Application of a method for engineering multivalent antibodies to substantially enhance functional affinity of clinical trial anti-SARS-CoV-2 antibodies

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Short title: Increasing functional affinity of anti-SARS-CoV-2 antibodies

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Abstract

Infection by the severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) causes COVID-19 disease. Therapeutic antibodies are being developed that interact with the viral spike proteins to limit viral infection of epithelium. We have applied a method to dramatically improve the performance of anti-SARS-CoV-2 antibodies by enhancing avidity through multimerization using simple engineering to yield tetrameric antibodies. We have re-engineered six anti-SARS-CoV-2 antibodies using the human p53 tetramerization domain, including three clinical trials antibodies casirivimab, imdevimab and etesevimab. The method yields tetrameric antibodies, termed Quads, that retain efficient binding to the SARS-CoV-2 spike protein and show up to two orders of magnitude enhancement in neutralization of pseudovirus infection. The tetramerization method is simple and general and its application is a powerful methodological development for SARS-CoV-2 antibodies that are currently in pre-clinical and clinical investigation.
Introduction

There is a continued need for new antibodies for use in clinical situations such as acquired disease and infections. This requirement highlights the associated costs of both individual antibody development and production of sufficient antibody for each efficacious dose. In theory, higher potency antibodies require lower dose which reflects on the cost of goods for patient implementation. We recently described a method for increasing functional affinity through multivalent antibody engineering \(^1\) to significantly improve potency by avidity effects. We have now applied this methodology to enhancing anti- SARS-CoV-2 antibodies including two that are in clinical use for COVID-19.

The advent of new viruses that can infect humans is a continuing threat to the population. The emergence of SARS-CoV causing severe acute respiratory syndrome occurred in 2002 and followed in late 2019 by a novel coronavirus SARS-CoV-2 are examples\(^2\) & PMID: 17934078. Methodologies to produce the best antibodies are a critical need for keeping ahead of new, emerging infectious agents. Controlling the SARS-CoV-2 pandemic requires development of rapid, sensitive testing to determine transmission but also effective therapeutic antibodies to prevent viral entry into cells that causes the subsequent COVID-19 symptoms. SARS-CoV-2 enters cells by binding to the angiotensin-converting enzyme 2 (ACE2) receptor via a spike protein displayed on the viral coat surface\(^3,4\). The severity of the SARS-CoV-2 pandemic has initiated a major international effort to develop antibodies that bind to the SARS-CoV-2 spike protein in the ACE2 receptor binding domain (RBD). These have been derived as monoclonal antibodies (mAbs) from infected people\(^5-12\), by animal immunizations with SARS-CoV-2 spike glycoprotein\(^13-17\) or phage screening methods\(^18-21\). At present more than a dozen anti-SARS-CoV-2 mAbs are in clinical development\(^22\) of which bamlanivimab (LV-CoV555, Lilly) and a cocktail antibody containing casirivimab (REGN10933, Regeneron) and imdevimab (REGN10987, Regeneron) have received Emergency Use Approval
(EUA) from the Food and Drug Administration. Etesevimab (CB6/Junshi, also known as JS-016 and LY-CoV016) is in phase I trial (NCT04441931) and is also being trialled in combination with bamlanivimab (NCT04427501).

One of the aims of the anti-SARS-CoV-2 antibody work is the development of high affinity molecules that can bind to the RBD and prevent the virus from attaching to its cell receptor ACE2. This needs to be a rapid process as the pandemic is unabated and new variant SARS-CoV-2 viruses begin to appear that could potentially need a new generation of antibodies. Antibody development can be a laborious process involving affinity maturation to generate the highest affinity molecules with the most effective virus neutralization potency. Given the unprecedented manufacturing scale-up challenges that would be required to produce and rapidly treat millions of people, it is pertinent that the most effective antibodies are prioritized and developed whereby antibody dose per patient can be minimized without compromising efficacy. With this in mind, we adapted a new method for increasing valency of antibodies by fusing the human p53 tetramerization domain (TD). The multivalent antibodies created by this method are self-assembling multimeric proteins that can be engineered into various different formats (Supplementary Fig. 1). This simple method can be applied to any antibody and is currently especially pertinent to anti-SARS-CoV-2 antibodies. We have applied this method to six published anti-SARS-CoV-2 antibodies, two of which are currently in clinical use for COVID-19. We show that all the tetramerized antibodies have dramatically enhanced binding properties against SARS-CoV-2 spike protein. The simple multimerization strategy described in this study can be rapidly adapted to reformat other anti-SARS-CoV-2 antibodies or antibodies against any other targets. The enhanced anti-SARS-CoV-2 antibodies are now available for clinical development.
Results

