

De novo design of an IL-4 antagonist and its structure at 1.9 Å

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An IL-4 antagonist was designed based on structural and biochemical analysis of unbound IL-4 and IL-4 in complex with its high-affinity receptor (IL-4R α). Our design strategy sought to capture a protein–protein interaction targeting the high affinity that IL-4 has for IL-4R α . This strategy has impact due to the potential relevance of IL-4R α as a drug target in the treatment of asthma. To mimic the IL-4 binding surface, critical side chains for receptor binding were identified, and these side chains were transplanted onto a previously characterized, *de novo*-designed four-helix protein called designed helical protein 1 (DHP-1). This first-generation design resolved the ambiguity previously described for the connectivity between helices in DHP-1 and resulted in a protein capable of binding to IL-4R α . The second-generation antagonist was based upon further molecular modeling, and it succeeded in binding IL-4R α better than the first-generation. This protein, termed DHP-14-AB, yielded a protein with a cooperative unfolding transition ($\Delta G^0_u = 8.1$ kcal/mol) and an IC_{50} of 27 μ M when in competition with IL-4 whereas DHP-1 had no affinity for IL-4R α . The crystal structure of DHP-14-AB was determined to 1.9-Å resolution and was compared with IL-4. This comparison revealed how design strategies targeting protein–protein interactions require high-resolution 3D data and the incorporation of orientation-specific information at the level of side-chains and secondary structure element interactions.

de novo protein design | helical bundle | IL-4 receptor | structure-based design

We sought to design an antagonist to compete with the high-affinity interaction between IL-4 and its receptor, IL-4R α . This protein–protein interaction is an important drug development target in the treatment of acute allergic asthma and other atopic conditions such as seasonal allergies, urticaria, and eczema (1, 2). Our purpose was to create a method for antagonist generation that could be instructive for curing these diseases and other diseases resulting from rogue protein–protein interactions. IL-4·IL-4R α forms a heterotrimeric receptor complex along with another cytokine receptor of either IL-13 receptor α chain 1 (IL-13R α 1) or γ common chain (γ C) to activate signaling across cell membranes in the immune system (3). IL-4 is in the short-chain helical cytokine family; this family lacks any significant sequence homology between members (4, 5). The IL-4 four-helix bundle has an up–up–down–down order and orientation between major helices (6–8).

The IL-4·IL-4R α complex formation ($K_d \approx 160$ pM) is one of the highest affinity interactions of a cytokine for its receptor (9). IL-4R α is a member of the hematopoietic receptor superfamily (10). Its association rate constant, $k_{on} \approx 1.8 \times 10^7$ M $^{-1}$ ·s $^{-1}$, for IL-4 to IL-4R α is nearly 100-fold as fast as the presumably diffusion-limited association rate constant for cytokine interactions with their first contact single receptor chain, such as human growth hormone·hGHR ($k_{on} \approx 3 \times 10^5$ M $^{-1}$ ·s $^{-1}$) (9, 11). This fast and high-affinity interaction has been attributed to “electrostatic steering” and to the complementary interface between the positively charged IL-4 and negatively charged IL-4R α

surfaces; the opposing charges attract each other, resulting in associate rates faster than diffusion, and coordinate each other in the complex (9).

The platform selected for IL-4 antagonist design was our own *de novo*-designed, four-helix bundle protein, designed helical protein (DHP) 1; previously, we showed that DHP-1 has stability like that of a natural protein and a structure exactly as intended (12). That design was based upon conjugation of four 24-aa amphipathic helical peptides that were built from a reduced set of the natural amino acids. The DHP-1 crystal structure did not resolve the connectivity between the helices, which resulted in a structural ambiguity. Both bundle topology and superhelical twist are described as “right-handed” or “left-handed.” The helical bundle topology refers to the macroscopic handedness (chirality) and is based on the backbone connectivity and unit direction vector of the helices (13). In contrast, the superhelical twist of a bundle refers to the wrapping of helices around the protein core axis regardless of connectivity between helices, for which most frequently bundles are left-handed (14). The DHP-1 helices have a left-handed superhelical twist. The DHP-1 structure without the loops had an ambiguity in the bundle topology that was resolved here by creating IL-4R α binding function.

