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A Multivalent and Thermostable Nanobody Neutralizing SARS-CoV-2 Omicron (B.I.I.529)

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Background: The severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) Omicron variants have risen to dominance, which contains far more mutations in the spike protein in comparison to previously reported variants, compromising the efficacy of most existing vaccines or therapeutic monoclonal antibodies. Nanobody screened from high-throughput naïve libraries is a potential candidate for developing preventive and therapeutic antibodies.

Methods: Four nanobodies specific to the SARS-CoV-2 wild-type receptor-binding domain (RBD) were screened from a naïve phage display library. Their affinity and neutralizing activity were evaluated by surface plasmon resonance assays, surrogate virus neutralization tests, and pseudovirus neutralization assays. Preliminary identification of the binding epitopes of nanobodies by peptidebased ELISA and competition assay. Then four multivalent nanobodies were engineered by attaching the monovalent nanobodies to an antibody-binding nanoplatform constructed based on the lumazine synthase protein cage nanoparticles isolated from the Aquifex aeolicus (AaLS). Finally, the differences in potency between the monovalent and multivalent nanobodies were compared using the same methods.

Results: Three of the four specific nanobodies could maintain substantial inhibitory activity against the Omicron (B.1.1.529), of them, B-B2 had the best neutralizing activity against the Omicron (B.1.1.529) pseudovirus (IC₅₀ = 1.658 μ g/mL). The antiviral ability of multivalent nanobody LS-B-B2 was improved in the Omicron (B.1.1.529) pseudovirus assays (IC₅₀ = $0.653 \mu g/mL$). The results of peptide-based ELISA indicated that LS-B-B2 might react with the linear epitopes in the SARS-CoV-2 RBD conserved regions, which would clarify the mechanisms for the maintenance of potent neutralization of Omicron (B.1.1.529) preliminary.

Conclusion: Our study indicated that the AaLS could be used as an antibody-binding nanoplatform to present nanobodies on its surface and improve the potency of nanobodies. The multivalent nanobody LS-B-B2 may serve as a potential agent for the neutralization of SARS-CoV-2 variants.

Keywords: COVID-19, bio-nanotechnology, self-assembly, nanobody multimers, thermal stability

Introduction

The global spread of coronavirus disease 2019 (COVID-19) has led to an unprecedented pandemic due to its high infectivity and its impact on human health and society. Given the increase in the number of infected individuals and the acceleration of the transmission speed, it provides sufficient natural selection opportunities for the virus, thus producing more mutations and providing a better breeding ground for virus mutation. The angiotensin-converting enzyme 2 (ACE2) receptor-binding domain (RBD) region of the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) spike (S) protein is the primary region of convergent mutations in circulating variants of SARS-CoV-2. Indeed, a variety of RBDmutated variants appeared, which accelerated the spread of SARS-CoV-2 or allowed it to evade antibody neutralization,

resulting in the decrease of antibody efficacy elicited by vaccines or the effect of protective neutralizing antibodies.^{1–5} The SARS-CoV-2 Omicron (B.1.1.529) variant which has an unprecedented number of mutations attracts worldwide attention, with 15 mutations in RBD, while the previous Beta (B.1.351) variant has 3 mutations (K417N, E484K, and N501Y) and Delta (B.1.617.2) variant has only 2 mutations in RBD (L452R and T478K).^{3,6} The amino acid site mutation of RBD, especially the receptor-binding motif (RBM), affects RBD interaction with ACE2, thus increasing the affinity of S protein with ACE2.^{4,7} Most neutralizing antibodies occupy the neck and shoulder epitopes of RBD, blocking the interaction between S protein and ACE2, thus preventing viruses from attaching to host cells.⁸ However, of the currently licensed or approved therapeutic monoclonal antibodies, only the antibody S309 and COV2-2196/COV2-2130 cocktails can neutralize the Omicron (B.1.1.529) variant, although the neutralization potency is also reduced by at least one-third.⁹ These antibodies, which retain neutralizing activity against Omicron (B.1.1.529) variant, can destabilize the S-trimer, and hardly block the interaction of ACE2 with RBD due to their antigen-binding sites being far away from the ACE2 binding site on the S protein.^{7,10}

Therefore, it is urgent to develop a multivalent antibody against multiple epitopes and more conserved epitopes of SARS-CoV-2. It would be able to target the highly conserved epitopes on S protein and use antibodies that recognize different epitopes on S protein in combination covering the neutralizing epitopes on S protein or RBD as much as possible.^{10,11} At present, there are several reports that SARS-CoV-2-neutralizing single-domain antibodies from naïve phage display library or immunization have a neutralizing effect on SARS-CoV-2, which can interfere with the interaction between RBD and ACE2.^{12–14} Also, existing nanobodies binding to highly conserved epitopes on SARS-CoV-2 RBD can cross neutralize SARS-CoV.¹⁵ These nanobodies are considered potential therapeutics for COVID-19 patients.

