



Full wwPDB EM Validation Report ⓘ

Mar 30, 2026 – 04:13 PM UTC

PDB ID : 6ZSE / pdb_00006zse
EMDB ID : EMD-11395
Title : Human mitochondrial ribosome in complex with mRNA, A/P-tRNA and P/E-tRNA
Authors : Aibara, S.; Singh, V.; Modelska, A.; Amunts, A.
Deposited on : 2020-07-15
Resolution : 5.00 Å(reported)

This is a Full wwPDB EM Validation 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/EMValidationReportHelp>
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:

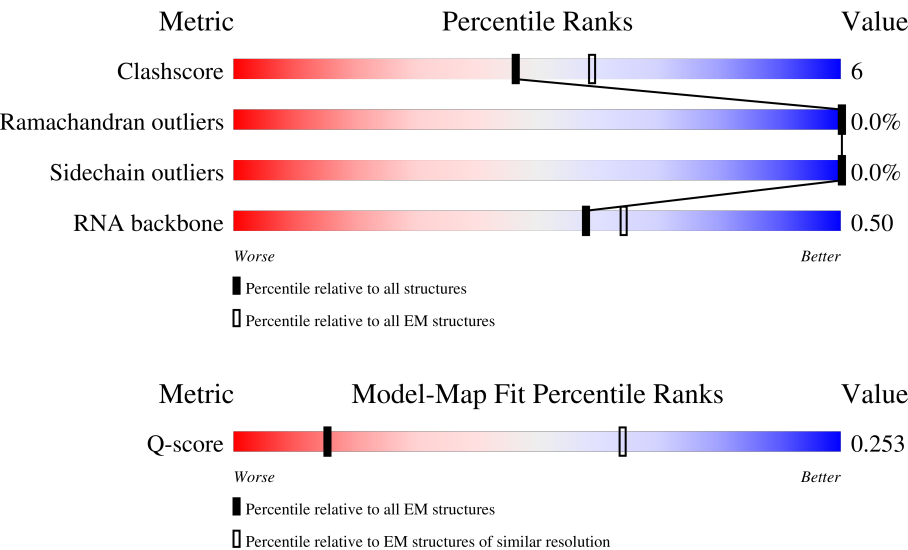
EMDB validation analysis : 0.0.1.dev132
Mogul : 2022.3.0, CSD as543be (2022)
MolProbity : 4-5-2 with Phenix2.0
Buster-report : wwPDB partial adaption of 1.1.7 (2018)
Percentile statistics : 20250101.v01 (using entries in the PDB archive January 1st 2025)
EM percentile statistics : 202505.v01 (Using data in the EMDB archive up until May 2025)
MapQ : 1.9.13
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP) : 2.49

1 Overall quality at a glance i

The following experimental techniques were used to determine the structure:
ELECTRON MICROSCOPY

The reported resolution of this entry is 5.00 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)	EM structures (#Entries)	Similar EM resolution (#Entries, resolution range(Å))
Clashscore	229148	23984	-
Ramachandran outliers	224038	23583	-
Sidechain outliers	223484	23102	-
RNA backbone	8273	3508	-
Q-score	-	25397	1057 (4.50 - 5.50)

The table below summarises the geometric issues observed across the polymeric chains and their fit to the map. The red, orange, yellow and green segments of the 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 EM map (all-atom inclusion $< 40\%$). The numeric value is given above the bar.

Mol	Chain	Length	Quality of chain
1	0	188	48%9%43%
2	1	65	6% 63%18%18%
3	2	92	39%11%50%

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Mol	Chain	Length	Quality of chain
4	3	188	
5	4	103	
6	5	423	
7	6	380	
8	7	338	
9	8	206	
10	9	137	
11	XA	1561	
12	A0	218	
13	A1	323	
14	A2	118	
15	A3	199	
16	A4	689	
17	AA	924	
18	AB	296	
19	AC	167	
20	AD	430	
21	AE	125	
22	AF	242	
23	AG	396	
24	AH	201	
25	AI	194	
26	AJ	138	
27	AK	128	
28	AL	257	

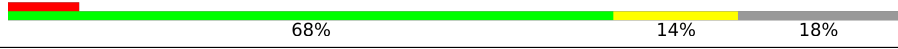
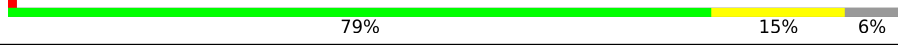
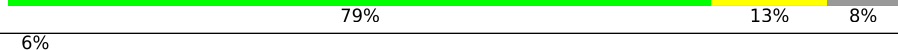
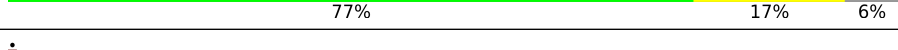
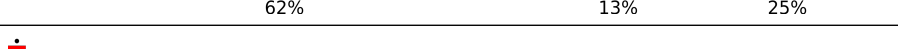
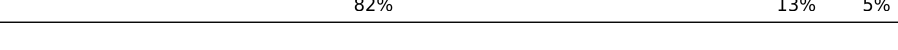
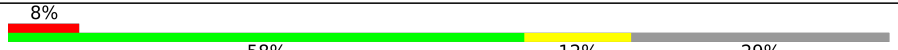
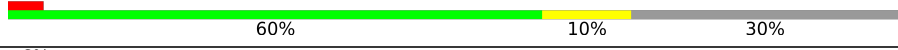
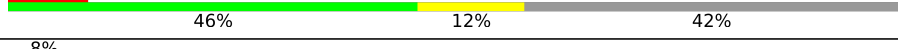
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Mol	Chain	Length	Quality of chain
29	AM	137	
30	AN	130	
31	AO	258	
32	AP	142	
33	AQ	87	
34	AR	360	
35	AS	190	
36	AT	173	
37	AU	205	
38	AV	414	
39	AW	187	
40	AX	398	
41	AY	395	
42	AZ	106	
43	XB	72	
44	XD	305	
45	XE	348	
46	XF	311	
47	XH	267	
48	XI	261	
49	XJ	192	
50	XK	178	
51	XL	145	
52	XM	296	
53	XN	251	

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Mol	Chain	Length	Quality of chain
54	XO	175	
55	XP	180	
56	XQ	292	
57	XR	149	
58	XS	205	
59	XT	206	
60	XU	153	
61	XV	216	
62	XW	148	
63	XX	256	
64	XY	250	
65	XZ	161	
66	a	142	
67	b	215	
68	c	332	
69	d	306	
70	e	279	
71	f	212	
72	g	166	
73	h	158	
74	i	128	
75	j	123	
76	k	112	
77	l	138	
78	m	128	

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Mol	Chain	Length	Quality of chain
79	o	102	
80	p	206	
81	q	222	
82	r	196	
83	r1	14	
84	r2	76	
85	r3	73	
86	s	439	
87	t1	198	
87	t2	198	
87	t3	198	
87	t4	198	
87	t5	198	
87	t6	198	
88	A	8	

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
83	Y5P	r1	50	-	-	X	-
84	P5P	r2	9	-	-	X	-
85	Y5P	r3	54	-	-	X	-

2 Entry composition

There are 92 unique types of molecules in this entry. The entry contains 313629 atoms, of which 143000 are hydrogens and 0 are deuteriums.

In the tables below, 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 39S ribosomal protein L32, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
1	0	108	Total	C	H	N	O	S	0	0
			1783	545	903	172	157	6		

- Molecule 2 is a protein called 39S ribosomal protein L33, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
2	1	53	Total	C	H	N	O	S	0	0
			919	281	480	84	72	2		

- Molecule 3 is a protein called 39S ribosomal protein L34, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
3	2	46	Total	C	H	N	O	S	0	0
			782	233	406	83	59	1		

- Molecule 4 is a protein called 39S ribosomal protein L35, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
4	3	95	Total	C	H	N	O	S	0	0
			1714	539	883	162	127	3		

- Molecule 5 is a protein called 39S ribosomal protein L36, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
5	4	38	Total	C	H	N	O	S	0	0
			703	217	362	72	48	4		

- Molecule 6 is a protein called 39S ribosomal protein L37, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
6	5	393	Total	C	H	N	O	S	0	0
			6405	2070	3201	559	564	11		

- Molecule 7 is a protein called 39S ribosomal protein L38, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
7	6	354	Total	C	H	N	O	S	0	0
			5787	1881	2840	525	532	9		

- Molecule 8 is a protein called 39S ribosomal protein L39, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
8	7	291	Total	C	H	N	O	S	0	0
			4737	1514	2372	401	432	18		

- Molecule 9 is a protein called 39S ribosomal protein L40, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
9	8	139	Total	C	H	N	O	S	0	0
			2377	747	1202	208	218	2		

- Molecule 10 is a protein called 39S ribosomal protein L41, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
10	9	124	Total	C	H	N	O	S	0	0
			1983	644	987	170	180	2		

- Molecule 11 is a RNA chain called 16S mitochondrial rRNA.

Mol	Chain	Residues	Atoms						AltConf	Trace
11	XA	1499	Total	C	H	N	O	P	0	0
			47992	14284	16159	5756	10294	1499		

- Molecule 12 is a protein called 28S ribosomal protein S34, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
12	A0	201	Total	C	H	N	O	S	0	0
			3369	1065	1685	322	292	5		

- Molecule 13 is a protein called 28S ribosomal protein S35, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
13	A1	275	Total	C	H	N	O	S	0	0
			4491	1414	2261	380	425	11		

- Molecule 14 is a protein called Coiled-coil-helix-coiled-coil-helix domain-containing protein 1.

Mol	Chain	Residues	Atoms						AltConf	Trace
14	A2	116	Total	C	H	N	O	S	0	0
			1889	574	964	181	162	8		

- Molecule 15 is a protein called Aurora kinase A-interacting protein.

Mol	Chain	Residues	Atoms						AltConf	Trace
15	A3	69	Total	C	H	N	O	S	0	0
			1292	393	682	130	86	1		

- Molecule 16 is a protein called Pentatricopeptide repeat domain-containing protein 3, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
16	A4	552	Total	C	H	N	O	S	0	0
			8955	2866	4485	756	820	28		

- Molecule 17 is a RNA chain called 12S mitochondrial rRNA.

Mol	Chain	Residues	Atoms						AltConf	Trace
17	AA	924	Total	C	H	N	O	P	0	0
			29592	8800	9964	3540	6364	924		

- Molecule 18 is a protein called 28S ribosomal protein S2, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
18	AB	218	Total	C	H	N	O	S	0	0
			3545	1135	1769	322	309	10		

- Molecule 19 is a protein called 28S ribosomal protein S24, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
19	AC	132	Total	C	H	N	O	S	0	0
			2170	699	1088	195	184	4		

- Molecule 20 is a protein called 28S ribosomal protein S5, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
20	AD	343	Total	C	H	N	O	S	0	0
			5499	1706	2783	515	482	13		

- Molecule 21 is a protein called 28S ribosomal protein S6, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
21	AE	122	Total	C	H	N	O	S	0	0
			1973	614	1001	177	177	4		

- Molecule 22 is a protein called 28S ribosomal protein S7, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
22	AF	201	Total	C	H	N	O	S	0	0
			3384	1069	1716	305	283	11		

- Molecule 23 is a protein called 28S ribosomal protein S9, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
23	AG	304	Total	C	H	N	O	S	0	0
			4995	1593	2490	444	454	14		

- Molecule 24 is a protein called 28S ribosomal protein S10, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
24	AH	135	Total	C	H	N	O	S	0	0
			2241	712	1136	187	203	3		

- Molecule 25 is a protein called 28S ribosomal protein S11, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
25	AI	136	Total	C	H	N	O	S	0	0
			2063	637	1052	192	178	4		

- Molecule 26 is a protein called 28S ribosomal protein S12, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
26	AJ	108	Total	C	H	N	O	S	0	0
			1725	521	887	169	142	6		

- Molecule 27 is a protein called 28S ribosomal protein S14, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
27	AK	101	Total	C	H	N	O	S	0	0
			1746	537	885	179	140	5		

- Molecule 28 is a protein called 28S ribosomal protein S15, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
28	AL	164	Total	C	H	N	O	S	0	0
			2855	883	1473	257	235	7		

- Molecule 29 is a protein called 28S ribosomal protein S16, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
29	AM	116	Total	C	H	N	O	S	0	0
			1871	582	951	182	150	6		

- Molecule 30 is a protein called 28S ribosomal protein S17, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
30	AN	107	Total	C	H	N	O	S	0	0
			1754	549	908	153	141	3		

- Molecule 31 is a protein called 28S ribosomal protein S18b, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
31	AO	185	Total	C	H	N	O	S	0	0
			3018	970	1490	285	267	6		

- Molecule 32 is a protein called 28S ribosomal protein S18c, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
32	AP	95	Total	C	H	N	O	S	0	0
			1561	493	796	132	132	8		

- Molecule 33 is a protein called 28S ribosomal protein S21, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
33	AQ	85	Total	C	H	N	O	S	0	0
			1483	455	749	149	123	7		

There is a discrepancy between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
AQ	50	ARG	CYS	variant	UNP P82921

- Molecule 34 is a protein called 28S ribosomal protein S22, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
34	AR	250	Total	C	H	N	O	S	0	0
			4134	1314	2074	353	385	8		

- Molecule 35 is a protein called 28S ribosomal protein S23, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
35	AS	133	Total	C	H	N	O	S	0	0
			2203	709	1103	196	194	1		

- Molecule 36 is a protein called 28S ribosomal protein S25, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
36	AT	162	Total	C	H	N	O	S	0	0
			2672	850	1342	231	238	11		

- Molecule 37 is a protein called 28S ribosomal protein S26, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
37	AU	173	Total	C	H	N	O	S	0	0
			2932	900	1471	294	263	4		

- Molecule 38 is a protein called 28S ribosomal protein S27, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
38	AV	349	Total	C	H	N	O	S	0	0
			5729	1841	2862	478	536	12		

- Molecule 39 is a protein called 28S ribosomal protein S28, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
39	AW	97	Total	C	H	N	O	S	0	0
			1551	486	785	137	139	4		

- Molecule 40 is a protein called 28S ribosomal protein S29, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
40	AX	348	Total	C	H	N	O	S	0	0
			5617	1802	2803	491	510	11		

- Molecule 41 is a protein called 28S ribosomal protein S31, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
41	AY	113	Total	C	H	N	O	S	0	0
			1868	621	912	157	176	2		

- Molecule 42 is a protein called 28S ribosomal protein S33, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
42	AZ	86	Total	C	H	N	O	S	0	0
			1465	467	734	131	129	4		

- Molecule 43 is a RNA chain called mitochondrial tRNA^{Val}.

