Lower Respiratory Tract Infection Induced by a Genetically Modified Picornavirus in Its Natural Murine Host

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Abstract

Infections with the picornavirus, human rhinovirus (HRV), are a major cause of wheezing illnesses and asthma exacerbations. In developing a murine model of picornaviral airway infection, we noted the absence of murine rhinoviruses and that mice are not natural hosts for HRV. The picornavirus, mengovirus, induces lethal systemic infections in its natural murine hosts, but small genetic differences can profoundly affect picornaviral tropism and virulence. We demonstrate that inhalation of a genetically attenuated mengovirus, vMC0, induces lower respiratory tract infections in mice. After intranasal vMC0 inoculation, lung viral titers increased, peaking at 24 h postinoculation with viral shedding persisting for 5 days, whereas HRV-A01a lung viral titers decreased and were undetectable 24 h after intranasal inoculation. Inhalation of vMC0, but not vehicle or UV-inactivated vMC0, induced an acute respiratory illness, with body weight loss and lower airway inflammation, characterized by increased numbers of airway neutrophils and lymphocytes and elevated pulmonary expression of neutrophil chemoattractant CXCR2 ligands (CXCL1, CXCL2, CXCL5) and interleukin-17A. Mice inoculated with vMC0 compared with those inoculated with vehicle or UV-inactivated vMC0, exhibited increased pulmonary expression of interferon (IFN-α, IFN-β, IFN-λ), viral RNA sensors [toll-like receptor (TLR)3, TLR7, nucleotide-binding oligomerization domain containing 2 (NOD2)], and chemokines associated with HRV infection in humans (CXCL10, CCL2). Inhalation of vMC0, but not vehicle or UV-inactivated vMC0, was accompanied by increased airway fluid myeloperoxidase levels, an indicator of neutrophilic airway inflammation. The picornavirus-induced neutrophilic airway inflammation should be useful for defining mechanisms of HRV pathogenesis in humans.

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Introduction

Infections with the picornavirus, human rhinovirus (HRV), are the most frequent cause of the common cold. However, HRV infections, which usually cause self-limiting upper respiratory tract illnesses, are also the leading cause of virus-induced asthma exacerbations [1], and HRV wheezing illnesses in the first few years of life are associated with increased risk for the development of childhood asthma [2–4]. The mechanisms by which a common cold virus can induce asthma exacerbations and contribute to the development of persistent lower airway sequelae in susceptible children remain to be elucidated [5,6].

There is considerable evidence that HRV can infect the lower respiratory tract [7–14] and that HRV infection stimulates the production of proinflammatory chemokines and cytokines by lower airway epithelial cells [15]. Neutrophils are the main first-line inflammatory cells recruited to the airways during HRV infections [16–18], and this neutrophilic inflammatory response has been associated with asthma symptoms and airway dysfunction [19–22]. However, the relationship between neutrophilic airway inflammation and HRV-induced airway disease is still largely undefined. The eventual outcome of HRV infection, a relatively uneventful upper respiratory tract illness versus a more severe lower respiratory tract illness, might be related to the balance between detrimental and beneficial effects of the neutrophilic inflammatory response in the airways, which might be influenced by host, viral, developmental, and environmental factors. The development of useful small animal models of picornavirus-induced neutrophilic airway inflammation could facilitate mechanistic studies to address these issues.

There are no known murine rhinoviruses, which has significantly hampered the investigation of the mechanisms governing the inflammatory responses to HRV infection and the subsequent development of airway sequelae. Experimental models using either minor receptor group HRV in wild-type mice or major receptor group HRV in mice that are transgenic for human intercellular
adhesion molecule-1 (ICAM-1; CD54), the receptor for major group HRV, have been developed recently [23–28]. A limitation of these useful models is that HRV titers exhibit a steep decline after inoculation of normal mice [23–26]. The development of rodent models in which picornavirus replication persists for several days in the airways of unmanipulated hosts, as in experimental and clinical HRV infections, could facilitate investigation of the relationships between viral replication and the development of airway inflammation and dysfunction. For this reason, we have explored the use of a murine picornavirus, which replicates efficiently in its natural hosts, to model picornavirus-induced respiratory infections.

