dpi all six guinea pigs, inoculated with HeV, developed signs of depression (reduced feed and water intake, weight loss, and inactivity). One guinea pig was found dead at 5 dpi with blood around the nares and another was found dead at 6 dpi. All surviving guinea pigs recovered at 9 dpi and remained clinically healthy until the end of the experiment. These observations were consistent with previous reports [14, 16]. Viral RNA was detected by HeV-M rRT-PCR in oral swabs from all six HeV-infected guinea pigs at 2-6 dpi, however, the level peaked at different days for each animal. Interestingly, the infection was fatal for guinea pigs with viral RNA at $6.7 \times 10^5$ copies/mL and higher in oral swabs, while guinea pigs with lower viral load survived until the termination of the experiment. Antibodies against HeV were detected in serum samples from infected guinea pigs at 8-25 dpi by BEI-HeV ELISA1.

3.2. Clinical observations in swine
Both of the infected Landrace pigs exhibited increased rectal temperatures of 40.0-40.7 °C and reduced feed intake at 4-5 dpi. Pig P1 was found in lateral recumbency with signs of severe depression and respiratory distress at 5 dpi and both pigs were euthanized that day. No clinical signs of disease were observed in the control pigs. According to the experimental schedule, inoculated minipigs MP6, MP5 and MP4 were sacrificed at 3, 5, and 7 dpi, respectively. A body temperature higher than 40 °C was observed only in minipig MP2 at 9 dpi. Minipig MP5 showed mild depression from 3 to 5 dpi (day of necropsy). Minipig MP2 developed signs of depression, cough, respiratory distress, and transient neurological signs (i.e., incoordination) at 7 dpi, from which it recovered the next day, although depression and cough lasted three additional days. Minipig MP2 and MP3 were clinically healthy after 10 dpi. Minipigs from the control groups showed no clinical signs.

3.3. Histopathology and immunohistology
At necropsy, pig P1 had petechial hemorrhages on kidneys and consolidated lungs. No remarkable changes were observed for pig P2. Minipig MP5 had petechial hemorrhages in multiple organs, including lungs, bronchial and submandibular lymph nodes. Minipig MP4 had areas of consolidation and dark red demarcated areas (hemorrhages) distally on both lung halves. The presence of histological lesions and their relationship to the detection of viral antigen is summarized in Table I. Histological lesions were minimal in minipig M6 at 3 dpi, and consisted of mild pulmonary edema and the presence of multinucleated syncytial cells in the submandibular lymph node. Viral antigen was detected by immunohistochemistry in a few respiratory epithelial cells of the nasal turbinates as well as endothelial cells, lymphocytes, macrophages and dendritic reticular cells in the submandibular lymph nodes. At 5 dpi (minipig MP5), mild to moderate edema was observed in most lung sections examined; however there was only one small focus of inflammation in the bronchiolar epithelium with an occasional formation of multinucleated syncytial cells. Antigen was detected in the bronchial and bronchiolar epithelium of the lung (Fig. 1A). Vasculitis was observed in arteries adjacent to these affected airways.

1 manuscript in preparation.
At 7 dpi in minipig MP4, mild to moderate pulmonary edema could still be observed in some sections of the lung but no other lung lesions were observed and no antigen was detected at this time point. Occasional patches of degeneration of the respiratory epithelial cells and several respiratory epithelial syncytial cells (Fig. 1B) with positive antigen staining were observed in the respiratory epithelium of nasal turbinates. Lesions in the submandibular lymph nodes consisted of scattered multinucleated syncytial cells in areas surrounding connective tissue trabecule. There was extensive immuno-staining concentrated around vessels and connective tissue trabecule within the lymph nodes (Fig. 1C). Most of the lesions in the infected Landrace pigs (P1 and P2), at 5 dpi, were similar to those observed in the minipigs. However, the severity of lesions and viral antigen distribution were more extensive within the lungs and bronchial lymph nodes than those in minipigs. Lesions in the lungs were characterized by a severe interstitial pneumonia. Alveoli were filled with numerous macrophages, fibrin, neutrophils, plasma cells and lymphocytes, as well as numerous necrotic syncytial cells. Spongiosis and necrosis with attenuation as well as bronchiolar syncytial cells were observed in the bronchiolar epithelium (Fig. 1D). Antigen was detected within alveolar macrophages, bronchiolar epithelial cells, alveolar and bronchiolar syncytial cells (Fig. 1E). Hemorrhage, neutrophil infiltration and numerous multinucleated cells were noted within bronchial lymph nodes, with antigen in macrophages, multinucleated cells and subcapsular lining cells. Significant lesions and/or antigen were not observed in the trachea, mesenteric and inguinal lymph nodes, spleen, cerebellum, medulla, forebrain, kidney, duodenum, colon, or thymus from any of the animals examined.

