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Single domain antibodies derived from ancient animals as broadly neutralizing agents for SARS-CoV-2 and other coronaviruses

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ABSTRACT

With severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) as an emergent human virus since December 2019, the world population is susceptible to coronavirus disease 2019 (COVID-19). SARS-CoV-2 has higher transmissibility than the previous coronaviruses, associated by the ribonucleic acid (RNA) virus nature with high mutation rate, caused SARS-CoV-2 variants to arise while circulating worldwide. Neutralizing antibodies are identified as immediate and direct-acting therapeutic against COVID-19. Single-domain antibodies (sdAbs), as small biomolecules with non-complex structure and intrinsic stability, can acquire antigen-binding capabilities comparable to conventional antibodies, which serve as an attractive neutralizing solution. SARS-CoV-2 spike protein attaches to human angiotensin-converting enzyme 2 (ACE2) receptor on lung epithelial cells to initiate viral infection, serves as potential therapeutic target. sdAbs have shown broad neutralization towards SARS-CoV-2 with various mutations, effectively stop and prevent infection while efficiently block mutational escape. In addition, sdAbs can be developed into multivalent antibodies or inhaled biotherapeutics against COVID-19.

1. Introduction

1.1. Trend of COVID-19

Recent outbreak of coronavirus disease 2019 (COVID-19) is attributable to severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). SARS-CoV-2 has high sequence homology (shares 96.2% sequence identity) with the bat coronavirus RaTG13, implying the virus may have originated in bats and latterly crossed to humans in 2019 [1, 2]. Parallel to severe acute respiratory syndrome coronavirus
(SARS-CoV) and Middle East respiratory syndrome coronavirus (MER- S-CoV), SARS-CoV-2 presents as respiratory infection with symptoms of fever, cough, and pneumonia, but in severe cases may be progressing into organ failure [3].

With SARS-CoV-2 as an emergent human virus since December 2019, the world population is susceptible by reason of the deficiency in proper treatments for COVID-19. Generic or repurposed drug candidates are in trials, as yet with unremarkable results. Hydroxychloroquine, as an antimarial drug, demonstrated antiviral activity against SARS-CoV-2 at micro molar concentrations in tissue culture, with clinical benefit in observational trials involving small number of patients; but showed no effect in reducing mortality for the large observational clinical trials [4–6]. Lopinavir–ritonavir, a human immunodeficiency (HIV)-1 protease inhibitor, demonstrated antiviral activity against SARS-CoV in tissue culture and infected patients; but showed no clinical benefit against SARS-CoV-2 [7]. Remdesivir, originally designed to treat Ebola virus infection, also displayed broad-spectrum antiviral activity against ribonucleic acid (RNA) viruses including SARS-CoV and MERS-CoV in tissue culture and animal models [8]. Remdesivir has been found to inhibit the RNA-dependent RNA polymerase of SARS-CoV-2, which improve recovery time in COVID-19 patients, nonetheless showed no effect in reducing mortality [9,10].

Active immunization has been considered, with multiple COVID-19 vaccines have been permitted under emergency use [11]. Sinovac’s CoronaVac, as inactivated vaccine, utilizing SARS-CoV-2 which has been killed by chemical or physical ways to trigger an immune response. AstraZeneca/Oxford’s AZD1222, as viral vector vaccine, utilizing non-replicating adenovirus as a vector containing genetic material of SARS-CoV-2 to trigger an immune response. Moderna’s mRNA-1273, as RNA-based vaccine made from the viral sequence of SARS-CoV-2, with the immune cells processing the mRNA to manufacture protein that triggers an immune response [12]. Nevertheless, the timeline for developing a safe, effective and widely available vaccine for SARS-CoV-2 remains tentative. Besides, vaccine may not be 100% effective for immunocompromised individuals, therefore with therapeutics would be beneficial [13]. Passive immunization via the trans-fusion of serum collected from COVID-19 convalescent individuals to the critically ill COVID-19 patients has led to better clinical outcomes, suggesting neutralization of virus by the existing antibodies is useful [14]. To date, specifically designed neutralizing antibody therapies such as Regeneron’s REGEN-COV (casirivimab with imdevimab), Lilly’s bamlanivimab with etesevimab, GlaxoSmithKline and Vir Biotech’s sotrovimab, AstraZeneca’s Evusheld (tixagevimab with cilgavimab), Lilly’s bebtelovimab have received authorization by the U. S. Food and Drug Administration (FDA) for emergency use in COVID-19 treatment [15–17].

SARS-CoV-2 was marked as one of the most transmissible coronaviruses that spreading rapidly and unceasingly throughout the world, adversely impacted human health while resulting in medical burden and lives lost. In March 2020, World Health Organization (WHO) declared COVID-19 as the first coronavirus pandemic in history, to be a public health emergency of international concern [18,19]. In course of prolonged infections, SARS-CoV-2 with escape mutants emerged. SARS-CoV-2 variants have been arisen in different countries due to the selection pressure across the worldwide spread, associated by the RNA virus nature with high mutation rate [20]. Till late 2021, five SARS-CoV-2 variants of concern have been identified: B.1.1.7 (Alpha), B.1.1617 (Gamma), B.1.617.2 (Delta), and B.1.1.529 (Omicron) [21–23]. Prevalent circulating variants of SARS-CoV-2 wane vaccine-elicited serologic responses and evade recognition and neutralization by clinical antibodies, as a daunting challenge that confront the development of therapeutics [24,25].

The pandemic caused by SARS-CoV-2 continues to spread, resulted in over 601.19 million infections and over 6.48 million deaths worldwide with the numbers still rising as of 2 September 2022 [26]. COVID-19 pandemic gradually leads to the collapse of healthcare systems, imposes a substantial social burden while causing tremendous economic losses worldwide. Hence, therapeutics that can stop as well as prevent SARS-CoV-2 infection in an effective manner are in crucial need. The development of therapeutics is actively in progress, with the recognition of therapeutic neutralizing antibodies as immediate and direct-acting antiviral agents accounted for short-to-medium term approach to combat COVID-19.

1.2. Characteristics of the cause for COVID-19: SARS-CoV-2

Coronaviruses can be categorized into four genera (α, β, γ, δ) which can infect a wide range of host organisms [27]. Thus far, seven types of coronaviruses that can cause disease in humans have been identified. HCoV-HKU1, HCoV-OC43, HCoV-NL63, HCoV-229E circulate seasonally and globally while causing mild respiratory disease [28]. SARS-CoV, MERS-CoV, and SARS-CoV-2 are zoonotic pathogens that have entered the human population over the last two decades, causing severe respiratory symptoms with high mortality and eventually leads to epidemics or pandemics [29–31]. SARS-CoV-2 which classified into the genus betacoronavirus in the family Coronaviridae, are more transmissible than previous coronaviruses [32,33]. In view of structural properties, SARS-CoV-2 is an enveloped virus, with positive-sense and single-stranded RNA genome (29,903 nucleotides in length) which encodes replicase and four major structural proteins: spike, envelope, membrane and nucleocapsid, like SARS-CoV [1,32,34]. Spike (S), envelope (E), and membrane (M) proteins are majorly incorporated into SARS-CoV-2 envelope lipid bilayer, enclosing a helical capsid formed by nucleocapsid (N) proteins bound to the RNA genome; hence formed the SARS-CoV-2 virion (Fig. 1) [11].

SARS-CoV-2 S glycoprotein forms homotrimers that protrude from envelope and then give rise to the coronal appearance, which can bind to the peptidase domain of human angiotensin-converting enzyme 2 (ACE2) as a host cell receptor on the cell membranes of type 2 pneumocytes, to invade susceptible cells for SARS-CoV-2 viral entry (Fig. 2) [11]. The trimeric complex composed of 1,273 amino acids while structurally belongs to the class I membrane fusion protein, where each protomer is functionally categorized into two distinctive subunits: N-terminal S1 subunit and C-terminal S2 subunit be parted by a furin cleavage site. The S1 region mainly includes the roughly 200-residue receptor-binding domain (RBD) for the interaction with the host cell receptor, while the S2 region holds the membrane fusion machinery, encompassing the hydrophobic fusion peptide and two heptad repeats, HR1 and HR2 that can interact to form six-helical bundle as a post-fusion structure [1,35–37]. Receptor binding by RBD in S1 subunit, proteolytic processing at the furin cleavage site between the S1 and S2 subunits via host cell transmembrane serine protease 2 (TMPRSS2) followed by S1 subunit shedding, S2 subunit structural rearranging into stable post-fusion conformation, the fusion of the viral membrane with the host cellular membrane are regarded as the key events for facilitating subsequent viral entry into the host cell [37–39].

