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NMR Structure of the Natural Killer Cell Receptor 2B4 (CD244):

Implications for Ligand Recognition†,,‡

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Abstract

2B4, a transmembrane receptor expressed primarily on natural killer (NK) cells and on a subset of CD8+ T cells, plays an important role in activating NK-mediated cytotoxicity through its interaction with CD48 on target cells. We report here the atomic-resolution structure of the ligand-binding (D1) domain of 2B4 in solution determined by nuclear magnetic resonance (NMR) spectroscopy. The overall main chain structure resembles an immunoglobulin variable (V) domain fold, very similar to that seen previously for domain 1 of CD2 and CD4. The structure contains nine *β*-strands assembled into two *β*-sheets conventionally labeled DEB and AGFCC'C". The six-stranded sheet (AGFCC'C") contains structural features that may have implications for ligand recognition and receptor function. A noncanonical disulfide bridge between Cys2 and Cys99 stabilizes a long and parallel *β*-structure between strand A (residues 3-12) and strand G (residues 100-108). A *β*-bulge at residues Glu45 and Ile46 places a bend in the middle of strand C' that orients two conserved and adjacent hydrophobic residues (Ile46 and Leu47) inside the *β*-sandwich as seen in other V domains. Finally, the FG-loop (implicated in ligand recognition in the CD2-CD58 complex) is dynamically disordered in 2B4 in the absence of a ligand. We propose that ligand binding to 2B4 might stabilize the structure of the FG-loop in the ligand complex.

> Natural killer $(NK)^1$ cells are a fundamental component of the innate immune system that play a vital role in the detection and destruction of virally infected and tumor cells (1,2). The cytolytic activity of NK cells is regulated by a dynamic interplay between activating and inhibitory signals transmitted by distinct classes of receptors found on their surface. The dominant signal received by an NK cell through its interaction with normal levels of major histocompatibility complex (MHC) class I on target cells is inhibitory. When expression of MHC class I is reduced through infectious or tumorigenic processes, this inhibitory signal is attenuated, and the NK cell is activated. In this way, cells with abnormal MHC class I expression become targets of NK lytic activity that results from loss of inhibition of NK cell activation. Inhibitory receptors specific for MHC class I include the killer immunoglobulinlike receptors (KIRs), members of the Ly49 family, and the CD94/NKG2D family (3). Activating receptors include 2B4, CD16, CD44, CD69, NKR-P1, NKp30, NKp44, and NKp46 (1,2). However, only for some of these stimulatory receptors are the physiological ligands known: CD48 for 2B4 (4), IgG Fc for CD16 (5), and Clr for NKR-P1 (6). The balance between positive signaling receptors (resulting in target cell lysis) and negative signaling receptors (preventing lysis) ultimately determines the outcome of NK cell-target cell encounters (1,2).

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The activating NK receptor 2B4 is a transmembrane glycoprotein expressed primarily on NK cells, and on a subset of CD8+ T cells, that is involved in activating NK-mediated cytotoxicity $(7-10)$. 2B4 comprises two Ig-like extracellular domains (D1 and D2) and a cytoplasmic portion containing a CxC/+ motif that may bind a complementary CxxC/-motif in the linker for activation of T cells (LAT) (11). By sequence homology, 2B4 is most closely related to the CD2 subgroup of the immununoglobulin (Ig) superfamily (Figure 1). The natural ligand for 2B4 is CD48 (4). As measured by surface plasmon resonance (SPR), soluble mouse CD48 binds to immobilized recombinant mouse 2B4 with a K_D of 16 μ M. The human counterpart of 2B4 was found to bind human CD48 with similar affinity $(K_D = 8 \mu M)$ (4). These affinities are approximately 5-fold higher than that of CD48 for CD2 (12,13).

In addition to LAT, the cytoplasmic tail of 2B4 binds the SH2 domain-containing protein SAP (14). This association is crucial for 2B4 to deliver activating signals to NK cells. Indeed, in the absence of SAP, 2B4 tranduces inhibitory signals that result in NK cell inactivation (7). This inhibition is mediated by the SHP-1 phosphatase, which competes with SAP for binding to 2B4. The 2B4 NK receptor has recently been associated with a severe, genetically inherited immune deficiency, termed X-linked lymphoproliferative disease (XLP), that is often characterized by Epstein-Barr virus (EBV)-induced lymphomas (7). In these patients, 2B4, upon binding CD48 on target cells, fails to activate, but instead inhibits, NK-mediated lysis of EBV-infected cells. The mutated gene responsible for XLP codes for SAP (15-17). In XLP NK cells, SAP is either truncated or missing and cannot associate with 2B4 (7). As a consequence, SHP-1 stably binds 2B4, contributing to the generation of a negative signal.

