



# wwPDB X-ray Structure Validation Summary Report ⓘ

May 16, 2020 – 03:14 pm BST

PDB ID : 3CF2  
Title : Structure of P97/vcp in complex with ADP/AMP-PNP  
Authors : Davies, J.M.; Delabarre, B.; Brunger, A.T.; Weis, W.I.  
Deposited on : 2008-03-01  
Resolution : 3.50 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](mailto: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.

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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 : 2.11  
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) : 2.11

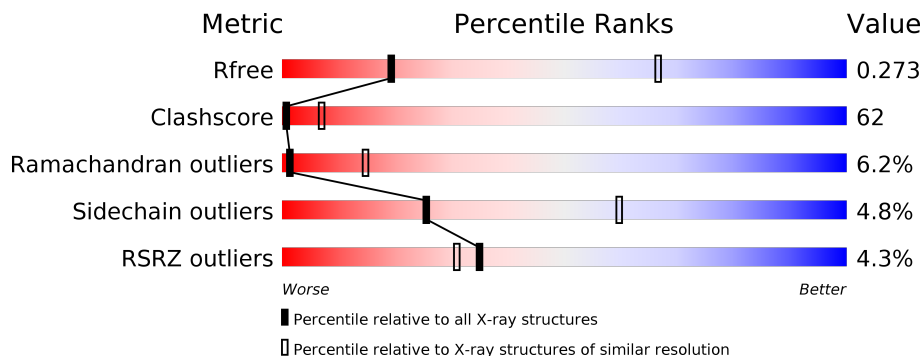
# 1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

*X-RAY DIFFRACTION*

The reported resolution of this entry is 3.50 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	1659 (3.60-3.40)
Clashscore	141614	1036 (3.58-3.42)
Ramachandran outliers	138981	1005 (3.58-3.42)
Sidechain outliers	138945	1006 (3.58-3.42)
RSRZ outliers	127900	1559 (3.60-3.40)

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  $\geq 3$ , 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	806	
1	B	806	
1	C	806	
1	D	806	

## 2 Entry composition [i](#)

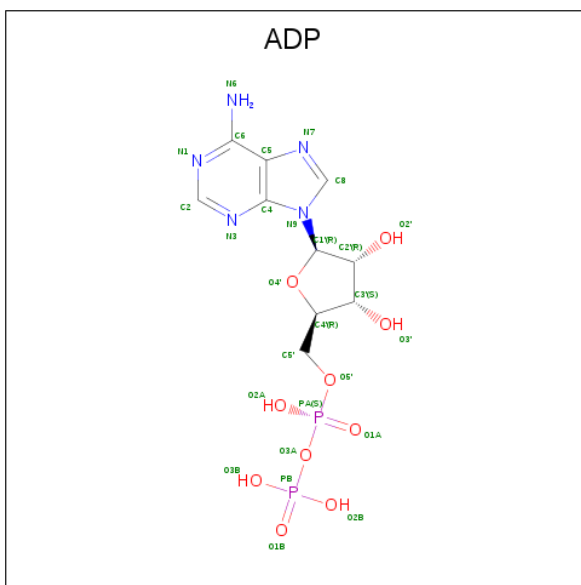
There are 3 unique types of molecules in this entry. The entry contains 20917 atoms, of which 0 are hydrogens and 0 are deuteriums.

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 Transitional endoplasmic reticulum ATPase.

Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
			Total	C	N	O	S			
1	A	659	Total 5172	C 3263	N 903	O 977	S 29	0	0	0
1	B	659	Total 5172	C 3263	N 903	O 977	S 29	0	0	0
1	C	659	Total 5172	C 3263	N 903	O 977	S 29	0	0	0
1	D	659	Total 5169	C 3260	N 903	O 977	S 29	0	0	0

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



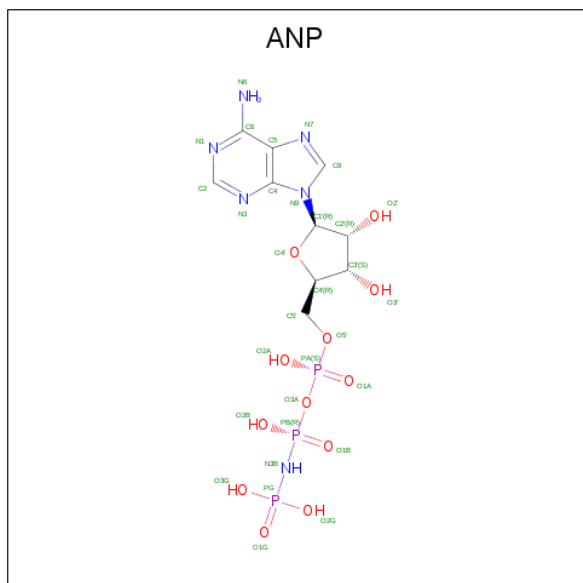
Mol	Chain	Residues	Atoms				ZeroOcc	AltConf	
			Total	C	N	O			P
2	A	1	Total 27	C 10	N 5	O 10	P 2	0	0
2	B	1	Total 27	C 10	N 5	O 10	P 2	0	0

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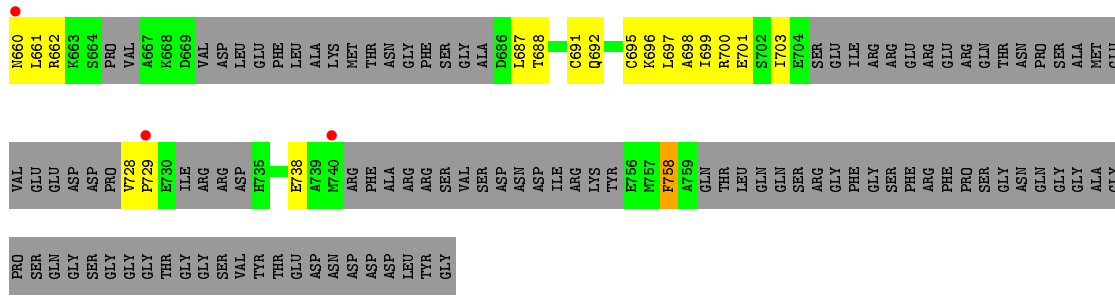
Mol	Chain	Residues	Atoms				ZeroOcc	AltConf	
2	C	1	Total	C	N	O	P	0	0
			27	10	5	10	2		
2	D	1	Total	C	N	O	P	0	0
			27	10	5	10	2		

- Molecule 3 is PHOSPHOAMINOPHOSPHONIC ACID-ADENYLATE ESTER (three-letter code: ANP) (formula:  $C_{10}H_{17}N_6O_{12}P_3$ ).

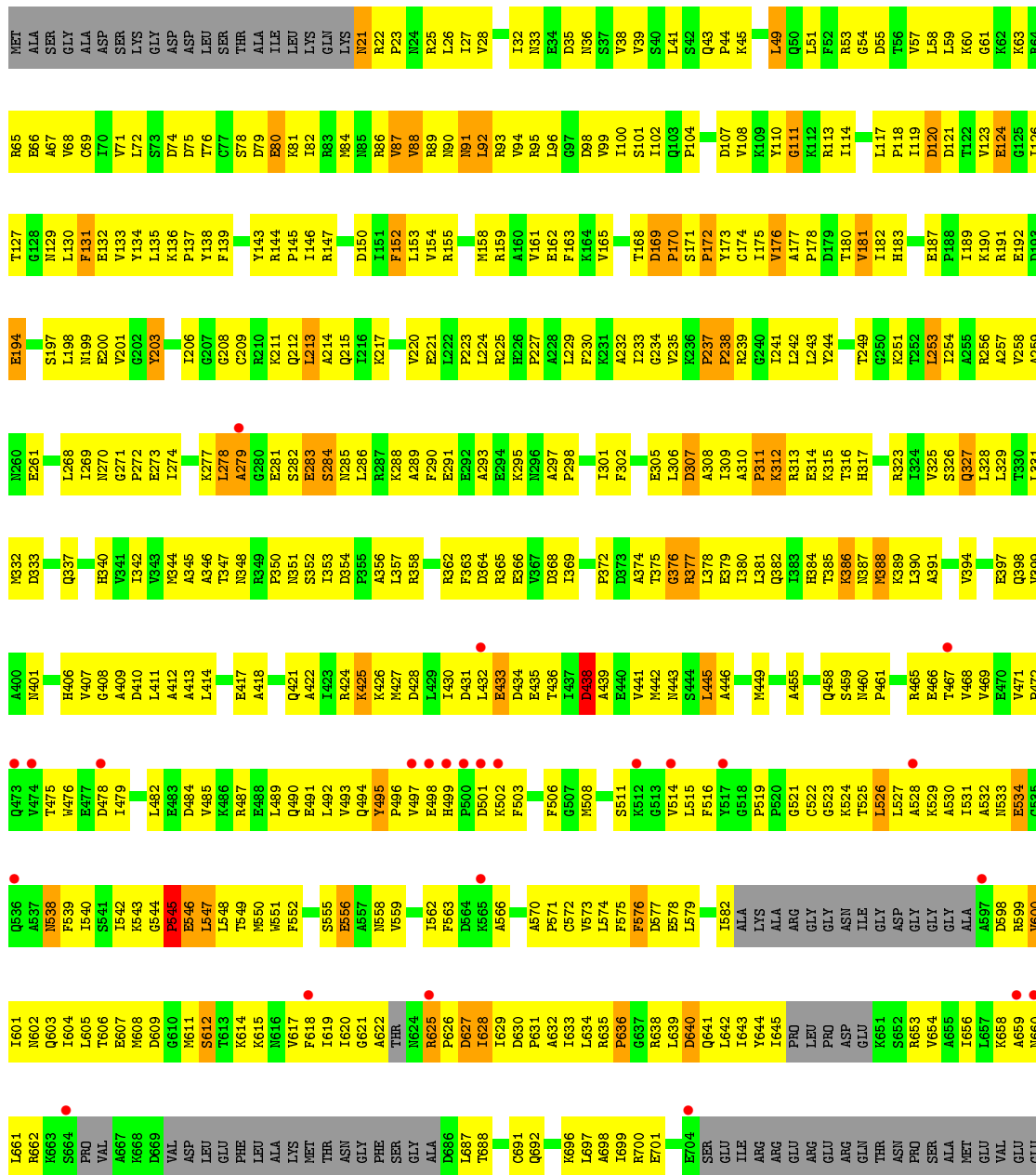


Mol	Chain	Residues	Atoms				ZeroOcc	AltConf	
3	A	1	Total	C	N	O	P	0	0
			31	10	6	12	3		
3	B	1	Total	C	N	O	P	0	0
			31	10	6	12	3		
3	C	1	Total	C	N	O	P	0	0
			31	10	6	12	3		
3	D	1	Total	C	N	O	P	0	0
			31	10	6	12	3		





