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Robust SARS-CoV-2 infection in nasal turbinates after treatment with systemic neutralizing antibodies

Highlights

- Systemic HuNAb or vaccine fails full SARS-CoV-2 infection prevention in nasal turbinate
- Post-exposure HuNAb suppresses SARS-CoV-2 in lungs but poorly in nasal turbinate
- Live SARS-CoV-2 persists in nasal turbinate for several days despite systemic HuNAb
- Robust SARS-CoV-2 infection in nasal turbinate is a mode to evade systemic HuNAb

In brief

SARS-CoV-2 infection in nasal turbinate determines viral transmissibility. Zhou and colleagues interrogate site-specific prevention of SARS-CoV-2 in hamsters. Prophylactic intraperitoneal or intranasal HuNAb or intramuscular DNA vaccination prevents infection effectively in lungs but not in nasal turbinate. Rapid postchallenge HuNAb suppresses infection significantly in lungs but poorly in nasal turbinate.
Robust SARS-CoV-2 infection in nasal turbinates after treatment with systemic neutralizing antibodies

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SUMMARY

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is characterized by a burst in the upper respiratory portal for high transmissibility. To determine human neutralizing antibodies (HuNAbs) for entry protection, we tested three potent HuNAbs (IC50 range, 0.0007–0.35 μg/mL) against live SARS-CoV-2 infection in the golden Syrian hamster model. These HuNAbs inhibit SARS-CoV-2 infection by competing with human angiotensin converting enzyme-2 for binding to the viral receptor binding domain (RBD). Prophylactic intra-peritoneal or intranasal injection of individual HuNAb or DNA vaccination significantly reduces infection in the lungs but not in the nasal turbinates of hamsters intranasally challenged with SARS-CoV-2. Although post-challenge HuNAb therapy suppresses viral loads and lung damage, robust infection is observed in nasal turbinates treated within 1–3 days. Our findings demonstrate that systemic HuNAb suppresses SARS-CoV-2 replication and injury in lungs; however, robust viral infection in nasal turbinates may outcompete the antibody with significant implications to subprotection, reinfection, and vaccine.

INTRODUCTION

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has resulted in about 80 million infections globally with nearly 1.7 million deaths by the end of 2020 since the discovery of the disease outbreak in December 2019 (Chan et al., 2020d; Zhu et al., 2020). The growing coronavirus disease 2019 (COVID-19) pandemic urgently requires the development of effective prophylaxis and treatment. While triple combination therapy (interferon-β1b, lopinavir/ritonavir, and ribavirin), remdesivir, and dexamethasone have all shown some clinical benefits in select patient groups (Bouware et al., 2020; Goldman et al., 2020; Hung et al., 2020b), the discovery of specific anti-SARS-CoV-2 agents with higher efficacy, better safety profiles, and bioavailability remain essential for improving the clinical outcome of COVID-19 patients. In addition to some drugs identified in large-scale drug repurposing programs (Riva et al., 2020), direct cloning of human neutralizing antibodies (HuNAbs) against SARS-CoV-2 has also been reported recently (Cao et al., 2020; Liu et al., 2020b; Robbiani et al., 2020; Shi et al., 2020; Sun et al., 2020; Wu et al., 2020a, 2020b; Zost et al., 2020). Furthermore, with the recent clinical trials of HuNAbs and urgent approval of COVID-19 vaccines for human use, it is necessary to determine whether or not HuNAbs may warrant sterile protection against live SARS-CoV-2 infection.

Unlike SARS patients who had peak upper respiratory tract (URT) viral loads at day 10 after symptom onset (Peiris et al., 2003), COVID-19 patients exhibited peak salivary or URT viral loads during the first week after symptom onset that declined over time; this phenomenon could account for the fast-spreading nature of the pandemic (Hung et al.,...
Regarding the humoral response, SARS-CoV-2-specific IgG and neutralizing antibody responses were quickly detectable in adult and children patients only 6 days after symptom onset (Liu et al., 2020c; Suthar et al., 2020; Zhou et al., 2020b). However, COVID-19 patients with higher amounts of anti-spike (S) and anti-nucleocapsid (NP) IgM and IgG tend to have poorer disease outcomes (Jiang et al., 2020; Tan et al., 2020). We and others also reported that COVID-19 patients with severe disease developed significantly more robust SARS-CoV-2-specific NAb responses (Liu et al., 2020a; Wang et al., 2020a, 2020b). Nevertheless, convalescent plasma with high NAb titers from recovered patients has been reported to be beneficial in the treatment of severe COVID-19 in small case cohorts (Duan et al., 2020). To replace convalescent plasma, which is not readily available in most countries, HuNAbS have been recently identified and showed promising results in preclinical studies (Cao et al., 2020; Liu et al., 2020b; Robbiani et al., 2020; Shi et al., 2020; Sun et al., 2020; Wu et al., 2020a, 2020b; Zost et al., 2020). However, the in vivo efficacy of anti-SARS-CoV-2 HuNAbS in protecting against URT infection in a physiologically relevant animal model has not been thoroughly investigated.

In this study, we identified a panel of candidate HuNAbS and conducted a thorough investigation of the lead candidate HuNAb ZDY20 at a dose of 10 mg/kg, which is much higher than its IC_{50} and IC_{90} values of 0.35 and 1 μg/mL, respectively, against live SARS-CoV-2 infection in both prophylactic and therapeutic settings in our established golden Syrian hamster model for COVID-19 (Chan et al., 2020a). We also tested the prophylactic efficacy of two much stronger RBD-specific HuNAbS 2–15 (IC_{50} and IC_{90} values of 0.0007 and 0.04 μg/mL) and ZB8 (IC_{50} and IC_{90} values of 0.013 and 0.031 μg/mL) and an S-based DNA vaccine to address the possible sterile protection in the same animal model.

RESULTS

Generation of a phage library displaying human antibody Fab of COVID-19 patients

To clone SARS-CoV-2-specific HuNAbS, we obtained peripheral blood mononuclear cells (PBMCs) from 12 convalescent COVID-19 patients in Hong Kong at a mean duration of 19 (±10.4) days after symptom onset (Table S1). These patients included seven females and five males with a mean age of 59 years (range, 21–75). Two patients had severe disease, seven had mild disease, and three were asymptomatic. The PBMCs of these 12 patients were pooled for the generation of a Fab phage library, as only small amounts of PBMCs were obtained from each patient. ELISA and pseudovirus neutralization assays were performed to measure the antibody titers in each patient’s serum prior to pooling, which confirmed that each study subject had SARS-CoV-2 RBD-specific neutralizing antibody activities (Figure 1B). Consistent with the observations that we and others reported previously, the two severe patients had higher NAb titers than the mild and asymptomatic patients (Liu et al., 2020a; Riva et al., 2020; Wang et al., 2020a). The mean NAb IC_{50} titer was 1:1753 with a range of 1:638–1:5701 (Figure 1B). Using the pooled PBMCs, we first set up a Fab phage library consisting of 3\times 10^6 clones (Figure S1). We subsequently developed an in-solution selection method for two rounds of panning (Figure 1C). We were able to pick up 384 single colonies (Figure 1D, color-coded) and name according to the clone numbers.
Specificity and neutralizing activities of cloned HuNAbs against SARS-CoV-2

