



wwPDB X-ray Structure Validation Summary Report ⓘ

May 29, 2024 – 04:27 PM EDT

PDB ID : 1AHU
Title : STRUCTURE OF THE OCTAMERIC FLAVOENZYME VANILLYL-ALCOHOL OXIDASE IN COMPLEX WITH P-CRESOL
Authors : Mattevi, A.
Deposited on : 1997-04-10
Resolution : 2.70 Å(reported)

This is a wwPDB X-ray Structure Validation Summary 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/XrayValidationReportHelp>

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 : 4.02b-467
Mogul : 1.8.5 (274361), CSD as541be (2020)
Xtrriage (Phenix) : **NOT EXECUTED**
EDS : **NOT EXECUTED**
buster-report : 1.1.7 (2018)
Percentile statistics : 20191225.v01 (using entries in the PDB archive December 25th 2019)
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP) : 2.36.2

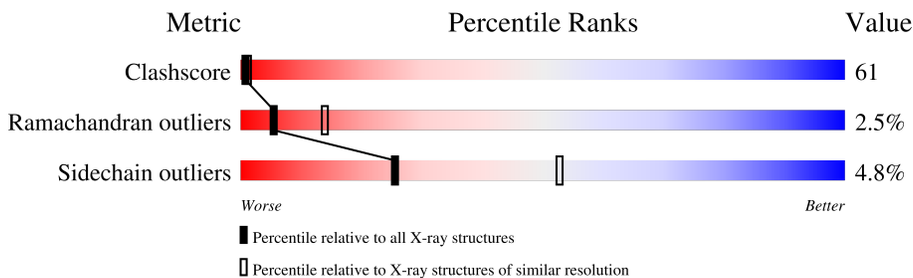
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

X-RAY DIFFRACTION

The reported resolution of this entry is 2.70 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	141614	3122 (2.70-2.70)
Ramachandran outliers	138981	3069 (2.70-2.70)
Sidechain outliers	138945	3069 (2.70-2.70)

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 of the lower bar indicate the fraction of residues that contain outliers for ≥ 3 , 2, 1 and 0 types of geometric quality criteria respectively. 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 $\leq 5\%$.

Note EDS was not executed.

Mol	Chain	Length	Quality of chain
1	A	560	 26% 56% 16% ..
1	B	560	 25% 57% 16% ..

2 Entry composition [i](#)

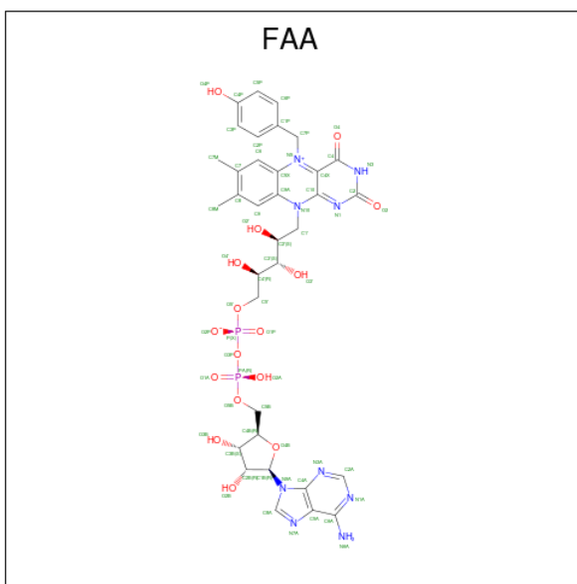
There are 3 unique types of molecules in this entry. The entry contains 9011 atoms, of which 0 are hydrogens and 0 are deuteriums.

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 VANILLYL-ALCOHOL OXIDASE.

Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
			Total	C	N	O	S			
1	A	555	4391	2817	751	799	24	37	0	0
1	B	555	4391	2817	751	799	24	37	0	0

- Molecule 2 is N5-(4-HYDROXYBENZYL)FLAVIN-ADENINE DINUCLEOTIDE (three-letter code: FAA) (formula: C₃₄H₃₉N₉O₁₆P₂).



Mol	Chain	Residues	Atoms					ZeroOcc	AltConf
			Total	C	N	O	P		
2	A	1	61	34	9	16	2	0	0
2	B	1	61	34	9	16	2	0	0

- Molecule 3 is water.

