Transmission-blocking strategies that slow the spread of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) and protect against coronavirus disease 2019 (COVID-19) are needed. We have developed an orally delivered adenovirus type 5–vectored SARS-CoV-2 vaccine candidate that expresses the spike protein. Here, we demonstrated that hamsters vaccinated by the oral or intranasal route had robust and cross-reactive antibody responses. We then induced a postvaccination infection by inoculating vaccinated hamsters with SARS-CoV-2. Orally or intranasally vaccinated hamsters had decreased viral RNA and infectious virus in the nose and lungs and experienced less lung pathology compared to mock-vaccinated hamsters after SARS-CoV-2 challenge. Naïve hamsters exposed in a unidirectional air flow chamber to mucosally vaccinated, SARS-CoV-2–infected hamsters also had lower nasal swab viral RNA and exhibited fewer clinical symptoms than control animals, suggesting that the mucosal route reduced viral transmission. The same platform encoding the SARS-CoV-2 spike and nucleocapsid proteins elicited mucosal cross-reactive SARS-CoV-2–specific IgA responses in a phase 1 clinical trial (NCT04563702). Our data demonstrate that mucosal immunization is a viable strategy to decrease SARS-CoV-2 disease and airborne transmission.

INTRODUCTION

The intramuscular severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) vaccines currently approved for clinical use are capable of protecting vaccinees from symptomatic infection, hospitalization, and death from coronavirus disease 2019 (COVID-19) (1–3). However, they do not completely prevent infection. mRNA-vaccinated individuals infected with the B.1.617.2 (Delta) and B.1.1.529 (Omicron) variants can shed viral RNA and infectious virus and potentially spread SARS-CoV-2 to others (4–6). The Omicron variant appears more capable of avoiding vaccine-induced immunity than the Delta variant (7) and caused a substantial winter 2021 surge in infections in the United States, creating an acute shortage of health care workers (8). Considering much of the world is underimmunized, including children, the possibility that a vaccinated individual with a postvaccination infection can spread SARS-CoV-2 to underimmunized family or community members poses a public health risk. There would be a substantial benefit to develop vaccines that protect against disease and reduce SARS-CoV-2 transmission from vaccinated to unvaccinated individuals.

Because the mucosal surface of the upper respiratory tract (URT) is the initial site of SARS-CoV-2 replication and primary site of infection (9), interventions that induce robust mucosal immune responses may have the greatest impact on reduction of SARS-CoV-2 transmission. We have created a replication-defective, shelf-stable oral adenoviral type 5 (Ad5) vector vaccine candidate expressing the spike protein from SARS-CoV-2 (r-Ad-S) that is designed to induce both systemic and mucosal immunity. More than 500 human participants have been administered these oral Ad5 vaccines, which have been well tolerated and able to generate robust humoral and cellular immune responses to the expressed antigens (10–13). Immune activation through the intestine may represent an important organ for oral immunization, as antibody-secreting plasmablasts and plasma cells can traffic from the gut to the nose, trachea, and lung (14, 15). Prior work in a human influenza challenge study with the same platform has shown an ability to limit viral RNA shedding of influenza virus (11), highlighting the utility of this vaccination strategy for respiratory viruses.

To study the potential impact of oral vaccination on transmission to naïve individuals, we used a hamster infection model and unidirectional air flow chambers. We vaccinated index hamsters with oral r-Ad-S, using intranasal r-Ad-S as a control for mucosal stimulation, intramuscular spike protein as a protein control, and oral phosphate-buffered saline (PBS) as a mock control. We then infected animals intranasally with a high titer of SARS-CoV-2 to replicate a postvaccination infection. One day after viral challenge, index hamsters were placed upstream of vaccine-naïve hamsters in a chamber that allowed airborne movement but not direct contact or fomite transmission. Here, we report the clinical and virological responses of both the vaccinated (SARS-CoV-2–infected) and naïve (SARS-CoV-2–exposed) hamsters. In addition, we present mucosal antibody data from participants in a phase 1 clinical trial using the same platform encoding the SARS-CoV-2 spike and nucleocapsid proteins. These data demonstrate that oral r-Ad-S immunization resulted in reduced disease and decreased SARS-CoV-2 transmission in a hamster model and can generate coronavirus cross-reactive, spike protein–specific immunoglobulin A (IgA) in the nose and mouths of humans.
RESULTS
Oral and intranasal r-Ad-S vaccination induced robust systemic and mucosal antibody responses