The finding that 2B4 can both activate and inhibit NK cytolysis, combined with the identification of CD48 as its biological ligand, makes 2B4 a compelling target for structural analysis. We present here the three-dimensional structure of the ligand-binding D1 domain of 2B4 by NMR methods. 2B4 adopts an immunoglobulin variable (V) domain fold, like that of domain 1 of CD2 and CD4 (Figure 1), but with several unique structural and dynamical features that could be related to ligand recognition and receptor function.

EXPERIMENTAL PROCEDURES

Protein Expression and Purification

A DNA fragment encoding the ligand-binding domain of rat 2B4 (residues 1-112, referred to throughout the text as simply 2B4) was generated by PCR from the full-length rat gene and cloned into the bacterial expression vector pT7-7 (Novagen). The protein was expressed as inclusion bodies in *Escherichia coli* BL21(DE3) cells (Stratagene). Isotopically enriched protein was produced by growing cells in M9 minimal medium containing 15N-labeled ammonium chloride and $13C$ -labeled glucose as sole nitrogen and carbon sources, respectively (18,19). The cells were cultured at 37 °C to an absorbance of 0.6-0.8 at 600 nm, and isopropyl- β -_D-thiogalactoside was added to a concentration of 1 mM. After incubation for 4 h, the cells were resuspended in 100 mM Tris-HCl (pH 8.0) containing 2 mM ETDA and 10 mM DTT and lysed by two passes in a French Press at 1000 psi. Inclusion bodies were washed several times with 50 mM Tris-HCl (pH 8.0), 0.5% (v/v) Triton-X 100, 100 mM NaCl, and 1 mM ETDA, then solubilized in 100 mM Tris-HCl (pH 8.0), 8 M urea, 10 mM EDTA, and 1 mM DTT.

For in vitro folding, solubilized 2B4 D1 was diluted to a final concentration of 60-100 *μ*g/mL into 0.8 M arginine, 100 mM Tris-HCl (pH 8.0), 0.3 mM reduced glutathione, 3 mM oxidized glutathione, and 2 mM EDTA. After 3-4 days at 4 °C, the folding mixture was concentrated, dialyzed against 25 mM Tris-HCl (pH 8.0) and 100 mM NaCl, and applied to a Superdex 75 HR size exclusion column (Amersham Biosciences). Further purification was carried out by anion exchange chromatography using a Mono Q column; the protein was eluted with a linear NaCl gradient. The buffer was exchanged to 100 mM sodium phosphate (pH 7.2) prior to NMR analysis.

NMR Spectroscopy

Samples for NMR analysis consisted of ¹⁵N-labeled or ¹³C/¹⁵N-labeled 2B4 (0.4 mM) in 0.3 mL of a 95% H₂O/5% $[^2H]$ H₂O solution containing 100 mM sodium phosphate (pH 7.2). All NMR experiments were performed at 25 °C on Bruker Avance 500 or 600 MHz spectrometers equipped with a four-channel interface and triple-resonance probe with triple-axis pulsed field gradients. The $15N-1H$ HSQC spectra (see Figure 2) were recorded on a sample of $15N$ -labeled 2B4 (in 95% H O, 5% $[^2H]H_2O$). The number of complex points and acquisition times were 256 and 180 ms (${}^{15}N$ (F₁)) and 512 and 64 ms (${}^{1}H$ (F₂)). All triple-resonance experiments were performed, processed, and analyzed as described (19,20) on a sample of ${}^{13}C/{}^{15}N$ -labeled 2B4 (in 95% H O, 5% $[{}^{2}H]H_{2}O$) with the following number of complex points and acquisition times: HNCO {¹⁵N (F₁) 32 and 23.7 ms, ¹³CO (F₂) 64 and 42.7 ms, ¹H (F₃) 512 and 64 ms}; HNCACB $\{^{15}N(F_1)$ 32 and 23.7 ms, $^{13}C(F_2)$ 48 and 6.3 ms, $^{1}H(F_3)$ 512 and 64 ms}; CBCACONNH {¹⁵N (F₁) 32 and 23.7 ms, ¹³C (F₂) 48 and 6.3 ms, ¹H (F₃) 512 and 64 ms}; HBHACONNH {¹⁵N (F₁) 32 and 23.7 ms, ¹H_{ab} (F₂) 64 and 21 ms, ¹H (F₃) 512 and 64 ms}. ¹⁵N-edited and ¹³C-edited NOESY-HSQC and TOCSY-HSQC experiments were performed as described previously (21,22).

