Intranasal administration of a virus like particles-based vaccine induces neutralizing antibodies against SARS-CoV-2 and variants of concern

Dominik A. Rothen1,2 | Pascal S. Krenger1,2 | Aleksandra Nonic1,2 | Ina Balke3 | Anne-Cathrine S. Vogt1,2 | Xinyue Chang1,2 | Alessandro Manenti4 | Fabio Vedovi4 | Gunta Resevica3 | Senta M. Walton5 | Andris Zeltins3 | Emanuele Montomoli4,6 | Monique Vogel1,2 | Martin F. Bachmann1,2,7 | Mona O. Mohsen1,2,5

Abstract

Background: The highly contagious SARS-CoV-2 is mainly transmitted by respiratory droplets and aerosols. Consequently, people are required to wear masks and maintain a social distance to avoid spreading of the virus. Despite the success of the commercially available vaccines, the virus is still uncontained globally. Given the tropism of SARS-CoV-2, a mucosal immune reaction would help to reduce viral shedding and transmission locally. Only seven out of hundreds of ongoing clinical trials are testing the intranasal delivery of a vaccine against COVID-19.

Methods: In the current study, we evaluated the immunogenicity of a traditional vaccine platform based on virus-like particles (VLPs) displaying RBD of SARS-CoV-2 for intranasal administration in a murine model. The candidate vaccine platform, CuMVTT-RBD, has been optimized to incorporate a universal T helper cell epitope derived from tetanus-toxin and is self-adjuvanted with TLR7/8 ligands.

Results: CuMVTT-RBD vaccine elicited a strong systemic RBD- and spike-IgG and IgA antibodies of high avidity. Local immune response was assessed, and our results demonstrate a strong mucosal antibody and plasma cell production in lung tissue. Furthermore, the induced systemic antibodies could efficiently recognize and neutralize different variants of concern (VOCs).

Conclusion: Our data demonstrate that intranasal administration of CuMVTT-RBD induces a protective systemic and local specific antibody response against SARS-CoV-2 and its VOCs.

KEYWORDS
COVID-19, intranasal, SARS-CoV-2, vaccine, virus-like particles
Till date, COVID-19 caused by SARS-CoV-2 is still considered a global pandemic that has wreaked havoc globally and put a heavy toll on public health and economy. The marketed vaccines such as mRNA, viral vector, and inactivated viruses have greatly reduced the number of COVID-19 mortality and hospitalization and continue to provide different levels of protections against the emerging variants of concern (VOCs).

Viral tropism depends, among other factors, on the susceptibility of a specific host cell. COVID-19 patients often present with respiratory illness that can progress to severe pneumonia. These observations suggested that the lung is the primary organ infected by SARS-CoV-2. The fact that lung epithelial cells express the angiotensin converting enzyme 2 (ACE2), the viral receptor, substantiates this observation. The primary port of entry to the body is alveolar epithelial cells, but vascular endothelial cells also express ACE2 and are a prominent place of viral replication. These cells may be considered the base for early infection and viral replication as well as long-term viral persistence in some cases.

The currently available marketed vaccines are administered intramuscularly (i.m.) producing systemic spike and RBD-specific antibodies (Abs) that can recognize and neutralize the virus. Given the tropism of SARS-CoV-2, recent research efforts have also been devoted toward the development of an intranasal (i.n.) COVID-19 vaccine. Seven intranasal COVID-19 vaccine candidates are currently in clinical trials. Intranasal vaccination route may offer several advantages over i.m. route including; needle-free administration, direct delivery to the site of infection, and most importantly, the induction of mucosal immunity in the respiratory tract. Secretory IgA (sIgA) is of major importance in the respiratory tract where it presents an efficient line of defence against respiratory infections. Furthermore, mucosal vaccination can result in resident B and T cell priming leading to long-lived Ab secreting cells or tissue-resident memory cells, which add in clearing the viral infection. This locally induced immune reaction has been shown to reduce viral replication and shedding in lungs and nasal passages leading to lower infection and transmission. The concept of i.n. vaccination goes back to 1960s based on observations with live-attenuated influenza vaccines (LAIV) that mimic a natural influenza infection and have shown to elicit a protective local and systemic antibody as well as cellular responses. Live-attenuated virus or viral-vector based vaccines need to infect cells for replication. Moreover, attenuated viruses may pose a small risk of retaining their replication ability, especially in people with weaker immune systems.

Efficient induction of mucosal immunity can be best achieved by vaccines that mimic mucosal pathogens. Virus-like particles (VLPs) constitute an efficient and safe vaccine platform as they lack genetic material for replication in vivo. Their particulate and repetitive surface structure enables them to stimulate innate and adaptive immune response and target the mucosa as well as the underlying dendritic cells (DCs). We have previously assessed the efficacy of Qß-VLPs as an i.n. vaccine platform. Our results indicated efficient induction of specific-IgGs in serum and lungs besides robust local IgA production. Here, we provide a proof-of-concept (PoC) in murine model proving the immunogenicity and efficacy of i.n. administration of a SARS-CoV-2 vaccine based on VLPs. The developed vaccine candidate is based on our optimized plant-derived VLPs (CuMV<sub>TT</sub>-RBD) displaying the receptor-binding domain (RBD) of SARS-CoV-2. Our results demonstrate that
CuMV_TT-RBD induces strong systemic and local B cell response including high levels of IgG and IgA, plasma cell (PC) formation as well as broad viral neutralization. Taken together, our vaccine constitutes an efficient candidate for the generation of Ab-based vaccine that can be administered mucosally in a needle-free manner.

## METHODS

### 2.1 Mice

All in vivo experiments were performed using (8–12-week-old) wild-type (wt) female BALB/cOlaHsd mice purchased from Harlan. All animal procedures were conducted in accordance with the Swiss Animal Act (455.109.1-5 September 2008) of University of Bern. All animals were treated for experimentation according to the protocols approved by the Swiss Federal Veterinary Office.