Mol	Chain	Residues	Atoms						AltConf	Trace
43	XB	59	Total	C	H	N	O	P	0	0
			1890	563	635	227	406	59		

- Molecule 44 is a protein called 39S ribosomal protein L2, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
44	XD	236	Total	C	H	N	O	S	0	0
			3738	1145	1896	373	315	9		

- Molecule 45 is a protein called 39S ribosomal protein L3, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
45	XE	304	Total	C	H	N	O	S	0	0
			4798	1539	2402	416	430	11		

- Molecule 46 is a protein called 39S ribosomal protein L4, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
46	XF	250	Total	C	H	N	O	S	0	0
			4057	1294	2044	365	348	6		

- Molecule 47 is a protein called 39S ribosomal protein L9, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
47	XH	95	Total	C	H	N	O		0	0
			1616	498	832	152	134			

- Molecule 48 is a protein called 39S ribosomal protein L10, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
48	XI	211	Total	C	H	N	O	S	0	0
			3474	1086	1783	303	291	11		

- Molecule 49 is a protein called 39S ribosomal protein L11, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
49	XJ	170	Total	C	H	N	O	S	0	0
			2658	825	1367	230	234	2		

- Molecule 50 is a protein called 39S ribosomal protein L13, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
50	XK	177	Total	C	H	N	O	S	0	0
			2899	934	1448	259	251	7		

- Molecule 51 is a protein called 39S ribosomal protein L14, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
51	XL	115	Total	C	H	N	O	S	0	0
			1830	559	941	171	154	5		

- Molecule 52 is a protein called 39S ribosomal protein L15, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
52	XM	287	Total	C	H	N	O	S	0	0
			4681	1472	2376	425	402	6		

- Molecule 53 is a protein called 39S ribosomal protein L16, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
53	XN	221	Total	C	H	N	O	S	0	0
			3586	1138	1808	325	305	10		

- Molecule 54 is a protein called 39S ribosomal protein L17, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
54	XO	152	Total	C	H	N	O	S	0	0
			2528	784	1283	239	215	7		

- Molecule 55 is a protein called 39S ribosomal protein L18, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
55	XP	143	Total	C	H	N	O	S	0	0
			2326	729	1162	223	207	5		

- Molecule 56 is a protein called 39S ribosomal protein L19, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
56	XQ	238	Total	C	H	N	O	S	0	0
			4000	1268	2022	352	349	9		

- Molecule 57 is a protein called 39S ribosomal protein L20, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
57	XR	140	Total	C	H	N	O	S	0	0
			2367	732	1214	231	186	4		

- Molecule 58 is a protein called 39S ribosomal protein L21, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
58	XS	160	Total	C	H	N	O	S	0	0
			2638	829	1354	226	225	4		

- Molecule 59 is a protein called 39S ribosomal protein L22, mitochondrial.

Mol	Chain	Residues	Atoms					AltConf	Trace	
59	XT	166	Total	C	H	N	O	S	0	0
			2778	875	1410	254	232	7		

- Molecule 60 is a protein called 39S ribosomal protein L23, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
60	XU	141	Total	C	H	N	O	S	0	0
			2335	743	1164	222	203	3		

- Molecule 61 is a protein called 39S ribosomal protein L24, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
61	XV	202	Total	C	H	N	O	S	0	0
			3304	1051	1656	294	295	8		

- Molecule 62 is a protein called 39S ribosomal protein L27, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
62	XW	111	Total	C	H	N	O	S	0	0
			1769	558	898	164	146	3		

- Molecule 63 is a protein called 39S ribosomal protein L28, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
63	XX	243	Total	C	H	N	O	S	0	0
			4089	1317	2054	351	362	5		

- Molecule 64 is a protein called 39S ribosomal protein L47, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
64	XY	178	Total	C	H	N	O	S	0	0
			3109	981	1575	295	254	4		

- Molecule 65 is a protein called 39S ribosomal protein L30, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
65	XZ	120	Total	C	H	N	O	S	0	0
			2008	626	1030	183	166	3		

- Molecule 66 is a protein called 39S ribosomal protein L42, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
66	a	97	Total	C	H	N	O	S	0	0
			1590	512	777	145	151	5		

- Molecule 67 is a protein called 39S ribosomal protein L43, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
67	b	148	Total	C	H	N	O	S	0	0
			2358	733	1180	229	213	3		

- Molecule 68 is a protein called 39S ribosomal protein L44, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
68	c	275	Total	C	H	N	O	S	0	0
			4437	1415	2220	383	410	9		

- Molecule 69 is a protein called 39S ribosomal protein L45, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
69	d	216	Total	C	H	N	O	S	0	0
			3501	1125	1743	305	315	13		

- Molecule 70 is a protein called 39S ribosomal protein L46, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
70	e	213	Total	C	H	N	O	S	0	0
			3463	1105	1732	304	317	5		

- Molecule 71 is a protein called 39S ribosomal protein L48, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
71	f	143	Total	C	H	N	O	S	0	0
			2313	737	1164	187	221	4		

- Molecule 72 is a protein called 39S ribosomal protein L49, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
72	g	132	Total	C	H	N	O	S	0	0
			2183	710	1086	191	194	2		

- Molecule 73 is a protein called 39S ribosomal protein L50, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
73	h	108	Total	C	H	N	O	S	0	0
			1748	560	866	154	165	3		

- Molecule 74 is a protein called 39S ribosomal protein L51, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
74	i	97	Total	C	H	N	O	S	0	0
			1684	532	857	165	126	4		

- Molecule 75 is a protein called 39S ribosomal protein L52, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
75	j	86	Total	C	H	N	O	S	0	0
			1367	426	678	134	127	2		

- Molecule 76 is a protein called 39S ribosomal protein L53, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
76	k	95	Total	C	H	N	O	S	0	0
			1477	456	745	139	132	5		

- Molecule 77 is a protein called 39S ribosomal protein L54, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
77	l	80	Total	C	H	N	O	S	0	0
			1327	427	654	118	125	3		

- Molecule 78 is a protein called 39S ribosomal protein L55, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
78	m	60	Total	C	H	N	O	S	0	0
			1025	309	525	104	85	2		

- Molecule 79 is a protein called Ribosomal protein 63, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
79	o	94	Total	C	H	N	O	S	0	0
			1601	501	804	165	128	3		

- Molecule 80 is a protein called Peptidyl-tRNA hydrolase ICT1, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
80	p	127	Total	C	H	N	O	S	0	0
			2141	661	1083	201	192	4		

- Molecule 81 is a protein called Growth arrest and DNA damage-inducible proteins-interacting protein 1.

Mol	Chain	Residues	Atoms						AltConf	Trace
81	q	164	Total	C	H	N	O	S	0	0
			2738	858	1359	267	249	5		

- Molecule 82 is a protein called 39S ribosomal protein S18a, mitochondrial.

Mol	Chain	Residues	Atoms						AltConf	Trace
82	r	152	Total	C	H	N	O	S	0	0
			2514	792	1267	239	208	8		

- Molecule 83 is a RNA chain called mRNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
83	r1	14	Total	C	N	O	P	0	0
			252	126	28	84	14		

- Molecule 84 is a RNA chain called A/P-tRNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
84	r2	76	Total	C	N	O	P	0	0
			1485	723	230	456	76		

- Molecule 85 is a RNA chain called P/E-tRNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
85	r3	73	Total	C	N	O	P	0	0
			1420	692	216	439	73		

- Molecule 86 is a protein called 39S ribosomal protein S30, mitochondrial.

Mol	Chain	Residues	Atoms					AltConf	Trace
86	s	370	Total	C	H	N	O	S	
			6058	1946	3022	542	534	14	0

- Molecule 87 is a protein called 39S ribosomal protein L12, mitochondrial.

Mol	Chain	Residues	Atoms					AltConf	Trace
87	t1	46	Total	C	H	N	O	2	0
			733	228	379	56	70		
87	t2	30	Total	C	H	N	O	0	0
			506	154	268	38	46		
87	t3	30	Total	C	H	N	O	0	0
			506	154	268	38	46		
87	t4	29	Total	C	H	N	O	0	0
			484	148	255	36	45		
87	t5	29	Total	C	H	N	O	0	0
			484	148	255	36	45		
87	t6	27	Total	C	H	N	O	0	0
			450	137	236	34	43		

- Molecule 88 is a protein called Quinupristin.

Mol	Chain	Residues	Atoms					AltConf	Trace
88	A	8	Total	C	H	N	O	S	
			140	53	67	9	10	1	0

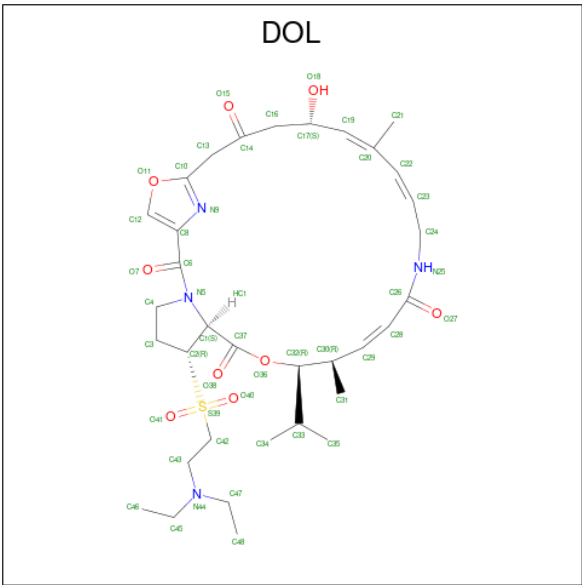
- Molecule 89 is ZINC ION (CCD ID: ZN) (formula: Zn).

Mol	Chain	Residues	Atoms		AltConf
89	0	1	Total 1	Zn 1	0
89	4	1	Total 1	Zn 1	0
89	AB	1	Total 1	Zn 1	0
89	AO	1	Total 1	Zn 1	0
89	AP	1	Total 1	Zn 1	0
89	AT	1	Total 1	Zn 1	0
89	r	1	Total 1	Zn 1	0

- Molecule 90 is MAGNESIUM ION (CCD ID: MG) (formula: Mg).

