



wwPDB X-ray Structure Validation Summary Report ⓘ

Mar 10, 2024 – 01:22 AM EST

PDB ID : 4EIW
Title : Whole cytosolic region of atp-dependent metalloprotease FtsH (G399L)
Authors : Suno, R.; Niwa, H.; Tsuchiya, D.; Yoshida, M.; Morikawa, K.
Deposited on : 2012-04-06
Resolution : 3.90 Å(reported)

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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 : 2.36
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) : 2.36

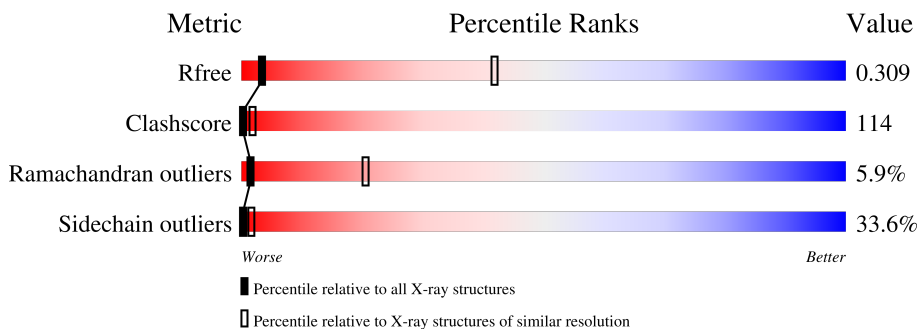
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

X-RAY DIFFRACTION

The reported resolution of this entry is 3.90 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))
R_{free}	130704	1002 (4.14-3.66)
Clashscore	141614	1004 (4.12-3.68)
Ramachandran outliers	138981	1021 (4.14-3.66)
Sidechain outliers	138945	1014 (4.14-3.66)

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 $\leq 5\%$.

Mol	Chain	Length	Quality of chain
1	A	508	16% (green), 49% (yellow), 23% (orange), 10% (red), 10% (grey)
1	B	508	9% (green), 53% (yellow), 22% (orange), 12% (red), 12% (grey)
1	C	508	16% (green), 49% (yellow), 21% (orange), 10% (red), 10% (grey)
1	D	508	9% (green), 52% (yellow), 23% (orange), 12% (red), 12% (grey)
1	E	508	17% (green), 49% (yellow), 22% (orange), 10% (red), 10% (grey)
1	F	508	10% (green), 52% (yellow), 23% (orange), 12% (red), 12% (grey)

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	B	2001	-	-	X	-
2	ADP	C	1001	-	-	X	-
2	ADP	D	2001	-	-	X	-
2	ADP	E	1001	-	-	X	-
2	ADP	F	2001	-	-	X	-

2 Entry composition [i](#)

There are 2 unique types of molecules in this entry. The entry contains 21429 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 ATP-dependent zinc metalloprotease FtsH.

Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
			Total	C	N	O	S			
1	A	458	3578	2245	658	662	13	0	0	0
1	B	446	3511	2206	641	651	13	0	0	0
1	C	458	3578	2245	658	662	13	0	0	0
1	D	446	3511	2206	641	651	13	0	0	0
1	E	458	3578	2245	658	662	13	0	0	0
1	F	446	3511	2206	641	651	13	0	0	0

There are 60 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
A	117	GLY	-	expression tag	UNP Q5SI82
A	118	PRO	-	expression tag	UNP Q5SI82
A	119	LEU	-	expression tag	UNP Q5SI82
A	120	GLY	-	expression tag	UNP Q5SI82
A	121	SER	-	expression tag	UNP Q5SI82
A	122	HIS	-	expression tag	UNP Q5SI82
A	123	MET	-	expression tag	UNP Q5SI82
A	124	GLY	-	expression tag	UNP Q5SI82
A	125	ALA	-	expression tag	UNP Q5SI82
A	399	LEU	GLY	engineered mutation	UNP Q5SI82
B	117	GLY	-	expression tag	UNP Q5SI82
B	118	PRO	-	expression tag	UNP Q5SI82
B	119	LEU	-	expression tag	UNP Q5SI82
B	120	GLY	-	expression tag	UNP Q5SI82
B	121	SER	-	expression tag	UNP Q5SI82
B	122	HIS	-	expression tag	UNP Q5SI82
B	123	MET	-	expression tag	UNP Q5SI82

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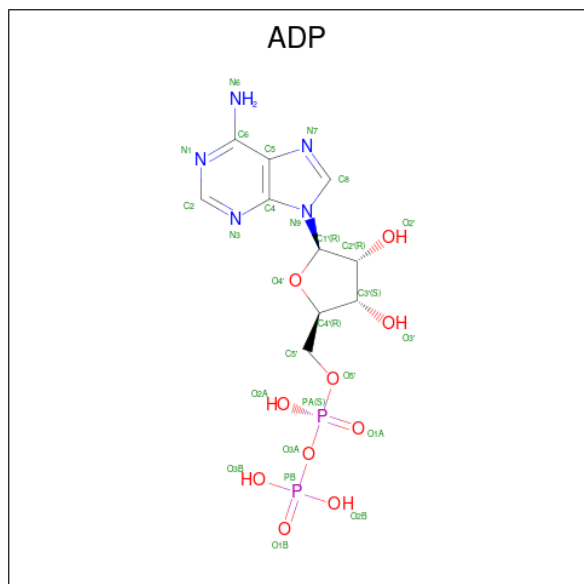
Chain	Residue	Modelled	Actual	Comment	Reference
B	124	GLY	-	expression tag	UNP Q5SI82
B	125	ALA	-	expression tag	UNP Q5SI82
B	399	LEU	GLY	engineered mutation	UNP Q5SI82
C	117	GLY	-	expression tag	UNP Q5SI82
C	118	PRO	-	expression tag	UNP Q5SI82
C	119	LEU	-	expression tag	UNP Q5SI82
C	120	GLY	-	expression tag	UNP Q5SI82
C	121	SER	-	expression tag	UNP Q5SI82
C	122	HIS	-	expression tag	UNP Q5SI82
C	123	MET	-	expression tag	UNP Q5SI82
C	124	GLY	-	expression tag	UNP Q5SI82
C	125	ALA	-	expression tag	UNP Q5SI82
C	399	LEU	GLY	engineered mutation	UNP Q5SI82
D	117	GLY	-	expression tag	UNP Q5SI82
D	118	PRO	-	expression tag	UNP Q5SI82
D	119	LEU	-	expression tag	UNP Q5SI82
D	120	GLY	-	expression tag	UNP Q5SI82
D	121	SER	-	expression tag	UNP Q5SI82
D	122	HIS	-	expression tag	UNP Q5SI82
D	123	MET	-	expression tag	UNP Q5SI82
D	124	GLY	-	expression tag	UNP Q5SI82
D	125	ALA	-	expression tag	UNP Q5SI82
D	399	LEU	GLY	engineered mutation	UNP Q5SI82
E	117	GLY	-	expression tag	UNP Q5SI82
E	118	PRO	-	expression tag	UNP Q5SI82
E	119	LEU	-	expression tag	UNP Q5SI82
E	120	GLY	-	expression tag	UNP Q5SI82
E	121	SER	-	expression tag	UNP Q5SI82
E	122	HIS	-	expression tag	UNP Q5SI82
E	123	MET	-	expression tag	UNP Q5SI82
E	124	GLY	-	expression tag	UNP Q5SI82
E	125	ALA	-	expression tag	UNP Q5SI82
E	399	LEU	GLY	engineered mutation	UNP Q5SI82
F	117	GLY	-	expression tag	UNP Q5SI82
F	118	PRO	-	expression tag	UNP Q5SI82
F	119	LEU	-	expression tag	UNP Q5SI82
F	120	GLY	-	expression tag	UNP Q5SI82
F	121	SER	-	expression tag	UNP Q5SI82
F	122	HIS	-	expression tag	UNP Q5SI82
F	123	MET	-	expression tag	UNP Q5SI82
F	124	GLY	-	expression tag	UNP Q5SI82
F	125	ALA	-	expression tag	UNP Q5SI82

