Analyses performed for DAOP well-defined residues.

Procheck analysis, RMSD calculation and structure superimposition are based on DAOP with S(phi)+S(psi)>=1.8:4A-60A

**NESG ID:** NAME

PDB ID:

**Deposition date:** 

**Common Name:** 

Class:

**Length (a.a.):** 61

Organism:
SwissProt /
TrEMBL ID:

# models: 5

**Oligomerization:** monomer

Molecular

weight: 6874

**Secondary Structure Elements:** 

alpha helices: 32A-41A

beta strands: 29A-30A, 21A-24A, 54A-57A

FIDs deposited in the BMRB? no

**Comparison of core atoms:** 

**DAOP > 1.8 Å:** A:4..A:5, A:7..A:60

FindCore2: A:1..A:61\*\*Selecte

**CYRANGE:** 9...59

RMSD All residues Ordered residues<sup>2</sup> Selected residues<sup>3</sup>

 All backbone atoms
 0.9 Å
 0.4 Å
 0.4 Å

 All heavy atoms
 1.2 Å
 0.6 Å
 0.6 Å

Ramachandran Plot Summary for selected residues<sup>3</sup> from Procheck

Most favoured regions Additionally allowed regions Generously allowed regions Disallowed regions

Ramachandran Plot Summary for selected residues<sup>3</sup> from Richardson Lab's Molprobity

Most favoured regions Allowed regions Disallowed regions View plot View model summary

92.3% 6% 1.8%

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Program	Verify3D	ProsaII (-ve)	Procheck (phi-psi) <sup>3</sup>	Procheck (all) <sup>3</sup>	MolProbity Clashscore
Raw score	0.07	N/A	-0.61	-0.37	N/A
Z-score <sup>1</sup>	-6.26	N/A	-2.08	-2.19	N/A

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

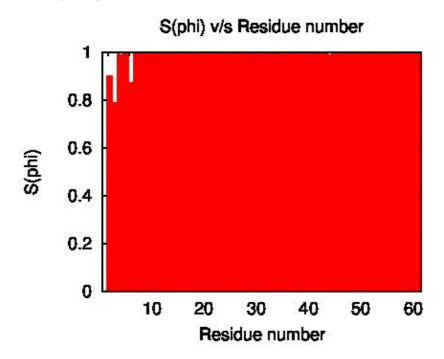
Number of close contacts (within 1.6 & Aring for H atoms, 2.2 & Aring for heavy atoms):

RMS deviation for bond angles:  $2.0^{\circ}$ 

RMS deviation for bond lengths:

0.011 Å

<sup>&</sup>lt;sup>3</sup> Selected residues DAOP with S(phi)+S(psi)>=1.8:4A-60A



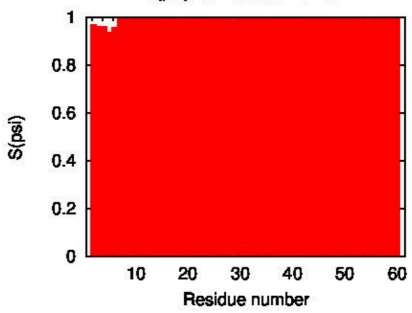
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 $<sup>^1</sup>$  With respect to mean and standard deviation for a set of 252 X-ray structures < 500 residues, of resolution <= 1.80 Å, R-factor <= 0.25 and R-free <= 0.28; a positive value indicates a 'better' score

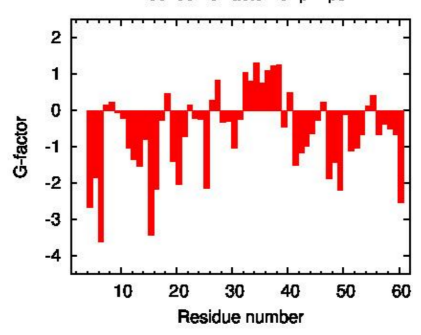
<sup>&</sup>lt;sup>2</sup> Ordered residues (DAOP): 4A-60A







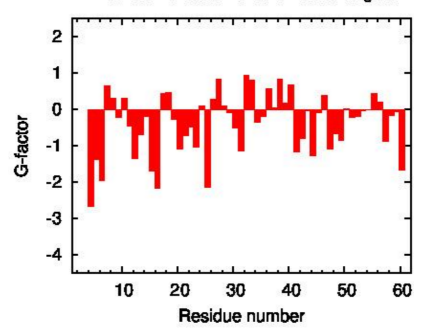
## Procheck G-factor for phi-psi

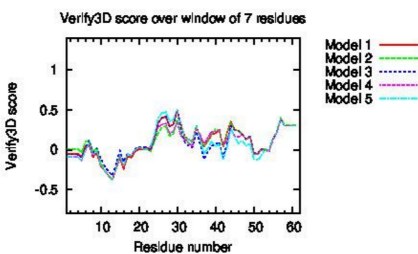


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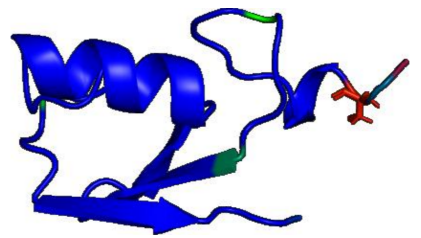
## Procheck G-factor for all dihedral angles





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

#### References:

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# Protein Structure Validation Suite (PSVS)



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## **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 6.35.040405 prekin Version 6.35.040406

reduce Version 2.14 Version 2.6 probe

## Other Software:

PERL convert PERL Version 5.16.3 ImageMagick 6.7.8 Ghostscript 9.25

htmldoc v1.8.28

gnuplot Version 4.6.2

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

Information updated on October, 2021. RTT

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