Crystal Structure of a Human Single Domain Antibody Dimer Formed through $V_H$-$V_H$ Non-Covalent Interactions

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Abstract

Single-domain antibodies (sdAbs) derived from human $V_H$ are considered to be less soluble and prone to aggregate which makes it difficult to determine the crystal structures. In this study, we isolated and characterized two anti-human epidermal growth factor receptor-2 (HER2) sdAbs, Gr3 and Gr6, from a synthetic human $V_H$ phage display library. Size exclusion chromatography and surface plasmon resonance analyses demonstrated that Gr3 is a monomer, but that Gr6 is a strict dimer. To understand this different molecular behavior, we solved the crystal structure of Gr6 to 1.6 Å resolution. The crystal structure revealed that the homodimeric assembly of Gr6 closely mimics the $V_H$-$V_L$ heterodimer of immunoglobulin variable domains and the dimerization interface is dominated by hydrophobic interactions.

Introduction

A typical antibody consists of two heavy-chains (H) and two light-chains (L) [1]. The N-terminal domains of both H- and L-chains are variable and are called variable regions [2], abbreviated as $V_H$ and $V_L$. Functionally, the antibodies are consisted of an antigen binding domain, Fab and an effector domain, Fc. The Fab is composed of light chain and heavy chain (reviewed by [3]). The antigen binding sites of conventional antibodies contain six complementary determining regions (CDRs), three of them from $V_H$ and three from $V_L$. The natural minimal antigen binding domain of such antibodies is composed of both $V_H$ and $V_L$. In cameldiae, a significant proportion of functional antibodies are heavy-chain antibodies which do not contain light chain [4]. The antigen binding domain of these heavy-chain antibodies is composed of only $V_H$ and is designated as $V_{H\text{H}}$ (reviewed by [5]). This discovery made it possible to isolate soluble and functional $V_H$-single domain antibody (sdAb) [6]. These sdAbs have many desirable properties from an antibody engineering point of view. They are relatively small in size with molecular weight of ~13 kDa and can be engineered to have very high affinities [7]. They can also be amplified and cloned easily because they are encoded by a single gene. In addition, these sdAbs have favorable refolding properties and biophysical stability [8]. Furthermore, they recognize epitopes that are inaccessible to conventional antibodies [9,10,11]. Finally, sdAb which is injected intravenously into mice localizes preferentially at the tumor site [12].

Similar types of sdAbs that are derived from human $V_H$ [13,14] are promising in particular for their potential use in immunotherapy because of their human origin. However, the solubility of these human sdAbs is one of the main problems. Several approaches have been reported to obtain soluble $V_H$ sdAbs [13,15], nevertheless structural information of such sdAbs is limited. The absence of light chain leads to exposure of the hydrophobic $V_H$-$V_L$ interacting interface which can cause aggregation [16]. Hence, the structural information of such human $V_H$ sdAbs is very limited. In this regard, we used a synthetic human $V_H$ library [17] to isolate a panel of soluble sdAb against human epidermal growth factor receptor-2 (HER2). The isolated sdAbs have affinities in the nanomolar range. We chose two sdAbs, Gr3 and Gr6, for further evaluation. The difference of the amino acid sequences between these sdAbs is restricted to their CDR1 and CDR3. Expression levels of both sdAbs as soluble protein were comparable. Size exclusion chromatography (SEC) analysis demonstrated that Gr3 exists as a monomer, whereas Gr6 is a dimer. To our knowledge, Gr6 is the first human-derived sdAb that is a strict homodimer. Therefore, we determined the crystal structure of Gr6 which showed that the structure mimics the $V_H$-$V_L$ pairing.

Results

Selection of HER2-specific sdAbs

A human $V_H$ phage display library [15,17] was used to select HER2-binding sdAbs as described [18] with the exception that the first two rounds of panning were performed on MDA-MB-231-Erb2 cells and the third and fourth round on the HER2/Fc protein. Ninety six randomly picked clones were tested on phage ELISA to identify clones displaying HER2-specific $V_H$, of which 25 scored positive. DNA sequencing of the 25 clones revealed 7

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different VHs, namely Gr1, Gr2, Gr3, Gr4, Gr5, Gr6 and Gr7. Gr1 was represented by 12 clones, Gr2 by 4 clones, Gr3 by 3 clones, Gr4 and Gr5 by 2 clones, and Gr6 and Gr7 by 1 clone.