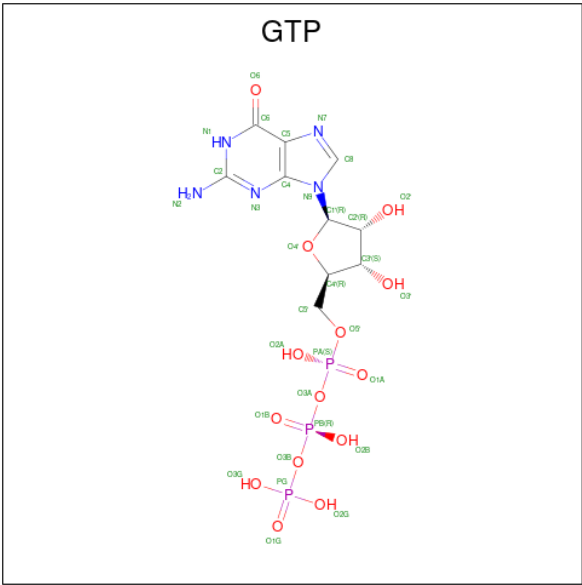
Mol	Chain	Residues	Atoms		AltConf
90	XA	141	Total 141	Mg 141	0
90	AA	46	Total 46	Mg 46	0
90	XD	1	Total 1	Mg 1	0
90	XE	1	Total 1	Mg 1	0
90	XI	1	Total 1	Mg 1	0
90	XM	2	Total 2	Mg 2	0
90	XW	1	Total 1	Mg 1	0
90	g	1	Total 1	Mg 1	0
90	o	1	Total 1	Mg 1	0

- Molecule 91 is 5-(2-DIETHYLAMINO-ETHANESULFONYL)-21-HYDROXY-10-ISOPROPYL-11,19-DIMETHYL-9,26-DIOXA-3,15,28-TRIAZA-TRICYCLO[23.2.1.00,255]OCTACOSA-1(27),12,17,19,25(28)-PENTAENE-2,8,14,23-TETRAONE (CCD ID: DOL) (formula: C₃₄H₅₀N₄O₉S).



Mol	Chain	Residues	Atoms					AltConf	
91	XA	1	Total	C	H	N	O	S	0
			98	34	50	4	9	1	

- Molecule 92 is GUANOSINE-5'-TRIPHOSPHATE (CCD ID: GTP) (formula: $C_{10}H_{16}N_5O_{14}P_3$).

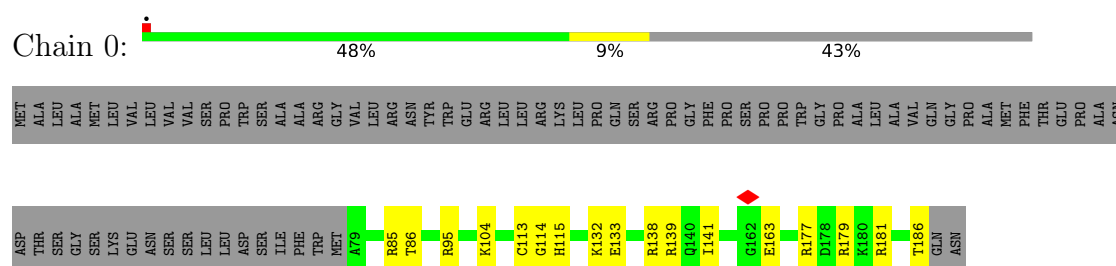


Mol	Chain	Residues	Atoms						AltConf
92	AX	1	Total	C	H	N	O	P	0
			42	10	10	5	14	3	

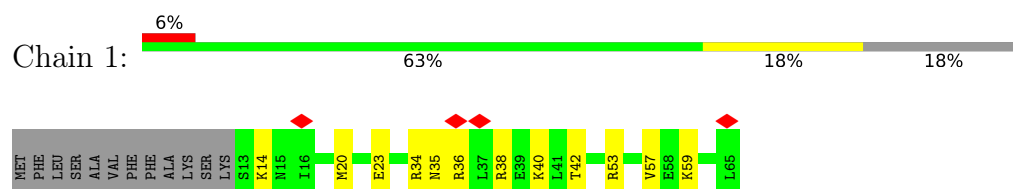
3 Residue-property plots

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 atom inclusion in map 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 diamond above a residue indicates a poor fit to the EM map for this residue (all-atom inclusion < 40%). 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.

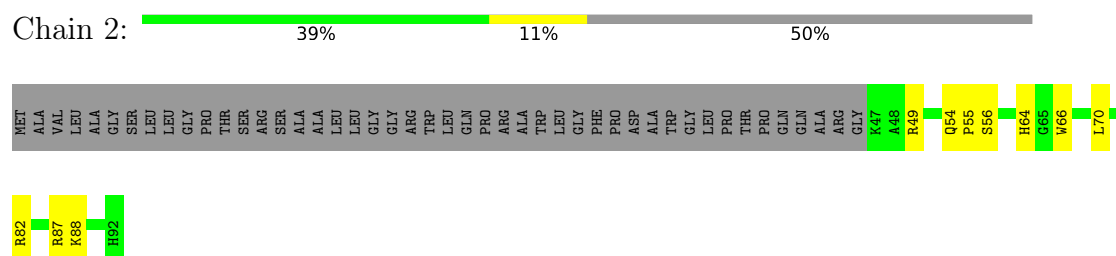
- Molecule 1: 39S ribosomal protein L32, mitochondrial



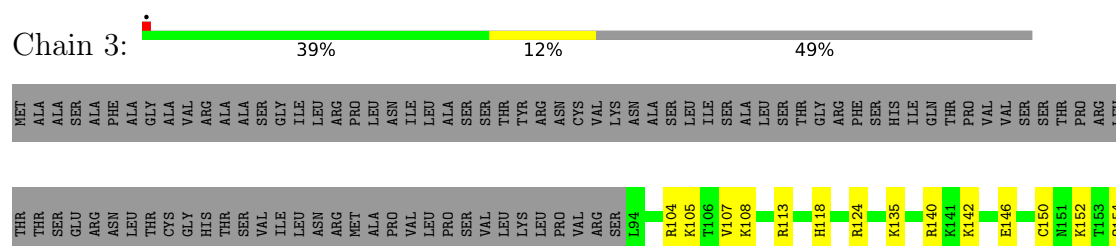
- Molecule 2: 39S ribosomal protein L33, mitochondrial



- Molecule 3: 39S ribosomal protein L34, mitochondrial



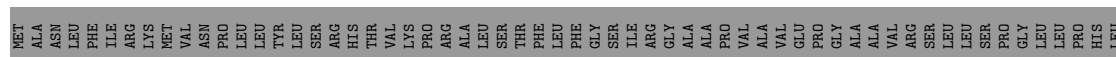
- Molecule 4: 39S ribosomal protein L35, mitochondrial





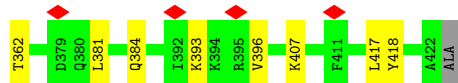
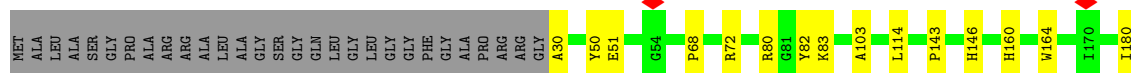
- Molecule 5: 39S ribosomal protein L36, mitochondrial

Chain 4: 30% 7% 63%



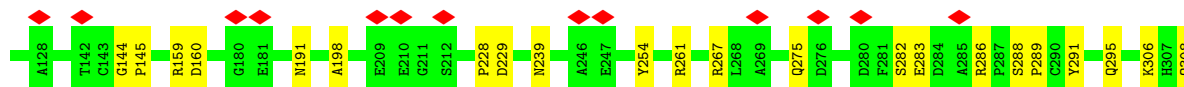
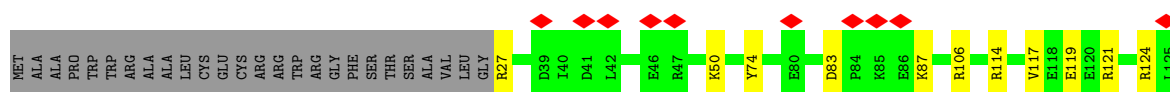
- Molecule 6: 39S ribosomal protein L37, mitochondrial

Chain 5: 81% 12% 7%



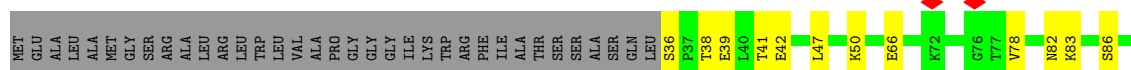
- Molecule 7: 39S ribosomal protein L38, mitochondrial

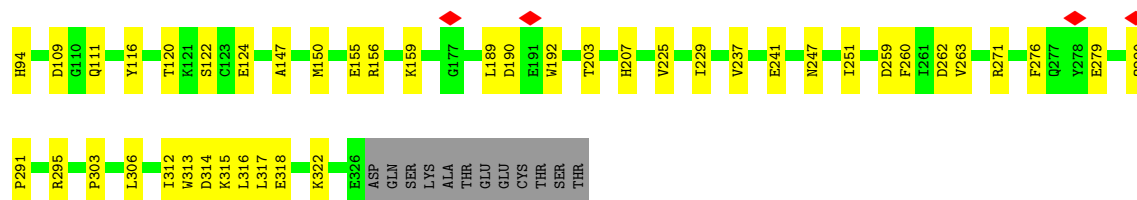
Chain 6: 6% 82% 11% 7%



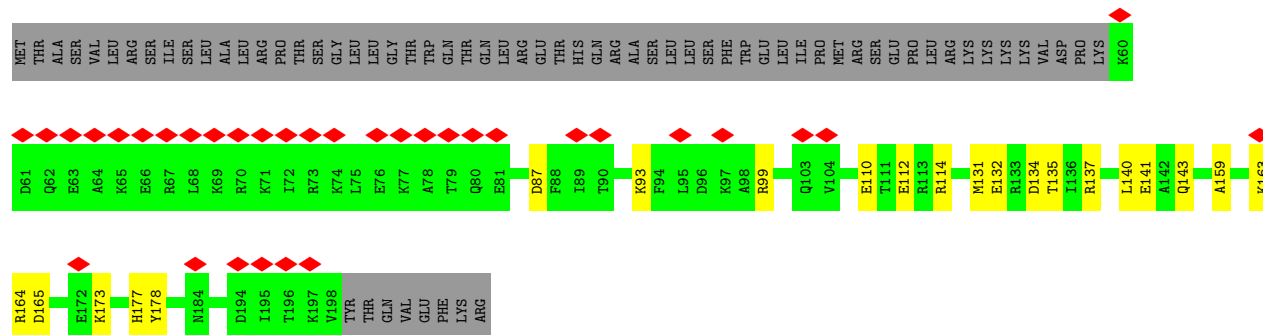
- Molecule 8: 39S ribosomal protein L39, mitochondrial

Chain 7: 70% 16% 14%

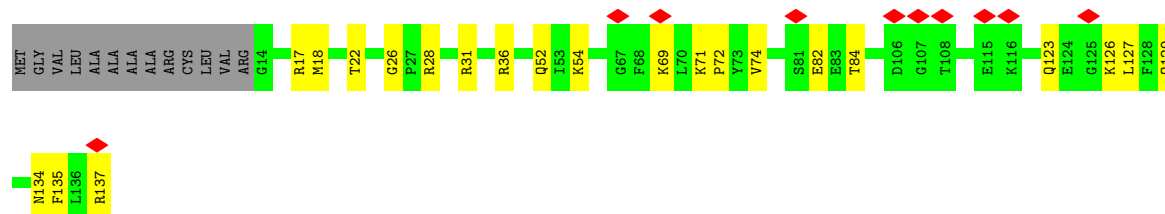
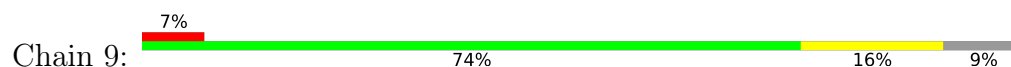




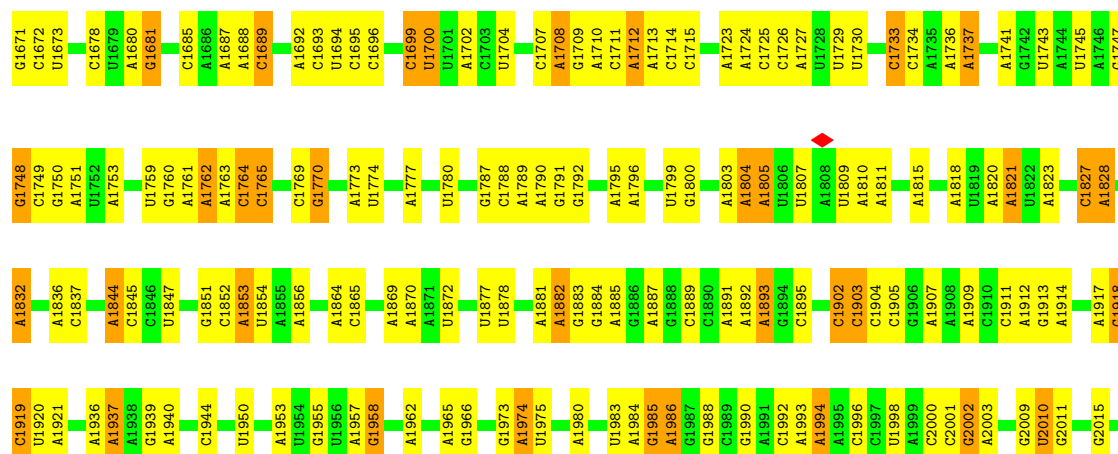
- Molecule 9: 39S ribosomal protein L40, mitochondrial