The DHP-1 structure showed that it had a pair of neighboring helices of approximately similar antiparallel orientation, size, and spacing as IL-4 (Fig. 1). This structural similarity was sufficient to initiate designing an antagonist to IL-4 onto the DHP-1 helices. The residues on IL-4, which interact with IL-4R α , were analyzed to select a subset for transfer. These residues on IL-4 were then modeled *in silico* onto DHP-1 to create the first-generation antagonists. Adding these residues was the first step toward resolving the topology question about DHP-1 by breaking its symmetry.

After the first generation of antagonist development, the so-called DHP-10 bundles, our proteins were assessed for stability in solution and functional binding to IL-4R α to ascertain which designed topology was represented. At the time when IL-4·IL-4R α coordinates became available, we were able to integrate the topology design results with analysis of steric clash-points revealed on IL-4R α that were suboptimal in the context of our first design. This observation led to the second-generation design to remove potential clashes from our IL-4 antagonist and to fit the target complex. This second-generation design, DHP-14-AB, was solved and compared with IL-4. The analysis of this structure along with IL-4 indicated that the crossing angle (Ω) between helices is likely to be as important as

Abbreviations: IL-4R α , IL-4 receptor α chain; IL-13R α 1, IL-13 receptor α chain 1; γ C, γ common chain; DHP, designed helical protein; MBP, maltose-binding protein; Rh, hydrodynamic radius; $\Delta\Delta G$, relative change in binding energy.

Data deposition: The atomic coordinates and structure factors for DHP-AB-14-MBP have been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 1Y4C).

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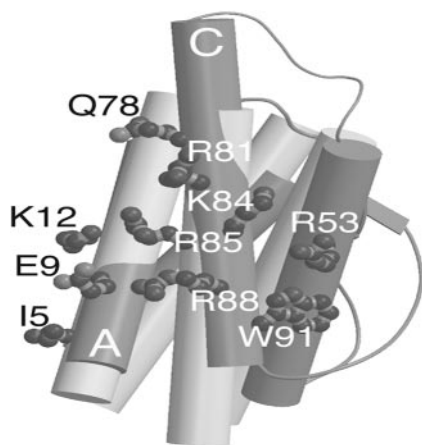


Fig. 1. DHP-1 (light gray) helices superimposed on unbound IL-4 (dark gray). IL-4 side chains are displayed in atom representation for mutations made onto DHP-1 in the first round design. Side chains fit onto the DHP-1 backbone structure except for R53, because the third helix of DHP-1 did not overlap well on the corresponding IL-4 B helix.

the overlap of the C α positions within the binding site and the energetic value of the side chain interactions with the receptor.

Materials and Methods

Molecular Modeling. The O suite software LSQMAN (15) was used to superimpose the molecules with 36 C α atoms included. Side chains were mutated by using the 3D modeling software MOLOC (16). Rotamers were selected from the library based on which side chain could fit on DHP-1 and be consistent with IL-4 unbound and later IL-4 bound to IL-4R α . First-generation models were subjected to energy minimization with CNS (17).

Gene Synthesis. Two genes were designed, synthesized (Integrated DNA Technologies, Coralville, IA), and subcloned into cloning and expression plasmids. The genes were based on the 108-aa sequence of DHP-1 (12) (Table 4, which is published as supporting information on the PNAS web site). Each gene was sequenced in the forward and reverse directions in both vectors.