Characterization of the sdAbs Gr3 and Gr6

The seven different VHs were sub-cloned in the expression vector pSJF2H [17]. After sequence verification, the sdAbs were expressed as 6xHIS-tagged soluble protein in the periplasm and purified by IMAC using a Ni-NTA column. This one-step purification resulted in more than 95% pure protein when assessed with SDS-PAGE (data not shown). The expression of Gr1, Gr2, Gr4, Gr5 and Gr7 was low and therefore were not used further. Like most of the soluble human sdAbs, Gr3 exists as a pure monomer seen as a single peak eluted at a volume corresponding to the apparent molecular mass of ~15 kDa in a SEC analysis using a Superdex™ 75 column (Fig. 1A). However,

**Figure 1. Characterization of Gr3 and Gr6 in solution.**
(A) Size exclusion chromatography of IMAC-purified Gr3 and Gr6 using a Superdex™ 75 column. (B) Amino acid sequences of Gr3 and Gr6 numbered according Kabat numbering [19]. (C) Binding of Gr3 and Gr6 to HER2/Fc analyzed by SPR.

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Gr6 which eluted at a volume that corresponds to the apparent molecular mass of ~30 kDa. This result demonstrated that Gr6 exists as a dimer in solution. Because of this difference, we chose Gr3 and Gr6, which differ only in CDR1 and CDR3 in their amino acid sequences (Fig. 1B) for further characterization.

The affinity of these sdAbs to HER2 was determined with surface plasmon resonance (SPR). Injection of sdAbs at concentration of 5 to 600 nM onto HER2-coupled surface revealed specific binding of the sdAb to the antigen. For Gr3, the actual association and dissociation curves fitted excellently to the 1:1 Langmuir binding model, giving an association rate constant ($k_a$) of $2.9 \times 10^{10}$ M$^{-1}$s$^{-1}$, a dissociation rate constant ($k_d$) of $2.0 \times 10^{-2}$ s$^{-1}$ and a dissociation equilibrium constant ($K_d$) of $6.8 \times 10^{-6}$ M (Fig. 1C). For Gr6, the $k_a$, $k_d$, and $K_d$ were $4.5 \times 10^{5}$ M$^{-1}$s$^{-1}$, $1.0 \times 10^{-1}$ s$^{-1}$ and $2.2 \times 10^{-7}$ M, respectively (Fig. 1C).

Circular dichroism (CD) profiles of Gr3 and Gr6 were determined to estimate the thermostability of the sdAbs. Thermo-induced denaturation of the protein was measured in the temperature range of 25 to 91 °C with 2 °C intervals. Plotting the CD value at 217 nm against temperature suggested a two phase denaturation (Fig. 2) with a calculated melting temperature ($T_m$) of 66 °C for Gr3 and 79 °C for Gr6, respectively.

Crystal structure of Gr6

Gr6 is a dimeric protein of 140-amino-acid residues and the crystal structure has been solved to 1.6 Å resolution by molecular replacement method (Fig. 3A). The majority of the electron density is well-defined which allows for unambiguously model building. The first three residues at the N-terminus in monomer A and the first four residues in monomer B are disordered in the map. In addition, the C-terminus of the last 12 residues in monomer A and 13 residues in monomer B, as well as the 6xHIS tag, could not be traced in the map. Kabat nomenclature [19] is used to number the positions of the amino acids and for the description of the structure. The crystallographic analysis showed that two molecules are present in the asymmetric unit in which one molecule is related to the other by a non-crystallographic 2-fold axis to form a dimer. The refined model includes residues Val2-Gly114 in monomer A and residues Gln3-Ser113 in monomer B as well as 266 water molecules. The residues for β-strand assignments are: 3-12 (strand A), 17-25 (strand B), 32-40 (strand C), 44-52 (strand C'), 56-60 (strand C''), 66-73 (strand D), 76-82b (strand E), 87-96 (strand F), and 102-112 (strand G). Validation of the structure by program SFCHECK [20] showed that the refined model is of good stereochemical quality (Table 1) and that no phi-psi angles are in the disallowed regions of the Ramachandran map.

