A Lethal Disease Model for Hantavirus Pulmonary Syndrome in Immunosuppressed Syrian Hamsters Infected with Sin Nombre Virus

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Sin Nombre virus (SNV) is a rodent-borne hantavirus that causes hantavirus pulmonary syndrome (HPS) predominantly in North America. SNV infection of immunocompetent hamsters results in an asymptomatic infection; the only lethal disease model for a pathogenic hantavirus is Andes virus (ANDV) infection of Syrian hamsters. Efforts to create a lethal SNV disease model in hamsters by repeatedly passaging virus through the hamster have demonstrated increased dissemination of the virus but no signs of disease. In this study, we demonstrate that immunosuppression of hamsters through the administration of a combination of dexamethasone and cyclophosphamide, followed by infection with SNV, results in a vascular leak syndrome that accurately mimics both HPS disease in humans and ANDV infection of hamsters. Immunosuppressed hamsters infected with SNV have a mean number of days to death of 13 and display clinical signs associated with HPS, including pulmonary edema. Viral antigen was widely detectable throughout the pulmonary endothelium. Histologic analysis of lung sections showed marked inflammation and edema within the alveolar septa of SNV-infected hamsters, results which are similar to what is exhibited by hamsters infected with ANDV. Importantly, SNV-specific neutralizing polyclonal antibody administered 5 days after SNV infection conferred significant protection against disease. This experiment not only demonstrated that the disease was caused by SNV, it also demonstrated the utility of this animal model for testing candidate medical countermeasures. This is the first report of lethal disease caused by SNV in an adult small-animal model.

Sin Nombre virus (SNV) and Andes virus (ANDV), both members of the genus Hantavirus within the family Bunyaviridae, are the predominant etiological agents of hantavirus pulmonary syndrome (HPS) in North and South America, respectively (1–3). Hantaviruses are negative-strand, single-stranded RNA viruses with three segments, denoted small (S), medium (M), and large (L). The S segment encodes the nucleoprotein (N), the M segment encodes the glycoproteins (Gn and Gc), and the L segment encodes the RNA-dependent RNA polymerase (RdRp) (4). Hantaviruses predominantly infect microvascular endothelial cells and create a vascular leakage-based disease by altering the barrier properties of the endothelium (5). This nonlytic infection renders the endothelium unable to regulate tissue fluid accumulation, leading to pulmonary edema, tachycardia, shock, and cardiac failure (6). The mechanism underlying this endothelium dysfunction and related pathogenic effects remains unknown.

Currently, Syrian hamsters infected with ANDV remain the only small-animal lethal disease model for hantaviruses that cause HPS (7). This disease model closely mimics human disease in incubation time, infection of the endothelium, and rapid onset of disease. Hamsters develop respiratory distress in the final 24 h prior to death approximately 10 to 14 days postinfection when challenged with 200 to 2,000 PFU of ANDV. Mportal virus, not known to cause disease in humans, causes a disease similar to HPS in Syrian hamsters, but with lower morbidity and mortality (8). Along with SNV, infection of hamsters with other hantaviruses (e.g., Hantaan [HTNV], Puumala [PUUV], Dobrava [DOBV], and Seoul [SEOV]) results in an asymptomatic infection that is rapidly cleared and is distinguishable only by the subsequent presence of neutralizing antibodies (9–12). The lack of disease associated with hamster infection with the aforementioned hantaviruses, except ANDV, limits our understanding of the pathogenesis of these viruses. Development of disease models for these viruses is an important step to expand the capability of evaluating potential therapeutics for hantavirus disease.

