Role of Permissive Neuraminidase Mutations in Influenza A/Brisbane/59/2007-like (H1N1) Viruses

Yacine Abed, Andrés Pizzorno, Xavier Bouhy, Guy Boivin*
Research Center in Infectious Diseases of the CHUQ-CHUL and Laval University, Québec City, Québec, Canada

Abstract
Neuraminidase (NA) mutations conferring resistance to NA inhibitors were believed to compromise influenza virus fitness. Unexpectedly, an oseltamivir-resistant A/Brisbane/59/2007 (Bris07)-like H1N1 H275Y NA variant emerged in 2007 and completely replaced the wild-type (WT) strain in 2008–2009. The NA of such variant contained additional NA changes (R222Q, V234M and D344N) that potentially counteracted the detrimental effect of the H275Y mutation on viral fitness. Here, we rescued a recombinant Bris07-like WT virus and 4 NA mutants/revertants (H275Y, H275Y/Q222R, H275Y/M234V and H275Y/N344D) and characterized them in vitro and in ferrets. A fluorometric-based NA assay was used to determine Vmax and Km values. Replicative capacities were evaluated by yield assays in ST6Gal1-MOCK cells. Recombinant NA proteins were expressed in 293T cells and surface NA activity was determined. Infectivity and contact transmission experiments were evaluated for the WT, H275Y and H275Y/Q222R recombinants in ferrets. The H275Y mutation did not significantly alter Km and Vmax values compared to WT. The H275Y/N344D mutant had a reduced affinity (Km of 50 vs 12 μM) whereas the H275Y/M234V mutant had a reduced activity (22 vs 28 μM/sec). In contrast, the H275Y/Q222R mutant showed a significant decrease of both affinity (40 μM) and activity (7 μM/sec). The WT, H275Y, H275Y/M234V and H275Y/N344D recombinants had comparable replicative capacities contrasting with H275Y/Q222R mutant whose viral titers were significantly reduced. All studied mutations reduced the cell surface NA activity compared to WT with the maximum reduction being obtained for the H275Y/Q222R mutant. Comparable infectivity and transmissibility were seen between the WT and the H275Y mutant in ferrets whereas the H275Y/Q222R mutant was associated with significantly lower lung viral titers. In conclusion, the Q222R reversion mutation compromised Bris07-like H1N1 virus in vitro and in vivo. Thus, the R222Q NA mutation present in the WT virus may have facilitated the emergence of NAI-resistant Bris07 variants.

Introduction
Influenza viruses are respiratory pathogens associated with significant public health consequences. Each year, influenza epidemics can be responsible for significant morbidity in the general population and excess mortality in elderly patients and individuals with chronic underlying conditions. Influenza A viruses of the H1N1 subtype have been associated with seasonal influenza epidemics for many decades and, in presence of immunological pressure, such viruses continue to evolve through genetic variability which is mainly confined to virus segments encoding surface glycoproteins i.e., the hemagglutinin (HA) and neuraminidase (NA) [1]. Consequently, viral strains to be used in annual influenza vaccines should be regularly updated to ensure optimal protection. Besides vaccines, neuraminidase inhibitors (NAI) including inhaled zanamivir, oral oseltamivir and intravenous peramivir provide an important additional measure for the control of influenza infections [2]. These antivirals target the active center of the influenza NA molecule, which is constituted by 8 functional (R-118, D-151, R-152, R-224, E-276, R-292, R-371, and Y-406; N2 numbering) and 11 framework (E-119, R-156, W-178, S-179, D-198, I-222, E-227, H-274, E-277, N-294, and E-425; N2 numbering) residues that are largely conserved among influenza A and B viruses [3]. However, the emergence of NAI-resistant viruses, as a result of drug use or due to circulation of natural variants, may compromise the clinical utility of this class of anti-influenza agents.

The H275Y (H274Y in N2 numbering) NA mutation conferring resistance to oseltamivir and peramivir has been detected with increasing frequency in seasonal A/H1N1 viruses since 2007 to the extent that almost all characterized A/Brisbane/59/2007-like (Bris07) (H1N1) influenza strains that circulated worldwide during the 2008–09 season were H275Y variants [4,5]. Interestingly, this drug-resistant strain seemed to have emerged independently of NAI use [6,7]. The rapid dissemination of the H275Y Bris07 variants in the absence of antiviral pressure suggests that the H275Y NA mutation may not compromise viral fitness and transmissibility in this recent H1N1 viral background. This contrasts with previous studies that analyzed the role of the H275Y mutation using older A/Texas/36/91 [8] and A/New Caledonia/99/01 [9] drug-selected H1N1 variants. Recent reports by our group and others have confirmed the differential impact of the H275Y mutation on viral fitness and enzymatic properties in the context of old and recent influenza H1N1 isolates.
Author Summary