Sequence analysis revealed that ZDY28, ZDY49, and ZDY95 share high similarity with the same immunoglobulin heavy chain variable region gene 3-30 (IGHV3-30) and immunoglobulin kappa chain variable region gene 1-39 (IGKV1-39) (Table S2). They share identical complementarity-determining region 3 (CDR3) heavy (H) and CDR3 light (L) chains but with a few amino acid differences in framework region 1 (FR1) (Figure S2). Since repeated sequencing gave identical results and the IGHV/IGKV fragments were cloned from the phage library derived from 12 patients, it is possible that the small amino acid differences between ZDY49 and ZDY95 might be derived from different donors. In contrast, ZDY20 has IGHV3-53 and IGKV1-33, which were recently shown to be associated with higher antiviral activity (Dalamaga et al., 2020; Yuan et al., 2020). ZDY20 has a longer CDR3H with three additional amino acids compared with the other three antibodies (Figure S2). These four antibodies all had low rates of somatic hypermutation (SHM) in both IGHV (0.35% or 0.69%) and IGKV (0.36% or 1.08%), except that the SHM rate in IGKV of ZDY20 reached 7.89% (Table S2). We next analyzed the binding activity of these four antibodies to recombinant SARS-CoV-2 RBD and S proteins by ELISA. In these solid phase ELISAs, ZDY28, ZDY49, and ZDY95 showed similar binding abilities to RBD (EC50 0.012 μg/mL) (Figure 2A) and to S (EC50 0.1 μg/mL) (Figure 2B), which were lower than those of ZDY20 (EC50 0.135 and 0.52 μg/mL, respectively) (Table S3). However, the anti-SARS-CoV-2 neutralization activity of ZDY20 (IC90 1.24 μg/mL) was better than those of ZDY28 (IC90 7.47 μg/mL), ZDY49 (IC90 17.08 μg/mL), and ZDY95 (IC90 13.35 μg/mL) in the pseudovirus assay (Figure 2C; Table S3). Moreover, ZDY20 exhibited, relatively, a better IC90 (1 μg/mL) than those of ZDY28 (1.28 μg/mL), ZDY49 (1.24 μg/mL), and...
ZDY95 (1.5 μg/mL) against live SARS-CoV-2 in Vero-E6 cells (Figure 2D; Table S3). Notably, none of these HuNAb showed cross-neutralization against the SARS-CoV pseudovirus (data not shown). Using a random yeast surface display of SARS-CoV-2 S fragments, we observed that all four HuNAb recognized conformational determinants instead of a small linear epitope because their binding domain on viral RBD required a minimum of 179 amino acid residues (337–524 for ZDY20, 336–523 for ZDY28, and 336–515 for ZDY49 and ZDY95) (Figure S3). Next, we measured the binding affinity of each antibody to the SARS-CoV-2 RBD or S trimer using surface plasmon resonance (SPR). Similar to the weaker binding activity observed by ELISA, ZDY20 Fab showed weaker affinity binding to SARS-CoV-2 RBD with a KD value of 74.9 nM, which was larger than those of the other three antibodies (ZDY28 19.2 nM, ZDY49 42.3 nM, and ZDY95 26.2 nM) (Table S4). The complete ZDY20 antibody, however, displayed the highest avidity binding to SARS-CoV-2 S trimer with a dissociation constant (KD) value of 8.8 nM, which was lower than those of the other three antibodies (ZDY28 10.2 nM, ZDY49 24.4 nM, and ZDY95 15 nM) (Figure 2E; Table S4). Furthermore, in the human cellular receptor angiotensin converting enzyme-2 (ACE2) competition assay by SPR, ZDY20 presented the most potent activity to block S binding to ACE2 followed by ZDY95, ZDY28, and ZDY49 (Figure 2F), although ZDY20 was not the strongest competitor to block soluble RBD binding to ACE2 (Figure 2G). These results suggested that the relative higher anti-SARS-CoV-2 neutralizing potency of ZDY20 is likely due to its stronger competitive blockade of S trimer binding to ACE2, which is probably associated with avidity interaction. Based on its relatively better activities of avidity binding KD to S, blocking S binding to ACE2, and neutralization, we chose ZDY20 for subsequent in vivo experiments.

SARS-CoV-2 robustly infects nasal turbinate despite systemic ZDY20 prophylaxis as compared with two stronger HuNAb and DNA vaccination

To determine the potential role of HuNAb as prophylaxis for SARS-CoV-2 infection, we first administered ZDY20 intraperitoneally to golden Syrian hamsters before virus challenge in our biosafety level-3 (BSL-3) animal laboratory. Syrian hamsters typically recover from SARS-CoV-2 infection with resolution of clinical signs and clearance of virus shedding within 1 week after infection, as we previously described (Chan et al., 2020a). Accordingly, the hamsters were sacrificed for analysis at 4 days post-infection (dpi) when viral loads and acute lung injury were consistently observed. In this prophylaxis study, each hamster received a single intraperitoneal injection of either 10 mg/kg (n = 4) high dose or 5 mg/kg (n = 3) low dose of ZDY20 (Figure 3A) that were much higher than its IC90 value of 1 μg/mL tested by live SARS-CoV-2 in Vero-E6 cells, respectively. Another group of hamsters (n = 4) was injected with a control HIV-1-specific HuNab VRC01 at a high dose of 10 mg/kg (Liu et al., 2020b). Twenty-four hours after the ZDY20 injection, each animal was challenged intranasally with 106 plaque-forming units (PFUs) of live SARS-CoV-2 (HKU-001a strain) (Chan et al., 2020a; Liu et al., 2020b). At 4 dpi, nasal turbinate, trachea, and lung tissues were harvested to quantify infectious viruses by plaque assay, viral loads by real-time PCR and infected cells by immunofluorescence (IF) staining. We found that infectious PFUs were readily detected in all tissue compartments of VRC01-controlled hamsters tested but not in 75% and 0% nasal turbinates, 75% and 67% trachea, and 50% and 33% lungs of hamsters that were given high and low doses of ZDY20, respectively (Figure 3B). The more sensitive RT-PCR further demonstrated that viral RNA copy numbers were reduced in the lungs by an average of 3 logs (range, 0.7–4.5) (Figure 3C). In contrast, there was no significant viral load reduction in nasal turbinates and trachea in both dose groups, suggesting less preventive efficacy by ZDY20 in the URT of viral entry. To determine the role of ZDY20 in protection, we measured the serum antibody concentration and neutralization titer at 0 and 4 dpi. Over 40 and 9 μg/mL ZDY20 were found in 50% high-dose and 67% low-dose animals at 0 dpi (Figure 3D, solid symbol; Table S5), together with mean IC50 values of approximately 1:236 and 1:99, respectively (Figure 3E, solid symbol). At 4 dpi, over 26 and 5 μg/mL ZDY20 were found in 50% high-dose and 67% low-dose animals (Figure 3D, open symbol; Table S5), together with mean IC50 values of approximately 1:55 and 1:45, respectively (Figure 3E, open symbol). These results demonstrated that most animals maintained peripheral amounts of ZDY20 much higher than its IC50 values during the entire course of the experiment.