RBD, as a globular domain positioned on the distal surface of SARS-CoV S, MERS-CoV S, and SARS-CoV-2 S, has undergone conformational rearrangements in a dynamic manner by interchangeably masking and presenting their receptor-binding interfaces as well as neutralizing epitopes to either host cells or potential neutralizing antibodies [36]. Initial cryogenic electron microscopy (cryo-EM) characterization of the SARS-CoV-2 spike in the pre-fusion conformation revealed two distinctive configurations adopted by the RBDs: in the ‘up’ state, RBD is away from the spike protein that it can engage ACE2 without steric clash; in the ‘down’ state, RBD is tightly packed against the top of the S2 subunit that it prevents ACE2 binding; with the similar phenomena observed as well in SARS-CoV S and MERS-CoV S [35,40,41]. In a receptor-binding event, the RBD would be trapped in the energetically unstable ‘up’ state, towards the gradual destabilization of S1 until S2 is eventually triggered to initiate membrane fusion [42]. Throughout the life cycle of virus, the spike trimmer exists in an equilibrium between an inactive, closed
conformation with all RBDs in the ‘down’ states and an active, open conformation with the RBDs in mixed ‘up down’ states. The SARS-CoV-2 S protein predominantly shown an asymmetrical homotrimer, with one RBD in ‘up’ state is ACE2-accessible while the other two RBDs in ‘down’ states are not [35].

SARS-CoV-2 RBD comprises residues 319 to 541, including the receptor-binding motif (RBM) spanning residues 438 to 506 which contains major ACE2-contacting residues: ACE2 interacts with residue F486 protruding from the 481 to 487 loop of SARS-CoV-2 RBD, for instance [37,43,44]. The SARS-CoV-2 variants of concern, B.1.1.7, B.1.351, P.1, B.1.617.2, and B.1.1.529, are known to carry several mutations in RBD: B.1.1.7 with N501Y mutation; B.1.351 with K417N, E484K, and N501Y mutations; P.1 with K417T, E484K, and N501Y mutations; B.1.617.2 with L452R and T478K mutations; B.1.1.529 with G339D, S371L, S373P, S375F, K417N, N440K, G446S, S477N, T478K, E484A, Q493K/R, G496S, Q498R, N501Y, and Y505H mutations [45–50]. Owing to viral components undergo structural changes, SARS-CoV-2 variants retain the same host cell receptor binding, however might no longer be recognized by the specific antibodies [51].

The RBD of SARS-CoV-2 spike protein involves in ACE2 receptor engagement to initiate viral infection. SARS-CoV-2 RBD-based immunogens were shown to stimulate the generation of neutralizing sera in animals, indicates that RBD contains immunodominant epitopes capable to elicit neutralizing antibodies that can provide protection against SARS-CoV-2 infection [52]. Taking these factors into account, RBD is a potential therapeutic target for the development of broadly neutralizing antibodies to stop and prevent SARS-CoV-2 viral infection.
1.3. Differences between SARS-CoV, MERS-CoV and SARS-CoV-2

COVID-19 caused by SARS-CoV-2 marks the third major coronavirus outbreak since last two decades, following severe acute respiratory syndrome (SARS) caused by SARS-CoV in 2002 and Middle East respiratory syndrome (MERS) caused by MERS-CoV in 2012 (Table 1) [30, 53]. SARS-CoV, first identified in Guangdong province of China on November 2002, resulted in the SARS epidemic with over 8,000 infections and a ~10% fatality rate. The outbreak of SARS disease was ended in 2004 [54]. MERS-CoV, emerged in Saudi Arabia on June 2012, resulted in the MERS epidemic with over 2,500 infections and a ~35% fatality rate as of July 2022 [55]. SARS-CoV-2, first identified in Wuhan, Hubei province of China on December 2019, resulted in COVID-19 pandemic with over 601.19 million infections and over 6.48 million deaths worldwide, with the numbers still rising as of 2 September 2022 [26]. It was observed that SARS-CoV-2 caused higher proportion of asymptomatic and mild symptomatic infections as compared to SARS-CoV and MERS-CoV [56]. With the active viral replication of SARS-CoV-2 in the upper respiratory tract (URT), leads to the viral shedding begins from incubation period then peaked during the time of symptom onset showing mild or no symptoms, in contrast to SARS-CoV and MERS-CoV with the viral shedding begins from the time of symptom onset then peaked in the second week after symptom onset [56,57]. As a result, pre-symptomatic transmission is rare for SARS-CoV and MERS-CoV, but plays roles in SARS-CoV-2 spread [56]. On the other hand, MERS-CoV with inefficient human-to-human transmission as compared to SARS-CoV and SARS-CoV-2, but leads to higher mortality [58].

SARS-CoV, MERS-CoV, and SARS-CoV-2 are classified into the genusBetacoronavirusin the family Coronaviridae: SARS-CoV and SARS-CoV-2 within the subgenus sarbecovirus, MERS-CoV within the subgenusmerbecovirus [33]. By analysing the spike protein sequence among SARS-CoV, MERS-CoV and SARS-CoV-2, SARS-CoV-2 showed greater sequence homology with SARS-CoV (76% identity, 87% similarity) than with MERS-CoV (42% identity, 58% similarity) [59]. Therefore, with the sharing of significant sequence identity between SARS-CoV and SARS-CoV-2 suggests the possibility of cross-reactivity. Nevertheless, there is lower sequence homology within spike N-terminal regions, taking account of the dissimilarity in regions including RBD while correlating to the difference in host cell receptors used [60]. The primary functional host cell receptor for SARS-CoV and MERS-CoV are ACE2 and dipeptidyl peptidase 4 (DPP4), respectively [61,62]. SARS-CoV-2, similar to its closest homolog SARS-CoV, utilizes ACE2 to enter host cell as well; however by 10 to 20-fold greater affinity (equilibrium dissociation constant, $K_D \approx 15 \text{ nM}$) than SARS-CoV [35,63]. E484, F486, Q493, and N501 are important RBD residues contribute to the stronger binding of ACE2 towards SARS-CoV-2 than that of SARS-CoV [43]. Hence, with a higher affinity towards ACE2 resulted from the sequence changes in RBD may supports SARS-CoV-2 in more efficient host cell entry, drives higher transmissibility of SARS-CoV-2 [38]. Besides, it was suggested that there is a difference in the immunogenicity of RBD between SARS-CoV and SARS-CoV-2 [64]. The glycans at residues N165, N234, N343 of spike protein shield SARS-CoV-2 RBD from the antibodies [65]. A more hidden RBD of SARS-CoV-2 than that of SARS-CoV leads to immune evasion as potential viral strategy [66].

| Feature                                | SARS-CoV       | MERS-CoV   | SARS-CoV-2         |
|----------------------------------------|----------------|------------|--------------------|
| Disease                                | SARS           | MERS       | COVID-19           |
| First identified                       | Guangdong, China (2002) | Saudi Arabia (2012) | Wuhan, China (2019) |
| Taxonomy                               | Betacoronavirus (Sarbecovirus) | Betacoronavirus (Mesbecovirus) | Betacoronavirus (Sarbecovirus) |
| Primary host cell receptor receptor    | ACE2           | DPP4       | ACE2               |
| Human-to-human transmission            | Efficient      | Inefficient | Efficient          |
| Viral shedding                         | Starts from the time of symptom onset. | Starts from the time of incubation period. | Starts from the time of symptom onset. |
| Proportion of asymptomatic and mild symptomatic infections | Low | Low | High |
| Pre-symptomatic transmission           | Rare           | Rare       | Plays roles in viral spread |
| Level of spread                        | Epidemic      | Moderate   | High               |

Table 1
A summary comparison of three highly pathogenic coronaviruses as the disease causative agents: SARS-CoV, MERS-CoV and SARS-CoV-2.

ACE2, angiotensin-converting enzyme 2; COVID-19, coronavirus disease 2019; DPP4, dipeptidyl peptidase 4; MERS, Middle East respiratory syndrome; MERS-CoV, Middle East respiratory syndrome coronavirus; SARS, severe acute respiratory syndrome; SARS-CoV, severe acute respiratory syndrome coronavirus; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2.
donors, moreover the quality of mAbs is easier to control. The development of therapeutic mAbs serves as the frontline in combating COVID-19 [78]. For instance, neutralizing mAb regdanvimab developed by Celltrion Inc. has been approved in South Korea to treat mild COVID-19 in patients aged above 50 years old with underlying health problem such as cardiovascular disease, diabetes or obesity, due to its potency to reduce the risk of progression to severe disease in COVID-19 patients [79,80].

