**INTRODUCTION**

Coronavirus disease 2019 (COVID-19), a disease caused by severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2), is continuously spreading worldwide and has become a serious threat to global health. Therefore, an effective vaccine against SARS-CoV-2 is urgently needed, while only a handful have already been approved. The SARS-CoV-2 genome encodes four structural proteins—spike (S), membrane (M), envelope (E), and nucleocapsid (N) (1). Similar to the S protein of SARS-CoV, the S protein of SARS-CoV-2 mediates the entry of the virus into the host cell. The S protein binds to the angiotensin-converting enzyme (ACE) receptor on the host cell through its receptor binding domain (RBD) (1, 2). As neutralizing antibodies against the RBD can block the binding of the virus to host cells, the RBD is considered the main target antigen for vaccines against SARS-CoV-2 (3, 4). Furthermore, RBD-based vaccines are expected to be safer than S protein–based vaccines, as they are not likely to induce T helper 2 (T_H2)–type immunopathology through antibody-dependent enhancement (5–7).

The N proteins of coronaviruses are highly immunogenic and are abundantly expressed throughout the course of the infection (8). According to a recent study, antibodies against the N protein are more sensitive indicators than anti–S protein antibodies for the detection of early SARS-CoV-2 infection (9). Moreover, the N protein of SARS-CoV-2 is believed to induce T cell responses. An adenoviral vector encoding the N protein induced T cell responses in rhesus monkeys, and mice vaccinated with the N protein exhibited T cell proliferation and cytotoxic T cell responses (10, 11). However, in vivo protection conferred by vaccines containing the N protein has not yet been experimentally validated. Thus, it is still ambiguous whether immunization with the N protein combined with RBD would manifest better or worse protective effects.

Currently, numerous SARS-CoV-2 vaccines based on different platforms are being developed owing to urgent demand and importance (12). Subunit vaccines are the most commonly used vaccines owing to their safety parameters. Many commercially approved vaccines against diseases caused by viruses such as human papilloma-virus, hepatitis B, influenza, and varicella zoster are being manufactured using this platform (13). However, because of their lower immunogenicity compared with that of live attenuated vaccines, subunit vaccines need adjuvants, carriers, or engineering of the proteins to boost their immunogenicity (14).

Here, we developed a previously unidentified SARS-CoV-2 subunit vaccine (RBD-P2) by fusing the P2 epitope of tetanus toxoid with the RBD of the spike protein to increase its immunogenicity. A previous study shows that the inclusion of the universal toxoid CD4+ T cell epitope P2 substantially increases the immunogenicity...
of a recombinant rotavirus subunit vaccine (15). Furthermore, we tested the efficacy of RBD-P2 with or without the N protein in mice, rats, and nonhuman primates (NHPs).

RESULTS
Characterization of recombinant RBD-P2 and N
We designed and synthesized RBD fused with the P2 epitope of tetanus toxoid (RBD-P2) as a vaccine candidate against SARS-CoV-2. The RBD-P2 and N proteins of SARS-CoV-2 were expressed and purified using High Five insect cells and the baculovirus-mediated Bac-to-Bac expression system. A human albumin signal peptide sequence was fused with the RBD-P2 and N protein sequences to enable the secretion of the proteins (Fig. 1A and fig. S1). Recombinant RBD-P2 and N proteins were successfully purified from the culture supernatant (Fig. 1B). The purity of the final protein preparation was approximately 90%, and the molecular weights of the purified RBD-P2 and N proteins were approximately 32 and 53 kDa, respectively (Fig. 1B). Both proteins were presumably glycosylated as their estimated molecular weights were higher than the theoretical molecular weights calculated on the basis of the amino acid sequences. The binding ability of RBD-P2 was measured using a functional enzyme-linked immunosorbent assay (ELISA). Immobilized RBD-P2 (2 µg/ml) bound to human anti–SARS-CoV-2 RBD-neutralizing antibody (ACROBiosystems, catalog no. SAD-S35) and mouse anti–SARS-CoV-2 spike-neutralizing antibody (Sino Biological, catalog no. MM57) (Fig. 1C). The median effective concentration (EC50) of RBD-P2 for the human antibody was 56.5 µg/ml, and that for the murine antibody was 20 µg/ml. These values were similar to those of RBD-positive protein (Sino Biological, catalog no. 40592-V08B; EC50: 49.4 µg/ml for human antibody and EC50: 22.3 µg/ml for murine antibody). The binding affinity of RBD-P2 for human ACE2 (hACE2) was calculated as follows: 

\[ K_D = \frac{K_{on}}{K_{off}} = \frac{1.1 \times 10^5 \text{ M}^{-1} \text{ s}^{-1}}{4.5 \times 10^{-4} \text{ s}^{-1}} = 2.4 \times 10^3 \text{ M}^{-1} \]