Triple resonance and NOESY spectra measured above were analyzed to determine resonance assignments and secondary structure of 2B4. The chemical shift index (see ref 23 for detailed description of the chemical shift index), $\frac{3}{{\cal J}_{\rm NH}}$ coupling constants, and Nuclear Overhauser Effect (NOE) connectivity patterns for each residue were analyzed and provided a measure of the overall secondary structure. Small ${}^{3}J_{\text{NH}\alpha}$ coupling constants (<5 Hz), strong NOE connectivities (NN $(i, i + 1)$ and α N $(i, i + 3)$), and positive chemical shift index are characteristic of residues in an α-helix. Conversely, large ${}^{3}J_{\text{NH}\alpha}$ coupling constants (>8 Hz), strong αN(*i*, *i* + 1) and weak NN(*i, i* + 1) NOE connectivities, and negative chemical shift index are

characteristic of residues in a *β*-strand. The results of the secondary structure analysis and topology of 2B4 are summarized schematically in Figure 1.

Structure Calculation

Backbone and side chain NMR resonances of 2B4 were assigned as described previously (20). Analysis of NOESY data determined nearly 1900 interproton distance relationships throughout the protein (19). The NMR-derived distances and dihedral angles then served as constraints (see Table 1) for calculating the three-dimensional structure of the protein using distance geometry and restrained molecular dynamics. Structure calculations were performed using the YASAP protocol within X-PLOR (24,25), as described previously (26). A total of 1891 interproton distance constraints were obtained as described (20) by analysis of 13 C-edited and 15N-edited NOESY-HSQC spectra (100 ms mixing time) of 13C/15N-labeled 2B4. In addition to the NOE-derived distance constraints, the following additional constraints were included in the structure calculation: 128 dihedral angle constraints (ϕ and ψ) and 170 distance constraints for 85 hydrogen bonds verified by identifying slowly exchanging amide protons in hydrogen-deuterium exchange experiments (18). Fifty independent structures were calculated, and the 15 structures of lowest energy were selected. The average total and experimental distance energies are 2969 and 249 kcal mol⁻¹. The average root-mean-square (rms) deviations from an idealized geometry for bonds and angles are 0.0074 Å and 2.25°. None of the distance and angle constraints were violated by more than 0.40 Å or 4° , respectively.

RESULTS AND DISCUSSION

The ¹H-¹⁵N HSQC NMR spectrum of ¹⁵N-labeled 2B4 exhibited close to the expected number of backbone amide resonances (101 out of 105), suggesting that the protein is stably folded (Figure 2). Pulsed field gradient NMR diffusion studies (27) confirmed a protein monomer in solution under NMR conditions. More than 90% of the NMR resonances had very sharp and uniform peak intensities and were assigned as indicated in Figure 2. The assigned peaks in the spectrum represent main-chain and side-chain amide groups that serve as fingerprints of overall conformation. Three-dimensional structures of 2B4 were calculated on the basis of NOE data, slowly exchanging amide NH groups, chemical shift analysis, and $3J_{\text{NH}\alpha}$ spin-spin coupling constants (see Experimental Procedures). The analysis of chemical shift index (CSI) (28), $3J_{\text{NH}\alpha}$ (29), and short-range NOEs (30) is illustrated in Figure 1A and revealed at least nine *β*-strands and one short α-helix. The final three-dimensional structures of 2B4 derived from the NMR data are superimposed and illustrated in Figure 3A (atomic coordinates have been deposited in the RSCB Protein Databank). Table 1 summarizes the structural statistics calculated for the15 lowest energy conformers.

