Mutations Designed by Ensemble Defect to Misfold Conserved RNA Structures of Influenza A Segments 7 and 8 Affect Splicing and Attenuate Viral Replication in Cell Culture

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Abstract

Influenza A virus is a significant public health threat, but little is understood about the viral RNA structure and function. Current vaccines and therapeutic options to control influenza A virus infections are mostly protein-centric and of limited effectiveness. Here, we report using an ensemble defect approach to design mutations to misfold regions of conserved mRNA structures in influenza A virus segments 7 and 8. Influenza A mutant viruses inhibit pre-mRNA splicing and attenuate viral replication in cell culture, thus providing evidence for functions of the targeted regions. Targeting these influenza A viral RNA regions provides new possibilities for designing vaccines and therapeutics against this important human respiratory pathogen. The results also demonstrate that the ensemble defect approach is an efficient way to test for function of RNA sequences.

Introduction

Influenza A virus is a segmented, single-stranded, negative sense RNA virus [1]. Every year, 15–20% of the world’s population is infected, and 250 to 500 thousand people are killed by influenza A virus [2]. Currently, the most widely used vaccine is the Inactivated Influenza Vaccine (IIV), which is given by intramuscular injection [3]. IIV is noninfectious, and thus recommended for all people above 6 months of age [4], but its immunogenicity is relatively weak [5, 6]. The other option for vaccination is the Live-Attenuated Influenza Vaccine (LAIV), which is given intranasally [7]. The live-attenuated virus includes five mutations within the A/Ann Arbor/6/60 H2N2 backbone [8, 9]. LAIV is temperature sensitive, so it only replicates in the cooler upper respiratory tract (33°C), but cannot damage the warmer lower respiratory tract (37°C). Because of safety concerns, it is only recommended for healthy, non-pregnant...
individuals 2–49 years of age [10, 11]. In the 2014–2015 influenza season, the overall vaccine effectiveness was as low as 19% [12], due to antigenic drifts in circulating H3N2 viruses. There is a huge demand for increasing both safety and protective efficacy of current influenza vaccine approaches. Structure targeted mutations in mRNA can potentially be used as a new approach to develop weakened virus for LAIV.

Two classes of therapeutics have been approved by the Food and Drug Administration (FDA) for use against influenza: adamantanes and neuraminidase inhibitors [13]. Both target essential proteins coded by influenza, the ion channel protein M2, and the neuraminidase protein, respectively [14]. Most circulating influenza viruses are already resistant to adamantanes [15, 16], and are starting to build up resistance to neuraminidase inhibitors [17]. No new anti-influenza drug has been approved in the US since 1999. Most new targets being investigated are still protein-centric [13]. Conserved viral RNA structures important for function are potentially new therapeutic targets.

Segments 7 and 8 of influenza A virus code for M1/M2 and NS1/NEP proteins, respectively [1], where M1 is a matrix protein connecting vRNAs to each other and to the viral envelope, NS1 is an interferon antagonist, and NEP is a nuclear export protein. Based largely on sequence comparison and predicted thermodynamics, conserved RNA structures were identified near or containing splice sites in segments 7 and 8 mRNAs [18, 19]. In segment 7, a three-way multi-branch loop (7MB) that is 79 nucleotides (nts) downstream from the 5′ splice site [20] (Fig 1A) and a pseudoknot/hairpin equilibrium (7PK) at the 3′ splice site [21] (Fig 1C) were confirmed in vitro by enzymatic and chemical mapping. In segment 8, a pseudoknot/hairpin equilibrium (8PK) at the 3′ splice site [19] (Fig 2A) was identified. In this work, each of these conserved structures was mutated by changing only two nucleotides predicted by an ensemble defect program [22] to completely change the secondary structure. In cell culture, the influenza A mutants affected mRNA splicing and attenuated viral replication. Similar effects were also observed upon disrupting a single UA base pair in a pseudoknot containing the 3′ splice site of segment 8. The results demonstrate that the ensemble defect approach can rapidly reveal function for regions predicted to fold into stable secondary structures.

