The effect of respiratory viruses on immunogenicity and protection induced by a candidate universal influenza vaccine in mice

Janelle Rowell\(^{1\text{a}}\), Chia-Yun Lo\(^{1}\), Graeme E. Price\(^{1}\), Julia A. Misplon\(^{1}\), Roberta L. Crim\(^{2}\), Priyanka Jayanti\(^{2\text{b}}\), Judy Beeler\(^{2}\), Suzanne L. Epstein\(^{1\text{*}}\)

1 Office of Tissues and Advanced Therapies, US Food and Drug Administration, Silver Spring, Maryland, United States of America
2 Office of Vaccines Research and Review, US Food and Drug Administration, Silver Spring, Maryland, United States of America

\(^{\text{a}}\) Current address: The MITRE Corporation, McLean, Virginia, United States of America
\(^{\text{b}}\) Current address: Systems Planning and Analysis, Alexandria, Virginia, United States of America

\(^{*}\) suzanne.epstein@fda.hhs.gov

Abstract

Current approaches to influenza control rely on vaccines matched to viruses in circulation. Universal influenza vaccines would offer the advantage of providing broad protection against diverse strains of influenza virus. Candidate universal vaccines are developed using model systems, often testing in naïve animals. Yet the human population is not naïve, having varied immune histories that include exposure to viruses. We studied a candidate universal influenza vaccine (replication deficient adenoviruses expressing the conserved influenza A antigens NP and M2 [A/NP+M2-rAd]) given intranasally, the route previously shown to be most effective. To model recipients exposed to viruses, we used mice given rhinovirus (RV1B), respiratory syncytial virus (RSV-A2), influenza B virus, or influenza A virus before or after universal influenza vaccine. Vaccine performance was assessed by measuring immune responses to NP and M2, and monitoring weight loss and survival following influenza A virus challenge. Prior influenza A virus infection enhanced the response to the vaccine by priming to conserved influenza A antigens. RSV-A2 or RV1B had no effect on antibody responses to NP and M2 in serum. None of the viruses inhibited the ability of the vaccine to protect against influenza A virus challenge. The study demonstrates that the usefulness of this universal vaccine is not confined to the immunologically naïve and supports possible use in a human population with a varied history of respiratory infections.

Introduction

Universal influenza vaccines have the potential to reduce the disease burden of seasonal and pandemic influenza. We have developed a candidate universal vaccine based on conserved influenza A virus (IAV) antigens nucleoprotein (A/NP) and matrix 2 (M2). Our previous studies demonstrated that DNA priming followed by boosting with a mixture of recombinant
adenoviruses expressing A/NP and M2 (A/NP+M2-rAd) [1, 2] or a single intranasal dose of A/NP+M2-rAd [3, 4] protect naïve animals against subsequent IAV challenge of diverse strains and subtypes, preventing death and severe weight loss.

Preclinical testing of candidate vaccines in animal models typically uses naïve animals. However, vaccines for human use would be administered to individuals previously exposed to a wide range of antigens, including infections and other vaccines. In an effort to generate models that more closely recapitulate adult human immune responses, mouse models using a variety of prior immune stimuli have been developed [5, 6]. One study showed that sequential viral and parasitic infections alter the mouse immune system, resulting in responses more closely resembling those of adult humans [6]. Other work evaluating sequential infections has identified cross-protection between viruses, which is termed heterologous immunity [7]. In this scenario, T-cells primed by the first pathogen provide cross-protection against a subsequent differing pathogen; the cross-protection is not necessarily reciprocal [8]. In this way, sequential infections with various pathogens can alter the T-cell memory pool and increase or decrease subsequent responses to other pathogens [9, 10]. Prior infection history may also affect progression of disease caused by other viruses. For example, influenza virus infection protects mice against RSV-induced lung pathology [11], while latent infection with mouse herpesvirus-68 protects against IAV infection [12]. In some cases, instead of improving outcomes, a prior infection with one virus can lead to worse outcomes following infection with a second virus, despite contributing to clearance [9].

In humans, the influence of previous or ongoing infections on subsequent immune responses has been investigated for various viruses and other pathogens [13–15]. For instance, cytomegalovirus infection may influence immune responses to influenza [16]. Similarly, T-cell responses to influenza virus epitopes can overlap with reactivity to hepatitis C virus [17] or Epstein-Barr virus [18–20]. The sequence of exposure to multiple IAV infections may also influence immune responses and outcomes. Studies suggest immune imprinting occurs with the first influenza virus encountered [21–23], influencing susceptibility to different IAV subtypes seen later in life [24].

Responses to vaccines can also be influenced by prior infections. Infections initiated early in life may alter the response to subsequent vaccinations, possibly reducing the ability to respond to conventional vaccines [15, 25–27]. We previously demonstrated that vaccination history influences performance of our universal influenza vaccine in mice, resulting in enhancement or partial inhibition of universal vaccine-mediated protection, depending on the nature of the previous vaccines used [28]. Thus, it may be important to consider immune history when evaluating new vaccines.

In the human population, it would not be feasible to catalogue an individual’s every infection and then assess the impact on vaccination. It would also be difficult to model the lifelong sequence of viral infections, which is unique to each individual. However, the impact of previous infections can be studied in animal models using examples of common pathogens to provide a more realistic model than naïve animals alone. In the present study, we analyze the effects of acute respiratory viral infection on the performance of a universal influenza vaccine, including protection from IAV challenge and immune responses to vaccine antigens.