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Chain	Residue	Modelled	Actual	Comment	Reference
F	399	LEU	GLY	engineered mutation	UNP Q5SI82

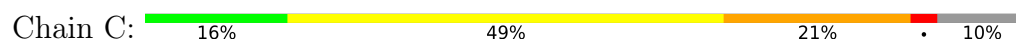
- Molecule 2 is ADENOSINE-5'-DIPHOSPHATE (three-letter code: ADP) (formula: $C_{10}H_{15}N_5O_{10}P_2$).



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

V237	V297	T357	H418	E479	K541	GLU
R236	M286	P358	E419	E480	R542	GLU
L240	A299	G359	A300	I481	T543	ARG
F241	A300	F360	G421	V482	D644	GLU
E242	T301	V361	H422	F483	V547	PRO
T243	R303	G362	A423	D484	R548	PRO
A244	P304	D364	L424	D485	R549	ARG
K245	F305	D364		V486	R549	VAL
R246	L306	E366	H427	T487	L550	VAL
H247	L307	E367	F428	T488	F551	VAL
E248	L307	N367	L429	V489	E552	PRO
A249	F308	L368	E430	A490	E553	LYS
P249	P309	L369	H431	E491	O554	VAL
C250	A310	N370	A432	N492	Y555	LYS
I251	L311	E371	D433	D493	Q556	PRO
V252	L312	A372	G434	F494	R557	GLY
F253	R313	A373	G435	R495	V558	GLY
I254	P314	L374	H436	K496	K559	ALA
D255	G315	L375	K437	A497	A560	LEU
E256	R316	A376	V438	L500	L561	GLY
I257	F317	R377	T439	L501	L562	ALA
A258	D318	R378	I440	A502	L563	ALA
D259	R319	E379	V441	R503	E564	SER
V260	Q320	G380	P442	M504	E565	THR
G261	L321	R381	R443	I505	R566	LYS
R262	A322	K382	G444	L506	E567	SER
K263	I323	K383	R445	T506	V568	R143
ARG	D324	I384	A446	L507	L569	A144
GLY	A325	T385	L447	W508	E570	R145
SER	P326	M386	G448	G509	R571	V146
GLY	D327	K387	F449	M510	V572	A210
VAL	V328	D388	M450	R511	L576	T148
GLY	K329	L389	M451	P512	L577	E149
GLY	G330	E390	P452	F513	L578	A150
GLY	R331	E391	R453	E514	R579	P151
D273	Q333	A392	R454	V517	E580	K152
E274	I334	A393	D456	A518	T581	V153
R275	L335	D394	M457	F519	L582	F154
E276	R336	R395	L458	A520	T583	F155
Q277	I337	V396	L459	V521	A584	K156
T278	H338	M398	H459	R522	E585	D157
L279	H339	A399	S461	R523	E586	E166
N280	R340	P400	R462	D524	F587	E167
Q281	K341	A401	K463	T525	Q588	L168
L282	K342	K402	R464	V526	R589	K169
L283	P343	K403	L465	L527	E590	E170
V284	L344	S404	L466	G528	V591	I171
E285	A345	L405	D467	G529	E592	V172
M286	E346	V406	Q468	Y530	G593	A173
D287	D347	L407	I469	D531	L594	E174
G288	V348	S408	A470	V532	P595	L175
F289	D349	R409	V471	R533	L596	K176
E290	R410	R410	R410	K534	E597	E177
K291	A351	D411	L473	S535	A598	V178
D292	L352	R412	A474	Y536	P599	F174
T293	L353	R413	G475	E537	E600	L175
A294	A354	L414	R476	GLU	ALA	K177
I295	K355	Y417	A478	ARG	ALA	M177
V296	R356					P178

● Molecule 1: ATP-dependent zinc metalloprotease FtsH



GLY	S179	A244	A310	E371	A432	R486	L561
PRO	R180	K245	L311	A372	D433	A497	L562
LEU	F181	R246	L312	A373	H436	L500	L563
GLY	H182	H247	R313	L374	K437	L501	E564
SER	E183	A248	R316	L375	V438	R502	K565
PRO	M184	P249	F317	A376	T439	R503	E566
ARG		C250	D318	A377	L440	M504	E567
VAL	I188	I251	R319	R378	V441	S505	V568
VAL	P189	V252	R319	E379	R442	T506	L569
ALA	K190	K190	Q320	G380	P442	E507	L570
ARG	G191	I254	L321	R381	R443	E508	F571
ASN	L192	I254	A322	R382	G444	V509	A572
GLY	L193	V260	L323	K383	R445	G509	A573
VAL	L194	G261	R324	I384	A446	M510	A574
PRO	V195	R262	A325	T385	L447	H511	V575
GLY	G196	K263	P326	M386	G448	L512	L576
GLY	P197	R264	D327	K387	F449	E513	L577
ALA	G199	G265	V328	R388	M450	F514	E578
ASP	G199	S266	K329	L389	M451	G515	R579
SER	V200	G267	G330	E390	P452	P516	E580
ALA	G201	V288	R331	E391	L458	V517	E581
PHE	K202		E332	A392	R454	A518	L582
SER	T203	G271	Q333	A393	E455	F519	L583
THR	H204	G271	I334	D394	D456	Y520	F584
LYS	L205	D273	L335	R395	M457	L527	V591
SER	A206	E274	R336	V396	L458	G528	E585
R143	R207	R275	I337	M398	H459	Q529	L584
A144	A208	R276	H338	N399	W460	W530	P595
R145	V209	Q277	A339	L399	S461	D531	P596
V146	A210	T278	R340	P400	R462	E537	L596
A210	G211	L279	G341	A401	K463	Q534	E597
T148	E212	N280	K342	K402	R464	O534	R598
E149	A213	Q281	P343	L405	L465	F535	L599
A150	R214	L282	L344	V406	L466	D536	L599
P151	V215	L283	A345	D467	Q468	R533	L599
K152	K152	M286	E346	L407	E346	E538	E597
V153	T154	D287	D347	S408	L469	R539	R599
F155	F155	G288	D349	R410	A470	A540	A598
K156	A220	F289	L350	D411	A472	ALU	GLU
D157	S221	E290	A351	R412	V482	R542	ALA
V158	G222	K291	L352	R413	F483	I543	ARG
A159	S223	D292	L353	I414	D484	L543	ARG
E163	D224	T293	A354	T415	D485	R548	VAL
A164	F225	A294	K355	A416	V486	E549	VAL
K165	V226	I295	R356	Y417	T487	L550	VAL
E166	M228	V296	T357	H418	V487	I551	LYS
E167	E167	V297	P358	E419	F483	E552	VAL
K168	L168	M298	G359	A420	D484	E553	LYS
K169	K169	A299	F360	H421	D485	O554	PRO
E170	E170	A300	V361	H422	V486	Y555	PRO
I171	I171	T301	G362	A423	T488	R557	GLY
V172	R236	R303	A363	L424	F488	V558	ALA
A173	V237	K302	D364	A425	G489	E586	ALA
E174	D239	P304	L365	A426	F493	L429	LEU
L175	L240	D305	E366	H427	F428	F494	LEU
K176	F241	I306	M367	F428	L428	M177	ALA
P178	T243	P309	L368	E430	F494	ALA	ALA
			N370	H431	R495	ARG	ARG

