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## Backbone structure of *Yersinia pestis* Ail determined in micelles by NMR-restrained simulated annealing with implicit membrane solvation

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### SUMMARY

The outer membrane protein Ail (attachment invasion locus) is a virulence factor of *Yersinia pestis* that mediates cell invasion, cell attachment and complement resistance. Here we describe its three-dimensional backbone structure determined in decyl-phosphocholine (DePC) micelles by NMR spectroscopy. The NMR structure was calculated using the membrane function of the implicit solvation potential, *eefxPot*, which we have developed to facilitate NMR structure calculations in a physically realistic environment. We show that the *eefxPot* force field guides the protein towards its native fold. The resulting structures provide information about the membrane-embedded global position of Ail, and have higher accuracy, higher precision and improved conformational properties, compared to the structures calculated with the standard repulsive potential.

### Keywords

Ail; *Yersinia pestis*; membrane protein; structure; NMR; implicit solvation

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*Yersinia pestis*, the bacterium that causes plague, is an extremely pathogenic organism with a long history of precipitating massive human pandemics (Perry and Fetherston 1997; Cornelis 2000; Brubaker 2004). *Y. pestis* is transmitted both via the bite of infected fleas (bubonic plague) and via aerosols (pneumonic plague) (Perry and Fetherston 1997), and can proliferate in diverse environments. This critical aspect of its virulence is governed by a number of virulence attributes that include Ail (attachment invasion locus), an outer membrane protein with established roles in cell invasion, cell attachment, complement resistance and virulence (Miller et al. 1990; Bartra et al. 2008; Felek et al. ; Hinnebusch et al. ; Kolodziejek et al. 2010).

Previous studies have shown that *Y. pestis* Ail binds the human extracellular matrix proteins fibronectin, laminin and heparan sulfate proteoglycans (Tsang et al. 2010; Yamashita et al.

2011; Tsang et al. 2012), as well as the complement regulatory component C4BP (Ho et al. 2014). The structure of Ail, crystallized from tetraethylene glycol mono-octyl ether (C<sub>8</sub>E<sub>4</sub>), has been determined at high resolution (Yamashita et al. 2011). The protein folds as an eight-stranded  $\beta$ -barrel with three intracellular loops (IL1-IL3) and four extracellular loops (EL1-EL4) formed by residues that are implicated in ligand binding (Miller et al. 2001; Tsang et al. 2013; Ding et al. 2015b). However, the functionally important extracellular loops EL2 and EL3 were incompletely resolved in the crystal structure.

Using NMR spectroscopy with Ail in decyl-phosphocholine (DePC) micelles, backbone resonances for all extracellular loop sites could be observed and assigned.

Heteronuclear <sup>1</sup>H/<sup>15</sup>N NOE experiments showed the presence of backbone flexibility in the extracellular loops, particularly EL2, consistent with their lack of resolution by crystallography (Ding et al. 2015b). Assignments of the backbone N, HN, CA and CB resonances were obtained for all Ail amino acids, with the exception of three residues in IL1 (Ile36, Phe37, Asn38) and two residues in IL2 (Glu83, Tyr84), and deposited in the Biological Magnetic Resonance data Bank (BMRB 25281). The NMR data showed that the Ail  $\beta$ -barrel is stabilized by a strong network of backbone amide hydrogen bonds that resist exchange with water, with stronger hydrogen bonds observed for residues located in a hydrophobic band around the  $\beta$ -barrel, and weaker, readily exchangeable hydrogen bonds, observed for sites that are water-exposed (Ding et al. 2015b). Furthermore, long-range amide hydrogen distances, derived from <sup>1</sup>H-<sup>1</sup>H NOE measurements were measured for both transmembrane and extracellular sites, between all neighboring  $\beta$ -strands, including key contacts between the first and last  $\beta$ -strand ( $\beta$ 1- $\beta$ 8) that define the barrel closure.

Here, we have used these NMR restraints to determine the backbone structure of Ail in DePC micelles. Two structures were calculated with XPLOR-NIH (Schwieters et al. 2003; Schwieters et al. 2006), one using the standard, purely repulsive form of the Van der Waals function (REPEL), with the default protein topology and parameter files (*protein.par* and *protein.top*), and the other using the implicit membrane function of the potential, *eefxPot*, with its dedicated protein topology and parameter files (*protein\_eef.par* and *protein\_eef.top*) (Tian et al. 2014, 2015). The structural coordinates and experimental restraints have been deposited in the Protein Data Bank (PDB: 2N2L, 2N2M).