Index hamsters were immunized at weeks 0 and 4 with oral r-Ad-S, intranasal r-Ad-S (mucosal positive control), intramuscular spike protein, or mock (oral PBS) before SARS-CoV-2 challenge at week 7 (Fig. 1A). To determine immunogenicity of these vaccines, serum was collected at weeks 0, 3, and 6 after immunization. Bronchoalveolar lavage (BAL) fluid was collected upon necropsy (day 5 after inoculation; Fig. 1A). Oral and intranasal r-Ad-S–vaccinated groups had significantly higher spike protein–specific IgG antibody titers in serum at week 3 compared to mock-dosed hamsters (P = 0.0165, oral; P < 0.0001, intranasal); this was not true in intramuscular spike protein–vaccinated hamsters (Fig. 1B). Serum spike protein–specific IgG at 6 weeks after immunization was also cross-reactive to the Beta and Delta variants in all groups (Fig. 1C). Using a surrogate virus neutralizing test, we found that serum from intranasal r-Ad-S hamsters had a greater ability to block binding of SARS-CoV-2 spike protein to angiotensin-converting enzyme 2 (ACE-2) after the booster vaccination (week 6) compared to serum from mock-vaccinated hamsters (Fig. 1D). Serum anti–spike protein IgA antibodies increased after oral and intranasal r-Ad-S vaccination but not in intramuscular spike protein– or mock-vaccinated groups (Fig. 1E). As expected from our oral immunization platform, oral r-Ad-S–vaccinated hamsters demonstrated similar spike protein–specific IgA concentrations in BAL fluid compared to the intranasal r-Ad-S–positive control group, suggesting similar stimulation of mucosal immunity (Fig. 1F). There were no differences in serum or BAL IgA responses between intramuscular spike protein– and mock-vaccinated hamsters, suggesting that IgA was not induced by systemic immunization with the intramuscular spike protein. These data demonstrate that oral r-Ad-S– and intranasal r-Ad-S–vaccinated hamsters generated robust systemic and mucosal humoral immunity.

Oral and intranasal r-Ad-S vaccination accelerated SARS-CoV-2 viral RNA clearance and protected against disease in hamsters

In index animals, SARS-CoV-2 RNA from nasal swabs at 3 and 5 days after inoculation was lower in oral and intranasal r-Ad-S–vaccinated animals compared to mock-vaccinated animals; this was not true for intramuscularly vaccinated animals (Fig. 2A). Nasal swabs of index animals immunized with intranasal r-Ad-S had significantly lower median tissue culture infectious dose (TCID_{50}) values compared to mock animals on day 1 (P = 0.0377; Fig. 2B), but all groups still had detectable infectious virus. Oral and intranasal r-Ad-S–vaccinated index animals had significantly lower viral RNA loads in the lungs at terminal collection (day 5 after inoculation) when compared to mock-vaccinated animals (P = 0.0198, oral; P = 0.0135, intranasal), which was not true for intramuscular spike protein–vaccinated animals (Fig. 2C). All vaccinees had significantly reduced lung infectious viral loads compared to mock (P < 0.0001), with all appearing below the limit of detection at day 5 (Fig. 2D). These data demonstrated that oral and intranasal r-Ad-S vaccination decreased SARS-CoV-2 after infection and accelerated viral clearance.

During vaccination, but before SARS-CoV-2 inoculation, all animals were gaining weight at a similar rate (fig. S1). After SARS-CoV-2 inoculation in index hamsters, we observed significantly greater weight loss in unvaccinated hamsters [area under the curve (AUC) = 767 ± 2.5] compared to uninfected hamsters (AUC = 809.6 ± 4.0; P < 0.0001; Fig. 3A), a characteristic of disease in this model. Oral and intranasal r-Ad-S–vaccinated animals lost less weight by the termination of the study compared to mock-vaccinated animals (Fig. 3B). To quantify pulmonary inflammation, lung weights were measured, and lungs were scored for gross pathology. In index animals, lung weights (normalized to total body weight; Fig. 3C) and average gross pathology scores (Fig. 3D) were decreased in both oral and intranasal r-Ad-S–vaccinated groups when compared to mock-treated hamsters.

Oral and intranasal r-Ad-S vaccination limited SARS-CoV-2 transmission to unvaccinated, naïve hamsters leading to decreased clinical evidence of disease

As a test of transmissibility, unvaccinated naïve hamsters were exposed to vaccinated, SARS-CoV-2–infected index animals at a 1:4 ratio of index to naïve, where each vaccine exposure group has 4 index animals and 16 vaccine naïve animals. Exposure was performed by putting an index animal in a chamber (index chamber) connected to a second chamber containing the four naïve animals ( naïve chamber), separated by a 5-inch connecting chamber (fig. S2). Screens at either end of the connecting chamber prevented direct contact. Air was circulated in the index chamber by a fan and was pulled into the naïve chamber by a vacuum. Exposure of naïve hamsters to air flow produced by index hamsters was performed for 8 hours before moving each hamster to an individual cage. On day 1 after infection, the amount of viral RNA loads in all index animals was above 7.8 × 10^7 gene copies per swab (Fig. 2A).