### 2.2 Protein expression and purification

RBD<sub>wt</sub> of SARS-CoV-2 and mutant RBDs (RBD<sub>K417N</sub>, RBD<sub>E484K</sub>, RBD<sub>N501Y</sub>, RBD<sub>K417N/E484K/N501Y</sub> and RBD<sub>L452R/E484Q</sub>) were expressed using Expi293F cells (Gibco, Thermo Fisher Scientific, Waltham, MA, USA). The amino acid (a.a.) sequence of each RBD was inserted into a pTWIST-CMV-BetaGlobin-WPRE-Neo vector (Twist Bioscience, San Fransisco, CA, USA). RBD-His Tag construct was further transformed into competent XL-1 Blue bacterial cells. After plasmid purification, 50 μg of the plasmid was then transfected into Expi293F cells (Gibco, Thermo Fisher Scientific, Waltham, MA, USA). Ninety-six hours later, the supernatant containing RBD was harvested and dialyzed with PBS. RBD protein was then captured using His-Trap HP column (GE Healthcare, Wauwatosa, WI, USA) or HiTrap TALON crude column (Cytiva, Uppsala, Sweden). Fractions were collected and concentrated. Buffer-exchanged to PBS using Vivaspin 20 5KDMWCO spin column (Sartorius, Goettingen, Germany). RBD proteins were removed using Amicon Ultra 0.5, 100 K (Merck Millipore, Burlington, MA, USA). VLP samples were centrifuged for 2 min at 12,000 g for measurement on ND-1000. RBD<sub>wt</sub>-SDS-PAGE was stained with InstantBlue<sup>TM</sup> Coomassie stain and image was obtained with Azure Biosystems c300 with a result of approximately 30%, meaning that there is about 60-RBD per one VLP. RBD<sub>wt</sub> coupling to CuMV<sub>TT</sub> was further analyzed by Western blot. For this purpose, RBD<sub>wt</sub>, CuMV<sub>TT</sub>, and coupled CuMV<sub>TT</sub>-RBD were separated on a 12% SDS PAGE. Using the Trans-Blot<sup>®</sup> Turbo™ Transfer System protein bands were transferred onto a 0.2 μM PVDF membrane (Biorad, Hercules, USA). The membrane was further processed by using the iBind<sup>™</sup> Flex Western Device (Invitrogen, USA) according to the manufacturer’s protocol. As primary antibody SARS-CoV-2 Spike RBD antibody (R&DSystems, MAB10540) at (1:1000) and as detecting antibody goat anti-mouse IgG conjugated to Horseradish Peroxidase (HRP) (Jackson ImmunoResearch, 115-035-071, West Grove, Pennsylvania) was added at (1:1000). Bound antibodies were detected by using SuperSignal™ West Pico PLUS Chemiluminescent Substrate (ThermoScientific, 34579). WB image was obtained with Azure Biosystem c300 with exposure time of 2s using chemiluminescence channel. Packaging of ssRNA in CuMV<sub>TT</sub>-VLPs and the developed vaccine CuMV<sub>TT</sub>-RBD was confirmed by % Agarose gel run at 50 V for 40 min and visualized using Azure Biosystems c300 with exposure time of 10s using UV302 channel. A DNA Ladder (Thermo Scientific, Cat. Nr. SM0242) was included.

### 2.3 CuMV<sub>TT</sub> expression and production

Expression and production of CuMV<sub>TT</sub> was described in detail in Zeltins et al.<sup>15</sup> The level of LPS is 10 endotoxin per mg of CuMV<sub>TT</sub> measured using LAL test (Pierce).

### 2.4 Development of CuMV<sub>TT</sub>-RBD vaccine

RBD<sub>wt</sub> was conjugated to CuMV<sub>TT</sub> using the cross-linker succinimidyl 6-(beta-maleimidopropionamido) hexanoate (SMPH) (Thermo Fisher Scientific, Waltham, MA, USA) at 7.5 molar excess to CuMV<sub>TT</sub> for 30 min at 25°C. The coupling reactions were performed with molar ratio RBD/CuMV<sub>TT</sub> (1:1) by shaking at 25°C for 3 h at 250 g on a DSG TiterTek (Flow Laboratories, Irvine, UK). Unreacted SMPH and RBD proteins were removed using Amicon Ultra 0.5, 100 K (Merck Millipore, Burlington, MA, USA). VLP samples were centrifuged for 2 min at 12,000 g for measurement on ND-1000. RBD<sub>wt</sub>-SDS-PAGE was stained with InstantBlue<sup>TM</sup> Coomassie stain and image was obtained with Azure Biosysstem c300 with a result of approximately 30%, meaning that there is about 60-RBD per one VLP. RBD<sub>wt</sub> coupling to CuMV<sub>TT</sub> was further analyzed by Western blot. For this purpose, RBD<sub>wt</sub>, CuMV<sub>TT</sub>, and coupled CuMV<sub>TT</sub>-RBD were separated on a 12% SDS PAGE. Using the Trans-Blot<sup>®</sup> Turbo™ Transfer System protein bands were transferred onto a 0.2 μM PVDF membrane (Biorad, Hercules, USA). The membrane was further processed by using the iBind<sup>™</sup> Flex Western Device (Invitrogen, USA) according to the manufacturer’s protocol. As primary antibody SARS-CoV-2 Spike RBD antibody (R&DSystems, MAB10540) at (1:1000) and as detecting antibody goat anti-mouse IgG conjugated to Horseradish Peroxidase (HRP) (Jackson ImmunoResearch, 115-035-071, West Grove, Pennsylvania) was added at (1:1000). Bound antibodies were detected by using SuperSignal™ West Pico PLUS Chemiluminescent Substrate (ThermoScientific, 34579). WB image was obtained with Azure Biosystem c300 with exposure time of 2s using chemiluminescence channel. Packaging of ssRNA in CuMV<sub>TT</sub>-VLPs and the developed vaccine CuMV<sub>TT</sub>-RBD was confirmed by % Agarose gel run at 50 V for 40 min and visualized using Azure Biosystems c300 with exposure time of 10s using UV302 channel. A DNA Ladder (Thermo Scientific, Cat. Nr. SM0242) was included.

### 2.5 Electron microscopy

Physical stability and integrity of the candidate vaccine CuMV<sub>TT</sub>-RBD were visualized by transmission electron microscopy (Philips CM12 EM). For imaging, sample-grids were glow discharged and 10 μl of purified CuMV<sub>TT</sub>-RBD (1.1 mg/ml) was added for 30s. Grids were washed 3x with ddH<sub>2</sub>O and negatively stained with 5 μl of 5% uranyl acetate for 30s. Excess uranyle acetate was removed by pipetting, and the grids were air dried for 10 min. Images were taken with 84,000x and 110,000x magnification.