(Fig. 1D). The binding
Analysis of humoral and cellular immune responses in mice immunized with RBD-P2 or RBD-P2/N

In this study, we chose alum as an adjuvant because of its strong safety record (16). First, we compared the immune responses of RBD-P2 and RBD. RBD was expressed and purified from insect cells using the Bac-to-Bac expression system as described for RBD-P2 (details are in Materials and Methods). After immunization in rats, RBD-P2 + alum induced a significantly higher neutralizing antibody titer than RBD + alum (fig. S2B). Therefore, all vaccine tests were conducted using RBD-P2. We confirmed the complex formation of RBD-P2 and RBD-P2/N with alum by using holotomography and dynamic light scattering (fig. S3A). As shown in fig. S3A, unlike alum alone, which had more scattered clusters, both RBD-P2 and RBD-P2/N with alum clustered in long rod-like shapes. We also used dynamic light scattering analysis to confirm the successful formulation of RBD-P2 and RBD-P2/N with alum. We could detect one main peak in both RBD-P2 + alum and RBD-P2/N + alum. This result indicates that most of the proteins formed a complex with alum (fig. S3B).

To test the potential of RBD-P2 as a SARS-CoV-2 vaccine candidate, mice were intramuscularly immunized twice with 30 μg of RBD-P2 + alum (mG2) 3 weeks apart. To confirm the effectiveness of the N protein, mice were immunized with 30 μg of RBD-P2 and 3 μg of N + alum (mG3), designated as RBD-P2/N. Mice immunized twice with alum without antigen were designated as mG1. All mice were euthanized 3 weeks after the booster dose (fig. 2A).

To assess humoral responses, serum samples were collected at the time of euthanization, and RBD-P2- and N-specific immunoglobulin G1 (IgG1) and IgG2c antibody levels were analyzed using ELISA. As shown in fig. 2 (B to E), RBD-P2 + alum–immunized mice exhibited induction of IgG1 and IgG2c against RBD-P2. The fusion of N with RBD-P2 (RBD-P2/N) slightly reduced the levels of RBD-specific IgG1 but not IgG2c (fig. 2, B to E). Understandably, N-specific antibodies were detected only in mG3.

Next, to determine cell-mediated immune responses, splenocytes from immunized mice were stimulated with a mix of SARS-CoV-2 RBD-P2 and N peptide mix (table S1) for 2 days to detect antigen-specific T cell responses. As shown in fig. 2 (F to H), the percentages of both CD4+ and CD8+ CD44hiCD62Lhi central memory type T cells (T CM ) were increased in mG2 and mG3 compared with those in mG1. The expression of CD103, a marker of tissue-residential memory T cells, was increased in the CD4+ T CM of mG2 and mG3 (fig. 2F). The expression of CD69, a T cell activation marker, was increased only in CD4+ T cells of mG3 and not in mG1 and mG2 (fig. 2G). To further evaluate the T H1/2 balance, interferon-γ (IFN-γ)– and interleukin-4 (IL-4)–producing cells were measured by enzyme-linked immunosorbent spot (ELISpot) assays. Notably, IFN-γ– and IL-4–producing cells were increased in mG3 compared with their numbers in mG2, even when splenocytes were stimulated with RBD alone (fig. 2H). To test whether immunization with the N protein could induce protective immunity, hACE2 mice were intramuscularly immunized twice with 30 μg of the N protein with alum at 2-week intervals and challenged with SARS-CoV-2 1 week after the second immunization (fig. S4A). Viral titers in the lungs were reduced in N protein–immunized mice, and parenchymal inflammation scores of mG2 were slightly reduced compared with those of mG1. Thus, immunization with N protein + alum could induce a certain degree of protective effect against SARS-CoV-2 (fig. S4, B to E).

Additional immunization with the N protein increased the levels of neutralizing antibodies in rats

To further assess the immune responses elicited by RBD-P2 or RBD-P2/N, rats were injected with 30 or 50 μg of RBD-P2 (rG2 and rG3) or in combination with N at a ratio of 10:1 (rG4 and rG5) together with alum. The immunization was performed twice (with a 3-week interval in between), and the rats were euthanized 6 weeks after the first injection (fig. 3A). Both anti–RBD-P2 IgG and anti-N IgG were detected 6 weeks after the first immunization. Immunization with 50 μg of RBD-P2 + alum induced slightly higher levels of RBD-P2–specific antibodies than that induced by 30 μg RBD-P2 + alum, although the difference was not statistically significant (fig. 3B). RBD-P2/N (rG4 and rG5) + alum immunization yielded an approximately 20 to 50% increase in reduction rates compared with those achieved using each of its RBD-P2 counterparts (rG2 and rG3, respectively) (fig. 3C). The number of IFN-γ–producing cells in the splenocytes of immunized rats increased in all immunized groups (rG2 to rG5) compared with that observed in rG1 after stimulation with RBD-P2, the peptide mix, or the P2 peptide (table S1). However, no significant differences were observed between RBD-P2/N and RBD-P2 groups (fig. 3D).