In naïve hamsters exposed to the oral and intranasal r-Ad-S–immunized groups, SARS-CoV-2 RNA was significantly lower on days 1 and 3 compared to hamsters exposed to mock-immunized animals (P = 0.0075, oral day 1; P = 0.0001, intranasal day 1; P = 0.0367, oral day 3; P = 0.0001, intranasal day 3; Fig. 4A). The number of vaccine naïve animals with nasal swab viral RNA above a threshold of 1 × 10^5 gene copies was also determined. On day 1 after exposure of the index and naïve animals, there were 3 naïve hamsters (3 of 16) exposed to the oral r-Ad-S index group that were above the threshold compared to 11 mock-exposed hamsters (11 of 16; P = 0.011, Fisher’s exact test; Fig. 4A and table S1). The naïve hamsters exposed to intranasal r-Ad-S index hamsters had 0 (0 of 16) animals above the threshold, which was not significantly different from oral vaccination but was significantly lower than naïve animals exposed to airborne transmission from mock vaccination animals (P = 0.22 and P < 0.0001 by Fisher’s exact test, respectively; Fig. 4A and table S1). On day 3, in the group exposed to orally vaccinated hamsters, 11 (11 of 16) had nasal swab viral RNA concentrations above the threshold compared to 16 (16 of 16) for the group exposed to unvaccinated index hamsters (P = 0.043 by Fisher’s exact test; Fig. 4A and table S1). The naïve animals exposed to intranasally vaccinated index animals had 7 of 16 animals with nasal swab viral RNA above the threshold, which was not significantly different from the animals exposed to orally vaccinated index hamsters (P = 0.285 by Fisher’s exact test; Fig. 4A and table S1). These data demonstrate decreased SARS-CoV-2 transmission from oral and intranasal r-Ad-S–vaccinated index to naïve animals. On day 5, nasal viral RNA did not differ between naïve hamsters exposed to orally vaccinated index and control index hamsters. However, nasal swab and lung viral RNA (P = 0.0062 and P = 0.0077, respectively) and infectious virus (P = 0.002) were lower in hamsters exposed to intranasal
r-Ad-S index hamsters compared to those exposed to unvaccinated index hamsters (Fig. 4, B to D).

Although oral and intranasal r-Ad-S vaccination did not completely prevent transmission, the vaccination likely reduced the effective dose reaching the naïve animals. In a prior virus titrating experiment, we demonstrated that animals lost more weight and had increased pulmonary inflammation when they received higher titers of SARS-CoV-2 inoculum, suggesting that the severity of disease acquired is dependent on the original infectious dose (fig. S3). Similar dose-dependent disease severity findings have been shown by others in hamsters, mice, ferrets, and nonhuman primates (16–20).

Reduction in effective dose may have led to a greater proportion of naïve animals exposed to oral r-Ad-S–vaccinated (14 of 16) and intranasal r-Ad-S–vaccinated (15 of 16) hamsters gaining weight by the end of the study (Fig. 4E). This is compared to naïve animals exposed to the intramuscularly (spike) vaccinated (8 of 16) and unvaccinated (10 of 16) hamsters (Fig. 4E), where fewer naïve animals increased in size. In addition, hamsters exposed to oral and intranasal r-Ad-S–vaccinated index hamsters had significantly lower lung weights ($P = 0.0012$, oral; $0.0002$, intranasal; Fig. 4F). Moreover, hamsters exposed to intranasal r-Ad-S–vaccinated index hamsters had lower pathology scores ($P = 0.0049$, intranasal; Fig. 4G) compared to unvaccinated hamsters. These data demonstrate that oral and intranasal r-Ad-S–vaccinated animals with a postvaccination SARS-CoV-2 infection transmit less infectious virus through the air to unvaccinated, naïve hamsters than infected...
but unvaccinated (or intramuscularly vaccinated) animals and that this difference in transmission resulted in reduced evidence of severe disease.

**Oral Ad5 vaccination in humans elicits cross-reactive IgA**

Using the same platform in an open-label phase 1 clinical study, 35 healthy individuals received either a single low \([1 \times 10^{10}]\) infectious units (IU); \(n = 15\) or high \([5 \times 10^{10}]\) IU; \(n = 15\) dose of the oral tablet vaccine candidate VXA-CoV2-1 (encoding the SARS-CoV-2 spike and nucleocapsid proteins); in addition, a small cohort \((n = 5)\) received two low doses. SARS-CoV-2–specific IgA was measured in the mucosal compartment using the Meso Scale Discovery (MSD) platform. Fifty-four percent of vaccinees \((19 of 35)\) had a twofold or higher increase in mucosal IgA in either saliva or nasal samples (Fig. 5, A and B). More specifically, 10 of 35 (29%) of vaccinees had a twofold or higher increase in IgA antibodies in their saliva, whereas 12 of 35 (35%) reached the same threshold in their nasal compartment by day 29 after vaccination (Fig. 5, A and B). The responses were similar to all three antigens measured: S, nucleocapsid, and the receptor binding domain (RBD). Furthermore, those that had a twofold increase in saliva or nasal virus–specific IgA also showed an increase in cross-reactive IgA, which bound to spike proteins from all four endemic coronaviruses, the Middle East respiratory syndrome coronavirus (MERS-CoV), and SARS-CoV-1 (Fig. 5, C and D). We did not observe differences between the two doses.