Protein Expression and Purification. DHPs were expressed in DH-5 α *Escherichia coli* by using the pMal-c2x vector, and levels were similar to DHP-1, with a yield of >25 mg/liter purified protein (12). DHP concentration was determined by absorption of ultra-violet light at 280 nm due to one tryptophan and the molar extinction coefficient of 5,500 M⁻¹·cm⁻¹ (18).

Helical Structure Monitored by CD. CD measurements were made by using a J-715 spectropolarimeter (Jasco, Easton, MD) with a temperature controlled holder and a 2-mm path length cuvette.

Unfolding Free-Energy (ΔG_0^U) Calculation. The unfolding transition was monitored at the α -helical signal minimum 222 nm by CD as a function of the denaturant, guanidinium hydrochloride (Gdn·HCl), concentration (M). The buffer was 10 mM sodium phosphate (pH 7.0) at 25°C; each component of the solution was weighted out in order to eliminate protein concentration variation. Samples were equilibrated overnight in the dark at 25°C. Gdn·HCl concentration was determined by using the difference in refractive index (Δn) between the buffer and the solutions with Gdn·HCl added (19). The data were fit to a two-state transition by using a nonlinear free-energy curve-fitting method (20) and the software KALEIDAGRAPH (Synergy Software, Reading, PA).

Table 1. ΔG_u^0 and unfolding midpoint (Gdn-HCl) was calculated by nonlinear extrapolation (20)

Protein	ΔG_u^0 , kcal/mol	Unfolding midpoint, M	IC ₅₀ , μ M
DHP-1	13.1	4.8	N.B.
DHP-10-AB	13.5	4.6	495
DHP-10-AD	4.3	4.4	970
DHP-14-AB	8.1	3.2	27

IC₅₀ for each protein were determined by competition with IL-4-HRP for IL-4R α . DHP-1 did not bind (N.B.) IL-4R α under these conditions.

Dynamic Light Scattering (DLS). Monodispersity, hydrodynamic radius (Rh), and molecular weight of DHPs were determined by DLS at 22°C by using the DynaPro MS/X Instrument and DYNAMICS 5.25.44 software (Proterion, Piscataway, NJ). The REGULARIZATION algorithm was used for analyzing the experimental autocorrelation function (channels 1–80), yielding the translational diffusion coefficient (Dt), Rh, and intensity distributions of particle diameter. Solvent refractive index and solvent viscosity were 1.333 and 1.019, respectively. Approximately 70% of the measurements were averaged and used to estimate Rh and molecular weight based on an empirical curve of known proteins. Before DLS analysis, DHPs at 3–5 mg/ml were dialyzed overnight against buffer (PBS, 10 mM sodium phosphate, pH 7.0, or 10 mM Tris, pH 7.4) and filtered.

Competitive Inhibition Assay. Each protein was incubated for 1 h at room temperature on a 96-well plate fixed with biotinylated IL-4R α and horseradish peroxidase (HRP)-conjugated IL-4. The loss of HRP activity due to displacement of HRP-IL-4 was monitored as a function of competing protein concentration. The curve-fit data K_i s were converted to IC₅₀s: IL-4, 2.3 $\pm$ 0.16 nM; DHP-14-AB, 64 $\pm$ 2.3 μ M; DHP-AB, 1.1 $\pm$ 0.20 mM; and DHP-AD, 2.2 $\pm$ 0.29 mM. The IC₅₀ for IL-4 was 2.3 nM, although it was typically 1 nM; therefore, division by 2.3 normalized each IC₅₀ for assay condition variation (Table 1).

Crystallization and X-Ray Diffraction Data Collection. Crystals were grown of DHP-14-AB, as an maltose-binding protein (MBP) fusion, by the hanging drop vapor diffusion method, with 1:1 ratio drops of protein solution (10 mg/ml) in 20 mM Tris (pH 7.4) with a well solution of 2 M ammonium sulfate, 50 mM sodium acetate (pH 4.6) at room temperature. Crystalline needles were used for seeding growth of rod-like crystals, which diffracted to 1.9 Å at Advanced Light Source (ALS) beamline 8.3.1 using glycerol in artificial mother liquor for cryoprotection. Diffraction data were processed by using ELVES (21) with MOSFLM (22) and CCP4 (23).

