Structural Insights into the Binding of Nanobodies Lam2 and Lam4 to the Red Fluorescent Protein mCherry

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Research Article

Keywords: nanobody, red fluorescent protein, mCherry, structure.

Posted Date: April 21st, 2021

DOI: https://doi.org/10.21203/rs.3.rs-424438/v1

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Abstract

Background

Red fluorescent proteins (RFPs) are widely used in molecular biology research, especially in deep tissues and animal models, because of their superior autofluorescence, light scattering, and phototoxicity to GFP. Although RFP can be easily monitored in vivo, improved manipulation of RFP is still desired. Using suitable nanobodies (Nbs) to bind to different epitopes of RFP is the most promising approach; thus, it is crucial to obtain structural information on how the different Nbs interact with RFP.

Results

We determined the crystal structures of the LaM2-mCherry and LaM4-mCherry complexes at 1.4 Å and 1.9 Å resolution. Our results showed that LaM2 binds to the side of the mCherry β-barrel, while Lam4 binds to the bottom of the β-barrel and does not interfere with the homo-oligomerization interface. The distinct binding sites of LaM2 and LaM4 were further verified by ITC, F-SEC and DLS assays. Our results also showed that LaM2 and LaM4 can bind simultaneously to mCherry, which is crucial for recruiting multiple operation elements to the RFP. The binding of LaM2 or LaM4 did not significantly change the chromophore environment of mCherry, which is important for fluorescence quantification assays, while several GFP Nbs significantly altered the fluorescence. Mutation of the residues of the LaM2 or LaM4 binding interface to mCherry significantly decreased the binding affinity of the Nb to mCherry.

Conclusions

Our results provided atomic resolution interaction information on the binding of Nbs LaM2 and LaM4 binding with mCherry, which is important for developing detection and manipulation methods for RFP-based biotechnology.

Introduction

Fluorescent proteins (FPs) are the most extensively studied and widely used genetic tools in molecular biology research. Fluorescent proteins can be easily expressed in almost all kinds of cells, and the fusion of fluorescent proteins generally does not affect the function of other proteins. Compared to jellyfish-derived green fluorescent proteins, red fluorescent proteins (RFPs) are ideal for in vivo imaging due to their long-wavelength excitation and their lower light scattering, autofluorescence, and phototoxicity [1–3]. Thus, for transgenic RFP or RFP fusion proteins in small animal models or deep tissues, it is easy to monitor the RFP signal by imaging approaches due to the low background noise and deeper in vivo imaging of tissues and organs. Although many genetically encoded RFP animal strains have been established to facilitate live observation, the manipulation of RFPs will remain problematic until the discovery and development of RFP-specific nanobodies (Nbs) [4, 5].

Nbs, first discovered by Hamers Casterman in 1993 [6], are single domain antibodies derived from the heavy chain variable regions (VHH) of Camelidae atypical immunoglobulins. Nbs are the smallest functional fragments derived from a naturally occurring immunoglobulin. Unlike monoclonal antibodies, Nbs can be easily produced in prokaryotic expression systems. Because of their small size (12–15 kDa) and very high stability and solubility, Nbs are widely used for industrial [7] and clinical in vitro diagnostic (IVD) applications [8]. The small size also allows Nbs to be genetically encoded as chimera proteins and delivered to cells by fusion plasmids. Typically, the long CDR3 region.
enables Nb binding to an antigen with similar high specificity and affinity [9, 10], including binding with the smooth PD-L1 protein surface [11], inserting into canyons on the HIV envelope that are not accessible to IgG [12] to neutralize a broad range of HIV-1 strains, and effectively blocking the entry of SARS-CoV-2 spike protein [13–15].