Experimental Methods

Cells

Madin–Darby canine kidney cells (MDCK; ATCC CCL-34), human lung epithelial carcinoma cells (A549; ATCC CCL-185), and human embryonic kidney 293T cells (293T; ATCC CCL-11268) were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Mediatech, Inc.) supplemented with 10% fetal bovine serum (FBS; Atlanta Biologicals) and 1% PSG (penicillin, 100 units/ml; streptomycin, 100 μg/ml; L-glutamine 2 mM; Mediatech, Inc.). Cells were grown at 37°C in air enriched with 5% CO₂.

Design of mutations using an ensemble defect program

The ensemble defect program was adapted from Zadeh, et al. [22]. Ensemble defect, n, is used here to choose a sequence, Φ, unlikely to fold into the wild type (WT) secondary structure, s. Ensemble defect uses thermodynamics to predict the average number of nucleotides paired differently in mutated sequence, Φ, relative to WT structure, s, at equilibrium over the ensemble of possible secondary structures:

\[ n(\emptyset, s) = \sum_{\sigma} p(\emptyset, \sigma) d(\sigma, s) \]

Here, Γ corresponds to the unpseudoknotted structural ensemble for the sequence Φ. p(Φ, σ) is
Fig 1. Conserved RNA structures identified in mRNAs of influenza A virus segment 7 and predicted folding of ensemble defect mutants. Ensemble defect designed mutations are indicated by arrows in A and C and by boxes in B and D. Wild-type (WT) sequences are from influenza A/Puerto Rico/1934 H1N1 (PR8) strain. Predicted free energies of folding at 37°C for WT and mutated sequences are provided below each structure. For hairpins and pseudoknots, respectively, \( \Delta G^\circ \) was predicted by RNAstructure [26] and ShapeKnots programs [27]. (A) The three-way multi-branch loop, 7MB, 79 nts downstream from the 5' splice site in segment 7 [20]. (B) The predicted folding of 7MB_ED. (C) Pseudoknot/hairpin equilibrium at the 3' splice site in segment 7 [21]. (D) The predicted folding of 7PK_ED.

WT \( \Delta G^\circ_{37} = -29.9 \) kcal/mol
mut \( \Delta G^\circ_{37} = -24.3 \) kcal/mol

\( \Delta G^\circ_{37} = -30.6 \) kcal/mol

WT \( \Delta G^\circ_{37} = -7.9 \) kcal/mol
mut \( \Delta G^\circ_{37} = -7.9 \) kcal/mol

\( \Delta G^\circ_{37} = -22.2 \) kcal/mol

\( \Delta G^\circ_{37} = -1.8 \) kcal/mol
mut \( \Delta G^\circ_{37} = -1.8 \) kcal/mol
Fig 2. Conserved RNA structures identified in mRNAs of influenza A virus segment 8 and predicted folding of ensemble defect mutants. (A) Pseudoknot/ hairpin equilibrium at the 3’ splice site in segment 8 [19]. The lower helix in the pseudoknot conformation may be extended by 4 additional base pairs and a single mismatch forming between nts 494–498 and 529–533 [19]. (B) The predicted folding of 8PK_ED. (C) Black arrow corresponds to mutation in 8PK_BP and blue arrow corresponds to compensating mutation in 8PK_BP_CP. Details are same as in Fig 1 caption.

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the equilibrium probability of any secondary structure $\sigma$ belonging to $\Gamma$ (i.e. a secondary structure that can be formed by the sequence $\Phi$) calculated using the partition function $[23, 24]$. $d(\sigma, s)$ is the distance between the WT secondary structure, $s$, and another secondary structure, $\sigma$, with $d$ defined as the number of nucleotides paired differently in the two structures. Thus, the goal is to computationally maximize $n$ to predict a sequence, $\Phi$, that will fold into a new secondary structure that is very different and much more probable than the WT structure.

