Epitope mapping of neutralising anti-SARS-CoV-2 monoclonal antibodies: Implications for immunotherapy and vaccine design

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Abstract
Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is the causative agent of the coronavirus disease 2019 (COVID-19) pandemic. This disease has currently affected more than 346 million people and resulted in more than 5.5 million deaths in many countries. Neutralising monoclonal antibodies (MAbs) against the SARS-CoV-2 virus could serve as prophylactic/therapeutic agents in COVID-19 infection by providing passive protection against the virus in individuals. Until now, no Food and Drug Administration/European Medicines Agency-approved neutralising MAb against SARS-CoV-2 virus exists in the market, though a number of MAbs have been authorised for emergency use. Therefore, there is an urgent need for development of efficient anti-SARS-CoV-2 neutralising MAbs for use in the clinic. Moreover, neutralising anti-SARS-CoV-2 MAbs could be used as beneficial tools for designing epitope-based vaccines against the virus. Given that the target epitope of a MAb is a crucial feature influencing its neutralising potency, target epitopes of neutralising anti-SARS-CoV-2 MAbs already reported in the literature and reactivity of these MAbs with SARS-CoV-2 variants are reviewed herein.

KEYWORDS
COVID-19, epitope mapping, immunotherapy, neutralising monoclonal antibody, RBD, SARS-CoV-2 virus

1 | SARS-CoV-2 VIRUS AND CORONAVIRUS DISEASE 2019 INFECTION

Coronavirus disease 2019 (COVID-19), caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), was first identified in Wuhan, China in December 2019.¹ Until 23rd of January 2022, the disease affected more than 346 million people and resulted in more than 5.5 million deaths in many countries.² Currently, there are several effective drugs licenced based on the results of randomized clinical trials. The Food and Drug Administration (FDA) approved antiviral drugs including remdesivir, Paxlovid, and molnupiravir for use in patients 12 years of age and older for the treatment of COVID-19.
COVID-19 requiring hospitalisation. The FDA and the World Health Organization (WHO) recommend several therapeutics for COVID-19 such as IL-6 receptor blockers (tocilizumab or sarilumab) as well as systemic corticosteroids in patients with severe or critical disease.\(^3\)\(^-\)\(^7\) Meanwhile, many efforts have been made to produce SARS-CoV-2 neutralising monoclonal antibodies (MAbs). Sotrovimab, REGN-CoV-2, and the cocktail of bamlanivimab and etesevimab have been authorised for emergency use as post-exposure prophylaxis for COVID-19 in adults and children at high risk for progression to severe COVID-19.\(^8\)\(^-\)\(^10\) Considering the emergence of new variants and lack of efficacy of a number of the neutralising MAbs against the newly emerged Omicron variant, there is imperative need for MAbs able to efficiently cross-neutralise various variants to be used as passive immunotherapy for the control SARS-CoV-2 infection and/or disease severity.

### 1.1 Structure of the SARS-CoV-2 virus

SARS-CoV-2 is a 29,881 bp single-stranded ribonucleic acid (RNA)-enveloped virus containing structural proteins including spike (S), envelope (E), membrane (M), and nucleocapsid (N) as well as non-structural proteins including 3-chymotrypsin-like protease, papain-like protease, and RNA-dependent RNA polymerase (Figure 1a). Structural proteins are involved in virus attachment to the host cell membrane and subsequent entry into cells, viral assembly, and release from host cells. Among the structural proteins, the S protein is responsible for the virus binding to the host cell receptors and subsequent entrance into cells. Three other structural proteins including E, M, and N proteins contribute to viral assembly resulting in formation of the virus whole particle. N protein also facilitates packaging of the viral genome into a helical ribonucleocapsid. On the other hand, non-structural proteins contribute to viral genome replication and transcription.\(^1\)

### 1.2 S protein structure and function

The S protein of SARS-CoV-2 virus (1273 amino acids (aa)) is a clover-shaped, type I Transmembrane (TM) protein and contains a large N-terminal extracellular domain (aa: 1–1212), a TM domain (TM; aa: 1213–1237), and a short C-terminal intracellular domain (aa: 1238–1273). S Protein consists of a signal peptide (aa: 1–13), S1 subunit (aa: 14–685), and an S2 subunit (aa: 686–1273).\(^1\)\(^2\) Moreover, the S1 subunit, responsible for binding the virus to host cell receptors, is composed of the N-terminal domain (NTD; aa: 18–305), the C-terminal receptor-binding domain (RBD; aa: 329–528), subdomain-1 (SD1; aa: 529–589), and subdomain-2 (SD2; aa: 590–686; Figure 1b).\(^1\)\(^2\) The Receptor binding domain (RBD) is composed of two sub-domains including core sub-domain composed of a β-sheet with 5 anti-parallel strands (β1, β2, β3, β4, and β7) in the inner side of the S protein and receptor-binding motif (RBM) from the outer side that extends from the core sub-domain and consists of β5 and β6 strands.\(^1\)\(^3\)\(^,\)\(^1\)\(^4\)

Angiotensin-converting enzyme 2 (ACE2), a membrane-bound zinc-containing enzyme expressed on many tissues including lungs, arteries, heart, kidneys, and intestines, has been demonstrated to be the main receptor for virus attachment to target cells.\(^1\)\(^5\) Moreover, transmembrane protease serine 2 (TMPRSS2) has been proposed for S protein priming.\(^1\)\(^6\) The RBM sub-domain of RBD, which forms a concave surface accommodating the N-terminal α-helix of the ACE2, is responsible for the virus binding to the ACE2.\(^1\)\(^2\) The SARS-CoV virus also employs a similar mechanism to bind to host cells. On the other hand, the S2 subunit mediates fusion of the viral membrane with the host cell membrane allowing virus entry into target cells.\(^1\)\(^2\) The S2 subunit consists of upstream helix (UH; aa: 687–819), N-terminal fusion peptide (FP; aa: 820–846), hepta peptide repeat sequence one (HR1; aa: 912–985), SD3 (aa: 1072–1139), stem helix (SH; aa: 1139–1163), HR2 (aa: 1163–1212), TM domain (aa: 1213–1237), and intracellular domain (aa: 1238–1273; Figure 1b).\(^1\)\(^2,\)\(^1\)\(^7\)

