Pathogenic Differences between Nipah Virus Bangladesh and Malaysia Strains in Primates: Implications for Antibody Therapy

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Nipah virus (NiV) is a paramyxovirus that causes severe disease in humans and animals. There are two distinct strains of NiV, Malaysia (NiVM) and Bangladesh (NiVB). Differences in transmission patterns and mortality rates suggest that NiVB may be more pathogenic than NiVM. To investigate pathogenic differences between strains, 4 African green monkeys (AGM) were exposed to NiVM and 4 AGMs were exposed to NiVB. While NiVM was uniformly lethal, only 50% of NiVM-infected animals succumbed to infection. Histopathology of lungs and spleens from NiVM-infected AGMs was significantly more severe than NiVM-infected animals. Importantly, a second study utilizing 11 AGMs showed that the therapeutic window for human monoclonal antibody m102.4, previously shown to rescue AGMs from NiVM infection, was much shorter in NiVB-infected AGMs. Together, these data show that NiVB is more pathogenic in AGMs under identical experimental conditions and suggests that postexposure treatments may need to be NiV strain specific for optimal efficacy.

Some 15 years ago, Nipah virus (NiV) emerged and was shown to be a previously unknown paramyxovirus, now classified along with Hendra virus and Cedar virus within the Henipavirus genus. NiV causes febrile encephalitis1 and severe respiratory disease1 in humans with a fatality rate as high as 100% in some outbreaks3. Pteropid fruit bats have been identified as the reservoir for NiV in nature4–6 although pigs served as an amplifying host during the first outbreak of NiV in Malaysia7. Additionally, there are numerous other mammalian species that are susceptible to NiV infection5,8–13.

Genetic analysis has identified at least two strains of NiV responsible for outbreaks in different geographical areas14. The Malaysia strain (NiVM) caused the initial outbreak of NiV from 1998–99 in Malaysia and Singapore in which over 270 people were infected with about 40% case fatality rate (CFR)7,14 with an additional 2014 outbreak in the Philippines with a CFR of ~52%, although the strain identification is based off a short read of the genome15 so it is not completely certain which strain of the NiV caused this outbreak. The Bangladesh strain (NiVB) however has caused repeated outbreaks, varying in number, in Bangladesh and northeast India with outbreaks occurring almost every year between 2001–20155,16–19. The outbreaks of NiVB have had higher CFRs averaging about 75%17 with human-to-human transmission also observed20,21. The observations that these two strains reportedly display differences in CFRs and human-to-human transmission are interesting as there is 91.8% nucleotide homology between the genomes14.

Clinical data from NiV outbreaks has revealed several key differences between patients infected with NiVM and NiVB. First, NiVB has a shorter average incubation period and a more narrow range for the incubation period than NiVM2,22,23. Second, most cases of NiV are presented with respiratory symptoms while few patients infected with NiVM presented with respiratory symptoms12,19. Third, few cases in the Bangladeshi and Indian NiV outbreaks reported myoclonus16, while a significant proportion of patients from the Malaysian outbreak presented with segmental myoclonus22,24 as well as the fatal cases in the Philippines presenting with an acute encephalitis...
syndrome. Fourth, the source of the virus in the Bangladeshi and Indian outbreaks is either unknown in some cases or has been traced to consumption of contaminated fruit or date palm sap, followed by human-to-human transmission and nosocomial spread, whereas the source of the virus in the Malaysian outbreak is known to be from pigs, which served as an amplifying host. Unlike examples found in NiV outbreaks, there were only two reported cases of potential transmission from human-to-human in the Malaysian outbreak, neither of which presented with symptoms during the outbreak, although there were some reported cases of documented human-to-human transmission in the Philippines outbreak. Fifth, there is an increased rate of vomiting with NiV infection compared with the NiVM stocks.

Despite the fact that there have been only one or two outbreaks of NiVM while repeated outbreaks of NiVB have occurred, almost all pathogenesis and vaccine research has utilized NiVM rather than the potentially more medically relevant NiVB strain. Recently, the important questions surrounding these two strains of NiV have been investigated, with particular interest in attempts to compare the strains in animal models that, to date, have accurately reflected the NiVM disease syndromes seen in humans. There are numerous animal models used to study NiVM (reviewed in ref. 31) with the hamster, ferret, and African green monkey (AGM) models most faithfully recapitulating NiV human disease. To date, studies comparing NiVM and NiVB in hamsters and in ferrets have been performed with conflicting results.

While the hamster and ferret models faithfully reproduce the neural and respiratory disease pathology seen in human cases of NiV, recent comparison of NiVM and NiVB in these model systems did not reflect the apparent differences between the strains as observed in humans. However, the observations of strain difference in hamsters, although opposite of the human cases, and the slight difference in oral shedding between the strains in the ferret model, suggested that differences seen in human cases could be due to the strain of virus. To advance an understanding of this issue, we examined the pathogenesis of NiVM and NiVB in the AGM model using virus stocks at the same passage, identical virus dose, with all animals challenged at the same time and by exactly the same route. Here, we report that infection of AGMs with NiVM and NiVB under identical experimental conditions led to similar observations as seen in the human cases between the strains with NiVB being more pathogenic for respiratory tissues, suggesting that one of the main differences in human outbreaks could be attributed to the strain of NiV. Additionally, we showed that the therapeutic window for NiV treatment of AGMs with the human monoclonal antibody m102.4 is shorter than for NiVM.

Results

Deep sequencing of NiVM and NiVB isolates. To compare the P2 NiV stocks used in this study to the published NiV nucleotide sequences, the viral stocks were deep sequenced. These data are presented in Supplementary Table 1. There were 10 differences of sufficient frequency to note between the P2 stock of NiVM and the reference sequence GenBank Assession number AJ627196.1. Of these, two were non-coding, seven were silent mutations, and one led to a single amino acid change in the N protein. There were four mutations of sufficient frequency to note between the P2 stock of NiVB and the reference sequence GenBank Assession number AY988601.1. Of these, one was non-coding, and the other three led to single amino acid changes: one in the M protein and two in the F protein. In this present study, the NiV strains were passage matched at P2 however, we were interested in a previous passage 3 NiVM stock used in an antibody therapeutic study. Deep sequencing was also used to analyze the NiVM P3 stock. Compared to the reference sequence GenBank Assession number AJ627196.1, there were two mutations at sufficient frequency to note. One of these was silent and one led to a single amino acid change in the Glycoprotein (G).

NiVM and NiVB growth kinetics in target endothelial cells. To assess the growth kinetics of NiVM and NiVB stocks, which were passaged in the same cell lines and same number of times, we infected brain (hCMEC/D3) or pulmonary endothelial cell lines (hpmeCs.t1.6r) or primary brain (HBCMEC) or pulmonary (HPMEC) cells as these mimic the assumed in vivo target cells. In the endothelial cell lines NiVM titers seemed to peak by 24 hpi whereas NiVB produced higher levels of infectious virus at 48 hpi in the pulmonary cell line (Fig. 1a, dark blue). This result was intriguing as NiVB outbreaks have more respiratory sequelae associated with human cases when compared to NiVM. When we further characterized the two strains in the primary cell lines, NiVB was found to grow to higher titers than NiVM in both the brain (Fig. 1b, light blue) and pulmonary (Fig. 1b, dark blue) primary endothelial cells suggesting that NiVB has better fitness in target endothelial cells. In addition, Vero E6 cells are used to propagate NiV isolates, and the peak titers observed at 48 hpi are comparable between NiVB and NiVM. While previous studies in hamsters revealed that NiVB had better fitness in BHK-21 cells and subsequently in hamsters, these data in human endothelial cells led us to compare the two NiV strains in the AGM model of disease as it best recapitulates the human NiV disease course.

