Analyses performed for DAOP well-defined residues.

Procheck analysis, RMSD calculation and structure superimposition are based on DAOP with S(phi)+S(psi)>=1.8: 10A-17A, 20A-45A, 47A-59A

NESG ID: NAME

PDB ID:

Deposition date:

Common Name:

Class:

Length (a.a.): 61

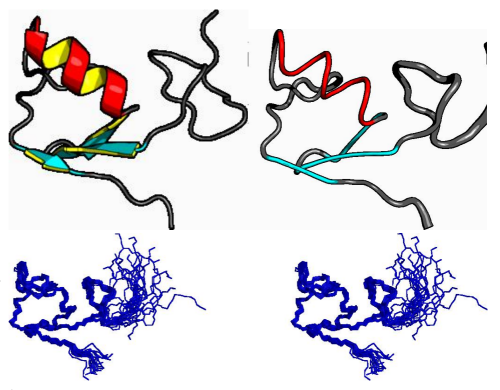
Organism: SwissProt / **TrEMBL ID:**

models: 20

Oligomerization: monomer

Molecular weight:

6874



Secondary Structure Elements:

alpha helices: 32A-38A

beta strands: 28A-30A, 21A-24A, 54A-57A

FIDs deposited in the BMRB? no

Comparison of core atoms:

DAOP > 1.8 Å: A:10..A:17, A:20..A:44, A:49..A:59

FindCore2: A:8..A:60 **CYRANGE:** 10..58

RMSD Ordered residues² Selected residues³ All residues

1.9 Å 0.5 Å0.5 ÅAll backbone atoms 2.3 Å 0.9~Å0.9 Å All heavy atoms

Ramachandran Plot Summary for selected residues³ from Procheck

Most favoured Additionally allowed regions Generously allowed regions Disallowed regions regions

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21.8% 0.5% 2.0%

Ramachandran Plot Summary for selected residues³ from Richardson Lab's Molprobity

Most favoured regions Allowed regions Disallowed regions View plot View model summary 86.4% 13.4% 0.2%

Global quality scores

Program	Verify3D	ProsaII (-ve)	Procheck (phi-psi) ³	$Procheck (all)^3$	MolProbity Clashscore
Raw score	0.05	0.20	-0.82	-0.47	10.08
Z-score ¹	-6.58	-1.86	-2.91	-2.78	-0.20

Close Contacts and Deviations from Ideal Geometry (from PDB validation software)

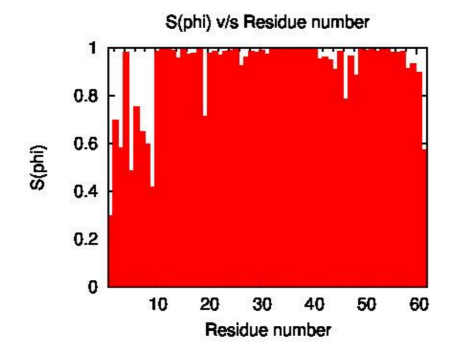
Number of close contacts (within 1.6 & Aring for H atoms, 2.2 & Aring for heavy atoms):

RMS deviation for bond angles: 1.3°

RMS deviation for bond lengths:

0.019 Å

³ Selected residues DAOP with S(phi)+S(psi)>=1.8:10A-17A, 20A-45A, 47A-59A

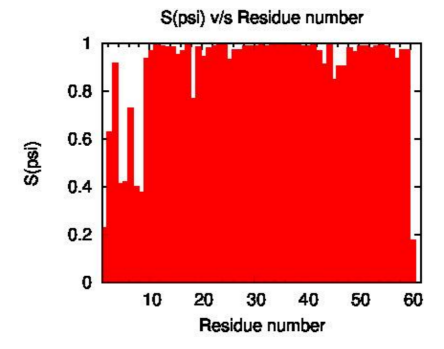


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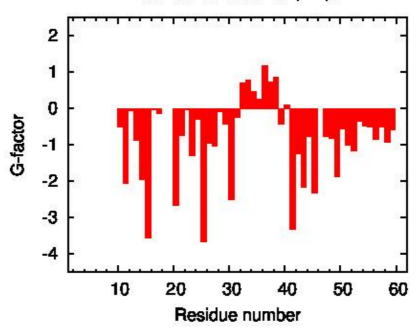
¹ With respect to mean and standard deviation for a set of 252 X-ray structures < 500 residues, of resolution <= 1.80 Å, R-factor <= 0.25 and R-free <= 0.28; a positive value indicates a 'better' score

