

# Full wwPDB NMR Structure Validation Report (i)

#### Jun 3, 2023 – 08:10 PM EDT

PDB ID : 2N0X BMRB ID : 25539

Title : Three dimensional structure of EPI-X4, a human albumin-derived peptide that

regulates innate immunity through the CXCR4/CXCL12 chemotactic axis and

antagonizes HIV-1 entry

Authors: Perez-Castells, J.; Canales, A.; Jimenez-Barbero, J.; Gimenez-Gallego, G.

Deposited on : 2015-03-18

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 (i) 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 (1)) were used in the production of this report:

MolProbity: 4.02b-467

Percentile statistics : 20191225.v01 (using entries in the PDB archive December 25th 2019)

wwPDB-RCI : v 1n 11 5 13 A (Berjanski et al., 2005)

PANAV : Wang et al. (2010)

 $\begin{array}{ccc} wwPDB\text{-}ShiftChecker &:& v1.2\\ BMRB \ Restraints \ Analysis &:& v1.2 \end{array}$ 

Ideal geometry (proteins) : Engh & Huber (2001) Ideal geometry (DNA, RNA) : Parkinson et al. (1996)

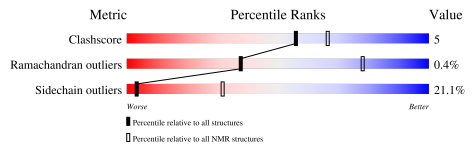
Validation Pipeline (wwPDB-VP) : 2.33

# 1 Overall quality at a glance (i)

The following experimental techniques were used to determine the structure:  $SOLUTION\ NMR$ 

The overall completeness of chemical shifts assignment is 53%.

Percentile scores (ranging between 0-100) for global validation metrics of the entry are shown in the following graphic. The table shows the number of entries on which the scores are based.



Metric	Whole archive $(\# \mathrm{Entries})$	$egin{array}{c} { m NMR \ archive} \ (\#{ m Entries}) \end{array}$	
Clashscore	158937	12864	
Ramachandran outliers	154571	11451	
Sidechain outliers	154315	11428	

The table below summarises the geometric issues observed across the polymeric chains and their fit to the experimental data. The red, orange, yellow and green segments indicate the fraction of residues that contain outliers for >=3, 2, 1 and 0 types of geometric quality criteria. A cyan segment indicates the fraction of residues that are not part of the well-defined cores, and a grey segment represents the fraction of residues that are not modelled. The numeric value for each fraction is indicated below the corresponding segment, with a dot representing fractions <=5%

Mol	Chain	Length	Quality of chain		
1	A	16	69%	19%	12%



# 2 Ensemble composition and analysis (i)

This entry contains 20 models. Model 4 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 mode					
1	A:1-A:14 (14)	0.90	4		

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 5 clusters and 1 single-model cluster was found.

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



# 3 Entry composition (i)

There is only 1 type of molecule in this entry. The entry contains 273 atoms, of which 145 are hydrogens and 0 are deuteriums.

• Molecule 1 is a protein called Serum albumin.

Mol	Chain	Residues	Atoms				Trace	
1	Λ	1.6	Total	С	Н	N	О	0
1	A	A   16	273	84	145	22	22	U



# 4 Residue-property plots (i)

#### 4.1 Average score per residue in the NMR ensemble

These plots are provided for all protein, RNA, DNA and oligosaccharide chains in the entry. The first graphic is the same as shown in the summary in section 1 of this report. The second graphic shows the sequence where residues are colour-coded according to the number of geometric quality criteria for which they contain at least one outlier: green = 0, yellow = 1, orange = 2 and red = 3 or more. Stretches of 2 or more consecutive residues without any outliers are shown as green connectors. Residues which are classified as ill-defined in the NMR ensemble, are shown in cyan with an underline colour-coded according to the previous scheme. Residues which were present in the experimental sample, but not modelled in the final structure are shown in grey.

• Molecule 1: Serum albumin



#### 4.2 Scores per residue for each member of the ensemble

Colouring as in section 4.1 above.