**Materials and methods**

**Viruses**

Human rhinovirus 1B, strain B632 (RV1B) was obtained from American Type Culture Collection (ATCC, Manassas, VA, USA). Virus was amplified and purified as previously described.
Briefly, RV1B was amplified in H1 HeLa cells (ATCC, CRL-1958). Cells were lysed by freeze-thaw, and then RV1B was precipitated using polyethylene glycol 6000. Virus was purified and concentrated using a centrifugal filtration device (Amicon Ultra 15 mL Filters (100,000 NMWL), MilliporeSigma, Burlington, MA). Fifty percent tissue-culture infectious dose (TCID<sub>50</sub>) was determined by titration in H1 HeLa cells. Respiratory syncytial virus, strain A2 (RSV-A2) was obtained from ATCC, then grown and prepared as previously described [30]. RV1B [31] and RSV-A2 [32] have been demonstrated to replicate in the respiratory tracts of mice.

Influenza A and B virus strains used were as follows: mouse-adapted A/Fort Monmouth/1/47 (H1N1) (A/FM) was provided by Earl Brown, University of Ottawa [33], A/Udorn/307/72 (H3N2) (A/Udorn) and B/Ann Arbor/1/86 (B/Ann Arbor) were obtained from Brian Murphy, National Institute of Allergy and Infectious Disease, National Institutes of Health. Viruses were prepared using embryonated hen’s eggs or lung homogenates of infected mice, as previously described [34].

**Recombinant adenoviral vaccines**

Replication-deficient (E1 and E3 deleted) recombinant adenovirus-5 (rAd) vectors expressing conserved IAV antigens A/NP or M2 have been previously described [35, 36]. A recombinant adenoviral vector expressing influenza B virus nucleoprotein (B/NP-rAd) [36] was used as a specificity control because it confers no protection against IAV challenge.

**Mice**

Female BALB/cAnNCR (BALB/c) mice were acquired from Charles River Laboratories (Ashburn, VA). Mice were 8–10 weeks of age at the beginning of experiments. All animal experiments were performed under ABSL-2 conditions in a facility accredited by the Association for Assessment and Accreditation of Laboratory Animal Care (AAALAC). Shelter, food and water were supplied in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. Caging was in filter top microisolator caging with filtered air and ad libitum access to reverse osmosis filtered water and to LabDiet Iso pro 3000 Irradiated #25 feed (St. Louis, MO). Environmental enrichment was supplied to all animals. The protocol (protocol number 1991–06) was approved by the FDA White Oak Campus Animal Care and Use Committee. In influenza challenge experiments causing disease, analgesics were not used to avoid interference in immune responses, but distress was reduced by using a 25% weight loss humane endpoint, and any mice reaching that endpoint were euthanized. Animals were monitored daily with increased monitoring at a minimum of twice daily during challenge infections. Euthanasia before tissue harvest was by ketamine/xylazine overdose, while euthanasia due to body weight loss or termination of study was by carbon dioxide inhalation in a chamber where the carbon dioxide was from a cylinder source delivered by Euthanex equipment (Palmer, PA).

**Respiratory infections**

For *in vivo* studies, mice received an intranasal dose of $10^4$ TCID<sub>50</sub> A/Udorn or $10^5$ TCID<sub>50</sub> B/Ann Arbor in 50 µL PBS; $5 \times 10^5$ TCID<sub>50</sub> RSV-A2 in 50 µL EMEM (Mediatech, Manassas, VA) containing 1% FBS, 100 mM MgSO<sub>4</sub>, and 50 mM HEPES [30]; $2 \times 10^6$ TCID<sub>50</sub> or $2 \times 10^7$ TCID<sub>50</sub> RV1B in 50 µL PBS. For RV1B, the two different doses both elicited immune responses in mice; the lower dose was used in all but one of the animal groups as noted in the text.
Vaccination and challenge
Mice were immunized intranasally under isoflurane anesthesia with $10^{10}$ virus particles (vp) of B/NP-rAd, or with A/NP+M2-rAd (a mixture of $5 \times 10^9$ vp A/NP-rAd and $5 \times 10^9$ vp M2-rAd). Four weeks later mice were challenged with A/FM, using doses noted in figure legends, and monitored for body weight and survival. As mentioned above, 25% weight loss was used as a humane endpoint, and any mice reaching that endpoint were euthanized.

T-cell responses to RV1B
Lung cells were re-stimulated with $10^6$ TCID$_{50}$ RV1B and interferon-γ response was determined by enzyme-linked immunospot (ELISPOT) as described previously [35].

Antibody responses to RSV-A2
Pre-immune sera and immune sera (three weeks following infection) were obtained from mice. Serum IgG antibodies to RSV nucleoprotein (RSV-N) were assessed by luciferase immunoprecipitation system (LIPS) using Renilla luciferase-tagged RSV-A2 nucleoprotein as previously described [37]. Serum samples from each animal experiment were tested in a single assay. The cutoff for a positive result was calculated for each assay based on 5 standard deviations above the mean value for pre-immune sera in that assay.