**DISCUSSION**

SARS-CoV-2 emerged in late 2019 and quickly spread around the globe, leading to hundreds of millions of cases and more than 4 million deaths. Despite reports of decreased viral RNA shedding from mRNA-vaccinated individuals compared to unvaccinated individuals \((5, 21)\), recent evidence has demonstrated that vaccinated individuals can get infected with SARS-CoV-2 and shed infectious virus, which can lead to onward transmission. Improved mucosal responses that stimulate local immunity at the site of SARS-CoV-2 replication
may have a much greater impact on human transmission. In this study, we show that oral and intranasal r-Ad-S vaccination decreased evidence of severe disease in vaccinated hamsters and reduced SARS-CoV-2 transmission to unvaccinated, naïve hamsters.

We observed robust anti–spike protein IgG responses after oral and intranasal r-Ad-S vaccination, as previously demonstrated in a different oral r-Ad-S hamster experiment (22). Furthermore, we could detect elevated IgA in the serum and the BAL fluid of mucosally immunized animals. The mucosally vaccinated animals had reduced airborne transmission to naïve animals during an 8-hour airborne exposure window. Specifically, this was assessed by reduced nasal swab viral RNA loads in naïve animals 1 and 3 days after transmission exposure to oral and intranasal r-Ad-S–vaccinated hamsters as compared to control exposed hamsters. We hypothesize that mucosal antibodies in the URT were able to enhance SARS-CoV-2 clearance in vaccinated animals, limiting the infectiousness of transmitted aerosols. Consistent with this hypothesis, anti–spike protein IgA in the BAL fluid of the oral and intranasal r-Ad-S–immunized animals was higher than intramuscular protein– or mock-vaccinated animals.

Other groups have also tested SARS-CoV-2 transmission in a hamster model. For example, Yinda and colleagues (18) determined that intranasally vaccinated hamsters were protected from SARS-CoV-2 when placed in the same cage as unvaccinated hamsters inoculated with $1 \times 10^4$ TCID$_{50}$ of virus. These experiments determined whether intranasally vaccinated animals were protected against transmission from unvaccinated animals. In our study, we tested whether mucosal vaccination reduces the ability of the virus replicating in vaccinated individuals from being transmitted specifically through the air to naïve animals. We used a relatively high dose of virus to induce a postvaccine infection ($1 \times 10^5$ TCID$_{50}$) and physically separated animals in the exposure chamber to remove the possibility of contact or fomite exposure as a means of transmission. We showed that mucosal vaccination could reduce SARS-CoV-2 transmission from vaccinated to unvaccinated animals. These data are relevant because of its implication in global public health, especially in areas in which a high percentage of the people are unvaccinated. In addition, we demonstrated that serum IgG from all vaccinated hamsters bound to spike protein of both the Beta and Delta variants of concern. This suggests that mucosal vaccination could induce
cross-protective antibodies against new variants of concerns. In support of that, we presented data that demonstrate that immunization with an oral tablet spike and nucleocapsid protein vaccine, VXA-COV2-1, generated robust anti–spike protein IgA in nasal swabs and saliva in a subset of individuals, which bound to the spike proteins of multiple coronaviruses, including the four endemic human coronaviruses (NL63, HKU1, OC43, and 229E) and other pathogenic coronaviruses (MERS-CoV and SARS-CoV-1). It is possible that mucosal vaccination against SARS-CoV-2 induces IgA antibodies at the mucosal surface that have superior cross-reactivity to coronaviruses when compared to systemic IgG antibodies. Reports have shown IgA to dominate the early neutralizing antibody response to SARS-CoV-2 (23) and may be more broadly cross-reactive against various coronaviruses (24, 25). In addition, it is suggested that anti-influenza IgA antibodies generated at the mucosal surface are more broadly reactive, likely resulting from enhanced avidity from their multimeric (mostly dimeric) structure (26–28). If additional variants, such as Omicron, arise that result in an increase in postvaccine infections, then broadly reactive IgA antibodies that contribute to reducing transmission even in vaccinated populations would help limit circulating virus. Mucosal vaccination could be considered for implementation not only because they protect the vaccinee but because they likely have a greater effect on the community as a whole. Reducing SARS-CoV-2 transmission to the
unprotected is likely to lead to decreased hospitalization and deaths.

