



wwPDB X-ray Structure Validation Summary Report i

Feb 28, 2014 – 07:02 AM GMT

PDB ID : 1HQF
Title : CRYSTAL STRUCTURE OF THE BINUCLEAR MANGANESE MET-ALLOENZYME ARGINASE COMPLEXED WITH N-HYDROXY-L-ARGININE
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Deposited on : 2000-12-16
Resolution : 2.90 Å(reported)

This is a wwPDB validation summary report for a publicly released PDB entry.
We welcome your comments at validation@mail.wwpdb.org
A user guide is available at <http://wwpdb.org/ValidationPDFNotes.html>

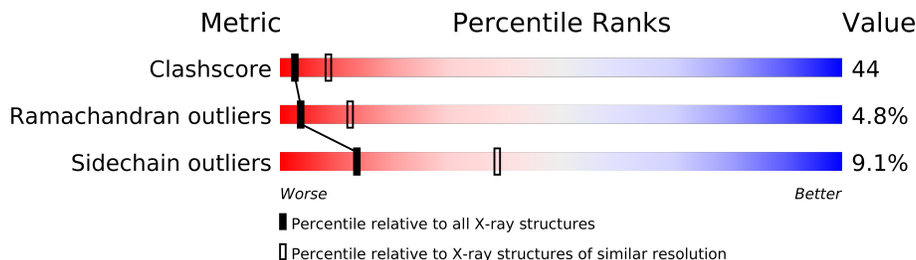
The following versions of software and data (see [references](#)) were used in the production of this report:

MolProbity : 4.02b-467
Mogul : 1.15 2013
Xtrriage (Phenix) : **NOT EXECUTED**
EDS : **NOT EXECUTED**
Percentile statistics : 21963
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et. al. (1996)
Validation Pipeline (wwPDB-VP) : stable22683

1 Overall quality at a glance

The reported resolution of this entry is 2.90 Å.

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 (#Entries)	Similar resolution (#Entries, resolution range(Å))
Clashscore	79885	1326 (2.90-2.90)
Ramachandran outliers	78287	1290 (2.90-2.90)
Sidechain outliers	78261	1292 (2.90-2.90)

The table below summarises the geometric issues observed across the polymeric chains and their fit to the electron density. The red, orange, yellow and green segments on the lower bar indicate the fraction of residues that contain outliers for ≥ 3 , 2, 1 and 0 types of geometric quality criteria. The upper red bar (where present) indicates the fraction of residues that have poor fit to the electron density.

Note EDS was not executed.

Mol	Chain	Length	Quality of chain
1	A	323	
1	B	323	
1	C	323	

2 Entry composition

There are 4 unique types of molecules in this entry. The entry contains 7245 atoms, of which 0 are hydrogen and 0 are deuterium.

In the tables below, the ZeroOcc column contains the number of atoms modelled with zero occupancy, the AltConf column contains the number of residues with at least one atom in alternate conformation and the Trace column contains the number of residues modelled with at most 2 atoms.

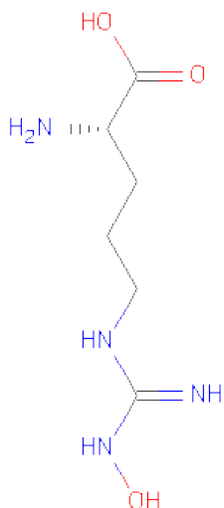
- Molecule 1 is a protein called ARGINASE 1.

Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
			Total	C	N	O	S			
1	A	314	2395	1528	405	455	7	0	0	0
1	B	314	2395	1528	405	455	7	0	0	0
1	C	314	2395	1528	405	455	7	0	0	0

- Molecule 2 is MANGANESE (II) ION (three-letter code: MN) (formula: Mn).

Mol	Chain	Residues	Atoms		ZeroOcc	AltConf
2	B	2	Total	Mn	0	0
			2	2		
2	A	2	Total	Mn	0	0
			2	2		
2	C	2	Total	Mn	0	0
			2	2		

- Molecule 3 is N-OMEGA-HYDROXY-L-ARGININE (three-letter code: HAR) (formula: C₆H₁₄N₄O₃).



Mol	Chain	Residues	Atoms				ZeroOcc	AltConf
			Total	C	N	O		
3	A	1	13	6	4	3	0	0
3	B	1	13	6	4	3	0	0
3	C	1	13	6	4	3	0	0

- Molecule 4 is water.