Mol	Chain	Residues	Atoms		ZeroOcc	AltConf
3	A	55	Total 55	O 55	0	0
3	B	52	Total 52	O 52	1	0

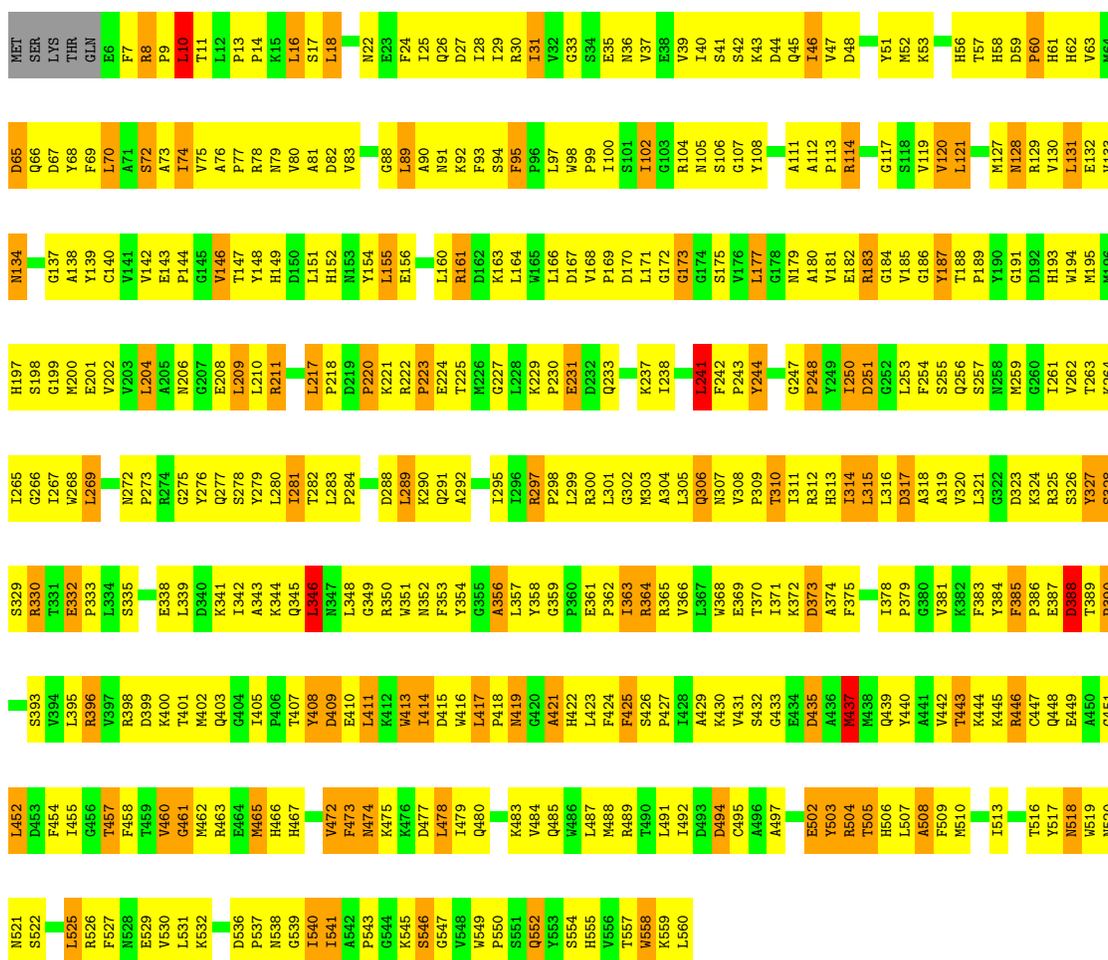
3 Residue-property plots

These plots are drawn for all protein, RNA, DNA and oligosaccharide chains in the entry. The first graphic for a chain summarises the proportions of the various outlier classes displayed in the second graphic. The second graphic shows the sequence view annotated by issues in geometry. Residues are color-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 outlier are shown as a green connector. Residues present in the sample, but not in the model, are shown in grey.

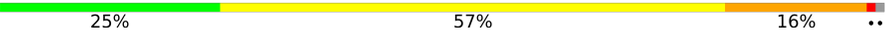
Note EDS was not executed.