NiVM and NiVB challenge and disease in AGMs. We previously described the development of a NiVM disease model in AGMs which recapitulated clinical signs and pathology observed in NiV-mediated disease in humans. Clinical signs in this model include severe depression, respiratory disease leading to acute respiratory distress, neurologic disease, reduced activity, and a time to death ranging from 9 to 12 days. To date NiVB challenge of AGMs has yet to be described; additionally, a direct comparison of the Malaysian and Bangladeshi strains of NiV in AGMs has not been examined. To explore the differences between these NiV strains we challenged two cohorts of four AGMs each by the i.n. and i.t. routes with either NiVM or NiVB and observed them over the course of 15 days post-challenge. After challenge, subjects from each cohort were observed and scored for clinical signs of disease based on depression, recumbency, respiratory quality, and neurological signs (Table 1). All four animals in the NiVB cohort succumbed to NiV-mediated disease on day 7 (Fig. 1c, blue); whereas two of the four animals in the NiVM cohort succumbed to NiV-mediated disease on day 10 and the two remaining animals survived until the end of the 15 day study (Fig. 1c, red) the results of which were significant by Log-rank (Mantel-Cox) test with
All animals from the NiVM cohort reached low, initial clinical scores on day 7, due to respiration quality and depression, with all animals scoring over the course of 4–7 days and two of the four animals (O8000 and O8023) reaching clinical scores that required euthanasia on day 10 post-challenge (Fig. 1c, red). As found in Table 1, each animal in the NiVM cohort experienced depression, fever, a loss of appetite, dyspnea, lymphopenia, thrombocytopenia, and hypoalbuminemia with noticeable nasal exudates from three of the four animals (O7912, O7935, and O8000) with epistaxis associated with O8000; additionally, two of the four animals experienced tremors (O7935 and O8000). The NiVB cohort also initially scored on day 7; however, these animals displayed a more severe clinical score due to depression, respiration quality, and recumbency ultimately succumbing to NiV-mediated disease on day 7 (Fig. 1c, blue). Animals in the NiVB cohort also experienced fever, depression, a loss of appetite (up to 3 days earlier than NiVM cohort, O8021 Table 1), severe dyspnea, lymphopenia, thrombocytopenia, hypoalbuminemia, noticeable nasal exudate with epistaxis in O8022, and tremors noted for O8046. Overall, the clinical scores for each group were associated with the period of observed clinical illness where animals infected with NiVM had a more protracted illness as seen in the clinical illness column in Table 1.

NiVM and NiVB load in swabs, blood, and tissues. To determine if there was a difference in virus replication between the NiVM- and NiVB-infected cohorts, shedding of virus was assessed by qRT-PCR on nasal, oral, and rectal swabs (Fig. 2a–c, respectively) with viremia also screened by qRT-PCR on whole blood samples (Fig. 2d). High levels of NiV GEq were detectable in nasal swabs for both cohorts starting at day 1 and through day 7 with the NiVM group having positive nasal swabs through day 15 (Fig. 2a). The NiVM oral swabs had detectable NiV GEq starting at day 3 and continuing through day 10 while the NiVB oral swabs were positive starting at day 5 and reaching up to two logs higher levels at day 7 (Fig. 2b). NiV GEq were detected at levels up to 3 logs higher in the blood of NiVB-infected animals at day 5 and 7 post-challenge when compared to the NiVM cohort. Infectious virus was isolated from all whole blood NiV RNA positive samples (Fig. 2d, +) while one NiVM whole blood sample was positive from O8023 on day 7 (Fig. 2d, +). Typically, detection of infectious virus from whole blood samples that are NiV RNA positive has been inconsistent in the NiVM model; however, the higher levels of circulating NiV in blood may have contributed to the consistent isolation of infectious virus from RNA positive samples.

To further examine the replication of the viruses in AGMs, we performed qRT-PCR on tissue samples collected after animals succumbed to NiV-mediated disease or at the end of the 15 day study (O7912 and O7935).
Table 1. Clinical findings and outcome for AGMs challenged with NiV. aSurvived to day 15. bO7936 and O8021 succumbed before sampling at day 7 could be achieved. cInfectious virus isolated from plasma on day(s) indicated. dDays after NiV challenge are in parentheses. Fever is defined as a temperature more than 2.5 °F over baseline or at least 1.5 °F over baseline and ≥103.5 °F. eLymphopenia and thrombocytopenia are defined by a ≥30% drop in numbers of lymphocytes and platelets, respectively. ALT- alanine aminotransferase; CRP- C reactive protein; AST- aspartate aminotransferase; BUN- blood urea nitrogen.

| Subject No. | Sex | NiV strain | Viral loada | Clinical illnessb | Clinical and gross pathologyd |
|-------------|-----|------------|-------------|-------------------|--------------------------------|
| O7912a      | Male | Malaysia   | 0°          | Fever (d7); Depression (d7–11); Loss of appetite (d7–11); Dyspnea (d7–12); Labored breathing (d7–12); Nasal exudate (d9). | Thrombocytopenia (d10); Lymphopenia (d7); Hypoalbuminemia (d10,15); >2-fold increase in ALT (d5); Increase in CRP (d7,15); Moderate congestion at gastroduodenal junction. |
| O7935a      | Female | Malaysia | ϕ | Fever (d7); Depression (d7–12); Loss of appetite (d7–13); Dyspnea (d7–12); Labored breathing (d7–12); Lethargy (d9–12); Tremors (d9–13); Nasal exudate (d5). | Thrombocytopenia (d10); Lymphopenia (d6,8); Hypoalbuminemia (d10,15); Increase in CRP (d7,10). |
| O8000      | Male | Malaysia | ϕ | Fever (d7); Depression (d7–10); Loss of appetite (d7–10); Dyspnea (d7–10); Labored breathing (d7–10); Lethargy (d9–10); Tremors (d10); Nasal exudate (d10); Epistaxis (d10). | Animal euthanized d10. |
| O8023      | Male | Malaysia | d7 | Fever (d7); Depression (d7–10); Loss of appetite (d7–10); Dyspnea (d7–10); Labored breathing (d7–10); Lethargy (d9–10); Hypoalbuminemia (d10,15); Increase in CRP (d7,10); Congestion of nares; Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Enlarged spleen darkened and mottled; Liver slightly darkened; Kidney pale and reticulated; Stomach with mild multifocal areas of congestion and small petechia on mucosal surface; Mild congestion at gastroduodenal junction; Prominent hemorrhages on mucosal surface of urinary bladder. |
| O8021b      | Male | Bangladesh | d5 | Loss of appetite (d4–7); Dyspnea (d7); Labored breathing (d6–7). Animal succumbed d7. | Lymphopenia (d5); Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Enlarged spleen darkened and mottled; Liver darkened and reticulated; Kidney pale and reticulated; Stomach with multifocal areas of congestion and small petechia on mucosal surface; Hemorrhages on mucosal surface of urinary bladder; Congestion of small blood vessels in the brain. |
| O7936c      | Female | Bangladesh | d5 | Loss of appetite (d6–7). Animal succumbed d7. | Thrombocytopenia (d5); Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; spleen darkened and mottled; liver darkened and reticulated; kidney pale and reticulated; hemorrhages on mucosal surface of urinary bladder. |
| O8022      | Male | Bangladesh | d5,7 | Fever (d7); Depression (d7); Loss of appetite (d7); Dyspnea (d7); Labored breathing (d7); Lethargy (d7); Nasal exudate (d7); Epistaxis (d7). Animal succumbed d7. | Thrombocytopenia (d5,7,10); Lymphopenia (d7); Hypoalbuminemia (d10); >2-fold increase in AST (d7); >3-fold increase in ALT (d10); Increase in CRP (d7,10); Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Enlarged spleen darkened and mottled; Liver slightly darkened; Kidney pale and reticulated; Stomach with mild multifocal areas of congestion and small petechia on mucosal surface; Mild congestion at gastroduodenal junction; Hemorrhages on mucosal surface of urinary bladder. |
| O8046      | Male | Bangladesh | d5,7 | Fever (d7); Depression (d7); Loss of appetite (d7); Dyspnea (d7); Labored breathing (d7); Lethargy (d7); Nasal exudate (d6–7); Tremors (d7). Animal succumbed d7. | Thrombocytopenia (d7); Lymphopenia (d7); Hypoalbuminemia (d7); >2-fold increase in AST (d5,7,10); Increase in CRP (d7,10); Frothy sanguineous nasal exudates; Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Spleen darkened and mottled; Liver darkened and reticulated; Kidney pale and reticulated; Stomach with multifocal areas of congestion and small petechia on mucosal surface; Hemorrhages on mucosal surface of urinary bladder. |