### 2.6 Binding ELISA assay

To test if the vaccine can bind the relevant human receptor ACE2, ELISA plates were coated with 2 μg/ml of ACE2 in PBS at a volume of 50 μl/well. The plate was incubated at 4°C overnight. The plate was washed with PBS+-Tween 0.01%. Added 50 μl/well of PBS-Casein 0.15% and incubated for 1 h at RT on a shaker. Flicked off the blocking solution and added 50 μl of the CuMV<sub>TT</sub>-RBD, CuMV<sub>TT</sub>, and
SARS-CoV-2 RBD<sub>wt</sub> from 50 µg/ml to the first row of the plate followed by 1:3 dilution. The plate was incubated for 1 h at RT, washed with PBS-Tween 0.01%. 50 µl of mouse anti-CuMV<sub>TT</sub> monoclonal antibody (clone 1-1A8/ batch 2) at a concentration of 1 µg/ml was added to each well as a secondary antibody and incubated for 1 h at RT on a shaker. The plate was washed and 50 µl of the detection antibody; HRP labeled goat anti-mouse IgG Fc gamma at a dilution of 1:1000 in PBS-Casein 0.15% was added to each well. The plate was incubated for 1 h at RT. The plate was developed and read at OD<sub>450</sub> nm (BioTek, USA).

2.7 | Vaccination regimen

Wild-type BALB/cOlaHsd mice (8–12 weeks, Harlan) were vaccinated intranasally (i.n.) with 40 µg with either CuMV<sub>TT</sub>-RBD vaccine or CuMV<sub>TT</sub> as a control in a volume of 40 µl without any adjuvants (20 µg in each nostril). The mice were boosted with an equal dose at day 28 and bled until day 49. Serum was collected on a weekly basis via tail bleeding and the serum was isolated using Microtainer Tube (BD Biosciences, USA). For Fluorospot samples and Bronchoalveolar lavage (BAL), wt BALB/cOlaHsd mice (8–12 weeks, Harlan) were vaccinated i.n. as indicated above. Serum samples were collected on day 35.

2.8 | Enzyme-Linked Immunosorbent Assay (ELISA)

To determine the total IgG Abs against the candidate vaccine CuMV<sub>TT</sub>-RBD in sera of vaccinated mice, ELISA plates were coated with SARS-CoV-2 RBD<sub>wt</sub> or spike protein (Sinobiological, Beijing, China) at concentrations of 1 µg/ml overnight. ELISA plates were washed with PBS-0.01% Tween and blocked using 100 µl PBS-Casein 0.15% for 2 h at RT. Sera from vaccinated mice serially diluted 1:3 starting with a dilution of 1:20 and incubated for 1 h at RT. After washing with PBS-0.01% Tween, goat anti-mouse IgG conjugated to Horseradish Peroxidase (HRP) (Jackson ImmunoResearch, West Grove, Pennsylvania) was added at (1:2000) and incubated for 1 h at RT. IgA Abs were detected by using a goat anti-mouse IgA POX (ICN 55549, 1:1000 dilution) detecting Ab. Plates were developed with TMB as described above and read at OD<sub>450</sub> nm.

2.9 | Avidity (ELISA)

To test the avidity of IgG and IgA Ab against RBD, the above-described protocol was expanded by an additional step as previously described. Following serum incubation at RT for 1 h, the plates were washed once in PBS/0.01% Tween, and then washed 3x with 7 M urea in PBS-0.05% Tween or with PBS-0.05% Tween for 5 min. After washing with PBS-0.05% Tween, goat anti-mouse IgG conjugated to Horseradish Peroxidase (HRP) (Jackson ImmunoResearch, West Grove, Pennsylvania) was added (1:2000) and incubated for 1 h at RT. IgA Abs were detected by using a goat anti-mouse IgA POX (ICN 55549, 1:1000 dilution) detecting Ab. Plates were developed with TMB as described above and read at OD<sub>450</sub> nm.

2.10 | Bronchoalveolar Lavage

Bronchoalveolar Lavage (BAL) samples were collected as described in Sun et al.éroth et al.

2.11 | Isolation of lymphocytes

2.11.1 | From lung samples

Lungs were perfused with 10 ml of 1 mM EDTA in PBS via the right ventricle of the heart to remove blood cells from the lung vasculature. Lungs were dissected and digested for 30 min at 37°C using RPMI media (2% FBS+Pen/Strep, glutamine, 10 mM HEPES) containing 0.5 mg/ml Collagenase D (Roche). The digested fragments were passed through a 70 µm cell strainer (Greiner bio-one, Art. Nr. 542070), and RBCs were lysed using ACK buffer. Lymphocytes were isolated using 35% Percoll gradient.

2.11.2 | From spleen samples

The spleen was collected from mice and transferred into 5 ml RPMI media (2% FBS+Pen/Strep, glutamine, 10 mM HEPES). A single cell suspension was prepared by passing the spleen through a 70 µm cell strainer. The suspension was collected and transferred into a falcon tube. The tube was centrifuged for 8 min at 4°C and 300×g. ACK lysis was performed, media added and centrifuged for 8 min at 4°C and 300×g. The pellet was resuspended in media.
2.11.3 | From bone-marrow (BM)

Tibia and femur were collected from mice and transferred into 5 ml RPMI media (2% FBS+Pen/Strep, glutamine, 10 mM HEPES). The BM cells were isolated using a syringe by rinsing the bones to flush out the cells. A single cell suspension was prepared by passing the spleen through a 70 μm cell strainer on a petri dish. The suspension was collected and transferred into a falcon tube. Petri dish was washed with 5 ml media and also added to the falcon tube. The tube was centrifuged for 8 min at 4°C and 300 g. ACK lysis was performed, media added and centrifuged for 8 min at 4°C and 300g. The pellet was resuspended in media.

2.12 | Fluorospot

Fluorospot assay was performed according to the manufacturer’s protocol (FluoroSpot-protocol.pdf (mabtech.com). Briefly, Fluorospot plate (Mabtech, Cat no. 3654-FL) was coated with 100 μl RBD (50 μg/ml) per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight. The next day, the plate was washed with PBS and blocked for 30 min at RT by the addition of 200 μl per well, 4°C overnight.