- Molecule 1: VANILLYL-ALCOHOL OXIDASE

Chain A:  26% 56% 16% ..



- Molecule 1: VANILLYL-ALCOHOL OXIDASE

Chain B:  25% 57% 16% ..



E449	T516	S326	T263	H197	M134	K64
A450	A450	Y327	K264	S196	G137	D65
G451	G451	S328	I265	G199	A138	Q66
L452	L452	S329	G266	M200	Y139	D67
D453	D453	R330	I267	E201	C140	Y68
F454	F454	T331	W268	V202	V141	F69
L455	L455	E332	L269	V203	V142	L70
G456	G456	P333	M270	L204	A171	A71
T457	T457	L394	P271	A205	E143	S72
F458	F458	S335	N272	N206	A73	A73
T459	T459	S335	P273	G207	G145	I74
V460	V460	E338	R274	E208	V146	V75
G461	G461	B340	G275	L209	T147	A76
M462	M462	K341	Y276	L210	Y148	P77
R463	R463	I342	Q277	R211	H149	R78
E464	E464	K343	S278	M214	D150	N79
M465	M465	A344	Y279	L217	L151	V80
H466	H466	K344	L280	L218	H152	A81
H467	H467	Q345	I281	P219	W153	D82
V472	V472	L346	T282	D219	Y154	V83
F473	F473	N347	L283	P220	L155	L89
N474	N474	L348	P284	K221	E156	A90
I540	I540	G349	D288	R222	L160	N91
K475	K475	R350	L289	R223	R161	K92
K476	K476	W351	K290	E224	D162	F83
D477	D477	N352	K291	M226	K163	S94
L478	L478	F353	Q291	G227	L164	F95
L479	L479	Y354	A292	L228	W165	P96
K480	K480	G355	I295	L228	L166	L97
K483	K483	A356	T296	K229	D167	W98
V484	V484	L357	R297	P230	V168	P99
Q485	Q485	Y358	P298	E231	P169	I100
W486	W486	G359	L299	D232	D170	S101
L487	L487	P360	R300	Q233	L171	I102
M488	M488	E361	L301	K237	G172	G103
M489	M489	F424	G302	I238	G173	R104
R489	R489	F425	M303	L241	G174	M105
T490	T490	S426	A304	F242	S175	S106
L491	L491	P427	L305	P243	V176	G107
L492	L492	T428	L306	Y244	L177	Y108
D493	D493	A429	Q306	G247	G178	A111
D494	D494	K430	N307	P248	M179	A112
C495	C495	E369	V308	Y249	V181	P113
A496	A496	T370	P309	R248	E182	R114
A497	A497	I371	I310	Y249	R183	G117
W500	W500	K372	I311	T250	G184	S118
G501	G501	D373	R312	D251	V185	V119
E502	E502	A436	H313	G252	G186	V120
Y503	Y503	F375	I314	L253	Y187	L121
R504	R504	L376	L315	T188	T188	L121
T505	T505	P379	L316	P254	P189	M127
H506	H506	Y440	D317	S255	P189	M128
L507	L507	A441	A318	Q256	P189	M129
A508	A508	V442	A319	Q256	Y190	M129
F509	F509	T443	V320	S257	G191	V130
M510	M510	K382	L321	M259	D192	M129
D511	D511	F383	G322	G260	H193	V130
L513	L513	Y384	K323	I261	W194	E132
		F385	K324	V262	M195	E132
		C447	R325		M196	V133
		E387				

4 Data and refinement statistics

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

Property	Value	Source
Space group	I 4	Depositor
Cell constants a, b, c, α , β , γ	128.82Å 128.82Å 130.79Å 90.00° 90.00° 90.00°	Depositor
Resolution (Å)	30.00 – 2.70	Depositor
% Data completeness (in resolution range)	95.4 (30.00-2.70)	Depositor
R_{merge}	(Not available)	Depositor
R_{sym}	0.97	Depositor
Refinement program	TNT 5E	Depositor
R, R_{free}	0.221 , 0.290	Depositor
Estimated twinning fraction	No twinning to report.	Xtrriage
Total number of atoms	9011	wwPDB-VP
Average B, all atoms (Å ²)	27.0	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: FAA

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	0.73	0/4511	1.74	110/6131 (1.8%)
1	B	0.73	0/4511	1.74	108/6131 (1.8%)
All	All	0.73	0/9022	1.74	218/12262 (1.8%)

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	2	0
1	B	2	0
All	All	4	0

There are no bond length outliers.