#### 4.2.1 Score per residue for model 1

• Molecule 1: Serum albumin



#### 4.2.2 Score per residue for model 2

• Molecule 1: Serum albumin





#### 4.2.3 Score per residue for model 3

• Molecule 1: Serum albumin

Chain A: 69% 19% 12%



### 4.2.4 Score per residue for model 4 (medoid)

• Molecule 1: Serum albumin

Chain A: 75% 12% 12%



#### 4.2.5 Score per residue for model 5

• Molecule 1: Serum albumin

Chain A: 62% 25% 12%



#### 4.2.6 Score per residue for model 6

• Molecule 1: Serum albumin

Chain A: 50% 31% 6% 12%



#### 4.2.7 Score per residue for model 7

• Molecule 1: Serum albumin

Chain A: 81% 6% 12%





#### 4.2.8 Score per residue for model 8

• Molecule 1: Serum albumin

Chain A: 62% 25% 12%



#### 4.2.9 Score per residue for model 9

• Molecule 1: Serum albumin

Chain A: 50% 25% 12% 12%



#### 4.2.10 Score per residue for model 10

• Molecule 1: Serum albumin

Chain A: 62% 25% 12%



#### 4.2.11 Score per residue for model 11

• Molecule 1: Serum albumin

Chain A: 56% 25% 6% 12%



### 4.2.12 Score per residue for model 12

• Molecule 1: Serum albumin

Chain A: 50% 31% 6% 12%





#### 4.2.13 Score per residue for model 13

• Molecule 1: Serum albumin





#### 4.2.14 Score per residue for model 14

• Molecule 1: Serum albumin

Chain A: 44% 38% 6% 12%



#### 4.2.15 Score per residue for model 15

• Molecule 1: Serum albumin

Chain A: 38% 44% 6% 12%



#### 4.2.16 Score per residue for model 16

• Molecule 1: Serum albumin

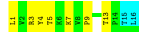
Chain A: 56% 31% 12%



### 4.2.17 Score per residue for model 17

• Molecule 1: Serum albumin

Chain A: 44% 44% 12%





### 4.2.18 Score per residue for model 18

• Molecule 1: Serum albumin



# 4.2.19 Score per residue for model 19

• Molecule 1: Serum albumin

Chain A: 62% 25% 12%



#### 4.2.20 Score per residue for model 20

• Molecule 1: Serum albumin

Chain A: 62% 19% 6% 12%





# 5 Refinement protocol and experimental data overview (i)



The models were refined using the following method: torsion angle dynamics.

Of the 100 calculated structures, 20 were deposited, based on the following criterion: structures with the least restraint violations.

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

Software name	Classification	Version
CYANA	structure solution	
CYANA	refinement	

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	126
Number of shifts mapped to atoms	126
Number of unparsed shifts	0
Number of shifts with mapping errors	0
Number of shifts with mapping warnings	0
Assignment completeness (well-defined parts)	53%



# 6 Model quality (i)

# 6.1 Standard geometry (i)

There are no covalent bond-length or bond-angle outliers.

There are no bond-length outliers.

There are no bond-angle outliers.

There are no chirality outliers.

There are no planarity outliers.

### 6.2 Too-close contacts (i)

In the following table, the Non-H and H(model) columns list the number of non-hydrogen atoms and hydrogen atoms in each chain respectively. The H(added) column lists the number of hydrogen atoms added and optimized by MolProbity. The Clashes column lists the number of clashes averaged over the ensemble.

Mol	Chain	Non-H	H(model)	H(added)	Clashes
1	A	113	127	129	1±1
All	All	2260	2540	2580	25

The all-atom clashscore is defined as the number of clashes found per 1000 atoms (including hydrogen atoms). The all-atom clashscore for this structure is 5.

All unique clashes are listed below, sorted by their clash magnitude.

Atom-1	Atom-2	Clash(Å)	$Distance(\mathring{A})$	Models	
Atom-1	Atom-2	Clash(A)	Distance(A)	Worst	Total
1:A:5:THR:HG23	1:A:13:THR:HG22	0.89	1.43	11	1
1:A:5:THR:OG1	1:A:13:THR:HG22	0.62	1.94	17	2
1:A:5:THR:HG22	1:A:14:PRO:HD3	0.58	1.75	15	3
1:A:1:LEU:HD21	1:A:4:TYR:CD2	0.56	2.35	17	1
1:A:1:LEU:HD21	1:A:9:PRO:CD	0.52	2.35	6	1
1:A:5:THR:HG23	1:A:7:LYS:O	0.50	2.06	20	4
1:A:1:LEU:HD12	1:A:1:LEU:O	0.50	2.07	15	1
1:A:1:LEU:HD22	1:A:9:PRO:HD3	0.50	1.82	9	2
1:A:1:LEU:HD13	1:A:9:PRO:HD3	0.49	1.84	12	2
1:A:1:LEU:HD13	1:A:9:PRO:HG3	0.46	1.87	16	2
1:A:12:SER:O	1:A:13:THR:HG23	0.45	2.11	15	4
1:A:1:LEU:HD11	1:A:9:PRO:HD3	0.44	1.89	15	1
1:A:1:LEU:HD21	1:A:9:PRO:HD3	0.42	1.91	6	1



# 6.3 Torsion angles (i)

#### 6.3.1 Protein backbone (i)

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	Tavoured Allowed		Percentiles		
1	A	13/16 (81%)	10±2 (75±14%)	3±2 (25±14%)	0±0 (0±2%)	38	78	
All	All	260/320 (81%)	194 (75%)	65 (25%)	1 (0%)	38	78	

All 1 unique Ramachandran outliers are listed below.