The H275Y neuraminidase (NA) mutation conferring resistance to oseltamivir was shown to impair old influenza H1N1 strains both in vitro and in vivo. By contrast, an oseltamivir-resistant A/Brisbane/59/2007 (Bris07)-like H1N1 H275Y NA variant emerged in 2007 and completely replaced the wild-type (WT) strain in 2008–2009. This discrepancy could be attributed to permissive NA mutations (R222Q, V234M and D344N) that were identified in most Bris07-like oseltamivir-resistant variants. To verify this hypothesis, we developed a reverse genetics system for a sensitive Bris07-like isolate (275H) whose NA protein contains the 3 permissive mutations (222Q, 234M, 344N). Using mutagenesis, we first introduced the H275Y then reverted codons at positions 222, 234 and 344. The resulting 5 recombinants (WT, H275Y, H275Y/Q222R, H275Y/M234V and H275Y/N344D) were compared with regard to NA enzyme properties, replicative capacities in vitro as well as infectivity and contact-transmissibility in ferrets. Among the studied permissive mutations, Q222R was associated with a significant reduction of both affinity and activity of the NA enzyme resulting in a virus with a reduced replicative capacity in vitro and decreased replication in lungs of ferrets. Thus, the R222Q mutation may have been the major permissive NA change that facilitated the emergence and spread of NAI-resistant Bris07 variants.

In an attempt to provide a molecular explanation for this observation, previous authors suggested that secondary NA mutations such as D344N that emerged in H1N1 variants isolated after the 2006–07 season were associated with higher NA activity and affinity and could have facilitated the emergence of the H275Y mutation [11,12]. Such drug-resistant mutants may have a better HA-NA balance than the susceptible viruses and indeed completely replaced them in a short period of time. In addition, Bloom and colleagues recently described two other secondary NA mutations at codons 222 and 234 that may have counteracted the compromising impact of the H275Y mutation [13]. In that study, the V234M and R222Q mutations were shown to restore the viral fitness of an A/New Caledonia/20/99 H1N1 variant containing the H275Y mutation [13].

To further investigate which secondary NA mutations may have facilitated the introduction of the H275Y mutation in contemporarily seasonal H1N1 viruses and allowed their dissemination, we developed a reverse genetics system using a clinical Bris07 (H1N1) isolate as genetic background and evaluated the impact of the H275Y oseltamivir resistance mutation as well as several potential compensatory NA mutations on enzyme activity, viral fitness and transmissibility.

Results

In the present study, five recombinant Bris07 influenza viruses were generated i.e., the WT virus (containing the putative permissive mutations) that briefly circulated during the 2007–08 season, the single H275Y oseltamivir-resistant variant and three double mutants containing the H275Y mutation as well as reversion of potential permissive mutations (H275Y/Q222R, H275Y/M234V and H275Y/N344D). NA enzymatic properties using equivalent titers of recombinants were first analyzed with determination of relative NA enzymatic activity (Vmax values), which reflects the total NA activity per virion, and Km values, which reflect the affinity for the substrate. As shown in Table 1, the single H275Y mutation had no significant impact on NA affinity and activity compared to the WT virus in the context of the Bris07 background. By contrast, the double H275Y/Q222R mutation was associated with a significant reduction of both NA affinity (Km of 40.31 vs 11.95 μM, P<0.001) and relative NA activity (7.01 vs 28.19 U/sec, P<0.001) compared to the WT (Table 1 and Fig. 1). The H275Y/M234V mutant had a Km value comparable to that of the WT, whereas its relative NA activity was significantly reduced (Vmax of 21.89 vs 28.19 U/sec, P<0.05). The H275Y/N344D mutant showed a significantly reduced affinity (Km of 50.77 vs 11.95 μM, P<0.001) with no change in NA activity compared to the WT. When comparing the double mutants to the single H275Y mutant, the Km values were significantly increased for the H275Y/Q222R and H275Y/N344D mutants (P<0.001) whereas only the double H275Y/Q222R mutant had a significantly lower relative NA activity (P<0.001).

Using recombinant NA proteins expressed in 293T cells, we further investigated the impact of NA mutations on the amount of NA activity at the cell surface. As shown in Fig. 2, all studied mutations were associated with a significant reduction of total surface NA activity compared to the WT with relative total surface activities of 66% (P<0.01), 9.72% (P<0.001), 32.07% (P<0.001) and 54.89% (P<0.01) for the H275Y, H275Y/Q222R, H275Y/M234V and H275Y/N344D mutant proteins, respectively. When compared to the single H275Y mutant, H275Y/Q222R (P<0.001), H275Y/M234V (P<0.001) and H275Y/N344D (P<0.05) double mutants also had significantly reduced surface NA activities. The differences observed in total surface NA activity between the different recombinant NA proteins may be due to a decreased number of NA molecules that reached the cell surface or to less activity per enzyme.