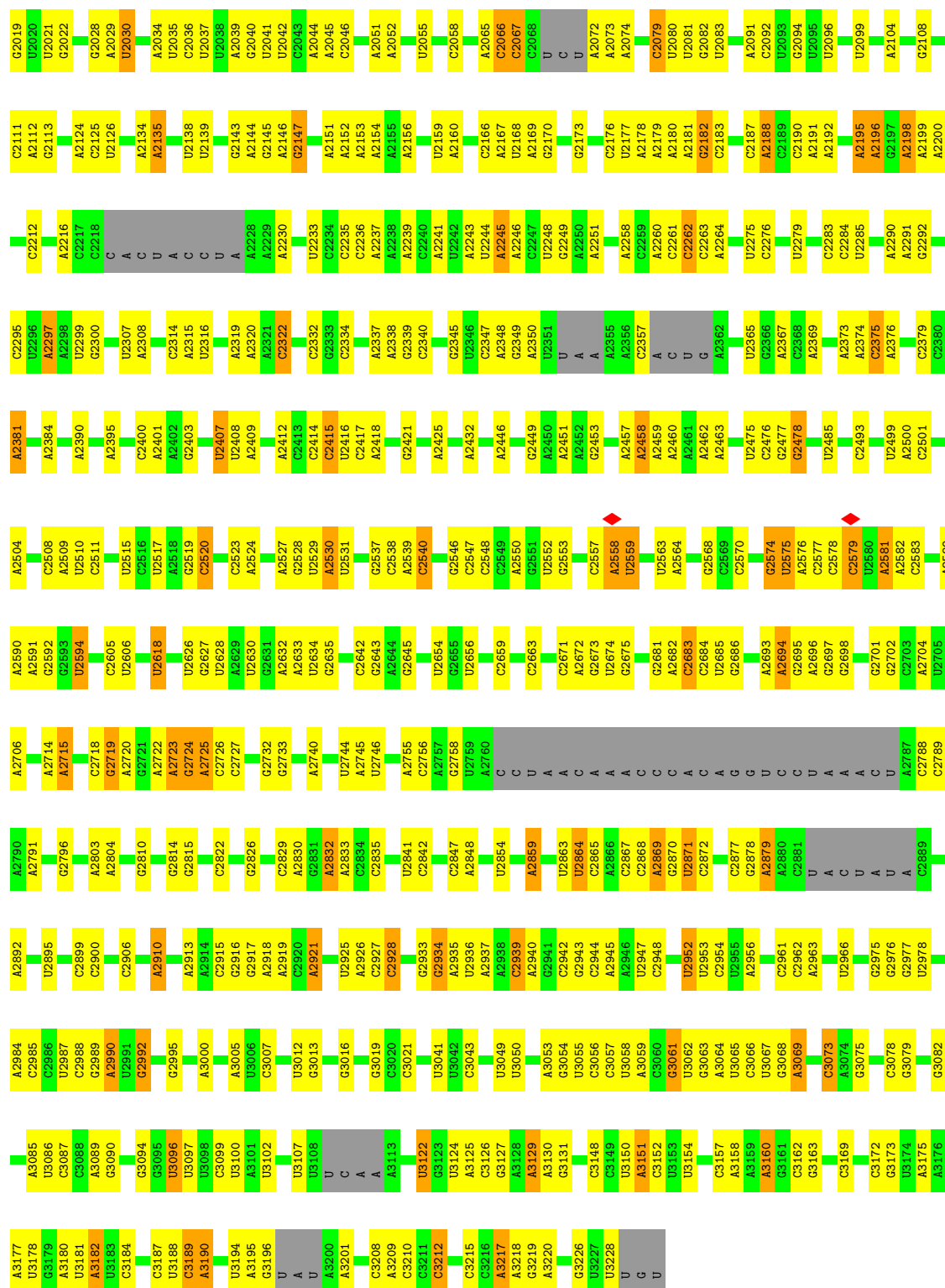


- Molecule 10: 39S ribosomal protein L41, mitochondrial

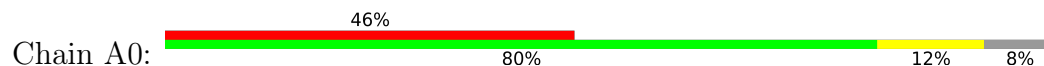


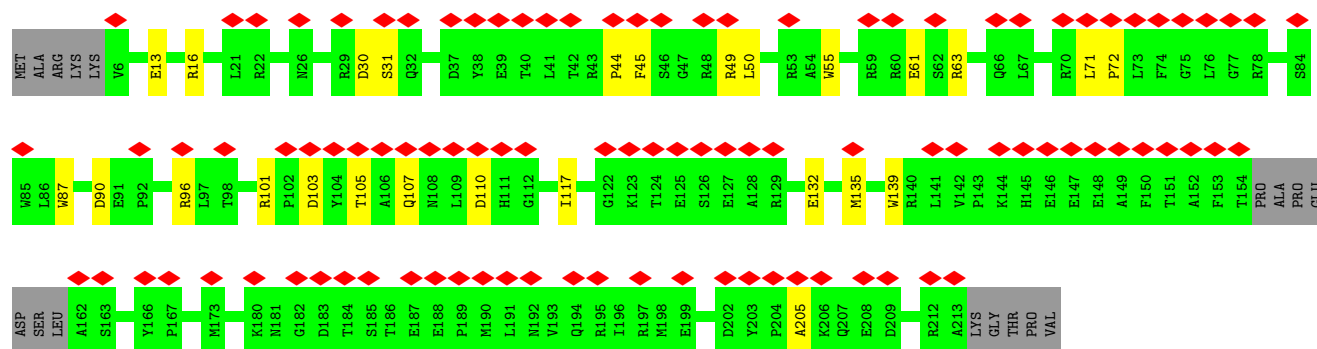
- Molecule 11: 16S mitochondrial rRNA



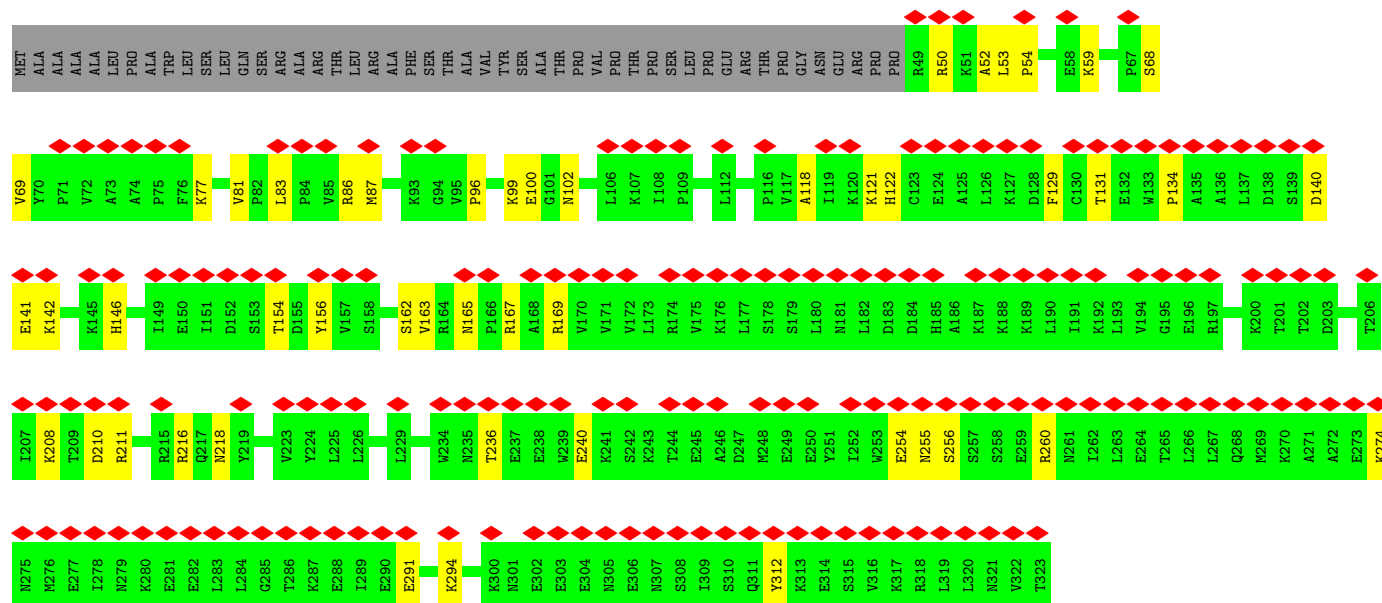


- Molecule 12: 28S ribosomal protein S34, mitochondrial

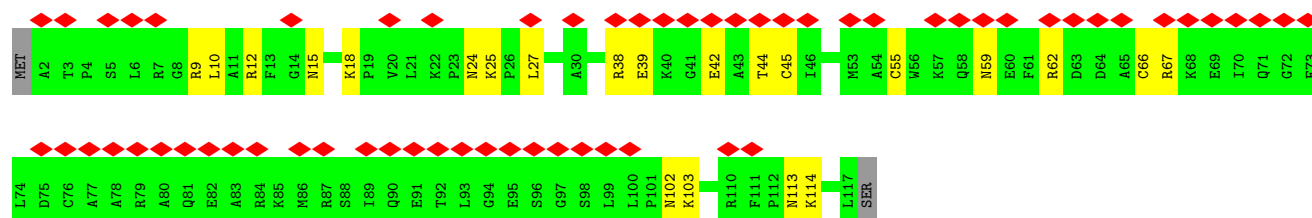
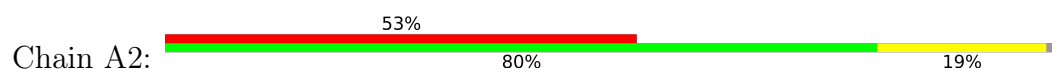




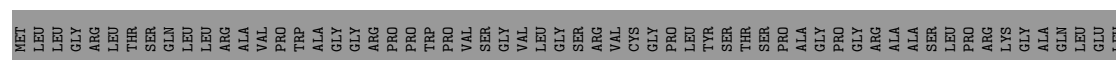
- Molecule 13: 28S ribosomal protein S35, mitochondrial



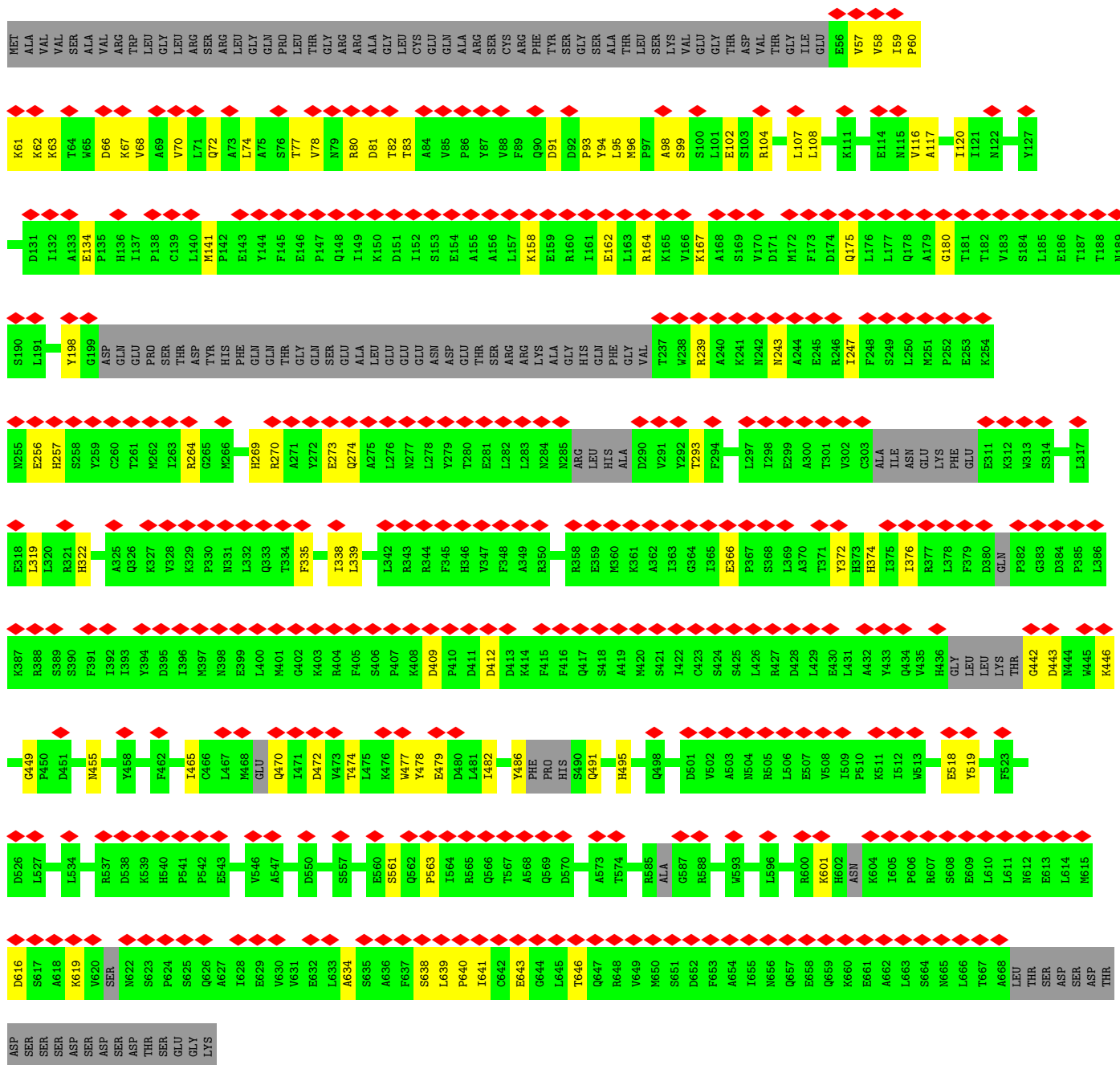
- Molecule 14: Coiled-coil-helix-coiled-coil-helix domain-containing protein 1



