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Virology

Pathogenic analysis of the pandemic 2009 H1N1 influenza A viruses in ferrets

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ABSTRACT. The pandemic 2009 H1N1 influenza A virus emerged in humans and caused the first influenza pandemic of the 21st century. Mexican isolates, A/Mexico/4108/2009 (H1N1) (Mex4108) and A/Mexico/InDRE4478/2009 (H1N1) (Mex4487) derived from a mild case and from a cluster of severe cases, showed heterogeneity in virulence in a cynomolgus macaque model. To compare the more pathogenic differences, we generated recombinant viruses and compared their virulence in ferrets. Ferrets infected with recombinant Mex4487 displayed a slightly higher rate of viral replication and severe pneumonia in the early stage of infection. In contrast, prolonged lower virus shedding of recombinant Mex4108 than that of recombinant Mex4487 was detected in throat swabs. Thus, Mex4487 induces severe pneumonia in infected individuals, whereas Mex4108 might have wide-spreading potential with mild disease.

KEY WORDS: ferret, pathogenesis, virus shedding

Influenza A virus infections in humans are typically associated with limited seasonal outbreaks of commonly circulating influenza virus strains (seasonal strains). A new virus strain occasionally emerges in humans, resulting in increased morbidity and mortality compared to those of seasonal influenza (pandemic strain) [29]. A novel H1N1 influenza A virus, the pandemic 2009 H1N1 influenza A virus (A(H1N1) pdm2009), caused the first human influenza pandemic of the new millennium [3, 4, 6]. In 2010, WHO announced that A(H1N1) pdm2009 had moved into the post-pandemic period [27, 28]. A(H1N1) pdm2009 has now replaced the classical seasonal H1N1 strains and is circulating globally as a current seasonal strain with a case fatality rate similar to that of classical seasonal influenza.

Human pandemic A(H1N1) pdm2009 infections appeared to be mild in general, and some infected individuals presented with symptoms atypical for seasonal influenza; however, severe illness was also reported, particularly in young, previously healthy individuals [2, 3, 21]. Several early isolates also caused severe diseases in experimentally infected animals, and pathogenicity factors were analyzed [9, 13, 16]. Interestingly, many severe cases of seasonal A(H1N1) pdm2009 were reported in Mexico during the 2013–2014 influenza season [14]. There is the possibility that antigenic change of HA is a reason for middle-aged adults being highly susceptible to seasonal A(H1N1) pdm2009 in the 2013–2014 influenza season [12]. Thus, several factors associated in virulence and human adaptation factors of viruses have identified, and it is still important to accumulate knowledge of the pathogenic potential and analyze the potential virulence factors of these influenza viruses.

In our previous study, infection of cynomolgus macaques with two genetically similar but clinically distinct human A(H1N1) pdm2009 strains, A/Mexico/4108/2009 (Mex4108) and A/Mexico/InDRE4487/2009 (Mex4487), isolated during the early phase of the pandemic [23], resulted in higher pathogenic potential of Mex4487. To investigate the potential virulence of Mexican isolates, we generated recombinant viruses between the two Mexican A(H1N1) pdm2009 isolates and evaluated their pathogenicity in the ferret model, a widely used and well-established model for studying both the pathogenicity and transmissibility of human influenza viruses [1, 26].
MATERIALS AND METHODS

Cells
Madin-Darby canine kidney (MDCK) cells were maintained in Eagle’s minimum essential medium (MEM) supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine and penicillin/streptomycin (Pen/St). Human lung carcinoma (A549) and human embryonic kidney 293T cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FBS, 2 mM L-glutamine and Pen/St.