A method for enhancement of anti-SARS-CoV-2 antibody potency by tetramerization

A method for increasing valency of antibodies \(^1\) was employed to reformat published anti-SARS-CoV-2 antibodies to increase the valency and augment functional affinity. The method is flexible because it simply relies on the p53 tetramerization domain to create self-multimerizing proteins. We have engineered various types of reformatted, engineered antibodies compared to IgG antibody as illustrated in Supplementary Fig. 1A). These include single chain variable fragment (scFv)-TD (Supplementary Fig. 1B), fragment antigen-binding (Fab)-TD (Supplementary Fig. 1C), whole immunoglobulin (Ig)-TD (Supplementary Fig. 1D) or whole Ig but lacking the hinge regions where disulphide bonding would normally occur (monomeric, mIg)-TD(Supplementary Fig. 1E).

We initially evaluated three anti-SARS-CoV-2 antibodies, viz. CR3022 isolated from a phage library \(^8\) and two mAbs obtained using data from a COVID-19 convalescent patient (H4 and B38) \(^12\) to evaluate the effect of multimerization on binding. We used CR3014 as a negative control as it binds SARS-CoV but not SARS-CoV-2 \(^8\). In these cases, the antibodies were prepared as tetramers with Fab (Fab-TD) or single chain variable fragment (scFv-TD) formats and the Quads purified to homogeneity by nickel chromatography and by size exclusion chromatography (SEC). The protein preparations were analyzed by SDS-PAGE to confirm the component chains (Fig. 1A, B).

In order to ensure maintained binding efficacy in the tetrameric forms, we evaluated the binding properties of the engineered antibodies by surface plasmon resonance (SPR) assays (Fig. 1C-I) with immobilized SARS-CoV-2 RBD protein. The original CR3022 IgG format (bivalent) and the engineered CR3022 Fab-TD (tetravalent) Quad have similar properties with \(k_{\text{off}}\) values at 0.00012 and 0.00042 s\(^{-1}\) respectively (Supplementary Table 1A). Further, the two COVID-19 patient-derived mAbs engineered...
as Fab-TD tetramers (H4 and B38, Fig. 1G, I) or as H4-scFv-TD (Fig. 1F) also maintained RBD binding and displayed slow off rates. The Fab versions (both CR3022 and H4) have more rapid off rate and subsequently a weaker Kd values (Fig. 1E, F). Comparing H4-Fab-TD and H4-scFv-TD with H4-Fab illustrates the improved k_{off} and subsequently K_d values (Fig. 1F-H, Supplementary Table 1B) that results from the increase of valency. Notably, the H4-Fab-TD out-performs the H4-scFv-TD, presumably because the tetravalent Fab has additional avidity effects on binding to the RBD on the SPR chip. The K_d of Fab version of the H4 mAb (14nM) is poor compared to the Quad proteins, mainly due to the relatively fast k_{off} (Supplementary Table 1B). These data show that Quad proteins maintain and improve binding to the coronavirus spike protein after tetramerization. No binding to SARS-CoV-2 RBD was observed with the CR3014-Fab-TD Quad as expected (Supplementary Table 1A). Therefore, this method for tetramerization of the antibodies confers advantageous valency and avidity of binding to the SARS-CoV-2 RBD protein.

**Augmented detection of SARS-CoV-2 spike protein using reformatted tetrameric antibodies**

We conducted direct enzyme-linked immunosorbent assays (ELISAs) to evaluate the effects of valency on the binding efficacy of the tetramerized Quad antibodies. We used spike protein-absorbed ELISA plates, with either SARS-CoV-2 S1 spike protein (Fig. 2A, comparing CR3022 formats and Fig. 2C comparing CR3022 with H4 and B38 Quads) or SARS-CoV-2 RBD (Fig. 2B, D with similar comparisons), and titrated antibody binding (detected by anti-HIS HRP antibody). CR3022-scFv or CR3022-Fab showed approximately 50% binding at about 1nM when interacting with either S1 or RBD, whereas whole IgG is closer to 0.1nM (Fig. 2A, B). The CR3022-Fab-TD or scFv-TD display the best functional affinity in this assay as the 50% binding of the tetramers CR3022-Fab-TD
or scFv-TD was closer to 0.05nM, due to increased valency and avidity effects of tetramerization.