• Molecule 1: Transitional endoplasmic reticulum ATPase





GLY
GLY
GLY
THR
GLY
GLY
SER
VAL
TYR
THR
GLU
ASP
ASN
ASP
ASP
ASP
ASP
THR
TYR
GLY

● Molecule 1: Transitional endoplasmic reticulum ATPase



MET	R65	T127	E194	R280	T330	Q398	V471	A532	V600	H660	GLU
ALA	E66	G128	S197	E261	L331	V399	P472	M533	I601	L661	GLU
SER	A67	M129	L198	L268	M332	A400	Q473	E534	M602	R662	GLY
GLY	V68	L130	M199	L269	D333	M401	V474	F539	Q603	R663	ASP
GLY	C69	E131	E200	M270	Q337	H406	T475	M538	I604	G664	GLY
ASP	I70	F432	E201	N270	H340	H407	E476	F540	L605	PRD	VAL
SER	V71	R133	V201	C271	V340	V407	E477	S541	T606	A667	VAL
VAL	L72	L135	G202	F272	V341	G408	D478	I542	E607	R668	R6739
LYS	S73	L136	Y203	E273	I342	D410	L479	K543	M608	ARG	ILE
GLY	S74	K136	G204	I274	I343	D411	L480	K544	D609	ARG	ARG
ASP	D74	P137	D205	I275	V344	L411	L481	G545	G610	ARG	ARG
ASP	D75	P137	D205	K277	M344	L412	E482	R545	M611	ASP	ASP
GLU	D76	P137	D205	L278	A345	A413	D484	E546	M612	LEU	LEU
ASP	D77	Y138	G208	A279	A346	L414	V485	L547	T613	ASP	GLY
ASN	C77	F139	C209	G280	T347	L414	V486	L548	R615	GLU	THR
ASP	S78	F139	R210	G280	E417	E418	F486	L549	M615	PHE	THR
ASP	D79	Y143	K211	E281	R347	A418	E488	T549	M616	LEU	THR
ASP	E80	R444	K212	E282	R349	A418	L489	T549	M616	LEU	THR
ASP	K81	P145	Q212	S283	R350	A418	E488	M550	M620	LEU	THR
LYS	I82	I146	L213	E283	M351	A418	E488	M551	M621	LEU	THR
LYS	I82	I146	L213	S284	S352	A418	E488	F552	M622	GLY	THR
GLN	R83	R147	A214	N285	I353	A418	E488	S555	M622	GLY	THR
LYS	M84	R147	Q215	R287	D354	A418	E488	E556	M624	SER	THR
N21	R85	D150	K216	K287	R425	A418	E488	A557	M625	GLY	THR
R22	R86	F152	K217	K288	R426	A418	E488	M558	M626	ALA	THR
P23	V87	L153	K217	F290	A356	A418	E488	V559	M626	ALA	THR
N24	V88	V154	K217	E291	R358	A418	E488	I562	M626	ALA	THR
L26	L89	R155	K217	E292	R362	A418	E488	F563	M627	ALA	THR
R25	R90	R155	K217	E293	F363	A418	E488	P500	M628	ALA	THR
L27	M91	M158	K217	E294	D364	A418	E488	A566	M629	ALA	THR
V28	R92	L158	K217	E295	R365	A418	E488	M570	M630	ALA	THR
I32	R93	A160	K217	E296	R366	A418	E488	F571	M631	ALA	THR
N33	V94	V161	K217	E297	R367	A418	E488	C572	M632	ALA	THR
E34	R95	E162	K217	F298	R368	A418	E488	L574	M633	ALA	THR
D35	G97	F163	K217	F299	R369	A418	E488	M508	M634	ALA	THR
R36	R98	K164	K217	E300	I369	A418	E488	F575	M635	ALA	THR
S37	V99	V165	K217	E301	F372	A418	E488	F576	M636	ALA	THR
V39	I100	S101	K217	E302	D373	A418	E488	D577	M637	ALA	THR
V39	S101	S101	K217	E303	A374	A418	E488	P910	M638	ALA	THR
S40	I102	D169	K217	E304	T375	A418	E488	M442	M639	ALA	THR
L41	Q103	P170	K217	E305	G376	A418	E488	M443	M640	ALA	THR
S42	L41	S171	K217	E306	R377	A418	E488	S444	M641	ALA	THR
Q43	P104	S171	K217	E307	L378	A418	E488	L445	M642	ALA	THR
P44	Q43	P172	K217	E308	R379	A418	E488	A446	M643	ALA	THR
K45	K45	C174	K217	E309	I380	A418	E488	P449	M644	ALA	THR
L49	L49	V175	K217	E310	L381	A418	E488	A455	M645	ALA	THR
Q50	Q50	V176	K217	E311	R382	A418	E488	Q458	M646	ALA	THR
L51	L51	H112	K217	E312	L383	A418	E488	S459	M647	ALA	THR
F52	F52	D179	K217	E313	H384	A418	E488	M460	M648	ALA	THR
B53	B53	I114	K217	E314	T385	A418	E488	P461	M649	ALA	THR
G54	G54	H181	K217	E315	K386	A418	E488	Q465	M650	ALA	THR
D55	D55	H182	K217	E316	R387	A418	E488	S469	M651	ALA	THR
T56	T56	H183	K217	E317	M388	A418	E488	M460	M652	ALA	THR
V57	V57	E187	K217	E318	R389	A418	E488	P461	M653	ALA	THR
L58	L58	I254	K217	E319	L390	A418	E488	R465	M654	ALA	THR
L59	L59	I189	K217	E320	I391	A418	E488	T467	M655	ALA	THR
K60	K60	T122	K217	E321	A391	A418	E488	A528	M656	ALA	THR
G61	G61	V123	K217	E322	V394	A418	E488	R529	M657	ALA	THR
R62	R62	E124	K217	E323	E397	A418	E488	V468	M658	ALA	THR
K63	K63	G125	K217	E324	L329	A418	E488	V469	M659	ALA	THR
R64	R64	I126	K217	E325	E397	A418	E488	E470	M660	ALA	THR



## 4 Data and refinement statistics

Property	Value	Source
Space group	P 3	Depositor
Cell constants a, b, c, $\alpha$ , $\beta$ , $\gamma$	144.90Å 144.90Å 164.40Å 90.00° 90.00° 120.00°	Depositor
Resolution (Å)	23.00 – 3.50 22.79 – 3.20	Depositor EDS
% Data completeness (in resolution range)	99.6 (23.00-3.50) 90.8 (22.79-3.20)	Depositor EDS
$R_{merge}$	0.12	Depositor
$R_{sym}$	(Not available)	Depositor
$\langle I/\sigma(I) \rangle$ <sup>1</sup>	1.22 (at 3.23Å)	Xtriage
Refinement program	CNS	Depositor
R, $R_{free}$	0.271 , 0.285 0.259 , 0.273	Depositor DCC
$R_{free}$ test set	6358 reflections (5.03%)	wwPDB-VP
Wilson B-factor (Å <sup>2</sup> )	75.9	Xtriage
Anisotropy	0.841	Xtriage
Bulk solvent $k_{sol}$ (e/Å <sup>3</sup> ), $B_{sol}$ (Å <sup>2</sup> )	0.28 , 72.8	EDS
L-test for twinning <sup>2</sup>	$\langle  L  \rangle = 0.44$ , $\langle L^2 \rangle = 0.27$	Xtriage
Estimated twinning fraction	0.369 for -h,-k,l 0.377 for h,-h-k,-l 0.369 for -k,-h,-l	Xtriage
$F_o, F_c$ correlation	0.89	EDS
Total number of atoms	20917	wwPDB-VP
Average B, all atoms (Å <sup>2</sup> )	148.0	wwPDB-VP

Xtriage's analysis on translational NCS is as follows: *The largest off-origin peak in the Patterson function is 6.63% of the height of the origin peak. No significant pseudotranslation is detected.*

<sup>1</sup>Intensities estimated from amplitudes.

<sup>2</sup>Theoretical 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: ANP, 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	0.48	2/5250 (0.0%)	0.67	1/7082 (0.0%)
1	B	0.48	2/5250 (0.0%)	0.67	1/7082 (0.0%)
1	C	0.48	2/5250 (0.0%)	0.67	1/7082 (0.0%)
1	D	0.48	2/5247 (0.0%)	0.67	1/7078 (0.0%)
All	All	0.48	8/20997 (0.0%)	0.67	4/28324 (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	2
1	B	0	2
1	C	0	2
1	D	0	2
All	All	0	8

The worst 5 of 8 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
1	B	425	LYS	C-N	-12.18	1.06	1.34
1	C	425	LYS	C-N	-12.14	1.06	1.34
1	D	425	LYS	C-N	-12.14	1.06	1.34
1	A	425	LYS	C-N	-12.14	1.06	1.34
1	B	438	ASP	C-N	-7.39	1.17	1.34

All (4) bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed( $^{\circ}$ )	Ideal( $^{\circ}$ )
1	D	538	ASN	O-C-N	5.67	131.78	122.70
1	B	538	ASN	O-C-N	5.66	131.75	122.70
1	C	538	ASN	O-C-N	5.65	131.74	122.70
1	A	538	ASN	O-C-N	5.64	131.72	122.70

There are no chirality outliers.