2.13 | BLI-based assay

Antibody competitive binding activity was measured on an Octet RED96e (FortéBio) instrument which allows real-time analysis due to the shift in the wavelength of the reflected light. Anti-Penta-HIS (HIS1K, Lot 2006292, FortéBio) biosensors were first loaded into a biosensor microplate and pre-hydrated in BLI assay buffer (PBS, 0.1% BSA, 0.02% Tween 20) for 10 min. 96-well microplates were loaded with 200 μl per well. The tips were immobilized with 15 μg/ml Sars-Cov-2 spike RBD containing a His-Tag (Sino Biological, USA). After, they were loaded with sera from mice, diluted 1:20 in BLI assay buffer. Next, association with 50 nM of human receptor ACE2 (Sino Biological, USA) diluted in BLI assay buffer was measured. To regenerate the tips, two additional steps with regeneration buffer (0.1 M glycine, pH 1.5) and neutralization buffer (BLI assay buffer) were performed.

2.14 | SARS-COV-2 wt and VOC live viruses

The SARS-CoV-2 2020-nCov/Italy-INMI clade V (Wuhan), the B.1.1.7 (UK VOC) named England/MIG457/2020, the B.1.351 (South Africa VOC) namedhCoV19/ Netherlands/NoordHolland_10159/2021, next strain clade 20H were all purchased from European Virus Archive (EVAg). SARS-CoV-2 strains were propagated in VERO E6 cells (ATCC—CRL 1586) in T175 Flasks using Dulbecco’s Modified Eagle’s-high glucose medium (DMEM) (Euroclone, Pero, Italy) supplemented with 2 mM L-glutamine (Lonza, Milano, Italy), 100 units/ml penicillin-streptomycin (Lonza, Milano, Italy, and 2% fetal bovine serum (FBS) (Euroclon, Pero, Italy). All viral growth and neutralization assay with SARS-CoV-2 live viruses were performed inside the VisMederi Bisecurity Level 3 laboratories.

2.15 | Neutralization Assay cytopathic-effect-based (CPE)

The neutralization assay was performed as previously reported by Manenti et al. Briefly, 2-fold serial dilutions of heat-inactivated mouse serum samples were mixed with an equal volume of viral solution containing between 25 TCID50 of SARS-CoV-2. The serum-virus mixture was incubated for 1 h at 37°C in a humidified atmosphere with 5% CO2. After the incubation time, 100 μl of the mixture at each dilution point was passed to a 96-well cell plate containing a sub-confluent VERO E6 (ATCC—CRL 1586) monolayer. The plates were incubated for 3 days (Wuhan strain) and for 4 days (B.1.1.7 and B.1.351) at 37°C in a humidified atmosphere with 5% CO2. The day of the read-out each well was inspected by means of an inverted optical microscope to evaluate the percentage of cytopathic effect (CPE) developed in each well. The neutralization titer has been reported as the reciprocal of the highest dilution of serum able to inhibit and prevent at least in 50% of cells the CPE.

2.16 | Statistical analysis

Data were analyzed and presented as mean ± SEM using Student’s t-test or one-way ANOVA as mentioned in the figure legend, with GraphPad PRISM 9. The value of p < .05 was considered statistically significant (*p < .05, **p < .01, ***p < .001, **** p < .0001).

3 | RESULTS

3.1 | CuMV TT constitute an efficient platform for vaccine development

In order to generate a vaccine-candidate against SARS-CoV-2 for i.n. administration, we have utilized our optimized plant-derived VLPs (CuMV TT) as a vaccine platform. RBD amino acid sequence (a.a. Arg319-Phe541) of SARS-CoV-2 was chemically coupled to CuMV TT using SMPH bifunctional cross-linker (Figure 1A). The generated...
vaccine candidate CuMV<sub>T</sub>-RBD is self-adjuvated with prokaryotic ssRNA (TLR7/8 agonist) which is packaged during expression and assembly in the bacterial <i>E. coli</i> system (Figure 1B). Efficiency of RBD coupling to CuMV<sub>T</sub> was confirmed by SDS-PAGE (Figure 1C) and Western blot (Figure 1D). The integrity of the VLPs following the coupling process was checked by electron microscopy and showed no signs of aggregation (Figure 1E). Finally, to confirm the antigenicity of CuMV<sub>T</sub>-RBD as well as the correct folding and confirmation of the RBD displayed on the particle’s surface, a receptor binding assay was performed. To this end, the human receptor ACE2 was coated on ELISA plate. CuMV<sub>T</sub>-RBD, CuMV<sub>T</sub>, and RBD were added. Anti-CuMV<sub>T</sub> antibodies were used as a secondary antibody to detect receptor bound VLPs. The results revealed that CuMV<sub>T</sub>-RBD can bind to ACE2 receptor indicating correct folding of RBD on the surface of the VLP while the control did not show any binding (Figure 1F).

3.2 | Intranasal administration of CuMV<sub>T</sub>-RBD induces a systemic RBD- and spike-specific IgG response of high avidity

To test the immunogenicity and the induction of a humoral immune response in murine models, BALB/c mice were i.n. primed on day 0 and boosted on day 28 with 40 μg of CuMV<sub>T</sub>-RBD vaccine or with 40 μg of CuMV<sub>T</sub> as a control without addition of adjuvants. Vaccination and bleeding regimen are shown in Figure 2A. Total systemic RBD and spike-specific IgG were measured by ELISA. Systemic RBD-specific IgG response was detected in the group receiving CuMV<sub>T</sub>-RBD seven days after the priming dose. Furthermore, the induced response increased by about 1000-fold following the booster dose on Day 35 (Figure 2B and C). Full-length spike protein responses remained low after priming but increased significantly by 30-folds upon a booster injection resulting in a stable IgG antibody titer (Figure 2D and E). ELISA plates were coated with 1 μg/ml of spike or RBD protein as mentioned in the method section, however, from a molar point of view, about 8-fold more RBD is coated than spike. This is a likely explanation for the higher reactivity of RBD compared with spike. No IgG response has been detected in the control group which received CuMV<sub>T</sub> only.

The avidity of an Ab is defined as the binding strength through points of interaction. It can be quantified as the ratio of <i>K_d</i> for the intrinsic affinity over the one for functional affinity of a multiple point interaction. High avidity Abs are formed upon affinity maturation in germinal centers (GCs) and are associated with protective immunity against SARS-CoV-2 infection. To assess the avidity of the induced IgG Abs against RBD, we carried out an avidity ELISA using day 49 sera. The obtained results indicated that about 40% of the systemically induced RBD-specific IgGs are of high avidity following the i.n. vaccination (Figure 2F and G).
Intranasal immunization with CuMV<sub>TT</sub>-RBD promotes isotype switching to IgA and leads to balanced IgG subclass responses

IgG subclasses are of major importance in the immunological response against viruses because of enhancing opsonization as well as immune effector functions. In additionally, IgA plays an important role in protection against respiratory viruses as it is found in mucosal tissue, the main entrance site for these kind of viruses. The ability of the CuMV<sub>TT</sub>-RBD vaccine to induce serum IgA and IgG subclasses was evaluated by performing ELISA against RBD with sera collected at day 42. All IgG subclasses were induced in vaccinated mice with IgG1, IgG2a, and IgG2b being the dominant ones. In contrast, no IgG subclasses were detected in the control group (Figure 3A). The vaccine was also able to induce isotype switching to IgA as shown in Figure 3B. Approximately, 20% of serum IgA Abs were of high avidity as shown in Figure 3C, D.