However, mAbs as large biomolecules are intravenously delivered with low efficiency across the plasma-lung barrier for pulmonary infection treatment, thus high administration doses of mAbs in several grams would be required for efficient neutralization [81–83]. The production of mAbs in large scale usually takes at the minimum of 3 to 6 months, thus would be difficult to achieve a timely production during pandemic [64]. As mAbs undergo post-translational modifications, the production of mAbs require eukaryotic expression system by using mammalian cells that are expensive to maintain [84]. In addition, mAbs can be degenerated due to the exposure to extreme ambient conditions such as humidity and high temperatures [67]. As there are diverse environmental conditions for varying countries, while there is the requirement for specific temperature to be maintained during storage and transportation of mAbs, widespread clinical use of mAbs may be limited. With the aim to improve properties in therapeutic applications, recombinant antibodies are generated using molecular lab techniques.

Antibody fragments or domains, such as Fab (antigen-binding fragment, 50 kilodaltons (kDa)), scFv (single-chain variable fragment, 30 kDa), and VH (heavy chain variable domain, 15 kDa) are appealing antibody formats to be used as smaller biomolecules for therapeutics [85].

In the early 1990s, camelids or cartilaginous fish were found to be possessing unconventional antibodies in their immune system: exists as a homodimer that is naturally devoid of light chains, thus known as heavy-chain only antibodies [86]. A single variable domain (VHH from camelid or VNAR from cartilaginous fish) represents the antigen-binding region of heavy-chain only antibodies, in lieu of two variable domains (VH and VL) that typically forms the antigen-binding region of conventional IgG (Fig. 3). VHH contains three hypervariable loops, denoted complementarity-determining region 1 (CDR1), CDR2 and CDR3; VNAR contains four hypervariable loops, denoted CDR1, hypervariable region 2 (HV2), HV4 and CDR3 [32,67]. CDR3 comprises the most variable region in antibody, also with at least 60 to 80% of the contact with the antigen, thus mainly contributes to the specific binding of antibodies towards target antigens [20,87,88]. The single variable domain from heavy-chain only antibodies can be expressed independently as a ~12 to 15 kDa antibody fragment, with the acquired specificity and affinity for target antigen is comparable to the conventional antibodies, therefore contain autonomous function as single-domain antibodies (sdAbs) (Fig. 4) [86,89,90].

Single-domain antibody, with a molecular weight of ~12 to 15 kDa,
is approximately one-tenth of the size of a conventional IgG in ~150 kDa [90,91]. As small biomolecules, sdAbs exhibit efficient tissue penetration [92,93]. sdAbs are less affected by steric hindrances that interfere with the binding for large conventional antibodies, as a result sdAbs have larger number of accessible epitopes [94,95]. In addition, with an extended antigen-binding region due to a long protruding CDR3 loop, sdAbs are capable to access cryptic epitopes [96,97]. Hence, sdAbs retain full antigen-binding capacity of antibody. Small size sdAbs allow for rapid kilogram-scale production with ease in prokaryotic expression systems, leads to high yield with relatively low production cost, consequently enable fast implementation during the outbreak [94,98]. As non-complex structure, sdAbs can be expressed in yeast and mammalian cells as well [99–101]. Since sdAbs do not bind light chains, with the absence of hydrophobic interface between VH and VL domain render the sdAbs a more hydrophilic surface, thus have high solubility for ease of downstream processing [102–104]. The intrinsic stability of sdAbs as exemplified by the inherent thermostability and chemostability, enables sdAbs to withstand prolonged storage [36,68,105,106]. For instance, sdAbs are with tolerance towards pH ranging from 3 to 11, also with resistance to chemical denaturant (0.35 - 8 M urea) [107–109,105,110]. Therefore, sdAbs can be reserved as a stockpile of therapeutic options for future epidemic.

In addition, the small size and non-complex structure of sdAbs allow flexible formatting according to the needs [68,98]. The origin of sdAbs from animals may limit their therapeutic application in humans, as there is immunogenicity risk. Thus, humanization techniques have been adopted, by modifying the animal-specific amino acid sequences within the framework into the human heavy chain variable domain as its counterpart, to reduce species heterogeneity without altering its antigen-binding affinity and solubility [94,111,112]. The monomeric nature of sdAbs has its drawback, such as their binding kinetics in terms of fast dissociation rates (k_off) may reduce neutralization potency. Therefore, sdAbs can be multimerized to enhance avidity, such that sdAbs that are designed in homo-dimeric or homo-trimeric form can increase valency to improve antiviral activities, while sdAbs that are designed in heterodimeric form can simultaneously target different epitopes to prevent virus mutational escape [113]. sdAbs have short serum half-life and rapid renal clearance due to their small size, as limitations for treatment and prevention of viral disease [67,114]. Hence, sdAbs can be fused with the crystallisable fragment (Fc) of IgG to become larger protein, to extend their blood residential time as well as prolong their circulation in the body [93,115].

In 2019, FDA has given approval to the first sdAb-based medicine, caplacizumab for the treatment of acquired thrombotic thrombocytopenic purpura, with an estimated cost of $270,000 [116,117]. The innovative nature and drug development account for the high cost of caplacizumab, nonetheless the novel therapy represents a major breakthrough [117]. Besides being used as injectable drug, the small and stable sdAbs may be nebulized and administered via inhalation directly to the airway epithelia, which can maximize their bioavailability and function by having high concentration of therapeutics at the respiratory sites infection [64,118]. It was reported that an inhaled sdAb, ALX-0171 for the treatment of respiratory syncytial virus has entered clinical trials [119]. Therefore, the use of sdAbs as biologics is an interesting approach, particularly for the treatment of respiratory infection. Similarly, with the generation of a neutralizing inhaler containing sdAbs offers a possibility for directly blocking viral replication in the upper airway during the early stages of COVID-19, meanwhile it helps to improve patient compliance by being a needle-free treatment.

2.1. Case studies on broadly-neutralizing single-domain antibody for SARS-CoV-2

Surface display technology has been utilized for the selection of sdAbs specific for the targeted antigen. There are several antibody surface display technologies, included phage display, ribosome display, yeast surface display, and bacterial surface display [13,120–122]. As exemplified by the widely employed phage display technology (Fig. 5), the gene encoding for sdAb is fused with the gene encoding for bacteriophage’s coat protein, giving rise to the display of sdAb on bacteriophage’s surface in which can be applied for the selection of antigen-specific binders [123]. Afterwards, a library of sdAbs genes are cloned into phagemid vectors, lead to the generation of a sdAb library with diversity. There are multiple types of sdAb library, inclusive of immune library and non-immune library such as naïve or synthetic library. sdAb library has the potential to be a rapidly accessed resource, which may bring about the fast-track discovery of neutralizing antibodies during an outbreak.
2.2. Camelid VHH against SARS-CoV-2 and other coronaviruses

According to the work by Wrapp et al. [36], SARS VHH-72, with high affinity to SARS-CoV RBD ($K_D = 1.2$ nM), was identified via phage-displayed sdAb library derived from a llama immunized with SARS-CoV and MERS-CoV S protein. Camelid VHH domains have high degree of homology with human type 3 VH domains, thus with the high conservation leads to low immunogenicity [124]. Crystal structure of SARS VHH-72 bound to viral target revealed that the epitope of SARS VHH-72 did not overlap with the ACE2 binding site on the SARS-CoV RBD. Instead, there were steric clashes between ACE2 and SARS VHH-72, possibly cause interferences towards ACE2 binding to RBD. SARS VHH-72 has shown the ability to cross-react with the SARS-CoV-2 RBD ($K_D = 39$ nM), as well as could interfere with the ACE2 binding. It is postulated that SARS VHH-72 binds onto the part of SARS-CoV RBD sharing low sequence variation with SARS-CoV-2 RBD, thus able to broadly bind towards SARS-CoV-like viruses.