5 of 8 planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	A	433	GLU	Peptide
1	A	545	PRO	Peptide
1	B	433	GLU	Peptide
1	B	545	PRO	Peptide
1	C	433	GLU	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	5172	0	5238	723	39
1	B	5172	0	5236	728	39
1	C	5172	0	5238	730	38
1	D	5169	0	5226	719	38
2	A	27	0	12	3	0
2	B	27	0	12	3	0
2	C	27	0	12	3	0
2	D	27	0	12	3	0
3	A	31	0	13	1	0
3	B	31	0	13	1	0
3	C	31	0	13	1	0
3	D	31	0	13	1	0
All	All	20917	0	21038	2616	77

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 62.

The worst 5 of 2616 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:A:699:ILE:HG12	1:B:506:PHE:CG	1.22	1.67
1:C:699:ILE:HG12	1:D:506:PHE:CG	1.29	1.66
1:A:549:THR:CB	1:B:602:ASN:HD22	1.13	1.57
1:C:549:THR:CB	1:D:602:ASN:HD22	1.14	1.57
1:A:550:MET:HG3	1:B:606:THR:CB	1.41	1.50

The worst 5 of 77 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:C:491:GLU:OE1	1:D:700:ARG:CD[2_765]	0.38	1.82
1:A:491:GLU:OE2	1:B:700:ARG:NH1[3_765]	0.38	1.82
1:C:491:GLU:OE2	1:D:700:ARG:NH1[2_765]	0.40	1.80
1:A:491:GLU:OE1	1:B:700:ARG:CD[3_765]	0.41	1.79
1:C:25:ARG:NH2	1:D:432:LEU:C[2_765]	0.80	1.40

## 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	641/806 (80%)	471 (74%)	130 (20%)	40 (6%)	1	15
1	B	641/806 (80%)	470 (73%)	131 (20%)	40 (6%)	1	15
1	C	641/806 (80%)	471 (74%)	130 (20%)	40 (6%)	1	15
1	D	641/806 (80%)	470 (73%)	131 (20%)	40 (6%)	1	15
All	All	2564/3224 (80%)	1882 (73%)	522 (20%)	160 (6%)	1	15

5 of 160 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	A	87	VAL
1	A	312	LYS

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Mol	Chain	Res	Type
1	B	87	VAL
1	B	312	LYS
1	C	87	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	565/678 (83%)	538 (95%)	27 (5%)	25	60
1	B	565/678 (83%)	538 (95%)	27 (5%)	25	60
1	C	565/678 (83%)	538 (95%)	27 (5%)	25	60
1	D	564/678 (83%)	537 (95%)	27 (5%)	25	60
All	All	2259/2712 (83%)	2151 (95%)	108 (5%)	25	60

5 of 108 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	B	556	GLU
1	C	152	PHE
1	D	534	GLU
1	B	625	ARG
1	C	21	ASN

Some sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 62 such sidechains are listed below:

Mol	Chain	Res	Type
1	B	533	ASN
1	C	212	GLN
1	D	406	HIS
1	C	91	ASN
1	C	285	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 carbohydrates in this entry.

## 5.6 Ligand geometry [i](#)

8 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	807	-	24,29,29	1.46	4 (16%)	29,45,45	1.66	3 (10%)
2	ADP	C	807	-	24,29,29	1.46	4 (16%)	29,45,45	1.66	3 (10%)
2	ADP	B	807	-	24,29,29	1.46	5 (20%)	29,45,45	1.65	3 (10%)
3	ANP	A	901	-	29,33,33	2.07	11 (37%)	31,52,52	1.71	5 (16%)
3	ANP	C	901	-	29,33,33	2.08	11 (37%)	31,52,52	1.71	5 (16%)
3	ANP	D	901	-	29,33,33	2.08	11 (37%)	31,52,52	1.71	5 (16%)
3	ANP	B	901	-	29,33,33	2.07	11 (37%)	31,52,52	1.71	5 (16%)
2	ADP	D	807	-	24,29,29	1.47	5 (20%)	29,45,45	1.66	3 (10%)

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	807	-	-	6/12/32/32	0/3/3/3
2	ADP	C	807	-	-	6/12/32/32	0/3/3/3
2	ADP	B	807	-	-	6/12/32/32	0/3/3/3
3	ANP	A	901	-	-	6/14/38/38	0/3/3/3
3	ANP	C	901	-	-	6/14/38/38	0/3/3/3
3	ANP	D	901	-	-	6/14/38/38	0/3/3/3
3	ANP	B	901	-	-	6/14/38/38	0/3/3/3
2	ADP	D	807	-	-	6/12/32/32	0/3/3/3

The worst 5 of 62 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
3	B	901	ANP	PB-N3B	-4.48	1.51	1.63
3	C	901	ANP	PB-N3B	-4.47	1.51	1.63
3	A	901	ANP	PB-N3B	-4.46	1.51	1.63
3	D	901	ANP	PB-N3B	-4.45	1.51	1.63
3	B	901	ANP	PG-N3B	-4.28	1.52	1.63

The worst 5 of 32 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
2	C	807	ADP	N3-C2-N1	-6.79	118.06	128.68
2	A	807	ADP	N3-C2-N1	-6.79	118.07	128.68
2	D	807	ADP	N3-C2-N1	-6.78	118.08	128.68
2	B	807	ADP	N3-C2-N1	-6.76	118.11	128.68
3	B	901	ANP	N3-C2-N1	-6.74	118.14	128.68

There are no chirality outliers.

5 of 48 torsion outliers are listed below:

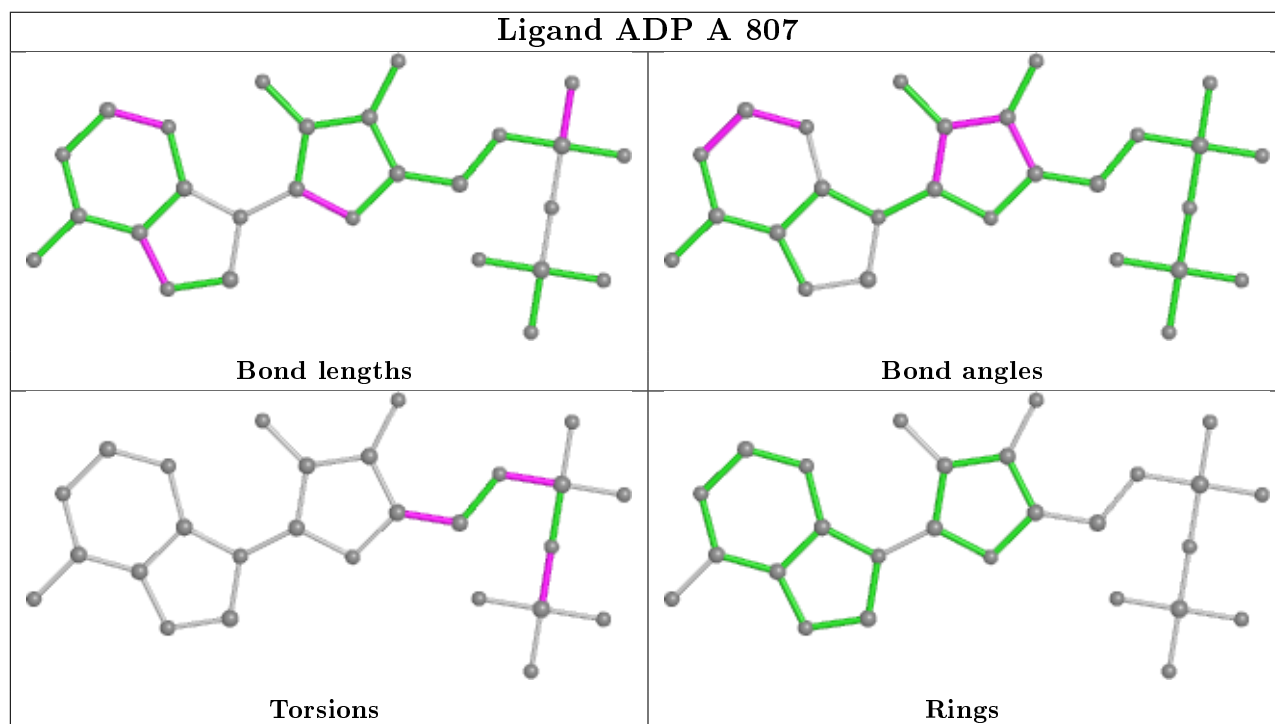
Mol	Chain	Res	Type	Atoms
2	A	807	ADP	C5'-O5'-PA-O1A
2	A	807	ADP	C5'-O5'-PA-O2A
2	A	807	ADP	C5'-O5'-PA-O3A
2	A	807	ADP	O4'-C4'-C5'-O5'
2	C	807	ADP	C5'-O5'-PA-O1A

There are no ring outliers.

