



wwPDB X-ray Structure Validation Summary Report ⓘ

Aug 20, 2023 – 01:52 PM EDT

PDB ID : 2IBM
Title : A novel dimer interface and conformational changes revealed by an X-ray structure of *B. subtilis* SecA
Authors : Zimmer, J.; Li, W.; Rapoport, T.A.
Deposited on : 2006-09-11
Resolution : 3.20 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)
Xtriage (Phenix) : 1.13
EDS : 2.35
buster-report : 1.1.7 (2018)
Percentile statistics : 20191225.v01 (using entries in the PDB archive December 25th 2019)
Refmac : 5.8.0158
CCP4 : 7.0.044 (Gargrove)
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP) : 2.35

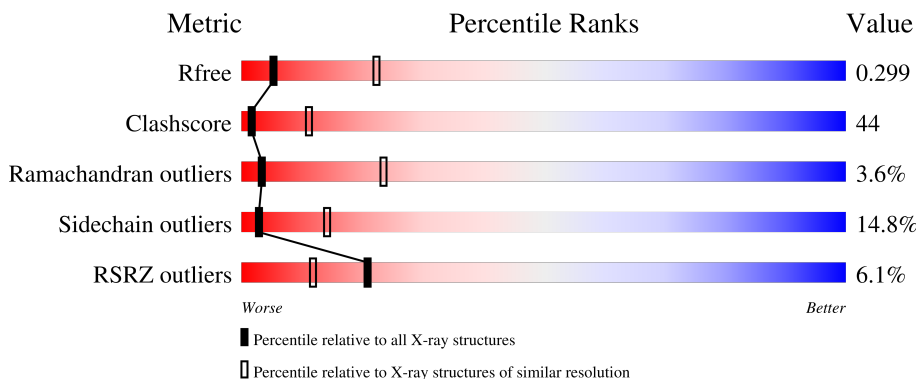
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

X-RAY DIFFRACTION

The reported resolution of this entry is 3.20 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))
R_{free}	130704	1133 (3.20-3.20)
Clashscore	141614	1253 (3.20-3.20)
Ramachandran outliers	138981	1234 (3.20-3.20)
Sidechain outliers	138945	1233 (3.20-3.20)
RSRZ outliers	127900	1095 (3.20-3.20)

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 $\leq 5\%$. The upper red bar (where present) indicates the fraction of residues that have poor fit to the electron density. The numeric value is given above the bar.

Mol	Chain	Length	Quality of chain
1	A	780	
1	B	780	

The following table lists non-polymeric compounds, carbohydrate monomers and non-standard residues in protein, DNA, RNA chains that are outliers for geometric or electron-density-fit criteria:

Mol	Type	Chain	Res	Chirality	Geometry	Clashes	Electron density
2	ADP	A	781	-	-	-	X

2 Entry composition [i](#)

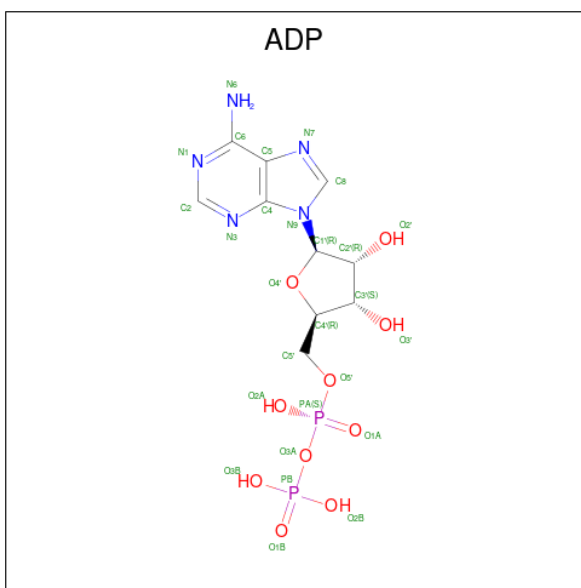
There are 2 unique types of molecules in this entry. The entry contains 12403 atoms, of which 0 are hydrogens and 0 are deuteriums.