Cyclophosphamide is a chemotherapeutic agent that suppresses B cell and T cell function (13–15) and also causes apoptosis in other cell types, including neutrophils, macrophages, and dendritic cells (16, 17). Dexamethasone is a glucocorticosteroid that acts as an immunosuppressant by interfering with NF-κB-dependent gene activation, inhibiting lymphocyte proliferation, and reducing inflammatory responses by reducing proinflammatory gene expression and inducing anti-inflammatory genes (18). Both cyclophosphamide and dexamethasone have been used as immunosuppressive agents to alleviate or create disease in animal models of infectious diseases, namely, West Nile virus (19), severe acute respiratory syndrome (SARS) (20), lymphocytic choriomeningitis virus (21), and adenovirus (22).

In the current study, we take an alternative approach to create a model of lethal hantavirus disease by evaluating the pathogenesis of a normally infectious yet nonpathogenic hantavirus in immunocompetent hamsters and in immunocompromised hamsters treated with dexamethasone and cyclophosphamide individually and in combination. This model of SNV-associated lethal disease in Syrian hamsters is the first report of an HPS-like disease caused by SNV in an adult small-animal model and should improve ef-
forts to develop vaccines and therapeutics to prevent and treat hantavirus disease in humans.

MATERIALS AND METHODS

Virus, cells, and medium. SNV strain CC107 (23) was propagated in Vero E6 cells (Vero C1008, ATCC CRL 1586). Preparation of twice-plaque-purified SNV stock has been described previously (7). Cells were maintained in Eagle’s minimum essential medium with Earle’s salts containing 10% fetal bovine serum, 10 mM HEPES, pH 7.4, penicillin-streptomycin (Invitrogen) at 1 X, and gentamicin sulfate (50 µg/ml) at 37°C in a 5% CO2 incubator.

Dexamethasone and cyclophosphamide administration. Water-soluble dexamethasone and cyclophosphamide monohydrate were purchased from Sigma-Aldrich. On the indicated days, anesthetized hamsters were injected intraperitoneally (i.p.) with the indicated dosages per kilogram of body weight of drug diluted in sterile phosphate-buffered saline (PBS), pH 7.4.

Challenge with hantavirus. Female Syrian hamsters 6 to 8 weeks of age (Harlan, Indianapolis, IN) were anesthetized by inhalation of vaporized isoflurane using an IMPAC 6 veterinary anesthesia machine. Once anesthetized, hamsters were injected with 2,000 PFU of virus diluted in PBS. Intramuscular (i.m.) (caudal thigh) injections consisted of 0.2 ml delivered with a 1-ml syringe with a 25-gauge, five-eighths-inch needle.

Blood chemistries. Blood samples were collected in lithium heparin capillary blood collection tubes. Concentrations of alanine aminotransferase (ALT), aspartate aminotransferase (AST) were determined using a comprehensive metabolic reagent disc and an Abaxis Piccolo xpress chemistry analyzer.

Hematology. Blood samples collected in lithium heparin capillary blood collection tubes were analyzed using an Advia 120 hematology analyzer using proprietary software version 3.1.8.0-MS. The dog setting was used for the complete blood count (CBC), and the guinea pig setting was used for the differential.

Plaque assay. Hantavirus plaque assays were performed as described (12).

PRNT. Plaque reduction neutralization tests (PRNT) were performed as previously described (7). Serum samples were gamma-irradiated on dry ice with 3 X 106 rad from a 60 Co source.

N-specific ELISA. The enzyme-linked immunosorbent assay (ELISA) used to detect N-specific antibodies (N-ELISA) was described previously (10, 24). The endpoint titer was determined as the highest dilution that had an optical density (OD) greater than the mean OD for serum samples from negative-control wells plus 3 standard deviations. The PUUV N antigen was used to detect SNV N-specific antibodies as previously reported (7).