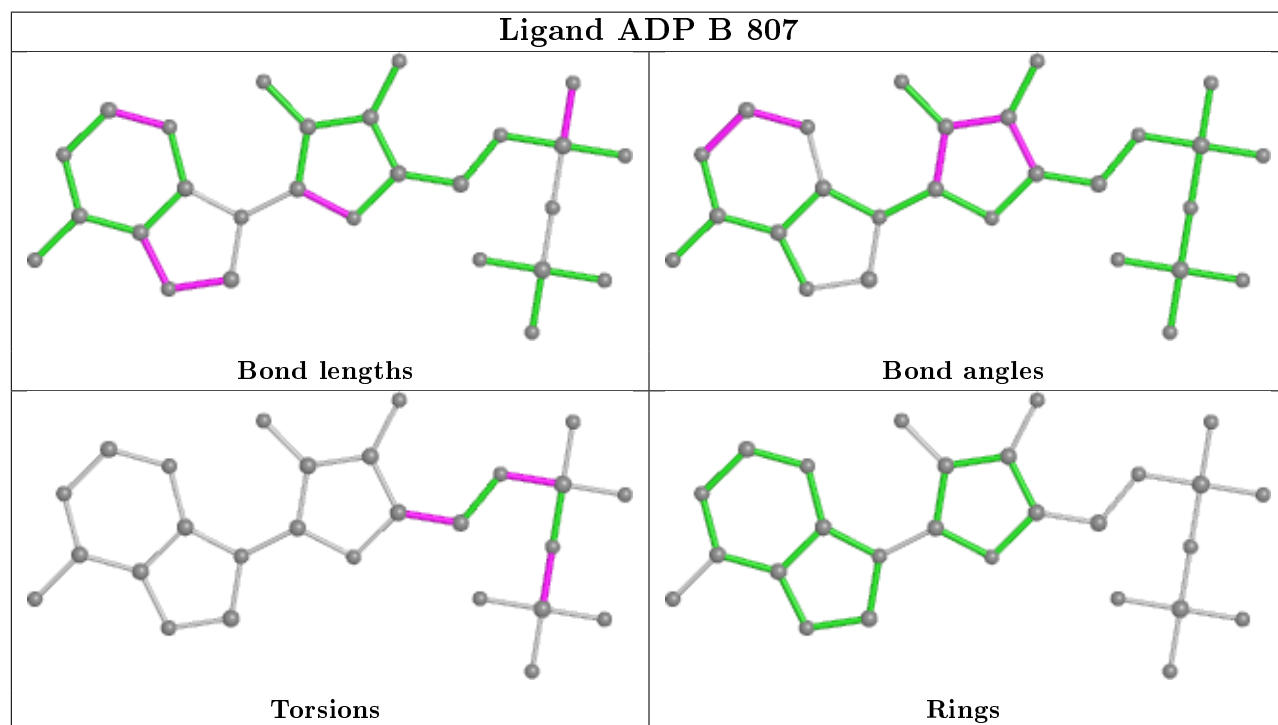
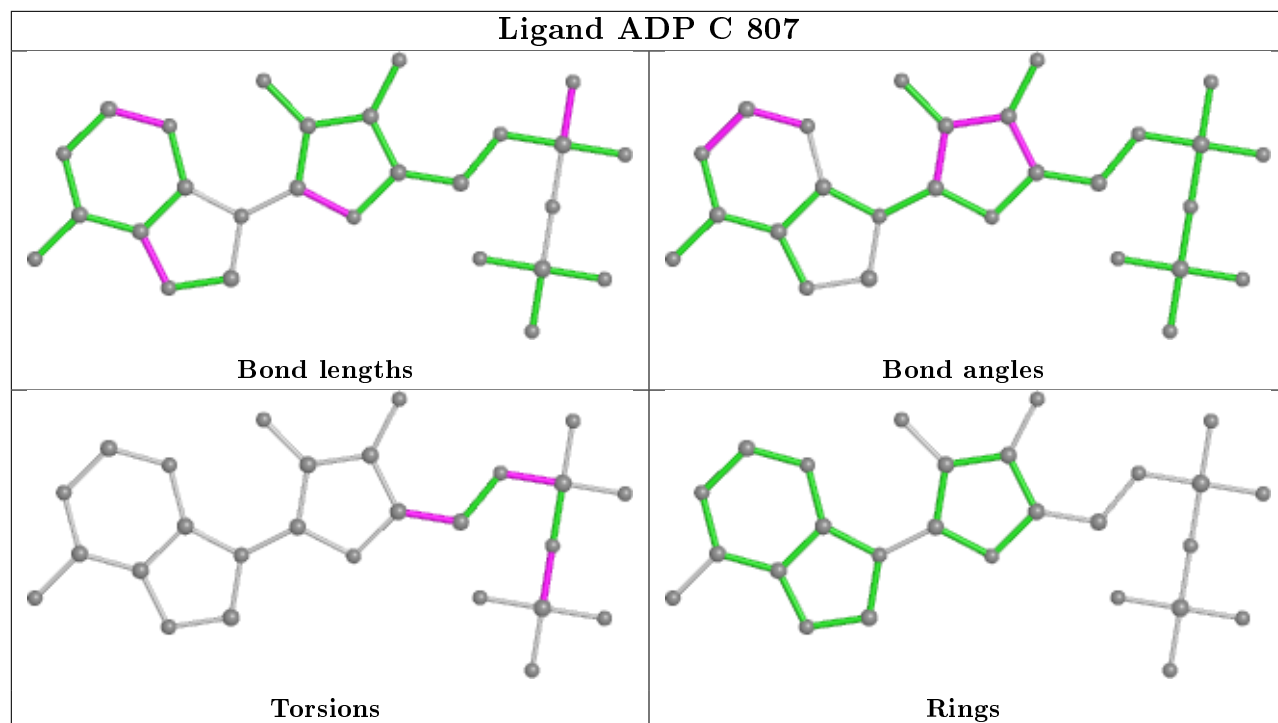
8 monomers are involved in 16 short contacts:

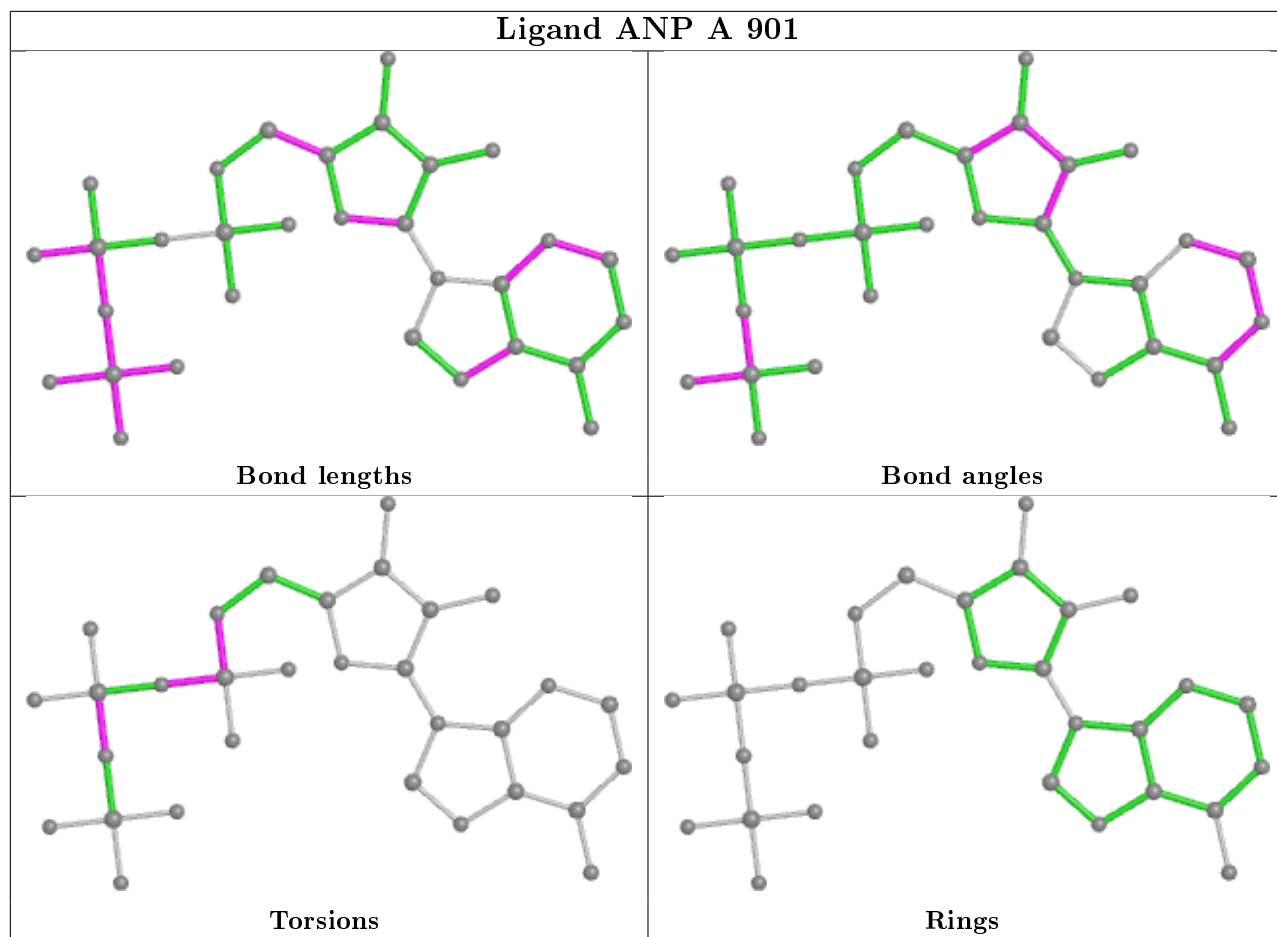
Mol	Chain	Res	Type	Clashes	Symm-Clashes
2	A	807	ADP	3	0
2	C	807	ADP	3	0
2	B	807	ADP	3	0
3	A	901	ANP	1	0
3	C	901	ANP	1	0
3	D	901	ANP	1	0
3	B	901	ANP	1	0
2	D	807	ADP	3	0