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 Preprotein translocase secA subunit.

Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
			Total	C	N	O	S			
1	A	780	Total 6228	C 3899	N 1084	O 1210	S 35	0	0	0
1	B	770	Total 6148	C 3846	N 1072	O 1197	S 33	0	0	0

- Molecule 2 is ADENOSINE-5'-DIPHOSPHATE (three-letter code: ADP) (formula: $C_{10}H_{15}N_5O_{10}P_2$).

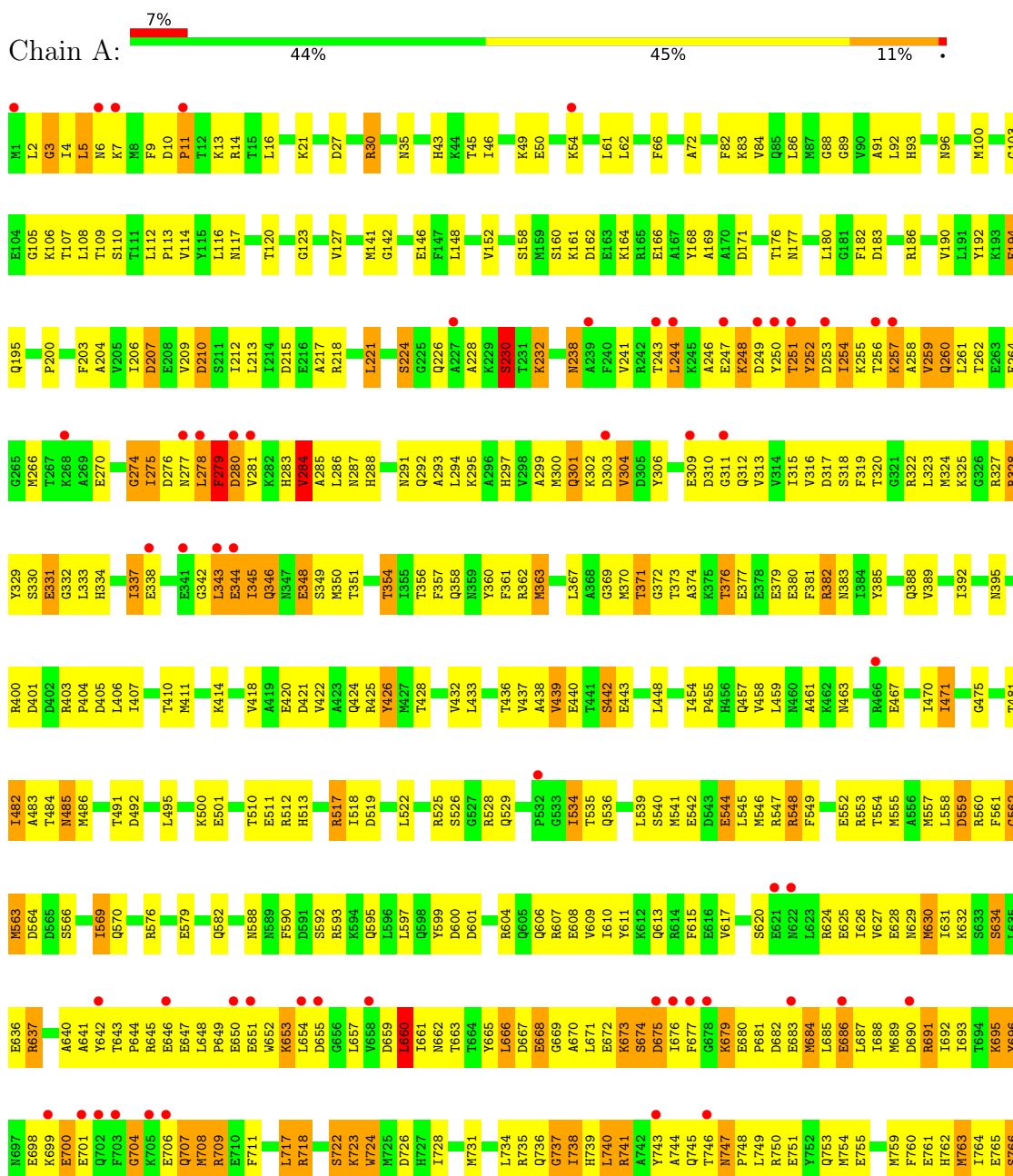


Mol	Chain	Residues	Atoms					ZeroOcc	AltConf
			Total	C	N	O	P		
2	A	1	Total 27	C 10	N 5	O 10	P 2	0	0

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 and electron density. 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. A red dot above a residue indicates a poor fit to the electron density ($RSRZ > 2$). 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.

● Molecule 1: Preprotein translocase secA subunit



I767
E768
D769
E770
V775
M776
K777
A778
E779
I780

• Molecule 1: Preprotein translocase secA subunit



MET	LEU	GLY	ILE	LEU	ASN	LYS	PHE	ASP	P11	T12	T13	R14	T15	L16	N17	R18	Y19	E20	Y33	E34	N35	L36	S37	D38	D39	A40	K42	T45	F48	L52	E53	K54	G55	L61	L62	V68	V69	R70	S73	R74	R75	K83	M87	G88	G89	V90	A91	L92	E99	M100	K101	T102	G103	P104	Q105	K106	T107	L108	T109	R110	T111	L112	P113	V114	D27	A28	L29	L213	G121	R122	V127	N287	L336	Y132	L133	A134	S135	E146	F147	L148	G149	V152	G153	L154	N155	S160	K164	L62	Y168	T172	S175	T176	D177	Y250	L180	L185	R186	D187	V190	L191	K192	K193	E194	Q195	M196	G197	Q198	R199	P200	L201	A204	G205	I206	V209	D210	S211	I212	L213	I214	D215	A216	A217	R218	T219	P220	L221	L222	Q226	L294	A227	A228	K229	S230	T231	K232	L233	M238	A239	F240	V241	R242	T243	L244	K245	A246	E247	K248	D249	D317	S318	T251	Y252	D253	I254	K255	T256	K257	Q260	