We next determined the phenotype of resistance to NAIs for the 5 recombinant viruses. As expected, the presence of the H275Y mutation was associated with resistance to oseltamivir (mean fold increase of 2627 in IC50 values) and peramivir (mean fold increase of 998) with no impact on zanamivir susceptibility (Table 2).

Interestingly, comparison of the levels of resistance for the double recombinant mutants versus the single H275Y mutant revealed a significant reduction in the level of resistance to peramivir for the double H275Y/Q222R mutant (IC50 of 35.25 nM vs 59.85 nM, P<0.01). A similar trend was observed for oseltamivir (IC50 of 651.86 nM vs 1024.54 nM) although, in this case, the difference between IC50 values was not statistically significant.

Viral fitness of recombinant A/Brisbane/59/2007-like viruses was assessed in vitro using ST6Gal1-MDCK cells. The double H275Y/Q222R mutant produced viral plaques with a significantly reduced area compared to the recombinant WT (0.13 mm2 vs 0.53 mm2, P<0.001) whereas the remaining recombinants generated plaques of comparable sizes (Table 1). Of note, the reduction in plaque size for the H275Y/Q222R mutant was also significant compared to that of the single H275Y mutant (P<0.001). In replication kinetics experiments, the peak viral titers for all recombinants were obtained at 36 h post-infection (PI) with viral titers ranging from 5.6×10⁶ PFU/ml (H275Y/Q222R) to 3.3×10⁷ PFU/ml (WT) (Fig. 3). The WT, the single (H275Y) and the double (H275Y/N344D) mutants had comparable viral titers at all time points. By contrast, and in accordance with plaque size data, the double H275Y/Q222R mutant was associated with a significant reduction in viral titers at 36 h (P<0.001) and 48 h (P<0.05) PI compared to the WT (Fig. 3). There was also a significant reduction in the viral titer obtained at 36 h PI for the double H275Y/M234V mutant compared to the WT (P<0.001). When compared to the single (H275Y) mutant, viral titers of the
double H275Y/Q222R and H275Y/M234V mutants were significantly lower at 36 h (P<0.001).

Intranasal inoculation of ferrets with the WT and two mutant (H275Y and H275Y/Q222R) Bris07 recombinant viruses resulted in a febrile response that peaked on day 2 PI (Fig. 4A). The area under the curve (AUC) of temperatures between days 0 and 6 PI was similar for the 3 groups of ferrets i.e. 6.81±1.19 for the WT virus, 5.99±1.9 for the H275Y/Q222R mutant and 7.26±0.55 for the H275Y mutant. There was no significant difference in body weight between the three groups of animals at any time points (data not shown). As shown in Fig. 5A, mean viral titers in nasal wash samples collected on day 2 PI from ferrets infected with the recombinant WT and the single H275Y mutant were comparable (4×10^5±2.9×10^5 PFU/ml for the WT and 2.6×10^5±8.7×10^4 PFU/ml for the H275Y mutant) whereas the H275Y/Q222R mutant had a reduced mean viral titer (4.6×10^4±4.2×10^3 PFU/ml; P<0.05 vs WT). Similarly, mean viral titers in nasal wash samples of ferrets infected with the H275Y/Q222R were significantly lower than those of the H275Y mutant (P<0.05) and WT virus (P<0.01) on day 4 PI (3.4×10^5±1.7×10^5, 1.1×10^4±6.7×10^3 and 1.5×10^4±9.6×10^2 PFU/ml, respectively). On the other hand, the three recombinants were associated with comparable mean viral titers on day 6 PI (2×10^5±4.6×10^4 PFU/ml for the WT, 1.1×10^5±5.8×10^4 PFU/ml for the H275Y/Q222R and 1.3×10^5±8.1×10^4 PFU/ml for the H275Y).

