

Full wwPDB NMR Structure Validation Report (i)

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PDB ID : 2FFK

Title: Solution structure of the complex between poxvirus-encoded CC chemokine

inhibitor vCCI and human MIP-1beta, minimized average structure

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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 (i)) 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)

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

PANAV : Wang et al. (2010)

ShiftChecker : 2.29

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

Validation Pipeline (wwPDB-VP) : 2.29

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 was not calculated.

There are no overall percentile quality scores available for this entry.

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	242	100%
2	В	69	100%



2 Ensemble composition and analysis (i)

This entry contains 1 models. Identification of well-defined residues and clustering analysis are not possible.



3 Entry composition (i)

There are 2 unique types of molecules in this entry. The entry contains 4601 atoms, of which 2247 are hydrogens and 0 are deuteriums.

• Molecule 1 is a protein called rabbitpox encoded CC chemokine inhibitor.

Mol	Chain	Residues			Atoms						
1	Λ	242	Total	С	Н	N	О	S	0		
1	A		3575	1104	1755	298	405	13			

• Molecule 2 is a protein called Small inducible cytokine A4.

Mol	Chain	Residues		_	Atom	ıs			Trace
9	D	60	Total	С	Н	N	О	S	0
2	Б	69	1026	337	492	83	109	5	U

There are 3 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference				
В	45	ALA	LYS	engineered mutation	UNP P13236				
В	46	ALA	ARG	engineered mutation	UNP P13236				
В	48	ALA	LYS	engineered mutation	UNP P13236				



4 Residue-property plots (i)

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: rabbitpox encoded CC chemokine inhibitor

Chain	A:																	10	00%	6																			
M1 P2 A3 S4	90 40	88 8	810	S11 S12	S13	C15	T16	E17	E19	N20	K21	H23	M24	G25 126	D27	V28	129	130 K31	V32	T33 K34	032	D36	T38	P39	T40	D42	K43	144 745	046	S47	V48 T49	E50	151	T52	254 S54	E55	S56	P58	D59 P60
K62 V62 E63 S64 F65	D66 D67	S68 T69	870	V71 E72	D73	V / 4 D75	P76	P77 T78	T79	Y80	Y81 S82	183	184	G85 G86	G87	L88	R89	M90 N91	F92	G93 F94	T95	K96	P98	660	1100	S102	1103	S104 E105	\$106	A107	D108	N110	T111	V112	N113 A114	R115	L116	\$118	V119 S120
P121 G122 Q123 G124 K125	D126 S127	P128	1130	T131 H132	E133	A135	L136	A137 M138	1139	K140	C142	E143	V144	S145 T146	D147	1148	R149	S151	E152	E153	K155	D156	D158	1159	K160 T161	H162	P163	V164	G166	\$167	N168 T169	8170	H171	K172	K173 V174	S175	Y176 E177	D178	1179 1180
G181 S182 T183 I184 V185	D186 T187	K188	V190	K191 N192	L193	F195	S196	V197 R198	1199	G200	D201	C203	K204	E205	S207	E208	L209	E210 V211	K212	D213 G214	F215	K216	V218	D219	G220	A222	S223	K224	A226	T227	D228	T230	S231	L232 T033	1233 D234	S235	T236 K237	L238	K239 A240
C241 V242																																							
• Mole	ecul	le :	2:	Sı	na	ıll	in	ıd	uc	cib	ole	c	yı	to	ki:	n∈	9 1	A 4	Į																				
Chain	В:																	10	00%	ó																_			
A 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	D6 P7	8 6 6	A10	C11 C12	F13	y 15	T16	A17	K19	L.20	P21	N23	F24	V25	D27	Y28	Y29	E30	S32	S33 1.34	C35	836	P38	A39	V40	F42	Q43	T44	A46	S47	A48	V50	C51	A52	D53	355	E56	W58	V59 Q60
61 62 63 64	99,	89																																					



Refinement protocol and experimental data overview (i) 5



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

Of the? calculated structures, 1 were deposited, based on the following criterion:?.

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

Software name	Classification	Version
DYNAMO	structure solution	3.1
DYNAMO	refinement	3.1

No chemical shift data was provided.



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	0	0	0	0
2	В	0	0	0	0
All	All	0	0	0	-

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 -.

There are no clashes.

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	Analysed Favoured Allowed			Percentiles
1	A	0	-	-	-	-
2	В	0	-	=	-	-
All	All	0	-	=	-	-

There are no Ramachandran outliers.



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	0	-	-	-
2	В	0	-	-	-
All	All	0	-	-	-

There are no protein residues with a non-rotameric sidechain to report.

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)

No chemical shift data were provided