Mice are the natural hosts for mengovirus, a picornavirus whose wild-type form causes infections that are more similar to systemic poliovirus infections than to HRV-induced airway infections [29]. The mengovirus genome has a poly(C) tract in the distal region of its 5' untranslated region, which has been shown to be an important virulence determinant that inhibits host type I interferon (IFN) responses [30–34]. Investigation of a mengovirus mutant, vMC0, in which the poly(C) tract had been deleted showed that vMC0 induces robust type I IFN responses and that vMC0 inoculation by intracerebral or intraperitoneal routes results in self-limited infections rather than the often lethal, systemic infections induced by wild-type mengovirus [30–34]. Wild-type mengovirus efficiently replicates in both epithelial and macrophage lineage cells; however, vMC0, like HRV, replicates well in epithelial cells but poorly in macrophage lineage cells [34,35]. These similarities between vMC0 and HRV led us to hypothesize that vMC0 could produce a respiratory tract infection in rodents akin to HRV infections in humans. Using rats, were able to demonstrate that inoculation of vMC0 by an inhalation route could induce infection of the lower airways and neutrophilic airway inflammation [36]. An effective HRV infection model has not been established in rats. Therefore, vMC0 should be useful for investigating picornavirus-induced airway infections in rats, and studies to explore the effects of vMC0 infection in an established rat model of allergic airway inflammation are underway. However, the much wider range of mutant strains and genetic tools available in the mouse also made it highly desirable to develop a robust murine model of vMC0-induced airway infection. In this paper, we demonstrate that inhalation of the genetically attenuated mengovirus, vMC0, induces an acute lower respiratory tract illness in mice, which is characterized by replication and persistent shedding of virus and neutrophilic airway inflammation with evidence of neutrophil activation and lung injury. This novel murine model of picornavirus-induced lower respiratory tract infection and inflammation should be useful for investigating mechanisms of HRV pathogenesis in humans.

Results

Viral replication and persistent viral shedding in the lungs after inhalation of attenuated mengovirus, vMC0

After an intranasal inoculation of 10⁶ plaque-forming units (PFU) of attenuated mengovirus, vMC0, a median of 5.4×10⁶ PFU were detected in whole lung homogenates from mice at 0.1 h postinoculation, with viral titers remaining relatively stable at 1

![Figure 1. Differences between attenuated mengovirus and HRV in the kinetics of lung viral titers.](http://www.plosone.org/article/fetchObject/v1/10.1371/journal.pone.0032061.g001)
and 3 h postinoculation (Fig. 1A). At 6 and 24 h postinoculation, there was a marked increase in viral titers to median levels of $5.3 \times 10^5$ and $6.9 \times 10^6$ PFU, respectively, which is indicative of substantial viral replication in the lung (Fig. 1A). Infectious vMC0 continued to be shed in the lungs for at least 5 days postinoculation (Fig. 1B). In contrast, after intranasal inoculation of $5 \times 10^6$ PFU of HRV-A01a, a minor receptor group HRV, a median of $3.4 \times 10^3$ PFU were detected in whole lung homogenates at 0.1 h postinoculation, and viral titers declined until they were undetectable at 24 h postinoculation (Fig. 1C). No virus was detected in vehicle-inoculated mice. These data show that inhalation of vMC0, compared with that of HRV-A01a, resulted in more robust viral replication and greater persistence of viral shedding in the lungs.

Type I and III IFN production in the lungs after inhalation of vMC0

The induction of type I and III IFN is a key component of the host response to HRV infection [37–39] and an indicator of viral replication. Intranasal inoculation with vMC0, but not UV-inactivated vMC0, resulted in significant increases in bronchoalveolar lavage (BAL) fluid levels of the type I IFNs, IFN-α and IFN-β, and type III IFN, IFN-λ, compared with those in vehicle-inoculated mice on days 1 and 2 postinoculation (Fig. 2). Thus, vMC0-induced IFN production required replication competent virus.

Pulmonary expression of viral RNA sensors after inhalation of vMC0

Pattern recognition receptors, toll-like receptor (TLR)3 and TLR7, have been implicated as viral RNA sensors in the host response to HRV infection [40,41]. NOD2 (nucleotide-binding oligomerization domain containing 2) is a pattern recognition receptor that detects bacterial peptidoglycan, but which has been shown recently to also function as a viral RNA sensor [42]. Inhalation of vMC0, but not UV-inactivated vMC0, resulted in significant increases in TLR3, TLR7, and NOD2 mRNA levels in the lungs on day 1 postinoculation compared with mRNA levels in vehicle-inoculated mice, indicating the stimulation of viral RNA sensor pathways (Fig. 3A–C). Pretreatment of mice with an anti-neutrophil monoclonal antibody (mAb) did not result in reduced pulmonary expression of TLR3, TLR7, or NOD2 mRNA in vMC0-inoculated mice on day 1 postinoculation compared with vMC0-inoculated mice that had been pretreated with a control mAb (Fig. 3D–F); indeed, a significant increase in TLR7 mRNA expression was observed in the anti-neutrophil mAb-treated group (Fig. 3E). Thus, the observed increases in TLR3, TLR7, and NOD2 mRNA levels after inhalation of vMC0 were apparently due, at least in part, to increased expression in resident lung cells rather than being solely due to the recruitment of neutrophils that might express these receptors. In vehicle-inoculated mice, pretreatment with the anti-neutrophil mAb resulted in a significant decrease in TLR3 mRNA levels compared with pretreatment with the control mAb (Fig. 3D).