Nanobodies are the variable regions of heavy chains of camelid antibodies and lack the first constant regions in conventional antibodies, so nanobodies are typically only 12–14 kDa in size, with dimensions in the nanometer range (about 2.5 nanometers in diameter and 4 nanometers in height).^{16,17} Due to the small size of the nanobodies, the complementarity determining region 3 (CDR3) is longer than that of humans, forming a finger-like structure, which can penetrate the cavity on the surface of the antigen, so that the nanobodies can bind to epitopes that traditional antibodies are not accessible.¹⁸ In addition to being the smallest unit that retains the antigenic affinity and specificity, nanobodies have an important advantage over traditional antibodies that they can be easily assembled into multimers. As monomers, especially nanobodies isolated from nonimmune synthetic libraries usually lack the high binding affinity required for therapeutic use or high sensitivity for bioanalytical applications. In many previous studies, the affinity of nanobodies can be improved by fusing the Fc domain of human immunoglobulin G1 or connecting nanobodies to form dimerization with flexible linkers.^{12,13,19} Moreover, nanobodies can be produced in large quantities in prokaryotic expression systems, with a short operation cycle and are usually very stable,²⁰ and can be atomized for direct lung delivery,^{19,21} which is a potential candidate for developing alternative drug delivery routes.

To further improve the affinity of nanobodies, we constructed an antibody-binding nanoplatform. Lumazine synthase isolated from the *Aquifex aeolicus* (AaLS) is a potential protein scaffold particle for the development of delivery and assembly vehicle (PDB:1HQK).²³ AaLS is an internal hollow dodecahedron composed of 60 identical subunits, with an outer diameter of about 16 nm and an inner diameter of about 8 nm. It has the advantage of being uniform in size, symmetrical in structure, and thermally stable, with a melting temperature (T_m) value of about 120 °C.²³ Due to the hollow spherical structure, AaLS is a remarkable vehicle with the surface presentation of ligands and encapsulation capability. In previous studies, modified AaLS can be used to encapsulate RNA, proteins,^{24,25} and also can bind small molecule drugs or antigens/antibodies to the outer surface to deliver them into cells.²⁶ To attach nanobodies to the surface of AaLS, after genetically modifying AaLS and nanobodies by introducing SpyTag/ SpyCatcher, we conjugated the heterogeneously produced nanobodies to the preassembled AaLS scaffold by using the spontaneous isopeptide formation between SpyCatcher and its peptide partner SpyTag. Both SpyTag and SpyCatcher can be located at different positions in the protein chain and can form isopeptide bonds under various conditions (pH, buffer, and temperature).^{27,28} This Plug-and-Display coupling assembly strategy can avoid proteins of antibodies or antigens misfolding caused by over-long gene co-expression. The isopeptide bond-based conjugation method has also been used to design and enhance the efficacy of different particle vaccines, such as influenza A virus, HIV, cancer cells (PD-L1), etc.^{29–32}

In summary, the study reports the development and characterization of four anti-RBD nanobodies isolated from a naïve phage display library with SARS-CoV-2 RBD. B-B2 of the four nanobodies had the most optimal neutralizing activity against pseudovirus Omicron (B.1.1.529). The attempt of attaching monovalent nanobodies to AaLS contributes to increased thermal stability and neutralizing activity of nanobodies. The results of peptide-based ELISA initially indicated that LS-B-B2 mainly reacted with some linear epitopes of the SARS-CoV-2 RBD conserved regions, LS-B-B2 could serve as a potential agent for the neutralization of SARS-CoV-2 variants.

Materials and Methods

Antigen

The antigen, SARS-CoV-2 wildtype (WT) and Omicron (B.1.1.529) spike RBD recombinant protein, were purchased from Sino Biological (Beijing. China). A DNA sequence encoding the SARS-CoV-2 WT RBD (YP_009724390.1) (Arg319-Phe541) without any terminal peptide tag. The WT RBD protein consists of 223 amino acids. A DNA sequence encoding the SARS-CoV-2 Omicron (B.1.1.529) RBD (YP_009724390.1, with mutation G339D, S371L, S373P, S375F, K417N, N440K, G446S, S477N, T478K, E484A, Q493R, G496S, Q498R, N501Y, Y505H) (Arg319-Phe541) with the Fc region of mouse IgG1 at the C-terminus. The Omicron (B.1.1.529) RBD consists of 457 amino acids. Two proteins were both expressed in HEK293 cells and dissolved in a phosphate buffer and stock at -20 °C. The experiments involving antigen, SARS-CoV-2 WT and Omicron (B.1.1.529) spike recombinant protein, only needed to be carried out under ordinary laboratory conditions.

Screening Nanobodies Against from a Naïve Phage Display Library

The phagemid vector pADL-10b was used to generate a VHH phage display library (AlpaLife) of about 2×10^9 cfu. *E. coli* SS320 cells containing the VHH library were infected with helper phage M13KO7 to produce phages displaying the encoded VHHs as fusion proteins. Phage displaying RBD (Sino Biological)-specific VHHs were biopanned as previously described.³³ After 3 rounds of panning, 96 individual clones were selected and inoculated into the $2 \times YT$ medium supplemented with 100 µg/mL ampicillin at 37 °C overnight. The cell-free phage supernatant was detected by ELISA with M13Bacteriophage Antibody (HRP) (Sino Biological). P (positive OD₄₅₀)/N (negative OD₄₅₀) greater than 3 P/N ratios were determined as positive clones. Positive candidates were sequenced (Sangon Biotech) and aligned with CDRs amino acid sequence. A 3% bovine serum albumin (BSA) was used as a negative control for each round.