To assess the immunogenicity and toxicity of RBD-P2 or RBD-P2/N in rats, the animals were immunized three times at 2-week intervals with RBD-P2 + alum (rG2) or RBD-P2/N + alum (rG3) according to the guidelines for SARS-CoV-2 vaccine development established by the Korean Food and Drug Administration (fig. S5A). In this experiment, rG3 produced higher levels of anti–RBD-P2 IgG than rG2 at 2 and 4 weeks after the first immunization in females (fig. S5B). However, 6 and 8 weeks after immunization, the levels of anti–RBD-P2 IgG produced in response to immunization with rG2 were markedly increased and reached levels similar to those of rG3. In addition, biochemical analysis revealed that neither RBD-P2 + alum nor RBD-P2/N + alum induced significant toxicity (fig. S5C).
were slightly higher than those of nG2 (Fig. 4B). Neutralizing antibody responses against SARS-CoV-2 were markedly increased in nG2 and nG3 at 6 weeks; however, they were reduced at 9 weeks in both nG2 and nG3. Although additional immunization with N increased the neutralizing antibody titers in NHPs, the increase was not statistically significant (Fig. 4C). Notably, the neutralizing antibody titers induced by RBD-P2 or RBD-P2/N were significantly higher than those of a panel of serum samples from convalescent patients [at 6 weeks, the geometric mean titer (GMT) of nG2 = 507.96 and the GMT of nG3 = 1015.93; at 9 weeks, the GMT of nG2 = 163, the GMT of nG3 = 320, and the overall convalescent serum of GMT = 109.4; all GMT values were calculated on the basis of plaque reduction neutralization test (PRNT<sub>50</sub>) values] (Fig. 4C).

To assess T cell responses, the number of IFN-γ–producing cells in peripheral blood mononuclear cells (PBMCs) were evaluated at 3 and 6 weeks after immunization. The average number of RBD-P2–specific IFN-γ–producing cells was slightly higher in nG3 than in nG2 at 3 and 6 weeks. As expected, IFN-γ–producing cells in response to the N protein were detected only in nG3, as the animals in this group were immunized with N (Fig. 4D). We also measured the levels of various cytokines in the supernatants of PBMCs from immunized NHPs; IFN-γ, IL-4, IL-5, IL-13, and tumor necrosis factor α (TNF-α) were significantly increased in nG3 compared with nG2 (Fig. 4E).

We further examined the cytokine responses at 3 and 6 weeks after immunization. The average number of IFN-γ–producing cells was significantly increased in nG3 compared with nG2 (Fig. 4D). We also measured the levels of various cytokines in the supernatants of PBMCs from immunized NHPs; IFN-γ, IL-4, IL-5, IL-13, and tumor necrosis factor α (TNF-α) were significantly increased in nG3 compared with nG2 (Fig. 4E).
factor-α production in PBMCs from nG2 and nG3 was increased compared to those of nG1 (Fig. 4E).

To confirm the protective effect of RBD-P2 + alum (nG2) and RBD-P2/N + alum (nG3), NHPs were challenged with a total of 10 ml of virus [2.6 × 10⁶, 50% tissue culture infectious dose (TCID₅₀)/ml] via intratracheal (4 ml), oral (5 ml), conjunctival (0.5 ml), and intranasal (0.5 ml) routes at 9 weeks after the first immunization. Of note, RBD-P2/N + alum–immunized NHPs showed slightly faster SARS-CoV-2 clearance in throat and lung samples than that induced by RBD-P2 + alum, although it was not statistically significant (Fig. 4, F and G). Also, live SARS-CoV-2 particles were not detected in the lungs of either nG2 or nG3 and were barely detectable in the nose and throat in postchallenges (Fig. 4, F and G).

Moreover, the body temperature of immunized NHPs (nG2 and nG3) was not increased compared with that of nG1 after the virus challenge (Fig. 4H). In addition, SARS-CoV-2 genomic RNA (gRNA) was reduced in total lung analyses of nG2 and nG3 compared with that in the lungs of nG1 (Fig. S6A). Further, nG3 showed lower gRNA levels than nG2 in throat swabs and some parts of lung tissue (fig. S6B). NHPs did not show any significant weight loss after challenge (fig. S7A). Hematological and biochemical analyses, including absolute lymphocyte counts, revealed no notable changes in the vaccinated groups (nG2 and nG3) compared with those in control NHPs (nG1) (fig. S7B).