Viruses
Influenza viruses, Mex4108 (kindly provided by the Centers for Disease Control and Prevention, Atlanta, GA, U.S.A.) and Mex4487 (kindly provided by Public Health Agency of Canada, Winnipeg, MB, Canada), as well as the recombinant viruses, were propagated in MDCK cells with MEM containing 2% FBS and 0.35 \( \mu \)g/ml of TPCK (tolylsulfonyl phenylalanyl chloromethyl ketone)-trypsin. Virus infectivity titers were determined using the 50% tissue culture infectious dose (TCID50) assay. For this assay, 10-fold dilutions of supernatants were used to infect MDCK cells. The virus-induced cytopathogenic effect (CPE) was scored at 3 days post-infection (dpi).

Generation of recombinant viruses
Genomic RNA of Mex4108 and Mex4487 was extracted from virus stocks and used to amplify the eight gene segments by reverse transcription-polymerase chain reaction (RT-PCR). Each PCR product was individually cloned into a polI-promoter plasmid (ppolI) [18]. The open reading frames coding for components of the influenza virus RNP complex, polymerase basic protein 2 (PB2), polymerase basic protein 1 (PB1), polymerase acidic protein (PA) and nucleoprotein (NP), were cloned into the expression plasmid pCAGGs (helper plasmids). All plasmids were sequence confirmed prior to use. To generate recombinant viruses, different combinations of the eight ppolI plasmids from either Mex4108 or Mex4487 were transfected together with the four helper plasmids into 293T cells. After 30 hr of incubation, transfection supernatants were removed, and OPTI-MEM with TPCK trypsin was added. At 48 hr post-transfection (hpt), the supernatant was collected, and the rescued virus was subsequently propagated in MDCK cells. All rescued recombinant viruses were sequence confirmed, and virus titers were determined by a standard TCID50 assay in MDCK cells (Table 1). Recombinant viruses, rgMex4108 and rgMex4487, rescued with titers similar to those of the original isolates.

In vitro growth kinetics of recombinant viruses
Confluent monolayers of A549 and MDCK cells were inoculated with rgMex4108, rgMex4487, rgM4108/M4487-HA.NP.M, rgM4108/M4487-PA.PB2, rgM4108/M4487-PB2 or rgM4108/M4487-PA at a multiplicity of infection (MOI) of 0.1 (A549 cells) or 0.001 (MDCK cells). Virus was allowed to adsorb for 1 hr, then unbound viruses were washed away, and DMEM or MEM with TPCK-trypsin was added. At determined time points, the supernatants were collected from 3 wells per virus. Virus titers were determined as TCID50 on MDCK cells.

Animal study
Groups of 12 ferrets (females, 4–12 months, weight range: 630 to 990 g) were inoculated with rgMex4487 or rgMex4108 (10\(^6\) TCID50/ferret) via the intranasal or intratracheal route, and the animals were monitored daily for body weight and signs of disease for 14 days. Four animals from each group were euthanized at 3 or 6 dpi, and tissue samples were collected for virology and histopathological evaluation. Tissues were placed in cassettes and fixed in 10% Neutral Buffered Formalin x2 changes, for a minimum of 7 days. Cassettes were processed with a Sakura Tissue-Tek VIP-6, on a 12 hr automated schedule, using a graded series of ethanol, xylene and Ultraffin-X paraffin. Embedded tissues are sectioned at 5 \( \mu \)m and dried overnight at 42°C prior to staining. Tissues were stained by hematoxylin and eosin (H&E) stain and scored as: 0=no, 1=minimal, 2=mild, 3=moderate, 4=marked and 5=severe lesion. Infectivity of the virus was determined as TCID50. All animal experiments were approved by the Institutional Animal Care and Use Committee of Rocky Mountain Laboratories and performed following the guidelines of the Association for Assessment and Accreditation of Laboratory Animal Care, International (AAALAC) by certified staff in an

Table 1. Recombinant viruses generated by a reverse genetics system

<table>
<thead>
<tr>
<th>Viruses</th>
<th>PB2</th>
<th>PB1</th>
<th>PA</th>
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<td>M4108/M4487-PB2.PA</td>
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<td>M4108/M4487-PA</td>
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White symbols: Mex4487-derived genes, black symbols: Mex4108-derived genes, striped symbols: common sequence of both Mex4487 and Mex4108, \(* \log \text{TCID50/ml} \)
**RESULTS**