- Molecule 15: Aurora kinase A-interacting protein

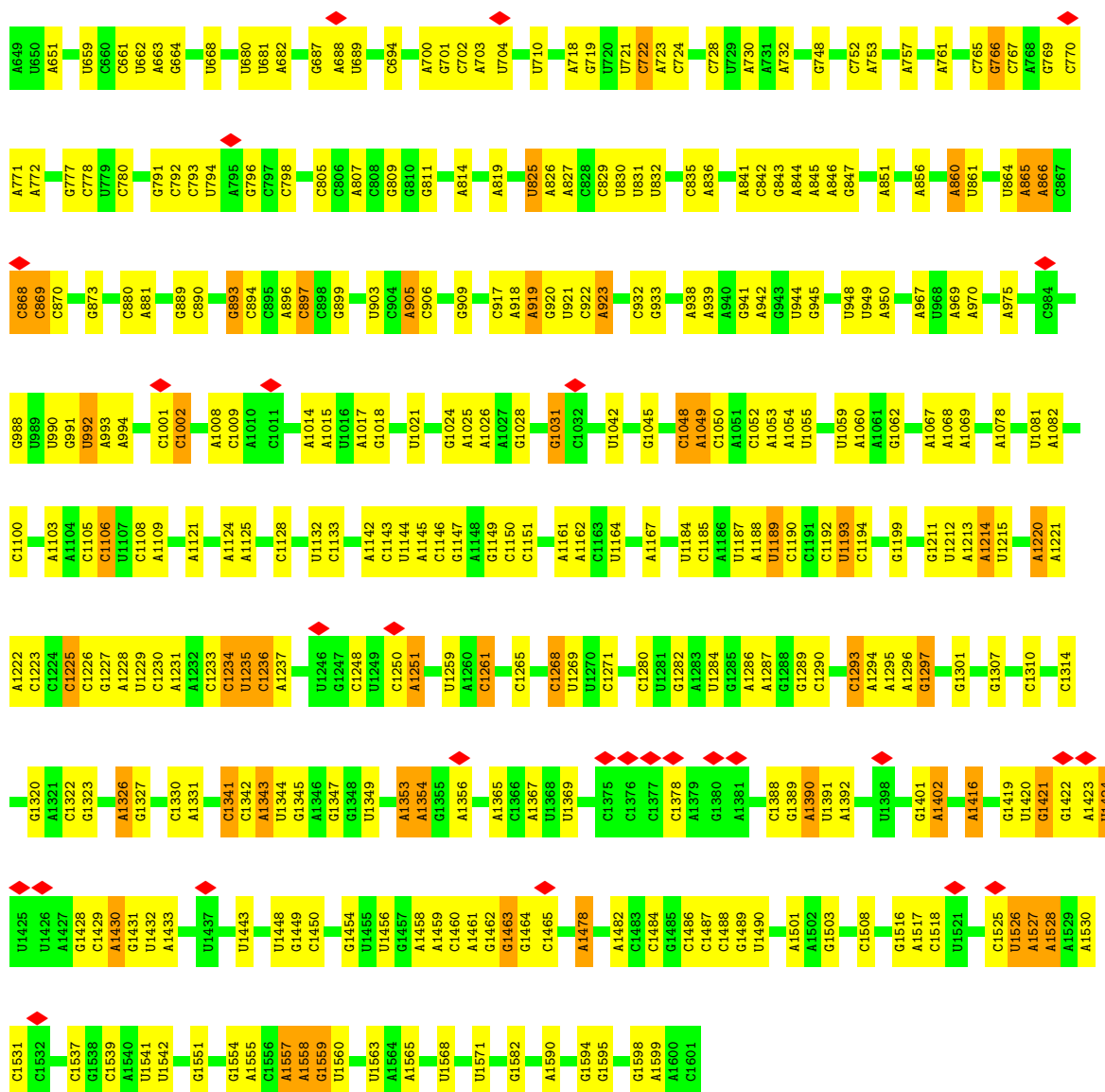


- Molecule 16: Pentatricopeptide repeat domain-containing protein 3, mitochondrial



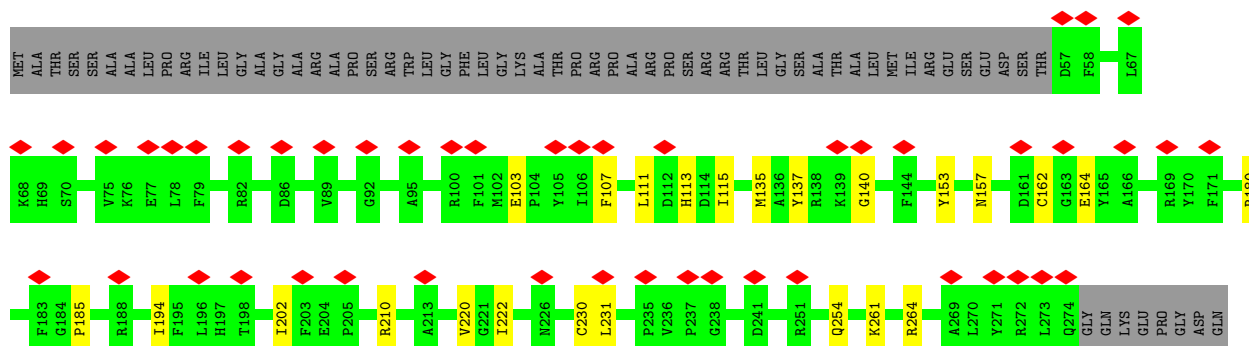
- Molecule 17: 12S mitochondrial rRNA

Chain AA: 



- Molecule 18: 28S ribosomal protein S2, mitochondrial

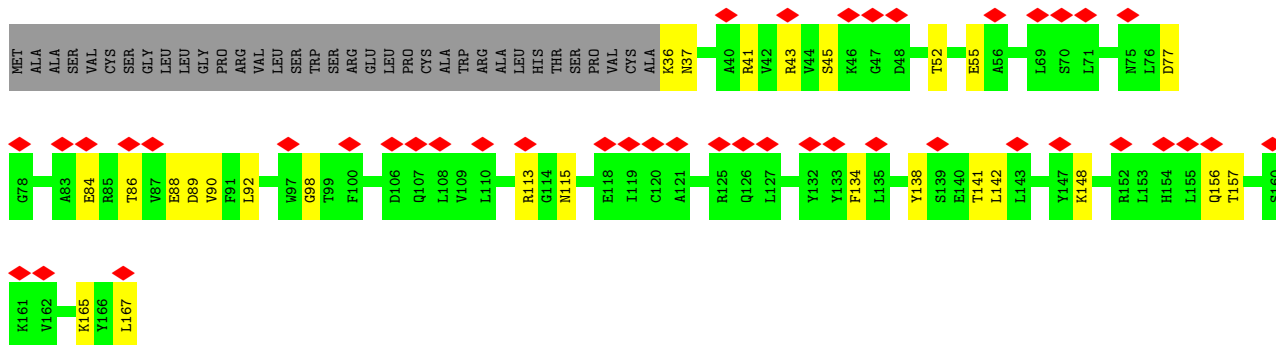
Chain AB: 



GLY
PRO
ALA
HIS
PRO
PRO
GLY
ALA
ASP
MET
SER
HIS
SER
LEU

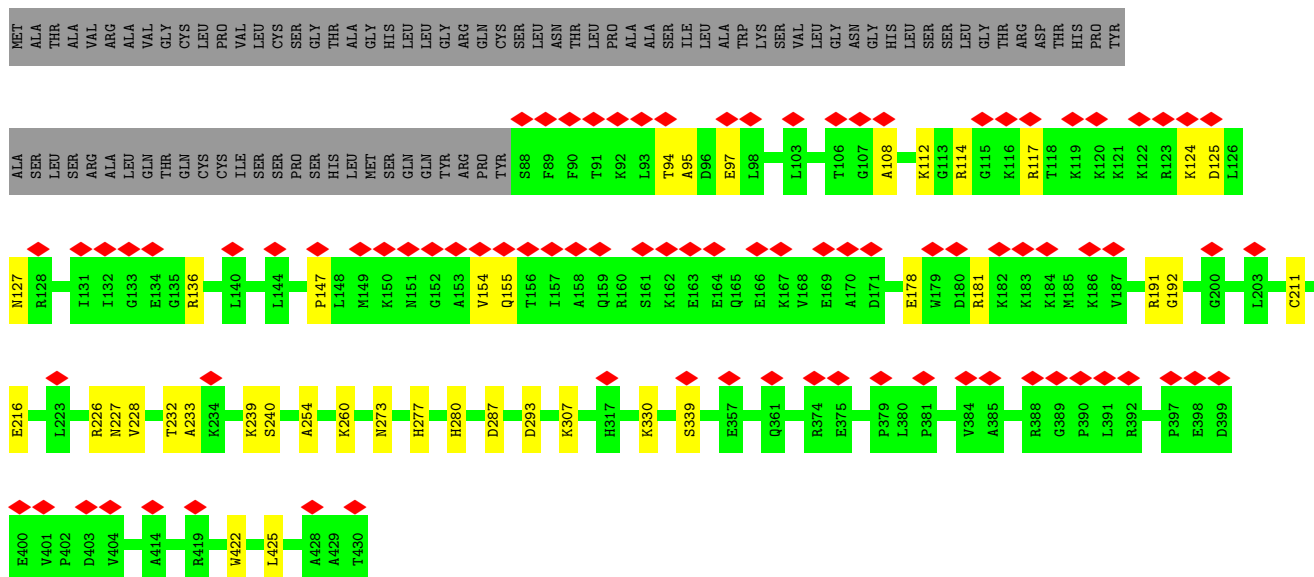
- Molecule 19: 28S ribosomal protein S24, mitochondrial

Chain AC: 



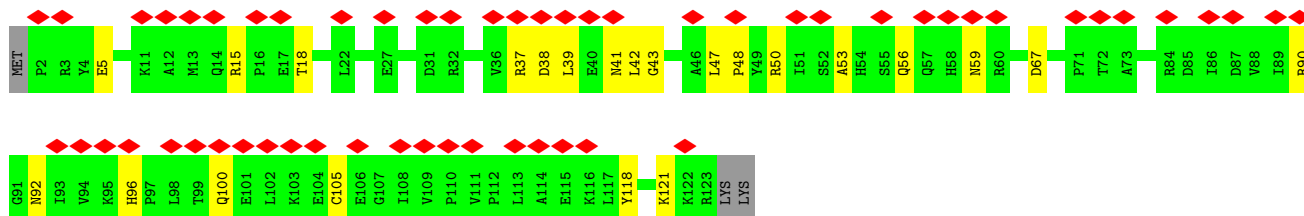
- Molecule 20: 28S ribosomal protein S5, mitochondrial

Chain AD: 