Strikingly, the detectable levels of NiV B GEq in the respiratory tissues were 3 logs higher from the bronchi to the lower lobes of the lungs when compared to the NiV M group (Fig. 3a). However, NiV GEq detected in neural tissue were similar among all neural tissues sampled (Fig. 3b). The results from the tissues in AGMs were similar to growth of NiVM and NiVB in microvascular lung endothelial cell and microvascular brain endothelial cell lines where NiV B grew to higher titers in the lung endothelial cells but the titers were equivalent in the brain endothelial cells (Fig. 1a) whereas NiV B grew to higher titers in both primary microvascular lung and brain endothelial cells (Fig. 1b). Examination of the lymphoid tissues revealed higher levels of NiV B GEq in the axillary and inguinal lymph nodes as well as in the spleen (Fig. 3c) while additional tissues with higher levels of NiV B included the liver and adrenal gland (Fig. 3d).

To determine if the isolation of infectious virus from tissues would reflect the difference observed between the groups in regard to the whole blood samples, tissues were selected for virus isolation from the qRT-PCR positive tissues from at least one tissue from each panel (Fig. 3). Infectious NiV B was isolated from all lung lobes for each animal (Fig. 3a, +) whereas one animal per lobe had infectious NiV M isolated including O7912 from the right upper lobe (Fig. 3a, +). Three animals (O7935, O8000, and O8023) from the NiV M cohort had infectious virus isolated from the cerebellum (Fig. 3b, +); two animals from the NiV B group were positive for infectious virus in the cerebellum as well (O7936 and O8022). Cervical spinal cord samples from one animal in each cohort had infectious virus isolated; O7912 from the NiV M group and O8022 from the NiV B group. Only animals from the NiV B group had infectious virus isolated from spleen samples (Fig. 3c, +) while one animal from the NiV M group (O8000) and two animals from the NiV B (O8021 and O8022) had virus isolated from the kidney (Fig. 3d, +).
Gross pathology, histopathology, and immunohistochemical analysis of NiV-infected AGMs. Typical gross lesions found in the urinary bladder and brain (Supplementary Fig. 1) of NiV-infected animals were similar between the NiVM and NiVB groups; however, examination of respiratory tissues revealed a characteristic failure of lung lobes to collapse with multifocal areas of hemorrhage and congestion (Supplementary Fig. 2a,b) with an excess of serosanguinous fluid in the pleural cavity (Supplementary Fig. 2c,d; asterisks) for both NiVM and NiVB. However, overall the NiVB lung lobes were more engorged with edema as the lobes were more rounded (Supplementary Fig. 2b, arrows) (i.e., inflated balloon) upon examination when compared to the NiVM lung lobes (Supplementary Fig. 2a, arrowheads) (i.e., collapsed balloon). Histopathological and immunohistochemical examination revealed interstitial pneumonia, edema, hemorrhage, and vascular pathology, indicated by the endothelial syncytial cell formation (Fig. 4d, asterisk/inset), were more severe in the NiVB cohort (Fig. 4a–d). Ordinal scoring of the lung lesions revealed a statistically significant difference between the NiVB cohort (blue) and all NiVM animals (dark red) as well as to the NiVM cohort that succumbed (bright red) (Fig. 5a). The NiVM cohort that succumbed also had a significant difference in lung lesions compared to the two animals that did not succumb, though not as significant as compared to NiVB (Fig. 5a). Additionally, the overall quantity and distribution of NiV antigen was more prevalent in the NiVB lung sections (Fig. 4a vs. c), with immunohistochemical ordinal scoring revealing a median of 0 for the NiVM cohort and 4 for the NiVB cohort. These observations suggest that the efficiency of oxygen exchange and blood flow in the lungs would be more reduced in the NiVB animals compared to the NiVM animals and indeed we observed an accentuation of the normal lobular pattern in the liver as indicated by pale multifocal to coalescing foci on the capsular surface of the liver (Supplementary Fig. 2d, inset). This observation coupled with such striking lung pathology is suggestive of reduced cardiovascular efficiency and passive hepatic congestion in which vulnerable zones of hepatocytes are subjected to periods of hypoxia and undergo vacuolar degeneration with adjacent hepatocyte zones having distended/congested sinusoids which is further suggestive of the apparent increase in lung pathology in the NiVB tissues.

Histopathological and immunohistochemical examination of the spleen and kidney lesions revealed differences that were also striking between the NiVM and NiVB animals that succumbed to disease. The lymphoid follicular architecture of the splenic white pulp was minimally altered in the NiVM animals (Fig. 4f), whereas partial to complete obliteration of the white pulp with hemorrhage, lymphoid depletion (Fig. 4h), and numerous, massive syncytial cells (Fig. 4h, asterisks/inset) were noted in the NiVB group. Ordinal scoring of the spleen lesions revealed a statistically significant difference between the NiVB cohort (blue) and all NiVM animals (dark red) as well as to the NiVM cohort that succumbed (bright red) (Fig. 5b). The NiVM cohort that succumbed did not have a significant difference in spleen lesions compared to the two animals that did not succumb (Fig. 5b). Strong immunoreactivity for NiV antigen was also more abundant in the spleens of NiVB animals (Fig. 4e vs. g), with immunohistochemical ordinal scoring revealing a median of 1 for the NiVM cohort and 4 for the NiVB cohort. In the kidney tissues the NiVM and NiVB had minimal interstitial lymphoplasmacytic infiltration, glomerular congestion, fibrin deposition, and immunolabeling for NiV antigen (Fig. 4i–l). However, the NiVB kidney tissues overall had stronger immunolabeling for NiV antigen (Fig. 4i vs. k) and more endothelial cell syncytial formation in the glomerulus and medullary vessels (Fig. 4l, asterisk).
Time of m102.4 treatment is crucial for protection of AGMs from NiV<sub>B</sub>-mediated disease.