CuMV<sub>TT</sub>-RBD induced immune sera is able to recognize VOCs

The mutation potential of SARS-CoV-2 is considered a burden in vaccine design and development, especially in terms of prolonged protection. Accordingly, we studied the capability of the induced RBD-specific IgG Abs in recognizing mutant RBDs of the different VOCs. Specifically, we have expressed and produced the following mutated RBDs: RBD<sub>K417N</sub>, RBD<sub>E484K</sub>, RBD<sub>N501Y</sub>, and RBD<sub>K417N/E484K/N501Y</sub>. Compared with RBD<sub>wt</sub>, RBD VOCs specific IgG levels were slightly lower (Figure 4). However, the difference observed...
between RBD VOCs and RBD\textsubscript{wt} IgG levels was statistically not significant (\(p = .28\)). These findings indicate a broad potential binding capacity of Abs induced by i.n. vaccination with CuMV\textsubscript{TT}-RBD.

### 3.5 RBD- and spike-specific IgG and IgA were detected locally in BAL, with RBD-specific IgG2b dominating the local subclass response

To test the ability of the CuMV\textsubscript{TT}-RBD vaccine to induce a humoral immune response in the lung mucosa; BAL was collected two weeks after the booster injection (Day 42) and assessed for RBD and spike protein specific IgG and IgA (Figure 5A–F). RBD- and spike-specific IgG Abs were detected at equal levels in the BAL (Figure 5A–C). However, IgA Abs in BAL were more abundant against RBD than against spike protein (Figure 5D–F). We have also assessed the induced RBD-specific IgG subclasses in BAL. Interestingly, the local mucosal IgG response was less balanced than the serum response and dominated by IgG2b (Figure 5G). Next, we tested the quality of the induced RBD-specific IgA in BAL samples, the results confirmed that about 60% of detected IgA Abs in BAL were of high avidity (Figure 5H–J).
FIGURE 5  RBD- and spike-specific IgG and IgA were detected locally in BAL, with RBD-specific IgG2b dominating the local subclass response (A–C) RBD- (A, C) and spike- (B, C) specific IgG titer in BAL for the groups vaccinated with CuMV<sub>TT</sub>-RBD vaccine or CuMV<sub>TT</sub> control, measured by ELISA, OD<sub>450</sub> shown in A, B, LOG<sub>10</sub> OD<sub>50</sub> shown in C. (D–F) RBD- (D, F) and spike- (E, F) specific IgA titer in BAL for the groups vaccinated with CuMV<sub>TT</sub>-RBD vaccine or CuMV<sub>TT</sub> control, measured by ELISA, OD<sub>450</sub> shown in D, E, LOG<sub>10</sub> OD<sub>50</sub> shown in F. (G) RBD-specific IgG1, IgG2a, IgG2b, and IgG3 titer in BAL for the groups vaccinated with CuMV<sub>TT</sub>-RBD vaccine or CuMV<sub>TT</sub> control, measured by ELISA, LOG<sub>10</sub> OD<sub>50</sub> shown. (H) RBD-specific IgA titer in BAL from mice immunized with CuMV<sub>TT</sub>-RBD vaccine, LOG<sub>10</sub> OD<sub>50</sub> shown. Plates in duplicates: treated with PBSTween or 7 M urea. (I) Avidity index of RBD-specific IgA titer. (J) Avidity indexes of RBD-specific IgA Abs found in BAL and serum after CuMV<sub>TT</sub>-RBD immunization. Statistical analysis (mean ± SEM) using Student’s t-test. Control group n = 6, vaccine group n = 6. One representative of 2 similar experiments is shown.
3.6 Intronasal administration of CuMV<sub>TT</sub>-RBD induced RBD-specific IgG and IgA plasma cells locally and systemically

In order to characterize the humoral immune response upon i.n. CuMV<sub>TT</sub>-RBD vaccination, specific plasma blasts were quantified in lymphoid organs and lung tissue. To this end, spleen, BM and lung were collected on day 42 and analyzed for the presence of RBD-specific IgG and IgA secreting plasma cells. As shown in Figure 6 and Figure S1, IgG secreting plasma cells were detected in all investigated tissues. Around 25 IgG secreting plasma cells were found per two million cells in spleen and BM. In lung, this ratio was ten-fold higher because the same amount of IgG plasma cells was observed while ten times less cells were seeded. IgA secreting cells were detected in all three tissues; however, at a lower level compared with IgG secreting plasma cells. RBD-specific IgA producing plasma cells in lung were thereby about ten times more abundant compared with spleen or BM (Figure 6A–C). In overall term, i.n. vaccination with CuMV<sub>TT</sub>-RBD induced a systemic humoral immune response which was accompanied by a potent local humoral immune response in the lung.

3.7 Abs induced by intranasal vaccination are capable of neutralizing SARS-CoV-2 and its variants

To test the ability of the sera of immunized mice to inhibit binding of RBD to ACE2, a biolayer interferometry competition assay was performed. Accordingly, RBD was immobilized onto anti-His biosensors and binding capacity of ACE2 to RBD in the presence of serum samples was quantified. As depicted in Figure 7A, the binding of ACE2 to RBD was reduced in the presence of sera from CuMV<sub>TT</sub>-RBD vaccinated mice. In contrast, no binding inhibition was observed in the presence of sera from CuMV<sub>TT</sub>-control mice. Percentage of ACE2 to RBD binding inhibition of individual mice is shown in Figure 7B. Interestingly, the binding inhibition correlated with RBD-specific IgG titers in serum (R value = 0.78) (Figure 7C), indicating higher RBD specific IgG titer in serum are more efficient at blocking RBD-ACE2 interaction.