Structure Determination and Refinement. A molecular replacement solution was found by using EPMR 2.5 (24) with MBP as the search model. Rigid body refinement and semiautomated building using ARP/WARP (25) produced interpretable maps with 77% of the asymmetric unit MBP and 23% DHP-14-AB (Table 2). Subsequent rounds of building were performed with MOLOC (16) and O (15), and refinement was done with CCP4 (23) and CNS suites (17).

Results

Transplantation of IL-4R α Binding Site onto DHP-1. Hage *et al.* (3) determined the structure of IL-4-IL-4R α and compared the atomic coordinates of IL-4 from the free IL-4 with those in the IL-4-IL-4R α complex. They found that the IL-4R α binding site is compact and composed exclusively of side-chains, most of which were derived from IL-4 helices A and C, and provided a predominantly positively charged interfacial surface of $\approx 800 \text{ \AA}^2$

Table 2. Summarized data collection and structure refinement statistics

Diffraction resolution, Å	1.9
Space group	$P2_12_12_1$
Unit cell, Å	$a = 69.56, b = 74.69, c = 103.69$
Final R -factor* (R_{free}), %	20.2 (22.4)
No. of residues†	
Protein	494 (14)
Waters	284
Maltose	1
rmsd angles, °	1.1
rmsd bonds, Å	0.0049

Complete data statistics are available in Table 5, which is published as supporting information on the PNAS web site. rmsd, rms deviation.

* R -factor = $\sum |F_{\text{obs}}| - |F_{\text{calc}}| / \sum |F_{\text{obs}}|$. R_{free} is computed in the same manner as the R -factor with the test set of reflections (5%).

†The values in parentheses are for number of missing residues.

of solvent accessible surface area on IL-4. Upon interaction with IL-4R α , IL-4 binding site helices shifted on average by <1 Å, and IL-4 accommodated by adjusting amino acid side-chains and tilting helices to form the interaction. For example, the distance between C α positions on IL-4 helices A and C within the binding interface decreased by 0.4–0.6 Å. More significant contractions in the range of 1–1.5 Å occurred between C α positions on IL-4 helices A and D. Helices A and D support the putative binding site for the second receptor (26). Our four-helix bundle design mimics only the high-affinity receptor binding site on IL-4 helices A and C, because it does not incorporate the putative second receptor binding site.

To design the interactions for binding to IL-4R α , the C α positions for two helices of DHP-1 were superimposed on IL-4 helices A and C. Upon superposition of these positions, the rms deviation between the structures is 0.96 Å using the binding site portion of the IL-4 structure and the analogous residues from two DHP-1 helices (Fig. 1). The crossing angle (Ω) between IL-4 A and C helices was broader by 2.6° than the DHP-1 crossing angle between helices A and B and broader by 5° than that for DHP-1 helices A and D. The inter-helix distances were within 1 Å between IL-4 and either unique pair of DHP-1 helices (27) (Table 3). Because the backbone helices superimposed within less than an angstrom and DHP-1 could place two helices facing

Table 3. Crossing angle (Ω) and closest approach distance between target helices for IL-4 antagonist template peptides and proteins and IL-4

Structure	Helix pair	Ω , °	Distance, Å
GCN4	1, 2	24.3	9.4
PD ₁	1, 2	18.0	8.6
	1, 4	17.4	10.2
DHP-1	A, B	21.0	8.5
	A, D	18.6	9.9
DHP-14-AB	A, B	18.1	8.2
	A, D	9.0	9.9
IL-4 free	A, C	23.6	9.6
	A, D	31.1	9.8
IL-4 bound	A, C	27.6	9.6
	A, D	38.1	10.4