The monomer structure of Gr6 has the same fold as the canonical immunoglobulin variable domain, consisting of two β-sheets with five and four strands. Gr6 shares 76% sequence identity with the camelized human antibody fragment cVH-E2, an inhibitor of the NS3 serine protease of hepatitis C virus [21] (PDB code 1OL0), and 62% identity with the VH domain of the human inhibitor of the NS3 serine protease of hepatitis C virus [21] (PDB code 1OL0). The CDR3 loop (residues 95-102) of Gr6 is similar to those of cVH-E2 and Fv-POT. The CDR3 loop of residues 95-102 of Gr6 exhibits a very different structure from the CDR3 loops of cVH-E2 and Fv-POT. A unique feature of the Gr6 structure is that CDR3 protrudes beyond the boundary of dimer interface and points to the other side of the other monomer.

Analysis of the dimer interface and the accessible surface area (ASA) was computed by PISA in CCP4 program package [23] and through the Protein Interactions Calculator website (http://crick. mbi.uiscernet.in/PI) [24]. The total buried ASA at the interface of approximately 2200 Å² is contributed from both monomers of ~1100 Å². The interactions between monomers are predominantly by hydrophobic contacts involving residues Val37, Leu45, Trp47, Ala50, Tyr58, Tyr91, Val93, Leu95, Pro96, Leu98, Ala101 and Trp103. Polar interaction in the interface includes 9 direct hydrogen bonds. No salt bridges and water molecules are found throughout the interface. The arrangement of the Gr6 homodimer resembles the structure of Fv-POT. Superimposition structure of Gr6 with structure of Fv-POT [22] (PDB code 1IGM) reveals that two VH domains in Gr6 structure maintain the same relative orientation between $V_H$ and $V_L$ in the structure of Fv-POT (Fig. 3C).

Discussion

Single-domain antibodies are very attractive tumor-targeting tools for their natural properties of small size, solubility and high
Figure 3. Structure of Gr6 and comparison of Gr6 with cVH-E2 and Fv-POT. (A) Ribbon diagram of the Gr6 structure shown in cyan (chain A) and red (chain B). CDRs are colored in green in chain A and orange in chain B. The β-strands are labeled by following the Kabat nomenclature. (B) Ribbon diagram of the superimposed Fv-POT Heavy chain onto the monomer of Gr6 (left) and the superimposed cVH-E2 monomer onto Gr6 monomer (right): cyan, Gr6; light pink, Fv-POT heavy chain; light blue, cVH-E2. (C) Ribbon diagram of the superimposed Fv-POT and Gr6 dimer: light pink, Fv-POT; cyan, Gr6. doi:10.1371/journal.pone.0030149.g003
permeability into tissues. An important feature of camelid sdAbs is their high thermodynamic stability [8]. This feature not only contributes to the high expression yield, but may also be required for their in vivo tumor targeting ability [25]. In this study, we characterize two anti-HER2 sdAbs, Gr3 and Gr6, from a semi-synthetic human VHlibrary. The Tms calculated from CD experiments for both Gr3 and Gr6 are in the same range as that of the camelid sdAb. Therefore, we presume that these human sdAbs are as stable as camelid sdAb. The single domain antibodies are also known for their monomeric behavior. In solution, Gr3 exists as a monomer; however, Gr6 has an unusual antibody fragment cVH-E2, a competitive and reversible inhibitor of the NS3 protease of the hepatitis C virus, is a concentration-dependent dimer which occurs in solution at protein concentration > 7 mg mL⁻¹ [21]. The dimerization of cVH-E2 should be the consequence of the selected CDR3 loop because the molecule was affinity-selected from a library in which CDR3 is randomized, implying that CDR3 should be involved in the dimerization [21]. However, in the crystal structure of cVH-E2 dimer, the CDR3 loops are at the opposite sides of the molecule and do not participate in the inter-domain interaction [21]. Therefore, the crystallographic structure of cVH-E2 is probably a result of crystal packing and may not represent the observed dimer in solution. In this work, the Gr6 crystal structure shows that the VH-VH interface mimics the classical association of VH-VL dimer, strongly suggesting that Gr6 is a strict dimer and that the structure does represent the dimerization in solution.