Previously, we showed that administration of the NiV neutralizing human mAb m102.4 beginning as late as 5 days after NiV<sub>M</sub> challenge of AGMs resulted in complete protection at a time when animals were viremic and showed signs of NiV disease.<sup>37</sup> Considering the higher rates of replication, increased lung pathology, and shorter time to death observed in AGMs challenged with NiV<sub>B</sub> when compared to NiV<sub>M</sub>, we were interested in determining whether or not initiating a similar post-challenge treatment regimen with m102.4 could protect AGMs against NiV<sub>B</sub>. An in vitro assessment of the neutralizing capability of m102.4 against NiV<sub>B</sub> revealed that m102.4 was at least as potent at neutralizing NiV<sub>B</sub> when compared to NiV<sub>M</sub> (Fig. 6a). With the encouraging m102.4 in vitro neutralization data, we proceeded to assess the protective efficacy of m102.4 against NiV<sub>B</sub> challenge. The m102.4 antibody was administered to AGMs i.v. twice, with the first dose given 1 day after infection and the second dose 3 days after virus challenge (Fig. 6b, black arrows). To determine if the m102.4 therapeutic window could be extended, we treated two additional groups: one group received m102.4 beginning at 3 days after challenge and again at 5 days after challenge (Fig. 6b, green arrows), and the third treatment group received m102.4 at 5 days after infection and again at 7 days after virus challenge (Fig. 6b, blue arrows). No adverse reactions were observed following m102.4 infusion in any of the 9 treated animals. The two control subjects, consistent with the pathogenesis study above, succumbed to NiV<sub>B</sub> disease on days 7 and 8 (Fig. 6c, red), respectively, and had signs of severe sustained behavior changes (depression, decreased activity), loss of appetite, fever (O7923), dyspnea, lymphopenia, thrombocytopenia, hypoalbuminemia, and changes in liver enzymes (Table 2). In contrast, subjects treated with m102.4 at day 1 after infection and again at 3 days later (D1/D3) survived challenge (Fig. 6c, gray) and showed little to no clinical signs of disease with normal hematology and minor clinical chemistry results (Table 2). The three animals treated at day 3 after infection and again two days later (D3/D5) survived NiV<sub>B</sub> challenge (Fig. 6c, green) and did not show any changes in behavior; these animals had mild changes in clinical pathology as compared to the D1/D3 treatment group (Table 2). The third treatment group differed as all three animals treated at day 5 after infection and again two days later (D5/D7) succumbed to NiV<sub>B</sub>-mediated disease.

Figure 3. Viral load in tissues. NiV<sub>M</sub> (red hues) and NiV<sub>B</sub> (blue hues) viral RNA genomic equivalents detected by qRT-PCR from tissues at study endpoint for each AGM in pulmonary associated tissues (a), neural tissues (b), lymphoid tissues (c), or other typical NiV-infected tissues (d). Plus sign denotes infectious virus isolation from sample; underlined plus sign denotes virus isolated from every AGM in group for that particular sample. R (right), L (left), RU (right upper), RM (right middle), RL (right lower), LU (left upper), LM (left middle), LL (left lower), LN (lymph node). Error bars represent standard deviation.
on day 8 post-challenge (Fig. 6c, blue); all three also had signs of severe sustained behavior changes (depression, decreased activity), loss of appetite, fever, dyspnea, lymphopenia, thrombocytopenia, hypoalbuminemia, and changes in liver enzymes as seen in the control group (Table 2). The surviving animals from the D1/D3 and D3/D5 groups all had detectable neutralizing antibody after treatment and through the end of the study unlike the control group and much higher than the observed levels for the D5/D7 group (Supplementary Table 2). These data show that in the AGM NiVB disease model, the window to therapeutically intervene is clearly shorter when compared to a previous NiVM study in which all D5/D7 (Fig. 6c, historical) had a significant effect on survival when compared to the current D5/D7 group (Fig. 6c, *37).

m102.4 treatment effect on NiV load and subsequent humoral response in AGMs. NiV genome replication was assessed by qRT-PCR on nasal, oral, and rectal swabs (Supplementary Fig. 3a, b and b, respectively) with viremia also screened by qRT-PCR on whole blood samples (Supplementary Fig. 3d). NiV genome equivalents were observed in nasal swabs in all groups starting on day 3 (Supplementary Fig. 3a) and varied between groups in oral swabs from the control (red), D3/D5 (green), and D5/D7 (blue) groups with the D1/D3 group (gray) only positive on day 3 (Supplementary Fig. 3b). Viral RNA was detected in blood samples at day 3 from the D1/D3 group (O8214), the D3/D5 group (O8149), and at high levels from day 5 for the D5/D7 (blue) and control (red) groups (Supplementary Fig. 3d). High levels of NiV RNA were also detected systemically in the tissues of all animals in the D5/D7 and control groups (Supplementary Fig. 3e). These data are consistent with the detection of viral antigen by immunohistochemistry where only the control and D5/D7 groups were positive for antigen in the major target organs lung and spleen (Fig. 7).

Considering we detected circulating NiV RNA in all groups examined, we wanted to assess the role of the host immune response in mediating protection for the D1/D3 and D3/D5 groups from NiV disease. To do this, we measured circulating levels of antibodies directed against the NiV F glycoprotein in sera of the NiV-infected AGMs (Fig. 8). Anti-NiV F antibodies were present in sera of all D1/D3 and D3/D5 m102.4-treated AGMs with antibody levels increasing by day 28 (O7522, O8149, and O8211) while the D5/D7 group and two control AGMs did not seroconvert to high levels.
Discussion

Two strains of NiV have been responsible for human outbreaks in different geographical regions\(^{14,15}\). Clinical data from these NiV outbreaks has revealed several key differences between patients infected with NiV\(_{M}\) and NiV\(_{B}\). Despite the fact that there have been only one or two outbreaks of NiV\(_{M}\) while repeated outbreaks of NiV\(_{B}\) have occurred, almost all pathogenesis, therapeutic, and vaccine research has utilized NiV\(_{M}\) models rather than the potentially more relevant NiV\(_{B}\) strain based on number of outbreaks and potential for human-to-human transmission. Recently, the important questions surrounding these two strains of NiV have been investigated, with particular interest in attempts to compare the strains in animal models that, to date, have accurately reflected the NiV\(_{M}\) disease syndrome seen in humans. Numerous animal models have been used to study NiV\(_{M}\) (reviewed in ref. 31) with the hamster\(^{11,32}\), ferret\(^{12,33}\), and AGM\(^{13,34}\) models most faithfully recapitulating NiV human disease.

To date, studies comparing NiV\(_{M}\) and NiV\(_{B}\) in hamsters\(^{35}\) and in ferrets\(^{36}\) have been performed with conflicting results. Comparison of NiV\(_{M}\) and NiV\(_{B}\) infection in hamsters revealed that NiV\(_{M}\) had more fitness in baby hamster kidney cells and caused a more rapid and pathogenic disease course \(^{35}\); however, respiratory tract lesions were found to be similar between the two strains in a separate study\(^{38}\). Interestingly, the higher fatality rate and more rapid disease progression of NiV\(_{M}\)-infected hamsters compared to NiV\(_{B}\) differ from what has been reported in human cases. Additionally, the hamster model has recently revealed a NiV\(_{B}\) food-borne infection and transmission model replicating the assumed outbreak conditions in India and Bangladesh\(^{39}\). From the hamster model data, it has been suggested that there are no intrinsic differences between the two strains of NiV\(^{38}\).