Further engineering of the cross-reactive SARS VHH-72 into a bivalent and monomeric human IgG Fc-fusion has conferred it the ability to readily neutralize SARS-CoV-2 pseudovirus, with an IC$_{50}$ of approximately 200 ng/mL. SARS VHH-72-Fc, through the fusion to human IgG1 Fc domain, can interact with Fc receptor (FcR) expressed on immune cells such as macrophages, B cells and monocytes. The engagement of FcR activates the immune cells to get rid of viruses inside the body, with Fc-dependent cytotoxic functions such as antibody-dependent cell-mediated cytotoxicity (ADCC), complement-dependent cytotoxicity (CDC) and antibody-dependent cellular phagocytosis (ADCP) [125–127]. The multi-valency approach aimed to enhance avidity, and to circumvent ADE of viral infection that can be triggered instead through Fc-FcR interaction due to the sub-optimal antigen-antibody interactions [128]. The production of SARS VHH-72-Fc has reached expression levels of ~300 mg/L in the industry’s standard Chinese hamster ovary (CHO) cell system, which is high-yielding. The SARS-CoV-2 variants of concern, B.1.1.7, B.1.351, P.1, B.1.617.2, and B.1.1.529 share one specific mutation within the spike protein, called D614G [49,129]. D614G mutant exhibits greater trimeric spike protein stability, results from the greater incorporation of spike protein into virions with less S1 shedding [130]. D614G mutant increases the viral infectivity without any effect on pathogenesis [131–133]. Nieto et al. [122] reported that the monomeric Fc fusion of SARS VHH-72 exhibiting modest neutralization for authentic SARS-CoV-2 wild type (D614) and D614G mutant (G614), with IC$_{50}$ of 1,287.75 nM and 1,233.90 nM, respectively. The hexavalent Fc fusion of SARS VHH-72 engineered by Zupancic et al. [134] potently neutralized SARS-CoV-2 pseudovirus variants B.1.1.7 and B.1.351, with IC$_{50}$ of 310 pM and 72 pM, respectively.

Xiang et al. [113] discovered that the serum of a llama immunized with SARS-CoV-2 RBD contained potent neutralizing sdAbs with picomolar to femtomolar affinities towards the RBD, such as Nb20 ($K_D = 10.4$ pM) and Nb21 ($K_D < 1$ pM) neutralized SARS-CoV-2 pseudovirus with IC$_{50}$ of 102 pM and 45 pM, respectively. Structural characterization through cryo-EM showed that Nb20 or Nb21 binds to epitope that partially overlapping with the ACE2 binding site on RBD, with their CDR1 and CDR3 would clash with the ACE2 α-1 helix containing major portion of residues responsible for the coordination of ACE2-RBD interaction, which can cause steric interference towards ACE2 binding when bound to the RBDs in ‘up’ states. Meanwhile, Nb20 or Nb21 can bind to the RBDs in ‘down’ states as well, indicates with the concurrent binding of three Nbs to all three RBDs in ‘down’ states may be locking the spike into an inactive conformation. The homo-trimeric constructs...
based on Nb20 and Nb21 have shown up to ~30-fold improvement of neutralization potency towards SARS-CoV-2 pseudovirus as compared to that of their monomeric form, with IC_{50} of 4.14 pM and 1.32 pM, respectively. Besides, Nb20 and Nb21 potently neutralized SARS-CoV-2 Munich strain (D614G mutant) with IC_{50} of 48 pM and 22 pM, respectively, while the homo-trimeric constructs of Nb20 and Nb21 have shown up to a ~6-fold increase of neutralization potency towards SARS-CoV-2 Munich strain with IC_{50} of 5.43 pM and 6.04 pM, respectively [113,135]. Nb20 and Nb21 in both monomeric and homo-trimeric form presented thermal stability ranged from 70 to 72.8 °C. Xiang et al. [113] also identified Nb34 and Nb95 which can only bind with at least two RBDS in ‘up’ states, with epitopes that do not overlap with the ACE2 binding site. Nb34 is fitted onto the top of spike trimer, locking the helices of S2 at pre-fusion conformation and thus prevent membrane fusion [5]. Nb95 is accommodated nearby to the firmly fixed N-terminal domain (NTD) of spike trimer, which may restrict spike flexibility. It was demonstrated that Nb34 and Nb95 can neutralize SARS-CoV-2 Munich strain as well, with IC_{50} of 1.125 nM and 5.105 nM, respectively.

The selection of sdAbs from a synthetic library requiring at least 2 to 3 weeks, as compared to the traditional generation of sdAbs with at least 6 weeks for animal immunization followed by within 3 to 4 months for antibody selection [136]. Schoof et al. [121] utilized a yeast surface-displayed synthetic llama sdAb library, where an anti-SARS-CoV-2 RBD Nb6 (K_{D} = 41 nM) that inhibited SARS-CoV-2 pseudovirus infection with IC_{50} of 2,000 nM was identified. Cryo-EM showed that Nb6 binds to spike in completely inactive conformation by recognizing RBD epitope that overlap with ACE2 binding site. It was observed that one Nb6 takes a straddle position at the interface between two ‘down’-state RBDS, with its CDR3 reach over to the neighbouring RBDS: indicates neutralization mechanism by locking the two RBDS into ‘down’ states while can pre-organize the binding site for a second and third Nb6 molecule, hence will stabilize the closed spike conformation that renders RBDS inaccessible to ACE2 binding. With affinity maturation of Nb6 by mutagenesis of CDR1 and CDR3 region, in addition with multivalency design, resulted in generating a matured Nb6 in trivalent form (mNb6-tri). mNb6-tri neutralized SARS-CoV-2 pseudovirus with IC_{50} of 120 pM, showing up to ~10-fold enhanced neutralization potency compared to that of Nb6. It is predicted that the neutralization mechanism of mNb6-tri involves conformational control of RBD accessibility, in which one mNb6-tri can simultaneously lock all three RBDS into ACE2-inaccessible ‘down’ states. Furthermore, the monovalent Nb6 and engineered mNb6-tri neutralized authentic SARS-CoV-2 (isolate France/IDF0372/2020, V367F mutant) with IC_{50} of 3,300 nM and 54 pM, respectively [121,137]. mNb6-tri retains function after the heat treatment for an hour at 50 °C, aerosolization, as well as lyophilization.

Gai et al. [20] have isolated Nb11-59 specific for SARS-CoV-2 RBD (K_{D} = 21.6 nM) while exhibited the most potent neutralizing activity against authentic SARS-CoV-2 with 50% neutralizing dose (ND_{50}) of 550 ng/mL, via phage-displayed sdAb libraries generated from camels immunized with SARS-CoV-2 RBD. Nb11-59 was shown to block the interaction between human ACE2 with the RBD of closely related beta-coronaviruses: bat-SL-CoV-WIV1 RBD and SARS-CoV RBD; as well as can block the interaction between ACE2 and eight SARS-CoV-2 RBD mutants, including Q321L, V341I, N354D, V367F, K378R, V483A, Y508H, and H519P circulated in China, England, France, and the United States. [138,139]. Humanized Nb11-59 (HunNb11-59) can be mass-produced using the methylotrophic yeast Pichia pastoris, with 99% purity and >20 g/L yield. High conservation of the HunNb11-59 has been proven, with a good stability profile at temperature ranged from 4 to 40 °C, also with a consistent post-nebulization stability profile showing merely small aggregates (0.23%) formed after nebulization.

Nieto and et al. [122] developed single-step sdAb selection using Escherichia coli surface-displayed sdAb library derived from an alpaca immunized with SARS-CoV-2 S protein, coupled with non-complex density gradient centrifugation. The bacterial surface-displayed system utilizes the high transformation efficiency of Escherichia coli, neither the infection by bacteriophages nor the shutting into yeast cells is required for the surface display of sdAbs [140]. W25, which targets SARS-CoV-2 RBD (K_{D} = 295 pM), efficiently competed against ACE2 for binding to RBD with an IC_{50} of 33 nM. W25 neutralized authentic SARS-CoV-2 wild type and D614G mutant, with IC_{50} of 9.28 nM and 5.09 nM, respectively. W25Fc, as a dimeric Fc fusion of W25, had a better neutralizing performance for authentic SARS-CoV-2 wild type and D614G mutant with IC_{50} of 7.39 nM and 3.69 nM, respectively. Interestingly, there was a slight enhancement in neutralization effect towards the D614G mutant. In addition, the effective conjugation by covalently labelled W25 to Horseradish Peroxidase (HRP) may be useful for the diagnostic development involving direct antigen detection.