Kirchhofer et al. first developed a series of GFP Nbs that can induce subtle opposing changes in the chromophore environment [16]. The GFP-specific Nbs GBP1 (GFP enhancer) and GBP4 (GFP minimizer) were suitable for monitoring protein expression, subcellular localization, and translocation. Our previous work also showed that the chimeric GFP Nb GFP-enhancer-(GGGGS)4-LaG16 increased the binding affinity of GFP and was suitable for GFP-tagged target protein purification [17]. Tang et al. developed a GFP Nb-based system for the selective manipulation of diverse GFP-labeled cells across transgenic lines [18]. Later, Tang et al. achieved direct optogenetic control of GFP expression in neurons by Cre/loxP recombination through the binding of the GFP-specific Nb Cre chimera protein to GFP[19]. Herce et al. designed a cell-permeable Nb system to label and manipulate intracellular antigens in living cells[20]. Simpson performed PROTAC degradation of a GFP fusion protein with an anti-GFP Nb conjugated to the Halo-tag [21].

Although many GFP Nb-related protein visualization and manipulation applications have been introduced, few RFP Nbs have been reported. Fridy et al. generated a series of Nbs (named LaMs) that bind specifically to mCherry through a high-throughput screening method based on MS and cDNA libraries [22]. Prole and Taylor developed methods to visualize and manipulate intracellular signaling through GFP and GFP Nbs[23].

To develop an in vivo RFP manipulation system, the design of two or more Nbs fused with other manipulating components that can interact with different epitopes of the RFP surface at the same time is required. However, the lack of structural information on the detailed interaction interfaces between RFP and specific Nbs hinders the design and application of manipulation of RFP or RFP fusion proteins by high-affinity antibodies. Here, we determined the crystal structure of the LaM2-mCherry and LaM4-mCherry complexes and clarified the details of the binding of these two Nbs to mCherry. We also verified the simultaneous binding of LaM2 and LaM4 to RFP by a series of orthogonal molecular biology assays. Our results provide crucial atomic resolution interaction information for the further development of methods to manipulate RFP or RFP fusion proteins in vivo.

Materials And Methods

Protein expression, purification and characterization

The coding sequences of LaM2 and LaM4 were optimized based on favored codon usage in E. coli and were synthesized by Genewiz (Suzhou, China). For crystallization and binding assays, DNA encoding LaM2 and LaM4 was subcloned into the pET28a-SUMO vector with an N-terminal 6xHis tag followed by a SUMO tag or a pET21a-derived vector with an N-terminal 10xHis tag, respectively. The plasmids were transformed into E. coli strain BL21 (DE3) for expression. The bacteria were cultured in LB medium at 37 °C until the OD600 reached 0.8. Recombinant protein expression was induced by the addition of 0.2 mM isopropyl-D-1-thiogalactopyranoside (IPTG) and incubation for an additional 18 h at 18 °C. The cells were harvested and resuspended in NiA buffer containing 20 mM imidazole, 5% glycerol, 150 mM NaCl, and 100 mM Tris-HCl, pH 7.5. The His10-tagged recombinant LaM2/LaM4 and their respective mutants were initially purified by Ni-NTA affinity purification using a HiTrap HP column (Qiagen) and eluted with NiB buffer containing 300 mM imidazole, 5% glycerol, 150 mM NaCl, and 100 mM Tris pH 7.5. For crystallization, the His-SUMO tag was removed by incubation with recombinant ULP1 overnight at 4 °C. The cleaved tag fragment and ULP1 were removed by passing through a HiTrap HP column. LaM2/LaM4 were further purified by size exclusion chromatography on a Superdex75 Increase column (Cytiva), and the buffer was exchanged to gel
filtration buffer: 10 mM HEPES, pH 7.4 and 100 mM NaCl. The purity and molecular weight of the target proteins were verified by SDS–PAGE. Detailed information on the expression and purification of recombinant proteins is provided in the supplementary data.

**Site-directed mutagenesis**

Site-directed mutagenesis was carried out by employing a PCR-based mutagenesis site-directed method (Vazyme 2x Phanta Master Mix) using His10-LaM2 and His10-LaM4 as the template. The sequences of the primers used to generate these mutants are displayed in supplementary Table S1. All site-directed mutagenesis constructs were confirmed by DNA sequencing (RuiDi, Shanghai, China).