Immune responses to influenza antigens
Three weeks after mice received A/NP+M2-rAd, lung and spleen cells from individual mice were assessed for IFN-γ production in response to peptides as follows: the dominant NP CD8 epitope in BALB/c mice NP$_{147-155}$ (NP147), the consensus sequence of the M2 ectodomain M2e$_{2-24}$ (M2e), and control SARS M$_{209-221}$ (SARS) by ELISPOT as previously described [2, 35]. Sera were tested for IgG antibodies to influenza A/NP, B/NP, and M2e by ELISA using plates coated with M2e peptide or recombinant NP protein from strain A/PR/8/34 or B/Ann Arbor [1, 2].

Statistical analysis
Statistical analyses were performed using SigmaPlot (Systat Software, San Jose, CA, USA). Survival data were analyzed by the log-rank test with pairwise comparisons using the Holm-Sidak test. Analysis of body weight following challenge was performed using One-Way ANOVA at a time-point (shown by an arrow in figures) when 100% survival was observed for all groups. Post hoc analyses used the Holm-Sidak method or Dunn’s method with the A/NP+M2-rAd group as the control. ELISPOT data were analyzed by two-way ANOVA or t-test, as stated in the figure legends. Multiple comparisons were made using the Holm-Sidak method with the A/NP+M2-rAd group as the pre-determined control for animal groups receiving different inocula, and SARS peptide as the control for different stimulating peptides within a single group of animals. P values less than 0.05 were considered statistically significant and are noted in the figures.

Results
Prior influenza A infection improves immune responses to A/NP+M2-rAd
The A/NP+M2-rAd universal vaccine candidate has previously been shown to provide broad cross-protection after a single dose. The best protection resulted from intranasal immunization, a route which generates mucosal T and B cell immunity efficiently [3] and is used
currently in humans for live attenuated influenza vaccines. Vaccination to A/NP+M2 induces both antibody and T-cell responses, as reviewed in [38, 39]. Intranasal immunization induces antibodies and T cell responses, both systemically and locally in the respiratory tract. There are cytotoxic T lymphocytes in the lungs that can kill influenza virus-infected targets [40] and NP_{147-155}-specific pentamer+ or tetramer+ CD8 T cells in the lungs [2, 3,40].

We first studied the impact of prior influenza infection on subsequent performance of the universal vaccine candidate (Fig 1A). A/Udorn and B/Ann Arbor replicate well in mouse lungs [41], but do not induce significant clinical symptoms in mice. Three weeks later, mice had produced serum IgG antibodies specific to A/NP or B/NP (Fig 1B and 1C). Four weeks after infection, mice were immunized with A/NP+M2-rAd or B/NP-rAd intranasally, and an additional four weeks later challenged with influenza A/FM.

Serum IgG antibody responses to vaccine antigens A/NP and M2e were comparable between mice with no prior infection and mice with prior B/Ann Arbor infection, but enhanced in mice exposed to A/Udorn before receiving the universal vaccine (Fig 1D). Anti-A/NP but not anti-M2e antibody was induced by A/Udorn followed by B/NP-rAd. Consistent with elevated antibody responses, T-cell mediated immunity was enhanced in mice previously infected with A/Udorn as determined by IFN-γ ELISPOT (Fig 1E). In lung and spleen, T-cell responses to NP147 were approximately 5-fold greater in mice with a history of A/Udorn infection compared to mice with no prior infection, while responses to M2e were not elevated. In contrast, mice previously exposed to B/Ann Arbor had T-cell responses to NP147 and M2e in lung and spleen similar to those observed in mice with no previous exposure.

Weight loss and survival curves are shown in Fig 1F. In control mice with no prior infection history, A/NP+M2-rAd alone protected from severe influenza disease, with 100% survival and minimal weight loss. Thus for survival as an endpoint, only inhibition but not enhancement can be assessed under these conditions. However, either enhancement or inhibition of vaccine-specific immune responses and weight loss would be detectable. Mice previously infected with A/Udorn followed by A/NP+M2-rAd immunization had minimal weight loss following challenge and 100% survived, an outcome as good as A/NP+M2-rAd in naïve mice. Infection with A/Udorn without the universal vaccine (A/Udorn followed by B/NP-rAd control) provided modest cross-protective immunity against the mismatched A/FM challenge virus. Mice exhibited significant weight loss and 40% survival. Prior exposure to B/Ann Arbor did not significantly affect outcomes of vaccination followed by influenza A challenge. Mice exposed to B/Ann Arbor and then A/NP+M2-rAd exhibited minimal weight loss following challenge, comparable to mice with no prior exposure, and all mice survived IAV challenge. Mice given B/Ann Arbor and the control immunization, B/NP-rAd, lost significant weight following A/FM challenge and all mice succumbed.