Body weight loss after inhalation of vMC0

Body weight reduction is a sensitive measure of viral respiratory illness in rodents [43]. Intranasal inoculation of vMC0 led to a significant loss of body weight on days 2 and 3 postinoculation compared with inoculation with either vehicle or UV-inactivated vMC0, which showed that the mice were experiencing a viral illness (Fig. 4). Mice inoculated with vMC0 also exhibited significant body weight loss compared with those inoculated with vehicle, but not UV-inactivated vMC0, on day 1 postinoculation and those inoculated with UV-inactivated vMC0, but not vehicle, on day 5 postinoculation (Fig. 4). Inoculation with UV-inactivated
vMC0 resulted in no significant body weight loss compared with inoculation with vehicle, indicating a requirement for replication-competent virus for the development of illness (Fig. 4).

**Inflammation in the lower airways after inhalation of vMC0**

Intranasal inoculation of vMC0 induced the recruitment of inflammatory cells into the lower airways. Total numbers of BAL cells were significantly elevated on days 1, 3, and 5 postinoculation in mice inoculated with vMC0 compared with those inoculated with UV-inactivated vMC0 or vehicle (n = 6 mice per group). Levels of (A, D) TLR3, (B, E) TLR7, and (C, F) NOD2 mRNA in lungs on day 1 postinoculation were determined by real-time quantitative RT-PCR and normalized to β-actin mRNA levels. * P < 0.01 (vMC0 vs. vehicle or UV-inactivated vMC0), ** P < 0.05 (anti-neutrophil mAb/Vehicle vs. control mAb/Vehicle), † P = 0.01 (anti-neutrophil mAb/vMC0 vs. control mAb/vMC0).

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with vehicle or UV-inactivated vMC0 (Fig. 7C and D). CXCR2 is presented as box plots. *

Elevated on days 1, 2, 3, and 5 postinoculation in mice inoculated with vMC0 compared with those inoculated with UV-inactivated vMC0, demonstrating the availability of these chemokines to mediate neutrophil recruitment.

Expression of CXCL10 and CCL2 in the lower airways after inhalation of vMC0

Because HRV infection induces high levels of CXCL10 and CCL2 expression [47–50], and CCL2 indirectly contributes to neutrophil recruitment to the lungs [51], the expression of CXCL10 and CCL2 was measured. Levels of CXCL10 and CCL2 mRNA in the lungs on day 1 postinoculation and BAL fluid levels of CCL2 on day 2 postinoculation were significantly increased in vMC0-inoculated mice compared with vehicle- or UV-inactivated vMC0-inoculated mice, indicating that the picornavirus-induced expression of these chemokines in the lower airways could be studied in this mouse model (Fig. 9).

Mucin expression in the lungs after inhalation of vMC0

HRV infection is associated with increased mucin expression [52,53]. Therefore, mucin mRNA expression was measured. MUC5B, but not MUC5AC, mRNA levels were significantly elevated on day 1 postinoculation in the lungs of mice inoculated with vMC0 as compared with vehicle-inoculated mice (Figure 10). However, periodic acid-Schiff staining of lung sections from vMC0-infected mice showed only sporadic staining for mucus-producing cells in the airways (not shown).

Lung edema after inhalation of vMC0

To examine whether the lower airway inflammation induced by vMC0 was associated with lung edema, wet/dry lung weight ratios were measured. Wet/dry lung weight ratios were significantly elevated on day 2 postinoculation in the lungs of mice inoculated with vMC0 as compared with vehicle- and UV-vMC0-inoculated mice, indicating the presence of acute lung injury (Fig. 11).

Effect of inhalation of vMC0 on pulmonary physiology

To examine whether infection of the lower airways with vMC0 induced changes in pulmonary physiology, mice received intranasal inoculations of vMC0, UV-inactivated vMC0, or vehicle, and pulmonary function was measured on day 2 postinoculation. No significant differences were observed among the groups with regard to respiratory system resistance (Rrs) or the input impedance variables, Newtonian resistance (Rn), tissue viscous (G), and elastance (H), either at baseline or in response to methacholine challenge (Fig. 12 and data not shown), indicating a lack of viral effects on pulmonary physiology and AHR.

Discussion

The establishment of useful and varied small animal models to study HRV pathogenesis continues to be an important goal. Diverse animal models can facilitate different types of mechanistic studies and the development of potential new therapeutic strategies [54]. Recently, murine experimental models of HRV infection have been established [23–28]. The development of these models represents a significant advance in the study of HRV-induced airway inflammation. However, these models have an important limitation. Although there is evidence of HRV replication in mice, HRV titers decline quickly [23–26]. Therefore, the development of small animal models with more robust viral replication could permit mechanistic studies of the role viral replication plays in HRV pathogenesis. To model the role of viral replication in HRV-induced lower respiratory tract...
infections of relevance to the development and exacerbation of asthma in humans, it would be desirable to develop small animal models where picornaviral replication occurs in the lower airways and viral titers increase, peak, and then wane while persisting for several days.