Our study does have limitations. First, we did not measure mucosal T cell responses, which have been shown to play a role in limiting SARS-CoV-2 infection at mucosal sites (29). In addition, the SARS-CoV-2 challenge dose that we used was above physiological dose likely to be picked up by an environmental exposure, as evidenced by the high viral RNA load in the nose at 1 day after infection in all index hamsters. It is likely that mucosal immunization would provide greater protection against SARS-CoV-2 transmission when lower doses of challenge virus are used. Our study was meant to be a stringent challenge to clearly identify advantages between vaccine groups. For practical reasons, such as the limitation of the number of airborne transmission chambers, lower-dose virus inoculation or evaluation at later time points were not done. Furthermore, future work should include challenging mucosally vaccinated hamsters with the Delta and Omicron variants and other variants of concern. In terms of future human studies, a clinical trial with the vaccine construct used in this hamster study is predicted to improve the mucosal antibody response rate because studies in nonhuman primates have shown improved antibody responses compared to the original vaccine tested in humans (30).

An orally delivered, temperature-stable SARS-CoV-2 vaccine is ideal for global vaccination, where adequate storage and qualified health care providers may be in short supply. Intranasal delivery has some of the same advantages, but translating intranasal SARS-CoV-2 vaccination efficacy in humans has proven to be more difficult than in animals (31, 32). Implementing oral vaccine campaigns around the world has been done, as evidenced by the rotavirus and poliovirus vaccination efforts (33, 34). We have previously demonstrated that our SARS-CoV-2 clinical candidate vaccine, VXA-COV2-1, generated robust humoral immune responses in mice (35). In addition, it was well tolerated and immunogenic in a phase 1 clinical trial where the oral vaccine was delivered to participants as tablets (NCT04563702) and protected hamsters from SARS-CoV-2 challenge when given by oral gavage (22). An additional phase 2 study using the vaccine candidate tested in this study (VXA-COV2-1.1-S) and given as a tablet has begun clinical studies (NCT05067933). In summary, the data presented here demonstrate that oral immunization is a viable strategy to decrease SARS-CoV-2 transmission and

Fig. 5. Cross-reactive anti–spike protein IgA in the mucosal and salivary compartments is elicited by oral Ad5 vaccination in a subset of participants. (A and B) Fold change (day 29 versus day 1) in spike (S)-, nucleocapsid (N)-, or receptor binding domain–specific IgA antibodies in nasal (A) and saliva (B) samples were measured by the MSD platform (n = 35). (C and D) The fold change in IgA specific to endemic human coronaviruses was measured in both nasal (C) and saliva (D) samples, respectively, in the responders (n = 12 or n = 7, respectively) from (A) and (B). The red dotted line represents a twofold change, by which individuals were classified as responders.
disease and should be considered for vaccination efforts that increase global immunity to SARS-CoV-2.

**MATERIALS AND METHODS**

**Study design**

Male Syrian hamsters (*Mesocricetus auratus*) about 12 to 14 weeks of age with a weight range of 106 to 136 g were sourced from Charles River Laboratories. Animal work was performed at Lovelace Biomedical, with approval from the Institutional Animal Care and Use Committee (Fy 20-117E-E5). Hamsters were singly housed in filter-topped cage systems and were supplied with a certified diet, filtered municipal water, and dietary and environmental enrichment. The study was powered to compare viral RNA loads in naïve hamsters exposed to vaccinated, SARS-CoV-2–infected index animals between vaccine groups, where \( \beta \) was set to 0.2 and \( \alpha = 0.05 \). Assuming an attack rate of 80% infected in the placebo group and a vaccine efficacy of 70%, an \( N = 15 \) was calculated with continuity correction (36). The study was rounded to 16 naïve and 4 index to maintain the 1:4 ratio. Although the study was not powered to directly compare index groups, many statistically significant differences were observed with \( N = 4 \).

All r-Ad-S vaccinations were given at a dose of \( 1 \times 10^9 \) IU [1:100 of a human dose (11)]. Oral vaccine was delivered by gavage in 300 μl of PBS subsequent to delivery of 300 μl of 7.5% bicarbonate buffer. Intranasal vaccination was delivered in PBS by pipette (25 μl per nostril and 50 μl per animal). The control group received PBS by oral gavage. One group received unadjuvanted recombinant SARS-CoV-2 spike protein, made in insect cells, by 100-μl intramuscular immunization at a dose of 0.1 μg per animal (BEI Resources, #NR-52308). All index animals were challenged by intranasal inoculation of SARS-CoV-2 at about \( 1 \times 10^2 \) TCID\(_{50}\) per animal in 200-μl (100 μl per nostril) volume 7 weeks after the initial vaccination. Index animals were then housed for 24 hours individually before placing them in an aerosol chamber. Each vaccine index group had 4 animals and matched with a corresponding \( N = 16 \) naïve exposed animals (1:4 ratio of index to naïve, with 1 index animal exposed to 4 naïve in a chamber setup). All animals were euthanized 5 days after inoculation (index) or aerosol chamber exposure (naïve) for terminal assays. For serum collection, animals were sedated with a mixture of ketamine (60 to 120 mg/kg) and xylazine (5 to 20 mg/kg) by intraperitoneal injection to restrain them for all blood draws. About 300 to 500 μl of blood samples were collected into yellow top serum separator tubes, allowed to clot for 30 min to an hour, and processed to serum by centrifugation at 2500g for at least 10 min at 2° to 8°C. Serum was aliquotted into a cryovial and stored at −80°C. BAL was collected by harvesting the right lung, centrifuging at 2000g for 5 min, discarding the pellet, and freezing at −80°C.