Pymm et al. [44] have identified the four most potent sdAbs against SARS-CoV-2 RBD using the phage-displayed sdAb libraries generated from alpacas immunized with spike protein from SARS-CoV-2 and RBD from SARS-CoV and SARS-CoV-2: WNb 2 (K_{D} = 360 pM), WNb 7 (K_{D} = 260 pM), WNb 15 (K_{D} = 140 pM), and WNb 36 (K_{D} = 430 pM). These sdAbs bound to RBD can be divided into two major groups: Cluster 1 sdAbs, as exemplified by WNb 7 and WNb 36 did not compete with Cluster 2 sdAbs, as exemplified by WNb 2 and WNb 7 and 15 for RBD binding. Structural characterization of sdAbs-RBD complex revealed that Cluster 1 sdAb and Cluster 2 sdAb bound simultaneously to two distinct antigenic sites on RBD, with each epitope overlapped with the ACE2 binding region on RBD at different degrees. Cluster 1 sdAb, with the epitope overlapping the binding position of ACE2 α-1 helix as the primary binding site for RBD, possibly contribute to ACE2 blocking. Cluster 2 sdAb, with the epitope overlapping the position of ACE2 α-10 helix considered as small binding overlap on the RBD, but can bind to RBD in an orientation that will cause steric clashes towards ACE2-RBD binding. Remarkably, the dimeric Nb-Fc fusions, WNbFc 7 and WNbFc 15 inhibited ACE2 interaction with SARS-CoV S1 at IC_{50} of 830 pM and 1.45 nM, respectively. N501Y mutation was found in the SARS-CoV-2 variants B.1.1.7, B.1.351, P.1 and B.1.1.529 [49,141-143]. N501, one of the six key ACE2-contacting residues inside the RBD, is important for ACE2-RBD interaction [144,145]. With the N501Y mutation, results in the higher binding affinity of RBD for ACE2, which leads to increasing viral transmissibility [146,147]. The two-antibody mixture combination, WNbFc 36 + 7 and WNbFc 2 + 15 neutralized authentic SARS-CoV-2-D614G N501Y mutant as well as the wild type SARS-CoV-2, with IC_{50} of ~100 pM and ~300 pM, respectively. Prophylactic administration of WNbFc 36 + 7 at a dose of 0.2 mg/kg has decreased the viral RNA load in the lung by up to 10^-fold in the SARS-CoV-2 D614G N501Y mutant-infected mice at 3 days‘ post infection (dpi), thus displayed the potential of antibody cocktails as prophylactic agents against SARS-CoV-2 in vivo.

Koenig et al. [148] have selected VH3 E (K_{D} = 2 nM) and VH3 Y (K_{D} = 9 nM) as high affinity binders specific for SARS-CoV-2 RBD via phage-displayed sdAb libraries generated from an alpaca and a llama immunized with SARS-CoV-2 RBD and inactivated SARS-CoV-2. VH3 E and VH3 Y neutralized SARS-CoV-2 pseudovirus with IC_{50} at 60 nM and 198 nM, respectively, likewise neutralized authentic SARS-CoV-2 wild type with IC_{50} at 48 nM and 142 nM, respectively. X-ray crystallography revealed two distinctive binding epitopes on the RBD: VH3 E binds to the ACE2 binding site on RBD, possibly block ACE2 binding; VH3 Y binds to RBD will cause steric clash with the ACE2 glycans at N322 and N546, possibly interfere with ACE2 binding. Cryo-EM revealed that the binding of VH3 E trapped the RBDS in the ‘up’ states, leads to the stabilization of spike in conformation with all three RBDS in ‘up’ states, as well as triggering activation of the fusion machinery in spike without host cell contact by ACE2 receptor, resulted in the spike undergo a premature transition from pre-fusion conformation into the non-reversible post-fusion conformation. The non-productive fusion caused the virions to be non-infectious. Engineered bivalentic VH3 E neutralized SARS-CoV-2 pseudovirus and authentic SARS-CoV-2 wild type with IC_{50} of 4.1 nM and 1.32 nM, respectively, displayed up to 50-fold improved neutralization potency compared to its monovalent
form. The simultaneous targeting on two different epitopes by VH VH, in addition with the aberrant activation of the spike fusion machinery as the mechanism of neutralization, suppressed the emergence of escape mutants during experimental evolution.

Xu et al. [149] demonstrated that mice can be engineered to produce cameld VH VHs, known as nanomice. The anti-SARS-CoV-2 RBD nanobodies, Nb12 (K<sub>D</sub> = 30 nM), Nb15 (K<sub>D</sub> = 8.15 nM), Nb30 (K<sub>D</sub> = 6.55 nM) and Nb56 (K<sub>D</sub> = 3.26 nM) were discovered via phage-displayed sdAb libraries generated from nanomice and llama immunized with SARS-CoV-2 RBD and S protein, shown to neutralize SARS-CoV-2 pseudovirus with IC<sub>50</sub> values ranging from 320 pM to 7.145 nM. Structural characterization of sdAbs-RBD complex revealed that Nb15 and Nb56 recognize the RBD-ACE2 interface, possibly neutralize by blocking ACE2 binding; Nb12 and Nb30 recognize a conserved region on RBD without overlapping with the ACE2 binding site, neutralize by stericly interfere with ACE2 binding. The trivalent Fc fusion of Nb12, Nb15, Nb56 and bivalent Fc fusion of Nb30 neutralized SARS-CoV-2 pseudovirus with IC<sub>50</sub> values ranging from 43 to 614 pM, as well as neutralized authentic SARS-CoV-2 wild type and variants B.1.1.7, B.1.351, Pt1 with IC<sub>50</sub> values ranging from 3 pM to 9.374 nM. Moreover, trivalent Nb12-Fc and bivalent Nb30-Fc neutralized SARS-CoV, Bat-SL-CoV (WIV1, WIV1, SCHO141, LYRA11, R5722, RS4084), Bat-CoV-RatG13, Pangolin-CoV-GD and Pangolin-CoV-GX pseudoviruses at IC<sub>50</sub> values below 423 pM. Based on informatics analysis, the binding epitopes for Nb12 and Nb30 are 54% and 79% conserved among sarbecoviruses, respectively, in comparison to the 23% conserved binding epitope for 51 RBD-directed human antibodies on average. The nanobodies have shown good stability profile, with their neutralization activity retained after nebulization, also with their integrity maintained after the heat treatment for 10 min at 98 °C.

2.3. Shark VNAR against SARS-CoV-2 and other coronaviruses

The first study on neutralizing sdAbs from shark origin, VNAR against SARS-CoV-2 was presented by Gauhar et al. [150]. Through the screening of phage-displayed semi-synthetic shark VNAR libraries, followed by further reformatting of the isolated VNARs into bivalent human IgG Fc-fusion, VNAR-hFc antibodies specific to SARS-CoV-2 RBD were obtained: 3ID10_16, 6ID10_75 and 3ID10_99. These antibodies blocked the interaction between ACE2 and wild type RBD at IC<sub>50</sub> values ranging from 2.5 to 130 nM, with neutralizing potential towards the authentic SARS-CoV-2 wild type. The SARS-CoV-2 variants B.1.351 and P.1 harbouring the E484K mutation, which connected to the immune escape from neutralizing antibodies induced by prior infection and SARS-CoV-2 reinfecion [151–153]. Evidently, 3ID10_16, 6ID10_75 and 3ID10_99 possessed the binding ability for two RBD mutants including E484K and N501Y. The discovery of shark VNAR as a novel class of sdAbs against SARS-CoV-2 has expanded the molecular toolkit of potential therapeutics for COVID-19.

Ubah et al. [154] described the identification of monomeric VNARs: 3B4 (K<sub>D</sub> = 17.2 nM) and 2C02 (K<sub>D</sub> = 63 nM) from phage-displayed synthetic shark VNAR library screened against SARS-CoV-2 RBD, to be discovered as the potent neutralizers of authentic SARS-CoV-2 wild type at IC<sub>50</sub> of 11.5 nM and 839 pM, respectively. The VNARs also effectively neutralized WIV1-CoV and SARS-CoV pseudoviruses, with IC<sub>50</sub> ranging from 7.93 to 71.1 nM. Of significance, 3B4 was capable of neutralizing MERS-CoV pseudovirus at IC<sub>50</sub> of 1,050 nM, suggesting that 3B4 bound to conserved region among beta-coronaviruses. Cryo-crystallographic analysis of 3B4 and 2C02 revealed that each VNARs recognizing distinctive epitopes on the RBD, neither of which overlaps with RBD-ACE2 interface. 3B4 binds distal to the ACE2 binding site on the ‘up’-state RBD, neutralizes by resulting in steric clash with ACE2 directly. It was observed that 2C02 can bind to the RBDs in either ‘up’ or ‘down’ state: 2C02 binds to the ‘up’-state RBD without close contact to ACE2, possibly neutralizes by causing allosteric effects towards ACE2 binding; 2C02 also binds within a cleft formed between protomer 1’s ‘down’-state RBD and protomer 3’s NTD, neutralizes by securing RBD in the ‘down’ state to block the access for ACE2. The study features shark VNARs as the useful therapeutic agents for beta-coronaviruses.