Helix packing interactions were calculated with WEBMOL (27) on these coordinates: IL-4 (1RCB), IL-4-IL-4R α (1IAR), DHP-1 (4HB1), GCN4 (2ZTA), and DHP-14-AB. Crossing angle values were added to 180° in order to make all values relative to parallel equal 0°. In order to represent the four-helix bundle of DHP-1, the DHP-1 asymmetric unit of two helices (A and B) had a twofold crystallographic symmetry operation applied to it.

toward the receptor in a similar orientation and position as IL-4 is to IL-4R α (Fig. 1), the first generation of mutations was designed onto the first two neighboring DHP-1 helices in the amino acid sequence.

To select critical binding site residues to incorporate onto DHP-1, the IL-4-IL-4R α structural analysis (3) and associated mutagenesis (28) were used to create a three-component scoring function to identify side-chains for transplantation onto DHP-1. Wang *et al.* characterized mutants in IL-4, which provided the relative changes in binding energy ($\Delta\Delta G$) associated with the loss of a side chain (28). The scoring function was based on three criteria: decrease in solvent accessible surface area ($\Delta SASA$) upon complex formation of >50 Å² (2 points), $\Delta\Delta G$ upon mutation to alanine of >0.3 kcal/mol (2 points), and a positively charged side chain (1 point). In total, fifteen residues were scored from IL-4 helices A and C: I5, T6, Q8, E9, K12, T13, Q78, R81, F82, K84, R85, R88, N89, W91, and G92. Side chain positions, which scored higher than 3 of the total 5 points, were selected for transplantation onto DHP-1.

Based on their score, seven residues were designed into DHP-1 corresponding to the following IL-4 residues: I5, E9, K12, K84, R81, R85, and R88. These residues represent 47% of the possible residues in the binding site on IL-4. W91 did not meet the three-point cut off, but was added to the design because it had a larger $\Delta\Delta G$ of 0.73 kcal/mol and it rearranged upon binding IL-4R α . Q78 was added to the design to remove a hydrophobic leucine side chain from the binding surface on DHP-1. Nine first-round mutations, 60% of the possible binding site residues, were made *in silico* onto two adjacent helices on DHP-1, hence termed DHP-9.

IL-4 helix B, the least interacting helix for IL-4R α binding, contributed one critical residue, R53, based upon a $\Delta SASA$ (decrease in solvent accessible surface area) upon binding of 50 Å², $\Delta\Delta G$ of 0.84 kcal/mol upon mutation to glutamine, and being positively charged (3, 28). R53 contributed to the receptor-binding interface through interaction with F41 of IL-4R α , which completes a hydrophobic collar around IL-4 R88 in the complex. Removal of IL-4 R88 resulted in a $\Delta\Delta G$ of $\approx$ 3.75 kcal/mol, which indicated its importance in binding and suggested that coordination of its position should be included in our design. Furthermore, IL-4 Y56, W91, and R53 rearrange upon binding to IL-4R α to bury IL-4 R88 in the interface. To simplify incorporation of R53, this side chain was placed in the model as close in 3D space as feasible, because DHP-1 did not have a helix that overlapped IL-4 helix B (Fig. 1). We expected that adding this approximation of R53 to our design of the binding interaction would improve mimicking of the most critical residue IL-4 R88. Thus, DHP-10 was designed to have a total of 10 IL-4R α binding mutations (Table 4).

Protein Design for Helical Bundle Topology. The goal of *de novo* protein design is to create a 3D model for a target fold with a corresponding linear amino acid sequence, synthesize it, and then determine its exact structure. The ambiguity of whether DHP-1 had right- or left-handed topology is due in part to the glycine linkers not being defined in the crystal structure and compounded by the repeating amino acid sequence of each helix. To resolve this topological uncertainty, two functional enantiomers were created toward a target with a specified fold, which was IL-4R α . DHP-1 and IL-4 both have the same four-helix core structures. Thus, IL-4 is compatible with our design efforts to resolve topology because wrapping around the DHP-1 and IL-4 four-helix bundle, each structure has the same inter-helical anti-parallel relationship. The IL-4R α binding function will require the side-chains to be displayed only on a unique set of adjacent helices in the DHP-1 template structure.