**Crystallization and data collection**

Concentrated LaM2 or LaM4 was mixed with mCherry at a molar ratio of 2:1 and incubated for 1 h at 4 °C. The mCherry/LaM2 and mCherry/LaM4 complexes were separated from the excess LaM2/LaM4 by size exclusion chromatography using a Superdex75 Increase column (Cytiva). The mCherry/Nb complex was then concentrated to 15 mg/ml in gel filtration buffer containing 10 mM HEPES, pH 7.4 and 100 mM NaCl. Crystals of the mCherry/LaM2 complex were obtained by the sitting-drop vapor diffusion method at 293 K in drops containing a mixture of 1 μl of protein solution and 1 μl of reservoir solution, which consisted of 0.2 M magnesium chloride hexahydrate, 0.1 M Tris-HCl, pH 8.5, 25% w/v polyethylene glycol 350. Crystals of the mCherry/LaM4 complex were obtained in 0.2 M lithium sulfate monohydrate, 0.1 M Bis-Tris, pH 5.5, and 25% w/v polyethylene glycol 3350.

Cryoprotection was performed by adding glycerol to the reservoir buffer at a 20% concentration. X-ray diffraction data were collected at 100 K in beamlines BL17U1 [24] and BL19U1 [25], Shanghai Synchrotron Radiation Facility, Chinese Academy of Sciences.

**Determination and refinement of protein structure**

Diffraction images were indexed and processed by HKL2000 [26]. The structures of mCherry/LaM2 and mCherry/LaM4 were obtained by molecular replacement using the Phaser program from the CCP4 crystallography package [27] with mCherry (PDB ID: 2H5Q) and a GFP Nb (PDB ID: 3K1K) as the search model. Structure refinement was performed by Refmac [28] and Phenix [29]. The model was refined by COOT [30]. The crystallographic parameters of mCherry/LaM2 (PDB ID: 6IR2; 1.39 Å) and mCherry/LaM4 (PDB ID: 6IRI; 1.92 Å) are listed in Table 1. The related figures were drawn by PyMOL [31].

**Isothermal titration calorimetry (ITC)**

The thermodynamic parameters of the binding of LaM2/LaM4 and their respective mutants to mCherry were determined by ITC using a VP-ITC calorimeter (MicroCal VP-ITC, Malvern). In a typical experiment, each titration was performed by injecting a 12 μl aliquot of protein sample (0.15 mM to 0.52 mM) into the cell containing another reactant (0.015 mM to 0.029 mM) at a time interval of 120 s to ensure that the titration peak returned to the baseline. Altogether, 23 aliquots were titrated in each individual experiment. The stoichiometry of binding (n), the association constant Ka, and the binding enthalpy H were evaluated using MicroCal Origin 7.0 software with a one-site binding model.

**Fluorescence-based Size Exclusion Chromatography (F-SEC)**
The oligomeric state of the tested samples in buffers were recorded by F-SEC. We used 100 μl 0.1 mg/ml mCherry as control. For LaM2-mCherry and LaM4-mCherry complex, 50 μl 0.2 mg/ml of mCherry (about 7 μM) and 0.2 mg/ml (about 14 μM) of LaM2 or LaM4 were mixed in equal volume (the final concentration of mCherry was 3.5 μM, and the final concentration of LaM2/LaM4 was 7μM) and incubated on ice for 1 hour. For LaM2-LaM4-mCherry complex, 50 μl 0.2 mg/ml mCherry (about 7 μM) was mixed with 25 μl 0.4 mg/ml LaM2 and 25 μl 0.4 mg/ml LaM4 and incubated on ice for 1 hour. After high-speed centrifugation, the supernatants were loaded onto a Superdex 200 Increase size-exclusion column (Cytiva) equilibrated with SEC buffer (20 mM HEPES pH 7.0, 150 mM NaCl). The fluorescence of each sample was recorded by a fluorometer (excitation, 587 nm; emission, 610 nm for mCherry fluorescence). The data were processed and normalized by FSECplotter software.