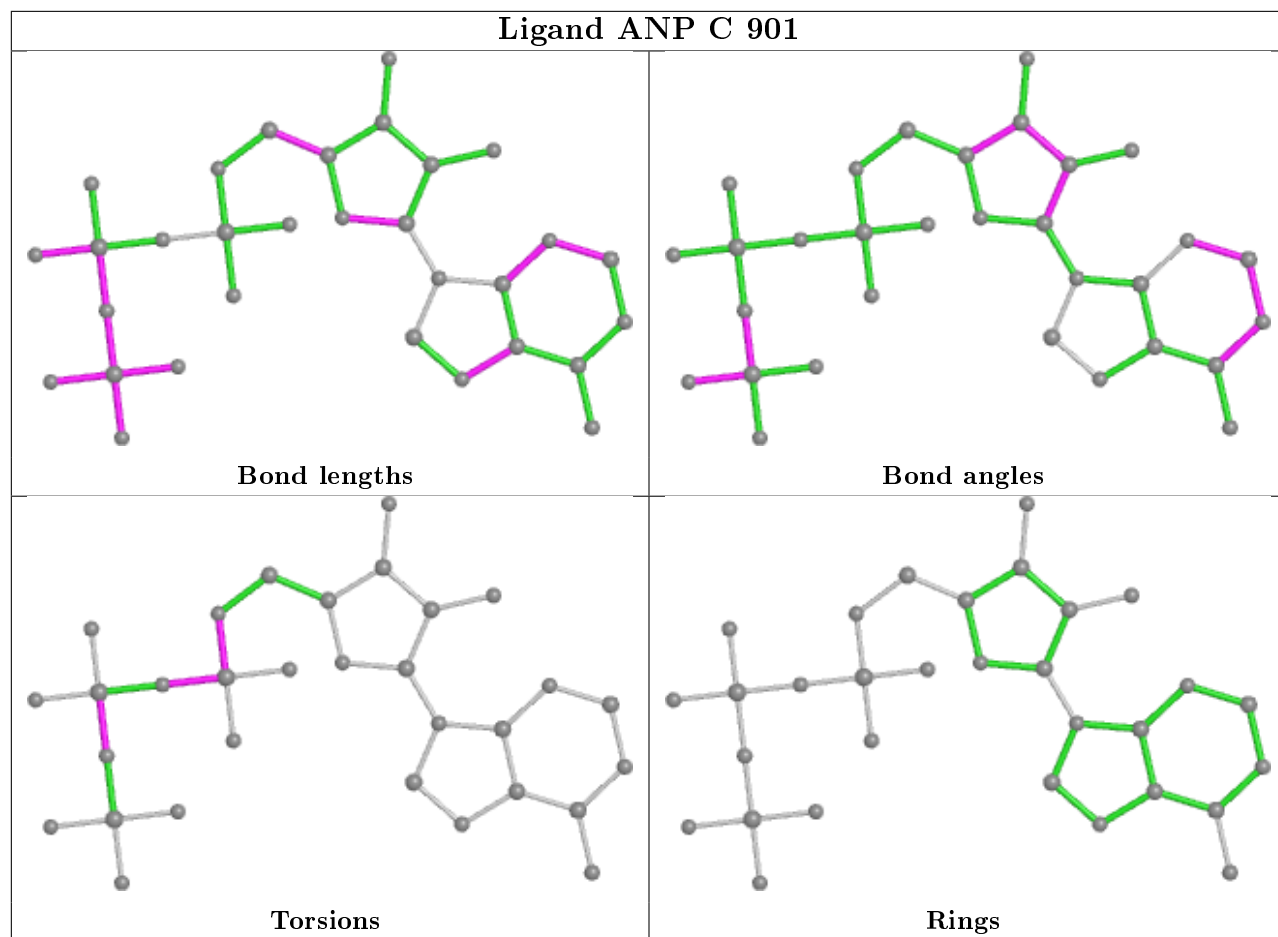
The following is a two-dimensional graphical depiction of Mogul quality analysis of bond lengths, bond angles, torsion angles, and ring geometry for all instances of the Ligand of Interest. In addition, ligands with molecular weight > 250 and outliers as shown on the validation Tables will also be included. 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.