The S protein possesses two distinct conformational states including prefusion and postfusion conformations. The prefusion state of the S protein, composed of three S1 subunits and three S2 subunits, exists in two conformations: (1) a closed conformation in which all three protomers of RBDs are hidden and thus preventing RBD-ACE2 interaction (down conformation of RBD or receptor inaccessible state), (2) an open conformation in which one protomer of RBD is exposed allowing for RBD-ACE2 interaction (up conformation of RBD or receptor accessible state; Figure 1c). Indeed, the up conformation of RBD provides the surface required for RBD interaction with the ACE2.\(^1\)\(^8\)\(^,\)\(^1\)\(^9\) Upon RBD binding to the ACE2, a conformational change in S protein structure occurs allowing for proteolytic cleavage of the protein at the S1-S2 boundary by host proteases. This converts S protein from the inactive prefusion state into the active postfusion state resulting in fusion of the viral membrane with the host cell membrane and entrance into the cell.\(^2\)\(^0\) Though TMPRSS2 has been proven as the protease responsible for cleavage of the S protein, other host proteases such as trypsin were also recognized for their function in cleavage of the S protein.\(^2\)\(^1\) Serum levels of antibodies specific for the spike RBD increase during the two weeks after onset of symptom. Higher levels of RBD-specific IgM have been shown in deceased COVID-19 patients rather than recovered patients. Also, a significant correlation was reported between RBD-specific IgG and IgM in both groups of patients.\(^2\)\(^2\)\(^,\)\(^2\)\(^3\) Furthermore, several studies found a positive correlation between serum neutralising capacity and disease severity in recovered patients with a wide range of disease severity (severe, moderate, mild, and asymptomatic).\(^2\)\(^4\)\(^,\)\(^2\)\(^7\)

## 2 NEUTRALISING ANTIBODIES

Passive protection against microbial agents, including viruses, could be achieved by neutralising antibodies (nAb). Neutralising antibodies bind to microbial structures responsible for binding to target cell receptors through their variable regions and consequently prevent
neutralise toxins of viral infection; and prevent entry of the virus.

Neutralising antibodies are a polyclonal pool of antibodies composed of a mixture of antibody clones recognising different epitopes of the corresponding antigen. These antibodies are prepared from human plasma of hyper-immunised individuals or convalescent patients. However, plasmas from infected patients with SARS-CoV-2 or convalescent patients have been shown to contain neutralising anti-SARS-CoV-2 antibodies with varying neutralisation capacity levels. The nAb titre required for prevention of COVID-19 infection in humans has not been determined yet. Using a non-human primate model of COVID-19 infection, prevention of clinical signs of the disease and reduced viral loads in bronchoalveolar lavage and nasal mucosa could be observed in re-infected NHPs compared with post-primary infection. The serum samples could potently neutralise the SARS-CoV-2 pseudovirus at a titre of 1:100 (ranging from 1:83-1:197) and the authentic SARS-CoV-2 virus at a titre of 1:35-1:326 on day 35 after rechallenge.

Polyclonal preparations of the antibodies have several limitations including insufficient level of neutralising potency in donor plasma, rapid decline of nAbs in convalescent patients, lot-to-lot variability, and lack of plasma donors, and possibility of transmission of microbial agents and adverse reactions to plasma proteins. Interestingly, neutralising MAbs targeting microbial antigens including COVID-19 lack these limitations and could therefore be considered as prophylactic/therapeutic alternative for the passive immunotherapy. Until now, no FDA/European Medicines Agency (EMA)-approved neutralising MAb for COVID-19 infection has entered in clinic, although a few number of the MAbs have been authorised for emergency use. Therefore, there is an urgent need for development of efficient neutralising anti-SARS-CoV-2 MAbs. Given that the target epitope of a MAb is a crucial feature influencing its neutralising potency, herein, epitope specificity of the neutralising antibody MAbs already reported in the literature is delineated and discussed.

3 | EPITOPE MAPPING OF NEUTRALISING ANTI-SARS-CoV-2 MAbs

Identification of the target epitope of an antibody molecule is instrumental in development of effective prophylactic therapeutics and epitope-based vaccines as well as molecular elucidation of MAb neutralising activities. In a recent study, we performed epitope mapping of RBD in COVID-19 patients' sera using a panel of linear
FIGURE 2 Prevention of virus binding to its receptor on target cell by neutralising antibodies. Binding of (a) non-neutralising antibodies and (b) neutralising antibodies to spike protein. ACE2: Angiotensin-converting enzyme two.

epitopes spanning RBD. Our results demonstrated involvement of mostly conformational disulfide bond-dependent epitopes in RBD-specific IgG antibody. Review of literature for neutralising anti-SARS-CoV-2 MAbS indicates that all reported nAbs recognise epitopes within only the S protein and most of them are directed against the RBD (Figure 3, Table 1).

3.1 Neutralising monoclonal antibodies recognising the Receptor binding domain fragment of the S protein

Given the crucial role of the RBD fragment of the S protein in binding of the virus to its receptor on target cells, it is not surprising that a major proportion of neutralising anti-SARS-CoV-2 antibodies are directed against RBD (Figure 3, Table 1). Analysis of MAbS isolated from 25 COVID-19-infected patients showed that a majority of the nAbs recognized the S1 subunit of the virus. Removal of anti-RBD antibodies from sera of patients abolished their neutralising activity, highlighting the dependency of the neutralising activity to anti-RBD antibodies. Accordingly, analysis of anti-SARS-CoV-2 MAbS for their neutralising activities against the virus showed that a large number of neutralising MAbS (67/70) recognise the RBD fragment. In line with these findings, none of the non-RBD-binding MAbS showed neutralising activities in a different study. Therefore, RBD seems to be the most crucial domain of the S protein for eliciting nAbS against the virus.

