

Article



# **Crystal Structure of a Chimeric Antigen Receptor (CAR) scFv Domain Rearrangement Forming a VL-VL Dimer**

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Abstract: Chimeric Antigen Receptor T-cell (CAR-T) immunotherapies are dependent upon designed transmembrane proteins to bind target antigens and stimulate an immune response. The success or failure of these CARs is only partially predictable, yet recent work has highlighted the importance of antigen binding scFvs driving distinct oligomerization states with varied CAR-T efficacy. Here, we sought to determine the extracellular structure of the anti-CD19 CAR 47G4-CD828Z. Unexpectedly, the resolved crystal structure revealed an IgVL homodimer bound along an inverted VL | VL interface. We found that the VL-VH linker, designed to be cleavage resistant, was cleaved, and the VH and CAR hinge domains were absent from the crystal structure lattice. Molecular Dynamics simulations revealed that the inverted VL | VL interface was more stable than the canonical VL | VL configuration. Our work substantiates the need to interrogate the scFv structure and CAR oligomerization state for optimal CAR-T design.

Keywords: CAR-T cell; immunotherapy; chimeric antigen receptor; VH-VL interface; Ig domains

## 1. Introduction

Naturally occurring tumor-reactive lymphocytes (T-cells) have a remarkable capacity to react to pathogens [1], and a central direction in cancer immunotherapy has been to harness patients' own (autologous) T-cells to fight their cancer through "Adoptive Cell Transfer" (ACT) [2,3]. One form of ACT, termed Chimeric Antigen Receptor (CAR) T-cell therapy, extends T-cells' ability to bind specific target antigens expressed on the surface of tumor cells and initiate a cascade of cellular events that results in clearing tumors [4,5].

CAR T-cell therapy involves isolating T-cells from the patient's blood (autologous T-cells) and genetically engineering them to express a CAR protein on its surface. These CAR-T cells (CAR-Ts) are expanded in large numbers in vitro and infused back into the patient to attack cancer cells. Moreover, as these CAR-Ts expand and persist in the patient they serve as a long-term "living drug" to keep the tumor in check [6].

A chimeric antigen receptor (CAR) chains together different domains of native cell surface receptor proteins of the immune system to integrate a set of specific functions from binding a target cell surface antigen to signal transduction and T-cell stimulation. Current "second generation" CARs [4] are composed of antibody variable domains (scFv) targeting a specific antigen, followed by a hinge (also called spacer) connecting to a transmembrane helical domain, both typically extracted from a single pass transmembrane protein such as CD8 or CD28, followed by an intracellular costimulatory domain extracted from CD28



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**Copyright:** © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). or 4-1BB, and lastly a signaling domain extracted from CD3z, as shown schematically in Figure 1. The optimization of CARs built from domains extracted from native cell surface receptors is a very active field of research [4,5].



**Figure 1.** 47G4-CD828Z CAR domains and the VL-VL dimer. Cartoon of CAR domains, represented as an ideal dimer. A single chain is composed of two immunoglobulin domains VL and VH from the 47G4 antibody, chained together in that order as an scFv followed by the CD8a hinge and transmembrane regions, and costimulatory domains extracted from CD28 and CD3zeta. The CAR name describes its composition in abbreviated form 47G4-CD828Z.

With the ability to recognize antigens expressed on the surface of tumor cells, anti-CD19 CAR-Ts target the cell surface CD19 molecule expressed on many B-lineage leukemias and lymphomas. They have been remarkably successful in treating relapsed/refractory disease, with two FDA approved therapies: CTL-019 (KYMRIAH, Novartis Pharmaceuticals Corp.) and axicabtagene ciloleucel (YESCARTA, Kite Pharma, Inc., Los Angeles, CA, USA) [4,7–10]. This success has inspired very strong interest in developing CARs against a broader range of cell-surface antigens, especially against solid tumors that represent 90% of cancers [11]. A total of 89 unique antigens (50 for hematological cancer, 57 for solid tumors, and 18 common to both) have been used as targets of CAR-Ts in over 800 ongoing clinical studies worldwide (https://clinicaltrials.gov/ct2/results/details?term=CAR-T+OR+%2528CAR-T+cell+therapy%2529&recrs=abdf, accessed on 19 April 2023), and more trials are projected in coming years [12].