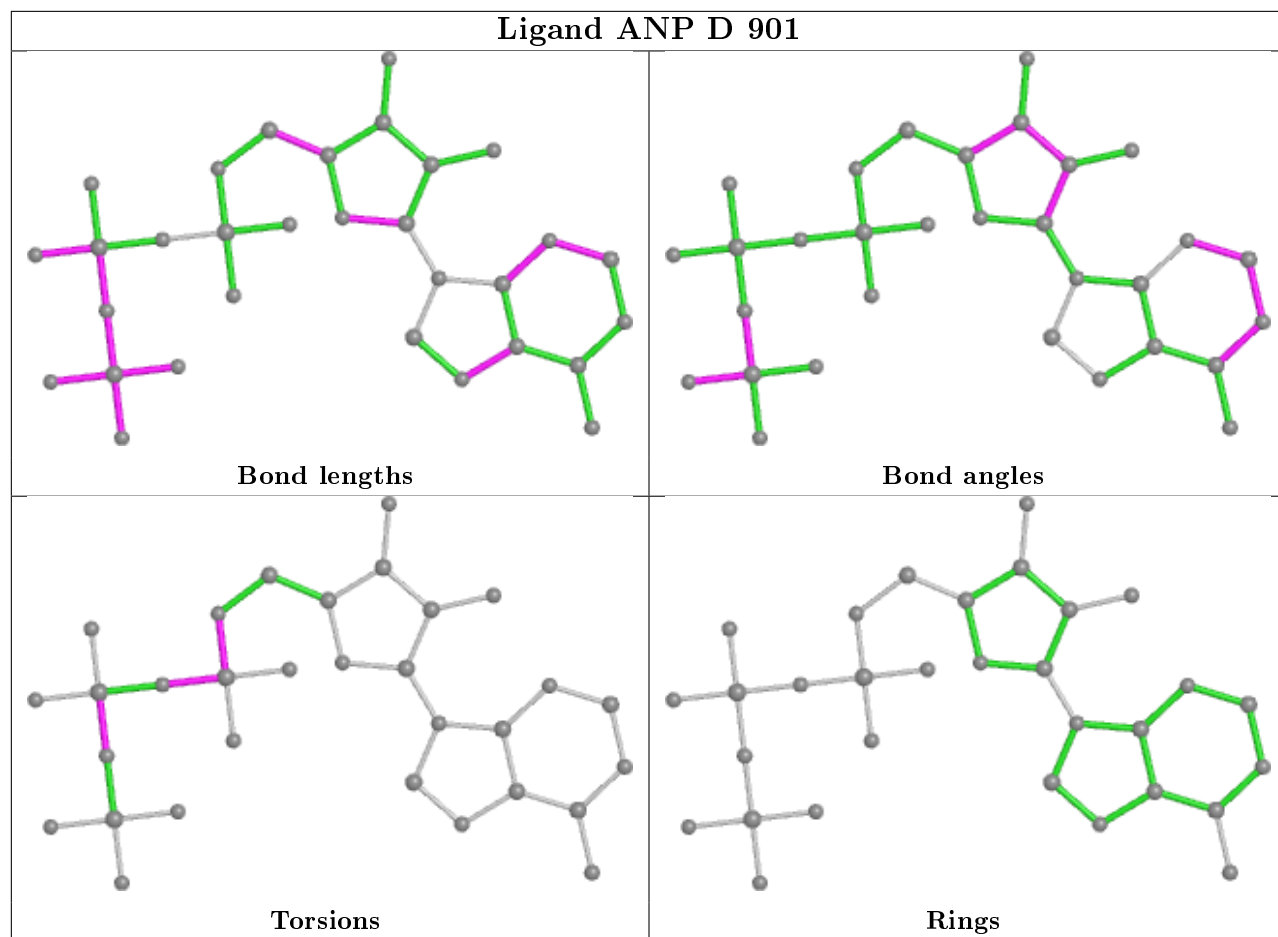


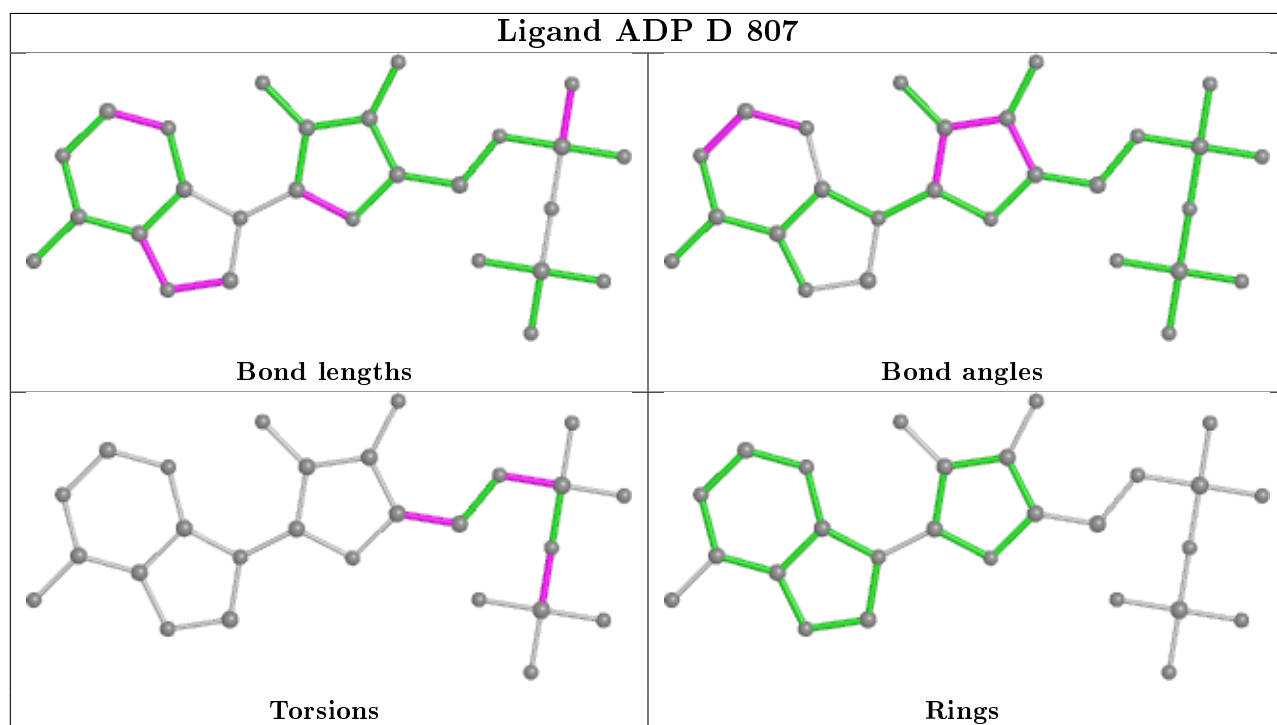
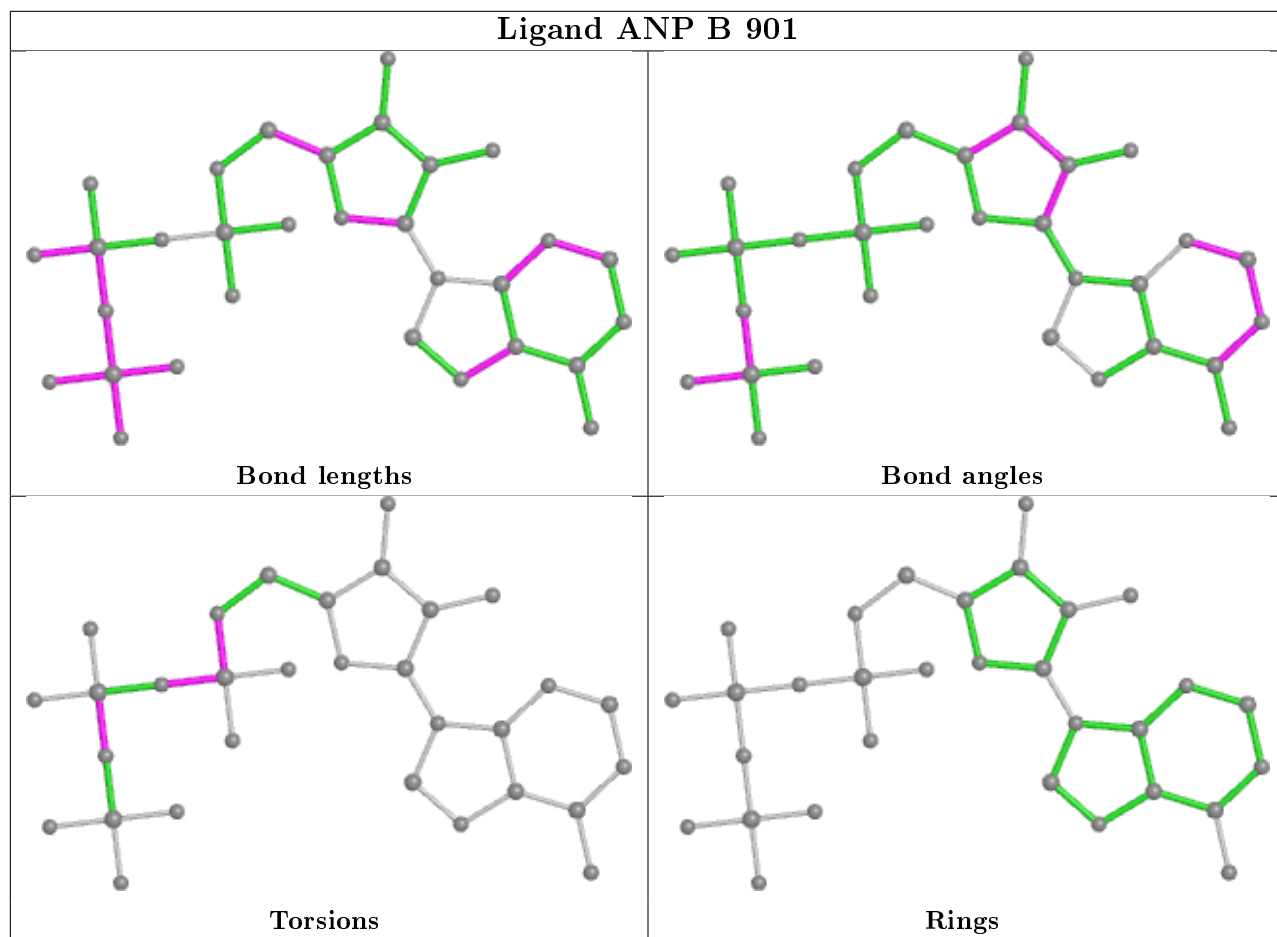












## 5.7 Other polymers [i](#)

There are no such residues in this entry.

## 5.8 Polymer linkage issues [i](#)

The following chains have linkage breaks:

Mol	Chain	Number of breaks
1	B	2
1	A	2
1	D	2
1	C	2

The worst 5 of 8 chain breaks are listed below:

Model	Chain	Residue-1	Atom-1	Residue-2	Atom-2	Distance (Å)
1	A	438:ASP	C	439:ALA	N	1.17
1	B	438:ASP	C	439:ALA	N	1.17
1	C	438:ASP	C	439:ALA	N	1.17
1	D	438:ASP	C	439:ALA	N	1.17
1	A	425:LYS	C	426:LYS	N	1.06

## 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, 95<sup>th</sup> 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 <sup>2</sup> )	Q<0.9
1	A	659/806 (81%)	0.06	28 (4%) 36 32	17, 128, 281, 412	0
1	B	659/806 (81%)	0.10	27 (4%) 37 33	17, 128, 281, 412	0
1	C	659/806 (81%)	0.04	25 (3%) 40 36	17, 128, 281, 412	0
1	D	659/806 (81%)	0.06	33 (5%) 28 25	17, 128, 281, 412	0
All	All	2636/3224 (81%)	0.06	113 (4%) 35 31	17, 128, 281, 412	0

The worst 5 of 113 RSRZ outliers are listed below:

Mol	Chain	Res	Type	RSRZ
1	B	660	ASN	7.8
1	B	498	GLU	6.2
1	A	498	GLU	6.0
1	D	618	PHE	5.8
1	B	478	ASP	5.6

### 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](#)

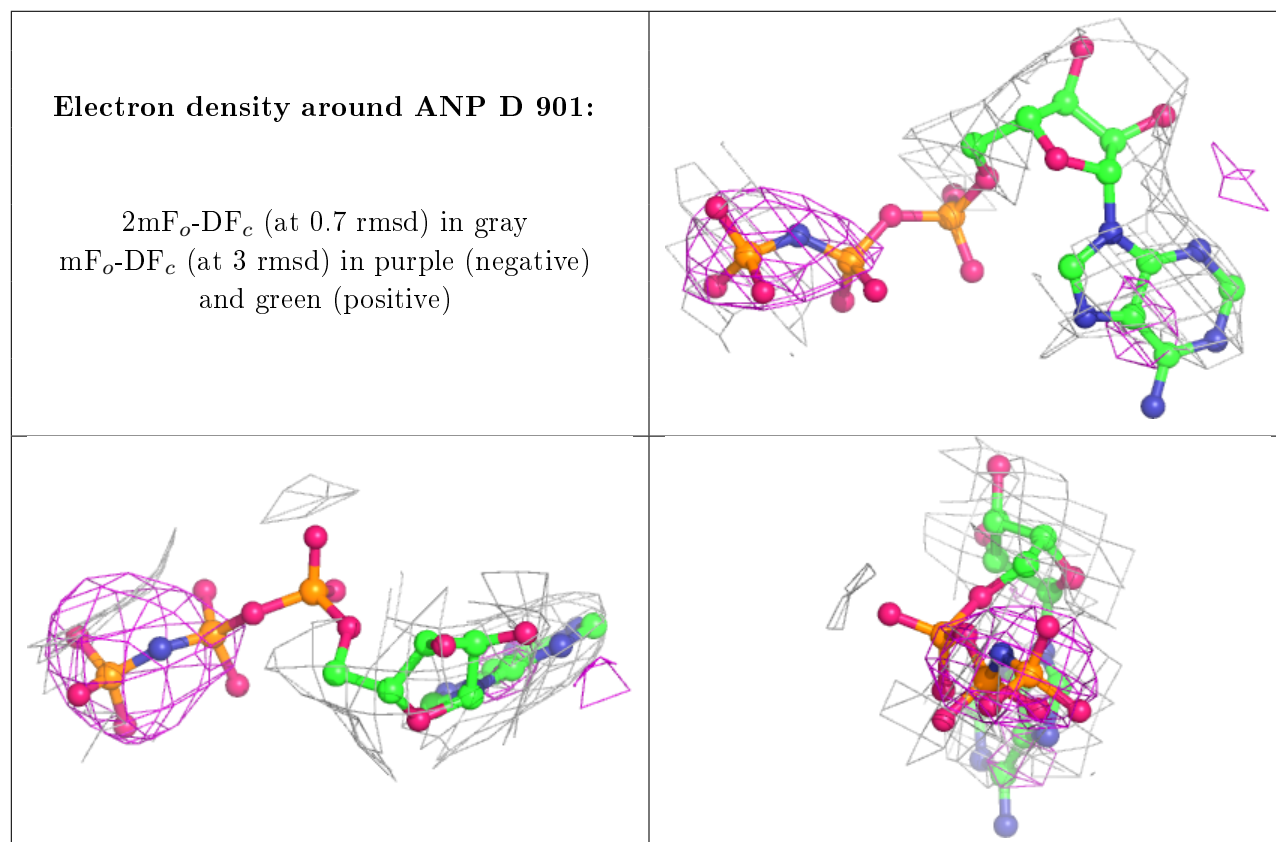
There are no carbohydrates in this entry.