Next, to understand whether or not ZDY20 protects against infection-induced lung injury in hamsters, we performed pathological analysis on lung specimens. Compared with uninfected healthy animals (Figure S4), hamsters treated with the high-dose ZDY20 showed mild interstitial alveolar inflammation with minor septal infiltration and congestion (Figure 3F, top left, arrows). There was no apparent peribronchial inflammation, and the bronchiolar epithelia appeared normal (open arrows). Hamsters treated with the low dose showed limited areas of bronchiolar epithelial cell swelling and detachment (Figure 3F, middle left, open arrows) together with mild peribronchial inflammation and mild alveolar septal infiltration. In contrast, control hamsters showed large patchy areas of alveolar wall and alveolar space involvement by inflammatory infiltrates and exudation (Figure 3F, bottom left, arrows). The two blood vessels (BVs) in the lung section showed vasculitis and endothelitis. Furthermore, we performed IF staining of viral NP antigen in both lung and nasal turbinate tissues compared with uninfected healthy animals (Figure S4). In the high-dose group, little viral NP expression was observed in lung sections (Figure 3F, top right). In the low-dose group, a small amount of viral NP was observed in localized areas of alveoli (Figure 3F, middle right, arrows) and in the epithelia of a few bronchiolar sections (thin arrows). In contrast, control hamsters showed diffuse NP expression in extensive areas of alveoli (Figure 3F, bottom right, thick arrows) and in the bronchiolar epithelia (thin arrows). The mean number of NP+ cells per 50× field in the lungs was significantly lower in the high-dose group than that in the control animals (Figure 3G). Remarkably, nasal turbinate tissues showed damage to the respiratory and olfactory epithelium with extensive submucosal immune cell infiltration, together with robust and diffuse viral NP expression in extensive areas lining the stratified squamous epithelia in both the high-dose and low-dose groups of ZDY20-treated animals, similar to untreated control hamsters (Figures 4A–4C). Moreover, no significant difference in NP+ cells per 50× field was found between treated and control animals when nasal turbinate tissues were analyzed (Figure 4D). Collectively, these results demonstrated that prior administration of 10 mg/kg ZDY20 did not prevent robust SARS-CoV-2 infection.
in nasal turbinates but suppressed productive infection in lungs in the Syrian hamster model.

Considering that HuNAb with greater inhibitory potency may be more effective in preventing viral replication in the nasal turbinates, we tested two such RBD-specific HuNAb, namely ZB8 and 2-15. ZB8 is a newly cloned HuNAb with IC50 and IC90 values of 0.013 and 0.031 mg/mL, whereas 2-15 has been previously reported with IC50 and IC90 values of 0.0007 and 0.04 mg/mL, all in the live virus neutralization assay (Table S6) (Liu et al., 2020b). ZB8 and 2-15 are therefore up to 32-fold (IC90) to 500-fold (IC50) stronger than ZDY20. Prophylactic intraperitoneal injection of 4.5 mg/kg ZB8 and 1.5 mg/kg 2-15 were tested in hamsters using the same protocol as ZDY20. We consistently observed significantly reduced viral loads and NP+ cells in the lungs mainly by ZB8 but still not in nasal turbinates of experimental hamsters at 4 dpi in both experiments. Abundant NP+ cells were readily found in nasal turbinate of each treated animal. In addition, we examined the efficacy of a systemic DNA vaccine that was

Figure 3. Efficacy of ZDY20 prophylaxis against live SARS-CoV-2 in Syrian hamsters
(A) Experimental schedule and color coding for three experimental groups. Two groups of hamsters received a single intraperitoneal injection of ZDY20 at doses of 10 mg/kg (n = 4) and 5 mg/kg (n = 3) one day before the live virus challenge (−1 dpi). Another group was given the control antibody VRC01 at 10 mg/kg (n = 4). On day 0, each hamster was intranasally inoculated with a challenge dose of 100 μL of Dulbecco’s modified Eagle medium containing 10^4 plaque-forming units of SARS-CoV-2 (HKU-001a strain, GenBank accession no: MT230904.1). The hamsters were sacrificed at 4 dpi for analysis.
(B) Infectious virions were tested by viral plaque assay in nasal turbinate, trachea, and lung tissue homogenates. PFUs per mg of tissue extractions were compared between different groups in log_{10}-transformed units.
(C) The relative viral concentration (normalized by β-actin) was determined in nasal turbinate, trachea, and lung tissue homogenates by the sensitive RT-PCR assay.
(D and E) Serum concentration (D) and neutralizing activity (E) of HuNAb ZDY20 were determined by ELISA and pseudovirus assays, respectively. Open symbols represent corresponding samples collected at 4 dpi in the same color-coding groups.
(F) Representative images of hamster lung tissues by H&E (100 x, left panel) and immunofluorescence (IF) staining (50 x, right panel). For viral NP antigen in the sections (50 x) stained by IF, the lungs of control hamsters showed diffuse NP expression in large areas of alveoli (thick arrows) and in bronchiolar epithelium (thin arrows). SARS-CoV-2 N protein (NP) was stained green, and cell nuclei were counterstained with DAPI (blue). No NP expression was observed in lung sections of hamsters treated with high-dose antibody (10 mg/kg). In low-dose antibody-treated hamsters, NP expression was occasionally observed in a small area of alveoli (thick arrow) and in the epithelium of a few bronchiolar sections (thin arrows).
(G) NP+ cells per 50 x field were compared in the lungs of three experimental groups. Data of individual animals were indicated by different symbols, and different colors represented different groups. Statistics were generated using ordinary one-way ANOVA and multiple comparisons test. *p < 0.05; **p < 0.01; ***p < 0.001. See also Figures S4 and S9 and Table S5.
codon-optimized and encoded the full-S protein as we previously described for SARS (Yi et al., 2005). The S protein was detected in transfected cells by western blot (Figures S8A and S8B) and presented with correct conformation by supporting pseudovirus infection (Liu et al., 2020a). Four injections of DNA vaccination induced both RBD binding and neutralizing antibodies in hamsters (Figures S8C–S8E). The mean NAb IC50 titer at 1 day before viral challenge sustained at 1:4419 with a range of 1:286–14175 comparable to most convalescent human subjects (Figure 1B) and vaccinees (Sadoff et al., 2021; Sahin et al., 2020; Xia et al., 2020). After the intranasal challenge, significantly reduced viral loads and NP+ cells were observed in the lungs but not in nasal turbinates of hamsters at 4 dpi (Figures 5Da and S8F–S8H). Our results therefore demonstrated that passive immunization and intramuscular DNA vaccination tested are unlikely effective for mucosal sterile protection due to robust viral infection in nasal turbinates outcompeting the systemic neutralizing antibodies.

**Postchallenge ZDY20 therapy reduces SARS-CoV-2 infection in the lungs but considerably less in nasal turbinate**

To explore the therapeutic potential of ZDY20, we infected four groups of hamsters (n = 4 per group) as described above. Three groups of hamsters were then treated intraperitoneally with a single high dose of 10 mg/kg at 1, 2, and 3 dpi (Figure 6A, color-coded). The fourth group was the VRC01 control using the same dose at 1 dpi. All animals were consistently sacrificed at 4 dpi for measurements of infectious virus titers and viral RNA copy numbers in lung, trachea, and nasal turbinate tissues (Figures 6B and 6C). At 1 dpi, ZDY20-treated hamsters showed no measurable infectious viruses in 100% nasal turbinates, 100% trachea, and 75% lungs by the plaque assay (Figure 6B, blue). The mean viral load reduction was 0.7 logs in both nasal turbinates (range, 0.1–1.0 logs) (p < 0.05) and trachea (range, 0.3–1.5 logs) but was 1.2 logs (range, 0.8–1.8 logs) in lungs (p < 0.01) compared with the control group (Figure 6C, blue). At 2 dpi, ZDY20-treated hamsters showed no measurable infectious viruses in 75% nasal turbinates, 75% trachea, and 50% lungs (Figure 6B, red). The mean viral load reduction was 1.0 logs (range, 0.3–1.5 logs) in the trachea (p < 0.05) but not significant in the nasal turbinate and lungs compared with the control group (Figure 6C, red). At 3 dpi, ZDY20-treated hamsters had no measurable infectious viruses in 25% nasal turbinates, 75% trachea, and 25% lungs (Figure 6B, green). The mean viral load reduction was not significant in the nasal turbinates, trachea, or lungs compared with the control group (Figure 6C, green).
These kinetics results demonstrated that earlier commencement of treatment within 2 dpi resulted in a more significant reduction of infectious viruses. For more significant viral load reduction, commencement of treatment within 1 dpi is likely necessary. To determine the role of ZDY20 in protection, we measured the antibody concentration and neutralization titer at 4 dpi. Over 40 μg/mL ZDY20 was found in 100% treated animals at 1 dpi, 75% treated animals at 2 dpi, and 50% treated animals at 3 dpi (Figure 6D; Table S5), together with mean IC_{50} values of 1:169, 1:62, and 1:69 in each corresponding group (Figure 6E), respectively.