**RSV-A2 or RV1B viruses do not affect vaccine protection**

We next examined the effect of two common respiratory pathogens, RSV-A2 and RV1B, on the performance of the universal vaccine. One month prior to immunization with A/NP+M2-rAd, mice received intranasal inoculations of 5 x 10^5 TCID_{50} RSV-A2 or 2 x 10^7 TCID_{50} RV1B (Fig 2A). These viruses have previously been shown to replicate in mouse lungs [31, 42]. To confirm exposure, immune responses to each virus were assessed. In mice exposed to RSV-A2, we detected serum antibodies to RSV-A2 nucleoprotein by LIPS, but none in pre-immune serum (Fig 2B). Infection with RV1B caused increased frequency of IFN-γ secreting lung cells in response to RV1B virus in comparison to cells from RSV-A2-infected mice (Fig 2C). Four weeks after viral exposure, mice received A/NP+M2-rAd or B/NP-rAd. Immune
Fig 1. Infection with Influenza A or Influenza B prior to immunization with A/NP+M2-rAd. A) Study timeline provides details of experimental design. Sera collected 3 weeks after A/Udorn or B/Ann Arbor infection were tested for IgG antibodies by ELISA as described in Materials and Methods. B) Serum IgG antibodies to influenza A/NP. C) Serum IgG antibodies to influenza B/NP. n = 7 per group of mice infected with virus that matched the antigen on the plate, n = 3 for test of non-specific binding. D) Sera collected 2 weeks after A/NP+M2-rAd or B/NP-rAd were tested by ELISA as described in Materials and methods for IgG antibodies to influenza antigens A/NP or M2e. n = 10 per group. E) Three weeks after A/NP+M2-rAd or B/NP-rAd, lung and
spleen cells from individual mice were assessed by ELISPOT for IFN-γ production in response to peptides NP147, M2e and SARS as specificity control. n = 2 for B/NP-rAd, n = 3 for all other groups. ELISPOT data were analyzed by two-way ANOVA using the Holm-Sidak method with A/NP+M2-rAd as control. A/NP+M2-rAd was predetermined as the comparator rather than comparing all pairs of groups, as explained in Materials and Methods. Significant differences between animal groups given different inocula are noted in the figure with horizontal brackets. Within an animal group, peptide comparisons are to SARS peptide as control and if significant indicated with asterisks *P<0.05 vs SARS. F) Left panel, percent initial body weight following challenge with 5.6 x 10^5 TCID_{50} A/FM. Weight loss data were analyzed by One-Way ANOVA on the day indicated by the arrow. Post hoc testing was performed using the Holm-Sidak method to determine significant differences relative to A/NP+M2-rAd as the predetermined control group. The three groups B/NP-rAd, B/Ann Arbor+B/NP-rAd, and A/Udorn+B/NP-rAd all differed significantly from the A/NP+M2-rAd control group, P<0.001. Right panel, survival until death or 25% weight loss endpoint. n = 8 for B/NP-rAd, n = 10 for all other groups. Survival data were analyzed by log-rank test, using the Holm-Sidak method for pairwise comparisons. Regardless of prior infection, all groups receiving A/NP+M2-rAd were significantly different (P<0.05) from groups receiving B/NP-rAd, but not from other groups receiving A/NP+M2-rAd P<0.05 B/NP-rAd vs. B/Ann Arbor+B/NP-rAd, A/Udorn+B/NP-rAd.

https://doi.org/10.1371/journal.pone.0215321.g001
Fig 2. Infection with RV1B or RSV does not affect performance of A/NP+M2e-rAd vaccine. A) Study timeline. B) Serum RSV-A2 Nucleoprotein antibodies. C) Lung T-cell response to RV1B. D) Serum A/NP IgG antibodies. E) Lung and Spleen IFN-γ secreting cells per million. F) % Initial Body Weight and % Survival.

Fig 2. Infection with RV1B or RSV does not affect performance of A/NP+M2e-rAd vaccine. A) Study timeline. B) Serum antibodies of individual mice to RSV-A2-N were assessed by LIPS assay. The cut-off for a positive result is noted by the dotted line above the X-axis. C) T cell response of individual mice to RV1B infection was assessed by ELISPOT. Whole purified RV1B virus was used for re-stimulation. n = 3. Significant differences were determined by Two-Way ANOVA using the Holm-Sidak method for post hoc testing of pairwise comparisons. D) Serum IgG antibodies to influenza antigens A/NP and M2e were assessed by ELISA as
Discussion

Immune history may have an impact on responses to a subsequent infection or immunization. Prior research has extensively described the phenomenon of immune responses to prior infections altering the responses to subsequent infection with a different pathogen [13]. Thus, testing candidate vaccines in animals with previous exposures can provide additional information relevant to human vaccination.

Our study tested the influence of exposure to various respiratory viruses on the performance of a universal vaccine designed to stimulate an immune response to conserved influenza A antigens. We gave respiratory viruses 28 days before immunization (A/Udorn, B/Ann Arbor, RSV-A2, and RV1B), 7 days before immunization (RSV-A2 and RV1B), and 28 days after immunization (RSV-A2 and RV1B). Subtle inhibition of protection might not be detected under the conditions used, because the vaccine alone gave such a high degree of protection (i.e., 100% survival), but no reduction in protection by the universal vaccine was demonstrated for the viral respiratory infections. The experimental conditions used also would have missed enhancement of protection, so the model might underestimate the potential of a vaccine, and in humans the vaccine might outperform expectations derived from naïve animal models. Protection against influenza challenge was maintained not only following respiratory infection one month or one week earlier, but also in the case of intervening infection. In some cases, prior infection enhanced immune responses to the universal vaccine. For example, A/Udorn infection before vaccination boosted recall immune responses to the vaccine (Fig 1D and 1E).

Infections at a longer interval before vaccination might have a different impact than recent infections. Future studies can address longer time intervals, to determine the impact on vaccination at different points in the development of immune memory to a preceding infection. Different pathogens might present different cross-reactive epitopes; we chose examples to study. The impact of infections could also depend on host genetics, and the dose of infecting pathogen or vaccine. Inflammatory effects and tissue damage due to previous infection could also play a role. Future studies can address these points. The findings so far provide encouragement that the NP+M2 vaccine is effective not only in pathogen-free responders, but in the more realistic setting of responders experienced with respiratory infection.