Mol	Chain	Res	Type	Models (Total)
1	A	3	ARG	1

#### 6.3.2 Protein sidechains (i)

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	14/16 (88%)	11±1 (79±9%)	3±1 (21±9%)	3	32
All	All	280/320~(88%)	221 (79%)	59 (21%)	3	32

All 7 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	13	THR	18
1	A	7	LYS	17
1	A	6	LYS	10
1	A	1	LEU	6
1	A	3	ARG	4
1	A	10	GLN	3
1	A	12	SER	1



#### 6.3.3 RNA (i)

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 monosaccharides 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 (i)

The completeness of assignment taking into account all chemical shift lists is 53% for the well-defined parts and 53% 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 (i)

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	126
Number of shifts mapped to atoms	126
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 (i)

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

# 7.1.3 Completeness of resonance assignments (i)

The following table shows the completeness of the chemical shift assignments for the well-defined regions of the structure. The overall completeness is 53%, i.e. 110 atoms were assigned a chemical shift out of a possible 208. 0 out of 4 assigned methyl groups (LEU and VAL) were assigned stereospecifically.

	Total	$^{1}\mathrm{H}$	$^{13}\mathbf{C}$	$^{15}\mathbf{N}$	
Backbone	25/66~(38%)	25/26~(96%)	0/28~(0%)	0/12 (0%)	
Sidechain	81/133~(61%)	81/87 (93%)	0/40 (0%)	0/6 (0%)	
Aromatic	4/9 (44%)	4/4 (100%)	0/5 (0%)	0/0 (%)	
Overall	110/208~(53%)	110/117 (94%)	0/73~(0%)	0/18 (0%)	

The following table shows the completeness of the chemical shift assignments for the full structure. The overall completeness is 53%, i.e. 126 atoms were assigned a chemical shift out of a possible 237. 0 out of 5 assigned methyl groups (LEU and VAL) were assigned stereospecifically.



	Total	$^{1}{ m H}$	$^{13}\mathbf{C}$	$^{15}{ m N}$	
Backbone	29/76 (38%)	$29/30 \ (97\%)$	0/32~(0%)	0/14 (0%)	
Sidechain	93/152 (61%)	93/100 (93%)	0/46 (0%)	0/6 (0%)	
Aromatic	4/9 (44%)	4/4 (100%)	$0/5 \ (0\%)$	0/0 (%)	
Overall	126/237 (53%)	126/134 (94%)	0/83 (0%)	0/20 (0%)	

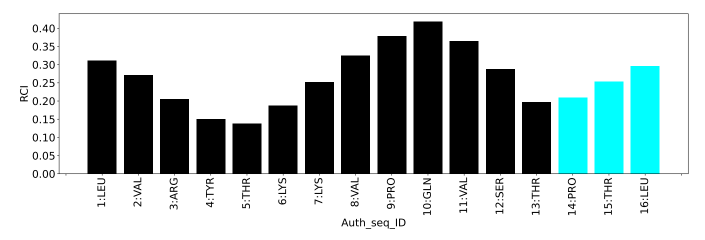
#### 7.1.4 Statistically unusual chemical shifts (i)

There are no statistically unusual chemical shifts.

#### 7.1.5 Random Coil Index (RCI) plots (i)

The image below reports random coil index values for the protein chains in the structure. The height of each bar gives a probability of a given residue to be disordered, as predicted from the available chemical shifts and the amino acid sequence. A value above 0.2 is an indication of significant predicted disorder. The colour of the bar shows whether the residue is in the well-defined core (black) or in the ill-defined residue ranges (cyan), as described in section 2 on ensemble composition. If well-defined core and ill-defined regions are not identified then it is shown as gray bars.