The NMR-derived 2B4 structures (Figure 3B,C) clearly showed a fold like that of an immunoglobulin V domain seen previously in the structures of domain 1 of CD2 (31, 32) and CD4 (33, 34). The 2B4 structure contains nine *β*-strands (see Figure 1A for residue definition of individual strands). The overall topology consists of two *β*-sheets of strands labeled DEB and AGFCC'C" (Figure 1B). The two *β*-sheets in 2B4 are similar to those of CD2 and CD4 and differ somewhat from the more conventional four-stranded (ABED) and six-stranded (A'GFCC'C") *β*-sheets observed in the majority of other V domains (35).

The three-stranded *β*-sheet (DEB) is somewhat shorter in length than that of CD2 and CD4. Strand B in 2B4 is flanked on both ends by proline (Pro16 and Pro21), which lacks a mainchain amide proton and destabilizes the hydrogen-bonding interaction with strand E. The length of the three-stranded sheet is also shortened by the disordered connecting loop between strands D and E (residues 68-72).

The six-stranded *β*-sheet (AGFCC'C") of 2B4 forms a curved (saddle-like) outer surface implicated in ligand binding in other V domains (32,36). In 2B4, strands A and G are connected in a parallel fashion by main-chain hydrogen bonds and a disulfide bridge between Cys2 and Cys99. Strands F and G form a hairpin *β*-structure with a connecting FG-loop (residues 92-97) that appears dynamically disordered (very broad NMR peaks) and whose structure is not accurately defined by the current NMR data. Preliminary NMR relaxation analysis of the FGloop suggested significant internal dynamics on the subnanosecond time scale (data not shown), consistent with a highly disordered main chain. Interestingly, this long and unstructured FG-loop contains residues that are believed to play a role in ligand binding based on the closely related crystal structure of the CD2-CD58 complex (36). Strands C and C' form a hairpin *β*-structure connected by a well-defined CC'-loop (residues 37-42) that might also play a role in ligand binding (32,36). Strand C' is interrupted by a *β*-bulge at Glu45 and Ile46, forming two shorter strands, C_1 ' and C_2 ' (Figure 1). The *β*-bulge disrupts the register of mainchain hydrogen bonds between strands C and C' (Figure 4). Strand C_1 ' connects to strand C by main-chain hydrogen bonds between Lys35 and Ile44. The *β*-bulge prevents hydrogen bonding between Trp33 and Ile46 and shifts the register of the two strands by one residue. As a consequence, strand C_2 ' connects to strand C by main-chain hydrogen bonds between Trp33 and Leu47 and between Val31 and Trp49. The *β*-bulge allows two adjacent hydrophobic side chains (Ile46 and Leu47) to point in the same direction, inside the *β*-sandwich (Figures 3C and 4) that otherwise would point in opposite directions in an extended main-chain conformation. A similar *β*-bulge in strand C' has been described for CD2 and CD4 and is believed to make the GFCC' sheet curved (35). Strands C_2 ' and C'' form a hairpin β -structure connected by a sharp, two-residue *β*-turn that is somewhat different from the longer C'C"-loop seen in some V domains. Finally, Asp1 at the N-terminus and the last three residues at the C-terminus (residues 110-112) are unstructured and not defined by the NMR data.

The two β-sheets (DEB and AGFCC'C") of 2B4 are connected by a short helix (residues 81-84) and three connecting loops (AB-, BC-, and C"D-loops). The structure of these intersheet regions is poorly defined by the NMR data, perhaps due to dynamics. Indeed, the NMR line widths and relaxation properties for residues in the connecting loops (AB-, BC-, and C"Dloops) suggest conformational fluctuations on the chemical shift time scale. The dynamical properties of the intersheet loops indicate conformational elasticity between the two *β*-sheets that might play a role in ligand recognition.

The overall main-chain structure of the *β*-sheet regions of 2B4 (DEB and AGFCC'C") is similar to that in domain 1 of CD2 and CD4. The rmsd alignment of main-chain atoms in the *β*-sheet regions is 2.4 Å, comparing 2B4 and CD2, and 2.7 Å, comparing 2B4 and CD4. The two *β*sheets of 2B4, CD2, and CD4 are somewhat different from the canonical four-stranded (ABED) and six-stranded (A'GFCC'C") *β*-sheets found in conventional V domains (35). Normally, the N-terminal region of strand A forms antiparallel *β*-structure with strand B in the four-stranded sheet, whereas the C-terminal end of strand A (A') forms a parallel interaction with strand G in the six-stranded sheet. The structures of 2B4, CD2, and CD4 completely lack the antiparallel interaction between strands A and B. Instead, strand A in 2B4 is quite long (nine residues in 2B4 vs five residues in CD4) and is held in an unusually long parallel *β*-structure with strand G due to a noncanonical disulfide bridge between Cys2 and Cys99. The long parallel *β*-structure between strands A and G may help stabilize the convex outer curvature of the six-stranded *β*sheet (AGFCC'C").