For the target sequences, all 2-nt mutations were computed and used to calculate the ensemble defects. Sequences with a maximized ensemble defect and minimal amino acid changes were manually selected. For the WT sequences that can fold into an equilibrium between a hairpin and a pseudoknot, the ensemble defects were calculated against the hairpin conformation. The free energy of folding at $37^\circ C$, $\Delta G^{37}$, was predicted by the RNAstructure $[25]$ and the ShapeKnots program $[26]$ for unpseudoknotted and pseudoknotted structures, respectively.

**Site-directed Mutagenesis**

QuickChange Site-Directed Mutagenesis kit (Agilent Technologies) was used to introduce mutations in pPolI M and NS plasmids $[27]$. Briefly, PCR reactions were assembled with two oligonucleotide primers containing the desired mutations, following manufacturer’s recommendations. The oligonucleotide primers, each complementary to opposite strands of the M and NS viral segments, were extended during temperature cycling by the Pfu Turbo DNA polymerase. Incorporation of the primers generates a plasmid containing the desired mutations. Following temperature cycling, Dpn I was added to the reaction mix and the sample was incubated for 1 h at $37^\circ C$ to digest the parental plasmids, leaving only the non-methylated plasmids generated by PCR. The resulting plasmids containing the desired mutations were then transformed into the XL1-Blue supercompetent cells. To ensure correct mutations, plasmids were sequenced (Genewiz, Inc.).

**Generation of influenza viruses from plasmid DNA**

WT influenza A/Puerto Rico/8/1934 H1N1 (PR8) virus $[27]$ was generated using plasmid-based reverse genetic techniques, as described previously $[28]$. Briefly, co-cultures (1:1) of MDCK/293T cells (6-well plate format, $10^6$ cells/well) were co-transfected with ambisense plasmids (pPDZ) containing the eight PR8 viral segments using Lipofectamine 2000 (Invitrogen). At 16 h post-transfection, the medium was changed to DMEM containing 0.3% bovine serum albumin (BSA), 1% PSG, and 1 $\mu$g/ml tosylsulfonyl phenylalanyl chloromethyl ketone (TPCK)-treated trypsin (Sigma). At 48 h post-transfection, tissue culture supernatants (TCS) were collected and used to infect fresh MDCK cells. At 48 h post-infection, recombinant viruses were plaque purified in MDCK cells. Virus stocks were generated by infecting confluent 10-cm dishes of MDCK cells at a multiplicity of infection (MOI) equal to 0.001. Viruses containing designed mutations were generated similarly using the pPolI M or NS plasmids containing the desired mutations instead of the WT pPDZ plasmids. After preparing the mutant virus stocks, the mutated genomic segment was amplified by RT-PCR and sequenced to ensure the presence of introduced mutations.

**Determination of virus titers using immunofluorescence assay (IFA)**

Influenza PR8 WT and mutant viral titers were measured by immunofocus assay as described by Baker, et al. $[29]$. Briefly, 10-fold serial dilutions of viruses were used to infect confluent monolayers of MDCK cells ($10^4$ cells/well, 96-well plate format) in triplicate. Cells were incubated with viruses for 1 h at room temperature to allow virus absorption. After that, cells were
washed with PBS, and incubated with DMEM containing 0.3% BSA, 1% PSG, and 1 μg/ml TPCK-treated trypsin at 37°C for 8 h. Then cells were fixed and permeabilized in 4% formaldehyde and 0.5% Triton X-100 for 15 min at room temperature. After washing with PBS, the cells were incubated in the blocking solution of 2.5% BSA overnight at 4°C, and then incubated with 1 μg/ml of an anti-influenza A NP antibody (HT103) for 1 h at 37°C. After washing three times with PBS, the cells were incubated with a fluorescein isothiocyanate (FITC)-conjugated rabbit anti-mouse IgG secondary antibody (Dako) and 1 mg/ml of 4′,6′-diamidino-2-phenylindole (DAPI; Research Organics) for 1 h at 37°C. Cells were visualized under a fluorescence microscope, and fluorescently-labeled NP-expressing cells were quantified to determine the virus titer (fluorescent focus forming units [FFU]/ml). The mean value and standard deviation were calculated using Microsoft Excel software.