The consequence of tetramerization was also tested with binding of the H4 and B38 mAbs, comparing interaction of H4-Fab-TD or H4-scFv-TD with CR3022-Fab-TD or CR3022-scFv-TD with the SARS-CoV-2 S1 spike protein (Fig. 2C) or the RBD (Fig. 2D). We observed the most potent Quad antibody was the H4-scFv-TD (50% binding about 0.02nM) which is about 5x better than CR3022 IgG, or compared with CR3022-scFv-TD (50% binding about 0.05nM). The H4 or B38 Fab-TD proteins did not have especially potent binding curves as for instance H4-Fab-TD displayed 50% binding at about 0.1nM, which parallels the whole IgG ELISA (Fig. 2A) and SPR (Fig. 1). B38-Fab-TD is the weakest of engineered proteins in its binding to the spike protein.

The increase in functional affinity in the reformatted H4 and CR3022 was emphasized by comparison of their scFv-TD forms with the H4-scFv or the CR3022-scFv and of Fab-TD with CR3022-Fab (respectively Supplementary Fig. 2A, B). The Quads saturate spike protein binding at lower molarity than the scFv or Fab counterparts. The antigenic epitopes on the SARS-CoV-2 spike protein for CR3022, H4 and B38 antibodies does not overlap. It was possible therefore that the enhanced mAb tetramers could give additive effects for detection of the virus protein. This was assessed in ELISA binding of mixtures of CR3022 Fab-TD, B38-Fab-TD and H4-Fab-TD to SARS-CoV-2 RBD or S1 compared to CR3022-Fab-TD alone (respectively Supplementary Fig. 2C and D). We did not observe additive effects, probably because of steric hindrance of the bulky antibody tetramers binding to the spike protein antigens in proximity.

We further evaluated the potential of employing anti-SARS-CoV-2 CR3022 mAb in viral antigen detection systems. The spike protein detection levels were determined in a sandwich ELISA using, as capture proteins, a dimeric version of the extracellular domain of SARS-CoV-2 receptor ACE2-Fc (Supplementary Fig. 2E) or a tetrameric version
(ACE2-Fc-TD)\textsuperscript{30} (Supplementary Fig. 2F). The CR3022 Fab-TD and scFv-TD tetramers displayed the highest signal performing with higher efficacy than the CR3022 IgG1 in various situations (Supplementary Fig. 2E-H). In all ELISA data, the non-binding CR3044 served as negative control, as predicted from published binding data\textsuperscript{8}.

**Potency threshold increase in virus neutralization by tetrameric antibodies**

The H4 and B38 mAbs bind different epitopes on the spike protein and both are neutralizing\textsuperscript{12}. The tetramer reformatted Quad versions of these were tested for their ability to interfere with coronavirus infection using a pseudotyped lentivirus-based neutralization assay. Interference with pseudotyped virus infection of HEK293T/17 cells stably expressing the ACE2 receptor and TMPRSS2 protease (HEK293T/17-A2-T2) by either H4-Fab-TD, H4-scFv-TD or B38-Fab-TD was compared to a commercially available neutralizing IgG mAb (SAD-S35) and to our tetramerized ACE2-Fc-TD protein\textsuperscript{30}. The most effective protein in this neutralization assay was the tetrameric scFv version of H4 that interferes with a 50% inhibitory concentration (IC\textsubscript{50}) 0.011 μg/mL compared to 0.23 μg/mL for SAD-S35 and 0.051 μg/mL for ACE2-Fc-TD. The two Fab-TD antibodies have IC\textsubscript{50} values of 0.15 (Fig. 3A). The IC\textsubscript{50} of H4-scFv-TD and of ACE2-Fc-TD are closer when determined from molar amounts, being respectively 0.084 nM and 0.113 nM (Fig. 3B). These data show that the mAbs reformatted as Quad tetramers have greater potency for virus neutralization.

As a means to expand on the flexibility of the tetramer valency increase method, we investigated anti-SARS-CoV-2 antibodies for which clinical trials have been reported. There are several such anti-SARS-CoV-2 antibodies under clinical trial investigation either as mono-therapy, for instance, LY-CoV555\textsuperscript{7} and CB6/Junshi, (etesevimab, also called LY-CoV016) are in phase I trials or as a cocktail REGN-COV2 which is a mixture of mAbs,
REGN10987 (imdevimab) and REGN10933 (casirivimab)\textsuperscript{31}. Accordingly, we reformatted the mAbs REGN10987, REGN10933 and CB6/Junshi with the p53 tetramerization domain into three types of Quad antibody format, namely Fab-TD, Ig-TD and mIg-TD (illustrated in Supplementary Fig. 1) to compare with the IgG antibody forms that are in the clinical trials.