L261	T262	E263	E264	G265	M266	R267	K268	A269	E270	I275	K340	D276	M277	L278	F279	D280	Q346	N347	E348	K282	H283	V284	A285	L286	N287	H288	H289	Q292	L293	L294	K295	A296	H297	M300	Q301	K302	D303	V304	D305	Y306	V307	F381	R382	M383	N387	Q388	V389	V390	T391	V399	R400	G321	R322	R404	D405	L406	M411	E412	G413	K414	F415	K416	A417	V418	A419	E420	D421	V422	A423	Q424	R425	Y426	H427	Q430	F431	Y432	L433	V434	V439	S442	E443	L444	L445	S446	K447	L448	L449	K450	L454	P455	H456	Q457	V458	L459	M460	A461	K462	M463	E467	L470	T473	E473	A474	G475	Q476	K477	T481	L482	A483	R484	M485	M486	A487	L493	K494	L495	G496	R500	G509	T510	E511	R512	H513	E514	S515	R516	G517	N588	L522	R523	G524	R525	S526	G527	R528	T534	T535	V602	Q536	F537	Y538	L539	S540	M541	E544	L545	M546	R547	R548	F549	G550	A551	E552	R553	T554	M555	A556	R557	D559	R560	F561	G562	M563	D564	D565	S566	T567	P568	L569	Q570	S571	K572	M573	V574	S575	E579	S580	S581	Q582	K583	R584	V585	L586	G587	G588	V589	D592	R593	K594	Q595	S596	L597	Q598	Y599	L602	L603	Q606	R607	E608	V609	I610	Y611	L612	K613	Q614	F615	E616	V617	I618	D619	S620	E621	N622	R623	E624	I625	V627	M630	I631	E636	R637	K638	A639	L639	A640	E641	Y642	T643	P644	E645	E646	E647	L648	P649	E650	E651	M652	K653	L654	D655	G656	L657	V658	D659	T663	T664	Y665	L666	D667	E668	K673	I676	K679	E680	P681	D682	E683	M684	L685	E686	L687	I688	I682	L693	T694	K695	Y696	E700	E701	Q702	F703	G704	R705	E706	M708	R709	E710	E711	E712	K713	V714	T715	V716	L717	R718	A719	V720	D721	S722	K723	W724	M725	D726	H727	I728	M731	D732	Q733	L734	R735	I738	H739	L740	R741	A742	Y743	A744	G745	T746	N747	P748	L749	R750	M754	E755	M759	F760	M763	T764
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4 Data and refinement statistics