For the evaluation of viral neutralization capacity, sera from CuMV<sub>TT</sub>-RBD and CuMV<sub>TT</sub> immunized mice were tested in a CPE-based neutralization assay. To this end, sera from immunized mice were assessed for their ability to prevent cytopathic effects of wt SARS-CoV-2 as well as VOCs on Vero cells in vitro. As shown in Figure 7D, all sera from CuMV<sub>TT</sub>-RBD vaccinated mice were able to neutralize the wt SARS-CoV-2 as well as SA and UK variants with high neutralization titers reaching to (1:600). In contrast, no neutralizing capacity was determined for sera from CuMV<sub>TT</sub> immunized mice.

4 DISCUSSION

The immune responses of the mucosal compartments are considered an early and essential line of defense against harmful pathogens such as SARS-CoV-2. The majority of mucosal vaccines have been administered via oral or nasal routes, with the rectal, ocular, sublingual, or vaginal routes being less often used. Ideally, an effective mucosal vaccine would induce both local and systemic responses including the distant mucosal tissues. Accordingly, i.n. administration of a vaccine appears to be a promising strategy.

Seven ongoing clinical trials are currently testing the efficacy of i.n. vaccination against COVID-19 and are based on live-attenuated virus, viral-vectors or protein subunits. Some drawbacks from using attenuated viruses or viral-vectors includes: pre-existing Abs that can impair the vaccine efficacy and the risk of reversion of the live-attenuated viruses especially in newborns or immunocompromised people. In the current study, we have tested the immunogenicity and efficacy of a conventional vaccine based on VLPs displaying RBD of SARS-CoV-2 for i.n. administration. The multi-protein VLP platform does not contain any genetic materials for replication and thus are considered a safe platform for vaccine development. Marketed vaccines against human-papilloma virus (HPV), hepatitis-B (HBV), hepatitis-E virus (HEV) and malaria are based on VLPs. Our immunologically optimized CuMV<sub>TT</sub> incorporates a universal T<sub>g</sub> cell epitope derived from tetanus toxin (TT) and are self-adjuvanted with prokaryotic ssRNA, a potent TLR7/8 agonist. We have shown in previous studies the essential role of TLR7 signaling in licensing the generation of secondary plasma cells as well as the production of systemic IgA Abs. Such processes are usually dependent on TLR7 expression in B cells; however, in contrast to systemic IgA, a successful induction of mucosal IgA requires TLR signaling in DCs. Our results here show effective induction of a systemic response of RBD-specific IgGs one week after the priming dose which increased significantly (p < .001) following the booster dose on Day 28 and antibodies showed a high degree avidity maturation. Moreover, a significant increase in RBD-specific IgA in serum was observed. Along with this, RBD-specific IgA and IgG secreting plasma cells were detected in spleen, BM, and within lung tissues. Especially, IgG secreting cells showed a high Ab secretion rate. Consequently, i.n. vaccination with CuMV<sub>TT</sub>-RBD is able to induce a strong systemic humoral immune response.

As COVID-19 presents a respiratory disease and the virus is invading through the respiratory system, an enhanced immunological local protection in the lung should be in the focus when it comes to vaccine design. However, all vaccines currently licensed are applied intramuscularly, thus ignoring this aspect. By applying CuMV<sub>TT</sub>-RBD i.n. instead of subcutaneously, we were able to induce RBD—as well as spike-specific IgA and IgG Abs in the lung. IgA localized in lung mucosa has previously been shown to be of major importance for SARS-CoV-2 neutralization. Furthermore, IgA may neutralize the virus in the lung without causing inflammation. The fact that, around 60% of the RBD specific IgA Abs in the lung were of high avidity confirms the quality of the local humoral immune response upon i.n. vaccination with CuMV<sub>TT</sub>-RBD. Interestingly, only 20% of the RBD specific IgA Abs were of high avidity in serum. This significant difference might be explained by the 2 different forms of IgA: While IgA Abs at mucosal sides are mostly found in a dimeric...
**FIGURE 6** Intranasal administration of CuMV\textsubscript{TT}-RBD induced RBD-specific IgG and IgA plasma cells locally and systemically. (A–C) Number of RBD specific IgG and IgA secreting cells per \(2 \times 10^6\) seeded cells in spleen (A), BM (B) or per \(2 \times 10^5\) seeded cells in lung (C) after immunization with CuMV\textsubscript{TT}-RBD vaccine or CuMV\textsubscript{TT} control, detected by Fluorospot. (D–F) Representative pictures of wells seeded with cells out of spleen (D), BM (E), lung (F). Statistical analysis (mean ± SEM) using Student’s t-test. Control group \(n = 6\), vaccine group \(n = 6\). One representative of 2 similar experiments is shown.

**FIGURE 7** Antibodies induced by intranasal vaccination are capable of neutralizing SARS-CoV-2 and its variants. (A) BLI-evaluation of ACE2 binding to RBD in the presence of vaccinated mice sera or controls. (B) Percentage of ACE2 to RBD binding inhibition of the individual mice vaccinated with CuMV\textsubscript{TT}-RBD, control mouse vaccinated with CuMV\textsubscript{TT} (C1), with no serum and naïve mouse. (C) Correlation of RBD-specific \(\log_{10} \text{OD}_{50}\) IgG titer with the percentage of ACE2 to RBD binding inhibition from individual mice sera. Statistical analysis (mean ± SEM) using Pearson Correlation test. (D) Neutralization titer against SARS-CoV-2 wt, SA and UK isolates. Statistical analysis (mean ± SEM) using Student’s t-test. Control group \(n = 1\) in a, b, c, \(n = 5\) for D, vaccine group \(n = 6\) for A, B, C, \(n = 5\) for D. One representative of 2 similar experiments is shown.
form, serum IgA is usually present monomeric form. In addition to IgA, IgG in the lung might also mediate protection. By passive transudation across alveolar epithelium, IgG can pass from blood into the lower lung and from there by the mucociliary escalator further be carried to the upper respiratory tract and nasal passages. However, only at high serum concentrations local protection through IgG is achieved in the lung.

While RBD-specific IgG subclass response in serum was well balanced, IgG2b was the most abundant subclass in BAL. Besides IgG2a, IgG2b is the only subclass that binds all three activating Fc receptors (FcγRI, FcγRIII, and FcγRIV) and the only inhibitory receptor (FcγRIIB) in mice. IgG2b Abs therefore mediate a wide variety of effector functions, which is of key importance in the maintenance of immune protection.