4 Data and refinement statistics i

Property	Value	Source
Space group	P 31 2 1	Depositor
Cell constants a, b, c, α , β , γ	146.15Å 146.15Å 349.06Å 90.00° 90.00° 120.00°	Depositor
Resolution (Å)	71.53 – 3.90 71.53 – 3.90	Depositor EDS
% Data completeness (in resolution range)	97.1 (71.53-3.90) 97.1 (71.53-3.90)	Depositor EDS
R_{merge}	(Not available)	Depositor
R_{sym}	(Not available)	Depositor
$\langle I/\sigma(I) \rangle$ ¹	7.46 (at 3.89Å)	Xtrriage
Refinement program	REFMAC 5.6.0117	Depositor
R, R_{free}	0.299 , 0.312 0.298 , 0.309	Depositor DCC
R_{free} test set	1967 reflections (5.01%)	wwPDB-VP
Wilson B-factor (Å ²)	99.1	Xtrriage
Anisotropy	0.272	Xtrriage
Bulk solvent k_{sol} (e/Å ³), B_{sol} (Å ²)	0.17 , 19.0	EDS
L-test for twinning ²	$\langle L \rangle = 0.27$, $\langle L^2 \rangle = 0.11$	Xtrriage
Estimated twinning fraction	0.237 for -h,-k,l	Xtrriage
F_o, F_c correlation	0.78	EDS
Total number of atoms	21429	wwPDB-VP
Average B, all atoms (Å ²)	112.0	wwPDB-VP

Xtrriage's analysis on translational NCS is as follows: *The largest off-origin peak in the Patterson function is 1.77% 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	0.60	3/3636 (0.1%)	0.95	16/4906 (0.3%)
1	B	0.63	3/3568 (0.1%)	1.01	18/4815 (0.4%)
1	C	0.59	2/3636 (0.1%)	1.03	24/4906 (0.5%)
1	D	0.60	2/3568 (0.1%)	0.98	17/4815 (0.4%)
1	E	0.61	6/3636 (0.2%)	0.96	13/4906 (0.3%)
1	F	0.58	2/3568 (0.1%)	0.96	12/4815 (0.2%)
All	All	0.60	18/21612 (0.1%)	0.98	100/29163 (0.3%)

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	2
1	B	0	5
1	C	0	1
1	D	0	6
1	E	0	1
1	F	0	5
All	All	0	20

The worst 5 of 18 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
1	B	586	GLU	CD-OE1	-12.04	1.12	1.25
1	C	214	ARG	CZ-NH2	8.16	1.43	1.33
1	E	214	ARG	CZ-NH2	-8.12	1.22	1.33
1	E	319	ARG	CZ-NH1	-7.39	1.23	1.33
1	D	586	GLU	CD-OE1	-6.72	1.18	1.25

The worst 5 of 100 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	E	319	ARG	NE-CZ-NH2	18.11	129.36	120.30
1	C	207	ARG	NE-CZ-NH1	-15.37	112.61	120.30
1	A	316	ARG	NE-CZ-NH1	-15.14	112.73	120.30
1	B	236	ARG	NE-CZ-NH1	-14.64	112.98	120.30
1	C	207	ARG	NE-CZ-NH2	13.93	127.26	120.30

There are no chirality outliers.

5 of 20 planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	A	213	ALA	Peptide
1	A	532	VAL	Peptide
1	B	244	ALA	Peptide
1	B	288	GLY	Peptide
1	B	382	ARG	Peptide

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	3578	0	3623	862	5
1	B	3511	0	3556	851	3
1	C	3578	0	3623	815	0
1	D	3511	0	3556	851	3
1	E	3578	0	3623	805	3
1	F	3511	0	3556	857	2
2	A	27	0	12	8	0
2	B	27	0	12	9	0
2	C	27	0	12	10	0
2	D	27	0	12	9	0
2	E	27	0	12	11	0
2	F	27	0	12	10	0
All	All	21429	0	21609	4895	8

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 114.

The worst 5 of 4895 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:E:416:ALA:CB	1:E:577:LEU:HD23	1.18	1.63
1:A:416:ALA:HB3	1:A:577:LEU:CD2	1.33	1.58
1:A:416:ALA:CB	1:A:577:LEU:HD23	1.15	1.55
1:F:376:ALA:CA	1:F:381:ARG:HD2	1.31	1.55
1:E:416:ALA:HB3	1:E:577:LEU:CD2	1.35	1.55

The worst 5 of 8 symmetry-related close contacts are listed below. The label for Atom-2 includes the symmetry operator and encoded unit-cell translations to be applied.

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:A:570:GLU:CG	1:D:382:ARG:NH2[3_564]	1.61	0.59
1:A:417:TYR:OH	1:D:382:ARG:NE[3_564]	1.74	0.46
1:A:177:ASN:OD1	1:E:214:ARG:NH2[6_665]	1.85	0.35
1:B:238:ARG:NE	1:F:378:ARG:NH2[6_665]	1.91	0.29
1:A:570:GLU:CB	1:D:382:ARG:NH2[3_564]	2.08	0.12

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	456/508 (90%)	323 (71%)	111 (24%)	22 (5%)	2	24
1	B	442/508 (87%)	289 (65%)	125 (28%)	28 (6%)	1	19
1	C	456/508 (90%)	323 (71%)	112 (25%)	21 (5%)	2	25
1	D	442/508 (87%)	288 (65%)	121 (27%)	33 (8%)	1	16
1	E	456/508 (90%)	324 (71%)	110 (24%)	22 (5%)	2	24
1	F	442/508 (87%)	288 (65%)	120 (27%)	34 (8%)	1	16
All	All	2694/3048 (88%)	1835 (68%)	699 (26%)	160 (6%)	1	20

5 of 160 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	A	153	VAL
1	A	511	HIS
1	B	153	VAL
1	B	274	GLU
1	B	379	GLU

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	365/402 (91%)	240 (66%)	125 (34%)	0	1
1	B	361/402 (90%)	242 (67%)	119 (33%)	0	1
1	C	365/402 (91%)	240 (66%)	125 (34%)	0	1
1	D	361/402 (90%)	241 (67%)	120 (33%)	0	1
1	E	365/402 (91%)	244 (67%)	121 (33%)	0	1
1	F	361/402 (90%)	240 (66%)	121 (34%)	0	1
All	All	2178/2412 (90%)	1447 (66%)	731 (34%)	0	1