All contact ferrets seroconverted for A/Brisbane/59/2007 when tested 14 days after contact, with geometrical mean hemagglutination inhibition (HAI) titers of 160±33, 145±119 and 95±55 for the WT, H275Y and H275Y/Q222R recombinant viruses, respectively. A febrile response could be observed on days 4 and 5 in the WT and the H275Y groups, respectively, but not in the H275Y/Q222R group (Fig. 4B). The AUC of temperatures between days 2 and 6 PI was similar between groups of ferrets infected with the recombinant WT (5.29±0.34) and its H275Y variant (4.54±0.19) whereas the AUC of the H275Y/Q222R group was significantly lower than that of the WT group (4.09±0.96; P<0.05). Viral titers in nasal wash samples collected on days 2, 4 and 6 PI are shown in Fig. 5B. Only the WT virus was detected on day 2 PI. Mean viral titers were comparable for the H275Y mutant and the WT virus on days 4 and 6 PI. In contrast,

Table 1. Neuraminidase enzymatic properties and plaque areas of recombinant A/Brisbane/59/2007-like (H1N1) viruses.

| Recombinants       | Km (µM)  | Relative NA activity (Vmax) (U/sec) | Vmax ratio vs WT | Plaque area (mm^2)  |
|-------------------|----------|------------------------------------|------------------|---------------------|
| WT                | 11.95±2.4| 28.19±0.74                         | 1.00             | 0.53±0.17           |
| H275Y             | 16.92±2.25| 23.81±1.95                         | 0.84             | 0.50±0.16           |
| H275Y/Q222R       | 40.31±5.3 | 7.01±0.11***                       | 0.25             | 0.13±0.06***        |
| H275Y/M234V       | 18.18±1.2 | 21.89±1.24*                        | 0.78             | 0.49±0.13           |
| H275Y/N344D       | 50.77±1.0 | 24.96±2.48                         | 0.89             | 0.50±0.15           |

*Values indicate mean Km and relative NA activity (Vmax) values of a representative experiment performed in triplicate ± standard deviations (SD).

**Values indicate mean plaque area (N = 16) ± SD.

P<0.05,

***P<0.001 compared to WT.

doi:10.1371/journal.ppat.1002431.t001

Figure 1. Neuraminidase (NA) enzyme kinetics of recombinant A/Brisbane/59/2007-like (H1N1) viruses. The rate of substrate conversion velocity (V0) by NA enzymes from a standardized dose of 10^6 PFU/ml of recombinant virus was determined. The fluorogenic substrate (MUNANA) was used at final concentrations of 0 to 3000 µM. Fluorescence was measured every 90 sec for 53 min at 37 °C using excitation and emission wavelengths of 355 and 460 nm, respectively. The data of one representative experiment performed in triplicate is shown.

doi:10.1371/journal.ppat.1002431.g001
Figure 2. Surface activity of recombinant A/Brisbane/59/2007-like (H1N1) neuraminidase proteins. 293T cells were transfected with pCAGGS-PA, -PB1, -PB2, and -NP plasmids in addition to plasmids expressing the WT or mutant A/Brisbane/59/2007-like neuraminidases (NA) proteins. At 24 h post-transfection, cells were treated with a non-lysing buffer and surface NA activity was measured by using the fluorogenic substrate (MUNANA). Percent surface NA activities were determined in triplicate experiments ± standard deviations. **P<0.01 and ***P<0.001 compared to the WT surface NA activity. doi:10.1371/journal.ppat.1002431.g002

Discussion

In this study, we used recombinant viruses derived from a clinical WT Bris07 strain to demonstrate using both in vitro and ferret experiments that the R222Q NA mutation was the main but possibly not the only permissive mutation that allowed the widespread dissemination of the oseltamivir-resistant H275Y mutant during the 2007–09 influenza seasons. Although such mutant seems to have disappeared since the emergence of the pandemic H1N1 virus in April 2009, understanding the mechanisms leading to the transmission of this unique virus is of great importance and could have an impact on the future use of NAIs.

The influenza NA protein plays a major role during the viral replication cycle. Its sialidase activity promotes virion release by removing sialic residues from viral glycoproteins and infected cells [14]. The NA enzyme also mediates virus penetration in the mucin layer of the respiratory tract, facilitating virus spread [15]. Importantly, the catalytic site of the NA enzyme has been shown to be conserved in all influenza A subtypes and influenza B viruses [3]. Therefore, the influenza NA protein has been considered as a suitable target for designing anti-influenza agents for both prophylactic and therapeutic purposes. Besides its functional role, the NA protein is a major structural surface glycoprotein that is exposed to the host immune pressure [14]. The NA gene, like the HA one, is therefore subject to more genetic variations than the rest of the influenza genome. Consequently, some amino acid (a.a.) changes, part of antigenic sites of the NA protein, may significantly contribute to the emergence of drifted variants, whereas certain substitutions located in or near the catalytic site may also affect the NA enzyme properties. For instance, Hensley and colleagues have recently identified NA mutations conferring resistance to zanamivir in variants of an influenza A/Puerto Rico/8/1934 H1N1 virus that was subjected to anti-HA monoclonal antibodies pressure [16].