## Structure calculations in the implicit membrane environment of *eefxPot*

NMR-restrained structure calculations are typically performed in a force-field where all nonbonded interactions are represented by a single, purely repulsive term, that does not include van der Waals attractive forces, electrostatic forces or solvation energy. This simplified potential is very successful for facilitating simulated annealing structure calculations from fully extended templates (Nilges et al. 1988; Clore and Gronenborn 1989), but does not provide a physically realistic solvent environment, especially for membrane proteins that reside in the highly heterogeneous and anisotropic environment of the lipid bilayer membrane.

The *eefxPot* potential, on the other hand, includes terms for Lennard-Jones van der Waals energy (both repulsive and attractive forces), electrostatic energy and solvation free energy.

The potential is based on the CHARMM effective energy function for implicit water and membrane solvation (Lazaridis and Karplus 1999; Lazaridis 2003). While the first version of *eefxPot* was limited to bulk water solvation (Tian et al. 2014), the most recent development includes the membrane and supports structure calculations of membrane proteins in an implicit membrane environment (Tian et al. 2015).

The membrane of *eefxPot* is modeled by a function that modulates the solvation and electrostatic parameters of all atoms according to their absolute distance along the membrane normal (coincident with the coordinate  $z$  axis) from the hydrophobic membrane center ( $z = 0$ ). This membrane profile function depends on the membrane hydrophobic thickness ( $T$ ) and a parameter ( $n$ ) that defines the size of the region over which the environment changes from hydrophobic to hydrophilic. The values of  $T=25$  Å and  $n=10$ , used in this study, give a membrane thickness that mimics the relatively thin bacterial outer membrane and a membrane profile similar to that derived from neutron and X-ray scattering (Luzzati and Husson 1962; Marsh 2013; Nagle and Tristram-Nagle 2000; Kucerka et al. 2011; Nagle 2013). The potential is accessible from the Python interface of XPLOR-NIH by calling *eefxPot* and its accessory module *eefxPotTools*. It may be downloaded from the web (<http://nmr.cit.nih.gov/xplor-nih/>) with the latest release of XPLOR-NIH (version 2.39).

Structures of Ail were calculated using two simulated annealing protocols, the first to fold 100 structures from an initially extended polypeptide conformation, and the second to refine 100 structures from the best folded structure. The final ensembles were generated by selecting the 10 best refined structures with lowest restraining energy terms, least number of violations and, in the case of *eefxPot*, proper membrane embedding.

The fold and refine protocols are based on the internal variable module (IVM) (Schwieters and Clore 2001) and share the same basic architecture comprising four stages: (i) torsion angle dynamics at high-temperature (3,500 K for fold, 3,000 K for refine), for a time of 32 ps or 32,000 timesteps (fold protocol), or a time of 20 ps or 20,000 timesteps (refine protocol); (ii) torsion angle dynamics with simulated annealing from the initial high temperature value to 25 K, in steps of 12.5 K, for a time of 0.4 ps or 200 timesteps per temperature step; (iii) 500 steps of Powell torsion angle minimization; and (iv) 500 steps of Powell Cartesian minimization.

Dihedral angle restraints were derived from analysis of the assigned chemical shifts with the program TALOS-N (Shen and Bax 2013) and imposed with the XPLOR term CDIH. Amide hydrogen bonds and amide hydrogen NOE distances were imposed with the distance restraint term *noePot*, side chain rotamers were restrained with the *torsionDB* statistical torsion angle potential (Bermejo et al. 2012), and standard potentials were implemented to define chemical bond lengths, bond angles, and atomic radii. In the high temperature stage, experimental restraints were applied with force constants of  $k_{CDIH}=100$  kcal mol<sup>-1</sup> rad<sup>-2</sup> (dihedral angles) and  $k_{DIST}=2$  kcal mol<sup>-1</sup> Å<sup>-2</sup> (distances). During simulated annealing,  $k_{CDIH}$  was set to 200 kcal mol<sup>-1</sup> rad<sup>-2</sup> and  $k_{DIST}$  was increased geometrically from 2 to 30 kcal mol<sup>-1</sup> Å<sup>-2</sup>. The *torsionDB* force constant was set to  $k_{tDB}=0.02$  kcal mol<sup>-1</sup> at high temperature stage and ramped geometrically from 0.02 to 2 kcal mol<sup>-1</sup> during simulated annealing.