**Emission spectrum measurements**

The emission spectra of mCherry (0.1 mg/mL) and mCherry-LaM2/LaM4 (mCherry 0.1 mg/mL with excess Nb) were recorded using a fluorescence spectrophotometer (Varian Cary Eclipse). The excitation wavelength was 587 nm. The emission spectrum was recorded between 550-700 nm. The spectra data were analyzed with Origin.

**Dynamic light scattering assay**

The particle size distribution of mCherry, LaM2-mCherry complex, LaM4-mCherry complex and LaM2-LaM4-mCherry complex were measured by Nano-size-Zeta potential analyzer (Malvern Instruments, ZS90-2026), the test temperature is 25 °C, the test angle is 90 degrees.

**Results**

**The overall structure of the mCherry/LaM2 and mCherry/LaM4 complexes**

To gain insight into the binding sites of Nbs to RFPs, we purified recombinant LaM2, LaM4 and the RFP mCherry and then determined the crystal structures of the mCherry/LaM2 and mCherry/LaM4 complexes. The crystal of the mCherry complex contains mCherry and LaM2 or LaM4 at a 1:1 stoichiometry. The overall structure of mCherry/LaM2 was refined to 1.39 Å resolution, and that of mCherry/LaM4 was refined to 1.92 Å resolution. The crystallographic data are shown in Table 1. The binding interface of CDRs 1-3 of LaM2/LaM4 and mCherry was well defined. LaM2-mCherry crystallized in the space group P2_12_12_1, and the asymmetric unit contained one LaM2 Nb and one mCherry molecule. The Matthews coefficient was approximately 2.11 Å^3/Da, and the solvent content was 41.58%. LaM4-mCherry crystallized in the space group C121, and the asymmetric unit contained one LaM4 Nb and one mCherry molecule. The Matthews coefficient was approximately 2.14 Å^3/Da, and the solvent content was 42.42%.

Figure 1A shows the overall structure of the LaM2-mCherry complex, and Figure 1B shows the overall structure of the LaM4-mCherry complex. The binding sites of LaM2 and LaM4 on mCherry were different. Figure 1C shows the superposed structures of LaM2-mCherry and LaM4-mCherry. LaM2 binds to the side of the β-barrel (the 4th and 5th β-sheets of the 11 total β-sheets), while LaM4 binds to the bottom of the β-barrel (both the amino and carboxyl termini of RFP are at the bottom). The binding modes of the Nbs themselves, the binding modes are also very different. Figure 1D compares the binding of LaM2 and LaM4. Although the Nb's constant domains are similar, the CDRs are totally different. CDR3 of Nb is longer than that in IgG, and therefore, while only a loop in the IgG secondary structure typically interacts with the antigen, an α-helix in the Nb may also emerge and provide an additional interaction mode with the antigen. CDR3 and CDR1 of LaM2 contain two α-helices: residues 123-126 (Ser-Glu-Asn-
Asp) and residues 42-45 (Thr-Phe-Ser-Asp). CDR3 of LaM4 contains an α-helix consisting of residues 110-112 (Gln-Arg-Leu). Additionally, the surface potentials of LaM2 and LaM4 are quite different; LaM2 has a large negative patch in CDR1 that contributes to salt bridge formation with mCherry, while the binding of LaM4 to mCherry does not involve a salt bridge (Figure 1E).