These Quad proteins were purified using protein A or Ni-HTA chromatography and purity and integrity assessed with SDS-PAGE in reducing and non-reducing conditions (Supplementary Fig. 3A-C). To ensure that the Quad protein formats bind to the SARS-CoV-2 RBD and to compare with the IgG forms, a plate-based surrogate neutralization assay measuring competitive binding of ACE2 with SARS-CoV-2 RBD was established (Supplementary Fig. 4). Anti-human IgG was used to capture ACE2-Fc and this was incubated with pre-mixed anti-SARS-CoV-2 competitor antibodies plus biotinylated SARS-CoV-2 RBD. Non-competed RBD that bound to ACE2-Fc was detected with HRP-conjugated streptavidin. In each case, we observed ACE2 binding neutralization was achieved at substantially lower concentrations of the tetramerized forms of mAb compared to the original IgG (Supplementary Fig. 4, panels A-C Fab-TD, panels D-F mlg and panels G-I Ig-TD). The fold potency change of the multimerized mAb versions was directly compared to the IgG formats and calculated based on the IC\textsubscript{50} values (Supplementary Fig. 4J, K). In this plate assay, the Fab-TD proteins were found to be consistently the most potent in neutralizing ACE2 interaction with SARS-CoV-2 RBD. The two larger molecular weight complexes of the Ig-TD and mlg-TD formats, yielded a similar fold enhancement of potency. Given all three Quad configurations are tetrameric molecules, it was interesting to observe that the size, shape and flexibility of these multimeric molecules appear to play an important role in potency modulation. Overall, the improvement trends in this plate-based analysis consistently shows that all three anti-SARS-CoV-2 Quad formats have greatly improved potency over the clinical stage mAbs.
The gold standard cell-based test for the efficacy of viral reagents is the capacity to neutralize the entry infection of viral particles into cells expressing the ACE2 receptor. We have used this approach to evaluate the H4 and B38 Quad proteins shown in Figure 3. Accordingly, we have determined the comparative efficiency that the Quad versions of the mAbs currently being evaluated in clinical trials to neutralize pseudotyped lentiviral infection into a permissive human cell line. Aligned to the plate competition assay, we compared the REGN10987, REGN10933 and CB6/Junshi IgG with Fab-TD, mlg-TD and Ig-TD formats. In addition, we compared these with the tetravalent H4 mAb in the form of an scFv tetrameric version (H4-scFv-TD) (Fig. 4). As a bench mark, Fig.4A displays the comparison of IgG proteins with the H4-scFv-TD showing that the two mAbs (REGN10987, REGN10933) and H4-scFv-TD have similar viral neutralization potencies. The CB6/Junshi mAb is the least potent (the calculated IC$_{50}$ values are in Fig. 4E). The relative neutralization potencies of the mAb IgG proteins are shown compared with the engineered tetramers, respectively Fab-TD (Fig.4B), mlg-TD (Fig.4C) and Ig-TD (Fig.4D) together with IC$_{50}$ values and fold change in neutralization (Fig.4E).

In all three tetramer formats of the mAbs, there is improved IC$_{50}$ compared to IgG in the neutralization of the pseudoviral infection and the potency trend in this assay is the Fab-TD, mlg-TD to Ig-TD. The Ig-TDs have the best IC$_{50}$ enhancement being 59 (REGN10987), 13 (REGN10933) and 83 (CB6/Junshi) fold change compared to IgG mAbs (Fig.4D, E). The scFv-TD engineered version of the H4 mAb has an IC$_{50}$ in the viral assay that approximates to those found for the Fab-TD and mlg-TD versions of the three clinical trial antibodies. Overall, we observed the greatest improvement for the CB6/Junshi tetramers and the greatest improved IC$_{50}$ being in the CB6/Junshi Ig-TD (Fig. 4D). Interestingly, the Fab tetramers were the least potent in the viral assay (Fig. 4) compared to the plate-based competition assays (Supplementary Fig. 4).
Discussion

A simple protein engineering method to increase valency and avidity of anti-SARS-CoV-2 antibodies

We show that implementing the tetramerization method generates antibodies with dramatically enhanced binding and importantly significantly enhanced neutralization properties against SARS-CoV-2 spike protein. These super-charged multivalent anti-SARS-CoV-2 mAbs are now available for further characterization and clinical investigation as the next-generation of anti-SARS-CoV-2 antibodies with superior binding and neutralization potency. This implies that lower dosage than currently used could be employed for therapeutic responses, which would be valuable for each treatment bolus but also for mass production as necessary amounts would be substantially lowered. Therefore, they have the potential to be developed as more effective treatment based on milligram scale doses as opposed to the gram-scale doses seen in the current EUA monoclonal antibody treatments \(^7,31\). 