During the *eefxPot* fold protocol, the 32 ps high temperature stage was divided into three blocks. This was done to prevent fatal atomic overlap in the early stages of calculations. Two initial 3 ps segments of dynamics were performed using REPEL with *eefxPot* turned off, the first with only CA-CA atomic interactions active, the van der Waals force constant set to  $C_{rep}=0.004 \text{ kcal mol}^{-1} \text{ \AA}^{-4}$  and the van der Waals radius scale factor set to  $k_{rep}=1.2$ , and the second with all atom-atom interactions turned on,  $C_{rep}=4 \text{ kcal mol}^{-1} \text{ \AA}^{-4}$  and  $k_{rep}=0.8$ . The third 26 ps segment of dynamics at high temperature, as well as the subsequent simulated annealing stage, were performed without REPEL, using exclusively *eefxPot* with its scale set to 0.004 at high temperature and then ramped from 0.004 to 1 during simulated annealing. The protein center of mass was positioned at the membrane center ( $z = 0$ ) before each block of the high temperature stage, while the protein position was left unrestrained during simulated annealing.

We have found that the timing and force constants of the fold protocol are particularly important for transmembrane  $\beta$ -barrels such as Ail, especially in cases where relatively few experimental restraints are available. In these proteins, sequential amino acids alternate in character between hydrophobic and hydrophilic and the membrane-spanning structural unit is formed by nonsequential residues far apart in the amino acid sequence. In nature,  $\beta$ -barrel folding and membrane insertion are assisted by dedicated machinery (Kleinschmidt 2015); here, they were assisted by extending the high temperature stage to allow attractive van der Waals forces to work, and imposing dihedral angle restraints more strongly in the early stages of the protocol. Once folded, the structure and membrane position of Ail were stably sustained by *eefxPot*.

## Structure of Ail in DePC micelles

Although the membrane model of *eefxPot* is intended to be used with restraints measured for membrane proteins in lipid bilayer membranes, it can also be applied to restraints measured in detergent micelles. In this study, the structure of Ail was determined in 170 mM DePC, at 45°C. This detergent gives the best solution NMR spectra (Ding et al. 2015b), but high quality solution NMR spectra can also be obtained for Ail incorporated in small lipid bilayer nanodiscs (Ding et al. 2015b), and larger nanodiscs reconstituted with Ail can be harvested by centrifugation as sedimented solutes to yield high quality solid-state NMR spectra (Ding et al. 2015a). Interestingly, even though the high detergent concentrations required for solution NMR spectroscopy are not compatible with the ligand binding activity of Ail (Ding et al. 2015b), it is notable that the solution and solid-state NMR spectra from micelles and nanodiscs all have very similar chemical shifts, indicating that the protein adopts the same overall fold in all three environments.

The NMR structure of Ail in micelles (Fig. 1) is also similar to the crystal structure (Yamashita et al. 2011). Both REPEL and *eefxPot* produce good structures, with acceptable conformational properties and statistics (Fig. 2; Table 1), but the *eefxPot* structures have higher accuracy and precision and also provide a view of the protein embedded in the membrane. The extracellular loops are more disordered than those of the crystal structure where they form an extended  $\beta$ sheet that extends out from the membrane. While the extent of loop dynamics will have to be assessed in detergent-free lipid bilayers, loop flexibility is

consistent with the versatile ligand-binding properties of Ail and with the incomplete observation of EL2 and EL3 electron density in the crystals.

In some of the REPEL structures the hydrophilic extracellular loops fold back towards the membrane and against the barrel body (Fig. 1A). Such conformations are not expected to exist in the native lipid bilayer membrane environment where the loops are known to engage ligands in the water phase. Previous NMR structure calculations of membrane proteins have sought to eliminate such anomalous conformations by establishing an artificial water-membrane boundary through the use of harmonic coordinate restraints (Teriete et al. 2007) or plane distance restraints (Xu et al. 2008; Fox et al. 2014; Gong et al. 2015), or with an empirical potential based on the preferred membrane insertion depth of each amino acid (Shi et al. 2009). However, these approaches do not provide a physical representation of the water-membrane environment. Notably, the structures calculated with *eefxPot* do not exhibit such unfavorable conformations, showing that the potential effectively guides the water-soluble extracellular loops away from the hydrophobic membrane interior and into the water phase.

