



Full wwPDB Integrative Structure Validation Report

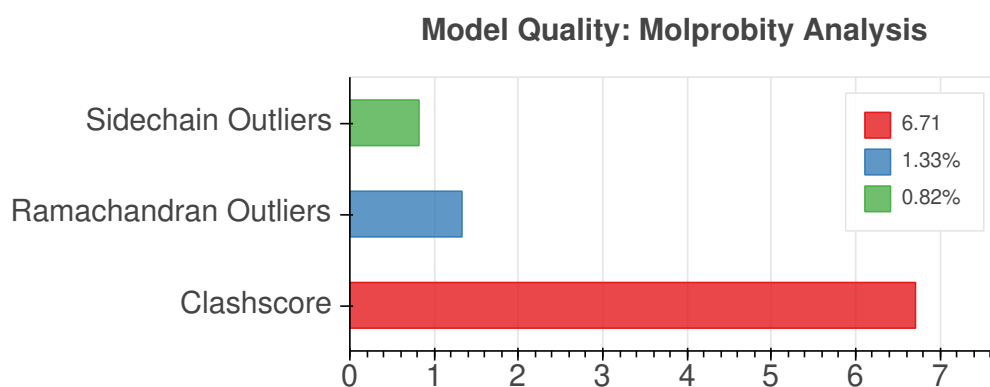
November 05, 2019 -- 04:07 PM

<i>PDB ID</i>	<i>PDBDEV00000005</i>
Molecule Name	Serum Albumin Domain A Structure
Title	Serum Albumin Domain Structures in Human Blood Serum by Mass Spectrometry and Computational Biology
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The following software were used in the production of this report:

Molprobability : Version 4.4
Integrative Modeling Validation Package : Version 1.0

1. Overall quality at a glance



2. Ensemble information

This entry consists of 0 distinct ensemble.

3. Model composition

3.1 Summary

This entry consists of 5 unique models, with 1 subunits in each model. A total of 2 datasets or restraints was used to build this entry. Each model is represented by 1 rigid bodies and 0 flexible or non-rigid units.

3.2 Entry composition

There are 5 unique types of models in this entry. These models are titled Best scoring model (domain A), 2nd best scoring model (domain A), 3rd best scoring model (domain A), 4th best scoring model (domain A), 5th best scoring model (domain A) respectively.

<i>Model ID</i>	<i>Subunit number</i>	<i>Subunit ID</i>	<i>Subunit name</i>	<i>Chain ID</i>	<i>Total residues</i>
1	1	1	HSA_A	A	197
2	1	1	HSA_A	A	197
3	1	1	HSA_A	A	197
4	1	1	HSA_A	A	197
5	1	1	HSA_A	A	197

3.3 Datasets used for modeling

There are 2 unique datasets used to build the models in this entry.

<i>ID</i>	<i>Dataset type</i>	<i>Database name</i>	<i>Data access code</i>
1	CX-MS data	PRIDE	PXD001692
2	unspecified	Not listed	None

4. Representation

This entry has only one representation and includes 1 rigid bodies and 0 flexible units.

<i>Chain ID</i>	<i>Rigid bodies</i>	<i>Non-rigid segments</i>
A	1-197:None.	-

5. Methodology and software

<i>Step number</i>	<i>Protocol ID</i>	<i>Method name</i>	<i>Method type</i>	<i>Number of computed models</i>	<i>Multi state modeling</i>	<i>Multi scale modeling</i>
1	1	Model-based search (MBS) in Rosetta	Conformational search	5000	False	True

There are 2 software packages reported in this entry.

<i>ID</i>	<i>Software name</i>	<i>Software version</i>	<i>Software classification</i>
1	Rosetta MBS	None	Model Building
2	EPC-map	None	Contact Predictor

6. Data quality

7. Model quality

7.1 Standard geometry

There are 7715 bond outliers in this entry.