HRV is a picornavirus family member and viral replication is often most efficient in the natural hosts of a virus. Therefore, we have focused on developing small animal models using a picornavirus, an attenuated form of mengovirus, whose natural hosts are rodents. Previously, we described a rat model in which the attenuated mengovirus, vMC0, caused a respiratory infection in rats with several days of viral shedding accompanied by a neutrophilic lower airway inflammatory response [36]. In the mouse model described here, we observed an approximately 3 log increase in vMC0 titers in the lungs from 0.1 h to 24 h postinoculation, with viral shedding persisting for at least 5 days after inoculation. The intranasal inoculum of 10^6 PFU of vMC0, an equivalent amount of UV-inactivated vMC0, or vehicle (n = 6 mice per group). Data are presented as box plots. † P < 0.05, * P < 0.01, ** P < 0.001, *** P < 0.0001 (vMC0 vs. vehicle or UV-inactivated vMC0).

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Figure 5. Recruitment of inflammatory cells to the lungs in response to inhalation of attenuated mengovirus. Numbers of (A) total cells, (B) neutrophils, (C) lymphocytes, and (D) macrophages in the BAL fluid harvested at the indicated times from the lungs of mice inoculated with 10^6 PFU of vMC0, an equivalent amount of UV-inactivated vMC0, or vehicle (n = 6 mice per group). Data are presented as box plots. † P < 0.05, * P < 0.01, ** P < 0.001, *** P < 0.0001 (vMC0 vs. vehicle or UV-inactivated vMC0).

Picornavirus-Induced Airway Infection in Mice

For both of our vMC0 and HRV-A01a inoculations, the amount of infectious virus detected at 0.1 h postinoculation by

For comparison, we inoculated mice with HRV-A01a, a minor group HRV, and found that infectious viral titers in the lungs declined until they were undetectable at 24 h postinoculation. Our HRV-A01 strain b, which showed a steep reduction in HRV titers over time in the lungs of healthy, immunocompetent mice [23–26]. In one study, after intranasal inoculation of 5 × 10^6 TCID_{50} of HRV-A01b, viral titers of approximately 10^5, 10^4, and 10^3 TCID_{50}/ml at 8, 16, and 24 h postinoculation, respectively, were reported [23]. In another study [25], after intranasal inoculation with 5 × 10^7 TCID_{50}, a 10-fold higher inoculum than used previously, lung viral titers were approximately 10^3 and 10^2 TCID_{50}/ml on days 1 and 4 postinoculation, although induction of substantial lung injury by repeated treatment with lipopolysaccharide and elastase before virus inoculation enhanced viral titers [25]. In a third study, lung viral titers were somewhat higher after intranasal inoculation with 4.5 × 10^6 TCID_{50} of HRV-A01b, with about 10^6 PFU/ml at 4 h postinoculation steadily declining to about 10^3 PFU/ml on day 4 postinoculation before becoming undetectable at day 4 postinoculation [26]. Overall, inhalation of HRV by mice led to some degree of viral replication accompanied by a consistent pattern of steadily declining levels of infectious virus in the lungs. In healthy, immunocompetent mice, low levels of infectious HRV shedding could be observed at 24 h postinoculation, and these levels continued to fall until little or no infectious virus was detected by day 4 postinoculation.
inactivated vMC0. Lungs were harvested on day 2 postinoculation.

hallmark of viral respiratory illnesses in rodents. Infection of the

signs of illness such as significant body weight loss, which is a

consistent with outcomes of experimental HRV infections in

methacholine concentration that was a half-log above the upper

subjects, but these small changes were only detectable at a

mRNA expression was not increased as well. In humans, HRV

expression of MUC5B mRNA. It is not clear why MUC5AC

plaque assay was substantially lower than the inoculum dose. This
discrepancy was probably due to deposition of some of the virus in
the nasopharynx and gastrointestinal tract as well as the eclipsing
of the virus in the lung during the early stage of the infection.
Importantly, the starting levels of virus in the lungs were similar for
vMC0 and HRV-A01a, but the viral replication and persistence were
greater with vMC0.

Along with the evidence of notable levels of viral replication and persistent viral shedding in the lower respiratory tract, this mouse model has many features in common with HRV infection in humans. Inhalation of live, but not inactivated vMC0, induced signs of illness such as significant body weight loss, which is a hallmark of viral respiratory illnesses in rodents. Infection of the lower respiratory tract with vMC0 resulted in the stimulation of host antiviral pathways much in the same way as HRV. There was a marked increase in Type I and III IFN proteins in the airway fluids as early as 1 day after virus inoculation, which also provides further evidence of viral replication in the lungs after vMC0 infection. Regulation of type I and III IFN production has been implicated as an important checkpoint in the host response to HRV infection [37–39]. In addition, pulmonary expression of the viral RNA sensors, TLR3, TLR7, and NOD2 were upregulated by vMC0 infection. The viral RNA sensors, TLR3 and TLR7, have been implicated in the host response to HRV infections [26,40,41]. NOD2 is an intriguing pattern recognition receptor because it not only detects bacterial peptidoglycan but can also function as a viral RNA sensor [42]. NOD2 has not been previously implicated in host responses to HRV. However, given its upregulation in the lungs in response to vMC0 infection, it will be of interest to examine its role in HRV infection. Depletion of neutrophils before the virus inoculation did not inhibit the subsequent upregulation of these viral RNA sensors, which was consistent with a vMC0-associated increase in expression in resident lung cells. In all cases, stimulation of these host antiviral pathways required active vMC0.