**Vaccine constructs**

r-Ad-S is a rAd5 vector containing the gene of the full-length SARS-CoV-2 stabilized spike protein, under control of the cytomegalovirus promoter. rAd5 vaccine constructs were created on the basis of the published DNA sequence of SARS-CoV-2 publicly available as GenBank accession no. MN908947.3. The published amino acid sequences of the SARS-CoV-2 spike were used to create recombinant plasmids containing transgenes cloned into the E1 region of Ad5 (rAd5) (37), using the same vector backbone used in prior clinical trials for oral rAd tablets (10, 11). All vaccines were grown in the Expi293F suspension cell line (Thermo Fisher Scientific) and purified by CsCl density centrifugation. The spike protein vaccine was provided by BEI Resources, NR-52308, spike glycoprotein (stabilized) from SARS-related coronavirus 2, recombinant from baculovirus, NR-52308.

**Transmission chamber**

The airborne transmission chamber was based on previous transmission work with aerosols (38) and adapted for hamsters. The chamber consists of multiple subchambers that support unidirectional flow. All chambers were fitted with access doors with air tight seals and appropriate safety features to ensure that animals could not interact with fans, sampling, flow, or other features. Approximate dimensions of the first chamber (chamber 1) were 4 inches by 10 inches by 9 inches (length by width by height). The unidirectional flow (5 liter/min) was controlled by regulated house exhaust flow from the third chamber. This drew room air into chamber 1 by a high efficiency particulate air (HEPA) filter (HEPA vacuum filter compatible with Kenmore 86880, EF-2, Panasonic MC-V194H vacuum cleaner). Chamber 1 was also fitted with a recirculating fan (ANVISION 40 mm by 10 mm DC 5 V USB brushless cooling fan, dual ball bearing, model YDM4010B05) to ensure homogeneity of the aerosol before transitioning into chambers 2 and 3. The connector chamber (chamber 2; approximate dimensions of 4 inches by 5 inches by 5 inches) connected chambers 1 and 3 to separate the hamsters but allows for air passage. Chamber 3 (about 10 inches by 10 inches by 9 inches) housed the naïve hamsters. A wire mesh screen with 0.25 inch by 0.25 inches holes was placed at each end of chambers 1 and 3 to prevent hamsters from moving to another chamber. In addition, chamber 2 was raised off the ground to prevent feces or urine from moving between chambers. No bedding was available in any of the three chambers.

**Assessment of infectious SARS-CoV-2 load in lung homogenates**

Lung tissue samples from euthanized hamsters infected with SARS-CoV-2 were collected, weighed, and homogenized with beads using a TissueLyser (QIAGEN). Each sample was serially diluted 10-fold in Dulbecco’s modified Eagle’s medium containing 2% fetal bovine serum and 1% penicillin-streptomycin solution. Vero E6 cell monolayers at ≥90% confluency in 96-well plates were rinsed with PBS. The plates were inoculated with 100 μl of each sample dilution in five technical replicates. Negative control wells contained dilution medium only. The plates were incubated at 37°C and 5% CO\(_2\) for 72 hours. Cytotoxic effect was scored after fixing the cell monolayers with 10% formalin and staining with 0.5% crystal violet. Viral load was determined by TCID\(_{50}\) per milliliter of lung homogenate using the Reed and Muench method (39). Infectious virus titers in infected lungs were expressed as TCID\(_{50}\) per gram of tissue.
solution, followed by the addition of 100 μl per well of supplied 3,3’, 5,5’-tetramethylbenzidine dihydrochloride (TMB) solution. Plates were developed for 15 min at room temperature in the dark before development was stopped with 50 μl per well of supplied stop solution. Optical densities (ODs) were measured at 450 nm with a SpectraMax M2 microplate reader.