3.1.1 Neutralising anti-RBD monoclonal antibodies that interfere with RBD-ACE2 interaction

Considering that the up conformation of RBD provides the surface required for RBD interaction with the ACE2, it is assumed that neutralising anti-RBD MAbS should recognise S protein in the up conformation. Barnes et al. classified neutralising anti-RBD MAbs into 5 groups: (1) MAbs that recognise the up conformation of the S protein and prevent RBD-ACE2 interaction, (2) MAbs that recognise both up and down conformations of the S protein and prevent RBD-ACE2 interaction, (3) MAbs that recognise the up conformation of the S protein, but the epitopes are located outside of the ACE2-binding site of RBD, (4) MAbs that recognise both up and down conformations of the S protein and do not bind to the ACE2-binding site of RBD, (5) MAbs that recognise the down conformation of the S protein and prevent RBD-ACE2 interaction. Therefore, majority of neutralising anti-RBD MAbs (4/5 groups) could recognise S protein in the up conformation. However, antibodies from group 5 unexpectedly bind to RBD epitopes that are solely available on the down conformation of the S protein. In efforts to explore neutralising mechanisms of these MAbs, the authors found MAbs binding to RBD epitopes on the down conformation locked S protein in the down conformation and consequently prevented accessibility of the ACE2-binding surface of RBD to the ACE2. Liu et al. reported a similar finding. They isolated a neutralising anti-RBD MAb (MAb 2-4) that bound to S protein in the down conformation and locked the protein in the receptor inaccessible state.

Robbiani et al. showed 54% of RBD-binding MAbs neutralised the virus. In accordance with this result, 46% of anti-RBD MAbs isolated by Kreye et al. showed neutralising activity. We have recently generated a panel of mouse MAbs against RBD and observed that less than half of these MAbs display neutralising activity in pseudovirus-based neutralising assays, suggesting that recognising RBD is not necessarily sufficient for virus neutralisation (unpublished data). These findings indicate that RBD contains potent neutralising epitopes even if not all RBD epitopes contribute to virus neutralisation. Robbiani et al. identified three distinct neutralising epitopes on RBD including C144 and C101 in group 1; C121 and C119 in group 2 and C135 in group 3. They showed that groups 1 and 2 antibodies could bind to the RBD immunocomplexed with group 3 antibodies. Of note, groups 1 and 2 displayed different properties in binding specificity, so that group 1 could bind to the RBD immunocomplexed with group 2, but not vice versa. Rogers et al. have also identified three distinct neutralising epitopes on RBD: the most potent neutralising MAbs were found to recognise the RBD-A epitope. Further analysis showed that the RBD-A epitope spans the ACE2-binding site in RBD. These RBD-A specific MAbs also efficiently neutralised the virus when administered prophylactically (antibody administration before virus challenge) in a Syrian hamster.
animal model of COVID-19 infection. Consistently, MAbs inhibiting RBD binding to ACE2 displayed the strongest neutralising activity. These MAbs also revealed in vivo efficacy when administered either prophylactically or therapeutically (the antibody administration after the virus challenge). BD23, another nAb in their panel that bound the "down" conformation of RBD also competed with ACE2. Moreover, the MAbs that interrupted RBD-ACE2 interaction imposed neutralising activity. Among the panel of human neutralising MAbs targeting the SARS-CoV-2 RBD isolated from patients at the acute phase, a subset of them inhibited binding to the human ACE2. Also, a large number of neutralising anti-RBD MAbs obtained by Zost et al. interfered with RBD-ACE2 interaction. The neutralising anti-RBD MAb, rRBD-15, inhibited binding of RBD to ACE2. Neutralising anti-RBD MAb LY-CoV555, that prevented RBD-ACE2 interaction, was successfully used in a phase two clinical trial conducted on outpatients with mild or moderate COVID-19 disease. A single dose (2800 mg) administration of this MAb, also known as bamlanivimab, significantly improved clinical outcomes in patients by reducing severity of symptoms and viral load. However, bamlanivimab was revoked by FDA because of increased risk for treatment failure due to continued development of SARS-CoV-2 escape variants. Altogether, these findings indicate that the most potent neutralising epitopes are the epitopes involved in RBD binding to the ACE2. Based on this notion, Liu et al. used an innovative approach to isolate neutralising anti-SARS-CoV-2 MAbs. In this approach, they initially isolated MAbs based on positive selection for RBD followed by negative selection of the isolated MAbs for a mutant RBD in which RBD residues that contribute to ACE2 binding were deleted. Indeed, these selections ensured isolation of neutralising MAbs including 4A2, 4A12, 4D5, and 4A10 accurately recognising RBD epitopes involved in ACE2 binding.