5 of 731 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	D	460	TRP
1	E	439	THR
1	D	533	ARG
1	D	459	HIS
1	E	264	ARG

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 37 such sidechains are listed below:

Mol	Chain	Res	Type
1	E	468	GLN
1	F	338	HIS

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Mol	Chain	Res	Type
1	E	496	GLN
1	F	281	GLN
1	B	431	HIS

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](#)

6 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	ADP	A	1001	-	24,29,29	1.10	2 (8%)	29,45,45	1.50	4 (13%)
2	ADP	C	1001	-	24,29,29	1.10	3 (12%)	29,45,45	1.56	4 (13%)
2	ADP	D	2001	-	24,29,29	1.03	3 (12%)	29,45,45	1.45	5 (17%)
2	ADP	B	2001	-	24,29,29	1.05	2 (8%)	29,45,45	1.49	4 (13%)
2	ADP	E	1001	-	24,29,29	1.06	2 (8%)	29,45,45	1.45	4 (13%)
2	ADP	F	2001	-	24,29,29	1.01	1 (4%)	29,45,45	1.52	4 (13%)

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	1001	-	-	2/12/32/32	0/3/3/3
2	ADP	C	1001	-	-	2/12/32/32	0/3/3/3
2	ADP	D	2001	-	-	2/12/32/32	0/3/3/3
2	ADP	B	2001	-	-	3/12/32/32	0/3/3/3
2	ADP	E	1001	-	-	2/12/32/32	0/3/3/3
2	ADP	F	2001	-	-	2/12/32/32	0/3/3/3

The worst 5 of 13 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
2	A	1001	ADP	C5-C4	2.78	1.48	1.40
2	F	2001	ADP	C5-C4	2.76	1.48	1.40
2	E	1001	ADP	C5-C4	2.73	1.48	1.40
2	C	1001	ADP	C5-C4	2.62	1.47	1.40
2	B	2001	ADP	C5-C4	2.59	1.47	1.40

The worst 5 of 25 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
2	A	1001	ADP	PA-O3A-PB	-4.26	118.21	132.83
2	C	1001	ADP	PA-O3A-PB	-4.20	118.40	132.83
2	E	1001	ADP	PA-O3A-PB	-4.05	118.92	132.83
2	B	2001	ADP	PA-O3A-PB	-3.99	119.13	132.83
2	C	1001	ADP	N3-C2-N1	-3.72	122.86	128.68

There are no chirality outliers.

5 of 13 torsion outliers are listed below:

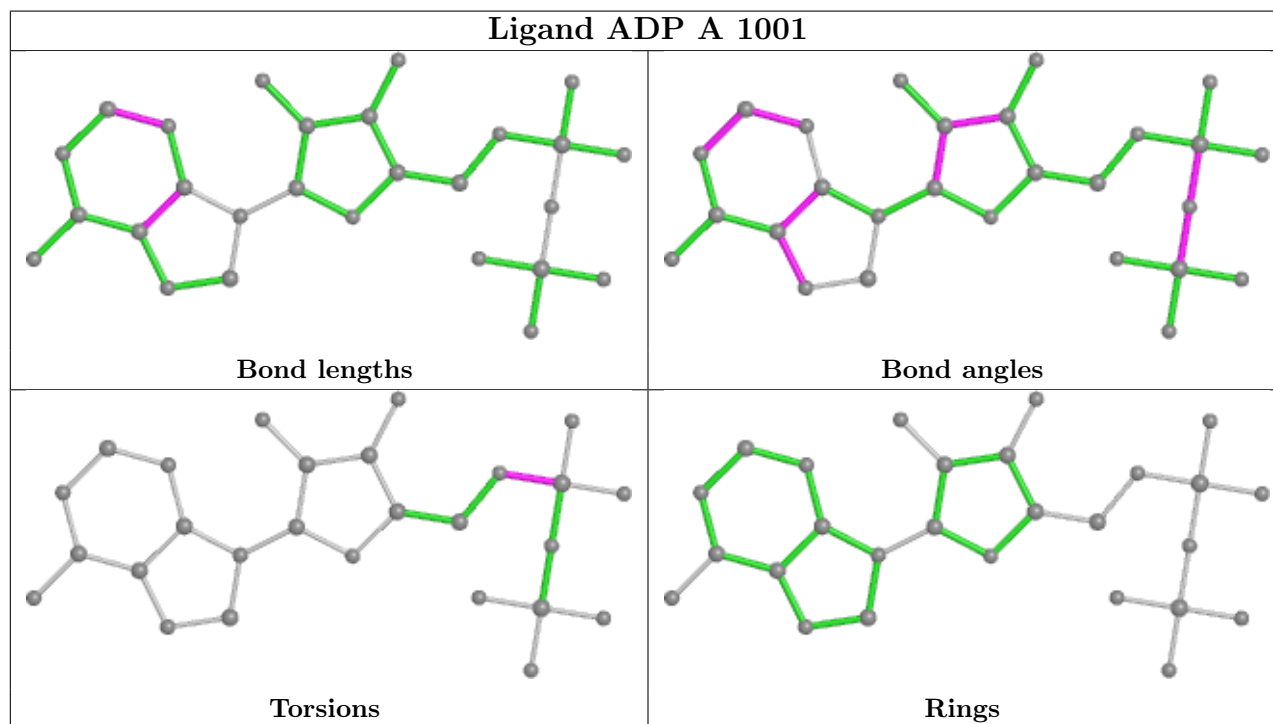
Mol	Chain	Res	Type	Atoms
2	A	1001	ADP	C5'-O5'-PA-O1A
2	B	2001	ADP	C5'-O5'-PA-O3A
2	C	1001	ADP	C5'-O5'-PA-O1A
2	C	1001	ADP	C5'-O5'-PA-O3A
2	D	2001	ADP	C5'-O5'-PA-O3A