Prokaryotic Expression and Purification of Nanobodies and AaLS

An N- terminal and a C- terminal $6\times$ His tag were added to the sequences of the nanobodies for further purification. Nanobodies are ready to be assembled with scaffolds with a $6\times$ His tag and flexible linker $(G_4S)_3$ -linked spytag003 (GenBank: MT945421.1) added at the C-terminal. The C-terminal of AaLS (PDB:1HQK) added a $6\times$ His tag and flexible linker (GGS)₄-linked spyCatcher003 (GenBank: QGX07219.1). The above sequences were all synthesized and subcloned into the pET-28a(+) vector (Generay Biotech). All the plasmids were transformed into *E. coli* BL21(DE3) (TransGen Biotech). Single colonies from agar plates were used to inoculate cultures of LB broth (5 mL) containing kanamycin (50 µg/mL) and grown at 37 °C overnight (250 rpm). This starter culture was then used to inoculate 200 mL LB cultures containing kanamycin (50 µg/mL) at 0.1 OD₆₀₀ and grown at 37 °C with shaking (250 rpm). The expressions of all proteins were induced overnight at 18°C (200 rpm) after adding IPTG with the final concentration of 0.5 mM when the cell density reached 0.6. All proteins were purified using nickel columns (Thermo) according to the instructions. Finally, quantitating the purified proteins in sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) using Image J software.

Indirect ELISA to Quantitate Initial Binding

The RBD proteins were coated on a 96-well plate (Corning) at a concentration of 0.5 μ g/mL, 100 mL/well at 4°C overnight, washed three times with PBST and blocked with 200 μ L/well of 3% BSA blocking solution diluted with PBST for one hour at room temperature (RT). After washing three times with PBST, every well was added 100 μ L of primary

antibody serially diluted with blocking solution and incubated at RT for one hour. After washing three times with PBST, 100 μ L/well of Anti-His tag mouse monoclonal antibody (HRP) (Atagenix) diluted 1:2000 was added and incubated for one hour at RT. After washing three times with PBST, 100 μ L TMB (TIANGEN) was added to each well. After 10 minutes of color development, an equal volume of 1M HCl was added to stop the reaction, and the absorbance was read at 450nm. The 3% BSA was used as the negative control.

Forming Multivalent Nanobodies and Observed by Negative-Stain Transmission Electron Microscopy (TEM)

SpyCatcher-AaLS was reacted with $2\times$ molar excess of nanobodies-Spytag diluted with 1M tris-buffered saline (TBS) buffer, pH 8.5, and incubated overnight at 4°C, performing SDS-PAGE to analyze the coupling efficiency. For protein shell observation, the 20 µL sample was dropped on the copper mesh grids (400 mesh), the filter paper was used to remove excess liquid from the edge of the copper mesh 3 minutes later, then stained with 3% phosphotungstic acid waited for 2–3 minutes for TEM (JEM-1400) observation. Data acquisition and processing were performed using NTA software 3.3. Every sample was recorded three times in 30s, with detection threshold 3.

Mapping Nanobody Epitopes

The ability of two nanobodies to recognize the same epitope was detected by an ELISA using RBD-coated plates. Briefly, one HRP-labeled nanobody and another nanobody without HRP were mixed, and the mixture was added to the RBD-coated plates. If the two nanobodies recognized the same epitope, they could compete to bind to the sites of RBD, thereby reducing the amount of nanobody-HRP bound to the bottom of the well and resulting in a lower OD_{450} value. The positive control comprised a mixture of the same labeled and unlabeled nanobodies, while the negative control was the labeled nanobody alone. Inhibition rate = $(1 - OD_{450} \text{ nm})/OD_{450}$ nm negative control × 100%. The inhibition rate was classified into four groups, $\geq 75\%$ relevant; 50% to 75%, partly relevant; 25% to 50% irrelevant; < 25%, totally irrelevant. What's more, performing molecular dynamics simulations between WT RBD and nanobody using Discovery Studio (DS). PDB files of RBD protein and four nanobodies were obtained by Phyre2.

A peptide-based ELISA was used to map the epitopes of the nanobodies. The 17 peptides were based on the protein sequence of the WT RBD, and a peptide of ZIKA EDIII (TLHGTVTVEVQYAGT) as the negative control. Each peptide ranged from 9–30 amino acids in length (GL Biochem). All peptides were provided as a lyophilized powder, reconstituted in double-distilled H₂O or dimethyl sulfoxide as required to a concentration of 1 mg/mL, and stored at -20 °C. Each peptide was coated at a concentration of 1µg/mL overnight at 4 °C. The primary antibodies were diluted to the concentration of 100 µg/mL with the blocking solution. Other details are as described in indirect ELISA.