Property	Value	Source
Space group	I 21 21 21	Depositor
Cell constants a, b, c, α , β , γ	125.83Å 166.83Å 211.99Å 90.00° 90.00° 90.00°	Depositor
Resolution (Å)	40.53 – 3.20 40.53 – 3.10	Depositor EDS
% Data completeness (in resolution range)	92.7 (40.53-3.20) 90.9 (40.53-3.10)	Depositor EDS
R_{merge}	(Not available)	Depositor
R_{sym}	0.07	Depositor
$\langle I/\sigma(I) \rangle$ ¹	2.18 (at 3.12Å)	Xtrriage
Refinement program	CNS 1.1	Depositor
R, R_{free}	0.321 , 0.323 0.293 , 0.299	Depositor DCC
R_{free} test set	3431 reflections (8.64%)	wwPDB-VP
Wilson B-factor (Å ²)	69.9	Xtrriage
Anisotropy	0.095	Xtrriage
Bulk solvent k_{sol} (e/Å ³), B_{sol} (Å ²)	0.35 , 39.2	EDS
L-test for twinning ²	$\langle L \rangle = 0.49$, $\langle L^2 \rangle = 0.32$	Xtrriage
Estimated twinning fraction	No twinning to report.	Xtrriage
F_o, F_c correlation	0.85	EDS
Total number of atoms	12403	wwPDB-VP
Average B, all atoms (Å ²)	47.0	wwPDB-VP

Xtrriage's analysis on translational NCS is as follows: *The largest off-origin peak in the Patterson function is 3.44% of the height of the origin peak. No significant pseudotranslation is detected.*

¹Intensities estimated from amplitudes.

²Theoretical values of $\langle |L| \rangle$, $\langle L^2 \rangle$ for acentric reflections are 0.5, 0.333 respectively for untwinned datasets, and 0.375, 0.2 for perfectly twinned datasets.

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: ADP

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	0.63	2/6316 (0.0%)	0.88	9/8492 (0.1%)
1	B	0.68	1/6235 (0.0%)	0.90	19/8384 (0.2%)
All	All	0.65	3/12551 (0.0%)	0.89	28/16876 (0.2%)

All (3) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
1	B	354	THR	CB-OG1	-13.18	1.16	1.43
1	A	279	PHE	C-N	6.88	1.49	1.34
1	A	701	GLU	N-CA	5.04	1.56	1.46

The worst 5 of 28 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	B	244	LEU	C-N-CA	10.24	147.31	121.70
1	B	247	GLU	CA-C-N	-9.45	96.40	117.20
1	B	353	ALA	N-CA-CB	8.80	122.42	110.10
1	B	244	LEU	CA-C-N	-8.14	99.29	117.20
1	A	11	PRO	N-CA-C	7.27	130.99	112.10

There are no chirality outliers.

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	6228	0	6227	545	0
1	B	6148	0	6139	578	0
2	A	27	0	12	3	0
All	All	12403	0	12378	1092	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 44.