**Growth kinetics of recombinant viruses**

A comparison of the two Mexican isolates, rgMex4108 and rgMex4487, revealed a difference in seven amino acids affecting five proteins [PB2 (amino acid position (aa) 82) and PA (aa275, aa581), HA (aa444), NP (aa100, aa373) and M2 protein (aa82)] [23]. We generated rgMex4108, rgMex4487 and other 4 recombinant viruses of which two were single-gene replacements (PB2, PA), a dual-gene replacement of PB2 and PA and a triple-gene replacement of HA, NP and M. The titers of the all rescued recombinant viruses ranged from $10^{6.0}$ to $10^{7.3}$ TCID50/ml (Table 1). *In vitro* growth kinetics was performed in A549 cells infected with a MOI of 0.1. As shown in Fig. 1A, rgMex4487 showed approximately 1 log higher replication compared to rgMex4108 at 12 h post-inoculation (hpi). Mex4108 backbone viruses of which PA and/or PB2 genes are from Mex4487 (white symbols) tended to higher replication than rgMex4108 or rgM4108/M4487-HA.NP.M (black symbols), however, no significant difference was observed. We also evaluated

AAALAC-approved facility.

**Biosafety**

All infectious *in vitro* and *in vivo* studies were performed in high biocontainment at the Integrated Research Facility (IRF) of Rocky Mountain Laboratories (RML), Division of Intramural Research (DIR), National Institute of Allergy and Infectious Diseases (NIAID), National Institutes of Health (NIH). Sample inactivation and removal from the containment facility was performed according to standard operating protocols approved by the local Institutional Biosafety Committee.

**Statistical analysis**

Statistical analyses were performed using a two-tailed Student’s *t*-test.
the growth kinetics in MDCK cells that was used for virus propagation. All recombinant viruses replicated similarly over a time course of 72 hr with no significant differences in titers and reached highest virus titer at 48 hpi as similar in Table 1 (Fig. 1B).

Pathogenicity of recombinant viruses in ferrets
Ferrets were inoculated with rgMex4487 or rgMex4108 via the intranasal (A, B) or intratracheal (C, D) route. Both groups developed only mild clinical symptoms, and all of the ferrets survived except for two ferrets inoculated with rgMex4108 via the intratracheal route. Those two ferrets died at 2 and 9 dpi with suspected secondary bacterial infection (Supplemental Fig. 1).

By the intranasal inoculation route, no weight loss was induced in either of the groups (Fig. 2A). Virus shedding in nasal and throat swabs was monitored every other day until 14 dpi. rgMex4487 reached the highest titer of 5 × 10^5 TCID50/ml at 1 dpi from throat swabs, and then, the virus titer decreased (Fig. 2B). In contrast, the titer of rgMex4108 in throat swabs remained at 10^4 TCID50/ml until 5 dpi. The virus titers in nasal swab were similar in the two groups. Ferrets inoculated with viruses via the intratracheal route exhibited mild weight loss until 7 dpi in both groups and subsequently recovered (Fig. 2C). As was observed in the groups with intranasal inoculation, the highest virus titer of rgMex4487 was detected at 1 dpi from throat swabs (Fig. 2D). The virus titer of rgMex4108 in throat swabs increased and reached 10^5 TCID50 at 5 dpi, but no significant difference was observed in all time points. The virus titers in nasal swab were similar in the two groups.