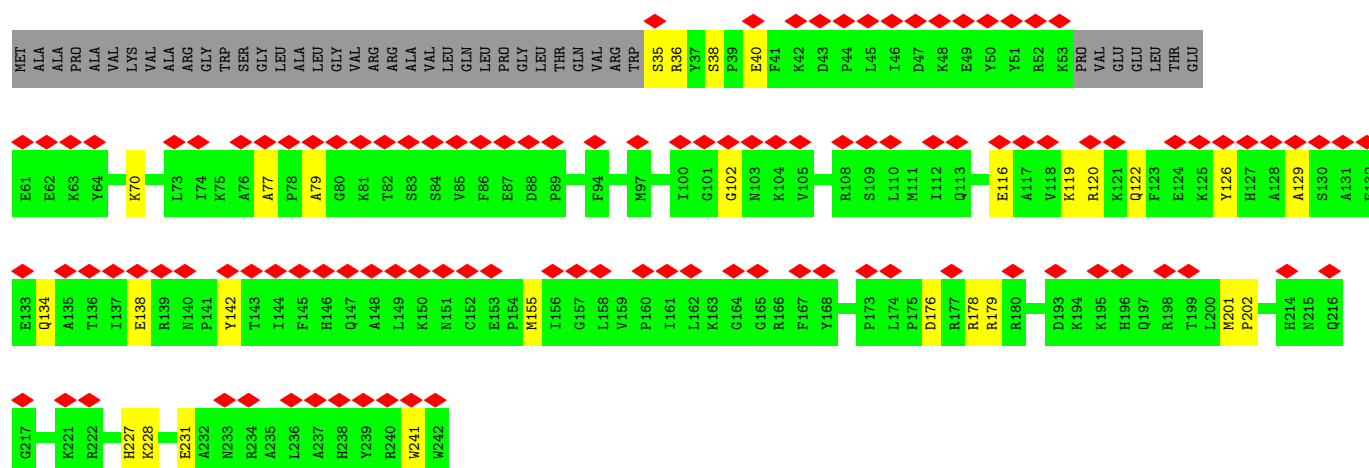
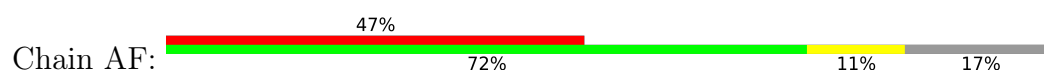


- Molecule 21: 28S ribosomal protein S6, mitochondrial

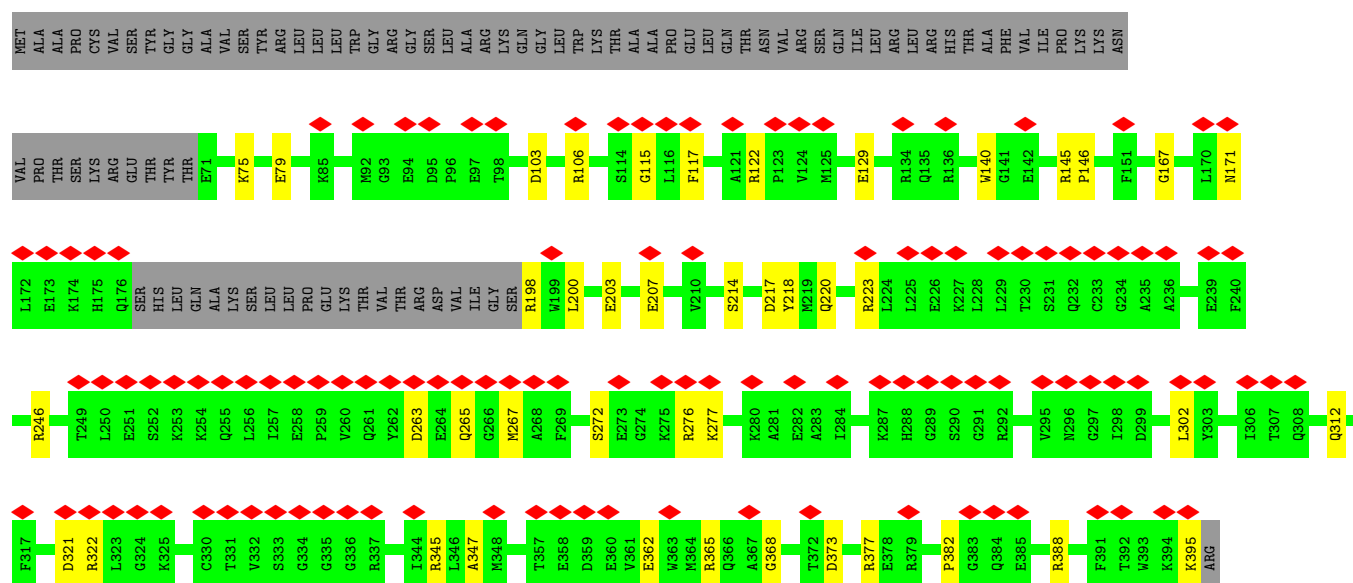
Chain AE: 



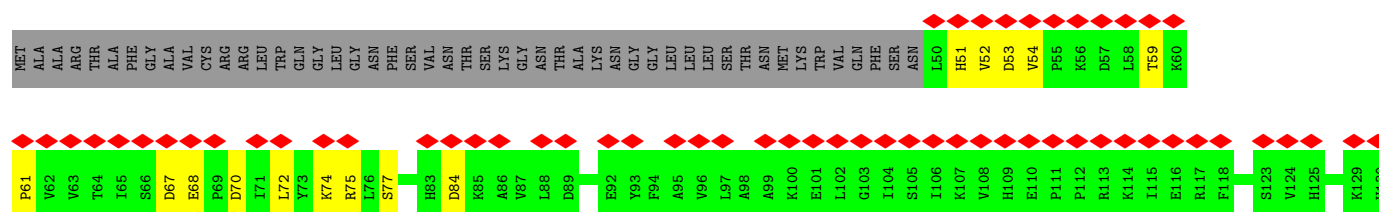
- Molecule 22: 28S ribosomal protein S7, mitochondrial

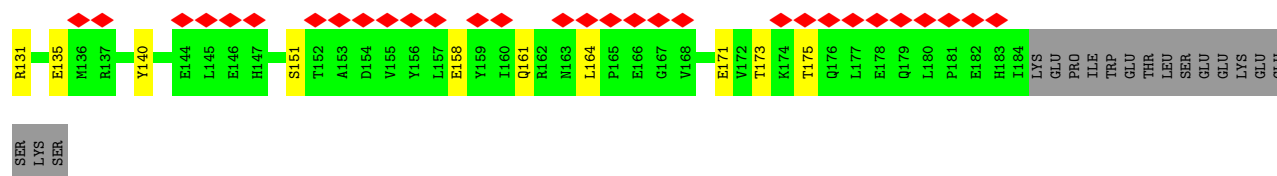


• Molecule 23: 28S ribosomal protein S9, mitochondrial

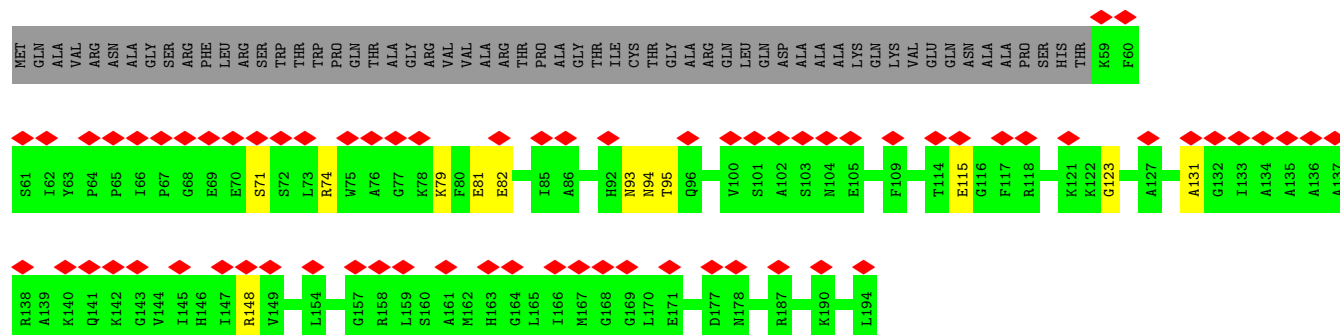


• Molecule 24: 28S ribosomal protein S10, mitochondrial

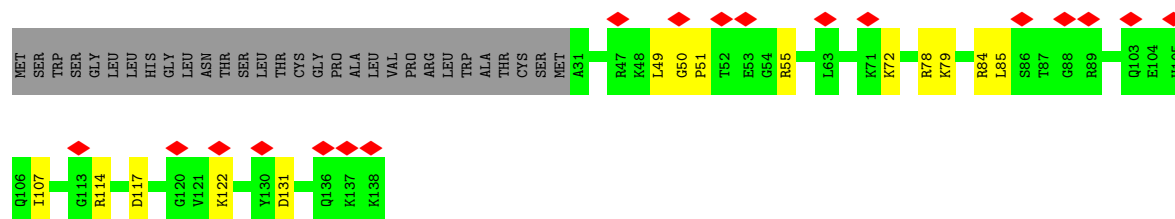




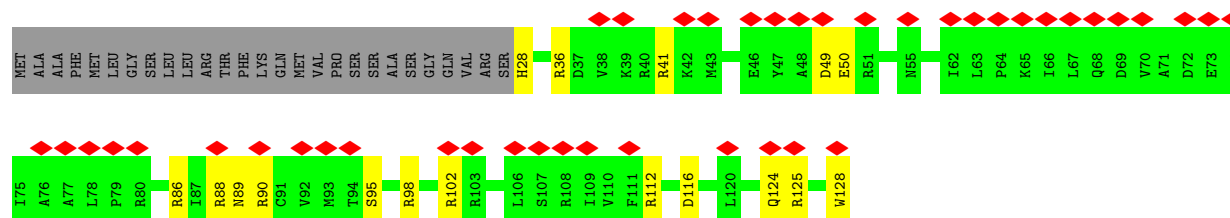
- Molecule 25: 28S ribosomal protein S11, mitochondrial



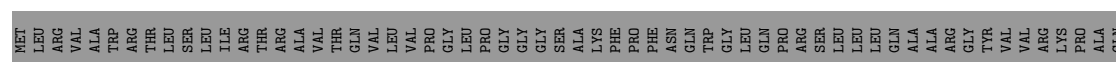
- Molecule 26: 28S ribosomal protein S12, mitochondrial



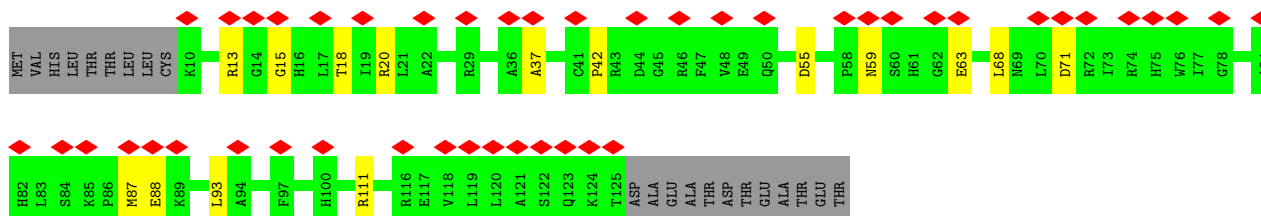
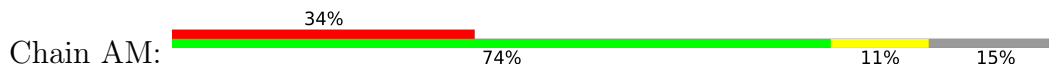
- Molecule 27: 28S ribosomal protein S14, mitochondrial



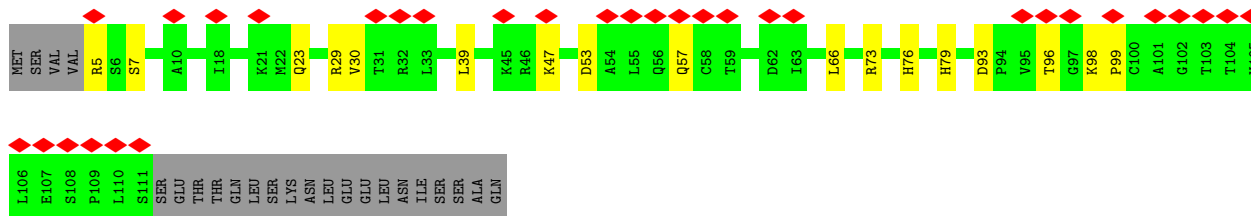
- Molecule 28: 28S ribosomal protein S15, mitochondrial