**DISCUSSION**

Currently, many types of vaccine candidates against SARS-CoV-2, including inactivated, vectored, nucleotide-based, and recombinant, are being tested in preclinical or clinical trials. Most of these candidates involve at least a portion of the spike protein, and it is still unclear whether immunization in combination with the N protein of SARS-CoV-2 would be beneficial for protection against SARS-CoV-2 challenges (17–19). In this study, we developed a previously unknown subunit vaccine against SARS-CoV-2 that involves the RBD linked to the P2 epitope of tetanus toxoid to enhance immunogenicity. Consistent with the previous finding, we noted higher neutralizing antibody titers against SARS-CoV-2 in rats when immunized with RBD-P2 than that with RBD. In addition, RBD-P2 + alum effectively induced a neutralizing antibody response in NHPs and provided protection against SARS-CoV-2 challenge. Thus, it seems that immunization with RBD-P2 + alum can induce effective and sufficient virus-neutralizing activity and provide protection. However, we could not ascertain whether RBD-P2 + alum induces long-term humoral immunity, as NHPs were challenged with SARS-CoV-2 6 weeks after boosting.

N proteins are considered highly immunogenic, and the gene encoding the N protein is more stable than that encoding the spike protein (20). A DNA vaccine encoding SARS-CoV N protein was reported to induce strong N-specific humoral and cellular immune...
responses in C57BL/6 mice and achieved a significant reduction in viral titers following challenge (10). On the contrary, Buchholz et al. reported that immunization with the N protein did not induce a neutralizing antibody response and failed to protect against infection in a SARS-CoV hamster model (21). To test whether the inclusion of the N protein as a vaccine antigen could generate more balanced immune responses, we compared the immune responses elicited by RBD-P2 and RBD-P2/N immunization in three different animals—mice, rats, and NHPs. Of note, RBD-P2/N + alum–immunized mice exhibited enhanced T cell–mediated immune responses. Immunization with the N protein and RBD-P2 increased CD69 expression in CD4+ T cells as well as IFN-γ and IL-4 production in mice. In NHPs, PBMCs in RBD-P2/N + alum–immunized animals produced IFN-γ when they were stimulated with the N protein; however, production of Th2-type cytokines (IL-4, IL-5, and IL-13) was very low. Consistent with these findings, the numbers of IFN-γ–producing cells were increased in the RBD-P2/N + alum–immunized NHPs in throat swab samples. Thus, increased T cell activation and Th1-prone responses may result in faster viral clearance and constrain viral spread.

In addition to inducing T cell responses, we expect that vaccines based on the N protein may provide effective protection against SARS-CoV-2 variants since the N protein is more conserved than the spike protein. Currently, SARS-CoV-2 spike protein variants have emerged, and it was reported that these variants are resistant to the sera of convalescent patients (22). Thus, the combination of
an RBD-P2/N immunization may be advantageous in overcoming escape mechanisms of SARS-CoV-2 because of gene mutations encoding the spike protein.

There were limitations to this study. We immunized NHPs with a relatively high dose of RBD-P2 (50 μg) and tested the efficacy of the RBD-P2 and N combination only at a 10:1 ratio. Thus, the effectiveness of the N protein could vary depending on the dose of RBD-P2. Further studies are needed to determine the optimal dose or ratio involving the N protein. Currently, we are evaluating the RBD-P2–based subunit vaccine in clinical trials (phase 1, no. NBP2001) because RBD-P2 and RBP-P2/N showed similar protective efficacy in NHPs.

In summary, our data demonstrated that RBD-P2 + alum immunization elicits a robust neutralizing antibody response and provides complete or near-complete elimination of live SARS-CoV-2 virus particles. Moreover, our findings provide insight into the potential use of the N protein in the development of SARS-CoV-2 vaccines.

**MATERIALS AND METHODS**

**Recombinant baculovirus production**

The Bac-to-Bac expression system (Invitrogen, Waltham, MA, USA; catalog no. 10359-016) was used to produce SARS-CoV-2 RBD-P2 and N proteins. The gene sequences of RBD and N were obtained from the genome sequence of the Wuhan seafood market pneumonia virus, Wuhan-Hu-1. The gene fragments were synthesized at GenScript (Piscataway, NJ, USA) and cloned into the pFastBac vector, and the recombinant plasmid was prepared using Escherichia coli DH5α cells. The recombinant pFastBac plasmid, which served as a donor plasmid, was transformed into E. coli DH10Bac competent cells ( Gibco, NY, USA; catalog no. 10361012) harboring a shuttle vector (bacmid). The recombinant bacmid was transfected into insect cells to produce recombinant baculoviruses. For virus production, Sf9 cells (ATCC CRL-1711TM, lot no. 62058694) derived from Spodoptera frugiperda were used. Cell culture supernatant was recovered 3 days after transfection to obtain recombinant baculoviruses.