3.4. Viral RNA shedding
Viral RNA was detected by rRT-PCR in all four types of swabs (oral, nasal, rectal and ocular) collected at 2-5 dpi from pigs P1 and P2, except for the ocular swab from pig P2 at 2 dpi (Figs. 2A-D). Greater variation in the presence of RNA in swabs was observed for minipigs, although they all had viral RNA present at some point (see Figs. 2A-D). Overall, viral loads in all four types of swabs were found in decreasing order for nasal, oral, rectal and ocular samples. Landrace pigs had notably higher viral loads in swabs than minipigs (Figs. 2A-D).

3.5. Viral RNA distribution in tissues
Tissues from HeV-infected pigs were screened by real-time RT-PCR for the presence of viral RNA, prior to attempts for virus isolation. Viral RNA was detected in nasal turbinates, trachea, bronchus, BALF, tonsil, submandibular and bronchial lymph nodes of both Landrace pigs P1 and P2, as well as in the tongue from pig P1 (Tab. II). Viral RNA was not detected in the remaining screened tissues, including different regions of the CNS. In minipigs, nasal turbinates and submandibular lymph nodes from all three pigs sacrificed on 3, 5, and 7 dpi (MP6, MP5, and MP4) were positive for viral RNA. Viral RNA was also found in the trachea, right apical and left cardiac lobes of lungs, tonsils and bronchial lymph nodes of minipig MP5, while in minipig MP6 only tonsils and olfactory bulb were positive for the presence of viral RNA. The olfactory bulb was the
only additional tissue with detectable levels of RNA in minipig MP4. An overview of the results is presented in Table II.

3.6. Virus isolation
Virus isolations were performed on samples positive during screening by HeV-M rRT-PCR, and the rRT-PCR was repeated at the time of the attempted virus isolation. Virus was consistently isolated from samples that yielded $3.0 \log_{10}$ copy number or higher per 12.5 µL on the repeat (Tabs. III and IV): namely from tonsils, BALF, oral, nasal and rectal swabs of the Landrace pigs; from lung and olfactory bulbs of the miniature pigs, as well as nasal turbinates, submandibular and bronchial lymph nodes of both breeds. It was noted that copy numbers on repeats were lower than on the screening (data in Fig. 2 and Tab. II), following the additional freeze-thaw, which may have decreased detectable infectivity in the individual samples.

3.7. Antibody development
Serum samples collected from HeV-infected pigs were analyzed by virus neutralization (PRNT) and indirect IgG ELISA using BEI-inactivated virus as antigen. The Landrace pigs P1 and P2 developed antibodies against HeV antigen at as early as 5 dpi. The sera from minipigs MP6 and MP5, sacrificed at 3 and 5 dpi, were negative for antibodies against HeV. The earliest time for detecting antibodies in the minipigs was at 7 dpi in the three remaining infected animals (MP4, MP3 and MP2), with increasing OD values thereafter (Fig. 3). Neutralizing antibodies with titres from 20 to 160 were found only in sera collected after 7 dpi (from minipigs MP3 and MP2). The sera cross-reacted with NiV at similar titers.