Our multivalent antibody method is a rapid technology that can boost antibody binding akin to affinity maturation. The method could easily be applied to other published anti-SARS-CoV-2 antibodies because it only needs inclusion of the p53 tetramerization domain. Further, the predicted structure of the Ig-TD, as we also noted for our tetrameric ACE2 protein \(^30\), is analogous to the structure of IgA antibodies which are tetrameric due to two IgA monomers being linked by the J chain. This IgA analogy may explain the potency enhancement in viral neutralization because potentially the Ig-TD can engage more than one viral particle, while the mIg-TD may be structurally constrained to function in the same way.

Super-charging antibodies with enhanced neutralization potency through multimerization
There is a remarkable increase in neutralization potency of the tetramerized formats with the Ig-TD versions having up to 80-fold change in virus neutralization compared to the IgG1 mAbs. Generating high potency antibodies can be slow for a number of technical reasons and the requirement for rapid and flexible methods for enhancement is required. Several anti-SARS-CoV-2 antibodies have been developed that show extremely promising efficacy as COVID-19 therapeutic reagents \textsuperscript{11, 31}. Three of these have been subject to the tetramerization method within this study and in all three, substantial increased potency has been achieved. Thus, using this simple tetramerization method enhanced viral neutralization can be achieved due to increased valency and avidity.

It is remarkable that the Ig-TD format displays the greatest effective potency on the viral interference assays (Fig. 4) while the Fab-TD formats perform better in the \textit{in vitro} competition assays (Supplementary Fig. 4). This difference may be influenced by size and flexibility of the Quad proteins but also due to the antigen availability in the two assays. In the plate assay, the ACE2 protein interactions with the SARS-CoV-2 spike RBD in a two-dimensional system and the tetramer antibodies bind the SARS-CoV-2 RBD in solution. In the virus neutralization assays, the extended putative structure of the Ig-TD (like IgA) could reach between virus particles extending the potency. It is also likely that steric hindrance effects may play distinct roles in the plate assay compared to the virus-based assay. Nonetheless, the choice of Quad format that appears to be more suitable for therapeutic development is the Ig-TD.

For potential clinical implementation, the Ig-TD that includes intact Fc regions, would have additional advantages, as it would be able to interact with Fc-receptors and invoke Fc-mediated effector function thus providing additional functionality. Further, the Fc would also be able to engage with neonatal Fc-receptor and could promote antibody recycling through rescue from normal lysosomal degradation and thus prolonged circulation half-life.
Finally, the standard human IgG form of antibodies has a bivalent interaction with antigens. Increasing the valency of antibodies by tetramerization \(^1\) or by other multivalency methods \(^{18, 32-34}\). This leads to higher functional affinity through valency and avidity effects \(^1, 35\). The implementation of our protein engineering method using the human p53 protein tetramerization domain is a simple and flexible method to increase valency \(^1, 30\). Since community transmission is a major problem with SARS-CoV-2 and the appearance of new variants that appear to have increased transmissibility \(^{23, 24, 26, 27}\) there is a need for effective rapid, sensitive testing. Employing the enhanced affinity tetrameric antibodies could be a valuable asset for lateral flow device development with reformatted antibodies that could produce enhanced signal and providing more sensitive reagents to be employed in lateral flow.

In conclusion, our results show that the method for tetramerization of antibodies points the way for the reformatting anti-SARS-CoV-2 antibodies to replace the existing versions for improved therapeutic applications (and possibly for viral antigen detection). The data further generalize the concept that mAbs reformatted as tetramers, using our flexible technology, will have greater potency. The technology can be applied to any antibody but also to all new forms of proteins, as we have shown for tetramerized ACE2 receptor for SARS-CoV-2 \(^{30}\). This will confer distinct advantages by invoking increased avidity for currently known viral targets, new viruses that may emerge in the future but also generally to any antigenic target.
Materials and Methods

Production and characterization of recombinant proteins

Anti-SARS-CoV-2 antibodies, antibody fragments and ACE2 proteins were based on published data for these molecules. Gene synthesis was used to generate clones for secretion and the human p53 tetramerization domain where appropriate for production of tetrameric Quads. All the sequences were cloned in pTT5 and expressed by secretion from Expi293F (Thermofisher) according to the manufacturer’s recommendations. The proteins were purified by nickel affinity chromatography and size exclusion as described elsewhere as Avi-tagged proteins and biotinylated using BirA enzyme. All proteins were characterized by SDS-PAGE and by size exclusion chromatography.