While CAR-Ts hold great promise in the clinic, the biochemical and biophysical basis for what makes an effective CAR protein remains largely unknown. In fact, no intact structure of any single pass membrane protein on which these chimeric molecules are based has been determined to this day [13]. We sought initially to obtain a 3D structure of a CAR extracellular domain using X-ray crystallography.

#### 2. Materials and Methods

#### 2.1. Protein Purification and Crystallization

DNA coding for the extracellular region of the 47G4-CD828Z CAR was synthesized (Genscript, Piscataway, NJ, USA) and cloned into the mammalian cell expression vector pcDNA3.1(+) between the BamH1 and Xho1 sites of the polylinker under the control of the human cytomegalovirus (CMV) promoter. The construct began with a native CD8 leader sequence at the N-terminus to direct secretion and ended with TEV protease-cleavable

octahistidine affinity tag at the C-terminus to aid in protein purification. The protein was expressed by transient transfection of Expi293 cells at high density using polyethylenimine [14]. Conditioned media containing secreted protein was harvested 4 days posttransfection, cleared of cells by centrifugation, and exchanged by cross-flow diafiltration with binding buffer containing 20 mM tris(hydroxymethyl)aminomethane (Tris) pH 7.5, 150 mM NaCl, and 40 mM imidazole. The 47G4-CD828Z CAR was purified with tandem nickel affinity and gel filtration chromatography using an AKTAexpress system (Cytiva, Marlborough, MA, USA). The elution buffer was of the same composition as the binding buffer except it contained 0.5 M imidazole and the gel filtration buffer contained 20 mM Tris pH 7.5 and 150 mM NaCl. Protein fractions were pooled and polyhistidine-tagged TEV protease [15] was used to cleave the C-terminal His<sub>8</sub> tag from the CAR. The cleaved protein was passed through Ni-NTA agarose resin (Qiagen, Hilden, Germany) to remove the protease and tag. Protein purity and integrity was verified using SDS-PAGE prior to concentration for sparse-matrix crystallization screening. Sitting-drop vapor diffusion experiments were performed using the JCSG Core I-IV (Qiagen) and Wizard Precipitant Synergy (Molecular Dimensions, Sheffield, England) screens and slow-growing crystals nucleated in one condition after 3 weeks of incubation at room temperature. Crystallization was further optimized in the conditions given in Table 1.

	47G4-CD828Z CAR
Method	Vapor diffusion, sitting drop
Plate type	Intelli-Plate 96-3 LVR
Temperature (K)	295
Protein concentration (mg mL $^{-1}$ )	11
Buffer composition of protein solution	20 mM Tris pH 7.5, 150 mM NaCl
Composition of reservoir solution	0.10 M Tris pH 8.5, 1.8 M ammonium sulfate, 6.5% ( $w/v$ ) PEG 400, 0.2 M magnesium sulfate
Volume and ratio of drop	400 nL total, 1:1 ratio
Volume of reservoir (µL)	60

Table 1. Crystallization.

#### 2.2. Data Collection and Processing, Structure Solution and Refinement

Crystals were flash frozen in liquid nitrogen after brief soaking in 3.4 M sodium malonate pH 8.5 (Hampton Research, Aliso Viejo, CA, USA) and screened in-house for diffraction. In order to improve diffraction, the crystals were annealed twice by blocking the cryostream for 1 s each annealing cycle prior to shipment to the synchrotron in the frozen state. Data were collected at beamline 19-ID (NYX) at the National Synchrotron Light Source II (Brookhaven National Laboratory, Upton, NY, USA), indexed and merged using HKL2000 [16], and converted to structure factors using CCP4 [17]. The structure was solved with molecular replacement using Phaser [18], searching independently with ensembles of isolated N- and C-terminal immunoglobulin domains from 8 different scFvs in the PDB (codes 6EJG, 4YJZ, 3AUV, 1KTR, 6I07, 1H8N, 5OGI, and 6J71). Two copies of only the N-terminal V<sub>L</sub>-derived domain of the extracellular domain of the 47G4-CD828Z CAR were found in the asymmetric unit. The model was automatically rebuilt using ARP/wARP [19] of the CCP4 package followed by manual model building in COOT [20] and refinement in Phenix [21]. Two molecules of PEG and a molecule of malonate were found ordered near crystal contacts. Data collection and processing, and structure solution and refinement statistics are listed in Tables 2 and 3, respectively.