Two molecules were designed to address the two-way ambiguity in the topology between the helices of DHP-1 (Fig. 2). The

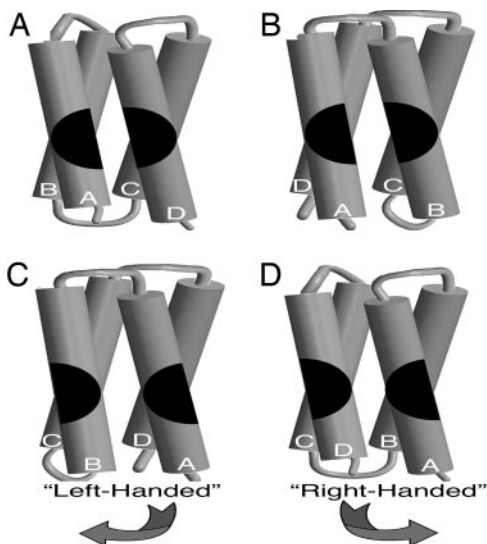


Fig. 2. Helical bundle scheme illustrates two possible topologies and the difference in the binding site depending upon the topology. As a matter of convention, orienting from the N-terminal helix, a left-handed bundle has the second helix in the bundle to the left; a right-handed bundle has the second helix in the bundle to the right (13). To imagine formation of a binding site for productive interaction with IL-4R α , two circle halves are split between neighboring helices in two different arrangements. When the circle is continuous and merging, the functional alignment of binding interactions is presented. When the circle is broken and diverging, the nonfunctional alignment of the binding interactions is presented. (A) The left-handed model forms a productive binding site when the circle halves could merge across the first and last helices. (B) The right-handed model forms a productive binding site when the circle halves could merge across the first and second helices. (C) The left-handed model does not form a productive binding site when the circle halves diverge across the first and last helices. (D) The right-handed model does not form a productive binding site when the circle halves diverge across the first and second helices.

spacing between helices in DHP-1, along with the designed length of loops between helices, targeted it to be, but did not confirm, a right-handed topology. DHP-1 itself was assembled from helices [termed PD₁ (29)], where short linkers of 3, 4, and 3-glycine residues were inserted between the four helices to maintain the inter-helical spacing observed in the PD₁ structure. The DHP-1 structure was maintained by its hydrophobic protein core and supported by “knobs into holes” side-chain packing interactions between these helices.

Creating IL-4R α binding function broke the DHP-1 sequence redundancy and structural pseudo 222-symmetry. DHP-10-AD carried the binding site on the adjacent helices A and D to represent the left-handed helical bundle (Fig. 2A); DHP-10-AB carried the binding site on the adjacent helices A and B to represent the right-handed helical bundle (Fig. 2B). The logic follows that, depending on which protein had stability and bound to IL-4R α , the DHP topology could be deduced. The protein with better stability and function would indicate that the binding site conformed to the restrictions of inter-helical packing interactions stabilizing the protein and that it was intact on one surface between adjacent helices.