Histopathological differences of recombinant viruses
We next compared the viral replication and histopathological changes in respiratory tissues. Four animals from each group were euthanized at 3 or 6 dpi, and tissue samples were collected. By the intranasal inoculation, the highest rate of virus replication
Fig. 3. Comparison of pathological differences in ferrets. Ferrets were infected with a dose of $10^6$ TCID50 of either rgMex4108 or rgMex4487 via the intranasal (A, C, E) or intratracheal (B, D, F) route. Four ferrets were euthanized at 3 or 6 dpi. (A, B) Virus titers of respiratory tissues are shown as TCID50 with S.D. (C, D) Histopathological data are shown as the average of pathological scores with S.D. *$P<0.05$. (E, F) Histopathological sections of lung tissues were stained with H&E stain, original magnification $\times 100$. Black symbols indicate Mex4108, and white symbols indicate Mex4487.
was detected in all respiratory tissues at 3 dpi (Fig. 3A). Although virus titers of rgMex4487 were slightly higher than those of rgMex4108 at 3 dpi in most of the respiratory tissues tested, no significant differences were observed. Virus replication decreased by 6 dpi in all respiratory tissues. Ferrets inoculated with rgMex4487 had developed severe inflammation and pneumonia on day 3 (Fig. 3C and 3E). Mild inflammation and pneumonia were still observed in all respiratory tissues at 6 dpi, even though the virus replication on day 6 was lower than on day 3. Ferrets inoculated with viruses via the intratracheal route showed more severe pneumonia at 3 dpi compared with ferrets inoculated with viruses via the intranasal route (Fig. 3D and 3F). The high rate of virus replication was detected in respiratory tissues at 3 dpi and continued through 6 dpi (Fig. 3B). Although moderate pneumonia was detected in both groups until day 6, rgMex4108 seemed to replicate well in respiratory tissues without severe inflammation, as compared to rgMex4487. All four groups of ferrets developed multifocal, moderate to marked bronchiointerstitial pneumonia; however, ferrets inoculated via the intratracheal route tended to have more severe lesions. Pulmonary changes are predominately centered on terminal bronchioles, but often extend into adjacent bronchi and surrounding alveoli (Fig. 3E and 3F).

**DISCUSSION**

Since the pandemic of 2009, A(H1N1) pdm2009 circulating worldwide as seasonal flu has caused mild respiratory illness in infected individuals. A(H1N1) pdm2009 has occasionally caused severe disease and related deaths depending on the virus or the patient’s condition [2, 4, 14, 19, 21]. In our previous study, we characterized two Mexican isolates, Mex4108 and Mex4487, derived from a mild case and from a cluster of severe cases, respectively. They also produced heterogeneity in virulence in a cynomolgus macaque model [23]. However, in this study, experimental infection with both rgMex4108 and rgMex4487 caused only mild and undistinguishable respiratory diseases in ferrets. Ferrets inoculated with original isolates also showed mild symptoms, as observed by the recombinant viruses (data not shown). In this experiment, we compared both intranasal and intratracheal inoculation routes, and ferrets inoculated via the intratracheal route showed slightly reduced body weight by day 7 post-infection and tended to have more severe lesions by histopathological analysis. However, ferrets showed similar mild symptom in all groups. We did not find the significant clinical differences between the two isolates that were observed in the cynomolgus macaque model. The ferret is a common animal model for an influenza virus study. However, animal models do not always seem to mimic symptoms that were observed in human cases. To evaluate the pathogenicity difference of two Mexican isolates, we may need to use the cynomolgus macaque model, since cynomolgus macaque demonstrated similar clinical differences reflecting human cases in the previous study. On the other hand, it has been reported that Mex4487-infected ferrets showed severe clinical signs and that approximately 50% of the animals succumbed to infection within 9 dpi [11, 15]. This difference indicates that the outcome of disease might be affected by rearing environments of infected hosts as well as technical differences of experiments. However, rgMex4108 yielded a higher and more prolonged virus shedding in throat swabs, and a higher virus replication was detected until 5 dpi in throat swab. Although there is also no significant difference between viruses in our ferret model, these results suggested that Mex4108 may have the potential for long and wide spreading of viruses. To compare the transmissibility of viruses, contact transmission experiments are required.