Cryo-electron microscopy studies showed that RBD interact with ACE2 through hydrogen and ionic bonds. Residues A475, N487, E484, and Y453 in RBD interact with residues S19, Q24, K31, and H34 of ACE2, respectively. Moreover, the residues Q498, T500, and N501 form hydrogen bonds of RBD interact with Y41, Q42, K353, and R357 of ACE2. In another study, RBD residues of Y505, Y449, G496, F497, and G502 bound to ACE2 residues including E37, D38, D39, K353, and G354, respectively. An ionic bond between P491 of RBD with K31 of ACE2 also contributed to RBD-ACE2 interaction. The other RBD residues, including T470, F486, Y489, and Q493, were also identified as crucial residues of RBD for interaction with the ACE2. Furthermore, the SARS-CoV-2 RBM provides a larger and more favourable contact interface with ACE2 in comparison to the SARS-CoV RBM. In sum, RBD residues, including Y449, Y453, L455, F456, T470, A475, E484, F486, N487, Y489, F490, P491, Q493, G496, F497, Q498, T500, N501, G502 and Y505, might be considered as ACE2-interacting residues of RBD, and all are located in SARS-CoV-2 RBM spanning from residue 438-506 of the S sequence (Figure 4).
| Ref. No. | MAb(s) designation | Target epitope | Inhibition of RBD-ACE2 interaction | In vivo neutralisation activity | Affinity constant (nM) | IGHV and IGLV genes usage |
|----------|---------------------|----------------|------------------------------------|--------------------------------|------------------------|--------------------------|
| 35       | 4A2, 4A12, 4D5, 4A10 | RBD            | Yes                                | NI                             | 1.03–5.82              | NI                       |
| 32       | C101, C119, C121, C135, C145 | RBD            | NI                                 | NI                             | NI                     | IGHV: Multiple IGHVs     |
|          |                     |                |                                    |                                |                        | IGLV: Multiple IGLVs     |
| 33       | -                   | RBD            | Yes                                | Prophylactic efficacy in syrian hamster (4.2 mg/kg) | NI                     | IGHV: IGHV1, IGHV3      |
|          |                     |                |                                    |                                |                        | IGLV: NI                 |
| 45       | -                   | RBD            | NI                                 | NI                             | NI                     | IGHV: Multiple IGHVs     |
|          |                     |                |                                    |                                |                        | IGLV: Multiple IGLVs     |
| 44       | -                   | RBD            | NI                                 | NI                             | NI                     | NI                       |
| 46       | BD23                | RBD            | Yes                                | Prophylactic efficacy in hACE2 transgenic mice (20 mg/kg) | <15.9                  | NI                       |
|          |                     |                |                                    | Therapeutic efficacy in hACE2 transgenic mice (20 mg/kg) |                        |                         |
| 51       | COV2-2196, COV2-2130 | RBD            | Yes                                | Prophylactic efficacy in hACE2 transgenic mice (10 mg/kg) | NI                     | NI                       |
| 52       | rRBD-15             | RBD            | Yes                                | NI                             | NI                     | NI                       |
| 49       | CV07-209, CV07-250  | RBD            | Yes                                | Prophylactic efficacy in syrian hamster (18 mg/kg) | 0.006–1.1              | IGHV: IGHV1-2, IGHV3-53, IGHV3-66 |
|          |                     |                |                                    | Therapeutic efficacy in syrian hamster (18 mg/kg) |                        | IGLV: IGVK1–33, IGVK2–14 |
| 60       | B38                 | RBD            | Yes                                | Therapeutic efficacy in hACE2 transgenic mice (25 mg/kg) | 1–100                  | IGHV: IGHV1, IGHV3      |
|          |                     |                |                                    |                                |                        | IGLV: IGVK1, IGVK2, IGVK3 |
| 67       | CB6                 | RBD            | Yes                                | Prophylactic efficacy in rhesus macaques monkey (50 mg/kg) | 249–68                 | NI                       |
|          |                     |                |                                    | Therapeutic efficacy in rhesus macaques monkey (50 mg/kg) |                        |                         |
| Ref. No. | Ref. No. | MAb(s) designation | Target epitope | Inhibition of RBD-ACE2 interaction | In vivo neutralisation activity | Affinity constant (nM) | IGHV and IGLV genes usage |
|---------|----------|---------------------|----------------|----------------------------------|-------------------------------|-----------------------|-----------------------------|
| 65 | MW05 | Yes | Prophylactic efficacy in rhesus macaques monkey (40 mg/kg) | 0.40–0.46 NI | | |
| 61 | P2B-2F6 | Yes | NI | 5.14 NI | |
| 70 | 47D11 | No | NI | NI | | |
| 71 | S309 | No (Fc-dependent effector functions, including ADCC and ADCP) | NI | NI | | |
| 63, 110 | REGN10933, REGN10987 | Yes | Prophylactic efficacy in rhesus macaques (25, 50 and 150 mg/kg) | NI | IGHV: IGHV3-53, IGHV3-66, IGHV2-70 | |
| | | Fc-dependent effector functions, including ADCC and ADCP | | | IGLV: IGKV1-9, IGKV1-33, IGKV1-39 | |
| 54, 53 | LY-CoV555 | Yes | Prophylactic efficacy in rhesus macaques (25 mg/kg) | NI | NI | |
| | | | Therapeutic efficacy in outpatients with mild or moderate COVID-19 disease (2800 mg) | | | |
| 111 | - | Yes | NI | NI | | |
| 112 | - | Yes | NI | NI | IGHV: IGKV3-53*01 | |
| 34 | MAb 2–4 | Yes | Prophylactic efficacy in syrian hamster (15 mg/kg) | NI | IGHV: IGHV3-30, IGKV3-20 | |
| Ref. No. | MAb(s) designation | Target epitope | Inhibition of RBD-ACE2 interaction | In vivo neutralisation activity | Affinity constant (nM) | IGHV and IGLV genes usage |
|----------|---------------------|----------------|----------------------------------|--------------------------------|----------------------|--------------------------|
| 50       | -                   | RBD            | Yes                              | NI                             | 1.8–15.6             | IGHV3-64: IGKV1-39       |
|          |                     |                |                                  |                               |                      | IGHV3-53: IGKV3-20      |
|          |                     |                |                                  |                               |                      | IGHV3-53: IGKV1-12      |
|          |                     |                |                                  |                               |                      | IGHV3-66: IGKV3-20      |
|          |                     |                |                                  |                               |                      | IGHV3-23: IGLV3-21      |
| 59       | CTP59               | RBD            | Yes                              | Therapeutic efficacy in ferret, hamster, and rhesus monkey (3 and 30 mg/kg) | 0.027                | IGHV2-70                |
| 77       | COV2-2676, COV2-4489| NTD            | Yes                              | Prophylactic and therapeutic efficacy in heterozygous K18-hACE2 c57BL/6J mice (10 mg/kg) | NI                   | IGHV4-39                |
|          |                     |                |                                  |                               |                      | IGKV1-69                |
| 33       | -                   | Non-RBD regions of S protein | NI                               | NI                             | NI                   | NI                       |
| 74       | 4A8                 | Non-RBD regions of S1 subunit | NI                               | NI                             | 92.7                 | NI                       |