Subsequently, we performed pathological analysis on lung specimens to understand whether ZDY20 protects against tissue injury in hamsters. The H&E-stained sections of control hamsters showed acute lung injury with peribronchiolar infiltration, bronchiolar epithelial cell death, diffuse alveolar wall thickening, patchy areas of alveolar space infiltration, exudation, and vasculitis (Figure 6F, top). In contrast, less severe histopathological changes were observed in all 3 groups of ZDY20-treated hamsters. There was no apparent peribronchiolar infiltration in hamsters treated at 1 dpi. The bronchiolar epithelia appeared normal, with only occasionally observed cell death in hamsters treated at 2 dpi. Moderate alveolar wall thickening and capillary congestion without alveolar space infiltration or exudation and limited vasculitis were observed in hamsters treated at 3 dpi. Corroboratively, abundant SARS-CoV-2 NP expression in the bronchiolar epithelium and alveoli was observed in the lungs of control hamsters (Figure 6F, bottom). In ZDY20-treated hamsters, NP expression was confined to small areas of alveoli at 1 dpi and was more frequently found in the bronchiolar epithelia at 2 and 3 dpi. The mean number of NP+ cells per 50 x 50 field of lung tissues at 1 dpi was significantly lower than that of the control group (Figure 6G). Importantly, when we performed IF staining of viral NP antigen in nasal turbinate tissues, robust and diffuse viral NP expression in extensive areas lining the stratified squamous epithelia was found in infected hamsters treated with 10 mg/kg ZDY20 either at 1, 2, or 3 dpi, similar to untreated control hamsters (Figure 7A). These results demonstrated that postchallenge treatment with 10 mg/kg ZDY20 suppressed robust SARS-CoV-2 infection in the lungs but not in the nasal turbinates in...
the hamster model (Figure 7B), suggesting the deficiency of systemic HuNAb to outcompete the viral replication within the nasal mucosal epithelia.

To further understand the difficulty of sterile protection in the URT, we sought to measure the amounts of intraperitoneally injected HuNAb in nasal wash and tissue homogenates. In contrast to high amounts of ZDY20, ZB8, and 2-15 detected in animal sera (Tables S5 and S6), no measurable amounts of antibodies were found in nasal wash specimens of hamsters tested. Various amounts of antibodies, however, were found in nasal turbinate and lung homogenates (Figure S9). When we included all hamsters from both ZDY20 prevention and treatment experiments together with 2-15 prevention (Figure S6) and intranasal prevention of ZDY20 and ZB8 (Figure S7) for analysis, the amount of peripheral HuNAb was negatively correlated with NP+ cells in the lungs but not in the nasal turbinates (Figure 7C). Furthermore, there was a significant negative correlation between the tissue HuNAb concentration and the number of infectious viral PFUs either in nasal turbinate or in lung homogenates (except that nasal turbinates of five animals in the 2-15 prevention experiment were not available for measuring PFUs) (Figure 7D). Therefore, HuNAb in tissue homogenates might account for the significant reduction of infectious viral particles in both nasal turbinates and lungs once the antibody encounters the virus for neutralization during the PFU measurement experiment.

DISCUSSION

ZDY20, ZB8, and 2-15 represent the most promising class of RBD-specific HuNAb that bind to the conformational
determinants of the SARS-CoV-2 RBD overlapping with the cellular entry receptor ACE2 binding site (Cao et al., 2020; Liu et al., 2020b; Robbiani et al., 2020; Shi et al., 2020; Sun et al., 2020; Wu et al., 2020a, 2020b; Zost et al., 2020). Both SARS-CoV-2 and SARS-CoV use ACE2 to initiate infection, although they share only 40% amino acid identity in the RBD external subdomain (Chan et al., 2020b; Sahin et al., 2020; Wan et al., 2020; Zhou et al., 2020a). Structural analysis indicated that residues in the SARS-CoV-2 RBD that are essential for ACE2 binding are highly conserved or share similar side chain properties with those in the SARS-CoV RBD (Lan et al., 2020). During the natural course of infection, there was cross-reactivity between anti-bodies to SARS-CoV-2 and SARS-CoV, but only antibodies from COVID-19 patients neutralized SARS-CoV-2 (Chen et al., 2005; Yi et al., 2005). In this study, we consistently found that RBD-specific ZDY20 displayed neutralizing activity through competition for the ACE2 binding site. Our in vivo analysis of ZDY20, together with data of the two stronger RBD-specific ZB8 and 2-15, should have important implications for other HuNAbS that have similar functional properties and are now under clinical development (Cao et al., 2020; Liu et al., 2020b; Robbiani et al., 2020; Shi et al., 2020; Sun et al., 2020; Wu et al., 2020a, 2020b; Zost et al., 2020).

Protection of the URT through passive immunization of HuNAb or intramuscular vaccination is a very important issue of SARS-CoV-2 prevention and vaccine development (Krammer, 2020). Ciliated nasal epithelial cells in the URT have the highest ACE2 and transmembrane serine protease 2 (TMPRSS2) expression to support SARS-CoV-2 infection (Hou et al., 2020), which is consistent to our findings in SARS-CoV-2-infected hamsters (Chan et al., 2020a). Several potent RBD-specific HuNAbS have been evaluated for SARS-CoV-2 prevention in various