The Gr6 structure present here not only provides information on how the dimer is associated, but also confirms that the dimerization of the sdAb cannot avoid the traditional VL-binding interface. More studies are required to understand whether Gr3/Gr6 can block the dimerization of HER2, whether Gr3/Gr6 can specifically target high HER2 expression cells, and how Gr3/Gr6 interacts with HER2. Although trastuzumab (Herceptin, Genentech Inc., San Francisco, CA) is an approved therapy for treating HER-2 dependent cancers, the development of trastuzumab resistance occurred within one year after the patients received the initial response to trastuzumab [30]. One of the problems with trastuzumab is that it does not efficiently block HER2 from dimerizing with other HER family members. Therefore, Gr3 and Gr6 could be alternative HER2-targeted antibodies that possess the advantage of accessibility to hidden antigens that are not easily reached by whole antibodies. The characterization and structural information obtained in this work therefore will be beneficial to the future design of such humanized sdAbs in the HER2-dependent cancer therapy.

### Materials and Methods

#### Selection of binders

The synthetic phagemid-based human VH library used in this study has been previously described [15,17]. VH repertoire was
Figure 4. Hydrophobic interface of Gr6 dimer. (A) View of side chain arrangement at the V\textsubscript{H}-V\textsubscript{H} hydrophobic interface of Gr6 homodimer. One Gr6 monomer is shown in cyan and the other one is shown in gray. (B) Surface representation of V\textsubscript{H}. Residues involved in dimer formation are highlighted in yellow. (C) Superposition of residues at V\textsubscript{H}-V\textsubscript{H} interface of Gr6 with that at V\textsubscript{H}-V\textsubscript{L} interface of Fv-POT to show the structural resemblances of the residues that are located at framework-2 involved in the dimer formation.

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expressed on phage after infection with M13K07 helper phage as described [18]. Specific V\(_{H}\) against HER2 were enriched by four rounds of in vitro selection. In the first two rounds MDA-MB-231-ErbB2 cells, a kind gift from Dr. Maria Jaramillo, NRC Canada, were used as the antigen. Three wells each with 1.5 million cells were incubated with 5\(\times\)10\(^{11}\) phages displaying V\(_{H}\). For the 3\(^{rd}\) and 4\(^{th}\) round of panning human HER2 (ErbB2)/Fc chimera (R&D Systems Inc, Minneapolis, MN) at the concentration of 100 \(\mu\)g mL\(^{-1}\) was used. 100 mM triethylamine at pH 11.0 was used to elute bound phage which were immediately neutralized with 1 M Tris-HCl at pH 7.4 and were used to infect exponentially growing TG1 cells.

Individual colonies obtained after fourth round of panning were tested against HER2 in a phage ELISA according to the standard procedures. Briefly, clones were grown in 2 \(\times\) YT medium + ampicillin (100 \(\mu\)g mL\(^{-1}\)) 0.1% glucose to OD\(_{600}\) = 0.3 - 0.5, and infected with M13K07 helper phage (37 \(^{\circ}\)C no shaking, 30 min) followed by addition of kanamycin (50 \(\mu\)g mL\(^{-1}\)) and incubation overnight (37 \(^{\circ}\)C with shaking). Cultures were centrifuged (4000 rpm, 20 min, 4 \(^{\circ}\)C) to pellet the cells. Subsequently 100 \(\mu\)L of supernatant containing recombinant phage particles were added to HER2 (ErbB2)/Fc chimera pre-coated microtiter plate wells. After a 2 h incubation at 37 \(^{\circ}\)C, microtiter plate wells were washed three times with PBS plus 0.1% (v/v) Tween 20 followed by addition of 1:5000 diluted anti-M13 HRP conjugate (GE Healthcare, Baie d’Urfe, QC). V\(_{H}\)-phages bound to HER2 were detected by the addition of 100 \(\mu\)L of HRP substrate. After a 15 min incubation the reaction was stopped by adding 100 \(\mu\)L of 1 M H\(_2\)PO\(_4\) and absorption at 405 nm was measured.