In this study, we focused on a.a. changes that occurred in the NA protein during the evolution of recent seasonal influenza H1N1 viruses and that may have been involved in the development and dissemination of resistance to NAIs. These changes included the well-known framework H275Y mutation, responsible for the resistance phenotype to oseltamivir and peramivir, as well as other substitutions (V234M, R222Q and D344N) that may have contributed to the emergence and dissemination of resistance by acting as permissive/compensatory mutations.

Phylogenetic analyses previously demonstrated that the V234M mutation was already present in oseltamivir-susceptible A/Solomon Islands/3/2006 (SI06) viruses [13]. In another report, NA enzyme properties of SI06 viruses were found to be similar to those of older oseltamivir-susceptible strains such as A/New Caledonia/99/2001 in terms of relative NA activity (Vmax) and affinity (Km) [11]. By contrast, the appearance of the R222Q and D344N mutations in H1N1 viruses isolated after 2007 was associated with a significant increase in NA affinity (decreased Km values) in both 275H and 275Y strains [11]. In accordance with these observations, we demonstrated a sharp impact for the Q222R and N344D reversion mutations on Km values using our Bris07 recombinants (Table 1). Besides its effect on NA affinity, the Q222R reversion mutation was also associated with a significant decrease in relative NA activity (Table 1 and Fig. 1) and total NA activity that was expressed on the cell surface (Fig. 2), in line with previously-reported results in another viral background [13]. As a result, the H275Y/Q222R mutant virus was significantly com-

Table 2. Susceptibility profiles of recombinant A/Brisbane/59/2007-like (H1N1) viruses against neuraminidase (NA) inhibitors as assessed by MUNANA NA inhibition assays.

| Recombinants | Oseltamivir* (nM) | Zanamivir* (nM) | Peramivir* (nM) |
|--------------|-----------------|----------------|----------------|
| WT           | 0.39±0.02       | 0.18±0.03      | 0.06±0.01      |
| H275Y        | 1024.54±114.10  | 0.27±0.06      | 59.8±2.42      |
| H275Y/Q222R  | 651.86±116.88   | 0.18±0.04      | 35.25±2.11**   |
| H275Y/M234V  | 1038.56±116.09  | 0.20±0.01      | 47.87±1.33     |
| H275Y/N344D  | 735.07±67.91    | 0.28±0.03      | 47.15±3.76     |

*Values indicated mean IC50 values of three experiments ± standard deviations.
**P<0.01 compared to the single H275Y mutant.
doi:10.1371/journal.ppat.1002431.t002
promised in vitro based on plaque size and replication kinetics patterns. Such decreased viral replication of the H275Y/Q222R mutant was also evident in vivo, resulting in lower viral titers in nasal wash samples and an absence of febrile response in contact ferrets. However, the H275Y/Q222R mutant was transmitted to all naive ferrets by direct contact meaning that the combination of several permissive NA mutations and/or mutations elsewhere in the viral genome may be necessary to recapitulate the epidemiological observations showing increased transmission of the oseltamivir-resistant Bris07 virus. Also, it should be noted that naive (non-immune) ferrets may not completely capture the fitness of Bris07 in humans with pre-existing immunity. Alternatively, the Q222R mutation could affect airborne transmission which has not been evaluated in our study. Of note, possibly due to the lower affinity of Q222R for MUNANA, less NAIs were required for competitive inhibition of the H275Y/Q222R mutant compared to the H275Y mutant. Residue 222 is located in the vicinity of the catalytic site of the N1 enzyme based on 3-D structure analysis [17]. Thus, substitution of a charged (R) by an uncharged (Q) a.a. at codon 222 may be the main change that dramatically altered the NA enzyme properties of recent seasonal H1N1 viruses. Of interest, only one NA substitution (R194G) was sufficient to restore the viral fitness of an influenza A/WSN/33 (H1N1) virus containing the compromising H275Y NA mutation [13].

In addition to the R222Q mutation, a permissive role was also suggested for V234M and D344N substitutions [11,13]. Interestingly, in a recent report on the evolution of influenza NA genes, positive epistasis (i.e., combination of mutations that are substantially more beneficial than single mutations alone) was detected in pairs of codons within the NA gene of the N1 subtype including 275–222, 275–234, and 275–344 [18]. In our study, although the M234V and N344D reversions were associated with decreased relative NA activity and affinity, respectively (Table 1 and Fig. 1), none of these mutations significantly altered the viral fitness in vitro. Nevertheless, a possible synergy between these mutations and Q222R cannot be completely excluded.