A(H1N1) pdm2009 lacks amino acid mutations previously identified as human adaptation signature, like PB2-E627K and D701N, and these known virulence associated mutations in PB2 increased reporter gene expression, but did not affect virus replication and transmission [8]. Whereas, several A(H1N1) pdm2009 strains possessing amino acid substitutions in PB2 and HA showed enhanced virulence and transmission in ferret models [10, 31, 34]. However, both Mex4487 and Mex4108 do not have identified amino acid sequence that was contributed to virulence and transmission of viruses in animal models. Only 7 amino acid differences were identified in the genome of two Mexican isolates, which may need to use the cynomolgus macaque model, since cynomolgus macaque demonstrated similar clinical differences reflecting human cases in the previous study. On the other hand, it has been reported that Mex4487-infected ferrets showed severe clinical signs and that approximately 50% of the animals succumbed to infection within 9 dpi [11, 15]. This difference indicates that the outcome of disease might be affected by rearing environments of infected hosts as well as technical differences of experiments. However, rgMex4108 yielded a higher and more prolonged virus shedding in throat swabs, and a higher virus replication was detected until 5 dpi in throat swab. Although there is also no significant difference between viruses in our ferret model, these results suggested that Mex4108 may have the potential for long and wide spreading of viruses. To compare the transmissibility of viruses, contact transmission experiments are required.

The results of in vitro growth kinetics of recombinant viruses suggested that PB2 or PA of Mex4487 is a potential factor for prolonged virus replication in human alveolar epithelial cells. rgMex4108 possessing PA and/or PB2 of Mex4487 showed a higher rate of replication than that of rgMex4108. PB2 and PA are components of the RNA-dependent RNA polymerase complex of influenza A virus, and mutations in PB2 [7, 32, 33] or PA [5, 22, 24] associated with host range or viral pathogenicity have been reported. To determine the functional differences of two Mexican isolates, we analyzed the polymerase activity and inhibitory effects of IFN induction of both RNP complexes derived from Mex4108 or Mex4487 using a reporter assay. However, no difference was observed in both assays tested (Supplemental Fig. 2). We evaluated the polymerase activity in human cell lines, A549 and 293, and polymerase activities were similar for any combination of the RNP complex derived from Mex4108 or Mex4487. As another function of PB2, we also compared the inhibition activity of MAVS-mediated IFN-β induction. From the IFN-β promoter-driven reporter assay, significant inhibition of MAVS-mediated IFN-β induction by both PB2 and PA was observed. However, again, no difference was observed in this activity between proteins derived from Mex4108 or Mex4487 (Supplemental Fig. 2). These results suggested that the polymerase activity and inhibitory effects of IFN induction of these proteins are not critical factors in pathological differences of Mex4108 and Mex4487 shown in human cases and a cynomolgus macaque...
model. Recently, several novel proteins encoded by polymerase genes and additional functions of these proteins have been reported, suggesting the presence of additional proteins and/or functions associated with pathogenesis [17, 25, 30]. To investigate the importance of PB2 or PA in virulence, the pathogenesis of other recombinant viruses switching of PA and PB2 genes should be examined in animal models. Also, the pathogenetic differences and molecular function of viruses will need to be determined.

In this study, we showed that Mex4487 tends to induce severe pneumonia in infected individuals, whereas Mex4108 has wide-spreading potential with mild disease. However, there is no significant difference in most experiments, despite two Mexican isolates showing heterogeneity in virulence in a cynomolgus macaque. It might be the limitation of the ferret model, because animal models do not always reflect human cases. Whereas the clinical outcome of two isolates was different, it may indicate that background and conditions of infected individuals are important for determining the virulence as well as molecular determinants. To understand the pathogenicity and transmission properties of viruses for preparation for future risks, it is necessary to observe and accumulate the information of potential factors as well as signature sequences to determine the phenotype of influenza viruses.

CONFLICT OF INTEREST. None to declare.

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