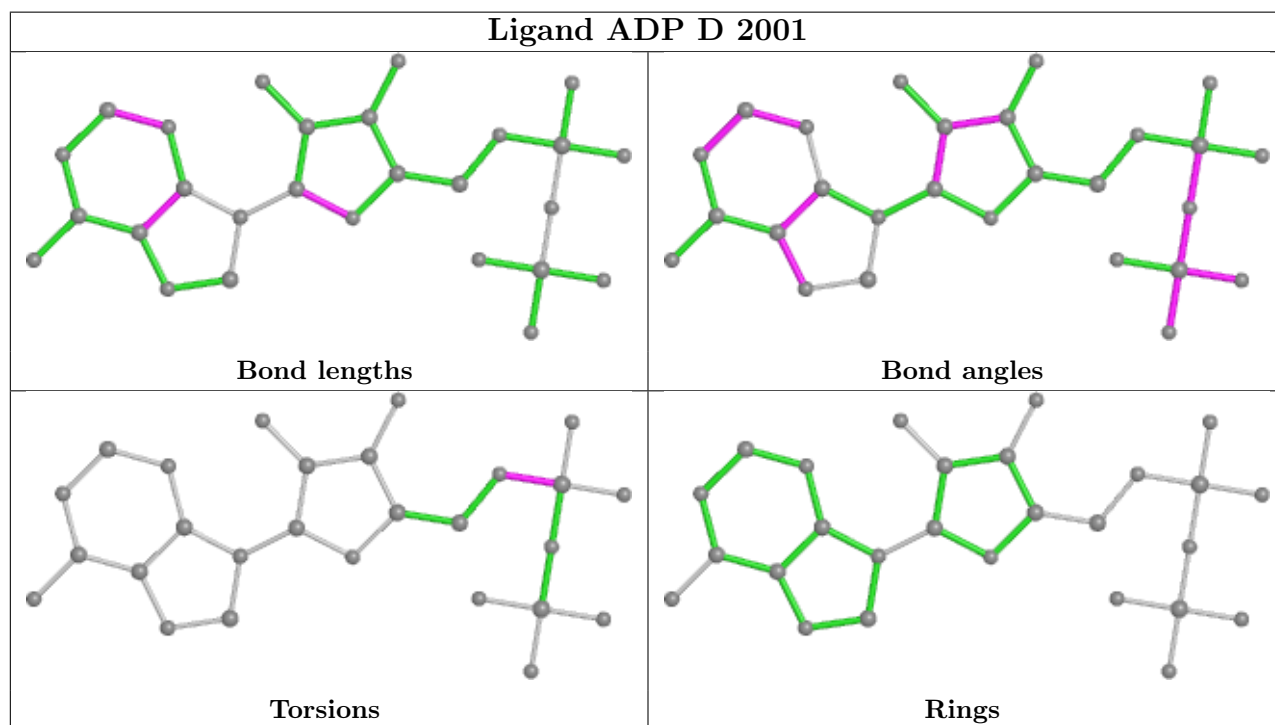
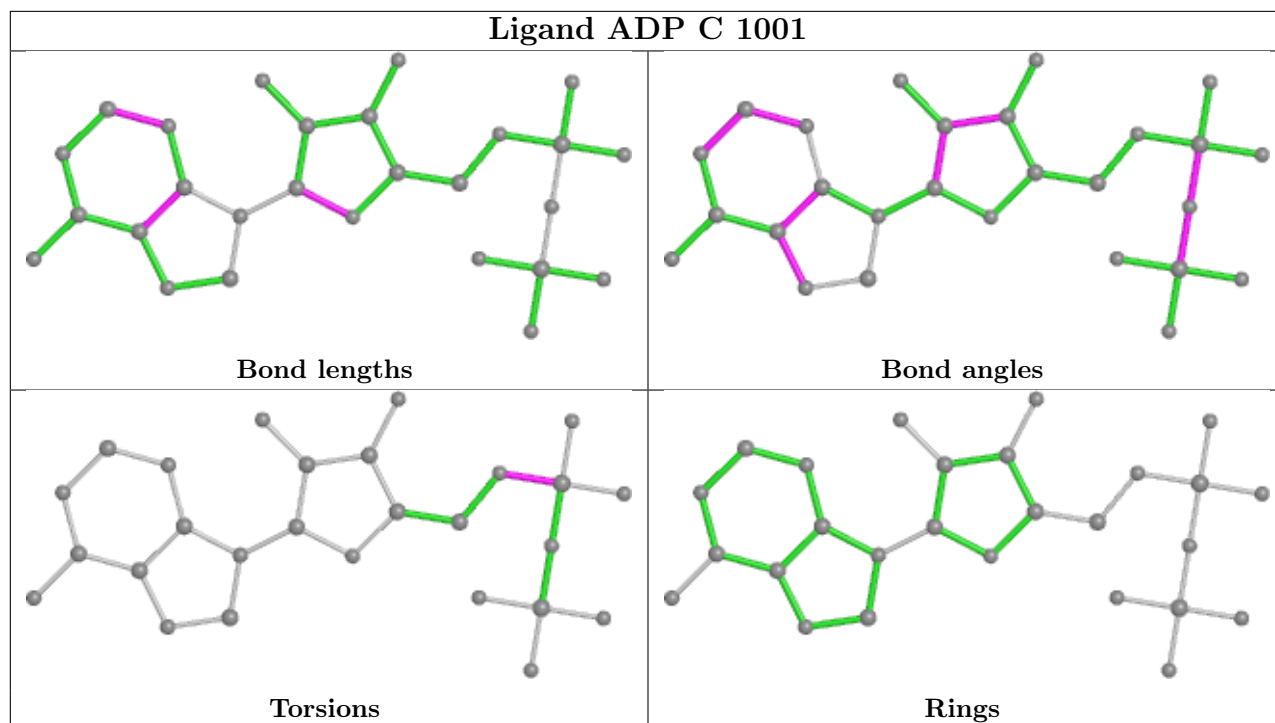
There are no ring outliers.