The worst 5 of 1092 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)
1:B:620:SER:HB3	1:B:625:GLU:CG	1.49	1.39
1:A:723:LYS:HD3	1:A:763:MET:SD	1.67	1.35
1:A:747:ASN:H	1:A:748:PRO:CD	1.39	1.31
1:A:10:ASP:HB3	1:A:11:PRO:CD	1.61	1.29
1:A:303:ASP:O	1:A:304:VAL:HG23	1.10	1.28

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	778/780 (100%)	647 (83%)	99 (13%)	32 (4%)	3	21
1	B	768/780 (98%)	626 (82%)	118 (15%)	24 (3%)	4	26
All	All	1546/1560 (99%)	1273 (82%)	217 (14%)	56 (4%)	3	23

5 of 56 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	A	230	SER
1	A	259	VAL

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Mol	Chain	Res	Type
1	A	275	ILE
1	A	280	ASP
1	A	304	VAL

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	670/670 (100%)	576 (86%)	94 (14%)	3	16
1	B	661/670 (99%)	558 (84%)	103 (16%)	2	12
All	All	1331/1340 (99%)	1134 (85%)	197 (15%)	3	14

5 of 197 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	B	266	MET
1	B	473	GLU
1	B	280	ASP
1	B	363	MET
1	B	514	GLU

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 35 such sidechains are listed below:

Mol	Chain	Res	Type
1	B	460	ASN
1	B	463	ASN
1	B	739	HIS
1	A	727	HIS
1	A	697	ASN

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](#)

1 ligand is 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	ADP	A	781	-	24,29,29	1.36	4 (16%)	29,45,45	1.56	6 (20%)

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	ADP	A	781	-	-	2/12/32/32	0/3/3/3

All (4) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
2	A	781	ADP	O3'-C3'	3.08	1.50	1.43
2	A	781	ADP	C5-C4	2.99	1.48	1.40
2	A	781	ADP	C2-N3	2.28	1.35	1.32
2	A	781	ADP	O4'-C1'	2.09	1.44	1.41

The worst 5 of 6 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
2	A	781	ADP	C3'-C2'-C1'	4.20	107.31	100.98
2	A	781	ADP	N3-C2-N1	-3.65	122.97	128.68
2	A	781	ADP	C4-C5-N7	-2.99	106.28	109.40
2	A	781	ADP	PA-O3A-PB	-2.37	124.70	132.83
2	A	781	ADP	O4'-C4'-C3'	2.35	109.76	105.11

There are no chirality outliers.

All (2) torsion outliers are listed below:

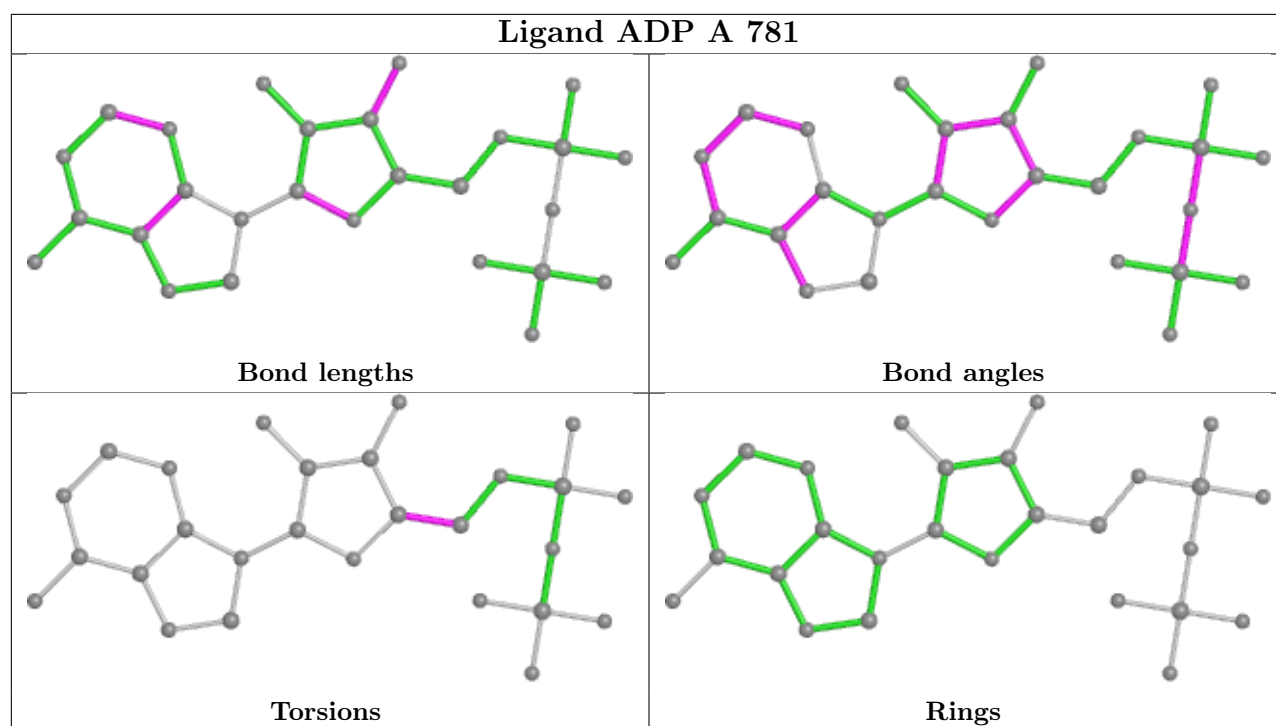
Mol	Chain	Res	Type	Atoms
2	A	781	ADP	O4'-C4'-C5'-O5'
2	A	781	ADP	C3'-C4'-C5'-O5'

There are no ring outliers.

1 monomer is involved in 3 short contacts:

Mol	Chain	Res	Type	Clashes	Symm-Clashes
2	A	781	ADP	3	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 [i](#)

6.1 Protein, DNA and RNA chains [i](#)

In the following table, the column labelled ‘#RSRZ > 2’ contains the number (and percentage) of RSRZ outliers, followed by percent RSRZ outliers for the chain as percentile scores relative to all X-ray entries and entries of similar resolution. The OWAB column contains the minimum, median, 95th percentile and maximum values of the occupancy-weighted average B-factor per residue. The column labelled ‘Q < 0.9’ lists the number of (and percentage) of residues with an average occupancy less than 0.9.

Mol	Chain	Analysed	<RSRZ>	#RSRZ>2	OWAB(Å ²)	Q<0.9
1	A	780/780 (100%)	0.35	55 (7%) 16 9	20, 45, 80, 187	0
1	B	770/780 (98%)	0.27	40 (5%) 27 15	18, 46, 61, 187	0
All	All	1550/1560 (99%)	0.31	95 (6%) 21 12	18, 46, 73, 187	0

The worst 5 of 95 RSRZ outliers are listed below:

Mol	Chain	Res	Type	RSRZ
1	B	622	ASN	7.7
1	A	622	ASN	7.3
1	B	623	LEU	5.6
1	A	654	LEU	5.2
1	B	261	LEU	5.1

