



Full wwPDB NMR Structure Validation Report ⓘ

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PDB ID : 2LQ2 / pdb_00002lq2
BMRB ID : 18289
Title : Solution structure of de novo designed peptide 4m
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Deposited on : 2012-02-22

This is a Full wwPDB NMR Structure Validation Report for a publicly released PDB entry.

We welcome your comments at validation@mail.wwpdb.org

A user guide is available at

<https://www.wwpdb.org/validation/2017/NMRValidationReportHelp>

with specific help available everywhere you see the ⓘ symbol.

The types of validation reports are described at

<http://www.wwpdb.org/validation/2017/FAQs#types>.

The following versions of software and data (see [references ⓘ](#)) were used in the production of this report:

MolProbity : 4-5-2 with Phenix2.0
Percentile statistics : 20250101.v01 (using entries in the PDB archive January 1st 2025)
wwPDB-RCI : v_1n_11_5_13_A (Berjanski et al., 2005)
PANAV : Wang et al. (2010)
wwPDB-ShiftChecker : v1.2
BMRB Restraints Analysis : v1.2
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP) : 2.49

2 Ensemble composition and analysis

This entry contains 20 models. Model 9 is the overall representative, medoid model (most similar to other models). The authors have identified model 1 as representative, based on the following criterion: *fewest violations*.

The following residues are included in the computation of the global validation metrics.

Well-defined (core) protein residues			
Well-defined core	Residue range (total)	Backbone RMSD (Å)	Medoid model
1	A:2-A:27 (26)	0.07	9

Ill-defined regions of proteins are excluded from the global statistics.

Ligands and non-protein polymers are included in the analysis.

The models can be grouped into 3 clusters and 3 single-model clusters were found.

Cluster number	Models
1	6, 7, 8, 9, 12, 13, 16, 18, 19
2	3, 4, 5, 10, 11, 17
3	1, 2
Single-model clusters	14; 15; 20

3 Entry composition [i](#)

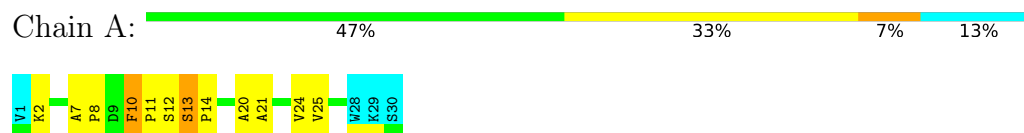
There is only 1 type of molecule in this entry. The entry contains 438 atoms, of which 222 are hydrogens and 0 are deuteriums.

- Molecule 1 is a protein called de novo designed antifreeze peptide 4m.

Mol	Chain	Residues	Atoms					Trace
1	A	30	Total	C	H	N	O	0
			438	139	222	37	40	

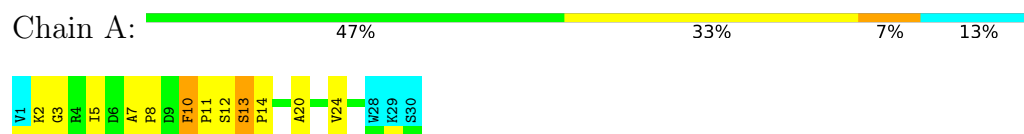
4.2.18 Score per residue for model 18

- Molecule 1: de novo designed antifreeze peptide 4m



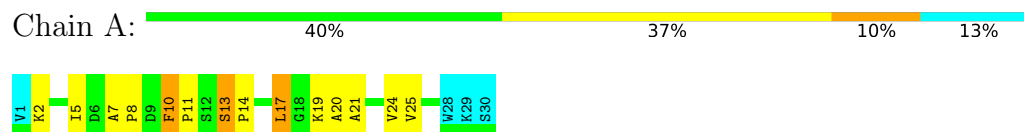
4.2.19 Score per residue for model 19

- Molecule 1: de novo designed antifreeze peptide 4m



4.2.20 Score per residue for model 20

- Molecule 1: de novo designed antifreeze peptide 4m



5 Refinement protocol and experimental data overview

The models were refined using the following method: *distance geometry*.

Of the 100 calculated structures, 20 were deposited, based on the following criterion: *target function*.

The following table shows the software used for structure solution, optimisation and refinement.

Software name	Classification	Version
DYANA	refinement	1.5

The following table shows chemical shift validation statistics as aggregates over all chemical shift files. Detailed validation can be found in section 7 of this report.