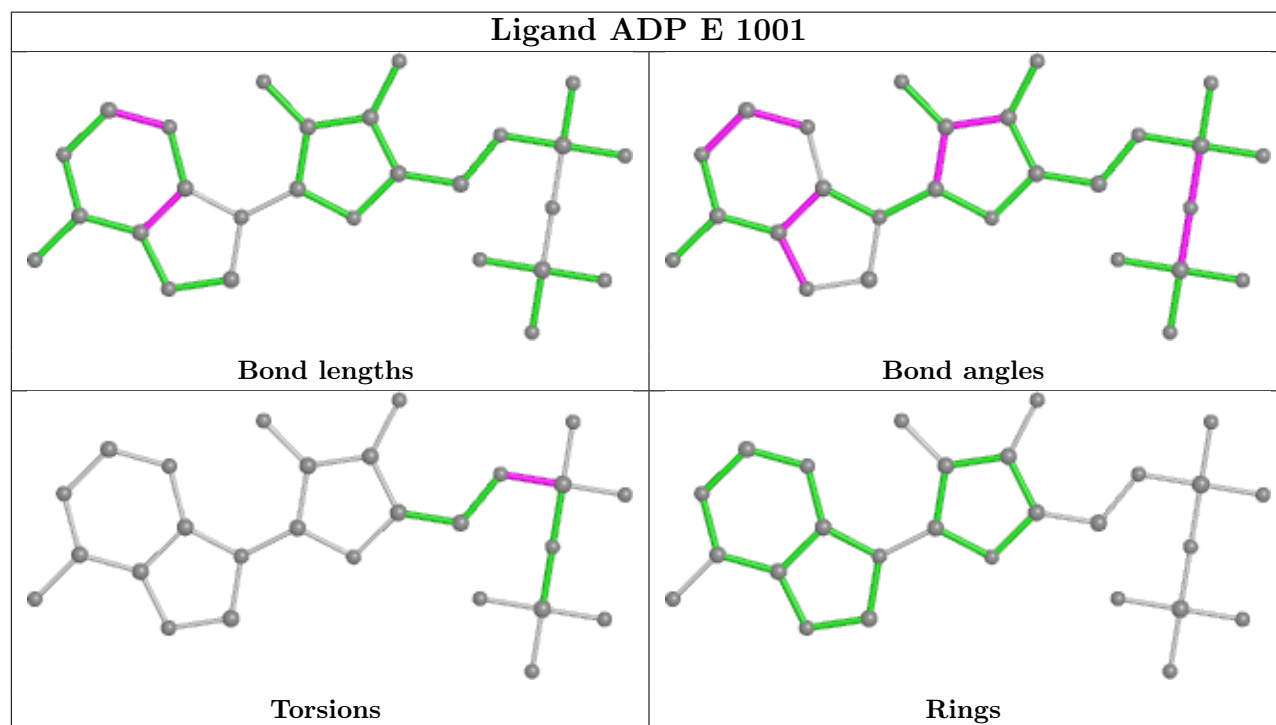
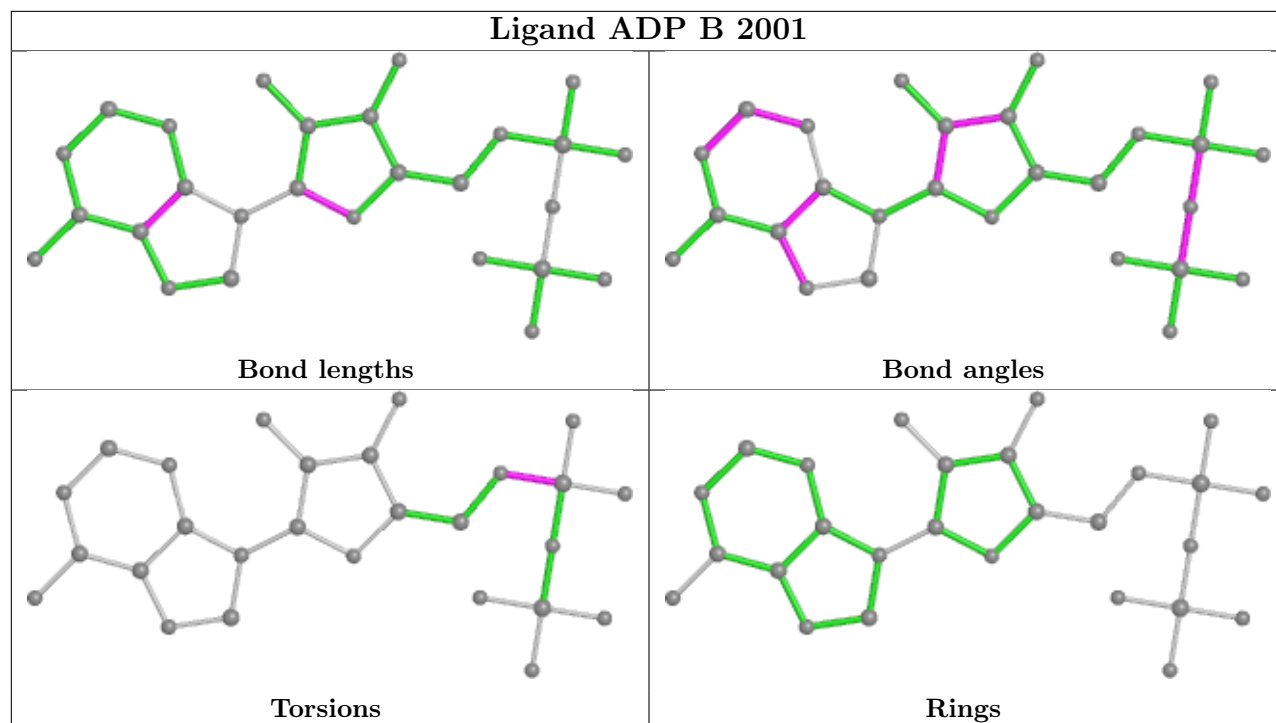
6 monomers are involved in 57 short contacts:

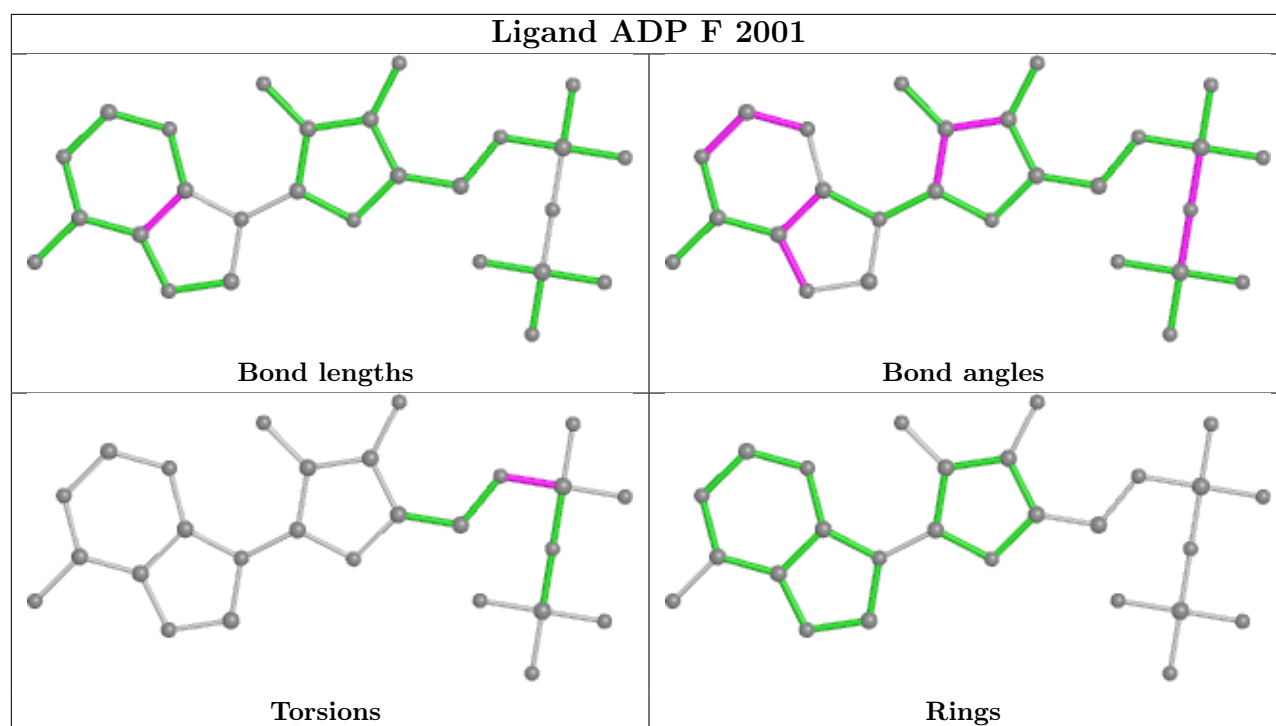
Mol	Chain	Res	Type	Clashes	Symm-Clashes
2	A	1001	ADP	8	0
2	C	1001	ADP	10	0
2	D	2001	ADP	9	0
2	B	2001	ADP	9	0
2	E	1001	ADP	11	0
2	F	2001	ADP	10	0

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 [\(i\)](#)

Unable to reproduce the depositors R factor - this section is therefore empty.

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

Unable to reproduce the depositors R factor - this section is therefore empty.