The worst 5 of 218 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	A	330	ARG	NE-CZ-NH1	-11.34	114.63	120.30
1	B	330	ARG	NE-CZ-NH1	-11.24	114.68	120.30
1	A	536	ASP	C-N-CD	-9.85	98.92	120.60
1	B	536	ASP	C-N-CD	-9.85	98.93	120.60
1	B	417	LEU	CA-CB-CG	-9.34	93.83	115.30

All (4) chirality outliers are listed below:

Mol	Chain	Res	Type	Atom
1	A	187	TYR	CA
1	A	332	GLU	CA
1	B	187	TYR	CA

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Mol	Chain	Res	Type	Atom
1	B	332	GLU	CA

There are no planarity outliers.

5.2 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 the 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 within the asymmetric unit, whereas Symm-Clashes lists symmetry-related clashes.

Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
1	A	4391	0	4330	531	0
1	B	4391	0	4330	548	0
2	A	61	0	35	13	0
2	B	61	0	35	13	0
3	A	55	0	0	9	0
3	B	52	0	0	9	0
All	All	9011	0	8730	1058	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 61.

The worst 5 of 1058 close contacts within the same asymmetric unit are listed below, sorted by their clash magnitude.

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
2:A:600:FAA:H51A	2:A:600:FAA:H8A	1.21	1.13
1:A:309:PRO:HG2	1:A:460:VAL:HB	1.32	1.11
2:B:600:FAA:H51A	2:B:600:FAA:H8A	1.21	1.10
1:A:555:HIS:HB3	1:A:559:LYS:HE3	1.32	1.07
1:A:507:LEU:HA	1:A:510:MET:HE3	1.37	1.06

There are no symmetry-related clashes.

5.3 Torsion angles [i](#)

5.3.1 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 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	553/560 (99%)	470 (85%)	69 (12%)	14 (2%)	5	14
1	B	553/560 (99%)	470 (85%)	69 (12%)	14 (2%)	5	14
All	All	1106/1120 (99%)	940 (85%)	138 (12%)	28 (2%)	5	14

5 of 28 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	A	44	ASP
1	B	44	ASP
1	A	30	ARG
1	A	46	ILE
1	A	328	SER

5.3.2 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 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	475/482 (98%)	452 (95%)	23 (5%)	25	53
1	B	475/482 (98%)	452 (95%)	23 (5%)	25	53
All	All	950/964 (98%)	904 (95%)	46 (5%)	25	53

5 of 46 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	B	128	ASN

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Mol	Chain	Res	Type
1	B	346	LEU
1	B	134	ASN
1	B	241	LEU
1	B	373	ASP

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 24 such sidechains are listed below:

Mol	Chain	Res	Type
1	B	91	ASN
1	B	240	HIS
1	B	197	HIS
1	B	277	GLN
1	A	277	GLN

5.3.3 RNA [i](#)

There are no RNA molecules in this entry.

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

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

5.5 Carbohydrates [i](#)

There are no monosaccharides in this entry.

5.6 Ligand geometry [i](#)

2 ligands are modelled in this entry.

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
2	FAA	A	600	1	60,67,67	0.78	2 (3%)	74,102,102	1.69	7 (9%)
2	FAA	B	600	1	60,67,67	0.77	2 (3%)	74,102,102	1.69	7 (9%)

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
2	FAA	A	600	1	-	11/34/54/54	0/7/7/7
2	FAA	B	600	1	-	11/34/54/54	0/7/7/7

All (4) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
2	A	600	FAA	C5X-N5	-2.41	1.36	1.40
2	B	600	FAA	C5X-N5	-2.36	1.36	1.40
2	B	600	FAA	C9A-N10	-2.06	1.37	1.41
2	A	600	FAA	C9A-N10	-2.00	1.37	1.41

The worst 5 of 14 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
2	A	600	FAA	C7P-N5-C5X	-11.26	102.71	119.14
2	B	600	FAA	C7P-N5-C5X	-11.26	102.72	119.14
2	B	600	FAA	C1P-C7P-N5	-4.14	106.81	113.02
2	A	600	FAA	C1P-C7P-N5	-4.11	106.85	113.02
2	B	600	FAA	C6-C5X-N5	-2.94	116.92	120.47

There are no chirality outliers.