In the *eefxPot* structure, the Ail  $\beta$ -barrel adopts a transmembrane tilt of  $7\pm 10^\circ$  (Fig. 1B) and a membrane insertion depth consistent with the  $^1\text{H}/^2\text{H}$  exchange data (Fig. 1D). Although the transmembrane tilt of the  $\beta$ -barrel will have to be confirmed experimentally, we note that *eefxPot* produces an accurate transmembrane position of the OmpX  $\beta$ -barrel, as assessed by cross validation with experimental  $^1\text{H}/^{15}\text{N}$  solid-state NMR orientation restraints (Tian et al. 2015). The transmembrane core of the Ail  $\beta$ -barrel is well defined by the  $^1\text{H}/^2\text{H}$  exchange data and delimited by two bands of aromatic residues, each located at the membrane-water interfaces. The membrane-embedded position of the barrel places a ring of positively charged Lys and Arg sidechains near the extracellular membrane-water interface (Fig. 1D). In the crystal structure, these sites can associate with sucrose octasulfate, a heparin analog small molecule (Yamashita et al. 2011). Their location at the membrane surface suggests that they can also play a role in mediating the interactions of Ail with other membrane components such as phospholipids and lipopolysaccharides.

## Effect of *eefxPot* on structural accuracy, precision and quality

The NMR structures of Ail calculated with *eefxPot* have significantly closer agreement to the crystal structure (Fig. 2A, Table 1). As transmembrane  $\beta$ -barrel conformations are relatively refractory, the crystal structure of Ail, which was determined with rather high resolution, is likely a good representative of the native conformation, thus, we infer that closer agreement to the crystal structure reflects higher structural accuracy. The precision of the *eefxPot* structural ensemble is also significantly higher than the REPEL ensemble (Fig. 2B, Table 1). Taken together, these results reflect the ability of *eefxPot* to effectively guide the conformation of Ail towards native fold, as observed for other soluble and membrane proteins (Tian et al. 2014, 2015).

Furthermore, although introducing an additional term in the target energy function can worsen the agreement between calculated structures and the other experimental and conformational energy terms, we find that the improvements in accuracy and precision

afforded by *eefxPot* are obtained without sacrificing agreement with either conformational or experimental restraints. Both *eefxPot* and REPEL structures have comparable RMSDs to the experimental restraints (Fig. 2C, Table 1), well within the acceptable range, with the *eefxPot* structures showing agreements that are better for distance restraints and only slightly lower for dihedral angles.

Assessment of the backbone conformations, side chain conformations and nonbonded atomic interactions by means of WHAT-IF (Vriend 1990; Doreleijers et al. 2012) and MolProbity (Chen et al. 2010) metrics shows that *eefxPot* yields structures with overall higher conformational quality (Fig. 2D, Fig. 2E), reflected in increased population of the favored regions of Ramachandran space and lower number of Ramachandran outliers. Furthermore, although both *eefxPot* and REPEL produce acceptable  $\chi^1/\chi^2$  rotamer normality scores, an effect due primarily to the *torsionDB* potential of XPLOR-NIH (Bermejo et al. 2012), the *eefxPot* structures appear to have fewer poor rotamers.

The validation metrics further show that *eefxPot* improves the quality of nonbonded atomic interactions. The structures of Ail calculated with *eefxPot* show improvements in both the WHAT-IF packing quality, which reflects the atomic distributions around different molecular fragments (Vriend and Sander 1993), and the MolProbity clashscore, which reflects the number of serious atomic overlaps per thousand atoms (Word et al. 1999). These key metrics provide database-independent estimates of the quality of nonbonded atomic interactions or atomic packing. The overall improvement is reflected in the lower MolProbity score - the lower the better – that is obtained with *eefxPot*.