According to the work by Feng et al. [155], 20G6 and 17F6 were isolated via phage-displayed shark VNAR library derived from a bamboo shark immunized with SARS-CoV-2 S protein. Structural characterization of VNAR-RBD complex revealed that 20G6 and 17F6 contain “WXGY” motif within the CDR3, that can bind to the residues 365 to 380 of RBD in ‘up’ state without overlapping with the ACE2 binding site, neutralize by causing steric hindrance towards ACE2-RBD interaction. Furthermore, the binding epitopes for 20G6 and 17F6 are highly conserved among sarbecoviruses. The dimeric Fc fusion of 20G6 and 17F6, 20G6-Fc (K<sub>D</sub> < 10 pM) and 17F6-Fc (K<sub>D</sub> < 10 pM) neutralized Pangolin-CoV-GD1 and Bat-CoV-RaTG13 pseudovirus, as well as SARS-CoV-2 pseudovirus wild type and variants B.1.351, B.1.617.2, B.1.617.2.1, B.1.617.1.1, C.37 at IC<sub>50</sub> values below 10 nM. Besides, authentic SARS-CoV-2 wild type and variants B.1.351, B.1.617.2 can be neutralized by 20G6-Fc at IC<sub>50</sub> ranging from 9.36 to 11.79 nM, as well as neutralized by 17F6-Fc at IC<sub>50</sub> ranging from 19.87 to 34.36 nM. Intranasal delivery of 20G6-Fc at 10 mg/kg conferred protection prophylactically and therapeutically, by reducing viral RNA load and lung pathology without significant weight loss in SARS-CoV-2-infected mice and SARS-CoV-2 variant B.1.351-infected mice at 3 dpi. High thermal stability of the VNARs has been proven, with strong binding activity retained after the heat treatment for an hour at 90 °C.

3. Future prospects and conclusions

sdAbs are the ideal building blocks of multivalent constructs; they allow the increasing in valency while retaining small molecular size. The study conducted by Moliner-Morro et al. [114] directed that multivalent constructs for the SARS-CoV-2 neutralizing sdAb can be generated using a combination of sortase-catalysed functionalization and click chemistry. Ty1, as a sdAb isolated from phage-displayed sdAb library derived from an alpaca immunized with SARS-CoV-2-S1-Fc and RBD, bound with high affinity (K<sub>D</sub> = 5-10 nM) to RBD while neutralized SARS-CoV-2 pseudovirus at an IC<sub>50</sub> of 770 ng/ml, in which three Ty1 monomers were required to bind with single trimeric spike protein [102]. With sortase A enzymatic approach to mediate the ligation of click chemistry functional groups (azide and cyclooctyne) site-specifically to the C-terminus of Ty1, followed by oriented assembly of the functionalized sdAbs via strain-promoted azide-alkyne cycloaddition (SPAAAC) or termed Cu-free click chemistry, to create C-to-C terminal Ty1 dimers and polyethylene glycol (PEG) tetramer armed with four Ty1 molecules [158]. In SARS-CoV-2 pseudovirus neutralization assay, Ty1-Ty1 and Ty1-PEG-Ty1 dimers were with similar performances at IC<sub>50</sub> within range of 125 pM. The dimeric Ty1 formulations increased neutralization potency by slightly as compared to Ty1-Fc, and by over 150-fold in comparison with the monomeric Ty1. Intriguingly, the 4-arm PEG-based tetrameric Ty1 construct substantially improved neutralization potency by 1,500-fold to an IC<sub>50</sub> of 13 pM. With an estimation of 30-40 nm linker length between sdAbs on the 4-arm PEG construct, implying that three Ty1 molecules can bind concurrently with one trimeric spike protein, while the fourth Ty1 molecule enables linking between the spike complexes on different virions, leads to the formation into large immune complexes. The main advantage of multi-valency approach as shown by Moliner-Morro et al. [114] over genetic fusion to Fc domain lies in the part on the oriented assembly of sdAbs via click chemistry, which can avoid possibly interfering linkers at the N-terminus near the CDRs of the sdAb. Besides, this approach allows rapid test on different combinations of sdAbs, where the individual cloning and expression are not required. Additionally, the conjugation of antibodies to PEG substantially prolong serum half-life [159,160].

Based on the previous study by Xiang et al. [113], the homo-trimeric construct of Nb21 effectively inhibited SARS-CoV-2 infectivity at pico-molar concentration in vitro. It was noteworthy that Nambulli et al.
translated great in vitro neutralization potency of Nb21 into therapeutic benefits in vivo, by generating homo-trimeric Pittsburgh inhalable Nb21 (PiN-21) whose therapeutic efficacy for SARS-CoV-2 was validated in vivo using animal models of infection. The SARS-CoV-2-infected Syrian hamsters modelled for moderate to severe COVID-19 have shown rapid weight loss up to 16% at 7 dpi. Intranasal delivery of PiN-21 at 0.6 mg/kg eliminated weight loss in SARS-CoV-2-infected hamsters, while having rapid and substantial suppression of viral replication in both upper and lower airways. Infectivity was insignificant in the URT include both nasal washes and throat swabs of PiN-21-treated hamsters at 2 and 4 dpi; while for lower respiratory tract (LRT), the viral titre in lung tissue has been reduced by 10^4-fold at 5 dpi. Aerosol delivery of PiN-21 at 0.2 mg/kg resulted in quick reverse of hamsters’ weight loss after infection, decreased viral titre in lung tissue by 10^6-fold at 3 dpi. According to histopathologic findings in SARS-CoV-2-infected hamsters, there were an abundance of S

Table 2
A summary on the characteristics of previously reported sdAbs against SARS-CoV-2, including binding affinity towards RBD (K_D) and neutralization potency are presented.

| sdAb | Source | CDR3 | K_D towards RBD (nM) | Potential binding epitope class | Potential neutralization mechanism | Potential broad neutralizing target |
|------|--------|------|----------------------|---------------------------------|-----------------------------------|-----------------------------------|
| SARS VHH-72 | Camelid | AGLTVVSEWDDYDDY | 39                  | Class 4                          | Steric interference towards ACE2 binding | SARS-CoV, SARS-CoV-2 D614G mutant, SARS-CoV-2 variants B.1.1.7 and B.1.351 |
| Nb20 | Camelid | RDIEATAEYY | 0.010 (<0.001) | Class 2                          | Steric interference towards ACE2 binding, or conformational control of RBD accessibility | SARS-CoV-2 D614G mutant |
| Nb21 | Camelid | SKDPYGSFPWTRSEFDDY | NA                  | Class 4                          | Lock the helices of S2 in pre-fusion stage to prevent conformational changes for membrane fusion | |
| Nb34 | Camelid | DDKVYVGYTSFPNEYEY | NA                  | Class 4                          | Restrict the flexibility of spike domains | SARS-CoV-2 V367F mutant |
| Nb95 | Camelid | DSDVYDGHSPNNDY | NA                  | Class 4                          | Conformational control of RBD accessibility | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| Nb6 | Camelid | DPASPAPGDY | 41                  | Class 2                          | Block ACE2 binding | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| Nb11-59 | Camelid | APSTTYGGSWVDPGD | 21.6               | Class 1/2                        | Block ACE2 binding | bat-SL-CoV-WIV1, SARS-CoV, SARS-CoV-2 Q321L, V341I, N354D, V367F, K378R, V483A, Y508H, H519P mutants |
| W25 | Camelid | LIKKNELGFLDY | 0.295              | Class 1/2                        | Block ACE2 binding | SARS-CoV-2 D614G mutant |
| WNB 2 | Camelid | IAATYYSGSYYFOCPHDGMDY | 0.36              | Class 1/2                        | Block ACE2 binding | SARS-CoV-2 D614G N501Y mutant |
| WNB 36 | Camelid | IAATYSSGTYYQOCPHYMDY | 0.43              | Class 1/2                        | Steric interference towards ACE2 binding | SARS-CoV-2, SARS-CoV-2 D614G mutant |
| WNB 7 | Camelid | DRLEGSSWPDRFGS | 0.26               | Class 1/2                        | Steric interference towards ACE2 binding | SARS-CoV-2, SARS-CoV-2 D614G mutant |
| WNB 15 | Camelid | DRMEGGSSWPDRFGS | 0.14               | Class 1/2                        | Block ACE2 binding | SARS-CoV-2, SARS-CoV-2 D614G mutant |
| VHH E | Camelid | TVGYTYGSONYHTGCDDMDY | 2                  | Class 1                          | Block ACE2 binding, or the aberrant activation of spike fusion machinery | Suppressed the emergence of resistant escape mutants in evolution experiments |
| VHH V | Camelid | EGSLKGGWGRDFGS | 9                  | Class 4                          | Steric interference towards ACE2 binding | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| Nb12 | Nanomice | AFYFGNSCVLDY | 30                  | Class 4                          | Steric interference towards ACE2 binding | SARS-CoV-2, Bat-SL-CoV (WIV1, WIV16, SH014, LVRa11, Rs7327, Rs40484), Bat-CoV-RaTG13, Pangolin-CoV-GD, Pangolin-CoV-GX, SARS-CoV-2 variants B.1.1.7, B.1.351, P.1 |
| Nb30 | Nanomice | DRGMGYYDFMDY | 6.55               | Class 4                          | Steric interference towards ACE2 binding | SARS-CoV-2 variants B.1.1.7, B.1.351, P.1 |
| Nb15 | Camelid | BPRGGRWDAHHDYNY | 8.15               | Class 1                          | Block ACE2 binding | SARS-CoV-2 variants B.1.1.7, B.1.351, P.1 |
| Nb56 | Camelid | PSYKGDPTSWNTRGDY | 3.26               | Class 1                          | Block ACE2 binding | SARS-CoV-2 E484K, N501Y mutants |
| 3D10_16 | Shark | NA | NA | Class 1/2 | Block ACE2 binding | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| 6D10_75 | Shark | NA | NA | Class 1/2 | Block ACE2 binding | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| 3D10_99 | Shark | NA | NA | Class 1/2 | Steric interference towards ACE2 binding | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| 3B4 | Shark | WSDTSQEPCHAWEQKMWGHV | 17.2             | Class 1                          | Allergenic interference towards ACE2 binding | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| 2C02 | Shark | LINTGKDCRNFHY | 63.0               | Class 3                          | Block ACE2 binding | SARS-CoV-2- SARS-CoV-2 V367F mutant |
| 20G6 | Shark | YSTTGDGERDRCRQGYI | NA                | Class 4                          | Steric interference towards ACE2 binding | Pangolin-CoV-GD1, Bat-CoV-RaTG13, SARS-CoV-2 variants B.1.351, B.1.617.2, B.1.617.2.1, B.1.617.1, G.37 |
| 17F6 | Shark | YSLASMWMCAWMGY | NA | Class 4 | Steric interference towards ACE2 binding | Pangolin-CoV-GD1, Bat-CoV-RaTG13, SARS-CoV-2 variants B.1.351, B.1.617.2, B.1.617.2.1, B.1.617.1, G.37 |