5 of 22 torsion outliers are listed below:

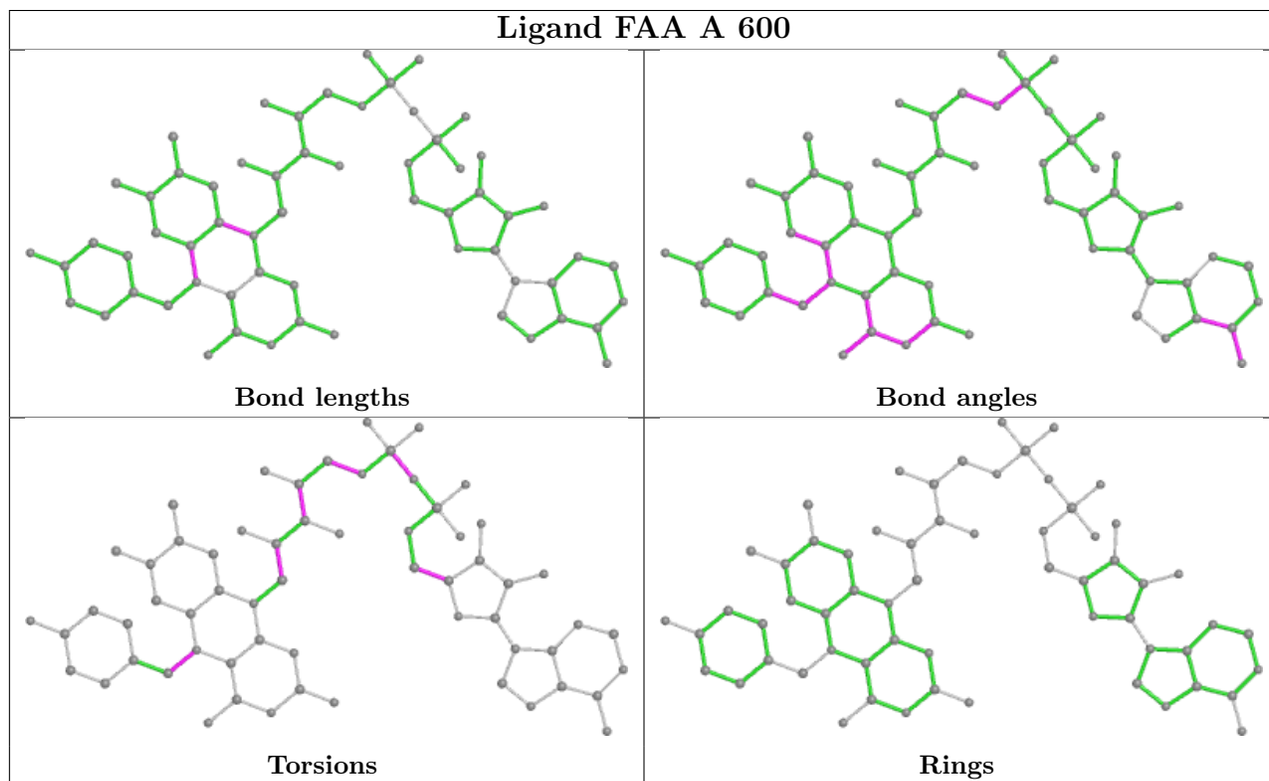
Mol	Chain	Res	Type	Atoms
2	A	600	FAA	C1P-C7P-N5-C4X
2	A	600	FAA	N10-C1'-C2'-O2'
2	A	600	FAA	N10-C1'-C2'-C3'
2	B	600	FAA	C1P-C7P-N5-C4X
2	B	600	FAA	N10-C1'-C2'-O2'