Abbreviations: ACE2: Angiotensin-converting enzyme 2, ADCC: Antibody-dependent cellular cytotoxicity, ADCP: Antibody-dependent cellular phagocytosis, IGHV: Immunoglobulin heavy chain variable region, IGLV: Immunoglobulin light chain variable region, NI: Not identified, nM: Nanomolar, NTD: N-terminal domain, RBD: Receptor-binding domain, S: spike.
Figure 4 Assignment of epitope residues of spike protein for binding to Angiotensin-converting enzyme 2 (ACE2) and SARS-CoV-2 neutralising monoclonal antibodies (MAbs). Epitope residues of (a) Receptor binding domain (RBD), (b) N-terminal domain (NTD), (c) Furin cleavage site, and (d) SH-targeting neutralising MAbs are highlighted in different colours. ACE2: Angiotensin-converting enzyme 2, SARS-CoV-2: Severe acute respiratory syndrome coronavirus 2; NTD: N-terminal domain; RBD: Receptor binding domain; SH: Stem helix.
Crystallographic analysis showed that neutralising anti-SARS-CoV-2 MAbs interfere with RBD-ACE2 interaction to different extents. In a panel of neutralising MAbs, the two highest neutralising MAbs, CV07–209 and CV07–250, highly interfered with RBD-ACE2 interaction. The RBD target epitope of MAb CV07–250 completely overlapped with the ACE2-binding site. On the contrary, CV07–270 binds from a different angle with partial overlapping with the ACE2 binding site. Neutralising MAb CTP59 (also known as regdanvimab) showed direct recognition of RBM by binding to 12 (K417, Y449, Y453, L455, F456, E486, Y489, F490, Q493, G496, and Y505) of the 21 residues of the RBD/ACE2 interface without conformational changes. Hence, the neutralising potencies of anti-SARS-CoV-2 MAbs are affected by the extent of their abilities to interfere with RBD-ACE2 interaction. This depends on the target residues of the epitope recognized by the corresponding MAb. For instance, MAb B38 completely abolished RBD-ACE2 interaction by binding to 86% of RBD/ACE2 interacting residues. A single dose of this MAb (25 mg/kg) administrated to hACE2 transgenic mice after a viral challenge significantly reduced the viral load and inhibited pathologic damage in the lung tissue.

Although a few MAbs have been investigated for their paratope-epitope interaction by cryo-EM, crystallographic and/or mutagenesis experiments, current findings suggest paratope sites of some neutralising anti-RBD MAbs interact with non-RBM residues of RBD. Zost et al. found two of the most potent neutralising MAbs, including MAb COV2–2196 (also known as tixagevimab or AZD8955) and MAb COV2–2130 (also known as cilgavimab or AZD1061), recognise a linear peptide (60 aa) on RBD interacting with ACE2 binding. Crystal structure of a potent neutralising MAb, designated P2B-2F6, recognising non-ACE2-interacting residues of RBD, including K444, G446, G447, N448, Y449, N450, L452, V483, E484, G485, F490, and S494, revealed interactions occurred between the MAb and the ACE2. Also, CV07–270 binds to a similar epitope as P2B-2F6. These interactions were detected between the light chain residues of the MAb, including R56, S58, G59, R63, S78, and G79 with D67, K68, A71, K74, E110, and K114 residues of ACE2, that prevented the efficient RBD-ACE2 interaction (Figure 5). Thus, neutralising effect of an anti-RBD MAb might be partially mediated via steric hindrance from the MAb on the ACE2 that consequently inhibits efficient ACE2 interaction with RBD. Recently, a cocktail of two fully human non-

**Figure 5** Assignment of epitope residues of (a) Receptor binding domain (RBD), (b) N-terminal domain (NTD), (c) SD2, and (d) SH-targeted by neutralising monoclonal antibodies (MAbs) illustrated in Figure 4. NTD: N-terminal domain; RBD: Receptor binding domain; RBM: Receptor-binding motif, SD2: Subdomain two; SH: Stem helix.
overlapping anti-RBD MAbs (REGN-COV2) received FDA EUA for treatment of mild to moderate COVID-19 non-hospitalised high-risk patients. REGN10933 (casirivimab) and REGN10987 (imdevimab) were isolated from the VelocImmune mouse platform and human B cells, respectively. Casirivimab binds to the spike-like loop region of RBD on one side of the ACE2 interface from above, while imdevimab can only target RBD from the front or the lower left edge, providing the probability of simultaneous binding of two MAbs to distinct regions of the RBD.

Mutagenesis experiments allowed identifying hot spot residues. For example, F486 or N487 residues were defined as hot spots for MAb COV2-2196 epitope binding. Mutagenesis analysis for MW05 revealed E484 as the hot spot residue of RBD. In addition, the paratope of the MAb occupied a non-ACE2-interacting residue of RBD: F490. MAb COV2-2130 also recognized non-ACE2-interacting hot spot residues of RBD including K444 and G447. These findings indicate that mechanism(s) other than direct competition for RBD residues responsible for ACE2 interaction might mediate neutralising activity. Steric hindrance of the paratope site on the ACE2-interacting surface of RBD may prevent efficient RBD binding to the ACE2 and, hence, partly explain the neutralising ability of the MAbs recognising non-ACE2/RBD-interacting domains.

Direct evidence supporting this notion comes from the crystallographic studies that investigated the epitope-paratope interaction of a neutralising MAb, CB6. Crystallography revealed steric hindrance of the paratope on the ACE2-binding surface of RBD. In addition, great overlap in ACE2-binding residues of RBD between the MAb and the ACE2 was identified indicating that the MAb also competitively prevented RBD-ACE2 interaction.

Altogether, these studies indicated that anti-RBD MAbs could neutralise the virus by preventing RBD-ACE2 interaction. Monoclonal antibodies may hinder RBD-ACE2 interaction through either direct competition of paratope for ACE2-interacting residues of RBD and/or steric hindrance on the ACE2-interacting surface of RBD. Alternatively, steric hindrance of the antibody paratope on the ACE2 could inhibit ACE2 interaction with RBD.