Cloning, protein expression and purification of multivalent formats of anti-SARS-CoV-2 clinical trial mAbs. Sequences of the Fab portion of the clinical stage mAbs REGN10987, REGN10933 and CB6/Junshi were reformatted into three different tetravalent types. Construction of the Fab-TD formats involved sub-cloning the p53 tetramerization domain (TD) linked onto the C-terminus of the immunoglobulin heavy chain CH1 domain devoid of the CH2-CH3 domains. For the Ig-TD format, the TD domain was linked directly to the C-terminus of the CH3 domain. The design of the monomeric Ig-TD (mIg-TD) format was identical to the Ig-TD format except the heavy chain core hinge region was removed. The corresponding light chains (LC) of the parental mAbs were kept intact. Expression constructs containing the modified heavy chains (HC) together with the corresponding LC constructs were expressed in Expi293F cells using Expifectamine293 Reagent according to the manufacturer’s recommendations (Thermo Fisher Scientific). Multimerized mAbs were purified directly from the culture supernatant by affinity purification. Fab-TD mAbs formats were purified using Ni-NTA and the Fc containing Ig-TD and mIg-TD mAbs formats were purified by Protein A affinity chromatography.
proteins were buffer exchanged and concentrated into PBS buffer using Amicon columns (Millipore) and aliquots were stored at 4°C or at -80°C for long-term storage.

**Surface plasmon resonance of anti-SARS-Cov-2 antibodies with SARS-Cov-2 RBD**

SPR experiments were carried out using Biacore T100 (Cytiva, formerly GE Healthcare) in PBS buffer, pH 7.4 containing 0.05% Tween-20. A streptavidin-coated SA chip (Cytiva) was washed with 3 x 30s injections of 1M NaCl/50mM NaOH at 10µL/min before biotinylated SARS-Cov-2 RBD (0.2µg/mL) was captured to target immobilisation level 500RU. Antibody binding to SARS-Cov-2 RBD was analysed at 25°C using multi-cycle injections of antibody at six concentrations at 30µL/min for 180s, followed by 420s dissociation time. Concentrations between 0.312 – 10nM were used for CR3022 and CR3014 antibodies, while concentrations between 0.78 – 12.5nM were used for analysis of H4 and B38 antibodies. The surface was regenerated after each antibody injection using 1M NaCl/50mM NaOH at 30µL/min for 20s contact time followed by 120s stabilisation time. Flow cell 1 was used as a reference and so did not contain any captured SARS-Cov-2 RBD and was subtracted from all sensograms before analysis. Data were fitted to a 1:1 kinetics model using Biacore T200 Evaluation Software version 2.0.

**Direct ELISA methods**

ELISAs were carried out in clear, flat-bottomed 96-well MaxiSorp plates obtained from Thermo-Fisher Scientific, UK. SARS-CoV-2 protein (either spike or RBD) was prepared at 2 µg per mL in PBS and 100 µL per well added to an ELISA plate alongside negative control wells with 100 µL PBS alone. Plates were then incubated for 16 hours at 4 °C and washed three times with 0.05% v/v Tween-20 in PBS (PBS-T) prior to addition of 150 µL of 5% w/v BSA dissolved in PBS and room temperature (RT) incubation for 4 hours. Blocked wells were then washed 2 times with PBS-T, followed by the addition of 100 µL of His-tagged antibody diluted in PBS-T and the plate incubated for 16 hours at 4 °C. Wells
were then washed 4 times with PBS-T and incubated with 100 µL of anti-His-HRP (mouse monoclonal anti-polyhistidine antibody (clone HIS-1) conjugated to peroxidase (anti-His-HRP) was obtained from Sigma-Aldrich) diluted 1:4000 with 1% w/v BSA in PBS for 2 hours at RT, followed by washing 4 times with PBS-T. 25 µL of TMB was added to each well, incubated for 20 minutes at RT, then 25 µL 3M HCl added and absorbance read at 450\text{nm}.

**Sandwich ELISA methods**

ACE2-Fc or ACE2-Fc-TD was prepared at 3 µg per mL in PBS and 100 µL per well added to an ELISA plate alongside negative control wells with 100 µL PBS alone. Plates were then incubated for 16 hours at 4 °C and washed three times with 0.05% v/v Tween-20 in PBS (PBS-T) prior to addition of 150 µL of 5% w/v BSA dissolved in PBS and room temperature (RT) incubation for 4 hours. Blocked wells were then washed 2 times with PBS-T, 100 µL of 0-300 nM RBD diluted in PBS-T added and the plate incubated for 16 hours at 4 °C, before wells were again washed 4 times with PBS-T. His-tagged antibodies diluted to 1 nM in PBS-T were then added and incubated for 3 hours at RT. Wells were then washed 4 times with PBS-T and incubated with 100 µL of anti-His-HRP diluted 1:4000 with 1% w/v BSA in PBS for 2 hours at RT, followed by washing 4 times with PBS-T. 25 µL of TMB was added to each well, incubated for 20 minutes at RT, then 25 µL 3M HCl added and absorbance read at 450\text{nm}.