Isolation of RNA and real-time PCR. Approximately 250 mg of lung tissue was homogenized in 1.0 ml TRIzol reagent using gentleMACS M tubes and a gentleMACS dissociator on the RNA setting. Serum samples were added directly to TRIzol reagent. RNA was extracted from TRIzol samples as recommended by the manufacturer. The concentration of the extracted RNA was determined using a NanoDrop 8000 instrument and raised to a final concentration of 10 ng/µl. Real-time PCR was conducted on a Bio-Rad CFX thermal cycler using an Invitrogen Power SYBR green RNA-to-CT on a Bio-Rad CFX thermal cycler using an Invitrogen Power SYBR green l. Real-time PCR was conducted

Dose(s) (mg/kg) (day[s] postinfection) Loading Maintenance

Dex | 16 (–3), 8 (–2, –1) | 4 (0–13) | NA
CyP | 140 (–3) | 100 (–1, 1, 4, 7, 10, 13) | NA
Dex/CyP | Same dosing as Dex and CyP alone | Same dosing as Dex and CyP alone | NA

No treatment | NA | NA

a Dexamethasone (Dex) and cyclophosphamide (CyP) were administered by i.p. injection. NA, not applicable.

b On days when both compounds were administered to the dexamethasone/cyclophosphamide group, compounds were combined into a single injection.

RESULTS

Dexamethasone and cyclophosphamide immunosuppress Syrian hamsters. In order to develop an immunosuppressed hamster model, groups of three hamsters were administered dexamethasone and cyclophosphamide, alone or in combination, according to the dosing schedule outlined in Table 1. On day 0, all hamsters were infected with 2,000 PFU of SNV i.m. WBC counts were monitored prior to and after virus infection to confirm immunosuppression in treatment groups (Fig. 1A). Dexamethasone and cyclophosphamide, alone and in combination, resulted in statistically significant reductions in WBC counts compared to no-treatment controls (for dexamethasone, P = 0.0428; for cyclophosphamide, P = 0.0010; and for dexamethasone and cyclo-
phosphamide, \( P = 0.0020 \)). The combination of dexamethasone and cyclophosphamide elicited the most immunosuppressive activity of the three treatment groups (3- to 4-fold reduction across all time points) based on WBC counts. Similarly, the combination of dexamethasone and cyclophosphamide also resulted in consistently reduced lymphocyte counts (for dexamethasone, \( P = 0.0155 \); for cyclophosphamide, \( P = 0.0115 \); for dexamethasone and cyclophosphamide, \( P = 0.0020 \)) (Fig. 1B). This combination did not reduce the number of peripheral blood neutrophils but did prevent neutrophil counts from increasing late after infection (Fig. 1C). In addition, liver panels demonstrate (based on the lack of a statistically significant increase in measured liver enzymes) that neither dexamethasone nor cyclophosphamide nor the combination caused toxicity in the hamster model compared to no-treatment controls (Fig. 1D to F).

SNV-infected, immunosuppressed hamsters exhibited clinical signs and mortality similar to what is exhibited by ANDV-infected hamsters. Hamsters in all groups were challenged with 2,000 PFU of SNV i.m. on day 0 (Fig. 2A). A single hamster succumbed on day 5 in the dexamethasone-treated group (suspected to be from complications due to repeated blood draws; no apparent toxicity was measured or observed); the remaining dexamethasone-treated hamsters survived to day 28 (\( P = 0.8900 \)). Hamsters in the cyclophosphamide-treated and dexamethasone/cyclophosphamide-treated groups displayed clinical signs including staggered gait, tachypnea, dyspnea, lethargy, and severe pulmonary edema. Clinical signs of disease in terminal hamsters were most notable within the 24 h preceding death. Clinical signs of disease were not observed in any surviving hamster. SNV was uniformly lethal for hamsters in the dexamethasone/cyclophosphamide-
treated group (P = 0.0001), resulting in a mean time to death of 13 days and a range of 10 to 14 days postchallenge. Half of the hamsters receiving cyclophosphamide alone developed HPS approximately 16 days postchallenge, with a range of 12 to 19 days postchallenge (P = 0.2034). Of note in the cyclophosphamide-treated group, two hamsters developed neurological symptoms, including holding their head to one side and circling inside the pan repeatedly. As previously reported, immunocompetent hamsters infected with SNV developed no clinical signs and survived to the end of the study (7). Serum from all surviving hamsters on day 28 was subjected to an N-ELISA to confirm infection (Fig. 2B). Survivors in the dexamethasone-treated and no-treatment groups all had ELISA titers of \( 3 \log_{10} \), confirming a productive infection and humoral response. Not surprisingly, the four surviving hamsters in the cyclophosphamide-treated group were unable to mount a robust immune response and had ELISA titers that were either at or below the level of detection for the assay (P = 0.0001).