According to standard recommendations, conventional influenza vaccines should not be administered if the intended recipient is ill, but those with recent infection remain eligible [43]. To model vaccination after recent infection, we assessed the performance of A/NP +M2-rAd given 7 days after respiratory infection. As determined by morbidity and survival following challenge, protection afforded by A/NP+M2-rAd was not diminished by respiratory viral infections occurring 7 days before immunization. We also observed comparable serum antibody and lung T-cell responses despite recent infections.
Fig 3. Recent infection with RV1B or RSV-A2 does not inhibit the performance of universal vaccine. A) Study timeline. B) RSV-N IgG antibodies measured in individual sera by LIPS assay. The dotted line indicates the cut-off for a positive result. C) IFN-γ T-cell response to RV1B virus in the lungs of individual mice (n = 3/group) measured by ELISPOT. Significant differences were determined by two-way ANOVA using the Holm-Sidak method for post hoc testing of pairwise comparisons. D) Serum antibody responses to A/NP (left) and to M2e (right) were assessed by ELISA. n = 10 per group. E) IFN-γ ELISPOT was performed as in Fig 1 stimulated by the indicated peptides, using lung (left) and spleen (right) cells. n = 1 for B/NP-rAd
Our findings demonstrate that prior infection with A/Udorn induces cross-protection, also termed heterosubtypic immunity. Heterosubtypic immunity has been described many times in the literature for mice and many other animal species [38], and is likely also induced in humans [44–47]. In this case A/Udorn (H3N2) partially protected against A/FM (H1N1) (40% of the mice survived challenge, Fig 1F). In contrast, A/NP+M2-rAd induced more potent immune responses, which were sufficient to protect all mice.

Antibody responses to M2e are weak and variable in mice following influenza virus infection or immunization with cold-adapted influenza virus [1, 4, 48–50]. Similarly, in humans, induction of an M2e-specific antibody response following influenza infection is highly variable [51–53]. In agreement with those findings, an M2e-specific antibody response was not induced by A/Udorn infection followed by B/NP-rAd immunization (Fig 1D). However, focused immunization, for example by M2-rAd, more effectively induces an antibody response. Previous work has shown that immunization strategies using engineered M2 conjugates or expression vectors are effective against influenza challenge [35, 50, 54]. While our previous studies showed that M2-rAd alone provides protection, A/NP+M2-rAd provides protection superior to either component alone [1, 3].

For adenovirus-vectored vaccines and gene therapies, interference by pre-existing immunity to the vector can indeed be a concern [55, 56]. Immunity to adenovirus serotype 5 (Ad5) is prevalent in the human population and can be a barrier to subsequent use of Ad5-based vectors, such as the A/NP+M2-rAd universal vaccine candidate used in this study. One way to overcome this barrier is to use an adenovirus vector to which humans are not exposed. In previous work, we used PanAd3, a nonhuman primate adenoviral vector in the same Ad species C as Ad5 [57], to construct a universal vaccine candidate expressing conserved influenza A antigens NP and M1. Humans have very little or no serum antibody to PanAd3, and a universal vaccine with this backbone protected mice from influenza challenge [58]. Also, mucosal administration of rAd (i.n. or aerosol) appears to circumvent blocking by prior immunity in some cases [59, 60].

The present study focused on viruses causing acute respiratory infections. Chronic infections with viruses, bacteria, and parasites can also have a major influence on host immune responses to subsequent vaccination [15, 25–27]. Given the high rate of pre-existing chronic infection in many parts of the world, their impact on immunizations is of major importance. Expanding upon the cases studied here, future research could examine vaccine performance in models of chronic infection.

Depending on virus and timing of infection, immune responses to the universal vaccine were unchanged, enhanced or modestly reduced compared to responses in mice without prior viral infection. Ultimately, the viral infections we tested did not abrogate protection elicited by the universal vaccine. These results are promising that the vaccine may perform well in humans despite widespread immunity to common respiratory viruses.
Fig 4. Intervening infection with RV1B or RSV-A2 does not affect universal vaccine protection. A) Study timeline. B) Intervening RSV-A2 infection was confirmed by antibody responses to RSV-N by LIPS assay as in Fig 2. In this test, the cut-off for a positive result is indicated by the dotted line above the X-axis. C) Intervening RV1B infection was confirmed by T-cell responses assessed by ELISPOT as in Fig 2. n = 3. Significant difference was determined by t-test. D) T cell response to influenza antigens was assessed following rAd vaccination using lung cells from mice infected with RSV-A2 (left panel) or RV1B (right panel).
Acknowledgments

We thank Mary Belcher and Anthony Ferrine of the FDA Vivarium for expert animal care and operation of the facility. We thank Andrew Byrnes and Maryna Eichelberger for insightful reviews of the manuscript.