**Production of RBD-P2 and N protein**

Sf9 cells were infected with a baculovirus containing the RBD-P2 gene and cultured in a flask. After 6 days, the baculovirus was harvested by centrifugation and filtration through 0.2-μm filters. An aliquot of the clarified baculovirus preparation was used for RBD-P2 antigen production. The titer of the baculovirus was obtained by rapid titration assay. High Five cells were cultured in a 10-liter bioreactor for RBD-P2 antigen production. The Hi-5 cells were infected with baculovirus harboring the RBD-P2 gene at a multiplicity of infection of 1 to 3 and cultured for 3 days. The cells were removed by centrifugation and filtration through 0.2-μm filters. The RBD-P2 antigen was purified using three columns—anion exchange, hydrophobic interaction, and multifunctional resins—from the separated cell culture supernatant. The RBD-P2 antigen was concentrated for formulation using the tangential flow filtration (TFF) system. The TFF system was used as an adjuvant at a 1:10 ratio. The N antigen was produced using the same baculovirus and insect system. Its blood vessel invasion and upstream processes were the same as those for RBD-P2. The N antigen was purified from the separated cell culture supernatant using an anion exchange column and concentrated for formulation using the TFF system. The formulation for the adjuvant was the same as that for RBD-P2.

**Mouse handling and care**

Female BALB/c and C57BL/6 mice aged 8 to 10 weeks were purchased from Dae-Han Bio-Link (Eumseong-Gun, Chungchonbuk-do, Republic of Korea) or Samtako (Osan-si, Gyeonggi-do, Republic of Korea). The mice were housed in the animal facility at the Catholic University of Korea under specific pathogen–free conditions at 21°C to 22°C and a 12/12-hour light/dark illumination cycle. The animal facility is fully certified by the Korean Association for Laboratory Animals, and the mice were handled in accordance with the protocols approved by the Catholic University of Korea. All experimental procedures performed on mice in this study followed the guidelines of the Institutional Animal Care and Use Committee of the Catholic University of Korea (approval nos. CUK-IBC-2020-001, CUK-IACUC-2020-014, and CUK-IACUC-2020-015).

**Immunization of mice and rats**

BALB/C or C57BL/6 mice were intramuscularly injected with 30 μg of RBD-P2 with or without 3 μg of N formulated with 300 μg of alum (Croda, Yorkshire, UK) twice at 3-week intervals. Sera were collected after the mice were euthanized. For the challenge, hACE2 transgenic mice were intramuscularly injected in the upper thigh once with 30 μg of N formulated with 300 μg of alum two times at 0 and 2 weeks. The hACE2 transgenic mice were challenged with the SARS-CoV-2 virus [5 × 10⁵ plaque-forming units (PFU) per mouse] at 3 weeks after the first immunization, and their body weight and survival rate were determined. All challenged mice were maintained in Biosafety Level 3 (BSL-3) facilities at the Korea Zoonomy Research Institute of Jeonbuk National University (JBN). All mouse experimental procedures conducted in this experiment followed the guidelines of the Institutional Animal Care and Use Committee of JBN (approval no. JBN2020-0155).

Rats aged ~8 to 12 weeks were purchased from Orient Bio Inc. (Seongnam, Republic of Korea) and were intramuscularly injected with 30 or 50 μg of RBD-P2 with or without N at a 1:10 ratio. The proteins were combined with 300 or 500 μg of alum (twice at 3-week intervals). Sera were collected from the rats at the indicated time points and at the time of sacrifice for the detection of serum antibodies against RBD-P2 and N.

**NHP care and study design**

Nine Cambodian-origin cynomolgus macaques (Macaca fascicularis; three males and six females; aged 6 to 7 years) were used in this study. All animals were housed in infrastructure facilities in the Korea National Primate Research Centre (KNPRC) at the Korea Research Institute of Bioscience and Biotechnology (KRIBB), under animal biosecurity level–2 (ABL-2) or ABL-3, as required. Macaques were anesthetized with a combination of ketamine sodium (10 mg/kg) and tiletamine/zolazepam (5 mg/kg) for immunization, viral challenges, swabs, and blood collection.