To further compare NiV\(_{M}\) and NiV\(_{B}\), the ferret model was employed for pathogenic examination. After inoculation, it was found that the disease course, clinical signs, and pathology of the two strains were similar in ferrets\(^{36}\). However, a few differences were noted, including a higher rate of virus shedding in oral secretions and a higher incidence of focal hepatic necrosis in NiV\(_{B}\)-infected ferrets, whereas there was a higher viral load in the blood at time of euthanasia and a higher rate of hemorrhagic diathesis in the NiV\(_{M}\)-infected ferrets. Interestingly, in contrast to the hamster model, the higher rate of virus shedding in oral secretions in NiV\(_{B}\)-infected ferrets suggested a potential mechanism for the higher rate of human-to-human transmission in NiV\(_{B}\)-infected humans.

In the current study, we compared the NiV\(_{M}\) and NiV\(_{B}\) strains with similar passage history in AGMs. The initial report on the NiV\(_{M}\) model in AGMs was reported as nearly uniformly lethal with various routes of infection and doses being examined and 7 of 8 AGMs succumbing. Subsequent studies have revealed the model to be less lethal than initially thought, depending on dose and route of infection\(^{15,35-40}\). When these studies are combined, all animals challenged with various combinations of i.t, i.n., intraperitoneal, and/or oral routes with various doses have only had 12 out of 19 (\(\sim 63\%\) lethality) total AGMs succumb, although all AGMs did show clinical disease. The lethality does not appear to be dependent on the various challenge doses used in these studies.

In the present study we challenged AGMs under identical conditions; virus passage history/cells, dose of inoculum, and route of inoculation. We found that AGMs challenged with NiV\(_{B}\) resulted in 100% mortality compared to 50% for NiV\(_{M}\), at least within the 15 day window of acute, respiratory-mediated disease; within these constraints we found the survival curves to be significantly different (Fig. 1c, \(*\) ). We cannot rule out the possibility that the surviving AGMs infected with NiV\(_{M}\) would not have succumbed later due to late-onset neurological...
disease, although some survivors in the previously mentioned studies survived to day 32 without developing neurological disease. In addition to the 6 AGMs infected with NiV₉ presented in the current study, we have 4 additional AGMs which have been infected with NiV₉ under identical conditions presented in the current study although at three different times (unpublished data). These animals all succumbed between 5–9 days p.i. Thus, including historical data, we see 10 of 10 AGMs infected with NiV₉ succumbing with only 12 of 19 AGMs infected with NiV₉₈ succumbing. Additionally, from these unpublished data, historical data, and the present study we see AGMs infected with NiV₉₈ succumbing between days 5–9, while AGMs infected with NiV₉ succumb between days 9–12. Specifically, in this current study, NiV₉₈ had a shortened time to death in AGMs by an average of 2.8 days; NiV₉ produced higher titers in target human pulmonary endothelial cell lines and primary cells, and higher virus load in the lungs and oral swabs of infected AGMs. A higher NiV₉ virus load was also observed in the blood, lymphoid tissue, and liver in AGMs when compared to NiV₉₈-infected AGMs. These data together suggest that there is an intrinsic difference between the two strains of NiV, particularly as the AGM model recapitulated the more severe respiratory disease and significant histopathology differences, shorter time to death, and higher mortality observed in NiV₉ human outbreaks when compared to outbreaks of NiV₉₈. These differences require

Figure 6. Human mAb m102.4 activity against NiV₉ in vitro and in vivo. (a) Percent neutralizing antibody dose response of m102.4 against NiV₉ (red) and NiV₉₈ (blue) infection in Vero cells. Error bars show standard deviation. (b) Diagram of the m102.4 treatment regimen and sampling days post NiV₉ challenge in AGMs. Black arrows; D1/D3 treatment. Green arrows; D3/D5 treatment. Blue arrows; D5/D7 treatment. * depicts the day of NiV₉ challenge. (c) Kaplan-Meier survival curves for the D1/D3 group (gray, n = 3), D3/D5 group (green, n = 3), the D5/D7 group (blue, n = 3), and the control group (red, n = 2). Log-rank (Mantel-Cox) Test; *p-value < 0.05 comparing the NiV₉ D5/D7 cohort with the previously reported data from the NiV₉₈ D5/D7 cohort.
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further characterization to elucidate which proteins and/or genomic elements are responsible for NiV<sub>B</sub> having a more acute and increased respiratory disease.

Recently we reported the 100% successful therapeutic human mAb m102.4 treatment of AGMs infected with NiV<sub>M</sub> showing clinical signs of disease along with detectable viremia. Additionally, m102.4 was employed in 2013 in the United States to treat a possible laboratory exposure to NiV<sub>B</sub> under a compassionate use protocol. This individual has never shown any evidence of henipavirus infection. Although this individual did not experience any NiV<sub>B</sub> disease manifestations considering the shortened disease course and higher viremia in the NiV<sub>B</sub> AGM model, we were interested in determining whether or not the therapeutic treatment window with m102.4 was similar to the NiV<sub>M</sub> model. Importantly, we found that the treatment window for NiV<sub>B</sub> is shorter as the D5/D7 treatment group succumbed to NiV<sub>B</sub>-mediated disease whereas this treatment regimen was completely protective against NiV<sub>M</sub>-mediated disease in AGMs with a significant difference in survival curve observed between the D5/D7 treatment groups. All AGMs in the D3/D5 group seroconverted in the NiV<sub>M</sub> study whereas only 1 AGM out of 4 in the NiV<sub>B</sub> D3/D5 study seroconverted. The average D5/D7 reciprocal neutralizing titers on day 7 NiV<sub>B</sub> post-challenge were much lower (426.6) compared to the NiV<sub>M</sub> study (5120) as well. All of these observed differences are most likely due to the more rapid and higher viral loads experienced by AGMs after exposure to NiV<sub>B</sub> versus NiV<sub>M</sub> but should be considered when assessing treatment options between someone exposed to NiV<sub>M</sub> or NiV<sub>B</sub>.

The exact mechanism of treatment failure in the D5/D7 group is not clear. The half-life of m102.4 in AGMs ranges from 10–12 days and therefore is not the cause of treatment failure in the D5/D7 group. Rather, the D5/D7 group had a noticeably decreased level of neutralizing antibody of the exogenously infused m102.4 compared to the other cohorts (320–640 compared to 1280–2560) at the time point 2 days after the first infusion (Supplementary Table 2). This may be due to higher loads of circulating virus on the day of initial treatment being bound by the m102.4, although since there was one animal (O8149) in the Day 3/5 group with a similarly high level of viremia on the day of its first treatment; that animal survived and did not have a low neutralizing titer. Perhaps the significant differences observed in the spleen histopathology between NiV<sub>M</sub> and NiV<sub>B</sub> (Fig. 5b) could account for this as these cells and germinal centers would also be needed to fully recover during a therapeutic treatment study. Alternatively, it may be due to loss of m102.4 through the leaky capillaries in the D5/D7 group as observed by the hypoalbuminemia observed in this group (Table 2). It is unknown if a higher dose of m102.4 treatments may have overcome these potential reasons for treatment failure in the D5/D7 group. There are however, at least two other considerations for the failure of the treatment for NiV<sub>B</sub>. We found no changes in the amino