**Time course ELISA methods**

ACE2-Fc-TD was prepared at 3 µg per mL in PBS and 100 µL per well added to an ELISA plate alongside negative control wells with 100 µL PBS alone. Plates were incubated for 16 hours at 4 °C and washed three times with 0.05% v/v Tween-20 in PBS (PBS-T) prior to addition of 150 µL of 5% w/v BSA dissolved in PBS and room temperature (RT) incubation for 4 hours. Blocked wells were then washed 2 times with PBS-T, 100 µL of 0-
300 nM RBD diluted in PBS-T added and the plate incubated for 10 minutes or 1 hour at RT or 37°C, before wells were washed 4 times with PBS-T. His-tagged CR3022-Fab-TD diluted to 1 nM in PBS-T was added and the plate incubated for 10 minutes or 1 hour at RT or 37°C, before wells were washed 4 times with PBS-T. 100 µL of anti-His-HRP diluted 1:4000 with 1% w/v BSA in PBS was added and the plate incubated for 10 minutes or 1 hour at RT or 37°C, before wells were washed 4 times with PBS-T. 25 µL of TMB was added to each well, incubated for 15 minutes at RT, then 25 µL 3M HCl added and absorbance read at 450 nm. For simultaneous addition experiments, the SARS-CoV-2 RBD, CR3022-Fab-TD and anti-His-HRP antibody were added together.

**Competitive ELISA with tetravalent mAb formats.**

As a surrogate neutralization assay, inhibition of ACE2 interaction with SARS-CoV-2 RBD was measured using a competitive ELISA protocol. High binding 96-well plates (Corning) were coated overnight at 4°C with 100 ng of anti-human IgG (Binding Site). Three washes were used between each subsequent step using PBS containing 0.1% Tween 20 and incubation at room temperature for 1 hour. Coated ELISA plates were blocked with 1% BSA followed by the addition of 100 ng of human ACE2-Fc. A mixture containing serially diluted anti-SARS-CoV-2 multivalent mAbs starting at 30 nM and a fixed amount of biotinylated SARS-CoV-2 RBD (75 pM) was pre-incubated at room temperature for 30 minutes prior to adding the mixture to the coated plate. Wells containing only the biotinylated RBD with no anti-SARS-CoV-2 antibody or wells containing ELISA assay buffer only (0.1% BSA) were used as control for calculating percentage neutralization. In the next step, HRP conjugated streptavidin (Abcam) diluted at 1 in 15,000 fold was added followed by 100 ul of TMB to generate the signal. The reaction was stopped using 50 ul of 1M sulphuric acid and the absorbance was measured at 450 nm using a CLARIOstar micro-plate reader (BMG Labtech).
SARS-CoV-2 lentiviral pseudotyped virus neutralization assay

SARS-CoV-2 pseudotyped lentivirus was produced in HEK293T/17 cells following a similar approach as described \(^{38}\) DOI: 10.6084/m9.figshare.13502580 \(^{39}\), a firefly luciferase reporter gene (pCSFLW) and the SARS-CoV-2 Spike gene (pCAGGS SARS-2-Spike) to produce virus containing supernatants. Neutralizing activity of antibodies was determined with 200 TCID50 of SARS-CoV-2 lentiviral pseudotyped virus using HEK293T/17 cells stably expressing ACE-2 and TMPRSS2 (HEK293T/17-A2-T2). These were plated at 20,000 cells/well in a 96-well plate for at least 2 hours at 37 °C, 5% CO\(_2\) before use. Mixtures of antibody and pseudotyped virus were incubated with target cells for 60 hours at 37 °C, 5% CO\(_2\) and infection determined by luciferase expression using the Promega Bright-Glo assay system and GloMax Navigator plate reader, following manufacturer’s instructions. Data are normalized to pseudotyped virus and cell only controls and non-linear regression analysis performed within GraphPad Prism, for IC\(_{50}\) values.
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Author’s contributions

Participated in research concept and design: THR
Conducted experiments: AL, AM, EB, JT, CM
Performed data analysis: All authors
Wrote or contributed to the writing of the manuscript: All authors

Conflict of interest statement

Terence Rabbitts is a stake holder in Quadrucept Ltd. None of the other authors have any conflicts of interest to declare.