4. DISCUSSION
In this inoculation study, pigs from the commercial Landrace breed and the Gottingen minipig breed were able to be infected with HeV thereby providing a proof in principle for the establishment of animal disease models. In both breeds, early in the infection, virus was detected mainly in tissues from respiratory and lymphoid systems which is consistent with previous findings in other animal species [15, 17]. Virus was isolated from nasal, oral, and rectal swabs, indicating the possible routes for virus shedding. Isolation from ocular swabs was unsuccessful, although low copy numbers of viral RNA were detected. Unlike previous findings in other infected species (horse, guinea pig, and cat) [15, 17], the virus was likely not excreted through the urinary tract in infected pigs, based on negative findings in the urine, bladder, or kidney. Virus was not detected in either serum or blood (real-time RT-PCR and virus isolation), spleen, brain, uterus, liver and heart from infected pigs (data not shown), in contrast to guinea pigs and cats [15, 17]. The presence of HeV in the olfactory bulb of infected minipigs indicates that the virus may invade CNS, possibly through the olfactory pathway, although virus (or viral RNA) was not detected in different sections of the central nervous system. At this point, it appears that HeV may not invade the CNS in pigs the same way as NiV [11]. The clinical signs in the two infected Landrace pigs were more severe compared to those observed in minipigs. The Landrace pigs were very sick at 5 dpi, with one developing severe depression, respiratory distress and possible neurological signs, requiring euthanasia. However, due to the small number of experimental animals, different ages (5
weeks for commercial pigs versus 5 months for minipigs), and different breeds it is
difficult to draw any conclusions.
The infection of five-week-old Landrace pigs with HeV appeared to lead to somewhat
more severe clinical disease compared to NiV infection. However due to the considerably
higher dose of HeV used for inoculation compared to NiV (in the range of $10^7$ compared
to $10^5$ PFU/animal, respectively), it is difficult to again draw conclusions at this point.
Since we were initially not sure whether the pigs would be susceptible to HeV, a higher
dose was chosen. Follow-up infection studies would be needed to further investigate the
virulence of HeV in pigs inoculated with a virus dose reflecting the level found in
secretions (e.g., nasal) during natural shedding.
On a preliminary level some differences were observed between HeV and NiV infections
in pigs (Tab. V), particularly in their ability to invade the CNS, which appears to be lower
for HeV. However HeV possibly demonstrated increased virulence for the respiratory
tract, illustrated by the more pronounced respiratory clinical signs than those observed in
experimental infections with NiV, as well as the more abundant presence of antigen in the
respiratory epithelial cells and more frequently observed syncytia namely in respiratory
epithelial cells than in NiV infected pigs [12]. Differences were also observed in virus
shedding, where HeV infected pigs also appear to shed virus via the rectal route.
This study demonstrated that pigs are susceptible to HeV infection through oro-nasal or
nasal inoculations, suggesting that pigs may play a role as an intermediate host for the
spreading of the virus to humans if exposed to infected bats which are considered as the
natural reservoir of henipaviruses. Since infected pigs may shed virus through oral, nasal,
and rectal excretions, possible transmission through close contact with these excretions
may occur. Lack of evidence for HeV infection in domestic swine [4], stresses the
importance of heightened on-farm biosecurity. However, our findings may reinforce the
concern about HeV infecting feral pigs if they are forced to share the same habitat as fruit
bats. This concern is based on the fact that Australia has over 23 million feral pigs living
in Queensland, New South Wales and the Northern Territory, according to reports by
Animal Control Technologies\(^2\).

Acknowledgements
This project was supported by CFIA internal funding. The authors would like to thank
Peter Marszal, Greg Smith, Brett Dalman, Brad Collignon, Shelley Ganske and Estella
Moffat for their technical assistance, Kevin Tierney and Kimberly Azaransky for the
animal care. The authors would also like to thank Marta Sabara for critical review of the
manuscript.

REFERENCES

Figure legends

Figure 1. Histopathological changes and the presence of viral antigen in pigs experimentally infected with Hendra virus.  
(A) Lung (minipig MP5): 5 dpi. Viral antigen was detected in epithelial cells of bronchioles (arrow).  
(B) Nasal Turbinate (minipig MP4); 7 dpi. Formation of epithelial syncytial cells (arrows).  
(C) Submandibular lymph node (minipig MP4); 7 dpi. Viral antigen was observed within reticular dendritic cells (arrows) as well as multinucleated syncytial cells (arrowhead).  
(D) Lung (Landrace pig P1); 5 dpi. A bronchiole shows attenuation and necrosis of the epithelium and formation of syncytial cells (arrow).  
(E) Lung (Landrace pig P1), 5 dpi. Positive immunostaining was observed within bronchiolar epithelial syncytial cells (arrows).  
(B, D) Hematoxylin and eosin (HE) stain; (A, C, E) immunohistochemistry/DAB/hematoxylin (IHC) staining.  
Bar = 20 µm. Magnification of B: × 100 objective.

Figure 2. Real-time RT-PCR analysis of swabs from HeV-infected pigs.  
Values presented as log₁₀ copy number per 12.5 µL of samples. Data displayed were averages from duplicates of each sample.  
2A: nasal swabs; 2B: oral swabs; 2C: rectal swabs; 2D: ocular swabs. Samples considered negative (2.0 log₁₀ copy number or less per reaction) are indicated with grey diamonds.