![Figure 7. Postchallenge ZDY20 therapy does not suppress SARS-CoV-2 replication in nasal turbinates considerably by IF staining and correlation analysis](image-url)
animal models (Liu et al., 2020b; Rogers et al., 2020; Shi et al., 2020; Zost et al., 2020). Most of these studies, however, have focused on viral load reduction in the lungs without detailed examination of viral replication in the URT tissues. In this study, we showed that systemic prophylaxis by ZDY20, ZB8, and 2-15 did not prevent SARS-CoV-2 infection significantly in nasal turbinates, although serum neutralization IC₅₀ values were maintained over 1:200 at the time of virus challenge and over 1:50 at the time of animal sacrifice. We only observed significantly reduced numbers of infectious viruses (PFUs) in homogenized nasal turbinates. We reasoned that the reduction was likely due to blood-derived potent HuNAb during tissue homogenization for the in vitro assay. Indeed, low amounts of ZDY20 and ZB8 were detected in homogenized nasal turbinates tissues. Moreover, the amount of HuNAb was negatively correlated with the PFU number of infectious viruses in both lung and nasal turbinates homogenates, suggesting that SARS-CoV-2 could be neutralized when it encountered the sufficient amount of neutralizing antibody. Since the amounts of ZDY20 and ZB8 were very low in nasal wash, our results suggested that the insufficient passive immunization for sterile protection is probably due to poor distribution of neutralizing antibodies on or within the viral entry portal of nasal turbinates to outcompete the robust viral infection there. This finding then brought the worry on vaccine-induced neutralizing antibodies for URT sterile protection. Based on experimental evidence, although the intramuscular DNA/electroporation vaccination induced high amounts of neutralizing antibodies, which were comparable to those of convalescent plasma or human vaccine recipients (Sadoff et al., 2021; Sahin et al., 2020; Xia et al., 2020), robust SARS-CoV-2 infection was still found in nasal turbinates tissues of experimentally infected hamsters. The subprotection may allow possibly HuNAb-resistant viruses to emerge. Collectively, these data highlight the need to explore vaccine-induced mucosal immunity (e.g., high amount of sIgA in URT) for the complete prevention of SARS-CoV-2 infection. Since COVID-19 patients have the highest viral load and viral shedding into saliva at or soon after presentation (To et al., 2020b), effective inhibition of SARS-CoV-2 replication in the URT would be critical for infection control. Mucosal protection faces a serious challenge because one-third of convalescent COVID-19 patients had serum IC₅₀ values less than 1:50 (Robbiani et al., 2020), and therefore, they may be susceptible to reinfection. Indeed, we have recently documented the world’s first reported case of SARS-CoV-2 reinfection in a Hong Kong patient whose serum NAb titer decreased from 1:40 to <1:10 within 140 days after an acute asymptomatic infection had virus shedding in the URT due to reinfection by another strain of SARS-CoV-2 (To et al., 2020a). Before we have an effective vaccine for sterile protection, our results strongly urge universal masking as the highest priority for SARS-CoV-2 prevention.

In our postchallenge treatment kinetics study, we showed that early use of ZDY20, preferably within 2 dpi, suppressed infectious virus, viral loads, and acute injury in the lungs of treated hamsters. In nasal turbinates, however, robust viral loads and NP⁺ cells were readily observed for treatment even at 1 dpi, except for the reduced numbers of infectious viruses in homogenized tissues. Since the amount of peripheral ZDY20 was negatively correlated with NP⁺ cells in the lungs but not in the nasal turbinates, the therapeutic efficacy of injected ZDY20 is likely limited in the URT. Several studies have evaluated the potency of RBD-specific HuNAbs for treating preinfected mice, hamsters, and monkeys (Baum et al., 2020; Cao et al., 2020; Shi et al., 2020; Wu et al., 2020b), but few have performed a treatment kinetics study. In hACE2-transgenic mice, a 20-mg/kg injection of BD-368-2 at 2 h post-infection (hpi) reduced the lung viral loads by approximately 3 logs (Cao et al., 2020). In another study, after a 25-mg/kg dose of a noncompeting pair of HuNAbs given at 12 hpi, viral loads were reduced for over 2 logs, yet there were still 10⁶ copies/g of lung tissues at 3 days after virus challenge in hACE2-transgenic mice (Wu et al., 2020b). In monkey models, a preprint article indicated that SARS-CoV-2 viral loads were detected in nasopharyngeal swabs at 4 dpi in animals that received a very high dose of 150 mg/kg of the REGN-COV2 antibody cocktail at 1 dpi (Baum et al., 2020). Moreover, an intraperitoneal dose of 50 mg/kg treatment with the RBD-specific CB6 antibody that has an IC₉₀ value of 1 μg/mL reduced but did not completely prevent histopathological changes in the lungs of SARS-CoV-2-infected monkeys (Shi et al., 2020). In addition to these HuNAb studies, for convalescent plasma therapy, we previously reported that convalescent sera of infected hamsters reduced SARS-CoV-2 replication but not acute lung injury (Chan et al., 2020a). Since the mortality rate of critically ill COVID-19 patients might be related to the IgG antibody levels in the transfused convalescent plasma (Joyner et al., 2020), it might be necessary to develop high-titer HuNAbs as specific antivirals for COVID-19 early treatment. In this study, by initiating ZDY20 treatment at the different time points of 1, 2, and 3 dpi, we demonstrated the benefits of commencing HuNAb treatment as early as possible in terms of better viral load reduction and alleviation of acute lung injury. Nevertheless, treatment started as late as 3 dpi still resulted in better histopathological changes in the lungs and therefore should still be considered in COVID-19 patients who presented later in the course of disease, as long as HuNAb did not promote acute lung injury (Liu et al., 2019).

In summary, we describe robust SARS-CoV-2 infection in nasal turbinates in the presence of potent systemic neutralizing antibodies in both prophylaxis and therapeutic studies in golden Syrian hamsters. Despite the presence of systemic serum HuNAb IgG, high viral loads and infected cells, as well as infectious viruses, may persist for at least 4 days in nasal turbinates tissues. Nevertheless, systemic ZDY20 is able to suppress SARS-CoV-2 replication and acute injury in the lungs of experimental hamsters. These findings have important implications for viral transmission, patient management, vaccine development, and control of reinfection.

Limitations of study
One limitation of study is that we could not determine the dose that RBD-specific HuNAb would be sufficient for sterile protection of the URT. Giving detectable viral loads in nasopharyngeal swabs at 4 dpi in monkeys that received a very high dose of 150 mg/kg of the REGN-COV2 antibody cocktail at 1 dpi (Baum et al., 2020), protection of the URT by IgG through systemic HuNAb or intramuscular vaccination can be quite difficult.
During our manuscript revision, Chen et al. published an RBD-specific HuNAb LY-CoV555 clinical study, examining three doses of 700, 2,800, and 7000-mg monotherapy (Chen et al., 2021). In this phase 2 trial, only the middle dose of LY-CoV555 appeared to accelerate the natural decline of viral loads in nasopharyngeal swaps over time, whereas the other doses had not by day 11. Systemic HuNAb is therefore unlikely sufficient to reach URT for immediate and complete viral suppression in humans, which is consistent with our findings in hamsters. For implications to vaccine development, three COVID-19 vaccines including the Pfizer-BioNTech BNT162B2 mRNA, the Moderna mRNA, and the Oxford-ChAdOx1 have released phase III trial results to support emergency use. Only the Oxford-ChAdOx1 vaccine has evaluated and indicated the low efficacy against asymptomatic infection (Voysey et al., 2021). In particular, the 3.8% efficacy among human vaccinees who received two standard doses was disappointing. Using the hamster model, our finding of robust viral replication in nasal turbinate outcompeting the vaccine-induced neutralizing antibodies has likely revealed a possible mechanism underlying the difficulty of the vaccine against asymptomatic infection. Another limitation of study is that we could not determine if viruses newly released from nasal turbinate tissues would be immediately neutralized by HuNAb because these viruses are difficult to separate from the inoculated viruses. The significantly reduced numbers of infectious viruses in homogenized nasal turbinate tissues measured by our plaque assay, however, provided evidence that viruses in the URT can be immediately neutralized when they encounter the HuNAb.

**STAR★METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.chom.2021.02.019.

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**AUTHOR CONTRIBUTIONS**

Z.C. and K.-Y.Y. supervised two collaborative teams in the study. D.Z., B.Z., J.F.-W.C., R.Z., and Z.C. designed the experiments, analyzed the data, and wrote the manuscript. D.Z. generated four ZDY HuNAbs. B.Z. generated ZB8. D.D.H. provided 2-15. J.F.I.W.C., R.Z., C.C.-S.C., V.K.-M.P., S.Y., H.X., K.K.-H.C., C.C.-Y.C., and J.C. conducted hamster, viral load, and live virus experiments. R.Z. and K.K.-W.T. prepared and studied clinical samples. K.-Y.K., Z.D., and T.T.-K.L. performed the immunofluorescence staining of the hamster tissue section. L.L. and A.J.Z. analyzed tissue pathology. S.L. performed yeast spike display and antibody epitope mapping. S.J.C., B.Z., and C.-Y.C. conducted antibody experiments. S.S., Q.Z., and LZ performed live virus neutralization.