## Conclusions

We conclude that the implicit membrane environment of *eefxPot* supports the native structure of the outer membrane protein Ail. The resulting structures have higher accuracy, higher precision and better overall quality. Although *eefxPot* is designed to be used with experimental restraints measured in detergent-free lipid bilayer samples, the results obtained with Ail show that it is compatible with measurements made in detergent micelles, where it is useful for imposing boundaries between the water and membrane environments and to establish proper protein membrane-embedded topology. The structure of Ail calculated with this implicit membrane model informs our understanding of the protein in the bacterial outer membrane.

It is also important to note that since the lipid bilayer membrane of *eefxPot* is modeled implicitly, its ability to reflect the structural and physical properties of natural membranes is limited (Tian et al. 2015). The principal limitations of the model are related to its inadequate representation of the hydrophobic matching phenomenon. In nature, match of a protein's amino acid sequence to the hydrophobic environment of the surrounding membrane is achieved by adjusting the protein's global position (tilt and rotation) in the membrane and/or adjusting the lipid bilayer's structure and order (Mouritsen and Bloom 1984; de Planque and Killian 2003). While the first mechanism is reproduced very well by *eefxPot*, the second is not, because membrane deformation is difficult to model implicitly.

Implicit membrane models cannot adequately reproduce the effects of lipid composition, aqueous phase composition, temperature, etc., on the structural and physical properties of lipid bilayers. Membrane proteins that form water-containing cavities also pose a challenge, although this can be addressed by mathematically shaping the implicit membrane to form a water-filled pore (Lazaridis 2005). In the case of detergent micelles, the structure of the hydrophobic environment departs even further from the continuous membrane model of *eefxPot*, as it is far from planar, much more fluid, and contains much more water than a lipid bilayer. A solution to this problem may be to perform calculations where the water phase is represented implicitly with *eefxPot*, and both the protein and the detergent molecules are treated in atomistic detail, as described for molecular dynamics (MD) simulations (Versace and Lazaridis 2015).

Nevertheless, while these limitations are significant in MD simulations performed free of the influence of experimental restraints, they become much less important when *eefxPot* is used for its intended purpose of assisting NMR-restrained structure calculations and the force field is not alone in determining protein structure. NMR structures calculated with *eefxPot*, in turn, can provide much improved starting points for MD simulations with all-atom force fields, designed to obtain atomic-level structure refinement and information about protein-lipid or protein-water interactions.

In all cases of soluble and membrane proteins tested so far (Tian et al. 2014, 2015), *eefxPot* outperforms the standard REPEL force field: it yields structures with higher accuracy, precision and conformational quality and is very effective at guiding proteins towards their native fold, even in the absence of large numbers of experimental restraints. Furthermore, as *eefxPot* is developed for XPLOR-NIH, it is fully compatible with all the experimental restraining terms available in the program, as well as with the full range of molecular parameters and topologies relevant for NMR, such as paramagnetic spin labels, for example.

These initial results are quite compelling in demonstrating that *eefxPot* is useful for NMR calculations. Indeed, there seems to be no reason not to use it, especially since it is as computationally efficient as REPEL and requires no or minimal additional setup. Further developments of the *eefxPot* membrane profile function to model water occlusion, surface charge, and micelle-like spherical or oblong structures will further extend its applications to NMR studies of membrane proteins.

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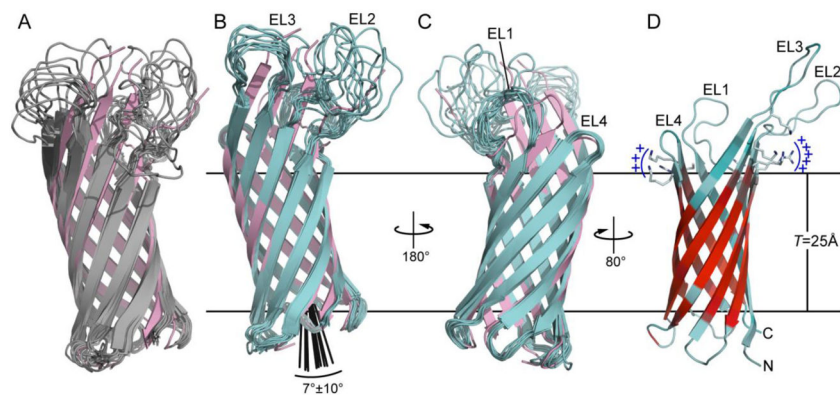
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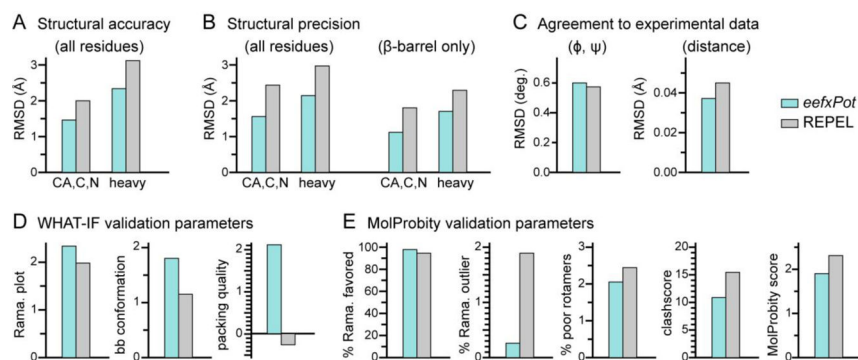
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**Figure 1. Structure of Ail determined by NMR in DePC micelles**