Random coil index (RCI) for chain A:





# 8 NMR restraints analysis (i)

# 8.1 Conformationally restricting restraints (i)

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	127
Intra-residue ( $ i-j =0$ )	56
Sequential ( i-j =1)	37
Medium range ( $ i-j >1$ and $ i-j <5$ )	10
Long range ( $ i-j  \ge 5$ )	24
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	7.9
Number of long range restraints per residue <sup>1</sup>	1.5

<sup>&</sup>lt;sup>1</sup>Long 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 (i)

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 (i)

Distance violations less than 0.1 Å are not included in the calculation. There are no distance violations

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

Dihedral-angle violations less than  $1^{\circ}$  are not included in the calculation. There are no dihedral-angle violations



# 9 Distance violation analysis (i)

# 9.1 Summary of distance violations (i)

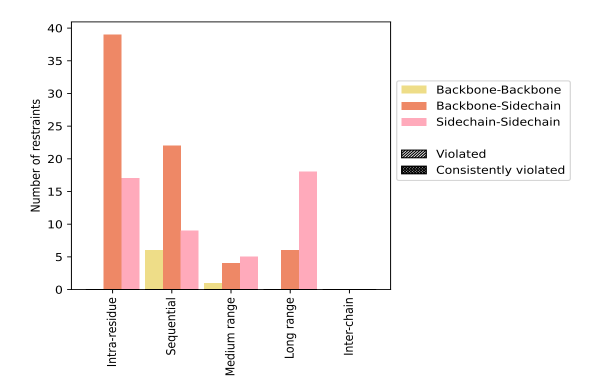
The following table shows the summary of distance violations in different restraint categories based on the sequence separation of the atoms involved. Each category is further sub-divided into three sub-categories based on the atoms involved. Violations less than 0.1 Å are not included in the statistics.

Dostusiuts tom s	Count	<b>%</b> <sup>1</sup>	${f Violated}^3$		Consistently		$\overline{ m Violated^4}$	
Restraints type	Count	70	Count	$\%^2$	$\%^1$	Count	$\%^2$	$\%^1$
Intra-residue ( i-j =0)	56	44.1	0	0.0	0.0	0	0.0	0.0
Backbone-Backbone	0	0.0	0	0.0	0.0	0	0.0	0.0
Backbone-Sidechain	39	30.7	0	0.0	0.0	0	0.0	0.0
Sidechain-Sidechain	17	13.4	0	0.0	0.0	0	0.0	0.0
Sequential ( i-j =1)	37	29.1	0	0.0	0.0	0	0.0	0.0
Backbone-Backbone	6	4.7	0	0.0	0.0	0	0.0	0.0
Backbone-Sidechain	22	17.3	0	0.0	0.0	0	0.0	0.0
Sidechain-Sidechain	9	7.1	0	0.0	0.0	0	0.0	0.0
Medium range ( $ i-j >1 \&  i-j <5$ )	10	7.9	0	0.0	0.0	0	0.0	0.0
Backbone-Backbone	1	0.8	0	0.0	0.0	0	0.0	0.0
Backbone-Sidechain	4	3.1	0	0.0	0.0	0	0.0	0.0
Sidechain-Sidechain	5	3.9	0	0.0	0.0	0	0.0	0.0
Long range ( $ i-j  \ge 5$ )	24	18.9	0	0.0	0.0	0	0.0	0.0
Backbone-Backbone	0	0.0	0	0.0	0.0	0	0.0	0.0
Backbone-Sidechain	6	4.7	0	0.0	0.0	0	0.0	0.0
Sidechain-Sidechain	18	14.2	0	0.0	0.0	0	0.0	0.0
Inter-chain	0	0.0	0	0.0	0.0	0	0.0	0.0
Backbone-Backbone	0	0.0	0	0.0	0.0	0	0.0	0.0
Backbone-Sidechain	0	0.0	0	0.0	0.0	0	0.0	0.0
Sidechain-Sidechain	0	0.0	0	0.0	0.0	0	0.0	0.0
Hydrogen bond	0	0.0	0	0.0	0.0	0	0.0	0.0
Disulfide bond	0	0.0	0	0.0	0.0	0	0.0	0.0
Total	127	100.0	0	0.0	0.0	0	0.0	0.0
Backbone-Backbone	7	5.5	0	0.0	0.0	0	0.0	0.0
Backbone-Sidechain	71	55.9	0	0.0	0.0	0	0.0	0.0
Sidechain-Sidechain	49	38.6	0	0.0	0.0	0	0.0	0.0

<sup>&</sup>lt;sup>1</sup> percentage calculated with respect to the total number of distance restraints, <sup>2</sup> percentage calculated with respect to the number of restraints in a particular restraint category, <sup>3</sup> violated in at least one model, <sup>4</sup> violated in all the models



#### 9.1.1 Bar chart: Distribution of distance restraints and violations (i)



Violated and consistently violated restraints are shown using different hatch patterns in their respective categories. The hydrogen bonds and disulfied bonds are counted in their appropriate category on the x-axis

# 9.2 Distance violation statistics for each model (i)

No violations found

# 9.3 Distance violation statistics for the ensemble (i)

No violations found

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

No violations found

# 9.5 All violated distance restraints (i)

No violations found



# 10 Dihedral-angle violation analysis (i)

No dihedral-angle restraints found