Recruitment of neutrophils and lymphocytes to the airways are common features of vMC0- and HRV-induced airway infections, and the chemokine milieu that was observed in the lung tissue and airway fluids in the vMC0-infected mice has marked similarities to that observed in HRV infections in humans. Increased pulmonary expression of the neutrophil chemoattractant CXCR2 ligands in response to vMC0 infection is consistent with the increased expression of CXCR2 ligands that is observed in response to HRV infection in humans and in mouse models [23,24,55–58]. We also observed the induction of pulmonary expression of CXCL10 and CCL2 in response to inoculation of vMC0, showing another similarity between this mouse model of picornaviral airway infection and HRV infection. CXCL10 and CCL2 expression have been shown to be markedly upregulated in response to HRV infection [23,24,47–50,59].

There was also evidence of acute airway injury, as shown by the development of lung edema, which might be related to the neutrophilic airway inflammation observed in the lower respiratory tract and the evidence for neutrophil activation, i.e., the release of the neutrophil granule protein MPO in the airway fluids. Although we only observed sporadic staining for mucus-producing cells in the airways, it is possible that mucin gene expression induced by vMC0 infection might have greater biological relevance in the exacerbation of existing airway disease in mice.

Similar to what we reported in our rat model of vMC0-induced airway infection and inflammation [36], inhalation of vMC0 by mice had no significant effect on baseline pulmonary function or AHR to methacholine challenge. As in our previous rat study, experimentally naïve adult animals without existing airway disease were used in these mouse studies. Similar to our rodent models, almost all reports of experimental HRV inoculations of healthy, nonasthmatic, nonallergic human subjects have resulted in no significant postinoculation changes in baseline pulmonary function or AHR [11,60–64]. One study showed small differences in AHR after experimental HRV infection of nonasthmatic, nonallergic subjects, but these small changes were only detectable at a methacholine concentration that was a half-log above the upper end of the typical dose range [65]. However, experimental HRV inoculation was shown to increase AHR in some individuals with asthma and/or allergic rhinitis in several studies [11,22,64,66,67] but not in others [60,61,63,65,68]. Consequently, the lack of AHR changes in healthy adult mice without existing airway disease was consistent with outcomes of experimental HRV infections in
humans with no underlying airway disease, such as asthma or allergic rhinitis. Overall, the absence of pulmonary function changes during vMC0-induced lower respiratory infection in mice without existing airway disease is consistent with observations in experimental HRV infections in healthy human volunteers. The absence of viral effects on AHR is also consistent with one mouse model of HRV infection in which no increase in AHR to methacholine challenge was observed after HRV infection unless the BALB/c mice had underlying experimental allergic airway inflammation [23]. However, in another model, a modest increase in AHR to methacholine challenge was reported after infection of C57BL/6 mice with HRV [24], which might be related to mouse strain differences. The underlying condition of the airways, whether related to genetic, developmental, and/or environmental factors, may have profound effects on host responses to picornaviral airway infection and the outcomes of these infections. In the future, it will be of interest to investigate the influence of age, genetic factors, and other environmental exposures, such as allergens, other respiratory viruses [69], or microbial factors, on host responses to vMC0-induced lower respiratory tract infection.

A potential limitation of using vMC0 in this mouse model of respiratory infection is that, unlike HRV, mengovirus is neurotropic. However, it is important to note that the HRV, as enteroviruses, are closely related to poliovirus, which is also neurotropic. Inhalation of the attenuated mutant of mengovirus, vMC0, induced a self-limited respiratory infection in rodents, demonstrating the plasticity of vMC0 with regard to its tissue tropism. After intranasal inoculation with vMC0, there was no evidence of systemic infection in homogenates of nonrespiratory organs, as tested by plaque assay (not shown), which was consistent with our previous work in rats [36]. We chose to use plaque assays for the detection of virus because our primary interest was in measuring titers of infectious virus. We cannot rule out the possibility that vMC0 RNA might have been detected in nonrespiratory organs if a highly sensitive PCR-based assay had been used. In fact, HRV viremia has been detected in some human patients by highly sensitive PCR-based assays [70,71].