Our study revealed that the H275Y NA mutation was not deleterious to fitness in the Bris07 genetic context in contrast to older H1N1 strains. However, this mutant did not have a replicative advantage compared to the WT as suggested by epidemiological studies. Indeed, the recombinant WT virus and its H275Y variant demonstrated similar replication kinetics during in vitro experiments. In addition, these recombinants had comparable infectivity and contact transmissibility in ferrets. Thus, the presence of the permissive mutations (R222Q, V234M and D344N) in the NA protein of our WT strain was apparently not sufficient to alter the viral fitness to the level that a compensatory change, such as the H275Y mutation, would be necessary. Therefore, we believe that changes in the NA gene alone may not provide a complete explanation for the emergence and spread of the oseltamivir-resistant H275Y Bris07 variant. Other changes in the genome might have been involved in this event. For instance, Yang and colleagues recently demonstrated that the dominant H275Y variant that emerged in Taiwan in 2007–2008 was a result of intra-subtypic reassortments between HA, NA, PB2 and PA genes from one clade (clade 2B) and the remaining 4 genes from another one (clade 1) [19]. Furthermore, the H275Y NA substitution and other changes in NA, HA, PB1 and PB2 proteins occurred in that background [19]. Thus, it would be also interesting to assess the effect of HA and particularly polymerase mutations that differed between WT and H275Y mutant clinical Bris07 isolates on replicative capacities and transmissibility.

Despite the fact that the secondary mutations described here were not investigated individually but in conjunction with H275Y, our study provides a comprehensive analysis of relevant permissive NA mutations in the contemporarily seasonal H1N1 background. This included in vitro characterization, assessment of viral fitness and contact transmission in ferrets as well as NA enzyme
Figure 4. Body temperatures of infected and contact ferrets. Body temperatures were recorded by rectal thermometer during 10 days post-inoculation in groups of 4 index ferrets infected with $1.25 \times 10^5$ PFU of recombinant A/Brisbane/59/2007-like wild-type (WT) virus as well as H275Y and H275Y/Q222R mutants (A) and in groups of 4 naive ferrets that were placed in direct contact with index ferrets 24 h later (B).

doi:10.1371/journal.ppat.1002431.g004
properties of recombinant mutants. In particular, our investigation clearly demonstrated the positive impact of one specific NA substitution (i.e. R222Q) in conjunction with the oseltamivir resistance H275Y mutation on enzymatic properties and viral fitness of the Bris07 H1N1 strain. Noteworthy, our results suggest that total NA activity was more likely predictive of in vitro and in vivo viral fitness than the enzyme affinity (Km) parameter. Whether the Q222R mutation is also deleterious in the absence of H275Y was not investigated here; however, in a previous work, influenza A/Paris/497/2007 (222Q/275H) and A/Solomon Islands/3/2006 (222R/275H) seasonal H1N1 isolates grew to comparable titers in in vitro kinetics experiments [11]. Although clinical 2009 pandemic H1N1 variants containing such permissive mutations have not been reported, a computational approach had recently led to the identification of R257K and T289M as potential secondary mutations in that context [20]. Thus, monitoring for resistance in influenza viruses should take into consideration not only NA resistance-mutations themselves but also permissive/secondary ones as the latter may significantly affect the clinical and epidemiological impacts of seasonal or pandemic influenza viruses.

Materials and Methods

Ethics Statement

All procedures were approved by the Institutional Animal Care Committee at Laval University according to the guidelines of the Canadian Council on Animal Care.

Rescue of recombinant viruses

Reverse transcription-PCR using universal influenza primers [21] was used to amplify the eight genomic segments of an oseltamivir-susceptible A/Quebec/15230/08 (H1N1) isolate whose HA and NA genes shared respectively 99.53% and 99.71% nucleotide identity with those of the influenza A/Brisbane/59/2007 vaccine strain [10]. All segments were cloned into the pJET plasmid (Fermentas, Burlington, ON, Canada) and sequenced. Sequence analysis

Figure 5. Mean viral titers in nasal wash samples of infected and contact ferrets. Mean viral titers ± standard deviations were determined in nasal washes by using standard plaque assays in groups of 4 index ferrets infected with 1.25 × 10⁶ PFU of recombinant A/Brisbane/59/2007-like wild-type (WT) virus as well as H275Y and H275Y/Q222R mutants (A) and in groups of 4 naïve ferrets that were placed in direct contact with index ferrets 24 h later (B). *P < 0.05 and **P < 0.01 for differences in viral titers when compared to the recombinant WT virus.