**IgG enzyme-linked immunosorbent assay**

Purified SARS-CoV-2 S1 protein (GenScript) or Beta and Delta variant S1 proteins (Acro biosciences) in carbonate buffer (pH 9.4; Thermo Fisher Scientific) were coated onto microtiter plates (MaxiSorp, Nunc) at 1 μg/ml and incubated overnight at 4°C before blocking with 100 μl of PBS–0.05% Tween (PBST) + 1% bovine serum albumin (BSA) for 1 hour. Serum samples were serially diluted in PBST. After a 2-hour incubation at room temperature, the plates were washed three times with PBST, followed by the addition of 100 μl per well of 1:3000 goat anti-hamster IgG-HRP (Thermo Fisher Scientific) in PBST + 1% BSA. Plates were incubated at room temperature for 1 hour before washing three times with PBST. Next, TMB substrate (Rockland) was added at a concentration of 50 μl per well. The plates were developed for 10 min and then stopped with 50 μl per well of 2 M sulfuric acid. ODs were measured at 450 nm with a SpectraMax M2 microplate reader.

**IgA MSD assay**

S1 protein was biotinylated according to the manufacturer’s instructions (EZ-link, Thermo Fisher Scientific) and was conjugated to a U-PLEX MSD linker (MSD). The linked S1 protein was coated on 2-spot U-PLEX 96-well plates (MSD) for a final concentration of 66 nM per well and incubated overnight. The next day, the plates were blocked with PBST for 1 hour before the addition of hamster serum or BAL fluid. Samples from each individual hamster acquired on days 0, 28, and 55 were quantified on the same plate. The serum samples were diluted to 1:200, and the BAL samples were added to a plate neat. After a 2-hour sample incubation, the plate was washed, and SULFO-TAG (MSD) anti-hamster IgA (Brookwood Biomedical) detection antibody was added. The plate was washed, and 1x MSD read buffer was added. Each plate was analyzed using MSD QuickPlex.

**SARS-CoV-2 challenge**

All animals were challenged by intranasal inoculation of 1 × 10⁵ TCID₅₀ per animal SARS-CoV-2 (isolate USA-WA1/2020) at 100 μl per nostril (for a total volume of 200 μl) 8 weeks after initial vaccination. SARS-CoV-2, isolate USA-WA1/2020, was sourced from BEI Resources. The complete genome has been previously sequenced for the original isolate (GenBank accession number MN985325), after one passage in Vero E6 cells (GenBank accession number MT020880), and after four passages in Vero E6 cells (GenBank accession number MT246667). It was propagated in Vero E6 African green monkey kidney cells (BEI Resources, catalog no. N596) at the University of Texas Medical Branch, and virus was stored in a biosafety level 3 compliant facility.

**Collection of nasal swabs**

Plasdent Dentistry Maxapplicators with 0.5-mm ultrafine nonlinting, nonabsorbable fiber head tips were used for nasal swab collections. Swabs were inserted about 2 to 4 mm into the naris of an anesthetized hamster, removed, and cut at the hinge point. Swabs were then placed into Safe-Lock Eppendorf tubes and flash-frozen at −80°C until RNA isolation.

**Detection of SARS-CoV-2 genomic RNA in nasal swabs and lung homogenates by quantitative reverse transcription polymerase chain reaction**

Lung samples were weighed and homogenized with beads using a TissueLyser (QIAGEN) in 1 ml of TRI Reagent before RNA was isolated and purified from tissue samples using the Direct-zol 96 RNA Kit (Zymo Research). Copies of the SARS-CoV-2 Nucleocapsid (N) gene were measured by quantitative reverse transcription polymerase chain reaction (qRT-PCR) TaqMan Fast Virus 1-step assay (Applied Biosystems). SARS-CoV-2–specific primers and probes from the 2019-nCoV RUO Assay Kit (Integrated DNA Technologies) were used: L Primer, TTACAAAACATTGGCAGAAA; R primer, GCCGCCACATTCCGAAGAA; and probe, 6FAM-ACAATTTGC-CCCACGGCTTCAG-BHQ-1. Reactions were carried out on a Stratagene MX3005P or Bio-Rad CFX384 Touch instrument according to the manufacturer’s specifications. A semilogarithmic standard curve of synthesized SARS-CoV-2 N gene RNA (LBRI) was obtained by plotting the Ct values against the logarithm of cDNA concentration and used to calculate SARS-CoV-2 N gene in copies per gram of tissue.

**Gross pathology scoring**

Gross necropsy observations of the lung were recorded in Provantis using consistent descriptive terminology to document location(s), size, shape, color, consistency, and number. Gross observations included a severity grade for red discoloration of the lung (likely to be associated with pneumonia) based on a 0 to 4 scale indicating percentage of whole lung affected: none (no grade), minimal (1), mild (2), moderate (3), and marked (4) correlating to 0%, 1 to 25%, 26 to 50%, 51 to 75%, and 76 to 100% affected, respectively.