Mol	Chain	Residues	Atoms		ZeroOcc	AltConf
			Total	O		
4	A	5	5	5	0	0
4	B	5	5	5	0	0
4	C	5	5	5	0	0

Chain C: 

RET	SER	SER	LYS	PRO	KG	R7	I8	E9	I10	I11	G12	A13	P14	F15	S16	K17	G18	G19	P20	R21	G22	G23	V24	V37	E38	K39	L40	K41	E42	Y45	N46	V47	R48	R49	H50	S51	D52	V56	D57	V58	D61	S62	P63	F64	O65	I66	V67	P70	R71	N77	E78	Q79	L80		
A81	V84	A85	E86	T87	N90	G91	T92	I93	S94	V95	G98	G99	S100	H101	S102	M103	A104	I105	G106	S107	I108	S109	G110	H111	L118	I121	W122	V123	D124	A125	H126	T127	D128	I129	N130	T131	P132	L133	T134	T135	S136	S137	G138	N139	L140	H141	G142	Q143	P144	V145	A146	F147	L148	L149	
K150	E151	L152	K155	F156	P157	D158	V159	P160	G161	F162	S163	W164	P167	C168	I169	S170	A171	I174	V175	Y176	I177	G178	L179	R180	D181	V182	D183	P184	G185	E186	H187	Y188	I189	I190	K191	T192	L193	G194	I195	K196	Y197	F198	S199	M200	T201	E202	T209	G209	K210	V211	M212	E213	E214	T215	F216
S217	Y218	L219	L220	K224	R225	P226	I227	H228	L229	S230	F231	D232	V233	D234	G235	L236	V239	F240	A243	T244	G245	T246	P247	V248	V249	G250	L251	L252	S253	E256	G257	L258	Y259	L260	T261	E262	E263	I264	Y265	K266	T267	G268	L269	L270	S271	G272	L273	D274	I275	M276	E277	V278	N279	P280	T281
L282	G283	K284	E287	E288	V289	T290	R291	T292	V293	W294	T295	K296	V297	A298	L299	T300	L301	S302	C303	F304	G305	T306	K307	R308	E309	G310	N311	H312	B317	Y318	L319	LYS	PRO	PRO	LYS																				

4 Data and refinement statistics (i)

Xtrriage (Phenix) and EDS were not executed - this section will therefore be incomplete.

Property	Value	Source
Space group	P 32	Depositor
Cell constants a, b, c, α , β , γ	88.00Å 88.00Å 112.00Å 90.00° 90.00° 120.00°	Depositor
Resolution (Å)	30.00 – 2.90	Depositor
% Data completeness (in resolution range)	(Not available) (30.00-2.90)	Depositor
R_{merge}	0.06	Depositor
R_{sym}	(Not available)	Depositor
Refinement program	CNS	Depositor
R, R_{free}	0.265 , 0.289	Depositor
Estimated twinning fraction	No twinning to report.	Xtrriage
Total number of atoms	7245	wwPDB-VP
Average B, all atoms (Å ²)	85.0	wwPDB-VP

5 Model quality i

5.1 Standard geometry i

Bond lengths and bond angles in the following residue types are not validated in this section: HAR, MN

The Z score for a bond length (or angle) is the number of standard deviations the observed value is removed from the expected value. A bond length (or angle) with $|Z| > 5$ is considered an outlier worth inspection. RMSZ is the root-mean-square of all Z scores of the bond lengths (or angles).

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	# Z >5	RMSZ	# Z >5
1	A	0.42	0/2448	0.71	1/3325 (0.0%)
1	B	0.42	0/2448	0.71	1/3325 (0.0%)
1	C	0.42	0/2448	0.71	1/3325 (0.0%)
All	All	0.42	0/7344	0.71	3/9975 (0.0%)

Chiral center outliers are detected by calculating the chiral volume of a chiral center and verifying if the center is modelled as a planar moiety or with the opposite hand. A planarity outlier is detected by checking planarity of atoms in a peptide group, atoms in a mainchain group or atoms of a sidechain that are expected to be planar.

Mol	Chain	#Chirality outliers	#Planarity outliers
1	A	0	1
1	B	0	1
1	C	0	1
All	All	0	3

There are no bond length outliers.

All (3) bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	A	99	GLY	N-CA-C	-6.78	96.16	113.10
1	C	99	GLY	N-CA-C	-6.77	96.18	113.10
1	B	99	GLY	N-CA-C	-6.76	96.21	113.10

There are no chirality outliers.

All (3) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	A	176	TYR	Sidechain

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Mol	Chain	Res	Type	Group
1	B	176	TYR	Sidechain
1	C	176	TYR	Sidechain

5.2 Close contacts

In the following table, the Non-H and H(model) columns list the number of non-hydrogen atoms and hydrogen atoms in the chain respectively. The H(added) column lists the number of hydrogens added by MolProbity. The Clashes column lists the number of clashes within the asymmetric unit, and the number in parentheses is this value normalized per 1000 atoms of the molecule in the chain. The Symm-Clashes column gives symmetry related clashes, in the same way as for the Clashes column.

Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
1	A	2395	0	2420	217	0
1	B	2395	0	2420	219	0
1	C	2395	0	2420	224	0
2	A	2	0	0	0	0
2	B	2	0	0	0	0
2	C	2	0	0	0	0
3	A	13	0	11	2	0
3	B	13	0	11	2	0
3	C	13	0	11	2	0
4	A	5	0	0	1	0
4	B	5	0	0	1	0
4	C	5	0	0	1	0
All	All	7245	0	7293	645	0

Clashscore is defined as the number of clashes calculated for the entry per 1000 atoms (including hydrogens) of the entry. The overall clashscore for this entry is 44.