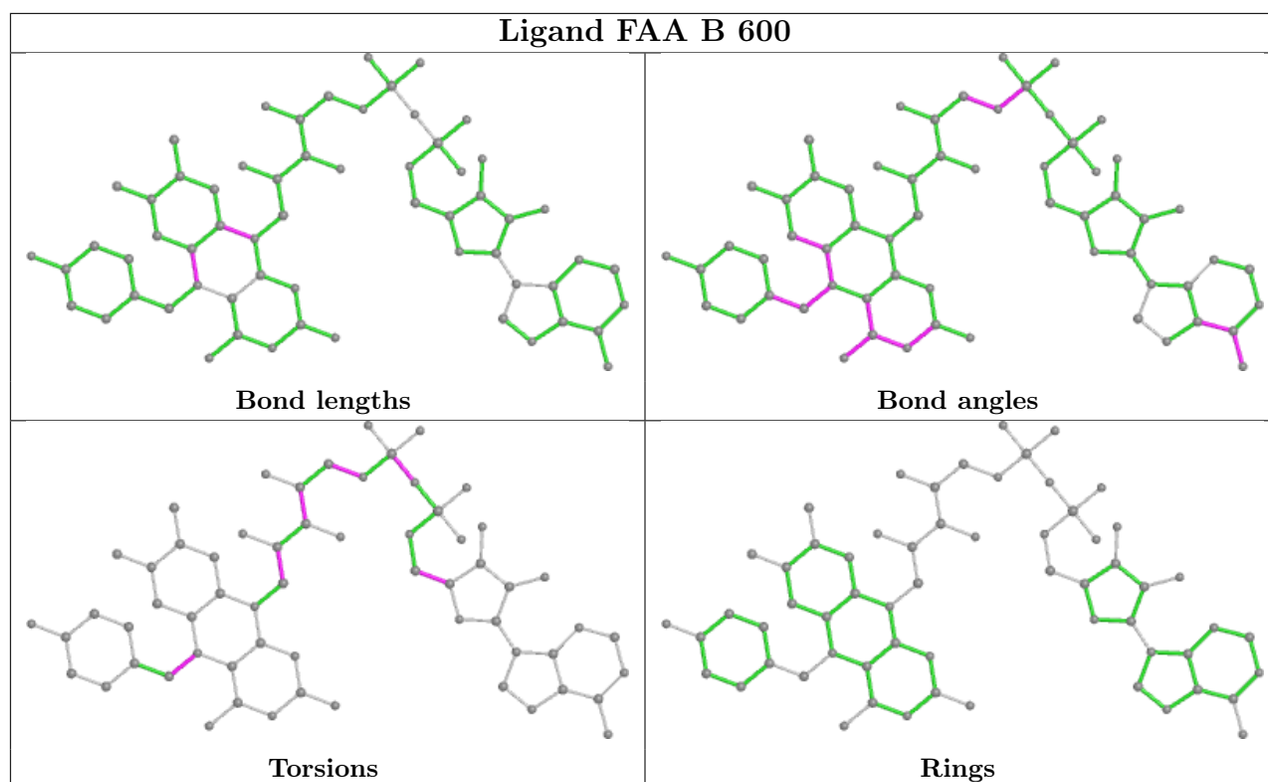
There are no ring outliers.

2 monomers are involved in 26 short contacts:

Mol	Chain	Res	Type	Clashes	Symm-Clashes
2	A	600	FAA	13	0
2	B	600	FAA	13	0

The following is a two-dimensional graphical depiction of Mogul quality analysis of bond lengths, bond angles, torsion angles, and ring geometry for all instances of the Ligand of Interest. In addition, ligands with molecular weight > 250 and outliers as shown on the validation Tables will also be included. For torsion angles, if less than 5% of the Mogul distribution of torsion angles is within 10 degrees of the torsion angle in question, then that torsion angle is considered an outlier. Any bond that is central to one or more torsion angles identified as an outlier by Mogul will be highlighted in the graph. For rings, the root-mean-square deviation (RMSD) between the ring in question and similar rings identified by Mogul is calculated over all ring torsion angles. If the average RMSD is greater than 60 degrees and the minimal RMSD between the ring in question and any Mogul-identified rings is also greater than 60 degrees, then that ring is considered an outlier. The outliers are highlighted in purple. The color gray indicates Mogul did not find sufficient equivalents in the CSD to analyse the geometry.





5.7 Other polymers [i](#)

There are no such residues in this entry.

5.8 Polymer linkage issues [i](#)

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 is therefore empty.

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

EDS was not executed - this section is therefore empty.

6.3 Carbohydrates

EDS was not executed - this section is therefore empty.

6.4 Ligands

EDS was not executed - this section is therefore empty.

6.5 Other polymers

EDS was not executed - this section is therefore empty.