doi:10.1371/journal.ppat.1002431.g005
confirmed the presence of histidine (H), glutamine (Q), methionine (M) and asparagine (N) residues at residues 275, 222, 234 and 344 (N1 numbering), respectively, of the NA protein. The PB1, PB2 and PA segments were sub-cloned into pLLBG whereas the HA, NA, NP, M1/M2 and NS1/NS2 segments were sub-cloned into pLLBA bidirectional expression/translation vectors as described [22]. The pLLBA plasmid containing the NA gene was used for the introduction of the H275Y mutation using appropriate primers and the QuickChangeTM Site-Directed Mutagenesis kit (Stratagene, La Jolla, CA). The resulting pLLB-NAH275Y mutant plasmid was then used for reverting potential compensatory mutations (Q222R, M234V or N344D) as described above. All recombinant plasmids were sequenced to confirm the absence of undesired mutations. The eight bidirectional plasmids were cotransfected into 293T human embryonic kidney cells using the LipofectamineTM 2000 reagent (Invitrogen, Carlsbad, CA) as previously described [23]. Supernatants were collected 72 h post-transfection and used to inoculate ST6Gal1-MDCK cells kindly provided by Dr. Y. Kawaoka, University of Wisconsin, Madison, WI. The recombinant wild-type (WT) and H275Y, H275Y/Q222R, H275Y/M234V and H275Y/N344D mutant viruses were subsequently sequenced and titrated by standard plaque assays in ST6Gal1-MDCK cells.

NA enzyme kinetics assays

A fluorometric based assay using MUNANA (Methylumbelliferyl-N-acetylmuramyl acid) (Sigma, St-Louis, MO) as substrate was performed to determine total NA enzymatic activity per infectious virus [24]. Briefly, recombinant viruses were standardized to an equivalent dose of 10^6 plaque forming-units (PFU)/ml and incubated at 37°C in 50-μl reactions with different concentrations of MUNANA. The final concentration of the substrate ranged from 0 to 3000 μM. Fluorescence was monitored every 90 s for 53 min (35 measures). The NA kinetic parameters (Km and Vmax) were calculated using standard HAI assays. To evaluate contact-transmissibility, plaque assays using ST6Gal1-MDCK cells. Serum samples were collected from animals on days 2, 4 and 6 PI. Virus titers from nasal wash samples were determined by plaque assays using ST6Gal1-MDCK cells. Serum samples were collected from each ferret before intranasal infection and on day 14 PI to evaluate specific antibody levels against the seasonal Bris07 strain using standard HAI assays. To evaluate contact-transmissibility, inoculated-contact animal pairs were established by placing a naive ferret into each cage 24 h after inoculation of the index ferret [27]. Contact animals were monitored for clinical signs and nasal wash and serum samples were collected as described above for determination of viral titers and serological status, respectively.

Statistical analyses

NA kinetic parameters (Km and Vmax values), NA IC50 values and viral titers in vitro and in nasal washes of ferrets were compared by one-way ANOVA analysis of variance, with the Tukey’s multiple comparison post test. The amount of NA activity on the cell surface and plaque sizes of the recombinants were compared to those of the WT virus and/or the H275Y mutant by the use of unpaired two-tailed t tests.

Author Contributions

Conceived and designed the experiments: YA GB. Performed the experiments: YA AP XB. Analyzed the data: YA AP GB. Contributed reagents/materials/analysis tools: AP XB. Wrote the paper: YA GB.

References

1. Reid AH, Fanning TG, Janczewski TA, Taubenberger JK (2000) Characterization of the 1918 "Spanish" influenza virus neuraminidase gene. Proc Natl Acad Sci U S A 97: 6785–6790.
2. Bolz DA, Aldridge JR, Jr, Webster RG, Govorkova EA (2010) Drugs in development for influenza. Drug 50: 1549–1562.
3. Colman PM, Hoyne PA, Lawrence MC (1993) Sequence and structure alignment of paramyxovirus hemagglutinin-neuraminidase with influenza virus neuraminidase. J Virol 67: 2972–2980.
4. Lackenby A, Hungnes O, Dudman SG, Meijer A, Paget WJ, et al. (2000) Emergence of resistance to oseltamivir among influenza A(H1N1) viruses in Europe. Euro Surveill 15: pii: 8026.
5. Hurt AC, Ernest J, Deng YM, Iannello P, Besselaar TG, et al. (2009) Emergence and spread of oseltamivir-resistant A(H1N1) influenza viruses in Oceania, South East Asia and South Africa. Antiviral Res 83: 90–93.
6. Hauge SH, Dudman S, Borgen K, Lackenby A, Hungnes O (2009) Osimertinib-resistant influenza viruses A(H1N1), Norway, 2007-08. Emerg Infect Dis 15: 155–162.
7. Dharn JN, Guibervara LV, Meyer JJ, Okomo-Adhiambo M, Mcclinton RC, et al. (2009) Infections with oseltamivir-resistant influenza A(H1N1) virus in the United States. JAMA 301: 1034–1041.
8. Hoehl MO, Trucon R, Elias S, Yen HL, Roberts NA, et al. (2004) Influenza viruses resistant to the antiviral drug oseltamivir: transmission studies in ferrets. J Infect Dis 190: 1627–1630.
9. Ives JA, Carr JA, Mendel DR, Tai CY, Lambkin R, et al. (2002) The H274Y mutation in the influenza A/H1N1 neuraminidase active site following oseltamivir phosphate treatment leave virus severely compromised both in vitro and in vivo. Antiviral Res 55: 307–317.