The picornaviral airway infection models in rodents involve a single viral inoculation, which is comparable to experimental HRV inoculation studies in human subjects. An HRV infection

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**Figure 7. Neutrophilic inflammation in the lungs in response to inhalation of attenuated mengovirus.** Mice received intranasal inoculations of $10^6$ PFU of attenuated mengovirus, vMC0, an equivalent amount of UV-inactivated vMC0, or vehicle. (A) Lung-associated MPO levels on day 1 postinoculation. MPO levels in lung tissue homogenates from mice inoculated with vehicle or vMC0 were determined by ELISA and normalized to total protein levels (n = 4 mice per group). (B) MPO release into airway fluids. BAL fluid was harvested on the indicated days, and MPO levels were determined by ELISA (n = 6 mice per group). Data are the total amount of MPO recovered (ELISA values were multiplied by the BAL fluid volume). MPO levels below the limit of detection were assigned a value of 1 ng for graphing purposes. (C) CXCR2 and (D) IL-17A expression in the lungs on day 1 postinoculation. Levels of CXCR2 and IL-17A mRNA were determined by real-time quantitative RT-PCR and normalized to β-actin mRNA levels (n = 6 mice per group). Data are presented as box plots. * P<0.05 (vMC0 vs. vehicle). ** P<0.01 (vMC0 vs. vehicle or UV-inactivated vMC0). doi:10.1371/journal.pone.0032061.g007
can trigger a wheezing illness or asthma exacerbation in children and adults [1], and experiencing an HRV wheezing illness early in life is associated with an increased risk for the subsequent development of childhood asthma [2–4]. Thus, these rodent models should be useful for relevant mechanistic studies. Some children appear to exhibit increased susceptibility to recurrent HRV infections, presumably caused by exposure to differing serotypes [72,73], but the precise role of recurrent HRV infections

**Figure 8. Increased pulmonary expression of CXCR2 ligands in response to inhalation of attenuated mengovirus.** Mice received intranasal inoculations of 10^6 PFU of attenuated mengovirus, vMCo, an equivalent amount of UV-inactivated vMCo, or vehicle (n = 6 mice per group). Levels of (A) CXCL1 (KC), (B) CXCL2 (MIP-2), and (C) CXCL5 (LIX) mRNA in lungs on day 1 postinoculation were determined by real-time quantitative RT-PCR and normalized to β-actin mRNA levels. Levels of (D) CXCL1, (E) CXCL2, and (F) CXCL5 protein in BAL fluid on day 2 postinoculation were determined by ELISA. Data are the total amount of chemokine recovered (ELISA values were multiplied by the BAL fluid volume). CXCL2 protein levels below the limit of detection (dotted line) were assigned a value of 4 pg for graphing purposes. Data are presented as box plots. * P<0.01, ** P<0.05 (vMCo vs. vehicle or UV-inactivated vMCo).

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in asthma pathogenesis remains to be elucidated. Recurrent picornaviral airway infections would be more challenging to study in animal models because of the limited serotypic diversity of viruses that have been experimentally tested in mice, but one possibility would involve sequential inoculation of \( vMC_0 \) and HRV because there should be limited immune crossreactivity between these picornaviruses.

A comparison between our mengovirus infection model and the HRV infection model in mice \([23–28]\) shows that they both induce a neutrophilic inflammatory response in the lower respiratory tract, which is consistent with the inflammatory responses induced by HRV infections in humans. By mimicking important aspects of natural HRV infections in humans, both models should facilitate mechanistic studies that would be difficult to perform in human subjects. The HRV infection model in mice allows for the effects of the actual human virus to be studied directly, but HRV is not a natural mouse pathogen and does not replicate efficiently in mice. An important distinction is that mengovirus, a natural mouse pathogen, replicates more efficiently in mice than does HRV. Our data show that after inhalation of \( vMC_0 \), viral titers in the lung increase, peak, and then wane while persisting for at least 5 days. Overall, both of these models have strengths and limitations, and the relative usefulness of each model will depend on the specific research questions that are being addressed.

In conclusion, we have developed a robust model of picornavirus-induced airway infection and inflammation in mice. One of this model's key strengths is that it employs a natural pathogen for rodents, mengovirus, which yields viral replication kinetics and magnitudes closer to natural viral respiratory infections. HRV infection models in mice are proving to be useful for the study of HRV-induced airway inflammation but have a weaker component of viral replication. The mengovirus airway infection model should complement these existing models, and should be especially useful for studies where viral replication is an important outcome.

**Materials and Methods**

**Ethics statement**

The mice were housed and all experimental procedures were performed in an American Association for Accreditation of Laboratory Animal Care-accredited laboratory animal facility at the University of Wisconsin School of Medicine and Public Health. The study was approved by the University of Wisconsin School of Medicine and Public Health Animal Care and Use Committee (protocol number M00582) and conformed to the Guide for the Care and Use of Laboratory Animals.

**Animals**

Female BALB/cAnNCr mice were purchased from the National Cancer Institute Animal Production Program (Frederick, MD) and used at 6–10 weeks of age for inoculation studies. The mice were housed in microisolator cages within HEPA-filtered isolation cubicles (Britz & Company, Wheatland, WY).

**Virus**

Recombinant \( vMC_0 \) has been described previously \([31]\). To prepare stocks of \( vMC_0 \), the plasmid containing the entire \( vMC_0 \) cDNA was linearized and transcribed \textit{in vitro} with T7 RNA polymerase. The viral RNA was transfected into HeLa cells, and plaques were isolated and eluted overnight. The viral plaque eluate was used to inoculate fresh plates of HeLa cells which were incubated for 24 h at 37°C. The plates were then frozen, thawed,
and scraped. The lysate underwent two additional freeze-thaw cycles before centrifugation to remove cellular debris. This lysate was used to inoculate HeLa cells (multiplicity of infection = 1) grown in suspension to a density of $4 \times 10^8$ cells/ml. Flasks were incubated at 37°C for 9 h in a shaking water bath and then frozen at $-80^\circ$C. Viral stocks were concentrated and purified by centrifugation through a 30% sucrose cushion. Titers were determined by duplicate plating on HeLa cell monolayers [74].