The worst 5 of 645 close contacts within the same asymmetric unit are listed below.

Atom-1	Atom-2	Distance(Å)	Clash(Å)
1:B:6:LYS:HD3	1:B:7:PRO:HD2	1.23	1.15
1:A:6:LYS:HD3	1:A:7:PRO:HD2	1.23	1.14
1:C:6:LYS:HD3	1:C:7:PRO:HD2	1.23	1.08
1:B:82:ALA:O	1:B:86:GLU:HG2	1.53	1.07
1:A:37:VAL:HG23	4:A:602:HOH:O	1.61	1.00

There are no symmetry-related clashes.

5.3 Torsion angles

5.3.1 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 X-ray entries followed by that with respect to entries of similar resolution. 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	312/323 (97%)	240 (77%)	57 (18%)	15 (5%)	4	12
1	B	312/323 (97%)	240 (77%)	57 (18%)	15 (5%)	4	12
1	C	312/323 (97%)	240 (77%)	57 (18%)	15 (5%)	4	12
All	All	936/969 (97%)	720 (77%)	171 (18%)	45 (5%)	4	12

5 of 45 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	A	64	PHE
1	A	233	VAL
1	B	64	PHE
1	B	233	VAL
1	C	64	PHE

5.3.2 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 X-ray entries followed by that with respect to entries of similar resolution. 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	264/273 (97%)	240 (91%)	24 (9%)	14	38
1	B	264/273 (97%)	240 (91%)	24 (9%)	14	38
1	C	264/273 (97%)	240 (91%)	24 (9%)	14	38
All	All	792/819 (97%)	720 (91%)	72 (9%)	14	38

5 of 72 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	B	141	HIS

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Mol	Chain	Res	Type
1	B	231	PHE
1	C	239	VAL
1	B	152	LEU
1	B	197	TYR

Some sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 16 such sidechains are listed below:

Mol	Chain	Res	Type
1	B	90	ASN
1	B	294	ASN
1	C	79	GLN
1	B	79	GLN
1	C	90	ASN

5.3.3 RNA ⓘ

There are no RNA chains in this entry.

5.4 Non-standard residues in protein, DNA, RNA chains ⓘ

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

5.5 Carbohydrates ⓘ

There are no carbohydrates in this entry.

5.6 Ligand geometry ⓘ

Of 9 ligands modelled in this entry, 6 are monoatomic - leaving 3 for Mogul analysis.

In the following table, the Counts columns list the number of bonds (or angles) for which Mogul statistics could be retrieved, the number of bonds (or angles) that are observed in the model and the number of bonds (or angles) that are defined in the chemical component dictionary. The Link column lists molecule types, if any, to which the group is linked. The Z score for a bond length (or angle) is the number of standard deviations the observed value is removed from the expected value. A bond length (or angle) with $|Z| > 2$ is considered an outlier worth inspection. RMSZ is the root-mean-square of all Z scores of the bond lengths (or angles).

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	# Z > 2	Counts	RMSZ	# Z > 2
3	HAR	A	906	2	12,12,12	1.44	3 (25%)	14,14,14	0.77	0
3	HAR	B	907	2	12,12,12	1.44	3 (25%)	14,14,14	0.76	0
3	HAR	C	908	2	12,12,12	1.44	3 (25%)	14,14,14	0.77	0

In the following table, the Chirals column lists the number of chiral outliers, the number of chiral centers analysed, the number of these observed in the model and the number defined in the chemical component dictionary. Similar counts are reported in the Torsion and Rings columns. '-' means no outliers of that kind were identified.

Mol	Type	Chain	Res	Link	Chirals	Torsions	Rings
3	HAR	A	906	2	-	0/12/13/13	0/0/0/0
3	HAR	B	907	2	-	0/12/13/13	0/0/0/0
3	HAR	C	908	2	-	0/12/13/13	0/0/0/0

The worst 5 of 9 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
3	B	907	HAR	O-C	2.91	1.32	1.22
3	A	906	HAR	O-C	2.91	1.32	1.22
3	C	908	HAR	O-C	2.90	1.32	1.22
3	A	906	HAR	CA-C	2.78	1.63	1.53
3	B	907	HAR	CA-C	2.78	1.63	1.53

There are no bond angle outliers.

There are no chirality outliers.

There are no torsion outliers.

There are no ring outliers.

5.7 Other polymers

There are no such residues in this entry.

5.8 Polymer linkage issues

There are no chain breaks in this entry.

6 Fit of model and data

6.1 Protein, DNA and RNA chains

EDS was not executed - this section will therefore be empty.

6.2 Non-standard residues in protein, DNA, RNA chains

EDS was not executed - this section will therefore be empty.

6.3 Carbohydrates

EDS was not executed - this section will therefore be empty.

6.4 Ligands

EDS was not executed - this section will therefore be empty.

6.5 Other polymers

EDS was not executed - this section will therefore be empty.