### Table 2. Clinical Description and Outcome of NiV<sub>B</sub>-Challenged and m102.4 Treated AGMs.

| Subject No. | Sex | Group | Clinical illness | Clinical and gross pathology |
|-------------|-----|-------|-----------------|-----------------------------|
| O7923       | Male| Control | Fever (d7); Depression (d7–8); Lethargy (d7–8); Loss of appetite (d6–8); Dyspnea (d8); Labored breathing (d5–8). Animal euthanized on d8. | Thrombocytopenia (d5–7,8); Lymphopenia (d5,7) Hypoalbuminemia (d8); >7-fold increase in ALT (d8); >2-fold increase in AST (d5,7); >10-fold increase in AST (d8); 2-fold increase in BUN (d8); 4-fold increase in CRE (d8); Increase in CRP (d7–8); Excess fluid in the pleural cavity; Lungs inflated and enlarged with coalescing multifocal areas of congestion and hemorrhage; Excess pericardial fluid; Enlarged adrenal glands; Liver darkened and reticulated; Congestion on mucosal surface of urinary bladder; Congestion of small blood vessels in the brain. |
| C4373       | Male| Control | Depression (d7); Lethargy (d7); Loss of appetite (d5–7); Severe dyspnea (d7); Labored breathing (d5–7); Nasal exudates (d7); Epistaxis (d7). Animal euthanized on d7. | Thrombocytopenia (d7); Lymphopenia (d3,5,7); Hypoalbuminemia (d7); >2-fold increase in AST (d5); >3-fold increase in AST (d7); Increase in CRP (d5, 7); Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Extensive fibrin mats along pleural surfaces; Liver darkened and reticulated; Multifocal areas of hemorrhage on mucosal surface of urinary bladder; Congestion of meninges in the brain. |
| O7698       | Female| D1/D3 | Mild abdominal breathing (d6–7); Mild dyspnea (d7). | None |
| O8215       | Male| D1/D3 | None | >2-fold increase in ALT (d5); >2-fold increase in AST (d7). |
| O8214       | Male| D1/D3 | Mild abdominal breathing (d6). | None |
| O7522       | Male| D3/D5 | None | Lymphopenia (d5,7); >2-fold increase ALT (d7); >2-fold increase AST (d7). |
| O8149       | Female| D3/D5 | None | Lymphopenia (d3, 4,7); Increase in CRP (d7). |
| O8211       | Male| D3/D5 | None | Lymphopenia (d7); >3-fold increase in AST (d5); >2-fold increase in AST (d7). |
| B4825       | Male| D5/D7 | Fever (d7); Depression (d8); Loss of appetite (d7–8); Dyspnea (d7–8); Labored breathing (d5–8); Tremors (d8). Animal euthanized on d8. | Thrombocytopenia (d7,8); Lymphopenia (d7,8); Hypoalbuminemia (d7,8); >2-fold increase in AST (d5); >3-fold increase in AST (d7); >10-fold increase in AST (d8); Increase in CRP (d5,7,8); Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Fibrin mats along pleural surfaces; Liver darkened and reticulated; Congestion of meninges in the brain. |
| B6484       | Male| D5/D7 | Fever (d7); Depression (d8); Loss of appetite (d6–8); Dyspnea (d5–8), Labored breathing (d5–8); Nasal exudates (d8). Animal euthanized on d8. | Thrombocytopenia (d8); Lymphopenia (d3,5,7); Hypoalbuminemia (d8); >2-fold increase in ALT (d5,7,8); >2-fold increase in AST (d5,7); >3-fold increase in AST (d8); >2-fold increase in BUN (d8); Increase in CRP (d7); Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Minimal fibrin mats along pleural surfaces; Congestion and fluid within the meninges of the brain. |
| C3882       | Male| D5/D7 | Fever (d7); Depression (d8); Loss of appetite (d6–8); Dyspnea (d5 and 8), Labored breathing (d5–8); Tremors (d8). Animal succumbed d8. | Thrombocytopenia (d3,7,8); Hypoalbuminemia (d8); >4-fold increase in AST (d8); >2-fold increase in BUN (d8); Increase in CRP (d7,8); Excess fluid in the pleural cavity; Lungs inflated and enlarged with multifocal areas of congestion and hemorrhage; Extensive fibrin mats along pleural surfaces; Liver darkened and reticulated; Multifocal areas of hemorrhage on mucosal surface of urinary bladder; Congestion of meninges in the brain. |
acids involved in the m102.4 binding pocket of the NiV glycoprotein as these are completely conserved between all isolates of NiVβ and NiVγ examined previously43 as well as in the isolates used in this study (Supplementary

Figure 7. Immunohistochemistry of AGM tissue after NiVβ challenge and m102.4 treatment. Lack of NiV antigen in representative m102.4 D1/D3 and D3/D5 treated tissues and localization of NiV antigen in representative control and D5/D7 treated tissues by immunohistochemical staining. Lung and spleen were labeled with an N protein-specific polyclonal rabbit antibody and images taken at 20X magnification.
media at 1 hour post-infection (hpi) as a baseline and from the same wells at 24 and 48 hpi.

and appropriate media (described above) was added back to cells. Samples were taken from initial addition of

software based on unpaired analysis of 125 bp overlaps. Torrent using 318-v2 deep sequencing chips. Sequence analysis was performed using DNA Star Seqman NGen

using Encore Ion Torrent library prep kit. Sequencing was performed by the UTMB Molecular Core on the Ion

using the NuGen Ovation RNA-seq 2.0 kit ultimately for the preparation of the double stranded DNA library

removal from the BSL-4. RNA was isolated from the Trizol LS/sample mixture using Zymo Research Direct-zol

which has implications for how aggressive treatment initiation should be undertaken after exposure to NiVB as it

is often difficult for decisions on what treatments should be given for drugs that do not have licensure.

Methods

Virus isolates. The isolate of NiV\textsubscript{M} used in the pathogenic comparison study was 199902916 and was

obtained from a fatal human case in 1999 and passed on Vero E6 cells twice making this a passage 2 virus. The

isolate of NiV\textsubscript{B} used in the pathogenic comparison and m102.4 study was 200401066 and was obtained from a

fatal human case during the outbreak in Rajbari, Bangladesh in 2004 and passed on Vero E6 cells twice making this

a passage 2 virus. Both strains of NiV used in this study were kindly provided by Dr. Thomas G. Ksiazek.

Each NiV challenge virus stock was assessed for the presence of endotoxin using The Endosafe\textsuperscript{TM}-Portable Test System (PTS) (Charles River). Virus preparations were diluted 1:10 in Limulus Amebocyte Lysate (LAL) Reagent Water (LRW) per manufacturer’s directions and endotoxin levels were tested in LAL Endosafe\textsuperscript{TM}-PTS cartridges as directed by the manufacturer. Each preparation was found to be below detectable limits while positive controls showed that the tests were valid.

Approximately 1 ml of NiV stock was removed from the seed vial and placed in 5 ml of Trizol LS and vortexed

3 times and allowed to sit for 10 minutes. The 6 ml were then placed into 2 separate 3 ml Nunc cryo-vials for

removal from the BSL-4. RNA was isolated from the Trizol LS/sample mixture using Zymo Research Direct-zol RNA mini-prep per manufacturer’s instructions. Approximately 150 ng of purified RNA were used to make cDNA

using the NuGen Ovation RNA-seq 2.0 kit ultimately for the preparation of the double stranded DNA library

using Encore Ion Torrent library prep kit. Sequencing was performed by the UTMB Molecular Core on the Ion

Torrent using 318-v2 deep sequencing chips. Sequence analysis was performed using DNA Star Seqman NGen

software based on unpaired analysis of 125 bp overlaps.