ACE2, angiotensin-converting enzyme 2; CDR, complementarity-determining region; K_D, equilibrium dissociation constant; NA, not applicable; RBD, receptor-binding domain; sdAb, single-domain antibody.
antigen as well as the complete absence of ACE2 within the cytoplasm of bronchiolar epithelium, and the interstitial and peribronchial infiltrates were made up of CD3ε+ T cells and CD68+ macrophages in large quantities. PIN-21 aerosols effectively mitigated the pathology of lung in SARS-CoV-2-infected hamsters, resulted in extremely sparse S antigen with the retention for ACE2 expression on bronchioles, along with minor interstitial and peribronchial mononuclear inflammation due to the declining in numbers of T cell and macrophage immune cell infiltrate. In addition, the fusion of PIN-21 to serum albumin-binding Nb generated a serum-stable construct (PIN-21_4L), with enhanced stability observed in the serum. PIN-21 aerosols treatment can provide cost-effective and more convenient drug administration, especially for mild COVID-19 patients constituted the majority in infected populations.

In conclusion, single-domain antibodies are presently developing into versatile research tools and cost-effective therapeutics targeting for SARS-CoV-2. Broadly neutralizing effect can be achieved by antibodies with diverse epitope engagement followed by potentially different neutralization mechanisms, contributes to the efficient blocking of SARS-CoV-2 mutational escape. The potent neutralizing activity shown by an antibody, in combination with its broad neutralizing ability towards SARS-CoV-2 mutants while possessing good developability profile provide a strong foundation as COVID-19 therapeutic agent.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

No data was used for the research described in the article.

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References

[1] P. Zhou, et al., A pneumonia outbreak associated with a new coronavirus of probable bat origin, Nature 579 (7798) (2020) 270–272.
[2] V.D. Menachery, et al., A SARS-like cluster of circulating bat coronaviruses shows potential for human emergence, Nat. Med. 21 (12) (2015) 1508–1513.
[3] C. Ronco, T. Reis, F. Husain-Syed, Management of acute kidney injury in patients with COVID-19, Lancet Respir. Med. 8 (7) (2020) 738–742.
[4] J. Liu, et al., Hydroxychloroquine, a less toxic derivative of chloroquine, is effective in inhibiting SARS-CoV-2 infection in vitro, Cell Discov. 6 (1) (2020) 1–4.
[5] P. Gautret, et al., Hydroxychloroquine and azithromycin as a treatment of COVID-19: results of an open-label non-randomized clinical trial, Int. J. Antimicrob. Agents 56 (1) (2020), 105949.
[6] M.A. Martinez, Clinical trials of repurposed antivirals for SARS-CoV-2, Antimicrob. Agents Chemother. 64 (9) (2020) e01101–e01120.
[7] B. Cao, et al., A trial of lopinavir-ritonavir in adults hospitalized with severe COVID-19, N. Engl. J. Med. 382 (19) (2020) 1787–1799.
[8] J. Parodi, et al., The journey of remdesivir: from Ebola to COVID-19, Drugs Context 2020 9, https://doi.org/10.1073/pjci.2020.4.14.
[9] J.H. Beigel, et al., Remdesivir for the treatment of COVID-19—preliminary report, N. Engl. J. Med. 383 (19) (2020) 1813–1826.
[10] Y. Wang, et al., Remdesivir in adults with severe COVID-19: a randomised, double-blind, placebo-controlled, multicentre trial, Lancet N. Am. Ed 395 (10262) (2020) 1569–1578.
[11] H. Ledford, D. Cyranoski, R. Van Noorden, The UK has approved a COVID vaccine—here’s what scientists now want to know, Nature 588 (7837) (2020) 205–206.
[12] O. Sharma, et al., A review of the progress and challenges of developing a vaccine for COVID-19, Front. Immunol. 11 (2020) 2413.
Biomedical Engineering Advances 4 (2022) 100054

H.T. Lim et al.