First Generation: Stability and Function. To determine stability of the designed proteins in solution, the free energy of unfolding (ΔG_u^0) was measured by monitoring the loss of helical structure as a function of increasing denaturant concentration (Fig. 3). DHP-10-AB had stability similar to DHP-1 with ΔG_u^0 equal to 13.5 and 13.1 kcal/mol, respectively. These values of ΔG_u^0 are considered equivalent because the data, collected previously and

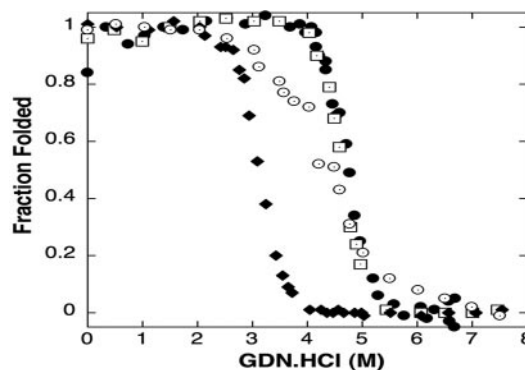


Fig. 3. Unfolding transition plot for each protein used to calculate ΔG_u^0 . DHP-10-AB (open squares), DHP-10-AD (open circles), and DHP-14-AB (filled diamonds) unfolding transitions were analyzed to determined free energy of unfolding, ΔG_u^0 . DHP-1 (filled circles) data were from Schafmeister *et al.* (12) and were realigned along with the new data for comparison.

reanalyzed, for DHP-1 had a narrower concentration range for the unfolded baseline >6 M Gdn·HCl as compared with DHP-10-AB and the unfolding transition overlaps. DHP-10-AD was less stable in solution, with $\Delta G_u^0 = 4.1$ kcal/mol (Table 1).

To access the aggregation state in solution, dynamic light scattering was used to measure the Rh and calculate the molecular mass for each protein. DHP-1 was mono-disperse with an average Rh of 1.79–1.80 nm and a molecular mass of 11.9–12 kDa, in agreement with the measured DHP-1 Rh determined by analytical ultracentrifugation (12). DHP-10-AB, which turned out to be the expected right-handed topology, was mono-disperse with an Rh of 1.78–1.95 nm and a calculated molecular mass of 11.9–12.7 kDa. In contrast, DHP-10-AD was poly-disperse at room temperature, which prevented an accurate determination of Rh and molecular mass.

To determine the IL-4R α binding activity, each protein was subject to competitive binding versus IL-4. DHP-1 did not compete with IL-4 for binding to IL-4R α over a concentration range from μ M to tens of mM. The IC₅₀ for DHP-10-AB was 495 μ M. The IC₅₀ for DHP-10-AD was 970 μ M (Table 1). Thus DHP-10-AB bound with approximately twice the affinity of DHP-10-AD but was still 5×10^5 times lower affinity than IL-4. This result is consistent with the right-handed topology.

Second Generation: Additional Mutations, Stability, and Function.

When the coordinates became available for the IL-4:IL-4R α complex, superimposing our model of DHP-10-AB with IL-4:IL-4R α , a number of residues in DHP-10-AB were found to overlap residues in IL-4R α (Fig. 6, which is published as supporting information on the PNAS web site). Therefore, additional mutations were added to the DHP-10-AB template to create DHP-14-AB. IL-4 residues T13, D87, N89, and G92 were incorporated into DHP-14-AB (Table 4). IL-4 G92 was mimicked with G92A within our model because glycine has the potential to destabilize the helix and alanine did not seem to overlap with IL-4R α in our model. On DHP-10-AB helix C, IL-4 R53 mimic was moved three residues toward the binding site, because the first R53 mimic pointed away from the rest of the IL-4R α binding site and its key residue analogous to IL-4 R88. DHP-14-AB was mono-disperse like DHP-10-AB with an Rh of 1.79–1.82 nm and a calculated molecular mass of 11.9–12.4 kDa. The unfolding free energy (ΔG_{U}^0) decreased from 13.5 kcal/mol to 8.1 kcal/mol, whereas the functional activity increased to IC₅₀ 27 μ M, 18 times the affinity of DHP-10-AB. DHP-14-AB was progressively improved as an IL-4 antagonist from DHP-1, which was stable and functionless, although still 2.7×10^4 times lower in affinity than IL-4 itself.

the knobs-into-holes packing of hydrophobic side-chains between the helices.