**Clinical protocol**

A phase I clinical study (https://clinicaltrials.gov/show/NCT04563702) was designed to evaluate the safety and immunogenicity of an oral spike and nucleocapsid protein vaccine (termed VXA-CoV2-1) in 35 individuals at two different doses (1 × 10¹⁰ and 5 × 10¹⁰ IU). Five sentinel volunteers were dosed first, and after a week of monitoring for vaccine-induced toxicities, the remaining volunteers in the treated cohort were randomized with four placebo controls. Only five individuals in the low-dose group were boosted; all other participants were given one dose of VXA-CoV2-1. In terms of demographics, 66% of participants were male, 57% were white, 20% were African-American, 14% were Hawaiian or Pacific Islander, and 9% were other. A total of 29% of participants reported being Hispanic or Latino. Participants were between the ages of 18 and 53.

**Study approval**

The study was conducted in accordance with applicable Good Clinical Practice guidelines, the United States Code of Federal Regulations, and the International Conference on Harmonization guidelines. Institutional Review Board (IRB) approval was obtained from the Aspire IRB (protocol no. 20202876) before study-specific screening and enrollment of participants. Informed consent was obtained from all participants after discussion of the study procedures and potential risks.
Measurement of vaccine-induced IgA antibody mucosal samples

Nasal samples were collected according to the package insert for the Nasosorption FX-I/SAM devices (Mucosal Diagnostics) and extracted by centrifugation in a 1× PBS with 0.02% azide solution. Saliva samples were collected by centrifugation after having each participant chew on a cotton swab for 45 s and transferred to a Salivette (Sarstedt). IgA was measured with a SARS-CoV-2-specific (panel 2) and coronavirus panel 3 MSD immunoassay. Plates were blocked, washed, and incubated with sample and detection antibody according to the manufacturer’s instructions. Samples were diluted at 1:10 and 1:100 in Diluent 100. Plates were read on a MesoQuickPlex instrument. Sample antibody concentrations were reported in arbitrary units per milliliter as calculated from a standard curve supplied with the kit.

Because of the variability in sampling the human mucosa, samples were normalized using an enzyme-linked immunosorbent assay (ELISA) for the detection of total IgA. Briefly, purified anti-human IgA monoclonal antibody (mAb) MT57 (Mabtech) was coated onto 96-well MaxiSorp plates (Thermo Fisher Scientific) at 2 μg/ml in PBS and incubated overnight at 4°C. Plates were washed with 1× PBS + 0.1% Tween-20 (PBST) and blocked with PBST + 1% BSA for 1 hour at room temperature. After a wash step, saliva and eluted nasal samples were diluted 1:100 in PBST and serially diluted 1:3 down the plate. Human IgA (Sigma-Aldrich) was used to create a standard curve starting at 200 ng/ml and serially diluted twofold seven times. The samples and standards were transferred to the coated plate and incubated for 2 hours at room temperature. After a wash step, a 1:1000 dilution of anti-human IgA mAb MT20 Alkaline Phosphatase (ALP) conjugate (Mabtech) was added to the nasal samples. The plates were incubated at room temperature for 1 hour followed by washing. Plates with nasal samples developed for 1 hour in the dark with para-nitrophenylphosphate substrate (Mabtech), and ODs were measured at 405 nm with a SpectraMax M2 microplate reader. Plates with saliva samples were developed in the dark with an HRP substrate (Rockland) and were stopped with sulfuric acid (Honeywell). ODs were measured at 450 nm. The concentration of total IgA in human mucosal samples was generated by a standard curve. A normalization factor was applied by dividing the postvaccination sample by the prevaccination sample. Data analysis was performed in GraphPad Prism Software.

Statistical analysis

All raw, individual-level data are presented in data file S1. The methods used for determining significance were one- or two-way analysis of variance (ANOVA) and Dunnett’s multiple comparisons test or Fisher’s exact test. For Fig. 1C, one-way ANOVA was used to create a standard curve starting at 200 ng/ml and serially diluted twofold seven times. The samples and standards were transferred to the coated plate and incubated for 2 hours at room temperature. After a wash step, a 1:1000 dilution of anti-human IgA mAb MT20 Alkaline Phosphatase (ALP) conjugate (Mabtech) was added to the nasal samples. The plates were incubated at room temperature for 1 hour followed by washing. Plates with nasal samples developed for 1 hour in the dark with para-nitrophenylphosphate substrate (Mabtech), and ODs were measured at 405 nm with a SpectraMax M2 microplate reader. Plates with saliva samples were developed in the dark with an HRP substrate (Rockland) and were stopped with sulfuric acid (Honeywell). ODs were measured at 450 nm. The concentration of total IgA in human mucosal samples was generated by a standard curve. A normalization factor was applied by dividing the postvaccination sample by the prevaccination sample. Data analysis was performed in GraphPad Prism Software.
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