Author Contributions

Conceptualization: Suzanne L. Epstein.
Data curation: Janelle Rowell, Chia-Yun Lo, Julia A. Misplon.
Formal analysis: Janelle Rowell, Chia-Yun Lo, Graeme E. Price, Julia A. Misplon, Roberta L. Crim, Priyanka Jayanti, Judy Beeler.
Funding acquisition: Judy Beeler, Suzanne L. Epstein.
Investigation: Janelle Rowell, Chia-Yun Lo, Graeme E. Price, Julia A. Misplon, Roberta L. Crim, Priyanka Jayanti.
Methodology: Janelle Rowell, Chia-Yun Lo, Graeme E. Price, Julia A. Misplon, Priyanka Jayanti, Judy Beeler, Suzanne L. Epstein.
Project administration: Judy Beeler, Suzanne L. Epstein.
Resources: Judy Beeler, Suzanne L. Epstein.
Supervision: Judy Beeler, Suzanne L. Epstein.
Validation: Janelle Rowell, Chia-Yun Lo, Graeme E. Price, Julia A. Misplon, Roberta L. Crim, Priyanka Jayanti, Judy Beeler, Suzanne L. Epstein.
Visualization: Janelle Rowell, Chia-Yun Lo, Graeme E. Price, Julia A. Misplon, Roberta L. Crim, Priyanka Jayanti, Judy Beeler, Suzanne L. Epstein.
Writing – original draft: Janelle Rowell.
Writing – review & editing: Janelle Rowell, Chia-Yun Lo, Graeme E. Price, Julia A. Misplon, Roberta L. Crim, Priyanka Jayanti, Judy Beeler, Suzanne L. Epstein.

References

1. Lo C-Y, Wu Z, Misplon JA, Price GE, Pappas C, Kong WP, et al. Comparison of vaccines for induction of heterosubtypic immunity to influenza A virus: Cold-adapted vaccine versus DNA prime-adenovirus boost strategies. Vaccine. 2008; 26(17):2062–72. https://doi.org/10.1016/j.vaccine.2008.02.047 PMID: 18378366

2. Price GE, Soboleski MR, Lo CY, Misplon JA, Pappas C, Houser KV, et al. Vaccination focusing immunity on conserved antigens protects mice and ferrets against virulent H1N1 and H5N1 influenza A
3. Price GE, Soboleski MR, Lo CY, Misplon JA, Quirion MR, Houser KV, et al. Single-Dose Mucosal Immunization with a Candidate Universal Influenza Vaccine Provides Rapid Protection from Virulent H5N1, H3N2 and H1N1 Viruses. PLoS ONE. 2010; 5(10):e13162. https://doi.org/10.1371/journal.pone.0013162 PMID: 20976273

4. Soboleski MR, Gabbard JD, Price GE, Misplon JA, Perez DR, et al. Cold-adapted influenza and recombinant adenovirus vaccines induce cross-protective immunity against pH1N1 challenge in mice. PLoS ONE. 2011; 6(7):e21937. https://doi.org/10.1371/journal.pone.0021937 PMID: 21789196

5. Beura LK, Hamilton SE, Bi K, Schenkel JM, Odumade OA, Casey KA, et al. Normalizing the environment recapitulates adult human immune traits in laboratory mice. Nature. 2016; 532(7600):512–6. https://doi.org/10.1038/nature17655 PMID: 27096360

6. Reese TA, Bi K, Kambal A, Filali-Mouhim A, Beura LK, Burger MC, et al. Sequential Infection with Common Pathogens Promotes Human-like Immune Gene Expression and Altered Vaccine Response. Cell Host Microbe. 2016; 19(5):2825–38. https://doi.org/10.1016/j.chom.2016.04.003 PMID: 27107939.

7. Welsh RM, Selin LK. No one is naive: The significance of heterologous T-cell immunity. Nature Reviews Immunology. 2002; 2(6):417–26. https://doi.org/10.1038/nri820 PMID: 12093008

8. Welsh RM, Selin LK. Heterologous immunity between viruses. Immunol Rev. 2010; 235(1):244 –66. https://doi.org/10.1111/j.0105-2896.2010.00897.x PMID: 20536568

9. Che JW, Daniels KA, Selin LK, Welsh RM. Heterologous immunity and persistent Murine Cytomegalovirus Infection. J Virol. 2011; 85(23):12953–60. https://doi.org/10.1128/JVI.01321-10 PMID: 21611074

10. Welsh RM, Selin LK. Heterologous immunity between viruses. Immunol Rev. 2010; 235(1):244 –66. https://doi.org/10.1111/j.0105-2896.2010.00897.x PMID: 20536568

11. Welsh RM, Selin LK. Heterologous immunity and persistent Murine Cytomegalovirus Infection. J Virol. 2011; 85(23):12953–60. https://doi.org/10.1128/JVI.01321-10 PMID: 21611074
23. Miller MS, Gardner TJ, Krammer F, Aguado LC, Tortorella D, Basler CF, et al.Neutralizing antibodies against previously encountered influenza virus strains increase over time: a longitudinal analysis. Sci Transl Med. 2013; 5(198):198ra07. https://doi.org/10.1126/scitranslmed.3006637 PMID: 23946196

24. Gostic KM, Ambrose M, Worobey M, Lloyd-Smith JO. Potent protection against H5N1 and H7N9 influenza via childhood hemagglutinin imprinting. Science. 2016; 354(6313):722–6. https://doi.org/10.1126/science.aag1322 PMID: 27846599

25. Frasca D, Díaz A, Romero M, Landin AM, Blomberg BB. Cytomegalovirus (CMV) seropositivity decreases B cell responses to the influenza vaccine. Vaccine. 2015; 33(12):1433–9. https://doi.org/10.1016/j.vaccine.2015.01.071 PMID: 25659271

26. Wald A, Selke S, Magaret A, Boechk M. Impact of human cytomegalovirus (CMV) infection on immune response to pandemic 2009 H1N1 influenza vaccine in healthy adults. J Med Virol. 2013; 85(9):1557–60. https://doi.org/10.1002/jmv.23642 PMID: 23852679

27. Muyanja E, Ssemaganda A, Ngauv P, Cubas R, Perrin H, Srinivasan D, et al. Immune activation alters cellular and humoral responses to yellow fever 17D vaccine. J Clin Invest. 2014; 124(7):3147–58. https://doi.org/10.1172/JCI75429 PMID: 24911151

28. Rowell J, Lo CY, Price GE, Misplon JA, Epstein SL, García M. Conventional influenza vaccines influence the performance of a universal influenza vaccine in mice. Vaccine. 2018; 36(7):1008–15. https://doi.org/10.1016/j.vaccine.2017.11.065 PMID: 29249542.