	47G4-CD828Z CAR
Diffraction source	NSLS II 19-ID
Wavelength (Å)	0.979
Temperature (K)	110
Detector	ADSC HF-4M
Crystal-to-detector distance (mm)	202
Rotation range per image (°)	0.2
Total rotation range (°)	360
Exposure time per image (s)	0.1
Space group	C2
<i>a, b, c</i> (Å)	79.3, 79.2, 40.6
α, β, γ (°)	90, 109.6, 90
Mosaicity (°)	0.6
Resolution range (Å)	38.27–1.40 (1.45–1.40)
Total No. of reflections	521,969 (44,484)
No. of unique reflections	46,404 (4550)
Completeness (%)	99.8 (98.2)
Multiplicity	11.2 (9.8)
CC <sub>1/2</sub>	1.0 (0.621)
Average $(I)/\sigma(I)$	20.2 (1.4)
R <sub>merge</sub> (%)	7.1 (126.3)
Overall <i>B</i> factor from Wilson plot ( $Å^2$ )	24.7

Table 2. Data collection and processing. Values in parentheses are for outer shell.

## 2.3. Structure Prediction

Alphafold2 predictions [22–24] were used to produce models of scFv monomers. Alphafold-multimer was used to produce models of scFv dimers and VL-VL conformations. The canonical VL-VL configuration structure was predicted by AlphaFold2 using a max template date of 10 May 2022. AlphaFold2 predicted 25 structural models; all but one (of low quality) were in the canonical VL-VL configuration (see Supplement File S3).

#### 2.4. Molecular Dynamics Simulations

Both the inverted and canonical VL-VL dimer configurations were studied by MD simulation. The inverted VL-VL configuration structure was taken from the PDB crystal structure in this study (PDBid: 7JO8). The canonical VL-VL configuration structure was selected as the top-ranked AlphaFold2 model. MD simulations were conducted using NAMD2 following a similar protocol used previously [25]. Briefly, both systems were solvated in TIP3P water boxes with 100 mM of NaCl concentration. Protein parameters were assigned from the CHARMM36 force field. Production simulations consisted of 100 ns NPT equilibrium MD simulations. The trajectory Root Mean Square Deviation of the atomic positions (RMSD) was computed using Gromacs tools.

## 2.5. Steered Molecular Dynamics Simulations

Constant velocity SMD simulations were conducted using NAMD2 software. We ran 20 SMD replicas for each system represented by 20 structures taken every 2.5 ns across the last 50 ns of the equilibrium MD simulation. The chain A VL domain was held fixed by  $C\alpha$  atoms of residues 23, 34, 38, 48, 63, and 89, while the chain B VL domain was pulled by all

heavy atoms at a constant velocity of 1 Å/ns with a pulling spring constant of 7 kcal/mol. The two VL-VL dimers were assumed to be unbound when the interaction energy reached 0 kcal/mol; we note that the same affinity trend existed when we defined the unbound state as an absence of native bound-state contacts ( $-64.9 \pm 0.8$  kcal/mol inverted vs.  $-46.4 \pm 1.2$  kcal/mol canonical).

**Table 3.** Structure solution and refinement.

	47G4-CD828Z CAR
PDB code	7JO8
Resolution range (Å)	38.3–1.40
No. of reflections, working set	46,442
No. of reflections, test set	2259
Final R <sub>cryst</sub>	0.155
Final R <sub>free</sub>	0.171
Cruickshank DPI	0.051
No. of non-H atoms	
Protein	1770
Ligand	36
Water	215
Total	2021
Solvent content (%)/Za	51.7/2
R.m.s. deviations	
Bonds (Å)	0.011
Angles (°)	1.28
Average <i>B</i> factors ( $Å^2$ )	24.7
Protein	23.1
Ligand	42.8
Water	34.7
Ramachandran plot	
Most favoured (%)	96.71
Allowed (%)	3.29
Outliers (%)	0.00