Virus multi-cycle growth kinetics
Multi-cycle viral growth kinetics were measured in MDCK and A549 cells (5 x 10^5 cells/well, 12-well plate format, triplicates) by infecting with WT or mutant viruses (MOI of 0.001 or 0.0001). After virus absorption for 1 h at room temperature, cells were washed with PBS, overlaid with DMEM containing 0.3% BSA, 1% PSG, and 1 μg/ml TPCK-treated trypsin, and incubated at 37°C. TCS were collected at indicated time points post-infection, and virus titers were measured using the immunofocus assay, as described above. Mean value and standard deviation (SD) were calculated using Microsoft Excel software.

Quantitative RT-PCR (qRT-PCR)
WT and mutant viruses were used to infect MDCK or A549 cells (5 x 10^5 cells/well, 12-well plate format, triplicates) with MOI of 5. An MOI of 5 was required to generate enough viral RNA for qRT-PCR. Cells were collected at 3 h post-infection, and total RNAs were extracted using RNeasy kit (Qiagen). One μg total RNA was reverse transcribed using oligo(dT) primers with SuperScript II reverse transcriptase (Invitrogen) to amplify mRNA. StepOnePlus Real-Time PCR System and Fast SYBR Green Master Mix (Invitrogen) were used for qPCR with the cDNAs as templates. Primers used for amplification are listed in S1 Table. The relative mRNA levels were analyzed using the Pfaffl method [30]. Amplification specificity was confirmed by melting curve analysis at the end of each program.

Results
Mutations designed to misfold three conserved RNA structures by maximizing ensemble defects
An ensemble defect program (see Methods) was used to predict sequences with nucleotides paired completely differently from the WT sequence. For three conserved RNA structures from segments 7 and 8, separate 2-nt mutations, 7MB_ED, 7PK_ED, and 8PK_ED, were predicted to maximize ensemble defect (Figs 1, 2A and 2B). Thus, the mutated sequences are forced to fold predominately into secondary structures different from WT.

In 7MB_ED and 7PK_ED, the mutants are predicted to fold into double hairpin conformations (Fig 1B and 1D). The free energy changes for folding at 37°C are more favorable by 6.3, 14.3, and 20.4 kcal/mole, respectively, compared with the WT structure of the multi-branch loop, the hairpin, and pseudoknot conformations if formed by the mutated sequences at 37°C. (Note that at 37°C, every change by 1.4 kcal/mol corresponds to a 10-fold change in equilibrium constant for folding.)
The mutations in 7MB_ED (A138C and G189C) are silent. The mutations in 7PK_ED (A724C and A726C) are 30 from the splice site, and change three amino acids: K242H in M1, and R12S plus N13T in M2 (Table 1). The same nucleotide and amino acid mutations exist in a natural strain, A/duck/Jiangsu/2012 (H5N2) [31].

In 8PK_ED, the mutations (A491U and U498A) enable the mutant to fold into a single hairpin conformation different from WT (Fig 2A and 2B). The predicted free energy is more favorable by 6.9 and 8.5 kcal/mole, respectively, compared with the WT hairpin and pseudoknot conformations if formed by the mutated sequence at 37°C. The mutations are 30 from the splice site, and change three amino acids: H169L in NS1, and I12L and L14Q in NEP (Table 1). That combination of amino acid changes has not been observed in natural influenza A strains.