UV-inactivated vMC0 stocks were prepared by exposing the virus to a germicidal UV lamp for 30 min. Plaque assays using HeLa cells were employed to verify that active virus was undetectable [<10 PFU/ml] in the UV-inactivated preparations. Purified HRV-A, genotype 1 (A01a) was provided by Dr. Wai-Ming Lee (University of Wisconsin-Madison, Madison, WI).

**Virus inoculation and measurement of viral titers**

Mice were inoculated intranasally with vehicle, $10^6$ PFU of vMC0, $10^6$ PFU equivalents of UV-inactivated vMC0, or $5 \times 10^6$ PFU of HRV-A01a under isoflurane anesthesia. For virus titration, lungs were removed from the chest cavity aseptically and weighed. The lungs were homogenized with an automated tissue homogenizer in an appropriate volume of phosphate-buffered saline (PBS) to yield a 10% w/v homogenate. Cell debris were removed by centrifugation, and the supernates were titered for virus by duplicate plating on HeLa cell monolayers [74].

**Depletion of neutrophils**

Purified anti-neutrophil mAb 1A8 (mouse Gr-1/Ly-6G-specific) and isotype-matched control mAb 2A3 (rat IgG2a; <0.18 and <0.63 endotoxin units/mg by Limulus amebocyte lysate test, respectively) were purchased from Bio X Cell (West Lebanon, NH). Anti-neutrophil mAb was administered to each mouse by both intraperitoneal and intranasal routes, which has been

**Figure 10. Mucin expression in the lungs in response to inhalation of attenuated mengovirus.** Mice received intranasal inoculations of $10^6$ PFU of attenuated mengovirus, vMC0, or vehicle (n = 5 mice per group). Levels of (A) MUC5B and (B) MUC5AC mRNA in lungs on day 1 postinoculation were determined by real-time quantitative RT-PCR and normalized to β-actin mRNA levels. * P < 0.01 (vMC0 vs. vehicle).

**Figure 11. Lung edema in response to inhalation of attenuated mengovirus.** Wet-dry lung weight ratios were measured for lungs harvested on day 2 after intranasal inoculation with $10^6$ PFU of attenuated mengovirus, vMC0, an equivalent amount of UV-inactivated vMC0, or vehicle (n = 9 mice per group). Data are presented as box plots; whiskers indicate the 10th and 90th percentiles. * P < 0.01 (vMC0 vs. vehicle or UV-inactivated vMC0).

**Figure 12. Effect of inhalation of attenuated mengovirus on pulmonary physiology.** Mice received intranasal inoculations of vehicle, $10^6$ PFU of attenuated mengovirus, vMC0 (n = 6–7 mice per group), or an equivalent amount of UV-inactivated vMC0 (n = 4 mice), and on day 2 postinoculation, pulmonary physiology measurements were obtained after exposure to aerosols of normal saline followed by escalating concentrations of methacholine. Values for respiratory system resistance (Rrs) are presented as the group means ± the standard error. There were no significant differences among the groups.
previously shown to effectively deplete neutrophils in the respiratory tract [75]. However, we used mAb 1A8 to deplete neutrophils rather than mAb RB6-8C5 because of its greater selectivity for neutrophils [76]. 1A8 was effective in depleting neutrophils, as a single intraperitoneal injection of 1A8 (500 μg) the day before virus inoculation reduced the day 1 postinoculation vMC0-induced increase in lung MPO protein levels by 81% in comparison with mice receiving 2A3 (500 μg) (n=4 mice per group). One day before intranasal inoculation with either vehicle or 10⁶ PFU of vMC0, mice received intraperitoneal injections of 500 μg and intranasal instillations (under isoflurane anesthesia) of 200 μg of either 1A8 or 2A3 in PBS. On the day of inoculation, the mice received an additional intraperitoneal injection of 500 μg of either 1A8 or 2A3 in PBS.

Measurements of pulmonary inflammation

Mice were anesthetized and exsanguinated by severing the dorsal aorta. After opening the chest, BAL was performed. The lungs were lavaged with 800 μl of PBS via a tracheal catheter. The lungs were then lavaged again in the same manner, and the BAL fluid samples were combined. The BAL fluid was centrifuged, and the cell pellet was resuspended in 0.2 ml of PBS. The total number of BAL leukocytes was determined with an automated cell counter (model Z1, Beckman Coulter, Hialeah, FL). Differential cell counts were determined by counting 200 leukocytes on cytopsin slides stained with Protocol® Wright-Giemsa stain according to the manufacturer’s protocol (Fisher Diagnostics, Middletown, VA). BAL fluid was stored at −80°C. For histological assessments of pulmonary inflammation, mouse lungs were filled with 10% formalin, tied off, removed from the chest cavity, and immersed in 10% formalin for 24 h. The tissue was then processed, embedded in paraffin, and cut into 5 μm sections. Sections were stained with a Giemsa stain.