ELISA-Based SARS-CoV-2 Surrogate Virus Neutralization Test (sVNT)

To preliminarily determine the ability of nanobodies to block the binding of RBD to ACE2, ELISA-based sVNTs were performed as recommended by the manufacturer (GenScript). The serially diluted nanobodies were mixed with equal volumes of HRP-conjugated RBD and the mixture was incubated at 37°C for 30 min. 100 μ L of the mixture was then added to a 96-well plate precoated with ACE2 and incubated at 37°C for 15 min. A competitive inhibitory reaction occurred between the HRP-conjugated RBD protein and the SARS-CoV-2 neutralizing antibodies. After thorough washing, the TMB substrate solution was added to the wells and incubated for 15 minutes. The reaction was stopped by adding an equal volume of stop solution and the color intensity was measured spectrophotometrically at 450 nm. The percentage of inhibition was calculated using the following formula: % inhibition = (1 – (OD₄₅₀ sample/OD₄₅₀ of negative control)) × 100.

Pseudovirus Neutralization Assay

All the SARS-CoV-2 pseudoviruses and the envelope recombinant plasmids are donated. The SARS-CoV-2 pseudovirus assays were performed on 293T-hACE2 cells as in the previous study.³⁴ The 293T-hACE2 cells in pseudovirus neutralization assays were purchased from GenScript (Nanjing. China). Nanobodies were diluted serially with

Dulbecco's modified eagle medium (Gibco). 50 μ L nanobodies were mixed with equal volume of 1.3×10^4 TCID₅₀/mL SARS-CoV-2 WT or Omicron (B.1.1.529) pseudoviruses were incubated for 1 h in the 37 °C incubators, supplied with 5% CO₂. The negative control group is composed of cells only. 100 μ L freshly trypsinized 293T-hACE2 cells were added to a 96-well plate. The plate was incubated at 37 °C for 24 h. The culture supernatant was aspirated gently to leave 100 μ L in each well, then, 100 μ L of luciferase substrate (Beyotime) was added to each well. Two min after incubation at RT, 150 μ L of lysate was transferred to white solid 96-well plates for the detection of luminescence using a multifunctional microplate detector (Synergy H1MF) to obtain the RLU to calculate IC₅₀. All the SARS-CoV-2 pseudovirus neutralization assays were operated in the biosafety level 2 (BSL- 2) laboratory.

Surface Plasmon Resonance (SPR) Assay

Install the COOH chip following the OpenSPRTM instrument standard operating procedure. Start running at a maximum flow rate (150 μ L/min) with HEPES assay buffer. After reaching the signal baseline, load 200 μ L isopropyl alcohol and run for 10s to remove bubbles. After reaching the baseline, flush the sample ring with buffer and empty it with air. After the signal reaches the baseline, adjust the buffer flow to 20 μ L/min. Load the EDC/NHS (1:1) solution to activate the chip. 200 μ L of ligand RBD diluted in activation buffer was loaded for 4 min, and the sample loop was rinsed with HEPES and evacuated with air. Load 200 μ L of Blocking Solution, rinse the sample loop with buffer HEPES and evacuated with air. Observe the baseline for 5 minutes to ensure stabilization. The analytes were diluted with buffer and loaded at 20 μ L/min, and the binding time of protein and ligand was 240 s; natural dissociation was 360 s. The level of protein coupling is 5ug/mL, NaAc pH 6.0, 1150 RU. Using 10mM Glycine-HCl pH1.5 as the regeneration conditions. The analysis software used for the experimental results was TraceDrawer (Ridgeview Instruments ab, Sweden), and the analysis method was the One-to-One analysis model.

Differential Scanning Calorimetry (DSC)

The DSC measurement of samples prepared in PBS was carried out on MicroCal PEAQ-DSC (Malvern) at a heating rate of 2.00 °C per minute. To evaluate the thermal stability, the software MicroCal PEAQ-DSC software (Malvern) was used to determine the temperature with the maximum heat capacity as the apparent T_m .

Statistical Analysis

Statistical analyses were carried out using GraphPad Prism 9 (San Diego, CA, USA). All the data were presented as the mean \pm SD with at least two replicates. Student's *t*-test was used for two groups comparison. The Dunnett *t*-test was used for the analysis of the peptide-based ELISA. Two-sided P-values < 0.05 were considered statistically significant. (*P<0.05; ****P<0.0001).

Results

Screening SARS-CoV-2 RBD Specific Nanobodies from a Naïve Alpaca VHH Library

At present, there are numerous reports that single-domain antibodies have a neutralizing effect on SARS-CoV-2, which are considered potential therapeutics for COVID-19 patients.^{12–14} Naïve VHH library can be used for rapid acquisition and exploitation of anti-SARS-CoV-2 nanobodies through the panning using the RBD.^{13,35–37} On this account, we used purified RBD protein of the SARS-CoV-2 WT to perform three rounds of panning in a naïve alpaca VHH library by in vitro phage display technology (Figure 1A). After three rounds of enrichment (Table 1), 96 clones (Figure 1B) were randomly selected by phage-ELISA, and the clones with relatively high OD values were sequenced. Multiple sequence alignment and phylogenetic analyses were performed to exclude clones with identical sequences (Figure 1C).

Four of the nanobodies with different amino acid sequences of CDR3 were expressed successfully by the prokaryotic expression system with *E. coli* strain BL21(DE3). SDS-PAGE showed the recombinant proteins were detected at the expected apparent molecular weights, which was 15~17 kDa (Figure 1D and E).