**Immunosuppression of SNV-infected hamsters allows detection of viremia.** We have previously reported that SNV-infected hamsters had undetectable levels of infectious virus in both the serum and whole blood and viral genome was not detectable in peripheral blood mononuclear cells (PBMCs) (12). In the current study, serum and lung tissue were collected from hamsters 10 days postinfection and evaluated for the presence of viral genome and infectious virus (Fig. 3). Increased levels of viral genome were detectable in the serum of immunosuppressed hamsters compared to those in the serum of immunocompetent hamsters (P = 0.0867) (Fig. 3A), and levels were highest, by approximately 2-fold, in hamsters treated with both dexamethasone and cyclophosphamide. Similarly, approximately 4-fold more infectious virus was detected in the serum of cyclophosphamide- and dexamethasone/cyclophosphamide-treated hamsters than in that of hamsters treated with dexamethasone and untreated hamsters (P = 0.0003) (Fig. 3B). In the lung, slightly higher, although not significantly different, levels of viral genome were detected in immunosuppressed hamsters than in immunocompetent hamsters, with SNV genome levels ranging from \( 10^5 \) to \( 10^6 \) (P = 0.0721) (Fig. 3C). However, there was a statistically significant increase in infectious virus in the cyclophosphamide and dexamethasone/cyclophosphamide hamster groups compared to untreated controls (P < 0.0001) (Fig. 3D).

**SNV infection of immunosuppressed hamsters results in marked pathology and widespread dissemination of virus in lung tissues.** Ten days postinfection, three hamsters per treatment group were euthanized and lung tissues were collected for histologic analysis and immunohistochemistry. H&E staining of lung sections showed minimal differences in dexamethasone-treated (Fig. 4A) and untreated (Fig. 4D) hamsters compared to uninfected controls (Fig. 4E). However, cyclophosphamide-treated (Fig. 4B) and dexamethasone/cyclophosphamide-treated (Fig. 4C) hamster lungs exhibited interstitial inflammation and vascular leakage consistent with changes normally seen in hamsters infected with ANDV (Fig. 4F). Immunohistochemistry performed on lung sections from each of the treatment groups showed widespread immunoreactivity in the alveolar septa and endothelial cells of larger vessels, with the most prominent staining noted in lung sections from the dexamethasone/cyclophosphamide treatment group (Fig. 5A to C). Hamsters in the untreated, SNV-infected group showed scattered, minimal immunoreactivity of infected endothelial cells, a finding which is consistent with previous results (Fig. 5D) (12).

**SNV isolated from immunosuppressed hamsters remains nonpathogenic in immunocompetent hamsters.** Despite the fact that repeated serial passaging of SNV through hamsters does not increase its pathogenicity in immunocompetent hamsters (28), we were interested in determining if passaging SNV through an
immunocompromised hamster can alter the virus to the point of creating disease in an immunocompetent host. To this end, 2,000 PFU of SNV isolated from the pleural effluent of a dexamethasone/cyclophosphamide-treated, SNV-infected hamster was used to infect immunocompetent hamsters. These hamsters all survived to day 28 with no signs of disease, but all developed antibodies to the SNV challenge as measured by ELISA, indicating productive but asymptomatic infection (data not shown).