As illustrated in Fig. 4A, the macaques were randomly assigned to three sex-balanced groups. The animals were intramuscularly immunized in the upper thigh at weeks 0 and 3 with one of the following formulations: (i) 500 μg of alum (Croda), (ii) 50 μg of RBD-P2 with 500 μg of alum, and (iii) 50 μg of RBD-P2 and 5 μg of N with 500 μg of alum. After 6 weeks from the last immunization, all animals were challenged with a total of 10 ml of the SARS-CoV-2
5% CO₂ for 16 to 18 hours. Six weeks after the first immunization, Vero cells were seeded in a six-well plate and incubated at 37°C and 5% CO₂ for 1 hour and then inoculated into Vero cells for 1 hour. Plasma samples were inactivated at 56°C for 30 min and 2-fold serially diluted from 20-fold diluted samples. Diluted plasma samples were mixed with SARS-CoV-2 (50 to 100 PFU per well) at 37°C and 5% CO₂ for 1 hour and then incubated into Vero cells for 1 hour under the same incubation conditions. After removing the plasma-virus mixture, 1.5% low–melting temperature agarose in DMEM, supplemented with 2% FBS, was added, and the cells were incubated for 3 days. The cells were fixed with 10% neutral formalin solution overnight and stained with 0.1% crystal violet solution. For each dilution, the number of plaques was counted, and the percentage of neutralization in test samples was compared with that in the control.

Isolation of PBMCs from NHPs

Blood samples were collected in EDTA-containing tubes, 0 and 2 weeks after immunization. PBMCs were isolated by centrifugation at 3500g for 20 min with gentle acceleration using Ficoll-Hypaque (Lymphoprep; Axis-Shield, 1114545). Red blood cells in the PBMCs were lysed in Ammonium-Chloride-Potassium (ACK) lysis buffer (Gibco, Gaithersburg, MD, USA) at 20°C to 25°C for 5 min. The PBMCs were washed once with flow cytometry staining buffer [1× phosphate-buffered saline (PBS) with 2% FBS] and resuspended in RPMI medium supplemented with 10% FBS and penicillin/streptomycin (1×).

Human blood samples

In this study, blood and serum samples were collected from 31 convalescent patients with COVID-19 enrolled from Seoul National University Hospital, Seoul Metropolitan Government–Seoul National University Boramae Medical Center, Kyungpook National University Hospital, Chungnam National University Hospital, and Chung-Ang University Hospital. The study was approved by the Institutional Review Boards of the mentioned hospitals (IRB nos. 2042-005-413, CNUH-2020-130, 2020-05-006, and 20–2020-41). Serum samples were heat-inactivated (20 min at 60°C) before using in binding or neutralizing assays.

Flow cytometry

Mouse splenocytes (1 × 10⁶) were stained in PBS containing 0.5% FBS for flow cytometry. In total, 0.5 μg of Fcy Receptor blocker (anti-CD16/CD32 antibody) was added 15 min before fluorochrome-conjugated antibody staining. The following antibodies were obtained from Invitrogen (Walham, MA, USA): anti-mouse CD4, CD8, CD44, CD62L, and CD103. Anti-mouse CD69 antibodies were purchased from BD Bioscience (San Diego, CA, USA). To distinguish live cells from dead cells, the cells were stained with Fixable Viability Dye eFluor 520 (Invitrogen). After washing, the cells were analyzed using a Cytæk Aurora Flow Cytometer (Cytæk, Fremont, CA, USA), and the results were interpreted using SpectroFlo software (Cytæk).

ELISpot assay

ELISpot assay for mice

Splenocytes of immunized mice were stimulated with 500 ng per well of RBD-P2 or 600 ng per well of N peptide mix for 48 hours. IL-4- and IFN-γ–secreting cells were detected using the mouse IL-4 ELISpot kit (R&D Systems, Minneapolis, MN, USA) and the mouse IFN-γ ELISpot kit (R&D Systems, Minneapolis, MN, USA) according to the manufacturer’s instructions.

ELISpot assay for rats

Splenocytes of immunized rats were stimulated with 300 ng per well of RBD-P2 or 300 ng per well of P2 peptides for 60 hours. IFN-γ–secreting cells were detected using a rat IFN-γ ELISpot kit (R&D Systems, Minneapolis, MN, USA) and the manufacturer’s instructions.

ELISpot assay for NHPs

PBMCs of immunized NHPs were stimulated with 300 ng per well of RBD or 300 ng per well of N or 1 μg per well of P2 peptides for 60 hours. IFN-γ–secreting cells were detected using primate IFN-γ ELISpot kit (R&D Systems) according to the manufacturer’s instructions. The resultant spots were counted on a computer-assisted AID EliSpot Reader System (AID, Strasbourg, Germany).