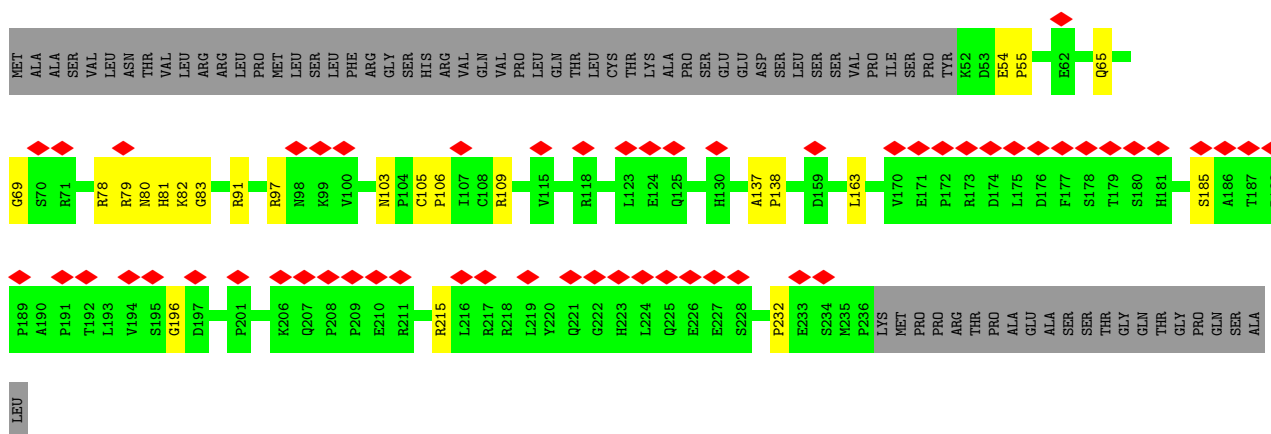
- Molecule 29: 28S ribosomal protein S16, mitochondrial



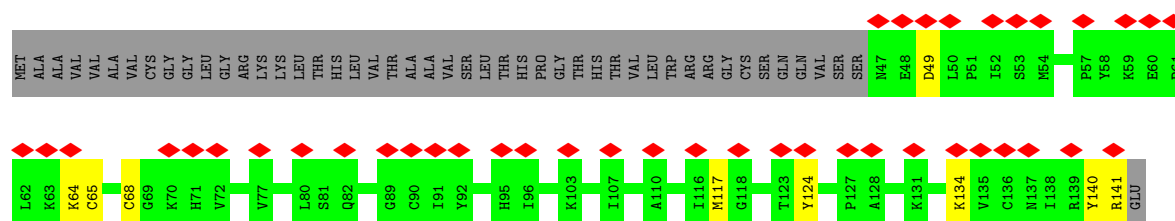
- Molecule 30: 28S ribosomal protein S17, mitochondrial



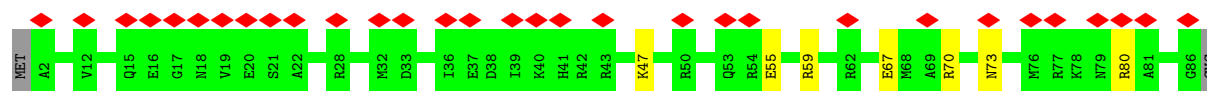
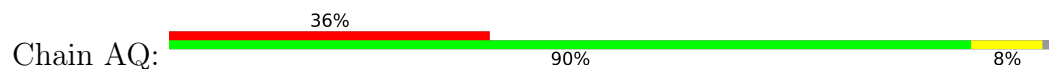
- Molecule 31: 28S ribosomal protein S18b, mitochondrial



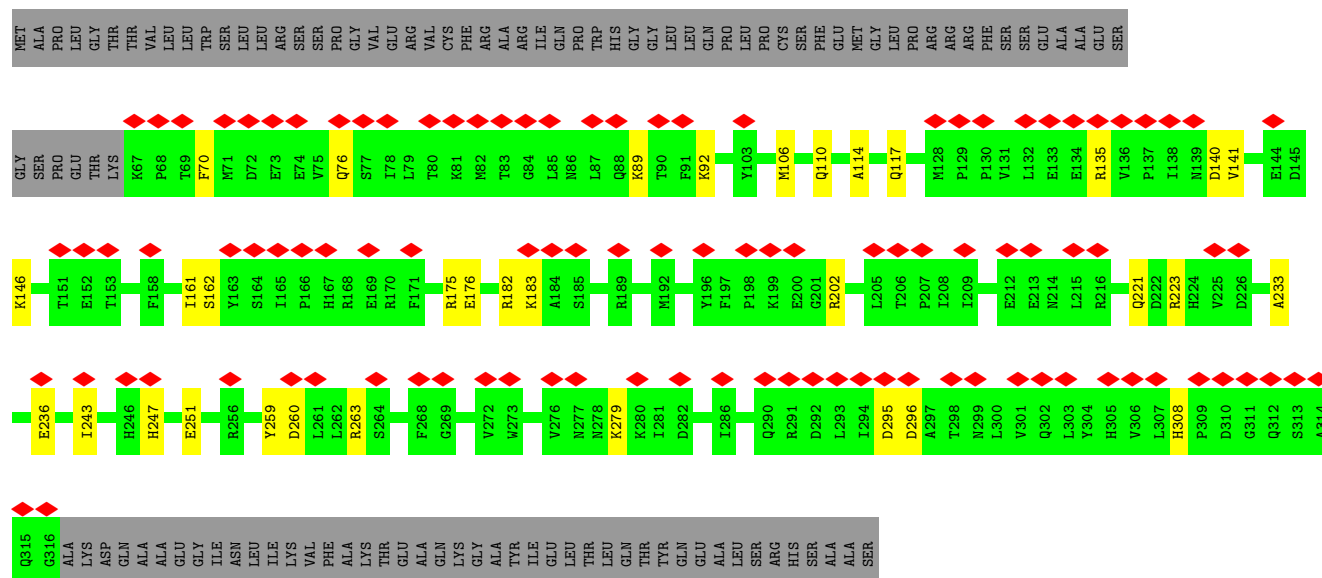
- Molecule 32: 28S ribosomal protein S18c, mitochondrial



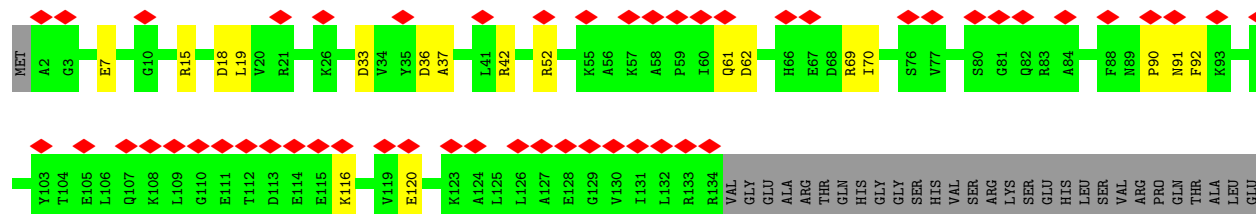
- Molecule 33: 28S ribosomal protein S21, mitochondrial



- Molecule 34: 28S ribosomal protein S22, mitochondrial



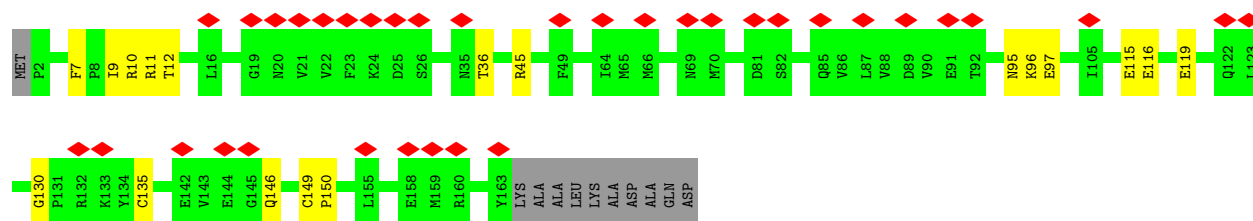
- Molecule 35: 28S ribosomal protein S23, mitochondrial



GLU ASN
GLU THR
GLN LYS
GLU VAL
PRO PRO
GLN ASP
GLN HIS
LEU LEU
GLU ALA
PRO PRO
ALA ALA
SER SER
GLN GLN
LEU LEU
PRO PRO

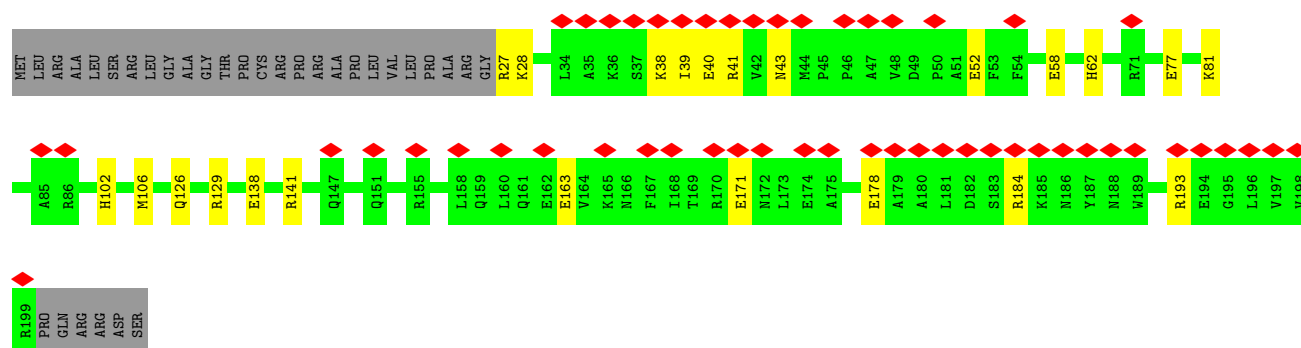
- Molecule 36: 28S ribosomal protein S25, mitochondrial

Chain AT:



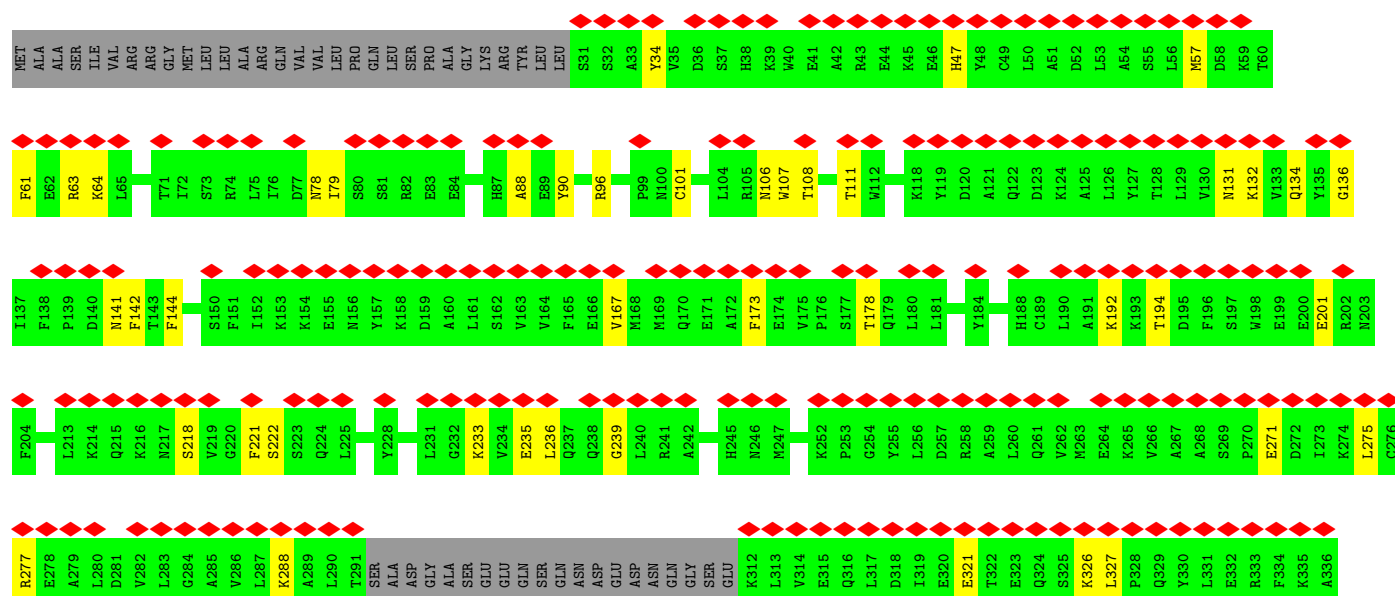
- Molecule 37: 28S ribosomal protein S26, mitochondrial

Chain AU:

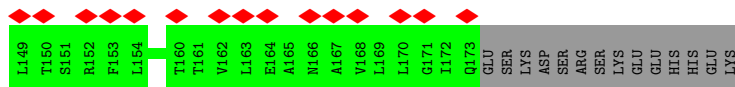
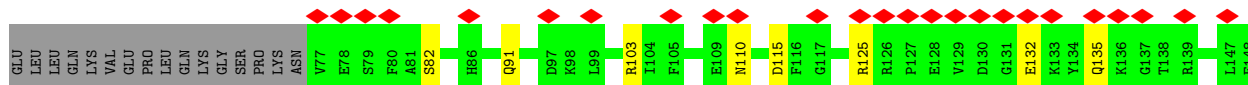
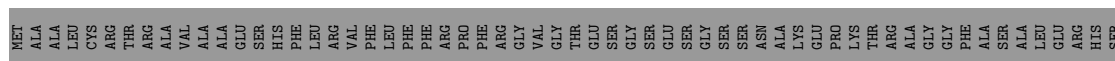