3.1.2 Neutralising anti-RBD monoclonal antibodies unable to interfere with RBD-ACE2 interaction

Evaluation of a panel of RBD-specific anti-SARS-CoV-2 MAbs for their abilities to inhibit RBD-ACE2 interaction showed that only 26% of MAbs prevented RBD binding to ACE2. Since RBD of SARS-CoV and SARS-CoV-2 shares 73% homology, several studies have evaluated the neutralising potential of SARS-CoV RBD-specific MAbs to cross-neutralise SARS-CoV-2 RBD. In this regard, Lindsley et al. reported cross reactivity of six SARS-CoV RBD-specific nAbs with SARS-CoV-2 RBD and showed that 18F3 and 7B11 cross-neutralised SARS-CoV-2 infection. 18F3 recognized epitopes containing residues D392 and V394 in SARS-CoV RBD which were conserved neutralising epitopes corresponding to residues D405 and V407 in SARS-CoV-2 RBD. 18F3 could not block binding between RBD and the ACE2 since its specific epitope did not overlap with the ACE2 binding site. 7B11 recognized epitopes containing I428, A430, and K439 in SARS-CoV RBD which were not fully conserved in SARS-CoV-2 variants. Most epitopes recognized by 7B11 were in proximity of the ACE2 binding sites and resulted in blockade of RBD and ACE2 binding. Moreover, a cross-neutralising anti-RBD MAb (47D11) did not inhibit RBD-ACE2 interaction.

These findings suggest that neutralisation of SARS-CoV-2 can be achieved by Abs without interfering with ACE2 interaction. Interestingly, the human SARS-CoV specific MAb 47D11 was able to cross-neutralise the SARS-CoV-2 virus. Given that the core subdomain of RBD, rather than RBM, is more conserved between SARS-CoV-2 and SARS-CoV viruses (aa identity of 86.3% for the core subdomain vs. 46.7% for RBM sub-domain), the target epitope of MAb 47D11 is probably localised on the core sub-domain of RBD of SARS-CoV-2. S309 is another cross-neutralising anti-RBD MAb (also known as sotrovimab or Vir-7831 which does not interfere with RBD-ACE2 interaction) which recognises the non-RBM region of RBD. Therefore, it might be assumed that neutralising anti-RBD MAbs, which are unable to inhibit RBD-ACE2 interaction, recognise non-RBM epitopes of RBD. It is not surprising, as the RBM sub-domain is responsible for the virus binding to ACE2. On the other hand, sotrovimab, which was recently authorised for emergency use by FDA, was found to enhance Fc-dependent effector mechanisms including antibody-dependent cellular cytotoxicity (ADCC) and antibody-dependent cellular phagocytosis (ADCP) that could increase clearance of the virus as well as the infected cells. This suggests that activation of immune cells, including natural killer cells, monocytes, and macrophages by the neutralising anti-RBD MAbs, could contribute to virus elimination.

 Moreover, comparison of neutralising activity of the intact antibody with its antigen-reacting Fab (Fab) showed higher neutralising activity by the intact IgG. This implies that cross-linking and subsequent virus aggregation that facilitates virus clearance from circulation might partly mediate neutralisation by MAbs. Intact forms of two other neutralising anti-RBD MAbs possessed higher neutralising activity compared to their Fab fragments, and thus, the MAbs also caused Fc-dependent effector functions including ADCC and ADCP. Interestingly, one of the MAbs recognized the ACE2-binding surface of RBD suggesting that neutralising anti-RBD MAbs that interfere with RBD-ACE2 interaction may also contribute to virus elimination by increasing Fc-mediated effector functions and virus cross-linking.

3.2 Neutralising anti-S monoclonal antibodies recognising non-RBD epitopes

Although a major proportion of the reported anti-SARS-CoV-2 neutralising MAbs are directed against the RBD fragment of the S protein, neutralising anti-S antibodies recognising non-RBD epitopes
of S protein have also been reported. Sera from 40% of COVID-19-infected patients contained both neutralising anti-S1 and neutralising anti-S2 antibodies, and only 4% of patients with neutralising activity developed only anti-S2 antibodies. Rogers et al. isolated anti-S MAbs that recognized non-RBD epitopes. However, in sharp contrast with anti-RBD MAbs, only a minority of these MAbs demonstrated neutralising activity.

On the contrary, Liu et al. identified that 52% of the neutralising MAbs recognise non-RBD epitopes, 42% of them were directed against NTD and 10% recognise neither RBD nor NTD epitopes. Anti-NTD MAbs possessed similar or slightly higher neutralising potency compared to anti-RBD MAbs indicating that anti-S MAbs recognising non-RBD epitopes could neutralise the virus as efficient as anti-RBD MAbs. Although it is currently unclear how antibody binding to the NTD fragment neutralises the virus, it is possible that anti-NTD MAbs prevent the prefusion-to-postfusion conversion of the S protein. It has been shown that coronaviruses NTD could bind to carbohydrate contents of target cells which facilitates conversion of the S protein from the prefusion state to the postfusion state. Accordingly, an anti-NTD MAb (designated 7D10) that recognized the MERS-CoV virus (a type of coronaviruses) inhibited prefusion-to-postfusion conformational change of the S protein. Given that the amino acid sequence of NTD is highly conserved between different coronaviruses, anti-NTD SARS-CoV-2 MAbs may neutralise the virus by inhibiting fusion of the viral membrane to the target cell. Cryo-electron microscopy analysis of the complex of an anti-NTD MAb (44A8) with NTD revealed that the MAb could restrain the conformational change of the S protein from prefusion to postfusion state upon binding to NTD loops, including N3 and N5 loops. Also, only two MAbs designated COV2-2676 and COV2-2489, belonging to a panel of human MAbs against different epitopes on the NTD of SARS-CoV-2 S protein, displayed neutralising activity via inhibiting a post-attachment step in the infection cycle. Interestingly, Cheng et al. found that there is a superantigen-like motif in the proximity of S1/S2 cleavage site that is similar to a staphylococcal enterotoxin B segment in sequence and structure. They reported that an anti-SEB MAb, designated 6D3, cross-reacts with this viral motif, especially the polybasic PRRA insert (aa: 681–684). This interaction resulted in prevention of infection through interfering with the proteolytic activity of TMPRSS2/furin and blocking the access of host cell proteases to the cleavage site.