Endothelial cell culture methods and NiV infection. The human brain microvascular endothelial cell line, hCMEC/D3, was a generous gift from Dr. Babette Weksler (Cornell, Ithaca, NY, USA) and was maintained in EBM-2 media (Lonza, Walkersville, MD, USA), containing 5% fetal bovine serum, 1.4 μM hydrocortisone, 5 μg/ml absorbic acid, 1 ng/ml basic fibroblast growth factor (Sigma), 1X chemically defined lipid concentrate, 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), and Penicillin-Streptomycin (Life Technologies). Media was changed every 2–3 days.

The human lung microvascular endothelial cell line, hpmec1.6r, was a generous gift from Drs Unger and Kirkpatrick (Johannes-Gutenberg University, Mainz, Germany). These cells were maintained in medium containing M199, 20% FCS, Glutamax (2 mm), Penicillin-Streptomycin (100U/100 μg/ml), heparin (50 μg/ml), and ECGS (50 μg/ml) and G418 (50 μg/ml, Life Technologies). Cells were grown in cell culture vessels precoated with gelatin (0.2%). Media was changed every 2–3 days.

Human pulmonary microvascular endothelial cells (HPMEC) were maintained in EBM-2 MV media supplemented with 5% fetal bovine serum and proprietary concentrations of human epidermal growth factor, fibroblast growth factor, vascular endothelial growth factor, insulin-like growth factor, hydrocortisone, and gentamicin all provided by Lonza. Human brain cerebral microvascular endothelial cells (HBCMEC) were obtained from Scientell and maintained in EBM-2 containing 5% fetal bovine serum, 1.4 μM hydrocortisone, 5 μg/ml absorbic acid, 1 ng/ml basic fibroblast growth factor (Sigma, St. Louis, MO, USA), 1X chemically defined lipid concentrate, 10 mM HEPES, and Penicillin-Streptomycin (Life Technologies). Media was changed every 2–3 days and cells were used up to passage 7.

To assess the growth kinetics of both NiV strains in target cells, ECs were infected with NiV\textsubscript{M} or NiV\textsubscript{B} at a MOI of 5 (MOI of 1 for primary cells) with rocking for 1 hour at 37 °C. Inoculum was removed, cells were washed, and appropriate media (described above) was added back to cells. Samples were taken from initial addition of media at 1 hour post-infection (hpi) as a baseline and from the same wells at 24 and 48 hpi.

![Graph showing the detection of NiV F specific antibodies from m102.4 treated and non-treated AGMs.](image)

**Figure 8. Seroconversion of AGMs post-challenge and m102.4 treatment.** Detection of NiV F specific antibodies from m102.4 treated and non-treated AGMs. D1/D3 group (gray, n = 3), D3/D5 group (green, n = 3), the D5/D7 group (blue, n = 3), and the control group (red, n = 2). Mean fluorescence intensities (MFI) are shown on the y-axis and represent binding of specific Ig (IgG, and IgM) to NiV F. Error bars represent the standard deviation of fluorescence intensity across 100 beads for each sample.

Table 1. Additionally, escape mutants were not the cause as we were unable to isolate any escape mutants from the blood or tissues recovered from the D5/D7 group at the terminal time points. Regardless of the reason for the D5/D7 treatment failure, we have shown that the efficacious treatment window is shorter for NiV\textsubscript{B} in AGMs which has implications for how aggressive treatment initiation should be undertaken after exposure to NiV\textsubscript{B} as it is often difficult for decisions on what treatments should be given for drugs that do not have licensure.
**Pathogenesis Challenge.** Eight, adult AGMs weighing 4 to 6 kg were used in this study and were randomized into two groups of 4 animals each (3 males and 1 female per group). Animals were inoculated with \(~5 \times 10^5\) PFU of either NiV\(_M\) or NiV\(_B\) with the dose being equally divided between the intratracheal (i.t.) and the intranasal (i.n.) routes for each animal. After challenge, animals were monitored for clinical signs of illness including temperature, respiration quality, blood count, and clinical pathology on days 0, 1, 3, 5, 7, 10, and 15 post-challenge.

**Challenge and m102.4 Treatment.** Eleven adult AGMs weighing 4–8 kg were inoculated by i.t. and i.n. routes with \(~5 \times 10^5\) PFU of NiV\(_B\) as above. Three animals were infused with m102.4 beginning 1 day after challenge and again 3 days after challenge (D1/D3); three animals were infused with m102.4 beginning 3 days after challenge and again 5 days after challenge (D3/D5); three animals were infused with m102.4 beginning 5 days after challenge and again 7 days after challenge (D5/D7). The two control animals were infused with saline. Each dose of m102.4 (~15 mg/kg) was administered intravenous (i.v.) Animals were anesthetized for antibody infusion and clinical examination including temperature, respiration rate/quality, blood collection, and swabs of nasal, oral, and rectal mucosa on days 0, 3, 5, 7, 10, 15, 21, and 28 post-challenge.

**Hematology and Serum Biochemistry.** Hematological analysis including total white blood cell counts, white blood cell differentials, red blood cell counts, platelet counts, hematocrit values, total hemoglobin concentrations, mean cell volumes, mean corpuscular volumes, and mean corpuscular hemoglobin concentrations were analyzed from blood collected in tubes containing EDTA using a laser based hematologic analyzer (Beckman Coulter). Serum samples were tested for concentrations of albumin, amylase, alanine aminotransferase (ALT), aspartate aminotransferase (AST), alkaline phosphatase (ALP), gamma-glutamyltransferase (GGT), glucose, cholesterol, total protein, total bilirubin (TBIL), blood urea nitrogen (BUN), creatinine (CRE), and C-reactive protein (CRP) by using a Piccolo point-of-care analyzer and Biochemistry Panel Plus analyzer discs (Abaxis).

**RNA Isolation.** Immediately following sampling, 100 μl of blood was added to 600 μl of AVL viral lysis buffer (Qiagen) for RNA extraction. For tissues, approximately 100 mg was stored in 1 ml RNAlater (Qiagen) for 7 days to stabilize RNA. RNAlater was completely removed, and tissues were homogenized in 600 μl RLT buffer (Qiagen) in a 2-mL cryovial using a tissue lyser (Qiagen) and ceramic beads. The tissues sampled included conjunctiva, tonsil, or/o nasopharynx, nasal mucosa, trachea, right bronchus, left bronchus, right lung upper lobe, right lung middle lobe, right lung lower lobe, right lung upper lobe, right lung middle lobe, bronchial lymph node (LN), heart, liver, spleen, kidney, adrenal gland, pancreas, jejunum, colon transversum, brachial plexus, brain (frontal and cerebellum), brain stem, cervical spinal cord, pituitary gland, mandibular LN, salivary gland LN, inguinal LN, axillary LN, mesenteric LN, urinary bladder, testes or ovaries, and femoral bone marrow. All blood samples were inactivated in AVL viral lysis buffer, and tissue samples were homogenized and inactivated in RLT buffer prior to removal from the BSL-4 laboratory. Subsequently, RNA was isolated from blood and swabs using the QiAamp viral RNA kit (Qiagen), and from tissues using the RNeasy minikit (Qiagen) according to the manufacturer’s instructions supplied with each kit.