NMR spectra for the 30 splice site pseudoknot in segment 8 of strain A/duck/Shanghai/13/01 [16] revealed coaxial stacking between the two helixes [19]. On the basis of sequence comparison, two mutants, 8PK_BP and 8PK_BP_CP, were designed to, respectively, disrupt and repair the coaxial stack (Fig 2C). Mutant 8PK_BP has a single silent mutation, U483C, that destabilizes the pseudoknot conformation by 2.6 (= 8.3–5.7) kcal/mol at 37°C (Fig 2A and 2C), while the hairpin conformation is predicted to maintain the same thermodynamic stability. The compensatory mutation, A522G, in 8PK_BP_CP restores the canonical base pair between nt 483 and nt 522, thus allowing coaxial stacking (Fig 2C). A522G changes an amino acid, E22G, in NEP. Both E and G exist widely at position 522 (Table 1). A single A522G mutation was not tested because in the pseudoknot, this would replace an AU pair with a GU pair, which is predicted to destabilize the helix by only 0.1 kcal/mol at 37°C [32]. Similarly, in the hairpin conformation, a GA pair would be replaced with a GG pair, which is predicted not to change folding.

### Mutated viruses affect splicing of segments 7 and 8 pre-mRNAs in cell culture

Plasmid-based reverse genetics techniques [28] were used to generate the recombinant mutated segment 7 and 8 PR8 viruses. All the mutant viruses were successfully rescued with titers similar to those of WT PR8. Sequencing showed that they retained the designed mutations (data not shown). For segment 7 mutants, total RNAs were extracted from MDCK and A549 cells 3 h after infection (MOI = 5) at 37°C. MDCK cells were used because they are common models for influenza studies [33]. Human lung epithelial A549 cells were used because they are more representative of cells infected in humans. The relative amounts of M1 and M2 mRNAs were measured using qRT-PCR. The results showed that M2 mRNA abundance was similar in WT and mutant viruses, while M1 mRNA abundance was raised about 1.4 to 2 fold in the mutant viruses as compared to WT (Fig 3A and 3B). Similar results were observed for both MDCK and A549 cells.

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**Table 1. Naturally occurring counts of WT and corresponding mutated amino acids for the listed mutant PR8 viruses.** Mutations in 7MB_ED and 8PK_BP are silent. All sequences were downloaded from the National Center for Biotechnology Information (NCBI) resource. Unique sequences were extracted from the alignment, and the total number of unique sequences is listed in the last column.

| mutant | protein | amino acid position | wild type sequence | occurrence | mutation | occurrence | total seqs |
|--------|---------|---------------------|--------------------|------------|----------|------------|------------|
| 7PK_ED | M1      | 242                 | K                  | 164        | H        | 1          | 201        |
| 7PK_ED | M2      | 12                  | R                  | 245        | S        | 1          | 302        |
| 7PK_ED | M2      | 13                  | N                  | 149        | T        | 39         | 302        |
| 8PK_ED | NS1     | 169                 | H                  | 402        | L        | 2          | 421        |
| 8PK_ED | NEP     | 12                  | I                  | 263        | L        | 1          | 280        |
| 8PK_ED | NEP     | 14                  | L                  | 26         | Q        | 24         | 280        |
| 8PK_BP_CP | NEP   | 22                  | E                  | 52         | G        | 140        | 280        |

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For segment 8 mutants, total RNAs were extracted from MDCK cells 3 h after infection (MOI = 5) at 37°C with WT, 8PK_ED, 8PK_BP, and 8PK_BP_CP. Measurements by qRT-PCR on 8PK_ED and 8PK_BP revealed that the relative amount of NEP mRNA was dramatically reduced as compared with WT, and even undetectable in 8PK_ED (Fig 3C). With the compensating base pair in 8PK_BP_CP, the lost NEP mRNA was restored to a level comparable with WT. The relative amount of the unspliced product, NS1 mRNA, was similar in WT and the mutant viruses (Fig 3C).