Mouse ELISA

Antigen-specific IgG1 and IgG2c levels in mouse serum were measured using ELISA. First, a 96-well plate (Corning, NY, USA) was coated with either S, N, or RBD-P2 (100 ng/μl) and incubated at 4°C overnight. After incubation, the wells were blocked with 100 μl of blocking buffer (PBS containing 1% bovine serum albumin) at room temperature for 1 hour. Diluted serum samples were added to the wells, and the plates were incubated at room temperature for 2 hours. After incubation, the wells were washed thrice with 200 μl of PBS containing 0.05% Tween 20 (PBS-T). Horseradish peroxidase–conjugated anti-mouse IgG1 (Bethyl Laboratories, Montgomery, TX, USA) and IgG2c (Novus Biologicals, Centennial, CO, USA) antibodies (diluted 1:1000 to 1:10,000 in PBS) were added to the plates, which were then incubated at room temperature for 1 hour. After three washes with PBS-T, tetramethylbenzidine substrate was added, and the plates were further incubated for 15 min before stopping the reaction with 2 N H₂SO₄. The optical density at 450 nm was measured using a GloMax Explorer microplate reader (Promega, Seoul, Republic of Korea).
Plaque reduction neutralization test
Sera of immunized mice, NHPs, and convalescent patients were serially diluted from 1:10 to 1:5120 with serum-free medium. Virus–serum mixtures were prepared by mixing 100 PFU of SARS-CoV-2 with the diluted serum samples, and the mixtures were incubated at 37°C for 1 hour. Vero cells were inoculated with the virus-serum mixtures, and the plates were incubated at 37°C in the presence of 5% CO₂ for 1 hour. After virus absorption, the agar overlay medium was added, and the plates were incubated at 37°C under 5% CO₂ for 3 days. The cells were stained with 0.1% crystal violet solution (Sigma-Aldrich, St. Louis, MO, USA), and the plaques were counted. The percentage of neutralization represented the reduction value, which was calculated as the number of plaques in 100 PFU of the virus-infected well per number of plaques in the virus-serum mixture–infected well (×100).

Viral quantification
The virus in nasopharyngeal and oropharyngeal swab samples and respiratory tissue, including bronchi and six lobes of the lungs, was quantified using reverse transcription quantitative polymerase chain reaction (RT-qPCR), and the TCID₅₀ was determined. Briefly, swab samples were centrifuged (1600g for 10 min), and the supernatants were filtered through 0.2-µm syringe filters. Lung and bronchus samples were homogenized using a Precellys Homogenizer (Bertin Instruments, Montigny-le-Bretonneux, France) in 10-fold (w/v) sterile PBS (pH 7.4). After centrifugation, the supernatants were directly inoculated into Vero cells and incubated for 3 days at 37°C to calculate the values of TCID₅₀/ml. Viral RNA was extracted from the supernatants of swab and tissue samples using a QIAamp Viral RNA Mini Kit (Qiagen, Germantown, MD, USA). RT-qPCR was conducted with a primer/probe set to detect the ORF1b gene as described previously (24). Viral RNA copy numbers from swab and tissue samples were calculated using the SARS-CoV-2 RNA standard sample, which was run in parallel with the RT-qPCR reaction.

Holotomography
Holotomography was performed to visualize and acquire images of protein complexes. A holotomography microscope (HT-2H, Tomocube Inc., Daejeon, Republic of Korea), which is a novel optical microscope that uses diffraction tomography to generate three-dimensional (3D) holographic images of label-free live cells, was used for this purpose. Holotomography provides a 3D refractive index distribution of the protein through which several 2D holographic images can be obtained from different angles to generate a 3D tomogram (25). The details of holotomography function and property can be found in the literature (26). The acquired data were processed and analyzed with Tomostudio software (Tomocube Inc., Daejeon, Republic of Korea). The protein samples, including RBD-P2, RBD-P2/N, RBD-P2 + alum (RBD-P2 in complex with alum), and RBD-P2/N + alum, were analyzed in this study (fig. S3). All samples were prepared at a final concentration of 30 ng/µl. To visualize the samples, a Tomicapping (Tomocube Inc., Daejeon, Republic of Korea) was used, and 25 µl of the sample was loaded into the 20 mm–by–20 mm well.

Dynamic light scattering measurements
RBD-P2 + alum complex, RBD-P2/N + alum complex, and alum alone were concentrated to a concentration of 0.1 mg/ml in 0.05 mM phosphate buffer [including 300 mM NaCl (pH 7.0)]. Dynamic light scattering (DLS) was measured on a ZetaSizer Nano ZS90 (Malvern Instrument, Worcs, UK) equipped with a 4-mW He-Ne laser. All the DLS measurements were performed at 4.0° ± 0.1°C and at a scattering angle of 90°. The apparent z-average hydrodynamic diameter (Dₐ) and polydispersity index (PDI; <D₄/D₉>²) were calculated using the Dispersion Technology Software provided by Malvern.