Figure I Screening and recombinant expression of nanobodies. (A) Schematic diagram of the process of screening Nanobodies from a naïve alpaca VHH library. (B) Reactions from the 96 clones binding with SARS-CoV-2 RBD protein. (C) The phylogenetic tree of nanobodies with higher RBD reactions constructed using MEGA5.2 software. (D) Soluble prokaryotic expression and purification of four recombinant nanobodies. The destined proteins were indicated by the red arrows. (E) Alignment of the amino acid sequences of specific nanobodies with different amino acid sequences of CDR3.

Affinity and Neutralizing Potency of Monovalent Nanobodies

Due to the potency of nanobodies being greatly affected by their affinity to antigen, 15,38 The binding kinetics of the four nanobodies were measured using SPR assays with the purified WT and Omicron (B.1.1.529) RBD, respectively. The results showed that the four nanobodies have affinities to WT RBD with K_D values ranging from 3.37 nM to 317 nM. Except for B2-3, the affinities of the other three nanobodies to Omicron (B.1.1.529) RBD maintained, while B-B2 has the highest affinity to Omicron (B.1.1.529) RBD, K_D value is 49 nM (Table 2).

To characterize the binding epitopes between the four nanobodies and RBD, we performed molecular docking between nanobodies and RBD using the computer molecular simulation software DS to predict and analyze the antigen-binding epitopes of nanobodies (Figure 2A). It can be seen from the red part of the RBD sequence in the prediction model that there is a binding epitope of up to 13 or 11 amino acids between B1-4 or B-B2 and RBD (Figure 2B). Except for B2-3, the predicted binding epitopes of these nanobodies are not cover the RBM region (marked in gray). As the results shown in the competitive assays

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Round of Screening	Input (pfu/Well)	Output (pfu/Well)	Recovery (P/Input)
lst round 2nd round 3rd round	1.28×10 ¹³ 2.4×10 ¹⁴ 2.0×10 ¹²	4.8×10 ⁶ 1.2×10 ⁶ 4×10 ⁶	3.75×10 ⁻⁷ 5.0×10 ⁻⁹ 2.0×10 ⁻⁶

 Table I Enrichment of Phage Particles Against SARS-CoV-2 RBD-Specific Nanobodies

 During Three Rounds of Panning

	WT RBD Affinity		Omicron (B.I.I.529) RBD Affinity		Affinity	
	KD(M)	Ka(M ⁻¹ s ⁻¹)	kd (s ⁻¹)	KD(M)	Ka(M ⁻¹ s ⁻¹)	kd (s ⁻¹)
BI-4	1.61×10 ⁻⁸	1.11×10 ⁵	1.79×10 ⁻³	1.10×10 ⁻⁸	1.53×10 ⁵	1.69×10 ⁻³
CI-5	3.17×10 ⁻⁷	1.83×10 ⁴	5.80×10 ⁻³	5.93×10 ⁻⁷	6.70×10 ⁴	3.97×10 ⁻²
B-B2	3.79×10 ⁻⁷	7.15×10 ³	2.71×10 ⁻³	4.90×10 ⁻⁸	5.11×10 ⁴	2.51×10 ⁻³
B2-3	3.37×10 ⁻⁹	1.20×10 ⁵	4.06×10 ⁻⁴	4.02×10 ⁻⁸	4.52×10 ⁴	1.82×10 ⁻³

 Table 2 The Results of Binding Affinity Between Nanobodies and RBD Using SPR

between nanobodies (Table 3), B-B2 and C1-5 were relevant, for which the inhibition rate was 77.22%. B-B2 and B1-4, B-B2 and B2-3 were partly relevant, with inhibition rates of 55.86% and 61.78%, respectively. The inhibition rate between the remaining combinations is less than 50%. We also tried to identify the binding epitopes of nanobodies by peptide-based ELISA, however, no statistically significant binding activities were observed between nanobodies and synthetic peptides (Figure 2C).

To assess the neutralizing activity of these nanobodies, we carried out the sVNTs that competed with hACE2 for RBD and the pseudovirus neutralization assays. The neutralizing activity of nanobodies against authentic SARS-COV-2 in vivo was not performed in the present study, because SARS-COV-2 live virus neutralization assays must be carried out in a biosafety level-3 laboratory and are time-consuming. The ELISA-based sVNT could be used to assess the ability of the nanobodies to block RBD-ACE2 interaction, and is more efficient and safer, without using viruses, and the detection can be completed in a short time under common experimental conditions. Moreover, the result of sVNTs is closely related to the cell-based neutralization assays.^{39,40} The pseudovirus neutralization assay based on the cell entry pattern similar to that of authentic viruses is also an important supplementary assay of antiviral activity.

The results of sVNTs showed that all four nanobodies could block WT RBD binding to ACE2 in a dose-dependent manner (Figure 2D). Then, we carried out the sVNTs using the Delta and Omicron (B.1.1.529) RBD, and found that the %inhibition could be maintained well (Figure 2D). We also carried out the SARS-CoV-2 WT and Omicron (B.1.1.529) pseudovirus neutralization assays with the nanobodies. The pseudovirus luciferase assay results indicated that four nanobodies neutralized WT SARS-CoV-2 pseudotyped virus with an IC₅₀ value at approximately 4.889 μ g/mL to 11.49 μ g/mL. Furthermore, although the previously dominant Omicron variant holds 15 unprecedented mutations in its RBD, the nanobodies neutralized Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ value at approximately 1.658 μ g/mL to 16.58 μ g/mL. B-B2 showed the highest activity with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus with an IC₅₀ at 1.658 μ g/mL to neutralize Omicron (B.1.1.529) pseudotyped virus be a table 4).