Mutations attenuate virus replication in cell culture

Multi-cycle growth kinetics at 37°C of WT and mutant viruses were measured in MDCK cells. WT and segment 7 mutants were also examined in A549 cells (Fig 4). Using MOI = 0.001, viruses with either 7MB_ED or 7PK_ED showed attenuated kinetics of replication in both cell lines, ranging from about 3–15 fold compared with WT virus at 12 and 24 h post-infection (Fig 4A and 4B). 8PK_BP mutation attenuated viral replication by about 26-fold in MDCK cells at 12 h post-infection, while 8PK_BP_CP and 8PK_ED showed similar growth kinetics as WT PR8 (Fig 4E).

With MOI = 0.0001, attenuation of the mutants was more dramatic. In MDCK cells, the viral titers of 7MB_ED and 7PK_ED were below detectable levels (20 FFU/ml) at 12 h post-infection, at least 58-fold attenuated relative to WT PR8 (Fig 4C). The differences were smaller in A549 cells, but were still more than 10-fold at 12 h post-infection (Fig 4D). 8PK_BP and
Fig 4. Viral replication was attenuated at 12 and 24 h post-infection in MDCK cells and A549 cells with MOI = 0.001 or 0.0001 at 37°C. Virus titers were quantified by immunofocus assay (FFU/ml). Data represent the means ± SDs of the results determined for triplicate wells. Replication of WT, 7MB_ED, and 7PK_ED were examined in (A) MDCK cells and (B) A549 cells with MOI = 0.001, and in (C) MDCK cells and (D) A549 cells with MOI = 0.0001. Replication of WT, 8PK_ED, 8PK_BP, and 8PK_BP_CP were examined in MDCK cells with (E) MOI = 0.001 and (F) MOI = 0.0001.

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8PK_ED were also attenuated with MOI = 0.0001, ranging from about 13–95 fold compared with WT virus at 12 or 24 h post-infection (Fig 4F). The attenuation with 8PK_BP was partially reversed by the compensatory mutant, 8PK_BP_CP, which replaces a UA pair with a CG pair expected to make coaxial stacking between two helices [19] more favorable [34–36]. All the mutated viruses were sequenced at 72 h post-infection. No revertants or additional mutations were observed (data not shown).

Discussion

Little is known about the biological function of RNA structure in influenza A virus [37], and no current vaccines or therapeutics are designed to target viral RNA structures. Conserved RNA secondary structures throughout protein coding regions of influenza A mRNAs were predicted by bioinformatics using a combination of thermodynamics and comparative sequence analysis, including suppression of synonymous codon usage [18, 19]. Structures predicted in mRNAs of segments 7 and 8 (Figs 1A, 1C and 2A) are most exciting because they locate near or containing splice sites. This suggests that these conserved RNA structures may be important for the splicing of segments 7 and 8 mRNAs. Enzymatic and chemical mapping, and nuclear magnetic resonance (NMR) provided experimental evidence for the predicted RNA structures [19–21]. In this work, mutational evidence is provided that is consistent with biological function for three of these structures. Sequences with only one or two mutations out of the 13.5 kb genome inhibited viral mRNA splicing and attenuated viral replication in cell culture. Evidently, the ensemble defect approach provides an efficient method to test for function of conserved structures.

The multi-branch loop and pseudoknot/hairpin region in segment 7 mRNA have function

An ensemble defect program [22] was used to design constructs, 7MB_ED and 7PK_ED, with 2-nt mutations predicted to misfold the multi-branch loop and pseudoknot/hairpin regions, respectively, in segment 7 mRNA (Fig 1). Both mutants increased the relative abundance of M1 mRNA compared with WT, while M2 mRNA levels remained the same (Fig 3A and 3B). This suggests that these conserved RNA structures affect the efficiency of segment 7 splicing, but a feedback loop regulates the amount of spliced mRNA. When production of M2 mRNA is inhibited, the virus may produce more M1 mRNA to ensure that enough M2 mRNA and thus enough essential M2 protein is produced. Both mutants also similarly reduced viral titers in both MDCK and A549 cells, with results most dramatic at 12 h post-infection at the lower MOI of 0.0001 (Fig 4A–4D).