## 3. Results

## 3.1. CAR Extracellular Construct

We generated constructs of the extracellular domains of the anti-human B-cell CD19 CAR 47G4-CD828Z [26] linking the VL and VH domains of the fully human anti-CD19 antibody [27], also called Hu19, to form an scFv in a VL to VH orientation connected by a linker peptide, followed by a CD8 hinge [10] to cover the entire extracellular region (See Figures 1 and 2E). Importantly, the linker peptide (GSTSGSGKPGSGEGSTKG) connecting VL to VH was originally designed for "reduced aggregation and enhanced scFv proteolytic stability" [28]. This construct produced crystals suitable for diffraction analysis. The crystal structure obtained unexpectedly formed a VL-VL homodimer with an inverted (flipped) interface, unlike the well-known Bence-Jones protein (1REI) that forms as a VL-VL homodimer with a canonical IgV dimer interface (see Figures 2 and 3 and Table S1 for sequence information).



MALPVTALLLPLALLLHAARPEIVLTQSPGTLSLSPGERATLSCRASQSVSSSYLAWYQ QKPGQAPRLLIYGASSRATGIPDRFSGSGSGTDFTLTISRLEPEDFAVYYCQQYGSSRF TFGPGTKVDIKGSTSGSGKPGSGEGSTKGQVQLVQSGAEVKKPGSSVKVSCKDSGGTFS SYAISWVRQAPGQGLEWMGGIIPIFGTTNYAQQFQGRVTITADESTSTAYMELSSLRSE DTAVYYCAREAVAADWLDPWGQGTLVTVSSFVPVFLPAKPTTTPAPRPPTPAPTIASQP LSLRPEACRPAAGGAVHTRGLDFACDENLYFQGHHHHHHHH

**Figure 2.** 47G4-CD828Z CAR domains and the VL-VL dimer. (**A**) VL-VL homodimer observed in the crystal with an inverted configuration. VL domains' CDR1 colored in purple, CDR2 in blue, and CDR3 in green. (PDB:7JO8 3D visualization link: https://structure.ncbi.nlm.nih.gov/icn3d/share. html?mMva3qCc76kn11LG6, accessed on 19 April 2023). (**B**) Cartoon of the scFv moiety, formed by linking a VL and a VH domain with a canonical VL | VH interface between the two Ig domains (**C**) Cartoon of the 47G4 VL-VL homodimer formed in the crystal with an inverted configuration. (**D**) Cartoon of a Bence-Jones VL-VL homodimer with a canonical interface (similar to a VL | VH heterodimer) for comparison (see Figure 3 for details). (**E**) Protein sequence of the extracellular region of the of the anti-CD19 47G4 CAR with the CD8 secretion signal at the N terminus (black), the VL domain (pink), the linker (red), the VH domain (blue), followed by the CD8a hinge (black) and the His8-tag.

The intact extracellular region was purified and used in crystallization; it is apparent that the flexible linker joining the VL and VH domains of the scFv portion of the CAR was proteolyzed during crystallization, despite its intended design [28], since the unit cell dimensions and lattice do not provide the space for the other regions of the extracellular domain to be present if they had been disordered. Subsequent sparse-matrix crystallization screening of purified protein supplemented with protease inhibitors failed to yield other crystallization hits.

The observed VL-VL structure, obtained after proteolysis of the linker, raises questions surrounding scFv linkers, variable domain swapping, and scFv oligomerization, and, therefore, regarding CAR oligomerization made possible through their IgV domains rearrangements. While the significance of this observed structure in the context of a full CAR, especially in vivo, has to be established, it points to possible domain swaps in scFv dimers.



Figure 3. Comparison of the inverted VL-VL dimer vs. the Bence-Jones VL-VL dimer. (A) Canonical dimer interface using the GFCC' sheet observed in the Bence-Jones protein (1REI): The observed quaternary axis of symmetry is vertical. This is the classical interface of antibody VH-VL domains with the CDRs on the same side. It is also found in the Bence-Jones protein as a VL-VL: 3D visualization of the Bence-Jones protein with a canonical interface iCn3D-1REI homodimer (VL-VL). (B) Inverted dimer interface using the GFCC' sheet observed in the CAR VL-VL dimer (7JO8): The observed quaternary axis of symmetry is horizontal coming out of the page plane. It corresponds to a flip (180 degrees rotation) of one VL domain vs. the other. In this case, CDRs are on opposite sides; the C terminus G strands are pointed in opposite directions. The figure shows the structure of a VL-VL dimer interacting through the GFCC' sheet in antiparallel iCn3D-7JO8 inverted homodimer. The second row (C,D) represents a schematic representation of the variable domains in both dimers. The third row (E,F) compares the interactions between the GFCC'C" sheet between the two dimers. The Bence-Jones protein symmetric dimer interface is similar to a canonical pseudosymmetric VH-VL interface, forming in particular H-bonds between residues Q38 in symmetric positions. The inverted VL-VL dimer interface forms a tighter symmetric interface with 4 residues forming H-bonds (green) (Q6-R96 Q39-Q90), aromatic interactions (blue) (Y88-F99, F99-F99), and aromatic cation interaction (red) (F97-K104). See 3D visualization: comparison of the two VLVL dimer interfaces. Coloring of Ig strands using the full rainbow spectrum (iCn3D: strands A/A': dark violet, strand B: light violet, strand C: dark blue, strand C': light blue, strand C'': dark green, strand D light green, strand E: yellow, strand F: orange, strand G: red. Loops in grey.