**Detection of NiV load.** RNA was isolated from blood or tissues and analyzed using strain specific primers/probe targeting the N gene and intergenic region between N and P of NiV M or NiVB for quantitative real-time PCR (qRT-PCR) with the probes used here being 6-carboxyfluorescein (6FAM)-5′-CGT CAC ACA TCA GCT CGT ACG A-3′ for NiV\(_M\), 6-carboxytetramethylrhodamine (TAMRA) for NiV\(_B\)-6FAM-5′-CGT CAC ACA TCA GCT CGT ACG A for NiV\(_B\) (Life Technologies, Carlsbad, CA). This strategy using the intergenic region allows for genome and anti-genome detection only without detecting contaminating viral mRNA. NiV RNA was detected using the CFX96 detection system (Bio-Rad) in One-step probe qRT-PCR kits (Qiagen) with the following cycle conditions: 50 °C for 10 minutes, 95 °C for 10 seconds, and 40 cycles of 95 °C for 10 seconds and 59 °C for 30 seconds. Threshold cycle (CT) values representing NiV genomes were analyzed with CFX Manager Software, and data are shown as genome equivalents (GEq). To create the GEq standard, RNA from NiV challenge stocks was extracted and the number of NiV genomes was calculated using Avogadro's number and the molecular weight of the NiV genome.

Virus titration was performed by plaque assay with Vero cells from all blood and tissue samples. Briefly, increasing 10-fold dilutions of the samples were adsorbed to Vero cell monolayers in duplicate wells (200 μl); the limit of detection was 25 PFU/ml.
Histopathology and immunohistochemistry analyses and scoring. Necropsy was performed on all subjects. Tissue samples of all major organs were collected for histopathologic and immunohistochemical examination and were immersion-fixed in 10% neutral buffered formalin for at least 21 days in BSL-4. Subsequently, formalin was changed; specimens were removed from BSL-4, processed in BSL-2 by conventional methods and embedded in paraffin and sectioned at 5 μm thickness. For immunohistochemistry, specific anti-NiV immunoreactivity was detected using an anti-NiV N protein rabbit primary antibody at a 1:5000 dilution for 30 minutes. The tissue sections were processed for immunohistochemistry using the Dako Autostainer (Dako). Secondary antibody used was biotinylated goat anti-rabbit IgG (Vector Laboratories, Burlingame, CA) at 1:200 for 30 minutes followed by Dako LSAB2 streptavidin-HRP (Dako) for 15 minutes. Slides were developed with Dako DAB chromagen (Dako) for 5 minutes and counterstained with hematoxylin for one minute. Non-immune rabbit IgG was used as a negative staining control.

To assess the differences in histopathology Principles for Valid Histopathologic Scoring in Research were used where a scoring system was 1) defined 2) repeatable and 3) produced meaningful results. The tissues for the study were group masked and scored in an ordinal scoring fashion, meaning that samples were assigned to a category in an ordered progression in severity; such as, 0 to 4 for immunohistochemistry and 0 to 5 on H&E based on an estimated percentage of the organ showing the lesion. For the ordinal scoring in this study the median was assessed as this is more appropriate when recording the central tendency. The slide sets were reviewed by one pathologist multiple times to reduce any interpretation consistency. Median values were calculated on selected subjects. Tissue samples of all major organs were collected for histopathologic and immunohistochemical examina-

NiV<sub>a</sub> serum neutralization assays. Neutralization titers against NiV<sub>a</sub> were determined by a conventional serum neutralization assay. Briefly, m102.4 or sera were serially diluted fivefold or twofold respectively, and incubated with ~100 PFU of NiV<sub>a</sub> for 1 h at 37°C. Virus and antibodies mixtures were then added to individual wells of 6-well plates of Vero cells. Plates were stained with neutral red 2 days after infection and plaques were counted 24 h after staining. The 50% neutralization titer was determined as the serum dilution at which at there was a 50% reduction in plaque counts versus control wells.

Measurement of circulating F glycoprotein specific antibodies. Antibodies to the fusion (F) glycoprotein were measured in NiV<sub>p</sub>-infected AGMs by including a recombinant soluble F (sF) glycoprotein-coupled microsphere in the assay by coupling of sF to microsphere #43 (Luminex Corporation)\(^{12}\). Plasma from NiV<sub>p</sub>-infected AGMs was inactivated by gamma irradiation (~5 mrad) prior to testing. Sera and plasma were assayed at 1:5,000 and 1:10,000 dilutions. Assays were performed on a Luminex\(^{8}\) 200 IS\(^{9}\) machine equipped with Bio-Plex Manager Software (v 5.0) (Bio-Rad Laboratories). Mean fluorescent intensity (M.F.I.) and the standard deviation of fluorescence intensity across 100 beads were determined for each sample.

Statistics. Conducting animal studies in BSL-4 severely restricts the number of animal subjects, the volume of biological samples that can be obtained and the ability to repeat assays independently and thus limit statistical analysis. Consequently, data are presented as the mean calculated from replicate samples, not replicate assays, and error bars represent the standard deviation across replicates.

Prism 5 software was used to calculate statistical significance throughout this study to perform ANOVA with Dunnett’s Multiple Comparison Test for viral growth kinetics, m102.4 neutralization, and viral load data; Log-rank (Mantel-Cox) Test for Kaplan-Meier survival curves; Mann-Whitney test for ordinal scoring of tissue pathology.

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Interpretations, conclusions, and recommendations are those of the authors and are not necessarily endorsed by the paper. All authors had access to all of the data and approved the final version of the manuscript. Opinions, gross pathologic, histologic, and immunohistochemical analysis of the data. C.E.M., B.A.S. and T.W.G. wrote assays. C.E.M., B.A.S., J.B.G., Y.-P.C., K.N.A., R.W.C., V.B., C.C.B. and T.W.G. analyzed the data. K.A.F. performed antibody assays and K.N.A. performed the anti-NiV antibody assays. C.E.M. and K.N.A. performed the PCR the Nipah virus infectivity and neutralization assays. Y.-P.C. and L.Y. developed and optimized the anti-NiV Laboratory. B.A.S., J.B.G. and K.N.A. performed the clinical pathology assays. J.B.G. and V.B. performed in vitro C.E.M., R.W.C. and T.W.G. conceived and designed the TWG and AI054715 and AI077995 to CCB and in part by the Intramural Research Program of the NIAID, NIH. This study was supported by the Department of Health and Human Services, National Institutes of Health, grants AI082121 to

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Author Contributions

C.E.M., R.W.C. and T.W.G. conceived and designed the in vitro work. C.E.M., C.C.B. and T.W.G conceived and designed the AGM study. R.W.C. propagated the primary human endothelial cells and participated in the in vitro work. C.E.M., J.B.G. and T.W.G. performed the Nipah challenge experiments at the Galveston National Laboratory. B.A.S., J.B.G. and K.N.A. performed the clinical pathology assays. J.B.G. and V.B. performed the Nipah virus infectivity and neutralization assays. Y.-P.C. and L.Y. developed and optimized the anti-NiV antibody assays and K.N.A. performed the anti-NiV antibody assays. C.E.M. and K.N.A. performed the PCR assays. C.E.M., B.A.S., J.B.G., Y.-P.C., K.N.A., R.W.C., V.B., C.C.B. and TWG. analyzed the data. K.A.F. performed gross pathologic, histologic, and immunohistochemical analysis of the data. C.E.M., B.A.S. and T.W.G. wrote the paper. All authors had access to all of the data and approved the final version of the manuscript. Opinions, interpretations, conclusions, and recommendations are those of the authors and are not necessarily endorsed by UTMB.
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