<i>Bond type</i>	<i>Observed distance (Å)</i>	<i>Ideal distance (Å)</i>	<i>Number of outliers</i>
CD2--HD2	1.089	0.93	129
CD1--HD1	1.089	0.93	99
CE2--HE2	1.088	0.93	99
CE1--HE1	1.089	0.93	129
CZ--HZ	1.088	0.93	64
NE--HE	1.01	0.86	49
NH2--HH21	1.01	0.86	49
N--H	1.008	0.86	939
NH1--HH11	1.009	0.86	49
NE2--HE2	1.009	0.86	29

NH1--HH12	1.008	0.86	49
NH2--HH22	1.008	0.86	49
NE2--HE21	1	0.86	34
NE2--HE22	0.999	0.86	34
ND2--HD21	1	0.86	34
ND2--HD22	0.998	0.86	34
CD--HD2	1.089	0.97	174
CG--HG3	1.088	0.97	324
CD--HD3	1.088	0.97	174
CB--HB3	1.087	0.97	859
CA--HA	1.088	0.97	964
CG--HG2	1.089	0.97	324
CB--HB2	1.087	0.97	859
SG--HG	1.328	1.2	59
CE--HE2	1.087	0.97	94
CD1--HD12	1.088	0.97	114
CG--HG	1.089	0.97	104
NZ--HZ2	1.009	0.89	84
CG2--HG22	1.089	0.97	104
OG1--HG1	0.958	0.84	44
OH--HH	0.959	0.84	34
CG1--HG11	1.089	0.97	49
CD2--HD21	1.089	0.97	104
CG2--HG23	1.088	0.97	104
NZ--HZ1	1.009	0.89	84
CD1--HD11	1.089	0.97	114
CG1--HG13	1.088	0.97	59

CB--HB1	1.089	0.97	114
CE--HE3	1.089	0.97	94
CB--HB	1.089	0.97	104
CD2--HD22	1.089	0.97	104
NZ--HZ3	1.008	0.89	84
OG--HG	0.959	0.84	24
CD2--HD23	1.089	0.97	104
CG1--HG12	1.088	0.97	59
CG2--HG21	1.088	0.97	104
CA--HA2	1.09	0.97	19
CE--HE1	1.09	0.97	9
CD1--HD13	1.088	0.97	114
CA--HA3	1.088	0.97	19
N--H3	1	0.89	4
N--H1	1	0.89	4
N--H2	1	0.89	4

There are 170 angle outliers in this entry.

Angle type	Observed angle (°)	Ideal angle (°)	Number of outliers
CA-CB-HB3	93.422	109	34
CB-CG-HG3	94.661	109	34
CB-CG-HG2	122.705	109	34
H2-N-H3	97.182	109.47	4
H1-N-H3	97.194	109.47	4
H1-N-H2	97.185	109.47	4
CZ-NE-HE	105.888	117.9	49

7.2 Too-close contacts

The following all-atom clashscore is based on a MolProbity analysis. All-atom clashscore is defined as the number of clashes found per 1000 atoms (including hydrogen atoms). The table below contains clashscores for all the models in this entry.

Model ID	Clash score	Number of clashes
Model 1	6.71	21
Model 2	3.84	12
Model 3	7.99	25
Model 4	6.39	20
Model 5	5.43	17

All 95 close contacts within the same asymmetric unit are listed below, sorted by their clash magnitude.

7.3 Torsion angles

7.3.1 Protein backbone

In the following table, Ramachandran outliers are listed. The Analysed column shows the number of residues for which the backbone conformation was analysed.

Model ID	Analyzed	Favored	Allowed	Outliers
1	195	183	9	3
2	195	179	13	3
3	195	187	7	1
4	195	180	12	3
5	195	178	14	3

Detailed list of outliers are tabulated below.

Model ID	Chain and res ID	Residue type
1	A:51	ALA
1	A:165	CYS
1	A:170	LYS
2	A:82	GLU
2	A:92	PRO

2	A:115	GLU
3	A:51	ALA
4	A:51	ALA
4	A:165	CYS
4	A:170	LYS
5	A:82	GLU
5	A:92	PRO
5	A:143	PRO

7.3.2 Protein sidechains

In the following table, sidechain outliers are listed. The Analysed column shows the number of residues for which the sidechain conformation was analysed.

Model ID	Analysed	Favored	Allowed	Outliers
1	170	165	1	4
2	170	168	1	1
3	170	166	4	0
4	170	166	3	1
5	170	168	1	1

Detailed list of outliers are tabulated below.

Model ID	Chain and res ID	Residue type
1	A:50	VAL
1	A:70	LEU
1	A:89	LYS
1	A:195	LYS
2	A:178	LEU
4	A:50	VAL
5	A:144	TYR

8. Fit of model to data used for modeling

9. Fit of model to data not used for modeling

10. Uncertainty of model