## 3.2. scFv Linkers and Oligomerization through Domain Swapping

CAR designs to date have assumed that scFvs are in a monomeric form within a single CAR chain in order to bind one target. Yet, scFvs are prone to dimerization and

aggregation, and the linker length between the VH and VL domains impacts scFv formation and, consequently, CAR formation. scFv oligomerization states are very sensitive to VL-VH linkers and can dimerize to form diabodies. Reducing the scFv linker length to 3–5 residues can lead to the formation of (dimeric) diabodies vs. (monomeric) scFvs through domain swapping [29–31] to form two canonical VL | VH interfaces to bind their targets. In the context of CAR-Ts, a recent study has demonstrated the importance of the scFv linker length in anti-CD22 CARs [32,33], differing only in the linker length with a drastic effect on the clinical outcomes [32,33]. The two CARs derived from the m971 anti-CD22 monoclonal antibody [34,35] differ in linker length, and it is known from X-ray crystallography that the short (5 residue) linker forms a diabody [36]. Short linkers are sometimes called Winter linkers [30].

We performed a size exclusion chromatography (AnSEC) study of the extracellular region 47G4-CD828Z CAR (Figure 2E) that confirms the formation of an oligomer, running primarily as a dimer with a small peak as a tetramer (see Figure S1). Yet, given that our crystals led to the formation of a single domain VL-VL dimer after cleavage of the linker, we are left with the question regarding the mode of dimerization for the 47G4 VL-linker-VH scFv.

#### 3.3. Predicted scFv Dimers in Domain-Swapped or Strand-Swapped Forms

In the absence of an experimental structure of an intact CAR or simply of an intact CAR extracellular region, we can interrogate monomeric or dimeric forms through either structure prediction and/or homology model building. Alphafold2 predictions [22–24] show swapped diabody forms of the scFvs, regardless of the linker length. These are canonical (parallel) IgV | IgV interfaces and are plausible from a structure prediction viewpoint. However, our observed crystal structure shows an inverted VL-VL interface (Figure 2).

On the other hand, the PDB database offers a variety of IgV interfaces, from a canonical parallel domain interface [37,38] to an inverted (antiparallel/flipped) interface [39,40], partially matching the one we observe in the present structure. A list of PDB files containing a variety of VL-VL interfaces observed in crystal structures is available in the Supplementary Material (File S1).

Alphafold2 models of scFv dimers of current CARs, especially with the 47G4 sequence, show the formation of swapped VH and VL domains in dimeric forms with long linkers and short linkers, yet long linkers allow for a variety of domain arrangements. In the case of the 47G4 scFv, the VL-to-VH sequence (Figure 2E) with a long linker (Figure 2E) leads to swapped domain models (see Supplement Files S2). This is consistent with observations that scFvs in VL-to-VH arrangements form diabodies with linkers longer than 5 residues while exhibiting higher flexibility [41] than in the VH-to-VL order. For the 47G4 sequence, however, four out of five Alphafold2 predictions exhibit a G-strand swapping of VH domains (https://www.ncbi.nlm.nih.gov/Structure/icn3d/full.html?type=icn3dpng&url=https:// zenodo.org/api/files/4c8a1713-cd80-4ee5-9e3c-93782c4b3193/5models\_Hu19\_scFv\_AF2\_icn3d\_ loadable.png, accessed on 19 April 2023). The highest-ranking model presents an antiparallel VL-VL arrangement, albeit not with the interface observed in the X-ray structure observed in this study (Figure 3). This strand swapping is sequence-dependent and does not happen in Alphafold2 models with all VH | VL sequence combinations. G-strand swapping may represent an aggregation mechanism and may point to the importance of pairing VL and VH domains to maximize their interface stability.