SARS-CoV-2 has the ability to mutate, albeit a proof-reading system is in place to keep the large genome of almost 30 kD genetically stable. Previously, we have described that a single N501Y mutation increased the binding affinity to ACE2, but could still be detected by convalescent sera. Contrary were the results with the E484K mutation, where no enhanced binding to ACE2 was shown but much lower recognition by convalescent sera. Triple mutant RBD (K417N/E484K/N501Y) exhibited both features: stronger affinity to ACE2 and much lower detection by convalescent sera. Since vaccines optimally mediate protection for many years, vaccine induced Abs should therefore be able to recognize new virus variants as well. In the present study, we could show that i.n. applied CuMVTT-RBD induced serum IgG Abs that are able to recognize wt RBD as well as numerous RBD VOCs.

A crucial milestone in vaccine development is effective neutralization of the virus. Sera induced after i.n. administration of CuMVTT-RBD could completely inhibit the cytopathic effect of wt SARS-CoV-2 as well as other VOCs, specifically SA and UK variants. This may be explained by the highly repetitive, rigid antigenic surface array of VLPs which are spaced by 5 nm and displaying RBD domains at a spacing of 5–10 nm. Such array of highly organized epitopes is considered a pathogen-associated structural patterns (PASPs) which are recognized by the immune system. In contrast, naturally induced Abs by SARS-CoV-2 are low in number and wane rapidly.

It has been shown that B and T cells are primed by mucosal vaccination or natural infection, express receptors which promote homing of these cells to mucosal sites as Ab-secreting cells or effector or tissue-resident T cells. We have shown in our previous studies that VLP-specific T cell response mediate specific B cell isotype-switch. Furthermore, the packaged RNA in VLPs can drive CD8+ T cells as well as Tγ1 responses. The role of T cells after i.n. vaccination using VLPs is an area we are currently investigating.

Collectively, we have shown in this study that our COVID-19 vaccine candidate CuMVTT-RBD is highly immunogenic and capable of inducing both mucosal and systemic IgG and IgA response against SARS-CoV-2 upon i.n. administration. The induced Abs could effectively recognize and neutralize wt as well as the emerging VOCs. The ability of the vaccine candidate to stop nasal viral shedding and transmission is currently under investigation. As COVID-19 pandemic continues to present a global threat to human health, it seems rational to further develop an i.n. vaccine based on conventional platform.

ACKNOWLEDGEMENT
This publication was funded by Saiba AG, the Swiss National Science Foundation (SNF grant 31003_185114 and 1ZRPZ0_194968). Open access funding provided by Inselspital Universitatssspital Bern. [Correction added on 14-May-2022, after first online publication: CSAL funding statement has been added.]

CONFLICT OF INTERESTS
M. F. Bachmann is a board member of Saiba AG and holds the patent of CuMVTT. Senta A. Walton works for Saiba AG. Mona O. Mohsen received payments by Saiba AG to work on the development of vaccines against Dengue and SARS-CoV-2. Martin F. Bachmann and Mona O. Mohsen are shareholder of Saiba AG.

AUTHOR CONTRIBUTIONS
MB, MM, PK, and DR involved in design of experiments. DR, PK, AN, IB, ACV, XC, AM, FV, GR, SW, and MM involved in methodology. DR, PK, MM, MB, ACV, AN, AM, FV, GR, EM, MV, and AZ involved in acquisition of data, and interpretation and analysis of data. DR, PK, MM, and MB involved in writing, revision, and editing of manuscript. EM and AZ involved in technical, material and tool support. MB, MM, and MV involved in study supervision. All authors read and approved the final manuscript.

ORCID
Ina Balke https://orcid.org/0000-0002-5171-7744
Xinyue Chang https://orcid.org/0000-0001-7027-3889
Monique Vogel https://orcid.org/0000-0002-5219-4033
Martin F. Bachmann https://orcid.org/0000-0003-4370-2099
Mona O. Mohsen https://orcid.org/0000-0003-3510-9148

REFERENCES
1. Swan DA, Bracis C, Janes H, et al. COVID-19 vaccines that reduce symptoms but do not block infection need higher coverage and faster rollout to achieve population impact. Sci Rep. 2021:11:1-9.
2. Peiris JSM, Chu CM, Cheng VCC, et al. Clinical progression and viral load in a community outbreak. Lancet. 2003;361:1767-1772.
3. Sun K, Gu L, Ma L, Duan Y. Atlas of ACE2 gene expression reveals novel insights into transmission of SARS-CoV-2. Heliyon. 2021;7(1):e05850.
4. Jia HP, Look DC, Shi L, et al. ACE2 receptor expression and severe acute respiratory syndrome coronavirus infection depend on differentiation of human airway epithelia. J Virol. 2005;79:14614-14621.
5. HarrisonAG,LinT,WangP.Mechanisms of SARS-CoV-2 Transmission and Pathogenesis. Trends Immunol. 2020;41:1100-1115.
6. Ziegler CG, Allon SJ, Nyquist SK, et al. SARS-CoV-2 receptor ace2 is an interferon-stimulated gene in human airway epithelial cells and is detected in specific cell subsets across tissues. Cell. 2020;181:1016-1035.
7. Krammer F. SARS-CoV-2 vaccines in development. Nature. 2020;586:516-527.
8. https://www.clinicaltrials.gov/
9. Renegar KB, Small PA, Boykins LG, Wright PF. Role of IgA versus IgG in the control of viral infection in the murine respiratory tract. *J Immunol*. 2004;173:1978-1986.

10. Rodriguez A, Tjärnlund A, Ivanji J, et al. Role of IgA in the defense against respiratory infections: IgA deficient mice exhibited increased susceptibility to intranasal infection with *Mycobacterium bovis* BCG. *Vaccine*. 2005;23:2565-2572.

11. Allie SR, Bradley JE, Mudunuru U, et al. The establishment of resident memory B cells in the lung requires local antigen encounter. *Nat Immunol*. 2019;20:97-108.

12. vanDoremalen N, et al. Intranasal ChAdOx1 nCoV-19/AZD1222 vaccination reduces shedding of SARS-CoV-2 D614G in rhesus macaques. *Serv Biol*. 2021. doi:10.1101/2021.01.09.426058

13. Mohn KGI, Smith I, Sjursen H, Cox RJ. Immune responses after vaccination reduces shedding of SARS-CoV-2 D614G in rhesus macaques. *Sci Transl Med*. 2021;13(577):eabf1555. doi:10.1126/scitranslmed.abf1555

14. Bessa J, et al. Efficient induction of mucosal and systemic immune responses by virus-like particles administered intranasally: Implications for vaccine design. *Eur J Immunol*. 2008;38:114-126.