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

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

6.3 Carbohydrates [i](#)

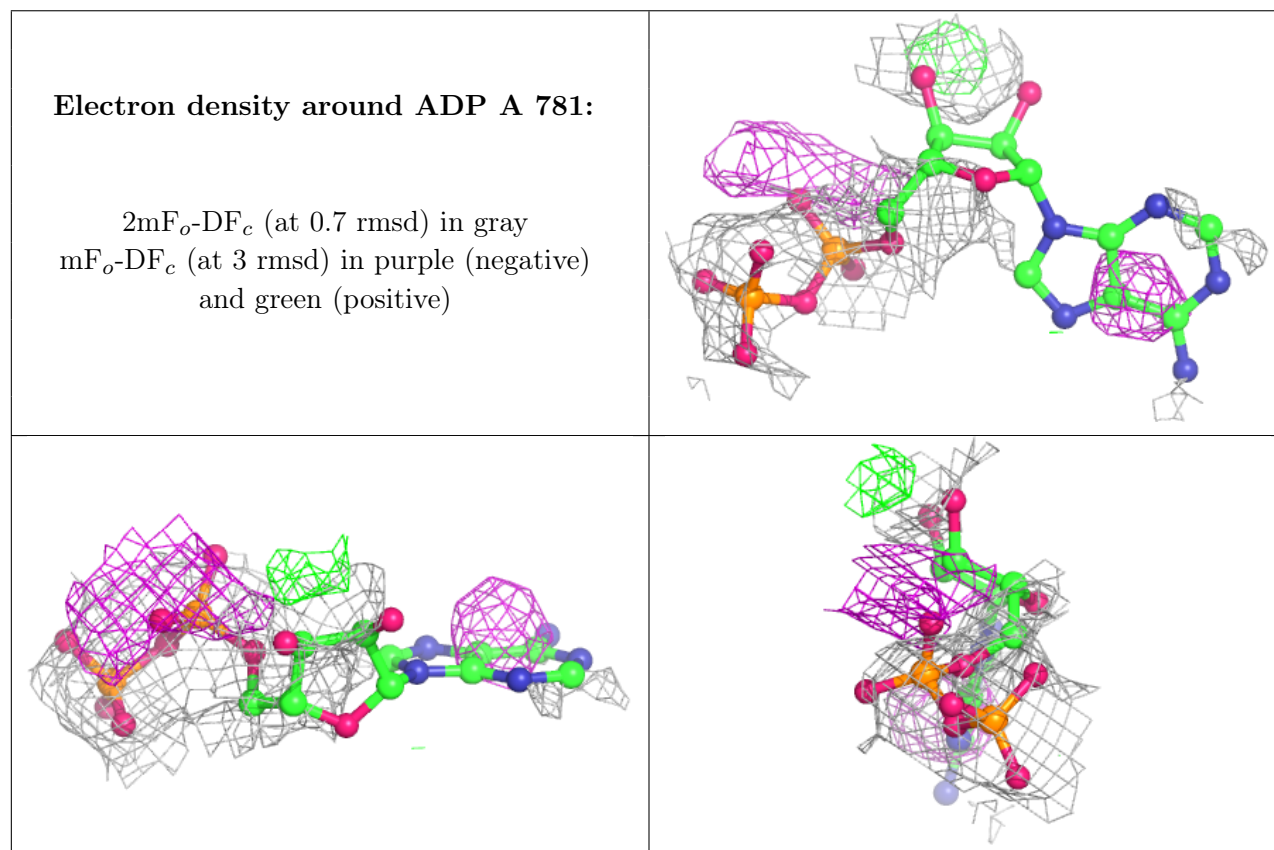
There are no monosaccharides in this entry.

6.4 Ligands [i](#)

In the following table, the Atoms column lists the number of modelled atoms in the group and the number defined in the chemical component dictionary. The B-factors column lists the minimum, median, 95th percentile and maximum values of B factors of atoms in the group. The column labelled ‘Q < 0.9’ lists the number of atoms with occupancy less than 0.9.

Mol	Type	Chain	Res	Atoms	RSCC	RSR	B-factors(\AA^2)	Q<0.9
2	ADP	A	781	27/27	0.62	0.63	47,108,108,108	0

The following is a graphical depiction of the model fit to experimental electron density of all instances of the Ligand of Interest. In addition, ligands with molecular weight > 250 and outliers as shown on the geometry validation Tables will also be included. Each fit is shown from different orientation to approximate a three-dimensional view.



6.5 Other polymers [i](#)

There are no such residues in this entry.