Passive transfer of α-SNV antibodies demonstrates viral specificity in the development of HPS. To confirm that SNV was the etiologic agent responsible for disease in immunosuppressed hamsters, we showed that SNV neutralizing antibodies can prevent or delay disease in these animals. Two groups of 8 hamsters each were immunosuppressed with dexamethasone and cyclophosphamide in combination according to the strategy detailed in Table 1. All hamsters were then infected with 2,000 PFU of SNV i.m. on day 0. Subsequently, on day 5 postchallenge, one group was administered 12,000 neutralizing antibody units (NAU)/kg of α-SNV antibodies in the form of sera from rabbits vaccinated with an SNV DNA vaccine (Fig. 6A) (29). All hamsters not receiving α-SNV antibodies developed disease and succumbed 12 to 13 days postchallenge. Three hamsters receiving α-SNV antibodies either succumbed or were euthanized on days 17 to 18, with the remaining 5 hamsters surviving to day 28 (P = 0.0001). SNV neutralizing antibodies delayed or prevented lethal disease in hamsters, confirming that HPS disease is specific to SNV infection. Lungs collected on day 28 from surviving hamsters were evaluated for the presence of viral genome by real-time PCR (RT-PCR) (Fig. 6B). Surviving hamsters had comparable levels of viral genome in lung tissue isolated on day 10 postinfection (Fig. 3C).

DISCUSSION
In this report, we describe the first SNV lethal disease small-animal model. To date, evaluations of potential SNV vaccines and immunotherapeutics have involved using either the deer mouse or hamster SNV infection model (29,30). Attempts to develop an SNV disease model have included experimental infections of rodents (i.e., deer mice and hamsters) using serially passaged virus that has resulted in increased viral replication in tissues, including the lung, liver, and spleen, but no signs of disease (28,31). In contrast, the immunosuppression methodology incorporated in this study allowed SNV to replicate and disseminate in the hamster host, creating an HPS disease similar to the disease observed in the ANDV hamster disease model and in human HPS cases. Both SNV and ANDV are New World hantaviruses that cause HPS in humans, and the kinetics and pathology of SNV in an immunocompromised hamster and ANDV in an immunocompetent hamster are remarkably similar. ANDV-infected hamsters will succumb to HPS disease in 10 to 14 days when challenged i.m. with 200 to 2,000 PFU of virus, and this is essentially the same time to death that was observed in the immunocompromised hamsters challenged i.m. with 2,000 PFU of SNV. Viremia can be detected in immunocompromised, SNV-infected hamsters, and these
approximately 106 S-segment RNA copies were detected in lung cells of multiple tissues, approaching 108 S-segment RNA copies in lung spread and prolonged viral dissemination to endothelial cells in hamsters with Syrian hamster-passaged SNV. Results in wide-tissue that were detectable by PCR (28). However, in both cases, the lungs of these animals. This may suggest that certain components of the cellular immune response (limited by cyclophosphamide treatment) regulate pathogenesis following SNV infection.

Cyclophosphamide has been used to create viral disease models in multiple rodent species, including Syrian hamsters (20, 21, 32–37). Cyclophosphamide is an alkylating agent which intercalates into the DNA of actively dividing cells, causing apoptosis resulting in lymphopenia, suppressed B cell activity and activation, suppressed regulatory T cell function, neutropenia, and decreases in macrophage and dendritic cell numbers (14–17, 38, 39). When used in viral infection models, this often leads to increased morbidity and mortality and enhanced viral replication and dissemination. Correspondingly, we did see significant decreases in total WBC counts (Fig. 1) as well as consistently significant decreases in lymphocyte cell numbers and increased titers of infectious virus in the serum and lungs of cyclophosphamide-treated animals. Somewhat surprisingly, the numbers of peripheral blood neutrophils in cyclophosphamide-treated animals were not dramatically different from numbers in untreated animals even though neutrophils, as well as macrophages, are sensitive to apoptosis due to cyclophosphamide (16, 17). However, the activation status of neutrophils, specifically NF-kB activation, can greatly influence neutrophil sensitivity to apoptosis-inducing stimuli (40). This argues that the decrease in WBC counts was a more direct result of decreased lymphocyte numbers (Fig. 1B). Other than a slight increase in neutrophil numbers on day 3 postinfection, treatment of animals with cyclophosphamide did not result in increased numbers of neutrophils compared to untreated animals (Fig. 1C).