Development and Characterization of Multivalent Nanobodies with Thermostable Scaffold AaLS

Previously, it is reported that the binding affinity of the monomer nanobody can be improved by tandem linking.^{13,35} To further improve the affinity and neutralizing activity of nanobodies, we tried to develop multivalent nanoantibodies with SpyCatcher-modified AaLS as shown in Figure 3A. AaLS were expressed in *E. coli* in soluble form and purified. To confirm that the AaLS can maintain its ability to self-assemble into dodecahedral nanostructures, we observed the structural characteristics of the SpyCatcher-conjugated nanoparticles by negative staining TEM. As shown in Figure 3C, the SpyCatcher-modified AaLS can still form a dodecahedral sphere.

Next, we tried to assemble the scaffolds and nanobodies at different molar ratios of 1:2, and 1:1 in TBS buffer. Taking the nanobody B1-4 as an example, the SDS-PAGE showed (Figure 3B) that when nanobodies were excessive, the number of remaining scaffolds decreases obviously and more scaffolds were bound. To allow the subunits of the AaLS to be decorated with nanobodies as much as possible, we used the reaction condition of a 2:1 molar ratio and then filtered out the nanobodies that were not bound to Alas by the ultrafiltration tube. The assembly products were observed by negative-stain TEM. As we expected, the diameter of the assembled product was larger than that of the AaLS without nanobodies, and the spherical structure was maintained, which indicated that nanobodies were successfully attached to the scaffolds (Figure 3C).



Figure 2 Affinity and binding sites of neutralizing nanobodies against RBD. (A) Structure docking model of nanobodies bound to RBD using DS. The antigen-binding epitopes are highlighted by yellow color. (B) RBD sequences of SARS-CoV-2 WT with a highlighted footprint of the four nanobodies (colored in light red). (C) The results of peptide-based ELISA of monomer nanobodies. (D) Exploring the RBD-ACE2 blocking activities (%inhibition) of four nanobodies to WT, Delta, and Omicron (B.1.1.529) RBD using sVNTs. (E) Measurement of the neutralization potency of nanobodies using pseudovirus neutralization assays.

The Multivalent Nanobodies are Highly Potent

We then characterize the affinity of the assembled products of scaffolds and nanobodies by using an indirect ELISA. Taking B1-4 as an example, we named the assembled products LS-B1-4. The indirect ELISA results indicated that the binding affinity of the multivalent nanobodies was significantly higher than that of monomer nanobodies (Figure 4A).

Nanobodies	BI-4	C1-5	B-B2	B2-3
BI-4	-	43.37	55.86	46.80
CI-5	-	-	77.22	28.79
B-B2	-	-	-	61.78
B2-3	-	-	-	-

 Table 3 Inhibition Rate Between Two Nanobodies

Table 4 The IC₅₀ of Monovalent and Multivalent Nanobodies in the Pseudovirus Neutralization Assays

	WT Pseudovirus Neutralization	Omicron (B.1.1.529) Pseudovirus Neutralization
	IC₅₀(µg/mL)	IC₅₀(µg/mL)
BI-4	6.198	5.751
LS-BI-4	1.150	2.001
CI-5	4.889	4.608
LS-CI-5	3.074	3.785
B-B2	9.912	1.658
LS-B-B2	1.609	0.653
B2-3	11.49	16.58
LS-B2-3	3.822	3.788

To compare neutralizing activity between monomer and multivalent nanobodies, sVNTs were carried out. The results showed that the %inhibition of multivalent nanobodies was higher than that of monomer nanobodies at the same concentration (Figure 4B). In the pseudovirus neutralization assays, all the multivalent nanobodies exhibited better neutralizing ability compared to the monovalent form. Results showed that LS-B-B2 exhibited the highest neutralizing



Figure 3 Conjugation and structural characterization of multivalent nanobodies. (A) Schematic diagram of multivalent nanobodies construction. (B) SDS-PAGE of the assembled products of AaLS and nanobodies at different molar ratios. (C) Negative-stain TEM images of the Spycatcher modified AaLS, Scale bar, 50 nm. Negative-stain TEM images of recombinant multivalent nanobodies, Scale bar, 50 nm.



Figure 4 Comparison of the affinity and neutralizing activity of multivalent nanobodies. (A) Measurement of the binding ability of monovalent and recombinant multivalent nanobodies using indirect ELISA. (B) RBD-ACE2 blocking activities of monomer nanobodies and their multivalent pattern characterized using sVNTs. (C) Measurement of the neutralization potency of multivalent nanobodies using pseudovirus neutralization assays. (D) The epitopes recognized by multivalent nanobodies, groups that are statistically different from the control group are marked with * (E) Sequence comparison of SARS-CoV-2 variants RBD. The peptides 5 and 16 were highlighted in yellow.

ability with an IC₅₀ at 0.653 μ g/mL against pseudotyped SARS-CoV-2 Omicron (B.1.1.529) (Figure 4C), was improved compared to the monovalent form B-B2 (IC₅₀ = 1.658 μ g/mL) (Figure 2E).