**Details of the binding sites of LaM2/LaM4 to mCherry**

Since the resolution of both Nb-mCherry complex crystals was high enough, the binding sites between LaM2/LaM4 and mCherry were clearly defined. The detailed interaction interfaces of LaM2 and LaM4 with mCherry are shown in Figure 2. In the LaM2-mCherry complex, all the CDRs 1-3 of LaM2 contributed to the binding to mCherry. In LaM2 CDR1, a hydrogen bond of 2.9 Å was formed between Ser44 and mCherry Lys92 and Lys182. In CDR2, there was a strong hydrogen bond between Ser66 and mCherry Glu94, a weak hydrogen bond between Ser68 and mCherry Glu94, and a weak hydrogen bond between His70 and mCherry Thr 180. In CDR3, the Trp119 and Tyr120 side chains were inserted in the pocket of mCherry, and a hydrogen bond formed between Tyr120 and mCherry Thr106. In the LaM4-mCherry complex, the main interactions were in the CDR3 region of LaM4. There was a strong hydrogen bond interaction between Asn103 and mCherry Lys84 (2.07 Å). There were also strong hydrogen bonds between Asn108 and mCherry Glu10 (2.00 Å), Gly102 and mCherry Tyr38 (2.02 Å), and Leu101 and mCherry Asp81 (2.13 Å). There was also a hydrogen bond between CDR1 Arg28 and mCherry Gln188 (2.16 Å).

**Validation of the thermodynamics and binding affinity of the nanobody to mCherry by site-directed mutagenesis**

To further clarify the detailed driving forces of the binding between the Nbs and mCherry, we performed structurally guided site-directed mutagenesis and studied the binding affinity of the mutated Nbs to mCherry. We first used isothermal titration calorimetry (ITC) to measure the binding affinity and thermodynamic parameters because it is a label-free and in-solution method and is regarded as the gold standard for protein-protein interactions (Figure 3 and Table 2). Both LaM2 and LaM4 showed high binding affinity to mCherry; the Kd of LaM2-mCherry was 3.02 nM, and that of LaM4-mCherry was 22.5 nM (Figure 3A and 3B). Then, we mutated some residues that contributed to the binding of mCherry. When the two residues of LaM2 CDR1 (Ser44) and CDR2 (Ser68) that form weak hydrogen bonds with mCherry were individually replaced by Ala, the binding affinity with mCherry was only slightly reduced (Figure 3A). However, when CDR3 Trp119 and Tyr120 were replaced by Ala simultaneously, the binding with mCherry was totally abolished (Figure 3A), indicating that this region was crucial for mCherry binding. In the LaM4-mCherry complex, replacement of CDR3 Asn104 with Lys, which reversed the charge, also completely abolished the binding to mCherry, indicating that the strong salt bridge between CDR3 Asn104 and mCherry Lys84 was crucial for the binding between LaM4 and mCherry. The replacement of Asn109 with Lys only slightly decreased the mCherry binding (Figure 3B).

**Validation of the simultaneous binding of LaM2 and LaM4 to mCherry**

The crystal structure of the LaM2-mCherry and LaM4-mCherry complexes showed that the binding regions of LaM2 and LaM4 to mCherry did not overlap, so we assumed that LaM2 and LaM4 could bind to mCherry simultaneously. We confirmed this assumption by ternary ITC and F-SEC experiments. The Kd of LaM2 titrated into the LaM4-mCherry complex obtained by gel filtration was similar to that obtained for LaM2 directly titrated into mCherry (Figure 4A and Table 3). Additionally, the Kd of LaM4 titrated into the LaM2-mCherry complex is similar to that of titration into mCherry alone, indicating that the binding of LaM2 with mCherry and the binding of LaM4 with mCherry do not significantly affect each other.
We also observed the formation of a ternary complex of LaM2-LaM4-mCherry by fluorescence-based size exclusion chromatography (F-SEC) [32], which can directly show the size of the biological macromolecule complex under physiological conditions. The F-SEC results also confirmed that a stable complex of 1:1:1 LaM2-LaM4-mCherry, 1:1 LaM2-mCherry and 1:1 LaM4-mCherry formed if the proper ratio of these proteins was mixed. It is worth noting that although LaM2 and LaM4 are similar in size, there was a certain difference in the position of the peak after binding with mCherry, which may be due to the different 3D shapes of the LaM2-mCherry and LaM4-mCherry complexes.