It is noteworthy that 7MB_ED and 7PK_ED have similar effects on M1 and M2 mRNA levels, and on viral replication. The mutations in 7MB_ED are silent, whereas those in 7PK_ED change two amino acids in M2 protein and one in M1 protein (Table 1). The results suggest both mutants affect splicing to a similar level regardless of amino acid changes. The changes in conserved RNA structures are most likely responsible for the observed phenotype. Thus, the identified multi-branch loop near the 5’ splice site and the pseudoknot/hairpin equilibrium at the 3’ splice site apparently play a role in the regulation of mRNA splicing.

The pseudoknot/hairpin region in segment 8 has function

Two mutants were constructed to change the structure of 8PK (Fig 2), and both exhibited a similar phenotype. The 8PK_ED (Fig 2A and 2B) and 8PK_BP (Fig 2C) mutants greatly reduced the amount of spliced mRNA (Fig 3C). At 37°C, using MOI = 0.001, 8PK_ED showed similar growth kinetics as WT virus, while 8PK_BP was attenuated at 12 h post-infection.
At 37°C with MOI = 0.0001, the replication of both 8PK_ED and 8PK_BP were attenuated by 13–95 fold at 12 and 24 h post-infection (Fig 4F). 8PK_BP_CP, which compensated the mutation in 8PK_BP, restored WT splicing, and increased the viral titers relative to 8PK_BP. With MOI = 0.001, the viral titer of 8PK_BP_CP was increased to a level comparable with WT while with MOI = 0.0001, the viral titer of 8PK_BP_CP relative to 8PK_BP was increased by about 6 and 17-fold at 12 and 24 h post-infection, respectively. Because segment 7 mutants have similar effects in MDCK and A549 cells, segment 8 mutants were only examined in MDCK cells.

The results suggest that the phenotypes observed with the mutants in segment 8 are due to changes in RNA structure. In the pseudoknot conformation, the U483-A522 pair is expected to be coaxially stacked with the adjacent U504-G523 pair [19]. 8PK_BP disrupts U483-A522, and thus the coaxial stacking, which may be important for regulating splicing. 8PK_BP_CP restores the observed phenotype because the mutant restores the base pairing between nt 483 and nt 522. A mutation that disrupts the canonical U483-A522 pair was observed in highly pathogenic H5N1 viruses, and caused the equilibrium to shift towards the hairpin conformation [19].

Amino acid changes are not expected to have large effects

In 7PK_ED, 8PK_ED, and 8PK_BP_CP, the mutations of the mRNA sequences also produce local mutations in amino acid sequences. The naturally occurring counts of the WT and the corresponding mutated amino acids for these three mutants are summarized in Table 1.

For 7PK and 8PK regions, the peptide motifs of M1, M2, and NEP corresponding to the conserved RNA structures are all in largely unstructured conformations [38–45], thus expected to be tolerant to mutations. The 7PK region corresponds to part of the C terminal domain of M1, which is known to be extended and partially flexible on the basis of circular dichroism and bioinformatics [39, 41–43]. The 7PK region also corresponds to part of the ectodomain of M2, which exists in a dynamic conformation [38, 44, 45]. The crystal structure of the M2 ectodomain in complex with a protective monoclonal antibody suggests that the M2 ectodomain may have different conformations, including a curled structure that can induce binding with specific antibodies [46]. Neither of the M2 WT amino acids, R12 and N13, mutated in 7PK_ED had direct contacts with the antibody, but they facilitated polar interactions between M2 and the antibody (Fig 5A). S12 and T13 in 7PK_ED may facilitate similar interactions. The similar phenotypes observed for 7PK_ED and the silent mutations in 7MB_ED (Figs 3A, 3B and 4A–4D) are consistent with the assumption that changes in protein sequence are not responsible for the phenotypes.