The peptide-based ELISA was used to explore the epitopes of four multivalent nanobodies binding to WT RBD (Figure 4D). A total of 17 WT RBD peptides (Table 5) were selected and a Zika EDIII peptide was used as a negative control. It was found that compared with the control peptide, LS-B1-4 mainly recognized peptides 10 and 16, LS-C1-5 mainly recognized peptide 16, and LS-B-B2 recognized peptides 5, 8, 12, and 16. Of note is that peptides 5 (S381-403 GVSPTKLNDLCFTNVYADSFVIR) and 16 (S519-543 HAPATVCGPKKSTNLVKNKCVNFNF) are conserved across the variants of SARS-COV-2 Delta and Omicron (B.1.1.529) (Figure 4E), which may initial clarify the mechanisms of these three nanobodies maintaining the neutralizing potency of SARS-COV-2 variants.

Improved Thermal Stability of Multivalent Nanobodies

While nanobody is often heat resistant with the T_m from 50 to 80°C, it is not heatproof,^{41–43} obstructing it as a superior reagent for detection, diagnostic, and therapeutic applications. Many nanobodies have been engineered to increase their T_m , making them stable until exposed to extreme temperatures. Since AaLS isolated from the hyper-thermophile has excellent robustness, the scaffold is expected to improve the thermal stability of monovalent nanobodies. Taking B2-3 as an example, DSC analysis showed that the T_m of B2-3 was about 50°C, and that of LS-B2-3 was about 70°C, being increased by 20 degrees (Figure 5A).

Number of Peptides	Sequence
1	VQPTESIVRFPNITNLCPF
2	NLCPFGEVFNATRFASVYAW
3	ATRFASVYAWNRKRISNCVADY
4	YSVLYNSASFSTFK
5	GVSPTKLNDLCFTNVYADSFVIR
6	NVYADSFVIRGDEVRQIAPGQTGKI
7	VIRGDEVRQIAPGQTGKIADYNYKL
8	QTGKIADYNYKLPDDFTGCVIAWNSNNLDS
9	IAWNSNNLDSKVGGNYNYLY
10	KVGGNYNYLYRLFRKSNLKPFERDISTEIY
11	FERDISTEIYQAGSTPCNGV
12	QAGSTPCNGVEGFNCYFPLQ
13	EGFNCYFPLQSYGFQPTNGVGY
14	SYGFQPTNGVGYQPYRVVVL
15	GYQPYRVVVLSFELLHAPATVCGPK
16	HAPATVCGPKKSTNLVKNKCVNFNF
17	VKNKCVNFNFNGLTGTGVLT
18 ^a	TLHGTVTVEVQYAGT

Table 5 17 Peptides of SARS-COV-2 WT RBD

Notes: ^aNegative control.

Abbreviations: SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; COVID-19, coronavirus disease 2019; RBD, receptor-binding domain; S, spike; ACE2, angiotensin-converting enzyme 2; RBM, receptor-binding motif; CDR3, complementarity determining region 3; AaLS, the lumazine synthase isolated from the hyperthermophile Aquifex aeolicus; T_m, melting temperature; WT, wildtype; BSA, bovine serum albumin; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis; RT, room temperature; TBS, tris-buffered saline; sVNT, surrogate virus neutralization tests; SPR, surface plasmon resonance; DS, discovery studio; TEM, transmission electron microscopy; DSC, differential scanning calorimetry.

While the DSC analysis could directly measure the temperature at which the nanobodies lose their secondary structure, it is necessary to measure the activity after exposure to high temperature, since the change of binding affinity is not parallel to the result of physical parameters in some cases.²⁰ Therefore, an indirect ELISA and SDS-PAGE were used to observe the binding affinity of nanobodies after one hour of treatment at different temperature conditions (no heat treatment, 50°C, 70°C, 90°C) for an hour (Figure 5B and C). The results of indirect ELISA showed that the binding ability of B2-3 degraded obviously with the increase of temperature (no heat treatment to 70°C), while the affinity of 90°C treated B2-3 was higher than that of B2-3 without heat treatment. Although the B2-3 at 90°C has been degraded partially as observed by SDS-PAGE. By contrast, 90°C did not significantly compromise the affinity of the multivalent nanobody, LS-B2-3, it was more thermally stable and retained good binding affinity, and the protein observed by SDS-PAGE did not been substantially reduced.

In summary, our research manifests that the nanobodies attached to scaffolds are endowed with higher T_m , greater ability to refold after denaturation, and ability to function after heat exposure.

Discussion

COVID-19 has brought unprecedented threats to human health and become a global concern. The variants of SARS-CoV -2 compromise the efficacy of most vaccines or therapeutic monoclonal antibodies. High-throughput naïve library can be used as an available platform for quickly obtaining antibodies against specific antigens. To date, some nanobodies screened from naïve nanobody libraries have neutralization potencies for SARS-CoV-2.^{35,44} Here, we developed four nanobodies specific to the SARS-CoV-2 RBD and used the prokaryotic expression system to express them successfully. Since the nanobodies were isolated from nonimmune synthetic libraries, the binding affinity and neutralizing activity needed to be further improved. We modified the nanobodies.