Expression and purification of sdAbs

The V\(_{H}\) genes of the clones that scored positive in phage ELISA were sequenced to determine the diversity of the isolated clones. They were re-cloned into the expression vector pSJF2H [30] using the restriction enzymes BssI and BamHI. The ligated plasmids were used to transform E. coli cells. After sequence reconfirmation, recombinant sdAbs proteins were expressed in the periplasm and purified by immobilized-metal affinity chromatography (IMAC). Briefly, clones were inoculated overnight in 25 mL LB with 100 \(\mu\)g mL\(^{-1}\) ampicillin at 37 \(^{\circ}\)C and 200 rpm. Twenty milliliter of the culture was transferred to 1 L of M9 medium (0.2% glucose, 0.6% Na\(_2\)HPO\(_4\), 0.3% KH\(_2\)PO\(_4\), 0.1% NH\(_4\)Cl, 0.05% NaCl, 1 mM MgCl\(_2\), 0.1 mM CaCl\(_2\)) supplemented with 0.4% casamino acids, 5 \(\mu\)L mg\(^{-1}\) vitamin B1 and 100 \(\mu\)g mL\(^{-1}\) ampicillin and incubated for 24 h. Hundred milliliter of 10 \(\times\) TB nutrients (12% Tryptone, 24% yeast extract and 4% glycerol), 2 mL of 100 \(\mu\)g mL\(^{-1}\) ampicillin and 1 mL of 1 M isopropyl-\(\beta\)-D-thiogalactopyranoside (IPTG) were added to the culture and incubation was continued for another 55 - 70 h at 28 \(^{\circ}\)C and 200 rpm. Cells were then centrifuged and the pellets were lysed with lysozyme. Cell lysates were centrifuged and supernatants were loaded onto 5-mL HiTrap\textsuperscript{TM} chelating HP affinity columns (GE Healthcare). After washing the columns with four column volume of wash solution (10 mM HEPES buffer, 500 mM NaCl, 20 mM imidazole, pH 7.5), His-tagged proteins were eluted with a linear gradient (10 to 500 mM) of imidazole and the eluted proteins were assessed for purity by SDS-PAGE and subsequently dialyzed in PBS buffer. The aggregation status of sdAbs was assessed by Superdex\textsuperscript{TM} 75 size exclusion chromatography (SEC) as described using PBS as the equilibration buffer [31].

Thermostability determination of sdAbs

To determine the circular dichroism (CD) profile and thermostability of the sdAbs, proteins were first buffer-exchanged in 10 mM phosphate buffer, pH 7.0, by a Superdex\textsuperscript{TM} 75 SEC. The peaks for the sdAbs were collected, and the proteins were used for CD analysis. CD spectra were collected from 250 to 200 nm at protein concentrations of 2.5 \(\mu\)M in a 10 mm quartz cuvette with a J-650 CD spectropolarimeter (JASCO, Easton, MD). Data were collected with a band width of 1.0 nm and a scanning speed of 50 nm min\(^{-1}\) with two accumulations of scans to determine the CD profile of the proteins. Under the same conditions but with a single data accumulation, CD spectra were automatically measured at 2°C intervals from 25-91°C to determine thermal denaturation of the protein at a temperature shift speed of 1°C/min. Ellipticity at 217 nm were plotted against temperature and melting temperatures (T\(_{m}\)) were calculated from Boltzmann sigmoidal equation using GraphPad Prism software.