Dexamethasone is commonly used to treat or prevent disease in animal models (41–44) and has been used, along with the related glucocorticosteroid methylprednisolone, in attempts to treat human hantavirus disease (45–47). Dexamethasone is a glucocorticoid which can downregulate proinflammatory cytokine expression, induce anti-inflammatory cytokines and proliferation, and induce apoptosis in multiple cell types (48, 49). Dexamethasone has been shown to be effective in downregulating type I interferon responses by regulating STAT1 activation (50); therefore, we hypothesized that treatment of hamsters with dexamethasone would allow SNV to disseminate and cause disease. Indeed, SNV was able to disseminate following dexamethasone treatment, but dexamethasone treatment alone resulted in a lack of viremia detected 10 days postchallenge and was insufficient to allow SNV to cause disease (Fig. 2 and 4). Dexamethasone alone also resulted in smaller decreases in lymphocyte numbers around the time of virus challenge and failed to control neutrophil proliferation on days 5 and 10. Increased neutrophil numbers may be a result of the inability of steroids to block NF-kB-induced neutrophil activation and proliferation resulting from a strong viral stimulus or the ability of glucocorticoids to inhibit neutrophil apoptosis. Several studies have demonstrated that dexamethasone delays or even prevents neutrophil apoptosis, thus prolonging neutrophil survival and functional responsiveness (51–54), due in part to the stabilization of the anti-apoptotic Mcl-1L protein (55). This, along with increased expression of neutrophil survival factors, such as granulocyte colony-stimulating factor (G-CSF) (56) and granulocyte-macrophage colony-stimulating factor (GM-CSF) (57), which are often induced during viral infections, may explain why dexamethasone-treated animals did not exhibit a more significant decrease in neutrophil numbers. Moreover, treatment of neutrophils with dexamethasone does not appear to inhibit their ability to degranulate and release various antimicrobial reactive oxygen species (51, 58). Earlier studies of ANDV infection of ham-

FIG 6 HPS disease is specifically caused by SNV infection in immunosuppressed hamsters. Groups of 8 hamsters were immunosuppressed with the combination of dexamethasone and cyclophosphamide according to Table 1. (A) On day 5, a single group of 8 hamsters was injected with 12,000 NAU/kg α-SNV antibodies. Both groups of hamsters were observed for survival. (B) Lung tissue isolated on day 28 postinfection was evaluated for viral genome by RT-PCR. ***, P value of <0.001 compared to immunosuppressed, infected controls.
sters suggest that hamsters become viremic approximately 6 to 7 days postchallenge (12), and the kinetics of disease are similar between immunocompetent hamsters infected with ANDV and immunocompromised hamsters infected with SNV. Correspondingly, we observed an increase in the number of circulating neutrophils beginning 5 days postchallenge. The increase in neutrophil numbers corresponds with the lower titers of infectious virus found in dexamethasone-treated animals and suggests the possibility that increases in activated neutrophils may protect against hantavirus disease by reducing live virus, possibly via the expression of proinflammatory cytokines, phagocytosis, or activation of endothelial cell responses.

Interestingly, the combination of cyclophosphamide and dexamethasone was uniformly lethal in hamsters infected with SNV whereas cyclophosphamide alone resulted in only 50% mortality and a prolonged disease course. This suggests that the presence of dexamethasone at the time of challenge probably allows the virus a better opportunity to replicate and disseminate by reducing both innately (e.g., skeletal muscle, smooth muscle, and endothelial cells) derived and cellulary (e.g., dendritic cells, NK cells, neutrophils) derived sources of type I interferon. The fact that cyclophosphamide alone still resulted in 50% mortality suggests that cellular sources of type I interferon are important in early clearance of SNV in immunocompetent hamsters.