29. Bartlett NW, Singanayagam A, Johnston SL. Mouse models of rhinovirus infection and airways disease. Methods Mol Biol. 2015; 1221:181–8. https://doi.org/10.1007/978-1-4939-1571-2_14 PMID: 25261315

30. Feldman SA, Audet S, Beeler JA. The fusion glycoprotein of human respiratory syncytial virus facilitates virus attachment and infectivity via an interaction with cellular heparan sulfate. J Virol. 2000; 74(14):6442–7. PMID: 10864656

31. Bartlett NW, Walton RP, Edwards MR, Aniscenko J, Caramori G, Zhu J, et al. Mouse models of rhinovirus-induced disease and exacerbation of allergic airway inflammation. Nat Med. 2008; 14(2):199–204. https://doi.org/10.1038/nm1713 PMID: 18246079

32. Mejias A, Chavez-Bueno S, Rios AM, Saavedra-Lozano J, Fonseca Aten M, Hatfield J, et al. Anti-respiratory syncytial virus (RSV) neutralizing antibody decreases lung inflammation, airway obstruction, and airway hyperresponsiveness in a murine RSV model. Antimicrob Agents Chemother. 2004; 48(5):1811–22. https://doi.org/10.1128/AAC.48.5.1811-1822.2004 PMID: 15105140

33. Smeenk CA, Brown EG. The influenza-virus variant A/FM/1/47-MA possesses single amino-acid replacements in the hemagglutinin, controlling virulence, and in the matrix protein, controlling virulence as well as growth. J Virol. 1994; 68(1):530–4. PMID: 8254767

34. Benton KA, Misplon JA, Lo C-Y, Brutkiewicz RR, Prasad SA, Epstein SL. Heterosubtypic immunity to influenza A virus in mice lacking either IgA, all Ig, NKT cells, or gd T cells. J Immunol. 2001; 166:7437–45. PMID: 11390496

35. Tompkins SM, Zhao ZS, Lo C-Y, Misplon JA, Liu T, Ye ZP, et al. Matrix protein 2 vaccination and protection against influenza viruses, including subtype H5N1. Emerg Infect Dis. 2007; 13(3):426–35. https://doi.org/10.3201/eid1303.070125 PMID: 17552096

36. Epstein SL, Kong WP, Misplon JA, Lo C-Y, Tumpey TM, Xu L, et al. Protection against multiple influenza A subtypes by vaccination with highly conserved nucleoprotein. Vaccine. 2005; 23(46–47):5404–10. https://doi.org/10.1016/j.vaccine.2005.04.047 PMID: 16011865

37. Kumari S, Crim RL, Kulkarni A, Audet SA, Mdluli T, Murata H, et al. Development of a luciferase immunoprecipitation system assay to detect IgG antibodies against human respiratory syncytial virus nucleoprotein. Clin Vaccine Immunol. 2014; 21(3):383–90. https://doi.org/10.1128/CVI.00594-13 PMID: 24403526

38. Epstein SL, Price GE. Cross-protective immunity to influenza A viruses. Expert Rev Vaccines. 2010; 9(11):1325–41. https://doi.org/10.1586/erv.10.123 PMID: 21087110

39. Epstein SL. Universal influenza vaccines: Progress in achieving broad cross-protection in vivo. Am J Epidemiol. 2018; 187(12):2603–14. https://doi.org/10.1093/aje/kwy145 PMID: 30084906.

40. García M, Misplon JA, Price GE, Lo CY, Epstein SL. Age Dependence of Immunity Induced by a Candidate Universal Influenza Vaccine in Mice. PLoS One. 2016; 11(4):e0153195. https://doi.org/10.1371/journal.pone.0153195 PMID: 27055234

41. Treanor JJ, Tierney EL, Zebedee SL, Lamb RA, Murphy BR. Passively transferred monoclonal antibody to the M2 protein inhibits influenza A virus replication in mice. J Virol. 1990; 64(3):1375–7. PMID: 2304147

42. Openshaw PJ. The mouse model of respiratory syncytial virus disease. Curr Top Microbiol Immunol. 2013; 372:359–69. https://doi.org/10.1007/978-3-642-38919-1_18 PMID: 24362699.
43. Grohskopf LA, Sokolow LZ, Broder KR, Olsen SJ, Karron RA, Jernigan DB, et al. Prevention and Control of Seasonal Influenza with Vaccines. MMWR Recomm Rep. 2016; 65(5):1–54. https://doi.org/10.15585/mmwr.rr6505a1 PMID: 27560619.