15. Zeltins A, West J, Zabel F, et al. Incorporation of tetanus-epitope into virus-like particles achieves vaccine responses even in older recipients in models of psoriasis, Alzheimer’s and Cat Allergy. *NPJ Vaccines*. 2017;2:1-12.

16. Wiuff C, Thorberg BM, Engvall A, Lind P. Immunochemical analyses of serum antibodies from pig herds in a Salmonella non-endemic region. * Vet Microbiol*. 2002;85:69-82.

17. Sun F, Xiao G, Qu Z. Murine bronchoalveolar lavage. *Physiol Behav*. 2017;176:139-148.

18. Manenti A, et al. Evaluation of SARS-CoV-2 neutralizing antibodies using a CPE-based colorimetric live virus micro-neutralization assay in human serum samples. *J Virol*. 2020;92:2096-2104.

19. Manenti A, Molesti E, Maggetti M, et al. The theory and practice of the viral dose in neutralization assay: Insights on SARS-CoV-2 ‘doublethink’ effect. *J Virol Methods*. 2020;297:114261.

20. Moisen MO, Augusto G, Bachmann MF. The 3Ds of virus-like particle based-vaccines: “Design, Delivery and Dynamics”. *Immunol Rev*. 2020;296:155-168.

21. Moisen MO, Balke I, Zinkhan S, et al. A scalable and highly immunogenic virus-like particle-based vaccine against SARS-CoV-2. *Allergy Eur J Allergy Clin Immunol*. 2022;77(1):243-257. doi:10.1111/all.15080

22. Moisen MO, Rothen D, Balke I, et al. Neutralization of MERS coronavirus through a scalable nanoparticle vaccine. *NPJ Vaccines*. 2021;6:1-9.

23. Bauer G. The potential significance of high avidity immunoglobulin G (IgG) for protective immunity towards SARS-CoV-2. *Int J Infect Dis*. 2021;106:61-64.

24. Hazenbos WL, et al. Murine IgG1 complexes trigger immune effector functions predominantly via Fc gamma RII (CD16). *J Immunol*. 1998;161:3026-3032.

25. Chang X, Augusto GS, Liu X, et al. BNT162b2 mRNA COVID-19 vaccine induces antibodies of broader cross-reactivity than natural infection, but recognition of mutant viruses is up to 10-fold reduced. *Allergy Eur J Allergy Clin Immunol*. 2021;76(9):2895-2998. doi:10.1111/ajr.14893

26. Russell MW, Moldoveanu Z, Ogra PL, Mestecky J. Mucosal immunity in COVID-19: A neglected but critical aspect of SARS-CoV-2 infection. *Front Immunol*. 2020;11:1-5.

27. Czerkinsy C, Holmgren J. Mucosal delivery routes for optimal immunization. *Curr Top Microbiol Immunol*. 2012;354:1-18.

28. Moisen MO, Zha L, Cabrál-Miranda G, Bachmann MF. Major findings and recent advances in virus-like particle (VLP)-based vaccines. *Semin Immunol*. 2017;34:123-132.

29. Krueger CC, Zabel F, Keller E, et al. RNA and toll-like receptor 7 license the generation of superior secondary plasma cells at multiple levels in a B cell intrinsic fashion. *Front Immunol*. 2019;10:1-13.

30. Bessa J, Jegerlehner A, Hinton HJ, et al. Alveolar macrophages and lung dendritic cells sense RNA and drive mucosal IgA responses. *J Immunol*. 2009;183:3788-3799.

31. Zha L, Chang X, Zhao H, et al. Development of a vaccine against sars-cov-2 based on the receptor-binding domain displayed on virus-like particles. *Vaccines*. 2021;9:1-14.

32. Wang Z, Lorenzi JCC, Muecksch F, et al. Enhanced SARS-CoV-2 neutralization by dimeric IgA. *Sci Transl Med*. 2021;13(577):eabf1555. doi:10.1126/scitranslmed.abf1555

33. Steffen U, Koeleman CA, Sokolova MV, et al. IgA subclasses have different effector functions associated with distinct glycosylation profiles. *Nat Commun*. 2020;11. doi:10.1038/s41467-019-13992-8

34. Gosselin EJ, Bitsaktsis C, Li Y, Iglesias BV. Fc receptor-targeted mucosal vaccination as a novel strategy for the generation of enhanced immunity against mucosal and non-mucosal pathogens. *Arch Immunol Ther Exp (Warsz)*. 2009;57:311-323.

35. Robson F, Khan KS, Le TK, et al. Coronavirus RNA proof-reading: Molecular basis and therapeutic targeting. *Mol Cell*. 2020;79:710-727.

36. Vogel M, Augusto G, Chang X, et al. Molecular definition of severe acute respiratory syndrome coronavirus 2 receptor-binding domain mutations: Receptor affinity versus neutralization of receptor interaction. *Allergy*. 2022;77(1):143-149. doi:10.1111/all.15002

37. Bachmann MF, Moisen MO, Zha L, Vogel M, Speiser DE. SARS-CoV-2 structural features may explain limited neutralizing-antibody responses. *NPJ Vaccines*. 2021;6:1-5.

38. Bachmann MF, Jennings GT. Vaccine delivery: A matter of size, geometry, kinetics and molecular patterns. *Nat Rev Immunol*. 2010;10:787-796.

39. Tay MZ, Poh CM, Rénia L, MacAry PA, Ng LFP. The trinity of the viral dose in neutralization assay: Insights on SARS-CoV-2 ‘doublethink’ effect. *Mol Immunol*. 2019;10:787-796.

40. Pizzolla A, Nguyen TH, Wakin LM, et al. Resident memory CD8+ T cells in the upper respiratory tract prevent pulmonary influenza virus infection. *Sci Immunol*. 2017;2:1-14.

41. Skibinski DAG, Hanson BJ, Lin Y, et al. Enhanced Neutralizing Antibody Titers and Th1 Polarization from a Novel Escherichia coli Derived Pandemic Influenza Vaccine. *PLoS One*. 2013;8:e76571.

**SUPPORTING INFORMATION**

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**How to cite this article:** Rothen DA, Krenger PS, Nonic A, et al. Intranasal administration of a virus like particles-based vaccine induces neutralizing antibodies against SARS-CoV-2 and variants of concern. *Allergy*. 2022;77:2446–2458. doi:10.1111/all.15311