Based on modeling with the complex coordinates available, two additional areas were targeted for second-generation mutations to remove what were perceived as clashes with IL-4R α (Fig. 6). Larger side-chains were replaced with IL-4-mimetic residues within the interface. The mimic of IL-4 R53 was moved closer to the middle of the putative IL-4R α binding site to create a smaller interaction surface. These mutations applied to DHP-10-AB grafted most of the binding site for IL-4R α and improved the affinity for IL-4R α to an IC₅₀ of 27 μ M, which was surprisingly below the IC₅₀ for IL-4 in the assay at 1 nM.

Because DHP-1 helices overlapped the IL-4R α binding helices on free IL-4 within a 1-Å rms deviation, we thought it would be straightforward to create a reasonably high-affinity antagonist for IL-4. However, calculation of the crossing angle between helices highlighted another feature of the IL-4 interaction with IL-4R α . DHP-1 helices A and B had a measured inter-helix crossing angle that was nearly 3° smaller than for free IL-4. If the proteins are considered rigid bodies, this 3° crossing angle difference meant that some residues would be in nearly perfect overlap whereas others would be significantly displaced from the binding site. Upon binding, changes in crossing angle between the IL-4 binding site helices (A and C) increased by 4° (Table 3), yet IL-4 A and C helices were observed to shift by $\approx$ 0.5 Å upon binding. DHP-14-AB had a crossing angle between binding site helices (A and B) of 18.1°, which is 5.5° smaller than free IL-4 and 9.5° smaller than bound IL-4. The DHP-14-AB helices have short glycine linkers between helices and symmetric knobs-into-holes packing interactions due to alternating leucine and alanine residues. DHP-14-AB may be structurally restricted by inter-helix packing that prevented the helices from shifting to accommodate binding to IL-4R α . In contrast, the other peptide mimic of IL-4 based on GCN4 helices had a crossing angle of 24.3°, which is a closer match to either free or bound IL-4 than DHP-14-AB (Table 3). Thus, DHP-14-AB may not bind IL-4R α like IL-4 because the helices are not interacting with one another as they do in IL-4 and it may not be able to adapt to the binding site movements required for the high-affinity interaction.

The extraordinarily high IL-4 affinity for IL-4R α seemed to be a result of “electrostatic steering” by highly charged surfaces as well as coordinated movements between helices presenting the charged interface (Fig. 5). In addition to increased crossing angles between IL-4 helices A and C, the distribution of the charge on IL-4 shifts from a cluster to a specific arrangement coordinated by hydrogen bonds to IL-4R α . Even though we designed the same number of charges into the IL-4R α binding interface, more atomic precision is required to match the IL-4 affinity than what we designed into DHP-14-AB.

De novo design principles were applied to mimic the presentation of the functional binding site on IL-4. By designing two enantiomers and testing their stability and function, we resolved the ambiguity in the four-helix design template topology as right-handed and achieved an IC₅₀ of 495 μ M for IL-4R α . A subsequent design was based on the later structure of the IL-4-IL-4R α complex analysis to remove potentially detrimental interactions that could not be foreseen when using the structure of IL-4 alone. The resulting designed protein, DHP-14-AB (the largest designed protein to date), is well ordered, yields an instructive crystal structure, and has an IC₅₀ of 27 μ M. The apparent affinity of DHP-AB-14 is $\approx$ 10⁴ lower than IL-4, which may be due to transplantation of only a subset of IL-4 binding residues as well as the geometry of the helices interacting with IL-4R α . *De novo* protein design of function requires atomic precision in the binding site as well as incorporating the entire structure. We observed a difference in crossing angle between DHP-14-AB and IL-4 free and bound of 5.5° to 9.5° for the binding site helices, respectively. Accounting for secondary structure interactions and dynamics between those elements as measured in the differences in crossing angle may increase the precision of and ability to optimize protein-protein interactions.

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