### 6.4 Ligands [i](#)

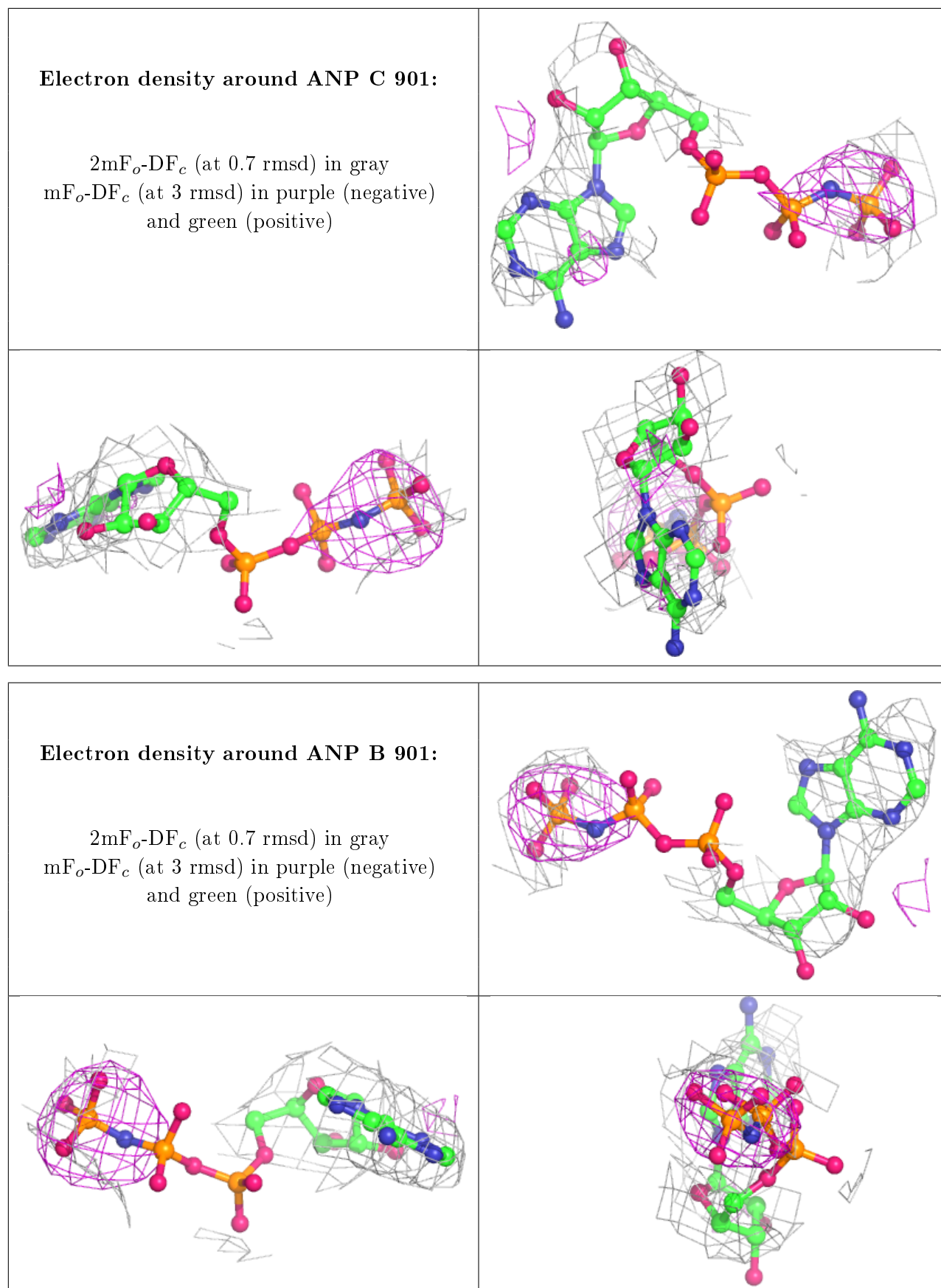
In the following table, the Atoms column lists the number of modelled atoms in the group and the number defined in the chemical component dictionary. The B-factors column lists the minimum, median, 95<sup>th</sup> percentile and maximum values of B factors of atoms in the group. The column labelled ‘Q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

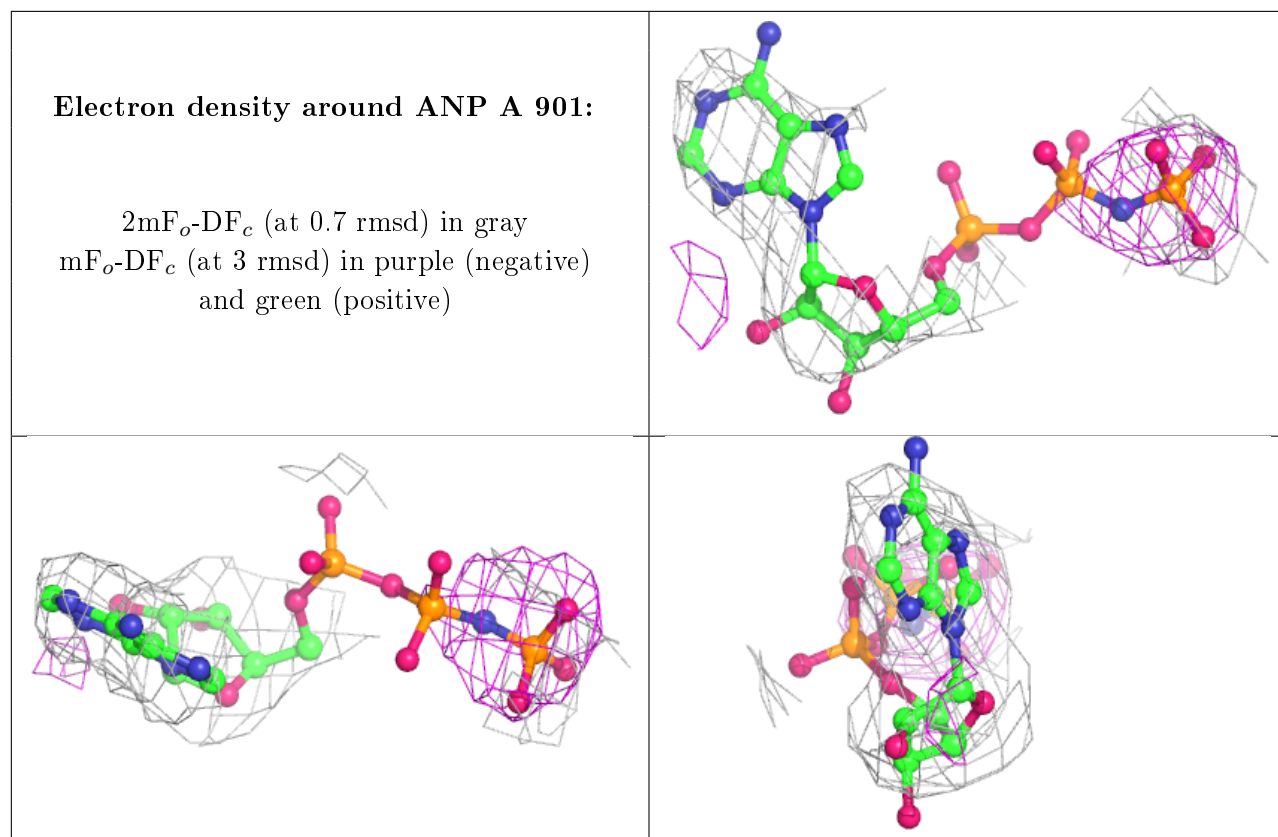
Mol	Type	Chain	Res	Atoms	RSCC	RSR	B-factors( $\text{\AA}^2$ )	Q<0.9
3	ANP	D	901	31/31	0.80	0.20	150,150,150,150	0
3	ANP	C	901	31/31	0.84	0.20	150,150,150,150	0
3	ANP	B	901	31/31	0.86	0.19	150,150,150,150	0
3	ANP	A	901	31/31	0.89	0.22	150,150,150,150	0
2	ADP	D	807	27/27	0.90	0.31	163,163,163,163	0
2	ADP	B	807	27/27	0.91	0.33	163,163,163,163	0
2	ADP	A	807	27/27	0.91	0.34	163,163,163,163	0
2	ADP	C	807	27/27	0.91	0.32	163,163,163,163	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.