Stem helix (aa: 1134–1151) is a highly conserved sequence within the S2 fusion subunit of β-coronaviruses. It forms a surface exposed membrane-proximal helical bundle and is critical for membrane fusion in the prefusion conformation of trimeric spike. It has been shown that it induces the antibody response during natural infection. A number of MAbs targeting SH demonstrated potent neutralising properties. CV3-25 identified an epitope in the SH and blocked membrane fusion. CC40.8 is another neutralising anti-stem helix MAb with neutralising effects against SARS-CoV-2 in vivo. Also, Pinto et al. reported that among the five nAbs recognising motif F1148KEELDKYF1156 of SH, S2P6 was the most broadly neutralising antibody against all β-coronaviruses through blocking membrane fusion as well as Fc-mediated effector functions.

Taken together, these findings indicate that non-RBD epitopes of the S protein might induce nAbs with comparable neutralisation potency as anti-RBD MAbs. This highlights importance of non-RBD epitopes of the S protein as additional neutralising epitopes for vaccine design as well as passive immunotherapy purposes. Genome wide analysis of virus variants showed that the RBD sequence is the most variable region prone to mutations. Currently, the exact mechanism(s) utilised by non-RBD-binding MAbs to neutralise the virus is not fully understood. Different neutralisation mechanisms have been proposed including inhibition of prefusion-to-postfusion conformational change of the S protein preventing virus membrane fusion with the host membrane, Fc-mediated effector functions, steric hindrance of Fab as well as Fc regions on the ACE2-binding site of the S protein and finally conformational changes in the ACE2 binding site leading to abrogation of the binding of the virus to its receptors on target cells.

Thus, vaccine designs based on the conserved regions in RBD and outside RBD are the favoured candidates for inducing protective immunity capable of neutralising the emerging pan-coronavirus variants.

4 | NEUTRALISING ACTIVITY OF MONOCLONAL ANTIBODIES AGAINST EMERGING SARS-CoV-2 VARIANTS

Since beginning of the COVID-19 pandemic, ongoing evolution of SARS-CoV-2 has led to emergence and circulation of genetic lineages around the world. Emerging variants are classified either as variants of interest or as variants of concern (VOC) by the WHO Virus Evolution Working Group. VOIs are no longer circulating or are detected at very low levels and do not confer a significant or critical risk for public health. SARS-CoV-2 variants harbouring genetic changes predicted or known to affect transmissibility, disease severity, immune escape, and diagnostic or therapeutic escape are considered as VOIs. On the contrary, VOCs have been demonstrated to be associated with one or more of the following changes: increased transmissibility, deleterious change in COVID-19 epidemiology, increased virulence and decreased effectiveness of public health measures or available diagnostics, vaccines, and therapeutics. Currently, the designated VOCs are as follows: Alpha variant (B.1.1.7), first identified in the United Kingdom, contains N501Y substitution in RBD. Beta variant (B.1.351), first detected in South Africa, contains three important mutations in RBD including N501Y, E484 K, and K417 N. Alpha and Beta variants are significantly more transmissible (43%–82% and 50%, respectively), due to N501Y substitution that enhances the accessibility of RBD and binding affinity to ACE2.85,87-89 Gamma variant (p.1) was first found in Brazil with biologically important mutations in the RBD region including N501Y, E484 K, and K417 N/T. Although K417 N/T substitutions found in Beta and Gamma variants decreased the binding affinity, N501Y and E484 K
Mutations enhanced the binding affinity of their RBDs to ACE2.\textsuperscript{90} Delta variant (B.1.617.2), first documented in India, harbours two substitutions in the RBD, including L452R and T478K associated with its 97% transmissibility and higher affinity and stability of S protein conformation.\textsuperscript{91-94} Omicron (B.1.1.529), which was first detected in South Africa, harbours 34 mutations, 15 of which are in the RBD region, leading to 4-fold increased infectivity compared to wild-type SARS-CoV-2. Lambda (C.37, Peru) and Mu (B.1.621, Colombia) variants are considered as VOIs.\textsuperscript{95} Each variant has heavily mutated spike proteins. Continuous evolution of SARS-CoV-2 can reduce MAb effectiveness if any of the mutations change epitopes targeted by the antibodies. Figure 6 shows spike mutations in VOC variants.

Several studies have evaluated the neutralising activity of MAbs authorised for emergency use by the FDA-emergency use (EU) in advanced clinical trials on new emerging variants in comparison with the prototype SARS-CoV-2 virus. Wang et al. performed an extensive study to assess the neutralising activity of several MAbs targeting outer side, RBM (including bamlanivimab and casirivimab), inner side of RBD (including imdevimab and sotrovimab) and NTD (including 4A8) against all current VOCs. Neutralising activity against B.1.1.7 was slightly reduced by a few MAbs such as sotrovimab because of N501Y substitution as well as NTD-directed MAbs due to Δ144. The B.1.351 variant was resistant to the neutralising activity of most M Abs specific for NTD and RBM. Accordingly, the neutralising activity of bamlanivimab and casirivimab was completely or markedly abrogated against B.1.351 because of E484K and K417N mutations and the neutralising activity of 4A8 was abolished due to Δ242-244 and/or R246I mutations. However, combination of casirivimab and imdevimab maintained much of the neutralisation activity against the B.1.351 variant.\textsuperscript{96} They also reported that the p.1 variant is relatively refractory to neutralisation by the FDA-EU authorised MAbs through adaptation of a conformation in trimer p.1 with one RBD in the “up” position, facilitating entry of the virus to target cells.\textsuperscript{97} In contrast, while bamlanivimab efficiently neutralised the B.1.1.7

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure6.png}
\caption{Spike amino acid mutations in Alpha, Beta, Gamma, Delta, and Omicron VOCs. VOCs: variants of concern (VOC)}
\end{figure}
variant, it lost the neutralisation effect against B.1.135 carrying the E484 K substitution. In another study, complete mapping of all mutations to RBD by bamlanivimab, and its cocktail combination with LY-CoV016 (etesevimab) was conducted. The results indicated that the E484 K substitution escapes bamlanivimab and K417N/T escapes etesevimab. Both mutations are present in B.1.351 and p.1 variants. In another study, casirivimab and imdevimab were tested against two VOCs including B.1.1.7 and B.1.351. Imdevimab maintains its neutralisation effectiveness against B.1.1.7 and B.1.351, but casirivimab lost reactivity against B.1.351 due to K417 N and E484 K mutations in RBD. B.1.1.7 and B.1.351 variants reduced neutralisation activity of six out of eight MAbs obtained from blood samples of COVID-19 convalescent patients. In accordance with these findings, the majority of our RBD-specific neutralising mouse hybridoma MAbs displayed significant neutralisation reduction to p.1 and B.1.353 variants in comparison with the wild-type spike protein (unpublished data).