Structures were calculated using *eefxPot* (cyan) or REPEL (gray), or taken from the crystallographic coordinates (pink; PDB 3QRA). The *eefxPot* membrane is depicted as horizontal lines separated by the membrane thickness  $T=25\text{\AA}$ . Each ensemble is aligned to its lowest energy structure. **(A-C)** Ensembles of 10 best structures calculated with REPEL (A) or *eefxPot* (B, C). The crystal structure is aligned to the lowest energy model of each ensemble. **(D)** Lowest energy structure from *eefxPot*, showing the four extracellular loops (EL1-EL4) and the two juxtaposed patches of positively charged residues near the extracellular membrane surface. The red-cyan color spectrum reflects the  $^1\text{H}/^2\text{H}$  exchange profile of amide hydrogens that resist (red) or undergo (cyan) exchange with solvent. Structures were rendered with PyMol (DeLano 2005)



**Figure 2.** Structure statistics and validation metrics. Bars represent results obtained with *eefxPot* (cyan) or REPEL (gray). Statistics were evaluated for the ten structures in each refined ensemble. **(A)** Structural accuracy evaluated as pairwise RMSD of backbone or heavy atoms between the calculated structures and the crystal structure (PDB: 3QRA). **(B)** Structural precision evaluated as average pairwise RMSD of backbone or heavy atoms, for all residues or  $\beta$ -barrel residues only. **(C)** Agreement between calculated structures and experimental dihedral angle ( $\phi$ ,  $\psi$ ) and distance (NOE and H-bond) restraints. **(D)** WHAT-IF validation statistics for Ramachandran plot appearance, backbone conformation and protein packing quality. **(E)** MolProbity validation statistics for percent of residues in favored regions of the Ramachandran plot, percent of residues in unfavored regions of the Ramachandran plot, percent of residues with poor sidechain torsion angles, clashscore, and overall Molprobity score. The MolProbity clashscore and MolProbity score are costs: the lower the better.

**Table 1**Structure and NMR restraints statistics.<sup>a</sup>

	<i>efxPot</i>	REPEL
<b>Experiment Restraints</b>		
dihedral angles	294	294
NOE distances (i-j>5)	146	146
H-bonds	77	77
<b>RMSD Agreement with Experimental restraints</b>		
dihedral angle (deg)	0.598±0.039	0.570±0.094
NOE (Å)	0.040±0.003	0.044±0.003
H-bond (Å)	0.028±0.006	0.047±0.003
<b>Idealized Covalent Geometry RMSD</b>		
bonds (Å)	0.995±0.001	1.036±0.000
angles (deg)	0.267±0.002	0.537±0.002
impropers (deg)	0.320±0.008	0.645±0.005
<b>Coordinate Precision as average RMSD from mean (Å)<sup>b</sup></b>		
backbone CA, C, N atoms	1.586	2.451
all heavy atoms	2.164	2.960
<b>Ramachandran Plot <math>\phi/\psi</math> Angle Statistics</b>		
most favored regions	92.9%	92.2%
additional allowed regions	5.8%	4.8%
generously allowed regions	0.8%	1.7%
disallowed regions	0.5%	1.4%

<sup>a</sup> Calculated for 10 structures, out of 100 refined structures, selected for lowest energy and lowest experimental restraint violations.

<sup>b</sup> Calculated for all 156 protein residues.