44. McMichael AJ, Gotch FM, Noble GR, Beare PAS. Cytotoxic T-cell immunity to influenza. N Engl J Med. 1983; 309:13–7. https://doi.org/10.1056/NEJM198307073090103 PMID: 6602294

45. Cowling BJ, Ng S, Ma ES, Cheng CK, Wai W, Fang VJ, et al. Protective efficacy of seasonal influenza vaccination against seasonal and pandemic influenza virus infection during 2009 in Hong Kong. Clin Infect Dis. 2010; 51(12):1370–9. https://doi.org/10.1086/657311 PMID: 21067351

46. Sridhar S, Begom S, Bermingham A, Hoschler K, Adamson W, Carman W, et al. Cellular immune correlates of protection against symptomatic pandemic influenza. Nat Med. 2013; 19(10):1305–12. https://doi.org/10.1038/nm.3350 PMID: 24056771

47. Hayward AC, Wang L, Goonetilleke N, Fragaszy EB, Bermingham A, Copas A, et al. Natural T Cell-mediated Protection against Seasonal and Pandemic Influenza. Results of the Flu Watch Cohort Study. Am J Respir Crit Care Med. 2015; 191(12):1422–31. https://doi.org/10.1164/rccm.201411-1988OC PMID: 25844934

48. Kim MC, Lee JS, Kwon YM, OE, Lee YJ, Choi JG, et al. Multiple heterologous M2 extracellular domains presented on virus-like particles confer broader and stronger M2 immunity than live influenza A virus infection. Antiviral Res. 2013; 99(3):328–35. https://doi.org/10.1016/j.antiviral.2013.06.010 PMID: 23811283

49. Feng JQ, Zhang MX, Mozdzanowska K, Zharkova D, Hoff H, Wunner W, et al. Influenza A virus infection engenders a poor antibody response against the ectodomain of matrix protein 2. Virol J. 2006; 3:102. https://doi.org/10.1186/1743-422X-3-102 PMID: 17150104

50. Neirynck S, Deroo T, Saelens X, Vanlandschoot P, Jou WM, Fiers W. A universal influenza A vaccine based on the extracellular domain of the M2 protein. Nature Med. 1999; 5(10):1157–63. https://doi.org/10.1038/nm.3350 PMID: 10502819

51. Black RA, Rota PA, Gorodkova N, Klenk HD, Kendal AP. Antibody response to the M2 protein of influenza A virus expressed in insect cells. J Gen Virol. 1993; 74(11):143–6.

52. Zhong W, Reed C, Blair PJ, Katz JM, Hancock K. Serum Antibody Response to Matrix Protein 2 Following Natural Infection With 2009 Pandemic Influenza A(H1N1) Virus in Humans. J Infect Dis. 2014; 209(7):986–94. https://doi.org/10.1093/infdis/jit811 PMID: 24325965

53. Lo CY, Strobl SL, Dunham K, Wang W, Stewart L, Misplon JA, et al. Surveillance Study of Influenza Occurrence and Immunity in a Wisconsin Cohort During the 2009 Pandemic. Open Forum Infect Dis. 2017; 4(2):ox023. https://doi.org/10.1093/ofid/ox023 PMID: 28730155

54. Lee YN, Kim MC, Lee YT, Kim YJ, Kang SM. Mechanisms of Cross-protection by Influenza Virus M2-based Vaccines. J Infect Dis. 2014; 209(7):986–94. https://doi.org/10.1093/infdis/jit811 PMID: 24325965

55. Black RA, Rota PA, Gorodkova N, Klenk HD, Kendal AP. Antibody response to the M2 protein of influenza A virus expressed in insect cells. J Gen Virol. 1993; 74(11):143–6.

56. Sumida SM, Truitt DM, Kishko MG, Arthur JC, Jackson SS, Gorgone DA, et al. Neutralizing antibodies and CD8+ T lymphocytes both contribute to immunity to adenovirus serotype 5 vaccine vectors. J Virol. 2004; 78(6):2673–73. https://doi.org/10.1128/JVI.78.6.2666-2673.2004 PMID: 14990686

57. Colloca S, Barnes E, Folgori A, Ammendola V, Capone S, Cirillo A, et al. Vaccine vectors derived from a large collection of simian adenoviruses induce potent cellular immunity across multiple species. Sci Transl Med. 2012; 4(115):115ra2. https://doi.org/10.1126/scitranslmed.3002925 PMID: 22218691

58. Vitelli A, Quirion MR, Lo CY, Misplon JA, Grabowska AK, Pierantoni A, et al. Vaccination to conserved influenza antigens in mice using a novel Simian adenovirus vector, PanAd3, derived from the bonobo Pan paniscus. PLoS ONE. 2013; 8(3):e55435. https://doi.org/10.1371/journal.pone.0055435 PMID: 23536756

59. Croyle MA, Patel A, Tran KN, Gray M, Zhang Y, Strong JE, et al. Nasal Delivery of an Adenovirus-Based Vaccine Bypasses Pre-Existing Immunity to the Vaccine Carrier and Improves the Immune Response in Mice. PLoS ONE. 2008; 3(10):e3548. https://doi.org/10.1371/journal.pone.0003548 PMID: 18958172

60. Song K, Bolton DL, Wei CJ, Wilson RL, Camp JV, Bao S, et al. Genetic immunization in the lung induces potent local and systemic immune responses. Proc Natl Acad Sci U S A. 2010; 107(51):22213–8. https://doi.org/10.1073/pnas.1015536108 PMID: 21135247