**DECLARATION OF INTERESTS**

J.F.-W.C. has received travel grants from Pfizer Corporation Hong Kong and Astellas Pharma Hong Kong Corporation Limited and was an invited speaker for Gilead Sciences Hong Kong Limited and Luminex Corporation. The other authors declare no conflicts of interest except for a provisional patent application filed for human monoclonal antibodies generated in our laboratory by the University of Hong Kong. The funding sources had no role in study design, data collection, analysis, interpretation, or writing of the report.

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### Key Resources Table

#### Antibodies

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Anti-FLAG M2 Mouse Monoclonal Antibody | Sigma-Aldrich | Cat# M2001; RRID: AB_2046826 |
| Anti-HA Mouse Monoclonal Antibody | Sigma-Aldrich | Cat# 1100; RRID: AB_2046827 |
| Anti-HA Mouse Monoclonal Antibody | Thermo Fisher Scientific | Cat# 302720 |
| Anti-HA Mouse Monoclonal Antibody | Thermo Fisher Scientific | Cat# 301200 |

#### Bacterial and virus strains

| NAME OF STRAIN | SOURCE | IDENTIFIER |
|----------------|--------|------------|
| TG1 Electrocompetent Cells | Lucigen | Cat# 60502-1 |
| M13KO7 Helper Phage | New England BioLabs | Cat# N0315S |
| SARS-CoV-2 HKU-001a strain | GenBank accession no: MT230904.1 |

#### Biological samples

| NAME OF SAMPLE | SOURCE | IDENTIFIER |
|----------------|--------|------------|
| Blood sample from COVID-19 patients | Queen Mary Hospital | N/A |

#### Chemicals, peptides, and recombinant proteins

| NAME OF CHEMICAL | SOURCE | IDENTIFIER |
|------------------|--------|------------|
| Biotinylated SARS-CoV-2 spike protein RBD | ACROBiosystems | Cat# SPD-C82E9-200ug |
| ACE2 protein | ACROBiosystems | Cat# AC2-H52H8-1mg |
| SARS-CoV-2 spike protein | ACROBiosystems | Cat# SPN-C52H8-500ug |
| Recombinant Protein G Agarose | Thermo Fisher Scientific | Cat# 15920010 |
| Trimethylamine (TEA) | Sigma-Aldrich | Cat# T47183-500ML |
| HEPES buffer (1M) | GIBCO | Cat# 15630080 |
| Anti-M13 Major Coat Protein Antibody-HRP | Santa Cruz Biotechnology | Cat# sc-53004 HRP |
| Goat anti-Human IgG Cross-Adsorbed Secondary Antibody, HRP | Invitrogen | Cat# 62-8420 |
| 3, 3¢,5, 5¢-Tetramethylbenzidine Liquid Substrate | Sigma-Aldrich | Cat# T4444 |
| Antigen Unmasking Solution (Citrate-Based) | Vector Laboratories | Cat# H3300; RRID: AB_2336226 |
| Protease Lys-C | Roche | Cat# 11047825001 |
| Protein A Sepharose | GenScript | Cat# L00210-50 |

#### Critical commercial assays

| NAME OF SYSTEM | SOURCE | IDENTIFIER |
|----------------|--------|------------|
| RNeasy Mini Kit | QIAGEN | Cat# 74106 |
| PrimeScript™ 1st strand cDNA Synthesis Kit | TaKaRa | Cat# 6210B |
| NEBuilder® HiFi DNA Assembly Master Mix | New England BioLabs | Cat# E2621L |
| Luciferase Assay System | Promega | Cat# E1501 |

#### Experimental models: cell lines

| NAME OF STRAIN | SOURCE | IDENTIFIER |
|----------------|--------|------------|
| HEK293T-hACE2 cells | In house | N/A |
| Vero-E6 cells | ATCC | Cat# CRL-1586 |
| Expi293F™ Cells | Thermo Fisher Scientific | Cat# A14527; RRID: CVCL_D615 |

#### Experimental models: organisms/strains

| NAME OF ORGANISM | SOURCE | IDENTIFIER |
|------------------|--------|------------|
| Golden Syrian hamster | Chinese University of Hong Kong Laboratory Animal Service Center | N/A |

(Continued on next page)
**RESOURCE AVAILABILITY**

**Lead contact**
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Zhiwei Chen (zchenai@hku.hk).

**Materials availability**
This study did not generate new unique reagents.

**Data and code availability**
The study did not generate any unique datasets or codes.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Patients**
A total of 12 patients with COVID-19 were recruited between February and March 2020. All patient cases were confirmed by reverse-transcription polymerase chain reaction (RT-PCR) as described previously (Chan et al., 2020c). Clinical and laboratory findings were entered into a predesigned database. Written informed consent was obtained from all patients. This study was approved by the Institutional Review Board of the University of Hong Kong/Hospital Authority Hong Kong West Cluster, Hong Kong East Cluster Research Ethics Committee, and Kowloon West Cluster Research Ethics Committee (UW 13-265, HKECREC-2018-068, KW/EX-20-038[144-26]).

**Syrian hamsters**
Animal experimental plan was approved by the Committee on the Use of Live Animals in Teaching and Research (CULATR 5359-20) of the University of Hong Kong (HKU). Male and female Syrian hamsters (aged 6–10 weeks) were purchased from the Chinese University of Hong Kong Laboratory Animal Service Centre through the HKU Laboratory Animal Unit (LAU). The animals were kept in Biosafety Level-2 housing and given access to standard pellet feed and water ad libitum following LAU’s standard operational procedures (SOPs). The viral challenge experiments were then conducted in our Biosafety Level-3 animal facility following SOPs strictly.

**Cell lines**
HEK293T-hACE2 and Vero E6 cells were maintained in Dulbecco’s Modified Eagle Medium (DMEM) (Thermo Fisher Scientific) containing 10% Fetal bovine serum, 2 mM L-glutamine and 100 U/mL penicillin and were incubated at 37°C in 5% CO2 setting (Liu et al., 2019). Expi293F™ cells were maintained in Expi293™ Expression Medium (Thermo Fisher Scientific) and were incubated in a 37°C incubator with 80% relative humidity and 5% CO2 setting on an orbital shaker platform at 125 ± 5 rpm (New Brunswick innova™ 2100) according to the manufacturer’s instructions.