While liver panels did not show any toxicity in any of the treatment groups in the hamster, demonstrating that the disease is specific to SNV is a key experiment in this study. Using passively transferred α-SNV neutralizing antibodies administered 5 days postinfection, we were able to show protection from lethal disease. This demonstrated that SNV was the etiologic agent that caused the lethal disease in the treated hamsters. This supports the idea that while neutralizing antibodies are not required for protection, they are sufficient to protect (59–65). Furthermore, this experiment demonstrates the practical utility of this new animal model for evaluating candidate medical countermeasures to prevent and treat hantavirus disease.

A key limitation to the use of this model is that immunosuppression precludes studying the contribution of the immune system in protection from hantavirus infection and vaccine efficacy in that dexamethasone and cyclophosphamide inhibit a wide range of immune cell types. This makes it difficult to directly evaluate a potential vaccine since these immune cells will be unable to secrete antibody or proliferate upon recognition of the viral antigen. Nevertheless, it is possible to indirectly evaluate vaccines that produce neutralizing antibodies, as illustrated in the passive transfer experiment shown in Fig. 6. Another limitation of the immunosuppression approach is that targeting immune modulation to enhance or prevent specific components of the immune response to hantavirus disease would not be possible due to this general immunosuppression methodology. However, this model would be invaluable for evaluating candidate antiviral therapies and investigating nonimmune mechanisms of disease pathogenesis. Disease models created by immunosuppressing animals also typically limit the extent to which disease pathogenesis can be studied, especially when disease is hypothesized to be mediated by components of the immune system. An interesting observation from this study is that lymphocyte numbers are significantly reduced in animals treated with the combination of cyclophosphamide and dexamethasone. Much has been made regarding the role of T cells in the pathogenesis of hantavirus disease. Indeed, activated hantavirus-specific T cells are found in human HPS cases (6, 66, 67), and correlations have been drawn between disease severity and T cell numbers (68). We and others have demonstrated that T cells are not required for disease pathogenesis in the hamster model of HPS (69, 70), in part due to the observation that cyclophosphamide treatment of ANDV-infected hamsters significantly reduced the number of T cells in the blood and lungs of hamsters but did not alter the course of disease in hamsters. Similarly, here we demonstrate that the combination of dexamethasone and cyclophosphamide significantly reduces lymphocyte numbers, and only when hamsters are immunocompromised can SNV uniformly cause disease. This argues for a disease mechanism that is independent of T cells.

Animal models of infectious disease are invaluable tools to study pathogenesis and to discover and test candidate medical countermeasures. Here, we have used a strategy whereby an immunologically competent animal (i.e., capable of mounting a normal immune response to a vaccine) can be transiently immunosuppressed to allow SNV to disseminate and cause disease that mimics HPS in humans. Further studies will be needed to elucidate the mechanism underlying the pathophysiology in hamsters and then to determine if the same mechanism accounts for the development of HPS in humans. The dexamethasone/cyclophosphamide approach described here differs from the use of transgenic knockout animals with defective innate and/or adaptive immune systems because the animals have a normal immune system up until the time when immunosuppression is induced. Furthermore, immunosuppression is transient and can be restored upon cessation of treatment. It will be of interest to determine if this same approach can be used to develop disease models for hantaviruses that cause hemorrhagic fever with renal syndrome (e.g., Hantaan virus) or for other viruses within the Bunyaviridae family for which there are no practical disease models. It will also be of interest to investigate whether this dexamethasone/cyclophosphamide approach can be used to develop disease models in other animal species that have useful attributes, such as mice for the abundance of immune reagents or nonhuman primates for their more human-like physiology.

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