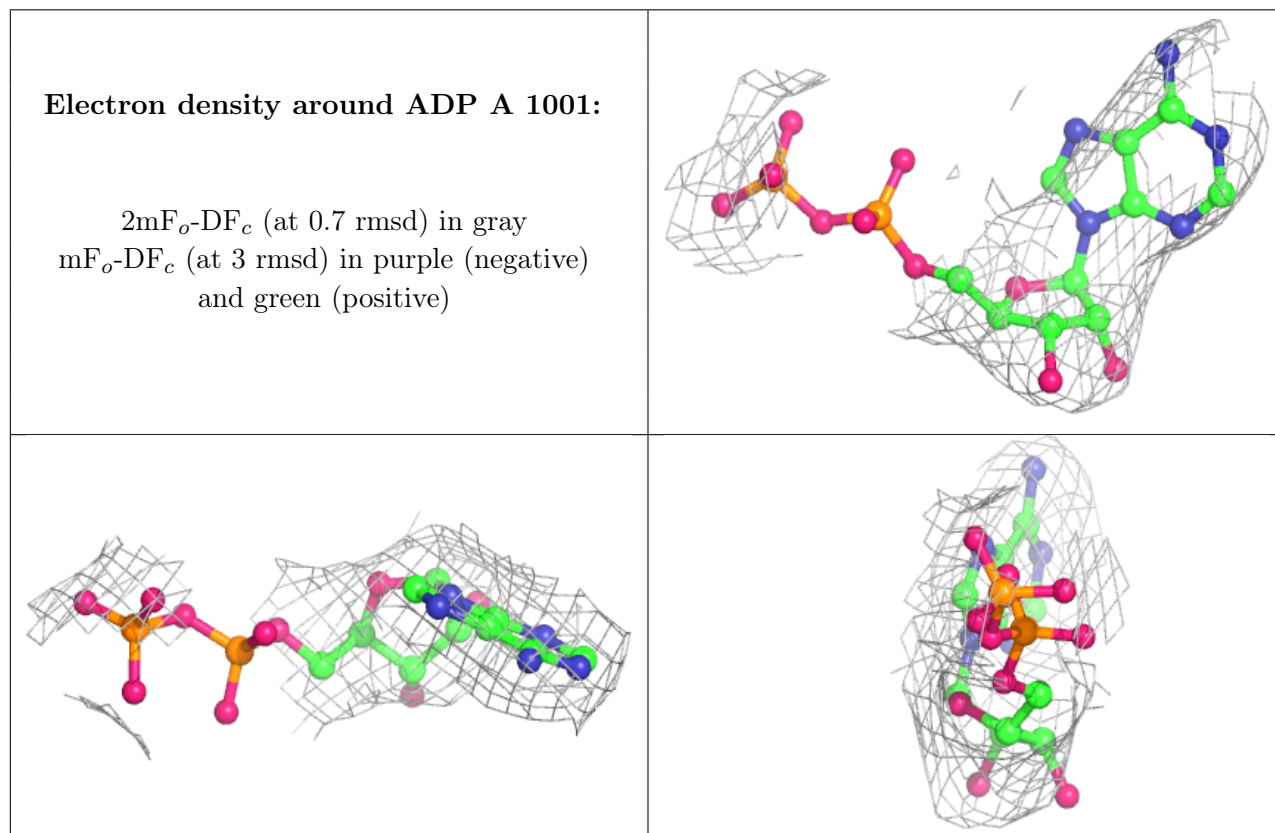
6.3 Carbohydrates [\(i\)](#)

Unable to reproduce the depositors R factor - this section is therefore empty.

6.4 Ligands [\(i\)](#)

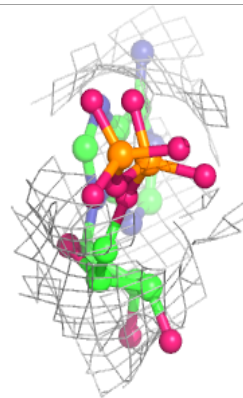
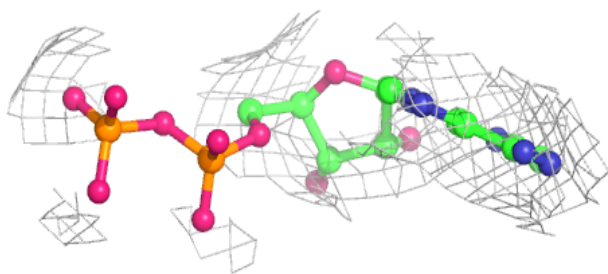
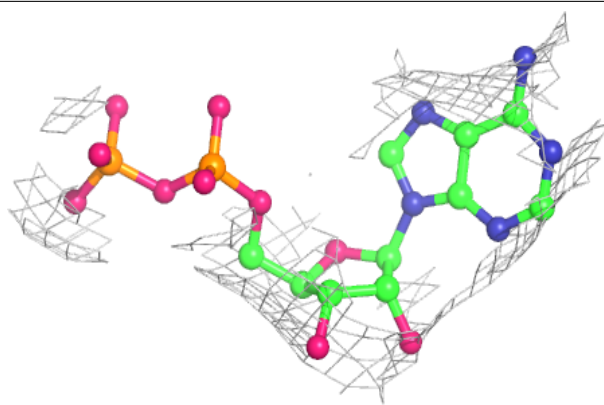
Unable to reproduce the depositors R factor - this section is therefore empty.

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.

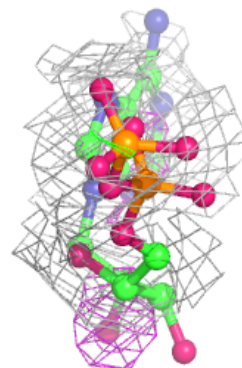
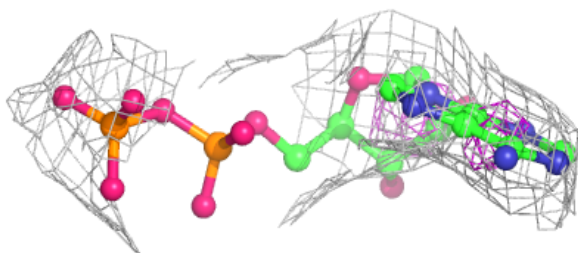
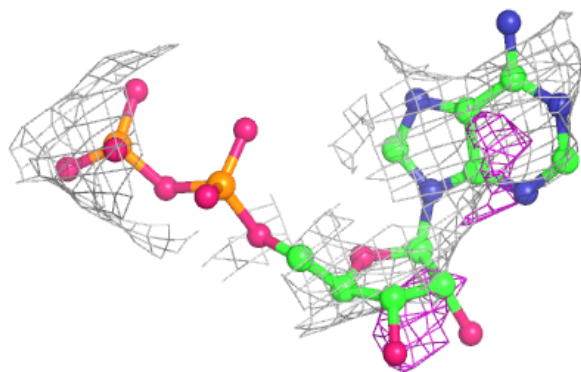


Electron density around ADP B 2001:

$2mF_o-DF_c$ (at 0.7 rmsd) in gray
 mF_o-DF_c (at 3 rmsd) in purple (negative)
and green (positive)