Affinity measurement of sdAb by surface plasmon resonance (SPR)

The binding of sdAbs to HER2 was determined by SPR using a Biacore 3000 (GE Healthcare). HER2 and ovalbumin (as reference protein, Sigma) were immobilized on research grade sensor chip CM5 (GE Healthcare). Immobilizations were performed with an amine-coupling kit (GE Healthcare) and carried out at 50 \(\mu\)g mL\(^{-1}\) HER2 in 10 mM acetate, pH 4.0 (GE Healthcare) and 50 \(\mu\)g mL\(^{-1}\) of ovalbumin in 10 mM acetate, pH 4.5. A volume of 120 \(\mu\)L of the sdAbs at concentration of 5 nM to 600 nM were injected over the surfaces at a flow rate of 40 \(\mu\)L/min. Analyses were carried out at room temperature in HBS-EP (10 mM HEPES pH 7.4, 150 mM NaCl, 3 mM EDTA and 0.005% surfactant P20 (GE Healthcare). Regeneration was performed with running buffer (HBS-EP). Data were analyzed with BIAnalyst software 4.1 (GE Healthcare).

Crystallization and data collection

Various commercial kits of Hampton Research and Emeralds BioSystems were used for initial screening of crystallization conditions for Gr6 protein, by vapor diffusion method performed with a Honeybee 961 robot (Digilab Genomic Solutions). Gr6 crystals were obtained in sitting drops mixing 0.5 \(\mu\)L of protein (30 mg mg\(^{-1}\) in 10 mM HEPES pH 7.0, 100 mM NaCl) with 0.5 \(\mu\)L of precipitant solution (100 mM Bicine buffer, pH 9.0, 20% (v/v) PEG 6000). Temperature was an important factor for the growth of Gr6 crystals. The plate-shaped crystals grew in the way that crystallization trays were incubated at 25°C for 2 days, and then shifted to 4°C. Crystals grew to maximum dimension ~ 0.30\(\times\)0.50\(\times\)0.03 mm within 2 weeks. Crystals in the mother liquor were flash-frozen in liquid nitrogen prior to data collection.

The native data set was collected to 1.6 \(\AA\) resolution with the wavelength \(\lambda=1.0 \text{\AA}\) at beamline 13BI of the National Synchrotron Radiation Research Center, Hsinchu, Taiwan. Crystals were kept at 100 K during data collection. Data were indexed, integrated and scaled with HKL2000 [32]. The space group is P2\(_1\)\(_1\)\(_2\)_1 with the unit cell dimensions a = 39.81 \(\AA\), b = 76.99 \(\AA\) and c = 81.56 \(\AA\). The Matthews’s coefficient is 2.34 \(\text{\AA}^3\text{Da}^{-1}\) with two molecules per asymmetric unit corresponding to solvent content of 47.4%. Crystal parameters and data collection statistics are summarized in Table 1.

Structure determination and refinement

The Gr6 structure was solved by molecular replacement with the program Phaser [33], using data between 20 and 2.5 \(\AA\). The monomer structure of the camelized human antibody fragment 4V5-E2 (PDB code 1OL0) [21] was used as the search model and two copies of the molecule were applied in the search. The best
solution from Phaser had a log-likelihood gain of 805.3. The model was rebuilt with the program Coot [34] with the guidance of the 2Fo-Fc and Fo-Fc electron density maps. Refinement of the model was carried out with the program CNS [35] without any non-crystallographic symmetry restraints. Multiple cycles of simulated annealing, positional, and individual isotropic B factor refinements against data to 1.6 Å resolution were performed with CNS, alternating with manual model rebuilding in Coot. 6.7% of the data were flagged for cross-validation during refinement. Total of 266 water molecules were included in the final model and final stage of the refinement. The refinement statistics are summarized in Table 1.

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**Author Contributions**

Conceived and designed the experiments: TBN SW JZ. Performed the experiments: TBN SYC. Analyzed the data: TBN SW. Contributed reagents/materials/analysis tools: SL JT MAG. Wrote the paper: TBN SW JZ.