Figure 3. Detection of antibodies against HeV in serum samples from HeV-infected pigs by indirect ELISA. MP2 to MP6: serum samples from MP2 to MP6; P1 and P2: serum samples from P1 and P2. Data displayed were averages of duplicates of each sample. The arrow indicated the day (13 dpi) when the animals were re-inoculated with HeV.
Table I. Presence of microscopic lesions (HE) and antigen distribution (IHC) in organs of Hendra virus infected pigs.

<table>
<thead>
<tr>
<th>Animal #</th>
<th>dpi</th>
<th>Lung HE/IHC</th>
<th>Nasal turbinates HE/IHC</th>
<th>Submandibular LN HE/IHC</th>
<th>Bronchial LN HE/IHC</th>
<th>Olfactory bulb HE/IHC</th>
<th>Tonsil HE/IHC</th>
</tr>
</thead>
<tbody>
<tr>
<td>MP6</td>
<td>3</td>
<td>+/-</td>
<td>-/+</td>
<td>+/-</td>
<td>+/-</td>
<td>+/-</td>
<td>-/-</td>
</tr>
<tr>
<td>P1</td>
<td>5</td>
<td>+/-</td>
<td>-/-</td>
<td>+/-</td>
<td>+/+</td>
<td>ns</td>
<td>+/-</td>
</tr>
<tr>
<td>P2</td>
<td>5</td>
<td>+/-</td>
<td>+/-</td>
<td>+/-</td>
<td>+/-</td>
<td>ns</td>
<td>+/-</td>
</tr>
<tr>
<td>MP4</td>
<td>7</td>
<td>+/-</td>
<td>+/-</td>
<td>+/-</td>
<td>+/-</td>
<td>+/-</td>
<td>+/-</td>
</tr>
</tbody>
</table>

*Presence of histological lesions/positive immunostaining for viral antigen; - (negative); + (positive); dpi (days post inoculation); ns (tissue not submitted), LN (lymph nodes).
Table II. Real-time RT-PCR screening of tissue samples from HeV-infected pigs.

<table>
<thead>
<tr>
<th>Animal #</th>
<th>MP6</th>
<th>MP5</th>
<th>P1</th>
<th>P2</th>
<th>MP4</th>
</tr>
</thead>
<tbody>
<tr>
<td>DPI</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nasal turbinates</td>
<td>4.6</td>
<td>5.0</td>
<td>4.0</td>
<td>3.3</td>
<td>2.1</td>
</tr>
<tr>
<td>Trachea</td>
<td>-</td>
<td>3.7</td>
<td>3.8</td>
<td>3.7</td>
<td>-</td>
</tr>
<tr>
<td>Lung</td>
<td>NA</td>
<td>NA</td>
<td>-</td>
<td>-</td>
<td>NA</td>
</tr>
<tr>
<td>Lung -right apical</td>
<td>-</td>
<td>2.6</td>
<td>NA</td>
<td>NA</td>
<td>-</td>
</tr>
<tr>
<td>Lung -left cardiac</td>
<td>-</td>
<td>3.6</td>
<td>NA</td>
<td>NA</td>
<td>-</td>
</tr>
<tr>
<td>BALF</td>
<td>NA</td>
<td>NA</td>
<td>6.0</td>
<td>5.3</td>
<td>NA</td>
</tr>
<tr>
<td>Submandibular lymph nodes</td>
<td>3.8</td>
<td>5.3</td>
<td>4.3</td>
<td>4.2</td>
<td>2.8</td>
</tr>
<tr>
<td>Bronchial lymph nodes</td>
<td>-</td>
<td>3.7</td>
<td>3.2</td>
<td>4.9</td>
<td>NA</td>
</tr>
<tr>
<td>Olfactory bulb</td>
<td>2.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>5.0</td>
</tr>
<tr>
<td>Tongue</td>
<td>NA</td>
<td>NA</td>
<td>2.7</td>
<td>-</td>
<td>NA</td>
</tr>
</tbody>
</table>

NA (not available); - (negative, detected but lower than 2.0).
Data displayed were averages of duplicates of each sample. Values shown are log_{10} copy number/12.5 µL of samples, i.e. 10% supernatants of tissue homogenates or undiluted samples of BALF.
Table III. Virus isolation on samples positive by real-time RT-PCR for HeV-M from infected Landrace pigs.