2B4 has unique structural features that may be important for ligand recognition. The structure of the CC' and FG loop regions differ substantially in 2B4 and domain 1 of CD2 (Figure 5A). These loops are not conserved in sequence or length. The CC' and FG-loops have been implicated previously in ligand binding (32, 36). The marked structural differences between 2B4 and CD2 in Figure 5A suggests that the CC' - and FG-loops might, in part, explain ligand

specificity. The FG-loop in 2B4 is dynamically disordered in our structure, perhaps because ligand binding is needed to stabilize its structure. The dynamics of the FG-loop might suggest an induced-fit mechanism for ligand recognition by 2B4. In addition, 2B4 has a noncanonical disulfide bridge at Cys2 and Cys99 that stabilizes a long and parallel *β*-sheet between strands A and G, causing a convex or saddle-like curvature to the outer surface of the six-stranded *β*sheet (AGFCC'C"). Finally, the C'-strand in 2B4 is interrupted by a *β*-bulge that causes two adjacent hydrophobic side chains (Ile46 and Leu47) to point in the same direction, inside the *β*-sandwich (Figure 3C,D).

Implications for Ligand Recognition

The structural interaction of 2B4 with its biological ligand, CD48 (D1-domain), was modeled on the basis of our current structure of 2B4 and the crystal structure of the CD2-CD58 complex (Figure 5B). The D1 domain of CD48 is ∼28% identical (45% homologous) in primary sequence to that of CD58 (Figure 1A) whose three-dimensional structure is known (36, 37). An approximate three-dimensional structure of the D1 domain of CD48 was calculated by homology modeling based on the known structure of CD58 using the InsightII/Discover/ Homology program (BIOSYM Technology, San Diego, CA). The modeled structure of CD48 was then docked onto our current structure of 2B4 by first overlaying (in a least squares fashion) the CD48 structure on top of CD58 in the CD2-CD58 complex (36), followed by a similar alignment of the 2B4 structure with respect to CD2 in the complex. This alignment procedure places 2B4 in close proximity to CD48 with the same relative orientation as defined in the CD2-CD58 complex. The six-stranded *β*-sheet (AGFCC'C") of 2B4 interacts face-to-face with the corresponding *β*-sheet of CD48. This interface lacks many of the intermolecular salt-bridge interactions between charged side chains that are evident in the CD2-CD58 complex. The charged, interfacial residues of CD2 (Asp31, Asp32, Lys34, Glu36, Lys43, Arg48, Lys51, and Lys82) and CD58 (Glu25, Lys29, Lys32, Asp33, Lys34, Glu37, Glu39, Glu42, Arg44, Glu76, Glu78, and Asp84) are not conserved in 2B4 and CD48, respectively. Therefore, the 2B4-CD48 interface does not possess many intermolecular electrostatic contacts, but instead is stabilized more by intermolecular hydrophobic and/or hydrogen-bonding interactions.

The modeled structure of the 2B4-CD48 complex suggests that the FG-loop of 2B4 is located very close to the CC'-loop of CD48 (Figure 5). The FG-loop of 2B4 (in the absence of ligand) is dynamically disordered in our NMR analysis, so the FG loop structure of 2B4 shown in Figure 5 represents the energy-minimized average structure in solution. We propose that the dynamical properties of the FG-loop would allow it to sample a wide range of conformations that rapidly select an optimum conformation for recognizing the CC'-loop of CD48. There is growing evidence that regions of structural flexibility may represent likely sites for receptor binding as demonstrated by the interaction of the NK receptor, NKG2D, with the MHC-like molecule, MICA. The receptor-binding site on MICA is disordered in the crystal structure of the free ligand but adopts an α -helical conformation upon binding to NKG2D (38).