Histological analysis
Sectioned lung tissue from experimental animals (mG1 to mG3) was submerged in 10% (v/v) neutral-buffered formalin, dehydrated, paraffin-embedded, and sectioned at a 5-µm thickness for histological examination. The histopathologists were blinded to group distribution when this analysis was conducted, and representative histopathological images were obtained and evaluated using Aperio ImageScope version 12.3 (Leica Biosystems Pathology Imaging, Buffalo Grove, IL, USA). The severity of histopathological changes was determined using a five-point scoring system as follows: 0 = no abnormality detected, 1 = minimal, 2 = mild, 3 = moderate, 4 = moderately severe, and 5 = severe. Distribution was recorded as focal, multifocal, and diffused. Recruitment of inflammatory cells into the lungs (bronchi or blood vessels) and morphological alterations of the tissue were assessed after hematoxylin and eosin staining under light microscopy.

Statistical analysis
Data are expressed as medians ± 95% confidence interval. Statistical analysis was performed using GraphPad Prism. Analysis was performed using a one-way analysis of variance (ANOVA) with Dunnett’s post hoc for multiple group comparisons and unpaired t test to compare between two groups. The type of analysis and statistical interpretation of the results are indicated in the figure legends. Differences were considered statistically significant when P < 0.05.

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/7/22/eabg7156/DC1

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Acknowledgments: We gratefully acknowledge M.-d. Oh and S. W. Park, Department of Internal Medicine, Seoul National University Hospital; S.-W. Kim, Department of Internal Medicine, School of Medicine, Kyungpook National University; K.-H. Park, Department of Infectious Disease, Chonnam National University Hospital; and J.-W. Chung, Department of Internal Medicine, Division of Infectious Diseases, Chung-Ang University for providing blood samples from convalescent patients with COVID-19. Funding: This work was supported by grants from RIGHT and SK Bioscience (grant nos. M-2020-D0731-0002 and M-2020-D0732-0003), as well as by the Ministry of Science and ICT and the Korea Research Institute of Bioscience (PRM1752011), Bio-industrial Core Technology Development Project from the Ministry of Trade, Industry, and Energy (20013101) and Biotechnology (KIRIBB) Research Initiative Program (KGM4572013). Author contributions: S.-M.L., J.J.H., and J.-H.N. conceived and supervised the research and designed the experiments. T.K., Hak Kim, K.-W.S., J.-H.K., S.L., and E.K. performed gene cloning, expression, protein purification, and vaccine formulation. H.O., S.H.B., G.K., B.-S.K., E.-H.H., and J.-H.P. performed the vaccinations in NHPs and challenged NHPs with live SARS-CoV-2. E.Y.O., K.W.K., and Y.K. performed the PRNT assay. H.W.K. measured serum antibody levels against RBD and N. S.-H.H. and H.W.K. performed flow cytometry analysis and interpreted the data. Hun Kim, Y.W.P., E.-H.H. assisted in the adjustments of directions and interpretation of the mechanistic aspects of the results. Hyoeng-Jun, Y.-S.L., Hyo-Jung, Y.-J.B., J.-Y.K., S.-H.B., S.-I.P., M.G., B.M.W., H.J., K.T.N., K.-A.H., and J.K. contributed to data acquisition. S.-H.H. and J.-H.N. contributed to writing the manuscript. Competing interests: K.-W.S. is an inventor on a pending patent related to this work filed by SK Bioscience (no.10-2020-0052855, filed 29 April 2020). The authors declare that they have no other competing interests. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

Submitted 23 January 2021
Accepted 9 April 2021
Published 28 May 2021
10.1126/sciadv.abg7156

Citation: S.-H. Hong, H. Oh, Y. W. Park, H. W. Kwak, E. Y. Oh, H.-J. Park, K. W. Kang, G. Kim, B.-S. Koo, E.-H. Hwang, S. H. Baeck, H.-J. Park, Y.-S. Lee, Y.-J. Bang, J.-Y. Kim, S.-H. Bae, S. J. Lee, K.-W. Seo, H. Kim, T. Kwon, J.-H. Kim, S. Lee, E. Kim, Y. Kim, J.-H. Park, S.-I. Park, M. Gonçalves, B. M. Weon, J. Jeong, K. T. Nam, K.-A. Hwang, J. Kim, H. Kim, S.-M. Lee, J. J. Hong, J.-H. Nam, Immunization with RBD-P2 and N protects against SARS-CoV-2 in nonhuman primates. *Sci. Adv.* **7**, eabg7156 (2021).