<table>
<thead>
<tr>
<th></th>
<th>DPI</th>
<th>P1</th>
<th>P2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RT</td>
<td>VI</td>
<td>RT</td>
</tr>
<tr>
<td>Oral swabs</td>
<td>2</td>
<td>3.9</td>
<td>8.0</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>4.2</td>
<td>62.8</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>3.8</td>
<td>8.0</td>
</tr>
<tr>
<td>Nasal swabs</td>
<td>2</td>
<td>4.8</td>
<td>10.0</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>5.2</td>
<td>444.5</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>4.0</td>
<td>8.0</td>
</tr>
<tr>
<td>Rectal swabs</td>
<td>4</td>
<td>-</td>
<td>ND</td>
</tr>
<tr>
<td>Nasal turbinates</td>
<td>2.7</td>
<td>ND</td>
<td>3.2</td>
</tr>
<tr>
<td>BALF</td>
<td>4.9</td>
<td>79.0</td>
<td>4.3</td>
</tr>
<tr>
<td>Tonsil</td>
<td>3.2</td>
<td>8.0</td>
<td>3.0</td>
</tr>
<tr>
<td>Submandibular lymph nodes</td>
<td>4.0</td>
<td>8.0</td>
<td>3.6</td>
</tr>
<tr>
<td>Bronchial lymph nodes</td>
<td>-</td>
<td>ND</td>
<td>3.9</td>
</tr>
</tbody>
</table>

RT (HeV-M rRT-PCR, \(\log_{10}\) copy number/12.5 µL of sample); VI (virus isolation, TCID\(_{50}\)/12.5 µL of sample); ND (not detectable); - (negative, detected but lower than 2.0). Samples were re-frozen and stored at -70 °C after the rRT-PCR screening (data in Tab. II). Samples with negative results on virus isolations for both infected Landrace pigs were not presented.
Table IV. Virus isolation on positive samples detected by real-time RT-PCR for HeV-M from infected minipigs.

<table>
<thead>
<tr>
<th></th>
<th>MP4</th>
<th></th>
<th>MP5</th>
<th></th>
<th>MP6</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RT</td>
<td>VI</td>
<td>RT</td>
<td>VI</td>
<td>RT</td>
<td>VI</td>
</tr>
<tr>
<td>Nasal turbinates</td>
<td>-</td>
<td>ND</td>
<td>4.9</td>
<td>108.3</td>
<td>4.8</td>
<td>62.8</td>
</tr>
<tr>
<td>Lung -left cardiac</td>
<td>ND</td>
<td>ND</td>
<td>3.7</td>
<td>8.0</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Submandibular lymph nodes</td>
<td>3.1</td>
<td>1.5</td>
<td>5.0</td>
<td>250.0</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Bronchial lymph nodes</td>
<td>ND</td>
<td>ND</td>
<td>4.7</td>
<td>62.8</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Olfactory bulb</td>
<td>4.8</td>
<td>62.8</td>
<td>ND</td>
<td>ND</td>
<td>2.3</td>
<td>ND</td>
</tr>
</tbody>
</table>

See notes in Table III. Samples with negative results on virus isolations for all infected minipigs were not presented.
Table V. Comparison of Hendra and Nipah virus infections in pigs.

<table>
<thead>
<tr>
<th>Virus</th>
<th>HeV</th>
<th>NiV [11,12]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pigs, age, route and dose of</td>
<td>Gottingen minipig, 5-month-old, IN</td>
<td>Landrace pig, 5-week-old, ON</td>
</tr>
<tr>
<td>inoculation</td>
<td>2.0 \times 10^7 PFU/pig</td>
<td>6.6 \times 10^7 PFU/pig</td>
</tr>
<tr>
<td>Clinical signs</td>
<td>Respiratory and possible neurological</td>
<td>Respiratory</td>
</tr>
<tr>
<td></td>
<td>syncytia</td>
<td>Frequently noticeable syncytia</td>
</tr>
<tr>
<td>Immuno-histology</td>
<td>Frequently noticeable syncytia</td>
<td>Rarely noticeable syncytia</td>
</tr>
<tr>
<td>Virus shedding</td>
<td>Nasal, oral, rectal, ocular</td>
<td>Nasal, oral, rectal, ocular</td>
</tr>
<tr>
<td>Virus distribution in tissues</td>
<td>Respiratory, lymphoid, (CNS? - olfactory</td>
<td>Respiratory, lymphoid, CNS</td>
</tr>
<tr>
<td></td>
<td>bulb)</td>
<td></td>
</tr>
</tbody>
</table>

CNS (central nervous system); IN (intra-nasal); ON (oral and nasal).
Figures
Figure 1
Figure 3

Days post-inoculation (dpi)

OD405

MP6
MP5
MP4
MP3
MP2
P1
P2