Another striking feature of the 2B4-CD48 complex (Figure 5) was the finding that the CC' loop of 2B4 is not located close to the FG-loop of CD48, unlike what is found in the CD2- CD58 complex. The *β*-bulge in 2B4 (Figure 4) places a bend in the middle of the C'-strand that displaces the CC'-loop of 2B4 downward with respect to the CC' loop of CD2 (Figure 5A). This downward disposition places the CC'-loop of 2B4 quite far from the FG-loop of CD48 in the complex. One interpretation is that ligand binding might reorient 2B4 as a whole in the complex that would place its CC'-loop in closer contact with the FG-loop of CD48. Alternatively, ligand binding might induce a local conformational change in 2B4 near the *β*bulge that would move the CC'-loop upward to adopt a structure more akin to that of CD2 in the complex.

In summary, we determined the three-dimensional structure of the ligand-binding D1 domain of 2B4 in solution by NMR (Figure 3) and proposed a modeled structure of the 2B4-CD48 complex (Figure 5). Our analysis revealed unique structural and dynamical properties of 2B4 (in the CC'- and FG-loops), perhaps implying ligand-induced conformational changes in 2B4 that might promote ligandspecific contacts. The implications of these results for ligand recognition will be tested more rigorously in the future by site-directed mutagenesis analysis (of the CC'- and FG-loops), ligand-induced chemical shift changes, and by ultimately determining the atomic-resolution structure of the 2B4-CD48 complex.

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Figure 1.

(A) Amino acid sequence alignment of immunoglobulin V domains for rat 2B4 (accession no. Q9JLM2), human CD2, human CD4, human CD48, and human CD58. Residue definition of *β*-strand regions are indicated by arrows. Conserved residues are highlighted in bold. The chemical shift index (CSI) of each residue $(-1, 0, +1)$ is represented by -, 0, or +, respectively. The $3J_{\text{NH}\alpha}$ coupling constants are denoted as "large" (l, >8 Hz), "medium" (m, 5-8 Hz), and "small" (s, <5 Hz). Sequential NOE connectivities $(d\alpha N(i, i + 1))$ are represented as strong, weak, or zero intensity in bar graphs. (B) Schematic representation of *β*-sheet topology.

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Figure 2.

 ${}^{1}H-{}^{15}N$ HSQC NMR spectrum of 2B4. Peaks corresponding to the NH₂-groups of the sidechain amides of Gln and Asn residues are connected by dotted lines. Sequence-specific assignments are indicated and deposited to the BioMagResBank Repository (BMRB accession no. 6538).

Figure 3.

Main-chain structure of 2B4 in solution determined by NMR². (A) Superposition of mainchain atoms of the 15 lowest energy calculated structures. (B) Ribbon representation of the energy-minimized average main-chain structure and side-chain atoms of cysteine residues. (C) A 180° rotation of B. The FG-loop (residues 92-97) is dynamically unstructured and not shown in A. *β*-strand regions are highlighted in yellow.

Figure 4.

Hydrogen bonds in the *β*-bulge region. The main-chain atoms of strands C and C' are depicted by a ball-and-stick model. Carbon, hydrogen, nitrogen, and oxygen atoms are colored white, yellow, blue, and red, respectively. Main-chain hydrogen bonds between strands C and C' are indicated by dotted lines. The *β*-bulge, formed by Glu 45 and Ile 46, disrupts the hydrogen bonding pattern between strands C and C'.

Figure 5.

(A) Overlay of main-chain structures of 2B4 (red) and CD2-CD58 complex (1qa9.pdb in blue). (B) Modeled three-dimensional structure of the 2B4-CD48 complex. The structure of CD48 was calculated on the basis of the known structure of CD58 (1qa9.pdb) using homology modeling. The modeled structure of CD48 was docked onto 2B4 by overlaying the CD48 and 2B4 structures on top of CD58 and CD2, respectively,in the complex.

Table 1 Structural Statistics for the Ensemble of 15 Calculated Structures of 2B4

NOE restraints (total)	1891
intra (i - $ i - j = 0$)	722
medium $(1 \leq i - j \leq 4)$	506
long $(i - j > 4)$	663
dihedral angle restraints (ϕ and ψ)	128
hydrogen bond restraints in β -sheet regions	170
rmsd from ideal geometry	
bond length (\AA)	$0.0074 + 0.0004$
bond angles $(°)$	$2.25 + 0.02$
Ramachandran plot	
most favored region (%)	70
allowed regions (%)	27
disallowed regions $(\%)$	3
rmsd of atom position from average structure	
β -sheet regions (main chain atoms) (\AA)	$0.4 + 0.05$
β -sheet regions (non-hydrogen atoms) (\AA)	1.1 ± 0.08