10. Baz M, Abed Y, Simon P, Hamelin ME, Boivin G (2010) Effect of the neuraminidase mutation H274Y conferring resistance to oseltamivir on the replicative capacity and virulence of old and recent human influenza A/H1N1 viruses. J Infect Dis 201: 740–745.

11. Ramezu-Welti MA, Enouf V, Cavalerie F, Jeannin P, van der Werf S (2008) Enzymatic properties of the neuraminidase of seasonal H1N1 influenza viruses provide insights for the emergence of natural resistance to oseltamivir. PLoS Pathog 4: e1000103.

12. Collins PJ, Haire LF, Lin YP, Liu J, Russell RJ, et al. (2009) Structural basis for oseltamivir resistance of influenza viruses. Vaccine 27: 6317–6323.

13. Bloom JD, Gong LI, Baltimore D (2010) Permissive secondary mutations enable the evolution of influenza oseltamivir resistance. Science 328: 1272–1275.

14. Hayden FG, Palese P (2002) Influenza virus. In: Douglas RWJ, Richman D, Hayden FG, eds. Clinical Virology. 2nd edition. Washington DC: ASM Press. pp 891–920.

15. Palese P, Compan RW (1976) Inhibition of influenza virus replication in tissue culture by 2-deoxy-2,3-dehydro-N-trifluoroacetylneuraminic acid (FANA): mechanism of action. J Gen Virol 33: 159–163.

16. Hendley SE, Das SR, Gibso JS, Bailey AL, Schmidt LM, et al. (2011) Influenza a virus hemagglutinin antibody escape promotes neuraminidase antigenic variation and drug resistance. PLoS One 6: e15190.

17. Russell RJ, Haire LF, Stevens DJ, Collins PJ, Lin YP, et al. (2006) The structure of H5N1 avian influenza neuraminidase suggests new opportunities for drug design. Nature 438: 45–49.

18. Kryazhimskiy S, Dushoff J, Baszkin GA, Plotkin JB (2011) Prevalence of epistasis in the evolution of influenza a surface proteins. PLoS Genet 7: e1001301.

19. Yang JR, Lin YC, Huang YP, Su CH, Lo J, et al. (2011) Reassortment and mutations associated with emergence and spread of oseltamivir-resistant seasonal influenza A/H1N1 viruses in 2005-2009. PLoS One 6: e18177.

20. Bloom JD, Nayak JS, Baltimore D (2011) A computational-experimental approach identifies mutations that enhance surface expression of an oseltamivir-resistant influenza neuraminidase. PLoS One 6: e22201.

21. Hoffmann E, Stech J, Guan Y, Webster RG, Perez DR (2001) Universal primer set for the full-length amplification of all influenza A viruses. Arch Virol 146: 2275–2289.

22. Liu Q, Wang S, Ma G, Pu J, Forbes NE, et al. (2009) Improved and simplified recombiningering approach for influenza virus reverse genetics. J Mol Genet Med 3: 225–231.

23. Abed Y, Baz M, Boivin G (2006) Impact of neuraminidase mutations conferring influenza resistance to neuraminidase inhibitors in the N1 and N2 genetic backgrounds. Antivir Ther 11: 971–976.

24. Duan S, Bolzt DA, Seiler P, Li J, Bragstad K, et al. (2010) Oseltamivir-resistant pandemic H1N1/2009 influenza virus possesses lower transmissibility and fitness in ferrets. PLoS Pathog 6: e1001022.

25. Pizzorno A, Bouhy X, Abed Y, Boivin G (2011) Generation and characterization of recombinant pandemic influenza A/H1N1 viruses resistant to neuraminidase inhibitors. J Infect Dis 203: 25–31.

26. Abed Y, Goyette N, Boivin G (2004) A reverse genetics study of resistance to neuraminidase inhibitors in an influenza A/H1N1 virus. Antivir Ther 9: 347–381.

27. Hamelin ME, Baz M, Abed Y, Couture C, Joubert P, et al. (2010) Oseltamivir-resistant pandemic A/H1N1 virus is as virulent as its wild-type counterpart in mice and ferrets. PLoS Pathog 6: e1001015.