#### 3.4. The Observed VL-VL Homodimer Interface

VL-VL homodimers have been observed originally in Bence-Jones proteins [42]. Unlike the Bence-Jones structure (PDBid:1REI), the 47G4-VLVL structure in this study exhibits an inverted, antiparallel VL-VL interface (see Figure 2). In fact, it has been shown that a single residue mutation in a VL domain can induce a rotation [43] or inversion [44,45] of the VL-VL domain interfaces (3D link showing an example of inversion due to a single mutation: 1LVE vs. 5LVE), and a previous survey in the literature found 61 VL-VL dimers in the PDB database [40] among which only nine formed a noncanonical quaternary arrangement. All these dimer complexes possess exact C2 symmetry (except for one with partial dissymmetry) but differ in the rotation angle and translational displacement between monomers. The 47G4-VL-VL dimer exhibits C2 symmetry. (A survey of the current PDB database for structures exhibiting significant VL-VL contacts in crystal structures can be found in Supplement Figure S1).

## 3.5. The Modeled vs. the Observed VL-VL Homodimer Interface: Binding Affinities

We hypothesized that preference for different VL-VL dimer configurations is due to more favorable binding affinity. To test this hypothesis, we conducted free energy calculations of VL-VL dimer formation for the solved inverted crystal structure, as well as the canonical interface of the same VL domain as predicted by AlphaFold2 [22,23]. As shown in Figure 4A, AlphaFold2 predicted the canonical interface for all 25 structural models but one model of low quality. It does not capture the inverted interface. This is surprising, as in many cases AlphaFold2 predicts oligomeric complexes with high accuracy [22,46].



**Figure 4.** Modeling of VL-VL binding interface. (**A**) Structural overlay of 25 VL-VL binding complex structures predicted by AlphaFold2 in the canonical configuration. Blue and pink coloring identifies different VL Ig domains. (**B**) RMSD profile of VL-VL complex showing stability across the Molecular Dynamics (MD) simulation. (**C**) Pulling work required to pull VL-VL complexes apart using Steered Molecular Dynamics (SMD). Shown are boxplots and points of all 20 replicas in different colors. (**D**) VL-VL binding free energies of SMD simulations from (**C**) computed using Jarzynski's inequality. Coloring denotes Canonical (blue) and Inverted (orange) conformation binding free energies. Error bars represent 95% CI.

However, here, we are asking AlphaFold2 to predict VL-VL homodimerization along two competitive interfaces, with one interface (canonical) being highly populated by sequence and structure libraries, that AlphaFold2 heavily relies upon for prediction. Reports in the literature have emerged that give a varying view of the ability of Alphafold2 to predict conformational diversity and/or quaternary configurations. Some studies show that the program can predict different monomeric vs. dimeric conformational states [47]. Other studies show that it does not accurately reproduce competing structures of fold-switching proteins [48] arguing that the lack of conformational diversity is due to its deep learning algorithm depending on pattern recognition rather than pure protein biophysics. Yet, the very success of AlphaFold2 lies in balancing deep learning and intrinsic knowledge of protein tertiary structure [23], but to address the complexity of quaternary interactions more biophysical knowledge is required.

We resorted to Molecular Dynamics (MD) methods to further study the observed dimeric interface. From 100 ns simulations (see Figure 4B), both the observed inverted crystal structure interface as well as the predicted canonical dimer interface remain bound, arguing that there is stability in both the predicted and the observed VL-VL interfaces. However, steered MD simulations (SMD) (see Methods) measuring the work required to pull the VL domains apart revealed that the inverted VL-VL dimer interface is more stable for the 47G4 VL domain (see Figure 4C). Likewise, when we use Jarzynski's equality to convert the pulling work to free energy values [49], we find that the inverted VL-VL dimer interface has a more favorable binding affinity than the canonical VL-VL interface:  $-66.8 \pm 1.1 \text{ kcal/mol vs.} -58.9 \pm 1.0 \text{ kcal/mol } (\pm95\% \text{ CI})$ ; although, from the SMD method we stress the relative qualitative trend rather than exact quantitative values. Ultimately, these modeling findings support our hypothesis that different VL-VL dimer configurations occur due to more favorable binding affinity.