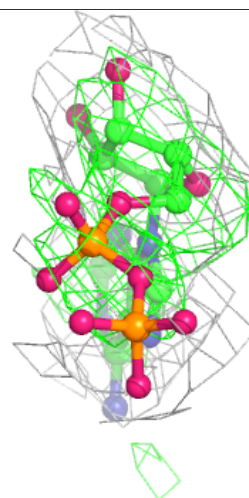
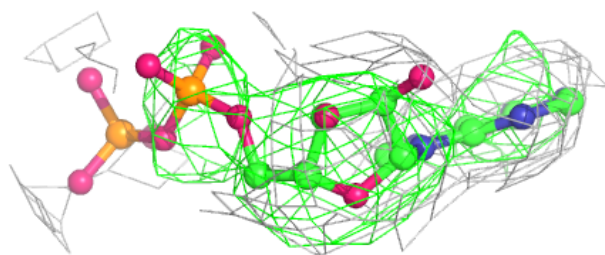
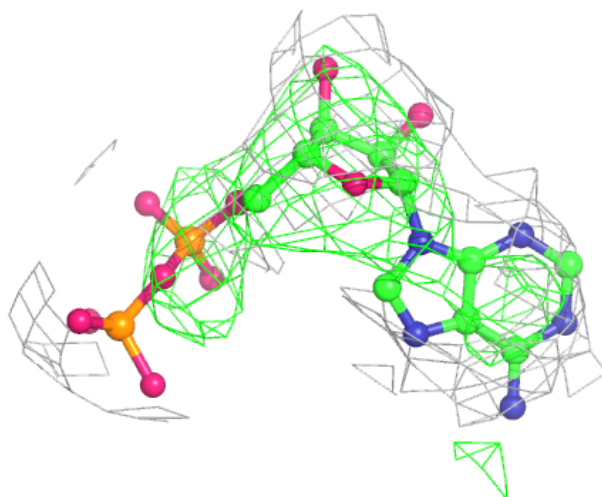






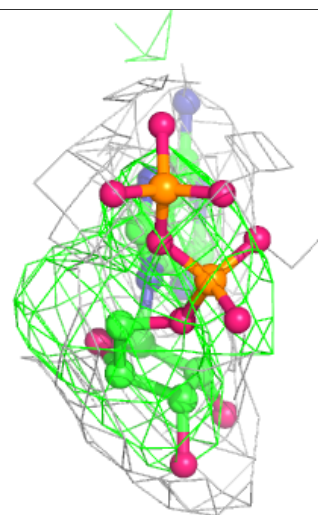
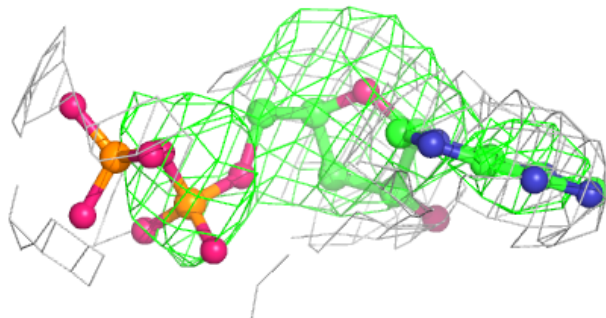
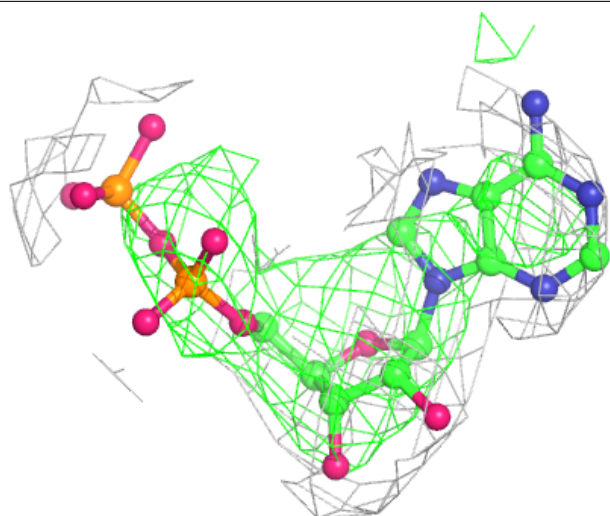
**Electron density around ADP D 807:**

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



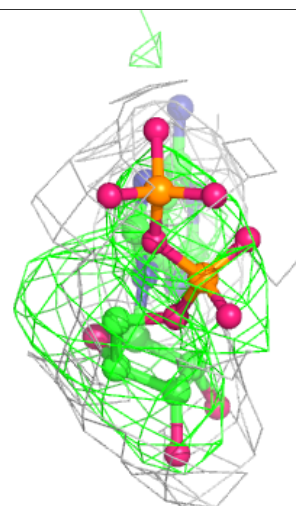
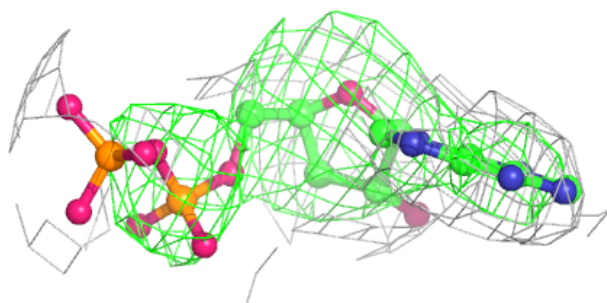
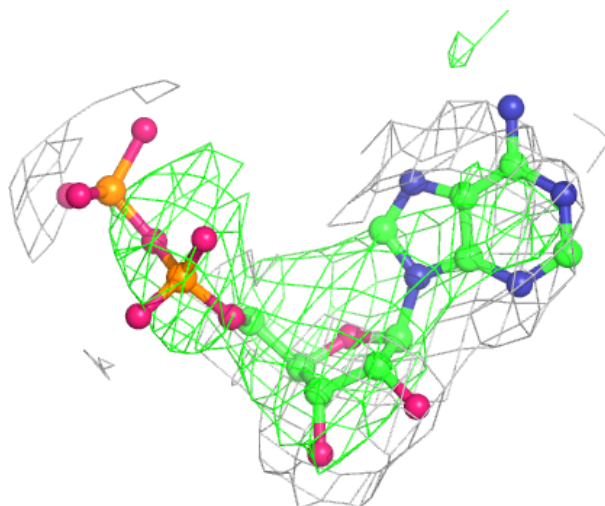
**Electron density around ADP B 807:**

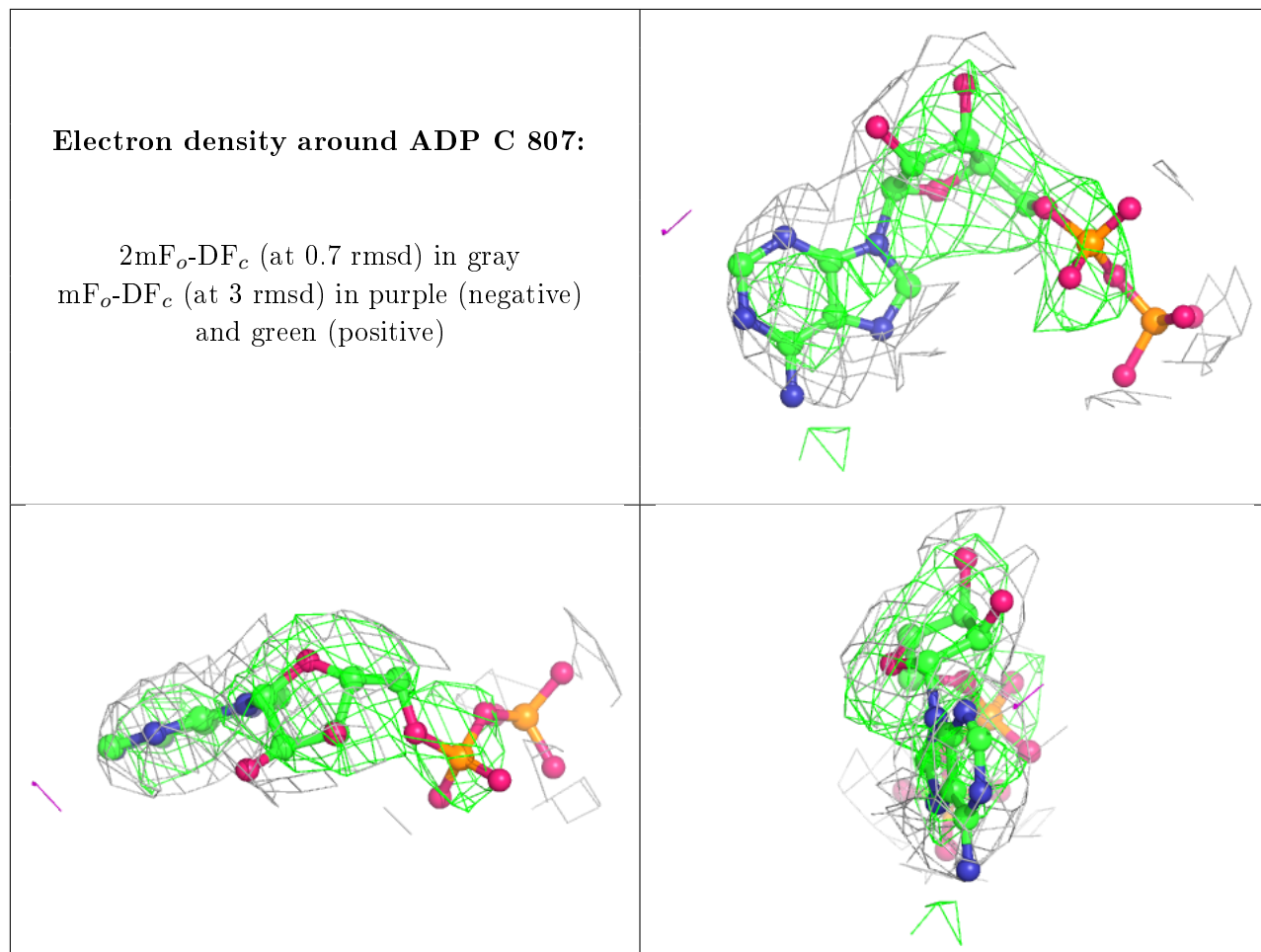
$2mF_o-DF_c$  (at 0.7 rmsd) in gray  
 $mF_o-DF_c$  (at 3 rmsd) in purple (negative)  
and green (positive)



**Electron density around ADP A 807:**

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





## 6.5 Other polymers [\(i\)](#)

There are no such residues in this entry.