The 8PK region corresponds to the N terminal domain of the nuclear export protein, NEP, which exists in a highly mobile and exposed state [40]. This flexibility may be important for the recognition of substrate proteins by the nuclear export machinery. The peptide motifs of NS1 corresponding to the RNA sequences mutated in 8PK are in an α-helical region [47–51] (Fig 5B). H169 locates at the end of the α-helix, thus mutation at this position has minimal effects in changing the propensities of forming α-helices [52]. It would be surprising if the mutated amino acids make a strong contribution to the observed phenotypes, but this cannot be ruled out.

Potential implications for developing safer vaccine

The temperature-sensitive (ts) and attenuated (att) phenotype of LAIV is imparted by five mutations within the viral replicative machinery: PB2 N265S; PB1 K391E, D581G and A661T; and NP D34G [8, 53]. Although this vaccine has an acceptable safety profile and is in general more effective than IIV, it is only recommended for use in healthy immunocompetent people because of possible complications in the very young and in patients with pre-existing conditions [54, 55].
Fig 5. (A) 3D structure of influenza M2 ectodomain in complex with monoclonal antibody Fab65 [46]. PDB ID: 4N8C. The light chain (light blue) and heavy chain (orange) of Fab65 are shown in cartoon presentation. R12 and N13 are labeled and shown in stick presentation. 7PK_ED has mutations R12S and N13T. The other residues of M2 ectodomain (dark blue) are shown in ribbon presentation. (B) 3D structure of NS1 C-terminal region [47]. PDB ID: 2GX9. The NS1 monomer (magenta) is shown in cartoon presentation. H169 is labeled and shown in stick presentation. 8PK_ED has mutation H169L.

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The mutations identified in this work that attenuated virus replication were made in segments 7 and 8, rather than in the segments currently mutated in LAIV. This suggests such mutations could potentially be introduced to LAIV to further decrease the virulence of the vaccine, and thus increase its safety profile [56]. Also, these mutations have the potential to serve as replacements if influenza evolves revertants to some currently used mutations.

**Potential implications for developing new treatments**

Because the conserved RNA structures are functionally important, oligonucleotides and small molecules targeting them may be designed as potential therapeutics against influenza A virus. For example, the M1 and M2 proteins coded by segment 7 are both essential, so inhibiting splicing is a potential therapeutic strategy. In general, a given mRNA or vRNA will produce many copies of protein, so therapeutic options targeting RNA structures would have to inhibit fewer copies of target than therapeutics targeting proteins.

**Refolding RNA structure by designing minimal mutations predicted to misfold structure by maximizing ensemble defects can be an efficient way to test regions of sequence for phenotype**

This is the first time that a program maximizing ensemble defects [22] was used to design sequences to refold conserved RNA structures for functional studies. As few as 2-nt mutations were required to test for a phenotype. This method provides an alternative to the traditional methods that destabilize specific base pairs. The method may also be applicable to unstructured regions because tight folding of an RNA structure in a mutant could inhibit dynamics and/or protein binding. Thus, it opens new avenues for targeting RNA sequences and structures for the development of safer vaccines and of therapeutics to control infections.

**Supporting Information**

S1 Fig. Multi-cycle growth kinetics of WT and mutant viruses plotted in log-scale. (DOCX)

S1 Table. Oligonucleotide primers used for qRT-PCR to amplify virus-specific and host-specific mRNAs. (DOCX)

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**Author Contributions**

Conceived and designed the experiments: TJ LMS DHT. Performed the experiments: TJ AN SFB. Analyzed the data: TJ. Contributed reagents/materials/analysis tools: LMS DHT. Wrote the paper: TJ LMS DHT.

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