[68] X. Chi, et al., Humanized single domain antibodies neutralize SARS-CoV-2 by
[65] Y. Watanabe, et al., Site-specific glycan analysis of the SARS-CoV-2 spike, Science
[63] J. Lan, et al., Structure of the SARS-CoV-2 spike receptor-binding domain bound
[61] W. Li, et al., Angiotensin-converting enzyme 2 is a functional receptor for the
[69] C. Newcombe, A.R. Newcombe, Antibody production: polyclonal-derived
[58] G. Zhou, Q. Zhao, Perspectives on therapeutic neutralizing antibodies against the
[57] S.J. Zost, et al., Potently neutralizing and protective human antibodies against
[56] B. Ju, et al., Human neutralizing antibodies elicited by SARS-CoV-2 infection, Nature
[55] W. Li, et al., Antibodies: the future of antibody-based therapies, Cancer Today. 8 (2) (2012) 57–62.
[54] S. Cunningham, et al., Nebulised ALX-0171 for respiratory syncytial virus lower respiratory tract infection in hospitalised children: a double-blind, randomised, placebo-controlled trial, Lancet Infect. Dis. 14 (11) (2014) 1017–1024.
[53] J. Steven, et al., In vitro maturation of a humanized shark VNAR domain to humanize and structural analysis, J. Biol. Chem. 288 (24) (2013) 17054–17064.
[52] M. Dumoulin, et al., Single-domain antibody fragments with high conformational stability, Protein Sci. 14 (9) (2005) 1624–1636.
[51] J. Holland, et al., Rapid evolution of RNA genomes, Science 215 (4540) (1982) 1577–1585.
[50] V. Papanikolaou, et al., From Delta to Omicron: S1-RBD/S2 mutation/deletion equilibrium in SARS-CoV-2 defined variants, Gene 814 (2022) 146134.
[49] CDC. Science brief: Omicron (B.1.1.529) variant. 2021 15 January 2022; Available from: https://www.cdc.gov/coronavirus/2019-ncov/science/science-briefs/science-brief-omicron-variant.html.
[48] S. Cherian, et al., SARS-CoV-2 spike mutations, L452R, T478K, E484Q and S477N and MERS to 2019-nCoV, J. Microbiol. Immunol. Infect. 53 (3) (2020) 351–254.
[47] C. Drosten, et al., Identification of a novel coronavirus in patients with severe acute respiratory syndrome, Nature 426 (6965) (2003) 450–454.
[46] N.-S. Zhong, G.W. Wong, Epidemicology of severe acute respiratory syndrome (SARS): adults and children, Paediatr. Respir. 5 (4) (2004) 270–274.
[45] WHO. Middle East respiratory syndrome. 2022 5 September 2022; Available from: http://www.who.int/health-topics/mers-cov/mers-outbreaks.html.
[44] Z. Wu, et al., The unique features of SARS-CoV-2 transmission: comparison with SARS-CoV, MERS-CoV and 2009 H1N1 pandemic influenza virus, Rev. Med. Virol. 31 (2) (2021) e217.
[43] R. Wölfel, et al., Virological assessment of hospitalized patients with COVID-2019, Nature 581 (7809) (2020) 465–469.
[42] P.-I. Lee, P.-R. Hsieh, Emerging threats from zoonotic coronaviruses-from SARS and MERS to 2019-nCoV, J. Microbiol. Infect. 53 (3) (2020) 365.
[41] J. Hicks, et al., Serologic cross-reactivity of SARS-CoV-2 with endemic and seasonal betacoronaviruses, J. Clin. Immunol. 41 (5) (2021) 906–913.
[40] S. Cherian, et al., SARS-CoV-2 spike mutations, L452R, T478K, E484Q and 20-00389, 2020.
[39] B. Beutin, et al., Identification of a novel coronavirus in patients with severe acute respiratory syndrome, Nat. Engl. J. Med. 348 (20) (2003) 1967–1976.
[38] V. Xiang, et al., Versatile and multivalent nanobodies efficiently neutralize SARS-CoV-2. Nat. Commun. 11 (1) (2020) 1.
[37] Y. Xiang, et al., Atypical antigen recognition mode of a shark non-structural protein 3, Nat. Immunol. 14 (10) (2013) 1052–1059.
[36] O.V. Kovalenko, et al., General strategy to humanize a camelid single-domain antibody targeting the spike receptor binding domain, Nat. Commun. 11 (1) (2020) 1.
[35] C. Vincke, et al., General strategy to humanize a camelid single-domain antibody targeting the spike receptor binding domain, J. Biol. Chem. 288 (24) (2013) 17054–17064.
[34] R. Shi, et al., A human neutralizing antibody targets the receptor-binding site of SARS-CoV-2, Nature 584 (7819) (2020) 115–120.
[33] O.C. Ubah, et al., Novel, anti-hTNF-α antibodies with enhanced neutralizing potency and multifunctionality, generated for therapeutic development, Front. Immunol. 8 (2017) 1780.
[32] L. Hanke, et al., An alpaca nanobody neutralizes SARS-CoV-2 by blocking receptor interaction, Nat. Commun. 11 (1) (2020) 1–9.
[31] P. Banas, J. Hambach, F. Koch-Nolte, Nanobodies and nanobody-based human heavy chain antibodies as antimitor therapeutics, Front. Immunol. 8 (2017) 1603.
[30] J.Y. Lee, et al., Effectiveness of regdanvimab treatment in high-risk COVID-19 patients to prevent progression to severe disease, Front. Immunol. 12 (2021) 436.
[29] J.W. Burger, W.A. Hess, Function of the rectal gland in the spiny dogfish, Science 131 (3401) (1660) 670–671.
[28] C. Vincke, et al., General strategy to humanize a camelid single-domain antibody and identification of a universal humanized nanobody scaffold, J. Biol. Chem. 284 (5) (2009) 3273–3284.
[27] O.V. Kovalenko, et al., Atypical antigen recognition mode of a shark immunoglobulin new antigen receptor (iNAR) variable domain characterized by humanization and structural analysis, J. Biol. Chem. 288 (24) (201) 17408–17419.
[26] S. Chaturvedi, Counting the cost of caplacizumab, Blood J. Am. Soc. Hematol. 137 (13) (2021) 1395–1396.
[25] A. Hofmeijer, et al., Nanobody- and VHH-based imaging of SARS-CoV-2 spike protein and viral nucleocapsid protein, Viruses 13 (5) (2021) 1331–1346.
[24] Y. Xiang, et al., Versatile and multivalent nanobodies efficiently neutralize SARS-CoV-2, J. Biol. Chem. 288 (24) (2013) 17054–17064.
[23] S.A. Godakova, et al., Camelid VHHs fused to human Fc fragments provide long-acting therapeutic nanobody for the treatment of respiratory syncytial virus infection, Biochem. 82 (2013) 775–783.
[22] S. Wadell, et al., Virological assessment of hospitalized patients with COVID-2019, N. Engl. J. Med. 384 (3) (2021) 238–248.
[21] J. Steven, et al., In vitro maturation of a humanized shark VNAR domain to exhibit superior penetration and treatment of solid tumors, Mol. Ther. 30 (3) (2022) 273–278.
[20] N. V. Bathula, H. Bommadevara, J. M. Hayes, Nanobodies: the future of antibody-based immune therapies, Cancer Ther. 19 (10) (2020) 2208–2215.
[120] T.F. Custodio, et al., Selection, biophysical and structural analysis of synthetic nanobodies that effectively neutralize SARS-CoV-2, Nat. Commun. 11 (1) (2020) 1–11.

[121] M. Schoof, et al., An ultrapotent synthetic nanobody neutralizes SARS-CoV-2 by stabilizing inactive spike, Science 370 (6523) (2020) 1473–1479.

[122] G.V. Nieto, et al., Potent neutralization of clinical isolates of SARS-CoV-2 D614 and G614 variants by a monoclonal, sub-nanomolar affinity nanobody, Sci. Rep. 11 (1) (2021) 1–14.

[123] J. McCafferty, et al., Phage antibodies: filamentous phage displaying antibody variable domains, Nature 348 (6301) (1990) 552–554.

[124] A. Klarenbeek, et al., Camelid Ig V Genes Reveal Significant Human Homology Not Seen in Therapeutic Target Genes, Providing for a Powerful Therapeutic Antibody Platform. In Mabs 7 (4), Taylor & Francis, 2015, pp. 693–706.

[125] C.L. Nigro, et al., NK-mediated antibody-dependent cell-mediated cytotoxicity in COVID-19, Nat. Rev. Immunol. 20 (6) (2020) 339–341.

[126] M.Z. Tay, K. Wiehe, J. Pollara, Antibody-dependent cellular phagocytosis in syncytial virus infection and disease, Front. Immunol. 10 (2019) 352.

[127] A. Iwasaki, Y. Yang, The potential danger of suboptimal antibody responses in antiviral immune responses, Front. Immunol. 10 (2019) 352.

[128] C. Chakraborty, et al., D614G Mutation Eventuates in all VOI and VOC in SARS-CoV-2, Infect. Genet. Evol. 81 (2021) 1667–1670.

[129] C.L. Nigro, et al., NK-mediated antibody-dependent cell-mediated cytotoxicity in solid tumors: biological evidence and clinical perspectives, Ann. Transl. Med. 7 (5) (2019).

[130] M.Z. Tay, K. Wiehe, J. Pollara, Antibody-dependent cellular phagocytosis in antiviral immune responses, Front. Immunol. 10 (2019) 352.

[131] B. Luan, H. Wang, T. Huynh, Enhanced binding of the N501Y-mutated SARS-CoV-2 spike protein to the human ACE2 receptor: insights from molecular dynamics simulations, FEBS Lett. 595 (10) (2021) 1454–1461.

[132] J.A. Plante, et al., Spike mutation D614G alters SARS-CoV-2 fitness, Nature 592 (7852) (2021) 116–119.

[133] E.A. Van Erp, et al., Fc-mediated antibody effector functions during respiratory syncytial virus infection and disease, Front. Immunol. 10 (2019) 548.

[134] B. Korber, et al., Tracking changes in SARS-CoV-2 spike: evidence that D614G increases infectivity of the COVID-19 virus, Cell 182 (4) (2020) 812–827, e149.

[135] P.-A. Koenig, et al., Structure-guided multivalent nanobodies block SARS-CoV-2 infection and suppress mutational escape, Science 371 (6530) (2021).

[136] J. Xu, et al., Nanobodies from camelid mice and llamas neutralize SARS-CoV-2 variants, Nature 595 (7866) (2021) 278–282.

[137] C. Klimstra, et al., D614G Mutation Eventuates in all VOI and VOC in SARS-CoV-2: is it Part of the Positive Selection Pioneered by Darwin? 26 Elsevier, 2021, pp. 827, e19.

[138] J. Xu, et al., Nanobodies from camelid mice and llamas neutralize SARS-CoV-2 variants, Nature 595 (7866) (2021) 278–282.

[139] A. Gauhar, et al., Single domain shark VNAR antibodies neutralize SARS-CoV-2 infection in vitro, FASEB J. 35 (11) (2021) e21970.

[140] Wise, J., COVID-19: the E484K mutation and the risks it poses. 2021, British Medical Journal Publishing Group.