We also determined the size distributions of mCherry alone, LaM2-mCherry, LaM4-mCherry and LaM2-LaM4-mCherry by dynamic light scattering (DLS) experiments. The results showed that mCherry alone showed very high uniformity, centered at approximately 6 nm, and formed complexes with the respective Nbs; the size increased to approximately 11-12 nm (Figure 5A).

In contrast to some GFP Nbs (GFP enhancer and minimizer) [16], the binding of Nbs LaM2 and LaM4 does not significantly affect the chromophore environment of mCherry, resulting in little effect on the fluorescence properties of mCherry (Figure 5B). This feature ensured that the optical activity of mCherry would not change significantly with the binding of LaM2/LaM4; thus, quantification by RFP fluorescence remains accurate when manipulated through the binding of LaM2/LaM4 chimeric operators.

Discusssion

The ITC results of LaM2 and LaM4 binding also provided thermodynamic information. For LaM2-mCherry binding, the total ΔG change is -19.85 kcal/mol, the main energy contribution is the change in enthalpy (ΔH -20.62 kcal/mol), and the entropy change -T ΔS is 0.78 kcal/mol. For LaM4-mCherry binding, the total ΔG change is -5.39 kcal/mol, which is smaller than that of LaM2-mCherry. Both the enthalpy change (-4.93 kcal/mol) and entropy change -T ΔS (-0.46 kcal/mol) contribute to the spontaneous formation of the LaM4-mCherry complex. Compared with LaM4, LaM2-mCherry binding is more specific because the enthalpy change is much more significant. If an application requires only one Nb, LaM2 is a better candidate than LaM4. LaM4 binding is mainly driven by hydrophobic interactions, which are relatively nonspecific compared to the hydrogen bonds and salt bridge involved in LaM2 binding. Further optimization of LaM4 may be performed to replace the entropy-driven hydrophobic interaction by designing novel specific hydrogen bonds to mCherry residues.

While the molecular weight of a Nb is only approximately one tenth that of IgG, Nbs still provide a relatively large binding interaction interface. We calculated and compared the buried surface areas of LaM2 and LaM4 to mCherry and 3 Nbs of GFP (GBP1 enhancer PDB ID: 3K1K, GBP4 minimizer PDB ID: 3G9A and LaG16, PDB ID: 6LR7) to GFP, in addition to a representative PD-L1 Nb KN035 (PDB ID: 5JDS) entering clinical trials, by PISA [11]. All of these complexes have similar buried surface areas of approximately 600-700 Å², which is comparable to that of IgG and provides high affinity and specificity. We also compared the buried surface areas of 2 haptane Nbs (CorNb-Cortisone, PDB ID: 6ITQ [33] and MTX Nb-MTX, PDB ID: 3QXV [34]). Since these haptane antigens are relatively small and cannot provide a large surface for binding, the buried surface areas are relatively small, between 300-400 Å²; however, in contrast to small buried surface areas of protein antigens, over 50% of the haptane total surface is buried, showing the effectiveness of their interactions with specific antigens.

In addition to the delivery of plasmids encoding Nbs, unlike IgG, Nbs can easily enter the cell membrane through a nonendocytic delivery system using a poly-Arg tag [20] and thus may have additional advantages over IgG-based chimeric manipulation systems.
Simulations based on the crystal structures show that LaM4 can bind to the DsRed tetramer, and the binding sites are not on the DsRed self-multimerization interface; thus, the binding of LaM4 does not affect the tetramerization. Therefore, it is possible to design chimeric proteins linking functional operation components with LaM4 and develop a self-assembling macromolecular machine based on the RFP tetramer.