Measurements of pulmonary protein expression

The levels of specific proteins in the BAL fluid or lung homogenates were measured by enzyme-linked immunosorbent assay (ELISA). Mouse CXCL1 (KC, keratinocyte-derived cytokine)-specific, CXCL2 (MIP-2, macrophage inflammatory protein-2)-specific, and IFN-β-specific ELISA kits with sensitivities of 7.8 pg/ml and Mouse IFN-α-specific kits with a sensitivity of 12.5 pg/ml were purchased from Invitrogen (Carlsbad, CA). Mouse CXCL5 (LIX, lipopolysaccharide-induced CXC chemokine)-specific and IFN-λ-specific ELISA kits with sensitivities of 7.8 pg/ml were acquired from R&D Systems (Minneapolis, MN). Mouse CCL2 (MCP-1, monocyte chemoattractant protein-1)-specific ELISA kits with a sensitivity of 7.8 pg/ml were purchased from BD Biosciences (San Diego, CA). Mouse MPO-specific ELISA kits with a sensitivity of 0.4 ng/ml were obtained from Cell Sciences Incorporated (Canton, MA). Total protein levels in lung homogenates were measured by the Coomassie Plus (Bradford) Protein Assay (Pierce, Rockford, IL).

Measurements of pulmonary gene expression

For RNA isolation, mouse lungs were harvested in an RNase-free manner and immediately flash-frozen in liquid nitrogen. Lungs were then powdered under liquid nitrogen with a chilled mortar and pestle. Total RNA was isolated from frozen lung powder with an RNeasy Mini Kit according to the manufacturer’s instructions (Qiagen, Valencia, CA). The total RNA was reverse transcribed into cDNA with SuperScript® III reverse transcriptase following the manufacturer’s instructions (Invitrogen, Carlsbad, CA). The cDNA product was diluted 1:2.5 fold and used as the template for quantitative real-time PCR, which was performed using an ABI 7500 Real Time PCR System (Applied Biosystems, Foster City, CA). TaqMan® primer and probe sets for TLR3, TLR7, NOD2, CXCL1, CXCL2, CXCL5, CXCL10 (IP-10, IFN-gamma inducible protein 10), CCL2, MUC5B, MUC5AC, and β-actin were purchased from Applied Biosystems. All values were normalized to the endogenous control, β-actin. Standard curves were generated by serial dilutions of cDNA from the lungs of a vMC0-inoculated mouse and used to determine relative expression levels.

Measurement of lung edema

To measure lung edema, wet/dry lung weight ratios were determined. Lungs were removed from the chest cavity and immediately weighed to obtain the wet lung weight. All lungs were place in an oven at 65°C for 4 days and then weighed again to determine the dry lung weight.

Measurements of pulmonary physiology

Mice were anesthetized with pentobarbital (Abbott, North Chicago, IL), intubated via tracheostomy, paralyzed with succinylcholine HCl (Sigma, St. Louis, MO), and ventilated mechanically (flexiVent, SCIREQ, Montreal, QC, Canada). Aerosol challenges were delivered by the ventilator via an inline nebulizer (Aeroneb, SCIREQ) with aerosolized normal saline being followed by methacholine HCl (Sigma) solutions in concentrations of 1, 3, 10, and 30 mg/ml. After each aerosol challenge, measurements of pulmonary physiology were performed by the flexiVent system, alternating measures of Rrs with measures of input impedance variables (Rn, G, and H). For each variable, the highest value occurring after each aerosol challenge was recorded as the response, referenced to the value obtained after saline challenge.

Statistical analysis

The numbers of total cells, neutrophils, lymphocytes, and macrophages in the BAL fluid were analyzed by an analysis of variance (general linear model), which was followed by planned pairwise comparisons using Fischer’s least significant difference test. A residual analysis was employed to test the adequacy of the models. Pulmonary physiology data were analyzed by an analysis of covariance, using the values from the methacholine dose 50 mg/ml as the dependent variable and baseline (after saline) values as the covariate to account for differences in baselines. Nonparametric tests were used to analyze all other data. The Kruskal-Wallis test was used for comparisons among three or more groups and was followed by planned pairwise comparisons using the Mann-Whitney test. For comparisons between two groups, the Mann-Whitney test was used. Box plots depict the median and the interquartile range between the 25th and 75th percentile, and whiskers show the 10th and 90th percentiles.

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Author Contributions

Conceived and designed the experiments: LAR RJS SPA YX ACP JEG RLS. Performed the experiments: LAR RJS SPA YX MRH RLS. Contributed reagents/materials/analysis tools: MRH ACP. Wrote the paper: LAR RJS ACP JEG RLS.
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