Chemical shift file(s)	working_cs.cif
Number of chemical shift lists	1
Total number of shifts	81
Number of shifts mapped to atoms	81
Number of unparsed shifts	0
Number of shifts with mapping errors	0
Number of shifts with mapping warnings	0
Assignment completeness (well-defined parts)	23%

6.3 Torsion angles

6.3.1 Protein backbone

In the following table, the Percentiles column shows the percent Ramachandran outliers of the chain as a percentile score with respect to all PDB entries followed by that with respect to all NMR entries. The Analysed column shows the number of residues for which the backbone conformation was analysed and the total number of residues.

Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
1	A	26/30 (87%)	26±0 (100±0%)	0±0 (0±0%)	0±0 (0±0%)	100	100
All	All	520/600 (87%)	520 (100%)	0 (0%)	0 (0%)	100	100

There are no Ramachandran outliers.

6.3.2 Protein sidechains

In the following table, the Percentiles column shows the percent sidechain outliers of the chain as a percentile score with respect to all PDB entries followed by that with respect to all NMR entries. The Analysed column shows the number of residues for which the sidechain conformation was analysed and the total number of residues.

Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
1	A	18/22 (82%)	14±1 (78±7%)	4±1 (22±7%)	2	29
All	All	360/440 (82%)	280 (78%)	80 (22%)	2	29

All 8 unique residues with a non-rotameric sidechain are listed below. They are sorted by the frequency of occurrence in the ensemble.

Mol	Chain	Res	Type	Models (Total)
1	A	10	PHE	20
1	A	2	LYS	19
1	A	13	SER	17
1	A	12	SER	9
1	A	17	LEU	6
1	A	5	ILE	5
1	A	4	ARG	2
1	A	19	LYS	2

6.3.3 RNA

There are no RNA molecules in this entry.

6.4 Non-standard residues in protein, DNA, RNA chains [i](#)

There are no non-standard protein/DNA/RNA residues in this entry.

6.5 Carbohydrates [i](#)

There are no oligosaccharides in this entry.

6.6 Ligand geometry [i](#)

There are no ligands in this entry.

6.7 Other polymers [i](#)

There are no such molecules in this entry.

6.8 Polymer linkage issues [i](#)

There are no chain breaks in this entry.

7 Chemical shift validation

The completeness of assignment taking into account all chemical shift lists is 23% for the well-defined parts and 21% for the entire structure.

7.1 Chemical shift list 1

File name: working_cs.cif

Chemical shift list name: *assigned_chem_shift_list_1*

7.1.1 Bookkeeping

The following table shows the results of parsing the chemical shift list and reports the number of nuclei with statistically unusual chemical shifts.

Total number of shifts	81
Number of shifts mapped to atoms	81
Number of unparsed shifts	0
Number of shifts with mapping errors	0
Number of shifts with mapping warnings	0
Number of shift outliers (ShiftChecker)	0

7.1.2 Chemical shift referencing

No chemical shift referencing corrections were calculated (not enough data).

7.1.3 Completeness of resonance assignments

The following table shows the completeness of the chemical shift assignments for the well-defined regions of the structure. The overall completeness is 23%, i.e. 72 atoms were assigned a chemical shift out of a possible 317. 0 out of 3 assigned methyl groups (LEU and VAL) were assigned stereospecifically.

	Total	¹ H	¹³ C	¹⁵ N
Backbone	46/126 (37%)	46/51 (90%)	0/52 (0%)	0/23 (0%)
Sidechain	26/181 (14%)	26/120 (22%)	0/56 (0%)	0/5 (0%)
Aromatic	0/10 (0%)	0/5 (0%)	0/5 (0%)	0/0 (—%)
Overall	72/317 (23%)	72/176 (41%)	0/113 (0%)	0/28 (0%)

The following table shows the completeness of the chemical shift assignments for the full structure. The overall completeness is 21%, i.e. 81 atoms were assigned a chemical shift out of a possible 378. 0 out of 4 assigned methyl groups (LEU and VAL) were assigned stereospecifically.

8 NMR restraints analysis

8.1 Conformationally restricting restraints

The following table provides the summary of experimentally observed NMR restraints in different categories. Restraints are classified into different categories based on the sequence separation of the atoms involved.

Description	Value
Total distance restraints	148
Intra-residue ($ i-j =0$)	42
Sequential ($ i-j =1$)	78
Medium range ($ i-j >1$ and $ i-j <5$)	28
Long range ($ i-j \geq 5$)	0
Inter-chain	0
Hydrogen bond restraints	0
Disulfide bond restraints	0
Total dihedral-angle restraints	0
Number of unmapped restraints	0
Number of restraints per residue	4.9
Number of long range restraints per residue ¹	0.0

¹Long range hydrogen bonds and disulfide bonds are counted as long range restraints while calculating the number of long range restraints per residue

8.2 Residual restraint violations

This section provides the overview of the restraint violations analysis. The violations are binned as small, medium and large violations based on its absolute value. Average number of violations per model is calculated by dividing the total number of violations in each bin by the size of the ensemble.