**Conclusion**

In summary, we have obtained the details of how Nbs LaM2 and LaM4 bind to mCherry's different epitopes at atomic resolution by structural biology. Additionally, our thermodynamic and molecular biology assays verified the crucial residues for the Nb-RFP interaction. The binding of LaM2 or LaM4 did not significantly change the fluorescence of mCherry, which is important for fluorescence quantification assays. LaM2 and LaM4 can bind simultaneously to mCherry, which is crucial for recruiting multiple operation elements to the RFP. These results provide important basic information for the development of a LaM2/LaM4-based RFP manipulation system and provide strategies to further optimize the binding affinity of Nbs to RFP.

**Abbreviations**

FP: fluorescent protein  
GFP: green fluorescent protein  
RFP: red fluorescent protein  
ITC: isothermal titration calorimetry  
MST: microscale thermophoresis  
DLS: dynamic light scattering  
Nb: nanobody

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and materials**

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Competing interests**

We declare that we have no conflicts of interest in the authorship or publication of this contribution.
Funding

This work was supported by grants 32070939, 91527305 and 82030106 from the National Natural Science Foundation of China, an innovative research team of high-level local universities in Shanghai and a key laboratory program of the Education Commission of Shanghai Municipality (ZDSYS14005).

Authors' contributions

YD conceived and supervised the study; ZW, LL, RH and RL performed protein purification, verification, crystallization, and crystal data analysis; ZW, PZ, YZ and SC performed the protein-protein interaction assay.

Acknowledgments

We thank Prof. M. Hattori of Fudan University for the help with the F-SEC experiment. This work was supported by grants 32070939, 91527305 and 82030106 from the National Natural Science Foundation of China, an innovative research team of high-level local universities in Shanghai and a key laboratory program of the Education Commission of Shanghai Municipality (ZDSYS14005).

References


Tables

Table 1 Data collection and refinement statistics. The values in parentheses are for the high-resolution shells.

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Table 2 The binding specificity of LaM2 and LaM4 series nanobodies to mCherry by ITC
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Figure 1

Structure of mCherry: mCherry-specific nanobody complex determined by X-ray crystallography. A. Overall structure of LaM2-mCherry complex. mCherry is shown in purple. LaM2 is shown in light blue, and CDR1-3s are shown in blue, green and red (PDB ID: 6IR2). B. Overall structure of the LaM4-mCherry complex. mCherry is shown in purple. LaM4 is shown in gray, and the CDRs are shown in blue (CDR1), green (CDR2) and red (CDR3) (PDB ID: 6IR1). C. Alignment of LaM2 (light blue)-mCherry and LaM4 (gray)-mCherry complex. D. Alignment of Nbs LaM2 and LaM4. E. Comparison of the surface potential of Nbs LaM2 and LaM4 (blue: positive, red: negative).
Figure 2

The detailed binding interfaces of Nbs to mCherry. A. The detailed interactions of LaM2 CDRs with mCherry. B. The detailed interactions of LaM4 CDRs with mCherry.
Validation of the interaction between nanobodies (including site-directed mutants) and mCherry by isothermal titration calorimetry at 25 °C. A. LaM2 series nanobodies (wild type, S44A, S68A and W119A/Y120A); B. LaM4 series nanobodies (wild type, N104K and N109K).
Figure 4

ITC and F-SEC assays confirmed that LaM2 and LaM4 can bind simultaneously to mCherry. A. The ITC result of LaM2 binding to the mCherry-LaM4 preformed complex. B. The ITC result of LaM4 binding to the mCherry-LaM2 preformed complex. C. The F-SEC results of mCherry (red), mCherry-LaM2 (light green), mCherry-LaM4 (dark green) and the premixed mCherry-LaM2 complex mixed with LaM4 (blue). The right panel shows an enlargement of 14-17 ml of F-SEC to clearly see the change in size.
Figure 5

The size and fluorescence intensity of mCherry binding with nanobodies. A. The size distributions of mCherry (black), LaM2-mCherry (red), LaM4-mCherry (blue) and LaM2-mCherry-LaM4 (green) by dynamic light scattering. B. The emission spectra of mCherry (black), LaM2-mCherry (red) and LaM4-mCherry (blue).

Supplementary Files

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- mCherrySuppl20210402.docx