**METHOD DETAILS**

**Construction of a Fab Phage Display Library**
Total RNA was extracted from peripheral blood lymphocytes of convalescent SARS-CoV2 patients by RNeasy Mini kit (QIAGEN), and both total RNA and mRNA were reverse transcribed into cDNA using PrimeScript 1st strand cDNA Synthesis Kit (Takara). Fd segment
This panning procedure was adapted from previously described procedures (Chames, 2012). For the first round of panning, 100 μl streptavidin magnetic beads (Thermo Fisher Scientific) were coated with 200 nM biotinylated SARS-CoV-2 spike protein RBD (ACROBiosystems), which were then blocked with 2% skimmed milk PBS for an h. At the same time, 1 × 10^12 phages from the Fab phage library were also blocked in 2% skimmed milk PBS. Then, the mixture of phage and biotinylated RBD on beads was incubated for an h at room temperature (RT) on a rotating wheel. Then, the beads were pulled out of solution and washed 5 times with 1 mL of PBS Tween 20 (0.1%) and 2 times with 1 mL of PBS. The remaining phage was eluted with 0.1 M trimethylamine (TEA) and 0.2 M glycine pH 2 sequentially and then immediately neutralized with 1 M Tris base. The eluted phage was amplified by infection of Escherichia coli TG1 cultures, which were followed by superinfection with helper phage M13K07 (NEB). At Round 2, 100 nM antigen was used with 50 μl beads. Then, 25 washes were performed at Round 2 to narrow down the Round 1 phage output to enrich for those that bind specifically to the target. Similarly, at Round 3, 25 washes were performed to narrow down the diversity and enrich for only specific, stronger binders from the phage library. Polyclonal phage ELISA was carried out at Round 2, and if phages were not sufficiently enriched at this stage, then a third round was carried out. All panning procedures were performed in solution using low protein binding 1.5 mL Eppendorf tubes.

### Polyclonal, monoclonal phage and recombinant protein ELISAs

The procedures of phage ELISA were adapted from previously described procedures (Chames, 2012). For polyclonal phage ELISA, the RBD proteins were coated on 96-well enzyme-linked immunosorbent assay plates at 50 ng/well. After blocking with 4% skimmed milk in PBS, 50 μL phage library or amplified eluted phage was added to the plates at serial dilutions starting from 1:5 and then incubated for 1 h at room temperature. The plates were washed, and the anti-M13 Major Coat Protein Antibody-HRP (Santa Cruz Biotechnology) was added prior to color development by 3,3¢,5,5¢-Tetramethylbenzidine liquid substrate (Sigma-Aldrich). Optical density at 450 nm was read on a Varioskan LUX multimode microplate reader (Thermo Fisher Scientific). For monoclonal phage ELISA, a single colony was picked from the plate of eluted phage and cultured in 96-deep-well plates with approximately 200 μL/well at 37°C and 250 rpm. When the OD600 reached 0.5, each well of the single colony culture was superinfected with helper phage M13K07 (NEB) and then incubated at 30°C and 250 rpm for 16-18 h. The supernatant of the single culture was used for ELISA as the polyclonal phage ELISA with a 1:1 dilution with 4% skimmed milk in PBS. The strong single binders were sent for sequencing. For recombinant protein ELISA, the RBD or spike proteins were coated on 96-well enzyme-linked immunosorbent assay plates at 50 ng/well. After blocking with 4% skimmed milk in PBS, 50 μL of HuNAbs was added to the plates at 1:5 serial dilutions starting from 10 μg/mL and then incubated for 1 h at 37°C. The plates were washed, and the secondary antibody anti-human IgG-HRP (Invitrogen) was added. 3,3¢,5,5¢-Tetramethylbenzidine liquid substrate (Sigma-Aldrich) was used for color development. Optical density at 450 nm was read on a Varioskan LUX multimode microplate reader (Thermo Fisher Scientific).

### Protein expression, purification and Fab production

The antibody VH/VL and constant region genes were amplified and cloned into the expression vectors AbvecIgG and AbvecIgKappa using the NEBuilder® HiFi DNA Assembly Cloning Kit (NEB). The plasmids of paired IgH and IgL genes were cotransfected into the Expi293F expression system (Thermo Fisher Scientific) following the manufacturer’s protocol to produce recombinant HuNAbs. Antibodies from cell culture supernatants were purified immediately by affinity chromatography using Recombinant Protein G Agarose (Thermo Fisher Scientific) according to the manufacturer’s instructions. The purified HuNAbs were concentrated by an Amicon ultrafiltration device (molecular weight cutoff, 50 kDa; Millipore) to a volume of 0.2 mL in PBS and stored at −80°C. To produce Fab fragments, antibodies were cleaved using Protease Lys-C (Roche) with an IgG to Lys-C ratio of 4000:1 (w/w) in 10mM EDTA, 100mM Tris-HCl, pH 8.5 at 37°C for approximately 12 h. Fc fragments were removed using Protein A Sepharose (GeneScript).

### Pseudovirus-based neutralization assay

The neutralizing activity of NABs was determined using a pseudotype-based neutralization assay as we previously described (Liu et al., 2020). The pseudovirus was generated through cotransfection of 293T cells with 2 plasmids, pVax-1-S-COVID19 and pNL4-3 Luc_Env_Vpr, carrying the optimized spike (S) gene (QHR63250) and a human immunodeficiency virus type 1 backbone, respectively (Liu et al., 2020). Viral supernatant was collected 48 h posttransfection and frozen at −150°C. The serially diluted NABs were incubated with 200 TCID₅₀ of pseudovirus at 37°C for 1 h. The HuNab-virus mixtures were subsequently added into preseeded HEK293T-ACE2 cells. After 48 h, infected cells were lysed to measure luciferase activity using a commercial kit (Promega, Madison, WI). Half-maximal (IC₅₀) or 90% (IC₉₀) inhibitory concentrations of the evaluated HuNAbs were determined by log (inhibitor) versus normalized response–Variable slope using GraphPad Prism 6 (GraphPad Software Inc.).
Neutralization activity of HuNAbs against live SARS-CoV-2

The SARS-CoV-2 focus reduction neutralization test (FRNT) was performed in a certified biosafety level 3 laboratory. Neutralization assays against live SARS-CoV-2 were conducted using a clinical isolate (HKU-001a strain, GenBank accession no: MT230904.1) previously obtained from a nasopharyngeal swab of an infected patient (Chu et al., 2020). Serial dilutions of testing antibodies were conducted, mixed with 50 μL of SARS-CoV-2 (1 × 10^6 focus forming units/ml, FFU/mL) in 96-well plates and incubated for 1 h at 37°C. Mixture were then transferred to 96-well plates preseeded with 1 × 10^5 well Vero E6 cells and incubated at 37°C. After 24 h, the culture medium of the plates was removed and air-dried in a biosafety cabinet (BSC) for 20 min. Cells were then fixed with 4% paraformaldehyde for 30 min and air-dried in the BSCs again. Cells were further permeablized with 0.2% Triton X-100 and incubated with cross-reactive rabbit anti-SARS-CoV-2-N IgG (Sino Biological, Inc.) for 1 h at RT before adding Alexa Fluor 488 goat anti-rabbit IgG (H+L) cross-adsorbed secondary antibody (Life Technologies). The numbers of SARS-CoV-2 foci were calculated using the Sapphire Biomolecular Imager (Azure Biosystems).

Antibody binding kinetics and competition with receptor ACE2 measured by SPR

The binding kinetics and affinity of HuNAbs to SARS-CoV-2 spike protein (ACROBiosystems) were analyzed by surface plasmon resonance (SPR, Biacore 8K, GE Healthcare). Specifically, spike protein was covalently immobilized to a CM5 sensor chip via amine groups in 10 mM sodium acetate buffer (pH 5.0) for a final RU of approximately 500. SPR assays were run at a flow rate of 30 μL/min in HEPES buffer. For conventional kinetic/dose-response, a concentration series of HuNAbs IgGs or Fabs were injected across the spike protein surface for 120 s, followed by a 600 s dissociation phase using a multicycle method. The remaining analytes were removed in the surface regeneration step with injection of 10 mM glycine-HCl (pH 2.0) for 2 × 30 s at a flow rate of 30 μL/min. Kinetic analysis of each reference subtracted injection series was performed using Biacore Insight Evaluation software (GE Healthcare). All sensogram series were fit to a 1:1 (Langmuir) binding model of interaction. Prior to evaluating competition between antibodies and the human ACE2 peptidase domain, both the saturated binding concentrations of antibodies and ACE2 protein (ACROBiosystems) with immobilized SARS-CoV-2 spike protein were determined separately. In the competitive assay, antibodies at the saturated concentration were injected onto the chip immobilized with spike protein for 120 s until binding steady-state was reached. ACE2 protein at the saturated concentration was then injected for 120 s, which was followed by another 120 s of injection of antibody to ensure the saturated binding reaction against the immobilized spike protein. The differences in response units between ACE2 injection alone and prior antibody incubation reflect the antibodies’ competitive ability against ACE2 in binding with spike protein.