McCarthy et al. assessed reactivity of 4A8 as an NTD-binding nAb and showed that it does not recognise the S protein with the following deletions: Δ69–70 + Δ144/145 (both found in the B.1.1.7 lineage and Δ69–70 which is key for increased infectivity of the B.1.1.7 lineage), Δ141–145, Δ144/145, Δ146, and Δ243–244 (found in the B.1.351 lineage). However, its binding to Δ210 and Δ69/70 alone remained unchanged, suggesting that the NTD deletions are not enough as the battalion of neutralising antibodies targeting different S epitopes. B.1.1.7 mostly conferred resistance to neutralisation by the NTD-directed nAbs, suggesting that developing nAbs against subdominant epitopes need to be considered against emerging variants. Thus, emergence of mutations similar to B.1.1.7 and B.1.351 is considered a critical challenge for therapeutic MAbs. B.1.351 was the most resilient variant to COVID-19 patient-derived MAbs, followed by p.1 and B.1.1.7 variants. This resistance is largely mediated by Δ144 and Δ242–244 mutations in NTD and K417 N/T, E484 K, and N501Y mutations in RBD. The Delta variant was refractory to neutralisation activity of bamlanivimab by impaired binding of the MAbs to the spike protein. An extensive in vitro and in vivo study of a panel of MAbs including COV2-2196, COV2-2130, sotrovimab, 47D11, casirivimab, imdevimab, bamlanivimab, and etesevimab was conducted against B.1.1.7, B.1.351, and B.1.617.1 variants. In vitro experiments showed no significant changes in neutralising activity of all MAbs against B.1.351, and B.1.617.1 variants, however, imdevimab and bamlanivimab displayed 10-fold decrease and complete loss of reactivity, against B.1.617.1, respectively. Low prophylactic doses of MAbs inhibited SARS-CoV-2 infection by tested variants in K18-hACE2 transgenic mice, 12952 immunocompetent mice and hamsters, except for bamlanivimab monotherapy and bamlanivimab and etesevimab combination therapy, which demonstrated complete loss of protective activity against B.1.351 and B.1.617.1. In another study, while sotrovimab showed 3-fold reduction and the combination of COV2-2130 and COV2-2196 demonstrated ~200-fold reduction in neutralisation activity against Omicron, other RBM-specific MAbs, including casirivimab, imdevimab, bamlanivimab, etesevimab, and CT-P59 completely lost antiviral activity. Also, Planas et al. reported that bamlanivimab, etesevimab, casirivimab, imdevimab, tixagevimab, and regdanvimab completely lost the neutralising potency against B.1.617.2 and Omicron variants. Interestingly, sotrovimab was the only antibody maintaining the neutralising potency with a relatively similar activity against these two variants. Although 85% of nAbs lost antiviral efficacy against Omicron, this variant showed less negative effect on nAbs with broad sarbecovirus (the viral subgenus containing SARS-CoV and SARS-CoV-2) neutralising activity. Gruell and colleagues have recently investigated the neutralising activity of a number of MAbs including bamlanivimab, etesevimab, casirivimab, imdevimab, P2B-2F6, and etesevimab against a variety of VOCs. They demonstrated that all antibodies maintain neutralising activity against B.1.1.7, B.1.351, and B.1.617.2, with the exception of bamlanivimab which lost its neutralising activity against B.1.351, and B.1.617.2. Notably, sotrovimab was the only antibody that maintained neutralising activity against the Omicron variant. Therefore, Omicron variant exerted substantial humoral immune evasion and nAbs recognising the sarbecovirus conserved region remain most effective. Altogether, ongoing major antigenic shifts and drifts and increased transmissibility and affinity of new emergent variants confer serious challenges to current therapeutic antibodies.

5 CONCLUSION

Neutralising anti-SARS-CoV-2 MAbs could serve as prophylactic/therapeutic agents in COVID-19 infection. Epitope mapping of the reported neutralising anti-SARS-CoV-2 MAbs has revealed that the neutralising epitopes of SARS-CoV-2 virus are mainly located on the RBD fragment of the S protein. Considering the crucial role of the RBD fragment in the virus binding to the ACE2, this is not a surprising finding. Inhibition of RBD-ACE2 interaction by MAbs might be either mediated through direct competition with RBD residues responsible for ACE2 interaction and/or steric hindrance on ACE2-interacting RBD residues mediated by the antibody paratope. Moreover, neutralising anti-RBD MAbs could enhance viral neutralisation by increasing antibody effector functions, including ADCC and ADCP as well as virus cross-linking.

Current studies also highlight that non-RBD epitopes of the S protein, including the NTD fragment, might elicit nAbs with neutralising potency comparable to anti-RBD antibodies. This indicates that non-RBD epitopes of the S protein could be considered as neutralising epitopes particularly with respect to the emerging SARS-CoV-2 variants. Although the exact mechanisms of virus neutralisation by these MAbs are not fully understood, several neutralisation mechanisms have been proposed and discussed in this review. Altogether, more studies are required to focus on neutralising MAbs directed against non-RBD regions of the S protein in order to generate MAbs with broad neutralising activity and to elucidate their possible neutralising mechanisms.
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CONFLICT OF INTEREST

The authors declare no conflict of interest.

AUTHOR CONTRIBUTION

Somayeh Ghotloo and Faezeh Maghsood wrote the initial draft of manuscript. Faezeh Maghsood designed and illustrated the figures. Forough Golsaz Shirazi, Mohammad Mehdi Amiri, and Christiane Moog edited the initial draft. Fazel Shokri conceived the study, edited and revised the manuscript. All the authors approved the final version of the manuscript.

DATA AVAILABILITY STATEMENT

Data sharing is not applicable to this article as no datasets were generated or analysed during the current study.

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