## 3.6. Hypothetical Models of Canonical and Inverted VL-VL Dimers of scFvs

While the inverted VL-VL structure obtained during crystallization can be considered an anecdote, we interrogated 47G4 scFv dimerization through an Alphafold-multimer (v2.2) [22]. The predicted models are essentially G-strand-swapped dimeric structures (see above), while the original version (Alphafold2.0) produced top-ranking models without dimerization, except for an unusual diabody pairing VL-VL and VH-VH domains; both domains have a canonical interface (Figure 5A and Supplement Files S4). Surprisingly, no classical VLVH diabody was predicted. In contrast, building a full-length dimerized scFv model, linking VL and VH domains based on the inverted VL-VL structure, would produce quasi free VH domains (Figure 5B). Machine Learning (ML) methods are still evolving, and oligomeric models can only be taken as suggestions at this stage. In this case, they only suggest possible mechanisms of oligomerization or aggregation. New functional chain architectures could be imagined using VL domains for dimerization, as long as VH domains would not aggregate and could bind antigen targets as single Ig domains, similar to nanobody VHH domains [50].



**Figure 5.** Comparison of VL-VL based scFv dimer. (**A**) VL-VL diabody: a canonical VL-VL dimer interface leads to a pairing of VH domains. Top: Cartoon, Bottom: AlphaFold2-predicted structure. (**B**) Inverted VL-VL diabody model leading to free VH domains. Top: Cartoon, Bottom: structural model built on the basis of the observed inverted VL-VL structure. Blue and pink coloring identifies different scFv domains. Models are available as supplemental data (Supplement Files S4). 3D models comparison link: https://www.ncbi.nlm.nih.gov/Structure/icn3d/full.html?type=icn3dpng&url=https: //zenodo.org/api/files/4c8a1713-cd80-4ee5-9e3c-93782c4b3193/VLVLDIAB\_VLINVDIAB\_icn3d\_ loadable.png, accessed on 19 April 2023).

## 4. Discussion

Our structural knowledge of single-pass transmembrane receptors is limited [13]. With the development of single-pass transmembrane chimeric antigen receptors for CAR-T cell immunotherapies, it is imperative that we gain knowledge of their structure for accurate rational design. Recent studies and clinical trials show the need to understand CAR oligomerization and aggregation on the T-cell surface to optimize T-cell stimulation and limit toxicities. These works point to CAR oligomerization and functional dependence on scFv constructs, especially the VH-VL linker and VH-VL stability, yet the determinants of scFv-driven CAR oligomerization have not been identified [33]. This study sought to determine the extracellular structure of the anti-CD19 CAR 47G4-CD828Z. We unexpectedly captured a homodimer of a single IgVL domain with an inverted VL-VL interface. Further investigation revealed that the VL-VH linker was cleaved in the crystal, despite its designed cleavage resistance, with the VH and CAR Hinge domains absent from the crystal structure lattice. Molecular Dynamics simulations support the finding that the inverted VL-VL interface is more stable (enhanced binding affinity) than the canonical IgV interface predicted by Alphafold, as in the Bence-Jones protein. We find that IgV domain pairing and interface stability is an important and nontrivial parameter in scFv structure and, consequently, CAR oligomerization and CAR-T cell function. We conclude that it is important to interrogate the pairing and stability of VH-VL interfaces in single-chain antibody fragments when optimizing scFvs for use as binding moieties of CARs.

While we are still left with the question surrounding the extracellular structure of the intact CAR including the VL-VH and hinge domains of CAR-T-cells in biological conditions, this work substantiates the need to interrogate the scFv structure and CAR oligomerization state for optimal CAR-T design and calls for further structural studies to reveal the full picture of CARs expressed at the surface of T-cells.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/cryst13040710/s1. Supplemental files can be found in the Zenodo repository: https://doi.org/10.5281/zenodo.7809232 (accessed on 19 April 2023) [51]. Any file in repository can be accessed as an iCn3D [52] from the Zenodo API record record: https://zenodo.org/api/records/7809232 (accessed on 19 April 2023) [53].

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