Figure 5 Nanobodies conjugated to the nanoplatform have good thermal stability. (A) DSC analysis of B2-3 and LS-B2-3. (B) SDS-PAGE of B2-3 and LS-B2-3 after heating at different temperatures. (C) Measurement of the binding ability of B2-3 and LS-B2-3 after heating at different temperatures using indirect ELISA.

Protein cage nanoparticles isolated from AaLS have highly uniform nano-size and symmetrical structure, and they have genetic and chemical plasticity amenable to simultaneously introducing multiple cell-specific targeting ligands at desired sites according to the purpose. At present, they have been widely explored and utilized as effective delivery nanoplatforms for diagnosis and treatment in the biomedical field. AaLS is the expected antibody-mediated targeted delivery nanoplatform for us to modify antibodies to increase affinity. Since the spike proteins naturally exist as a trimer on the surface of SARS-CoV-2, there is the possibility that the multimerized nanobodies could target more than one RBD antigen at the same time, and the multimeric nanobody-antigen structure could be more stable than the monomeric nanobody-antigen complex. Consistent with the assumption, in the present study, we observed the LS-nanobodies showed better potency than monoclonal antibodies in indirect ELISA, the sVNTs, and the pseudovirus neutralizing assays.

In addition, since the nanoplatform is isolated from hyperthermophile *Aquifex aeolicus*, it may contribute to the thermal stability of the nanobodies. In the study, we observed that the nanobodies attached to AaLS can maintain binding affinity after being treated at different temperatures. The affinity of monovalent nanobodies for antigen decreased with the increase of treatment temperature. However, the affinity of monovalent nanobody B2-3 treated after 90°C is a special case. It is possible that the spatial structure of B2-3 was reconstructed due to heat-induced denaturation that exceeded its melting point, which transformed the conformational space of a CDR, opened the hidden side of its binding to the antigen, and increased the binding affinity.²² In a word, the AaLS could be used as a promising candidate for multifunctional delivery nanoplatform. Its advantages will further increase its application in disease diagnosis, prevention, and treatment.

The ACE2 receptor-binding motif (RBM) contains the majority of RBD mutations found in Omicron (B.1.1.529). The RBM binders, usually competing with ACE2 to bind to RBD, lose (or weaken) their binding and neutralizing activities to the SARS-CoV-2 variants, especially the most recent Omicron.⁴⁵ Previous studies have revealed the presence of conserved epitopes located on SARS-CoV-2 RBDs and the Omicron neutralizing potency of non-RBM antibodies.^{46,47} In this study, we explored the conservation of the nanobodies epitope on the RBD using DS molecular simulation docking. The results showed that the binding sites of B-B2 do not cover the RBM region. The peptide-based ELISA indicated that it mainly recognized the peptides 5 (S381-403) and 16 (S519-543), and these two peptides were 100% conserved across the variants of SARS-COV-2, suggesting the nanobodies recognize them might not lose most of the affinity. These results preliminarily indicated the region of antigen recognized by B-B2 were conserved epitopes, might also be a powerful agent for creating effective broad-spectrum antiviral agents. Besides, though the K_D of B2-3 was the highest in SPR using WT RBD, it did not show a better neutralizing activity. However, the neutralizing activity is not only simply determined by the affinity of the antibody for antigen binding, it is also affected by a variety of factors, such as the heterogeneity of the antibody, the antigenic determinant of the antibody targeting, and the property of binding to complement.⁴⁸

There are some limitations in this research. Firstly, the SARS-COV-2 live virus neutralization assay is the gold standard for measuring antibody neutralization ability. However, it must be carried out in a biosafety level-3 laboratory. Therefore, here, we performed the pseudovirus neutralization assay, which is also suitable for simulating the inhibition of nanobodies that inhibits the virus entry. Secondly, the mechanism of these three nanobodies maintaining pseudovirus neutralizing activity to variant strains deserves further investigation. However, the pattern of antigen-antibody interaction is very complex. In our study, the binding site prediction of the nanobody-RBD complex relies on peptide-ELISA and DS simulation docking, which lacked cryo-electron microscopy or X-ray diffraction to investigate the crystal structure parameters.

Conclusion

In summary, our study developed four nanobodies against SARS-COV-2 WT RBD from a non-immunized alpaca VHH library which may target epitopes of conserved regions of antigens, and improved the nanobodies using AaLS. Of them, LS-B-B2 exhibited the highest neutralizing ability against pseudotyped SARS-CoV-2 Omicron (B.1.1.529). Our study indicated that the AaLS could be used as an antibody-binding nanoplatform to present nanobodies on its surface and improve the potency of nanobodies. The multivalent nanobody LS-B-B2 may serve as a potential agent for the neutralization of SARS-CoV-2 variants.

Ethics Approval

Ethics committee approval was not required as IRB approval is not a requisite for this study.

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Disclosure

Dr Yuying Lu reports a patent ZLTA20220905 pending. The authors report no other conflicts of interest in this work.

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