8.2.1 Average number of distance violations per model

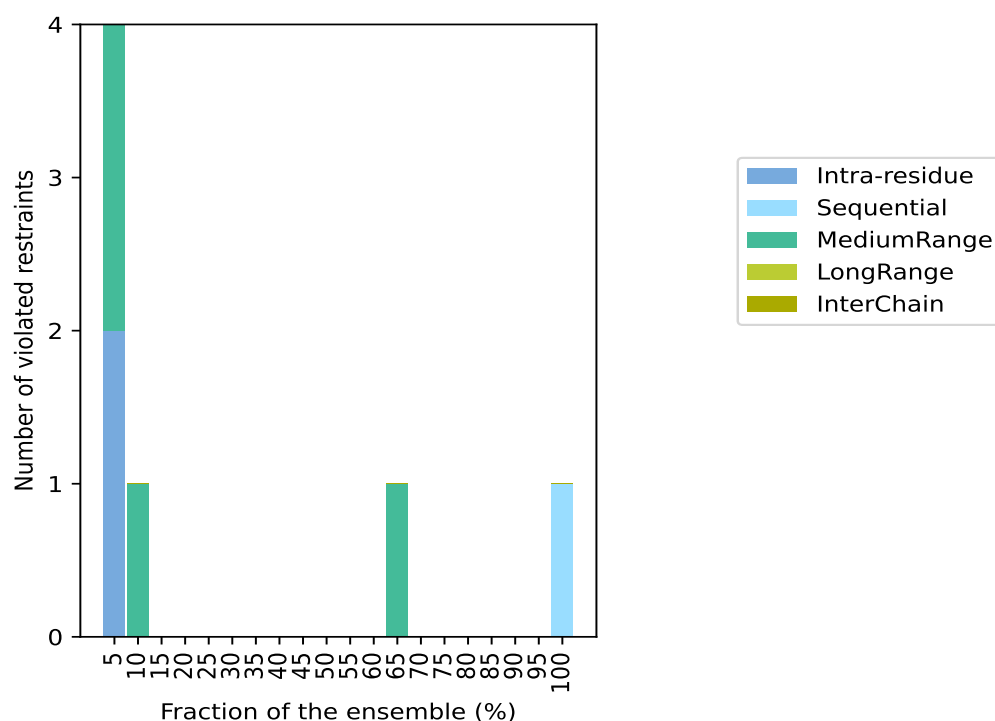
Distance violations less than 0.1 Å are not included in the calculation.

Bins (Å)	Average number of violations per model	Max (Å)
0.1-0.2 (Small)	1.6	0.19
0.2-0.5 (Medium)	None	None
>0.5 (Large)	None	None

8.2.2 Average number of dihedral-angle violations per model [i](#)

Dihedral-angle violations less than 1° are not included in the calculation. There are no dihedral-angle violations

9.3.1 Bar graph : Distance violation statistics for the ensemble [i](#)



9.4 Most violated distance restraints in the ensemble [i](#)

9.4.1 Histogram : Distribution of mean distance violations [i](#)

The following histogram shows the distribution of the average value of the violation. The average is calculated for each restraint that is violated in more than one model over all the violated models in the ensemble

Continued from previous page...

Key	Atom-1	Atom-2	Model ID	Violation (Å)
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	6	0.12
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	8	0.12
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	10	0.12
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	12	0.12
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	14	0.12
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	15	0.12
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	17	0.12
(1,6)	1:2:A:LYS:HA	1:3:A:GLY:H	18	0.12
(1,104)	1:25:A:VAL:HB	1:25:A:VAL:H	19	0.11
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	6	0.11
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	7	0.11
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	9	0.11
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	13	0.11
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	16	0.11
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	18	0.11
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	19	0.11
(1,73)	1:16:A:ILE:HA	1:19:A:LYS:H	2	0.1
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	8	0.1
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	12	0.1
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	15	0.1
(1,70)	1:15:A:ALA:HA	1:18:A:GLY:H	17	0.1
(1,61)	1:13:A:SER:HA	1:17:A:LEU:H	14	0.1
(1,34)	1:5:A:ILE:HA	1:9:A:ASP:H	20	0.1

10 Dihedral-angle violation analysis ⓘ

No dihedral-angle restraints found