Additional information: Supplementary Information accompanies this paper.
Figure 1. Characterization of anti-SARS-Cov-2 antibodies with different valencies by surface plasmon resonance
The binding potencies of the anti-SARS-Cov-2 tetrameric Quad proteins was determined using surface plasmon resonance. Purified proteins were captured using immobilized SARS-Cov-2 RBD, carried out with a Biacore T200 instrument.

(A) Anti-SARS-Cov-2 antibodies were expressed by transfection and secretion from Expi293 suspension cells and characterized by SDS-PAGE following purification by Ni-affinity chromatography and gel filtration. Gels were stained with InstantBlue Ultrafast Protein Stain and Mw size markers included (left hand lanes).

(C-I) Antibody binding to SARS-Cov-2 RBD was evaluated using SPR. Biotinylated SARS-CoV-2 RBD was captured on a streptavidin chip and antibodies flowed over the surface at different concentrations. CR3022 and CR3014 antibody concentrations were 10, 5, 2.5, 1.25, 0.625 and 0.312 nM, while those for H4 and B38 antibodies were 12.5, 6.25, 3.125, 1.56 and 0.78 nM. Sensograms representative of two independent experiments are shown for (C) CR3022-Fab-TD, (D) CR3022-IgG, (E) CR3022-Fab, (F) H4-Fab (G) H4-Fab-TD (H) H4-scFv-TD and (I) B38-Fab-TD. Kinetic parameters are shown in Supplementary Table S1. Tetramerization decreases dissociation rates for the multimeric species compared to the monomeric proteins. This results in increased affinity for SARS-Cov-2 RBD for CR3022-Fab-TD compared to CR3022-Fab, and for H4-Fab-TD and H4-scFv-TD compared to H4-Fab.
Figure 2. Direct ELISA immunoassays with anti-SARS-CoV-2 antibodies detecting viral spike proteins

The ability and potency of engineered anti-SARS-CoV-2 antibodies to bind to the viral spike proteins was compared using Enzyme-linked immunosorbent assays (ELISAs). SARS-CoV-2 spike protein as either S1 (panel A and C) or RBD (panel B and D) was adsorbed onto immunoassay plates at 2 µg per µL for 16 hours at 4 °C. Wells were thoroughly washed, blocked, and various concentrations of the indicated His-tagged antibodies were added to each well and incubated for 16 hours at 4°C, followed by washing and incubation with anti-His-HRP antibody for 2 hours at room temperature. Colour was developed using TMB and absorbance read at 450 nm. All samples were run in triplicate.
Figure 3. Tetramerized H4 and B38 anti-SARS-CoV-2 antibodies are potent inhibitors of virus infection

The viral infection neutralizing potency of tetrameric versions of the mAbs H4 and B38 was examined in SARS-CoV-2 pseudovirus infection assays. Increasing concentrations of H4-scFv-TD, H4-Fab-TD or B38-Fab-TD were incubated with HEK 293T/17-A2-T2 cells after 1 hour. The cells were incubated for 60 hours before luciferase activity, indicative of viral entry, was determined. The efficacy was compared to the neutralization effects of a commercial anti-SARS-CoV-2 mAb SAD-S35 (AcroBiosystems) and a tetramerized ACE2-Fc-TD. Data are represented in µg/mL (panel A) and nM (panel B). Calculated IC₅₀ values are tabulated below each set of curves. The assays were performed twice, in duplicate with error bars indicating the standard error.
The potency of clinical stage mAbs that were engineered as tetramer formats was measured in SARS-CoV-2 pseudovirus infection assays. Increasing concentrations of
antibodies were incubated with HEK 293T/1A2-T2 cells after 1 hour. The cells were incubated for 60 hours before luciferase activity was determined to show viral entry. The data are split to different panels for clarity. Panel A: The neutralization potency of H4-scF-TD Quad was compared to clinical stage REGN10987, REGN10933 and CB6/Junshi IgG1. Panel B: The neutralization potency of REGN10987, REGN10933 and CB6/Junshi IgG1 was compared to REGN10987 Fab-TD, CB6/Junshi Fab-TD and REGN10933 Fab-TD. Panel C: The neutralization potency of the REGN10987, REGN10933 and CB6/Junshi IgG1 was compared to REGN10987 mlg-TD, REGN10987 mlg-TD and CB6/Junshi mlg-TD. Panel D: The neutralization potency of the REGN10987, REGN10933 and CB6/Junshi IgG1 was compared to REGN10987 Ig-TD, CB6/Junshi Ig-TD and REGN10933 Ig-TD. The assays were performed twice and each point in duplicate. The error bars indicate the standard errors. Computed IC\textsubscript{50} data (nM and \(\mu\text{g/ml}\)) and fold change based on nM are shown in panel E.