Epitope mapping through a yeast surface display system

The antigen library displayed on the yeast surface was constructed by methods described previously (Guo et al., 2015; Zuo et al., 2011). The size of the full-length SARS-CoV-2 spike gene library is approximately 10^6. The conditions of yeast culture and antigen expression induction were described previously (Chao et al., 2006; Guo et al., 2015; Zuo et al., 2011). Galactose-induced yeast library cells (10^9-10^7 cells/test) were blocked by FACS buffer (2% FBS in PBS) for 30 min at 4°C and then incubated with 2 μg screened anti-RBD (SARS-CoV-2) HuNAbs in 50 μL FACS buffer at 4°C for 1 h. After washing twice, yeasts were incubated with AF546-conjugated goat anti-human IgG secondary antibodies (Invitrogen) for another 1 h at 4°C (0.2 μg/test in 50 μL FACS buffer). The fluorescence-positive (PE channel) cells (50k-100k) were sorted into yeast culture media, recultured and reinduced by galactose. The induced yeasts were stained with the same antibodies under the same conditions. The sorted yeast single clones from the second round were verified by FACS. The displayed gene fragments in the plasmids of positive yeasts were amplified by yeast colony PCR and then sequenced. The sequence data were analyzed by Sequencher 5.4.6. (GeneCodes Corp.).

Syrian hamster experiments

In vivo evaluation of HuNAb ZDY20 in an established golden Syrian hamster model of SARS-CoV-2 infection was performed as described previously with slight modifications (Chan et al., 2020a). Briefly, 6- to 8-week-old male and female hamsters were randomized from different litters into experimental groups. The hamsters were obtained from the Chinese University of Hong Kong Laboratory Animal Service Centre through the HKU Centre for Comparative Medicine Research. The experiments were performed in compliance with the relevant ethical regulations. For prophylaxis studies, 24 h before live virus challenge, hamsters were intraperitoneally administered one dose of HuNAb ZDY20 in phosphate-buffered saline (PBS) at 10 mg/kg (n = 4), 5 mg/kg (n = 3), or ZB8 at 4.5 mg/kg (n = 4) or 2-15 at 1.5 mg/kg (n = 4) or control antibody VRC01 at 10 mg/kg (n = 4). On day 0, blood samples were collected from each hamster. Hamsters were then intranasally inoculated with a challenge dose of 100 μL of Dulbecco’s modified eagle medium containing 10^5 PFU of SARS-CoV-2 (HKU-001a strain, GenBank accession no: MT230904.1) under intraperitoneal ketamine (200 mg/kg) and xylazine (10 mg/kg) anesthesia. For intranasal administration of HuNAb, each hamster was received 100 μL of PBS containing 10 mg/kg ZDY20 (n = 4) or 4.5 mg/kg ZB8 (n = 4) under intraperitoneal ketamine (200 mg/kg) and xylazine (10 mg/kg) anesthesia at 12 h before 10^5 PFU of SARS-CoV-2 challenge. For the treatment study, each hamster received one 10 mg/kg dose of intraperitoneal HuNAb ZDY20 at 24 h, 48 h, or 72 h (n = 4 per group) after virus challenge. The hamsters were monitored twice daily for clinical signs of disease. Syrian hamsters typically clear virus within one week after SARS-CoV-2 infection (Chan et al., 2020a). Accordingly, animals were sacrificed for analysis at day 4 after virus challenge. Half nasal turbinates, trachea, and lung tissues were used for viral load determination by quantitative SARS-CoV-2-specific RdRp/Hel reverse-transcription polymerase chain reaction assay (Chan et al., 2020c) and infectious virus titration by plaque assay (Chan et al., 2020a). The viral load was normalized by β-actin while the plaque-forming units (PFU) was normalized by mg of tissue extractions based on hamster’s bodyweight. For measuring antibody
concentrations in nasal wash, hamsters were treated with Isoflurane lightly, then 200 μL of PBS was injected into one nasal opening while collecting the turbid wash from the other one without any blood contamination.

**DNA vaccination in Syrian hamsters**

To determine vaccine-induce protection against live SARS-CoV-2, we generated a DNA vaccine using pVAX vector encoding the complete spike protein as we previously described for SARS DNA vaccines (Yi et al., 2005). A dose of 150 μg DNA was used to immunize a group of 6–8 weeks old hamsters (n = 4) at three-week intervals for four time through intramuscular electroporation delivery. The voltage of electroporation was pre-set 60 V in the TERESA DNA Delivery Device (Shanghai Teresa Healthcare Sci-Tech Co., Ltd). Another group of hamsters (n = 3) received equal amount of pVAX vector as a negative control. Serum samples were collected for measuring neutralizing antibodies three weeks after the final vaccination. The vaccinated animals were then challenged intranasally with the same dose as mentioned above at eight weeks after the final vaccination. Animals were consistently sacrificed for endpoint analysis at day 4 after virus challenge.

**Histopathology and immunofluorescence (IF) staining**

The lung tissues collected at necropsy were fixed in zinc formalin and then processed into paraffin-embedded tissue blocks. The tissue sections (4 μm) were stained with hematoxylin and eosin (H&E) for light microscopy examination as previously described with modifications. For identification and localization of SARS-CoV-2 nucleocapsid protein (NP) in organ tissues, immunofluorescence staining was performed on deparaffinized and rehydrated tissue sections using rabbit anti-SARS-CoV-2-N protein antibody together with FITC-conjugated donkey anti-rabbit IgG (Jackson ImmunoResearch, PA, USA). Briefly, the tissue sections were first treated with antigen unmasking solution (Vector Laboratories) in a pressure cooker. After blocking with 0.1% Sudan black B for 15 min and 1% bovine serum albumin (BSA)/PBS at RT for 30 min, the primary antibody rabbit anti-SARS-CoV-2-N antibody (1:4000 dilution with 1% BSA/PBS) was incubated at 4°C overnight. This step was followed by FITC-conjugated donkey anti-rabbit IgG (Jackson ImmunoResearch) for 30 min and then mounted with 4',6-diamidino-2-phenylindole (DAPI). All tissue sections were examined, and the images were captured with an Olympus BX53 semimotorized fluorescence microscope using cellSens imaging software (Olympus). NP+ cells per 50 × magnification field were quantified based on the mean fluorescence intensity (MFI) using the ImageJ (https://imagej.nih.gov/ij/download.html).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistical analysis was performed using PRISM 6.0 or SPSS 26.0. Ordinary one-way ANOVA and multiple comparisons were used to compare group means and differences between multiple groups. Unpaired t test was used to compare group means between two groups only. A P-value < 0.05 was considered to be significant. The figure legends and Methods Materials denotes the number of independent experiments performed, number of animals in each group and specific details on statistical tests.