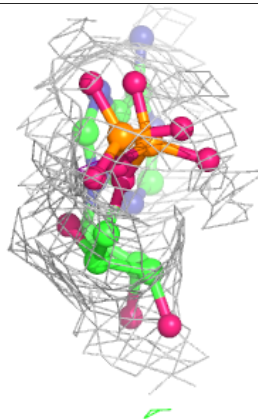
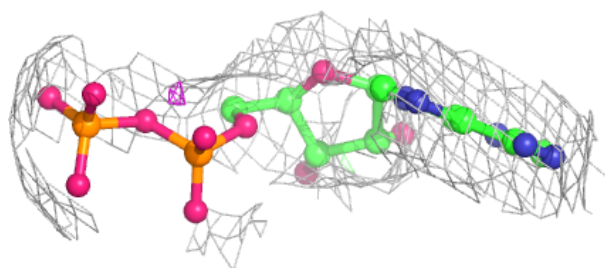
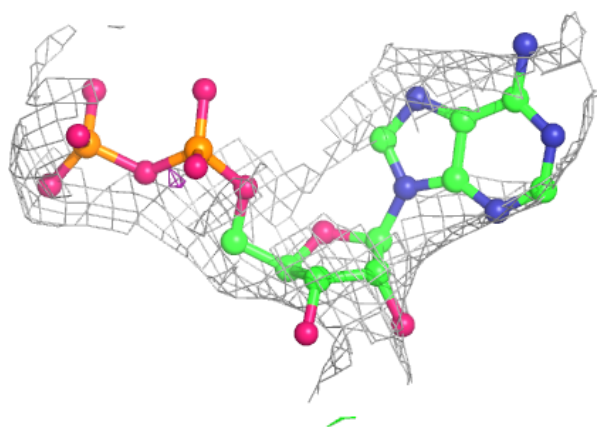
**Electron density around ADP C 1001:**

$2mF_o-DF_c$ (at 0.7 rmsd) in gray
 mF_o-DF_c (at 3 rmsd) in purple (negative)
and green (positive)

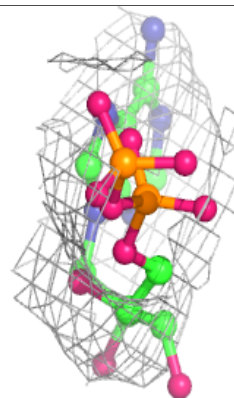
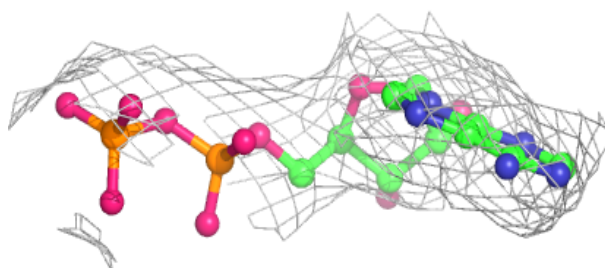
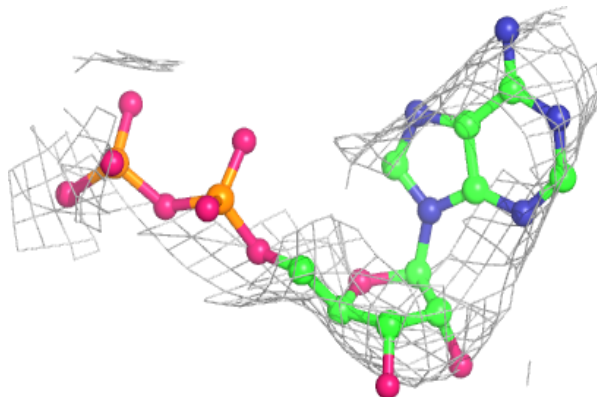


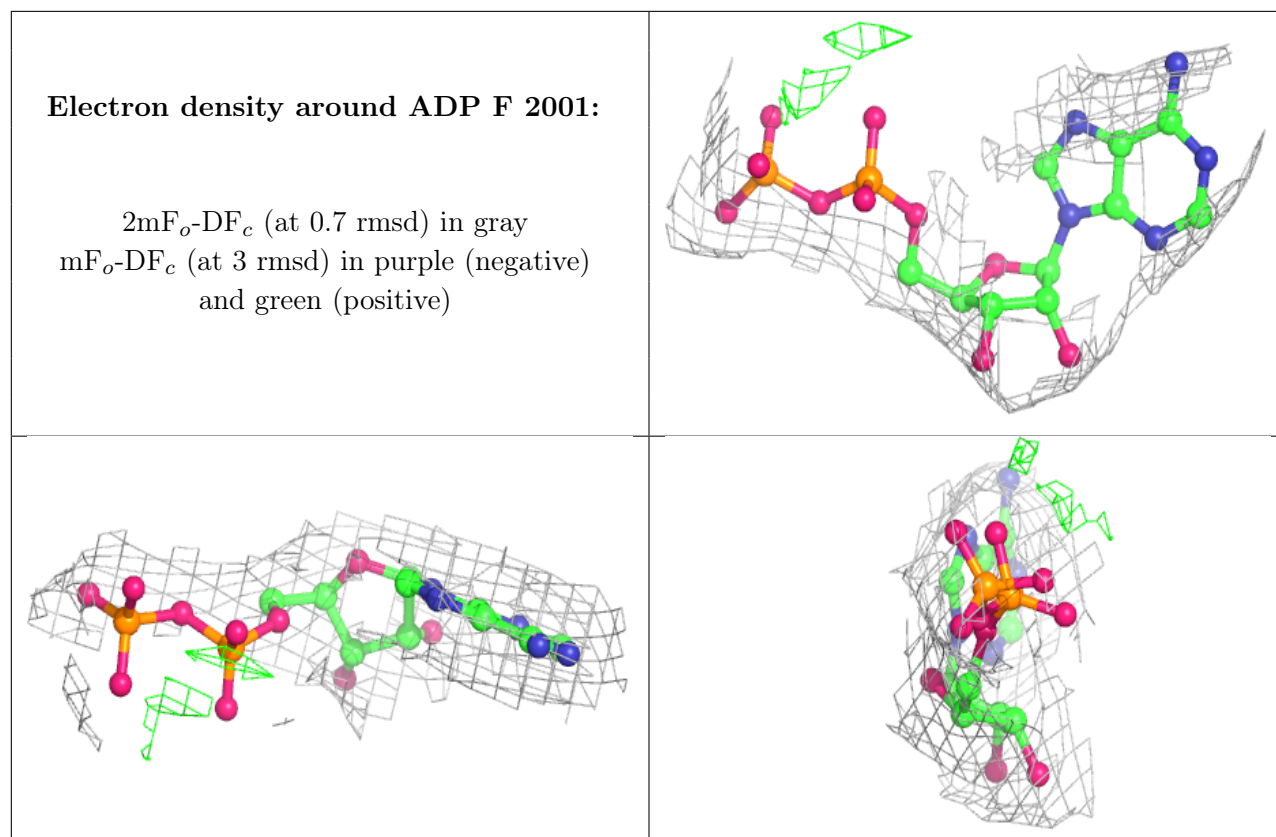
Electron density around ADP D 2001:

$2mF_o-DF_c$ (at 0.7 rmsd) in gray
 mF_o-DF_c (at 3 rmsd) in purple (negative)
and green (positive)

**Electron density around ADP E 1001:**

$2mF_o-DF_c$ (at 0.7 rmsd) in gray
 mF_o-DF_c (at 3 rmsd) in purple (negative)
and green (positive)





6.5 Other polymers [i](#)

Unable to reproduce the depositor's R factor - this section is therefore empty.