² Ordered residues (DAOP): 10A-17A, 20A-45A, 47A-59A





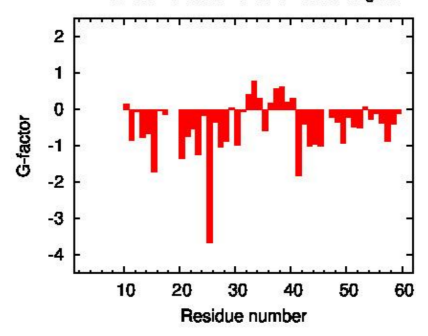
Procheck G-factor for phi-psi

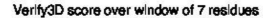


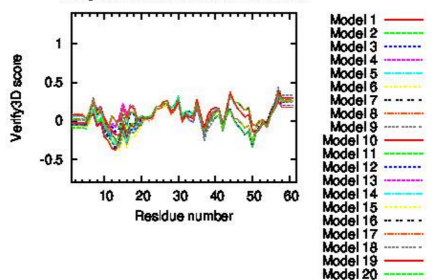




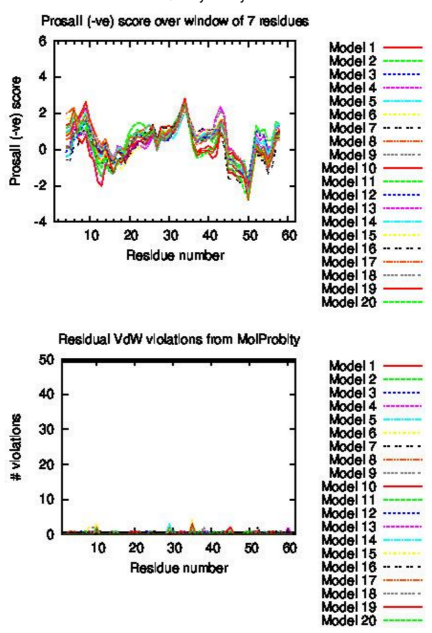
Procheck G-factor for all dihedral angles



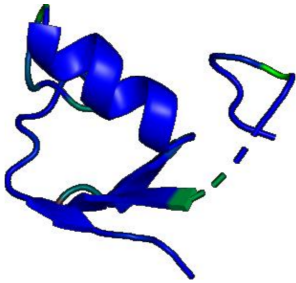












Residue Plot of Ramachandran anlysis(based on data from Richardson Lab's Molprobity)

References:

- 1. Bhattacharya A, Tejero R and Montelione GT, "Evaluating protein structures determined by structural genomics consortia". Proteins 2007, 66:778-795
- 2. Tejero R, Snyder D, Mao B, Aramini JM and Montelione GT, "PDBStat: a universal restraint converter and restraint analysis software package for protein NMR". J Biomol NMR 2013, 56:337-351
- 3. Luthy R, Bowie JU and Eisenberg D, "Assessment of protein models with three-dimensional profiles". Nature 1992, 356:83-85
- 4. Bowie JU, Luthy R and Eisenberg D, "A Method to identify protein sequences that fold into a known three-dimensional structure". Science 1991, 253:164-169
- 5. Sippl MJ, "Recognition of errors in three-dimensional structures of proteins". Proteins 1993, 17:355-362
- 6. Sippl MJ, "Calculation of conformation ensembles from potentials of mean force". J Mol Biol 1990, 213:859-883
- 7. Laskowski RA et al, "AQUA and PROCHECK_NMR: programs for checking the quality of proteins structures solved by NMR". J Biomolec NMR 1996, 8:477-486
- 8. Laskowski RA et al "PROCHECK: a program to check the stereochemical quality of protein structures". J Appl Cryst 1993, 26:283-291
- 9. Word JM et al, "Exploring steric constrains on protein mutations using MAGE / PROBE". Prot Sci 2000, 9:2251-2259 10. Word JM et al, "Asparagine and glutamine: using hydrogen atom contacts in the choice of side-chain amide orientation". J Mol Biol 1999, 285:1735-1747
- 11. Word JM et al, "Visualizing and quantifying molecular goodness-of-fit: small-probe contact dots with explicit hydrogens". J Mol Biol 1999, 285:1711-1733
- 12. Luthy R, McLachlan AD and Eisenberg D, "Secondary structure-based profiles: use of structure-conserving scoring tables in searching protein sequence databases for structural similarities". Proteins 1991, 18:229-239
- 13. Richardson DC, Richardson J S, "The kinemage: a tool for scientific communication". Prot Sci 1992, 1(1):3-9
- 14. Guntërt P, Mumenthaler, C & Wüthrich, K "Torsion angle dynamics for NMR structure calculation with the new program DYANA". J. Mol. Biol 1997, 273:283-298
- 15. Lovell SC et al, "Structure validation by Calpha geometry: phi,psi and Cbeta deviation". Proteins 2003, 50:437-450
- 16. Kabsch W, Sander C, "Dictionary of protein secondary structure: pattern recognition of hydrogen-bonded and geometrical features". Biopolymers 1983, 22:2577-2637
- 17. Bagaria A, Jaravine, V, Huang YJ, Montelione, GT, and Guntert, P "Protein structure validation by generalized linear model root-mean-square deviation prediction". Protein Sci 2012) 21:229-238.



Protein Structure Validation Suite (PSVS)





Software Environment

Software for structure quality evaluation:

DSSP DsspCMBI-April-2000 PdbStat-5.20.8 Version pdbstat

AutoAssign AutoAssign Version 2.4.0 (uses only AVS scripts) RPF analysis ASDP-2.3

PDB validation Version 8.061 Verify3D Version 1.0 corrected by Aneerban

Verify3D Version 1.0 c ProsaII Prosa2003 PROCHECK Version 3.5.4

MolProbity programs:

cluster 1999

clashlistcluster 1999 (corrected by Aneerban)

mage Version 0.35.040406

Version 6.35.040406

reduce Version 2.14 Version 2.6 probe

Other Software:

PERL Version 5.16.3 PEKL convert ImageMagick 6.7.8 Ghostscript 9.25 ps2pdf

v1.8.28 htmldoc

gnuplot Version 4.6.2

netpbm-progs 10.79.00 jpegtopnm netpbm-progs 10.79.00 pnmcrop pnmtojpeg netpbm-progs 10.79.00

Information updated on October, 2021. RTT

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