- Molecule 38: 28S ribosomal protein S27, mitochondrial

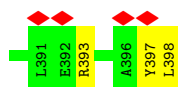
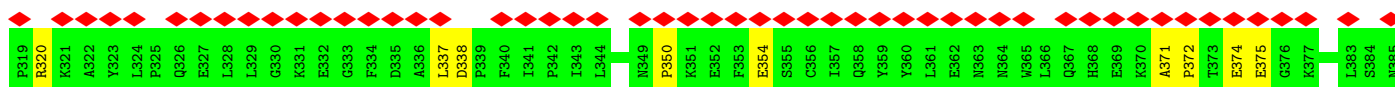
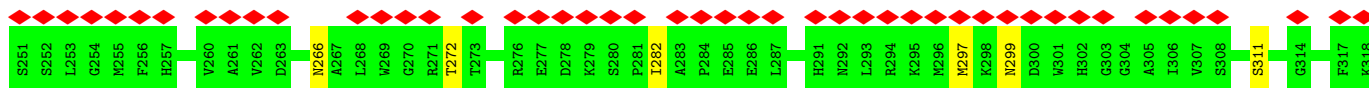
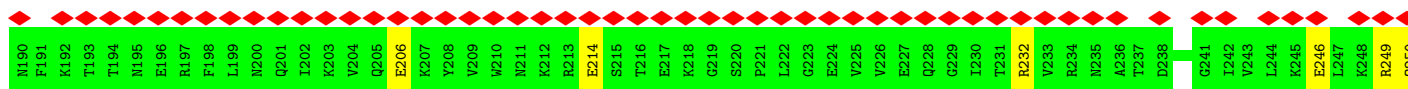
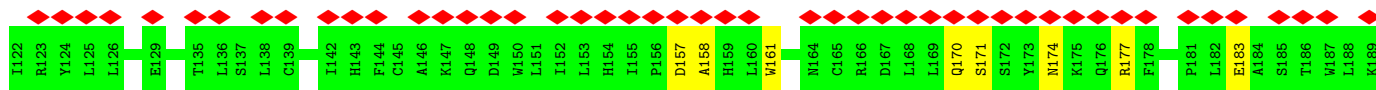
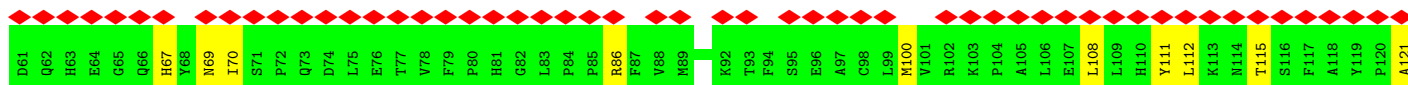
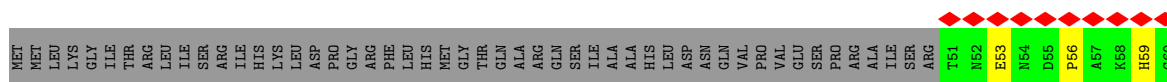
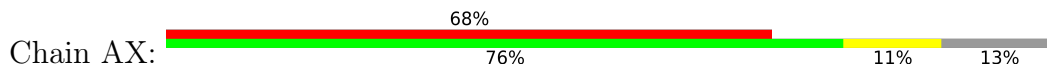
Chain AV:



- Molecule 39: 28S ribosomal protein S28, mitochondrial



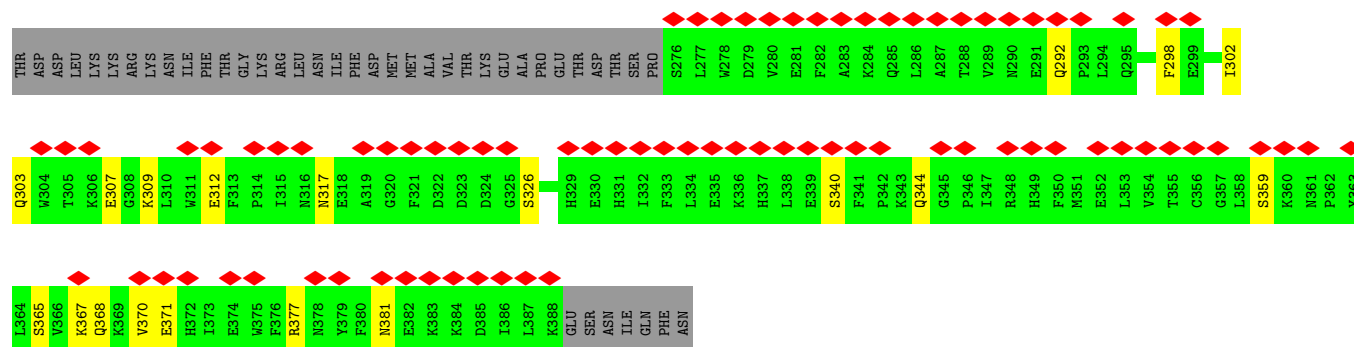
- Molecule 40: 28S ribosomal protein S29, mitochondrial



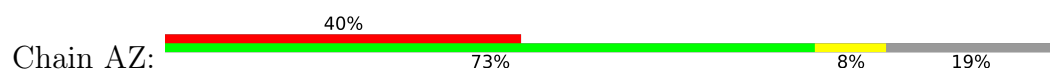
- Molecule 41: 28S ribosomal protein S31, mitochondrial



LEU	GLN	GLN	HIS	GLU	GLU	GLU	GLU	ARG	ALA	GLN	ARG	ASP	ALA	LYS	PRO	ARG	PHE	SER	ASN	ILE	ILE	SER	ASP	MET	LYS	VAL	ALA	ARG	ALA	VAL	ARG	PRO	GLU	ARG	GLN	ASP	PHE	GLN	GLY	TYR	ASP	ASN	TYR	PRO	GLY	GLN	GLY	GLU	GLY	PRO	PHE				
PRO	LYS	ARG	ARG	PRO	LEU	LYS	SER	LEU	GLU	ALA	THR	LEU	GLY	LEU	ARG	ARG	GLU	THR	PRO	LYS	THR	GLU	TYR	ALA	THR	VAL	GLU	ALA	ALA	SER	LEU	PRO	VAL	ASP	GLN	PHE	ASP	LYS	THR	LYS	SER	GLU	LEU	LEU	LEU	SER	GLN	PRO	THR						
GLY	THR	ASN	SER	VAL	ILE	CYS	SER	LYS	ASP	LYS	GLY	GLN	VAL	ARG	THR	GLU	THR	GLU	THR	GLY	SER	GLU	THR	GLY	GLU	THR	LYS	ASP	ALA	ALA	LEU	ASN	THR	VAL	GLY	ILE	ILE	LYS	GLY	LEU	GLY	THR	THR	LEU	ALA	SER	THR	TYR	THR						
MET	PHE	PRO	ARG	VAL	SER	THR	PHE	LEU	PRO	LEU	ARG	LEU	SER	PRO	HIS	PRO	LEU	SER	SER	GLY	SER	PRO	GLU	THR	ALA	ALA	ALA	LEU	LEU	THR	VAL	ARG	HIS	GLY	THR	VAL	ARG	TYR	ARG	SER	SER	ALA	LEU	LEU	ALA	ARG	THR	LYS	ASN	ASN	ILE	GLN	ARG	TYR	PRO



• Molecule 42: 28S ribosomal protein S33, mitochondrial



MET	SER	SER	LEU	S5	E6	Y7	A8	F9	R10	M11	S12	A16	R17	L18	F19	G20	E21	V22	T23	R24	P25	T26	N27	S28	K29	S30	M31	K32	V33	V34	K35	L36	F37	S38	E39	L40	P41	L42	T48	Y49	D50	P53	N54	H55	Y58	A59	E60	L65	R66	F67	L70	Y71	R72
D73	E74	D77	D80	E81	Q82	K87	L88	R89	G90	LYS	GLU	LYS	PRO	LYS	LYS	GLY	GLU	GLY	LYS	ARG	ALA	ALA	LYS	ARG	LYS	LYS	A1643	G1644	A1645	U1646	U1648	C1649	A1650	A1651	C	U	U	A	A	C	U1658	U1659	G1660	A1661	C	C1663	A1670	C	C	A			

• Molecule 43: mitochondrial tRNAVal

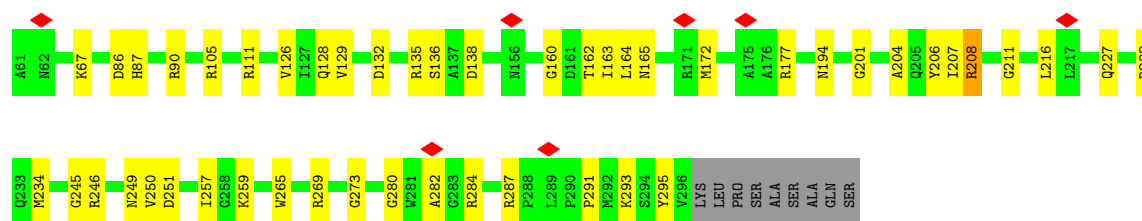


C	A1603	U1607	G1608	U1609	A1610	G1611	U1614	A1615	A1616	A1618	C	A1620	A1621	A1622	C1623	A1624	A1625	A1630	C1635	U1639	A1640	A1643	G1644	A1645	U1646	U	U1648	C1649	A1650	A1651	C	U	U	A	A	C	U1658	U1659	G1660	A1661	C	C1663	A1670	C	C	A
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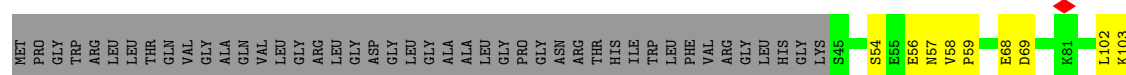
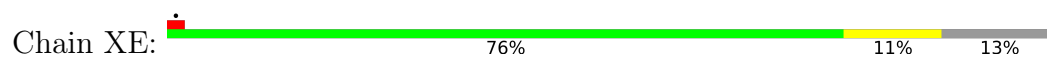
• Molecule 44: 39S ribosomal protein L2, mitochondrial



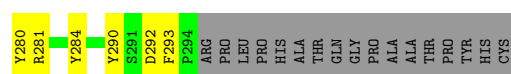
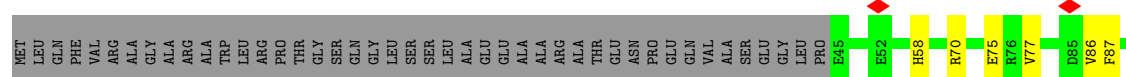
MET	ALA	LEU	CYS	ALA	LEU	THR	ARG	ALA	ARG	SER	LEU	ASN	ALA	PRO	THR	VAL	ALA	ALA	PRO	PRO	SER	LEU	PHE	PRO	ALA	ALA	GLN	MET	MET	ASN	ASN	GLY	LEU	LEU	GLN	PRO	PRO	SER	ALA	LEU	MET	LEU	LEU	PRO	PRO	CYS	ARG	PRO	VAL	VAL	THR	SER	VAL	ALA	LEU	ASN
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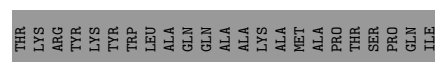
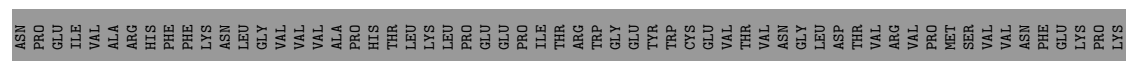
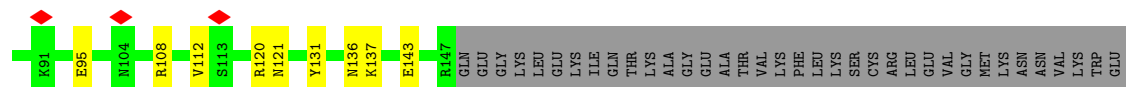
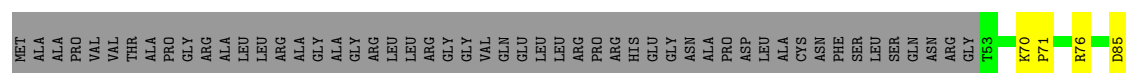
- Molecule 45: 39S ribosomal protein L3, mitochondrial



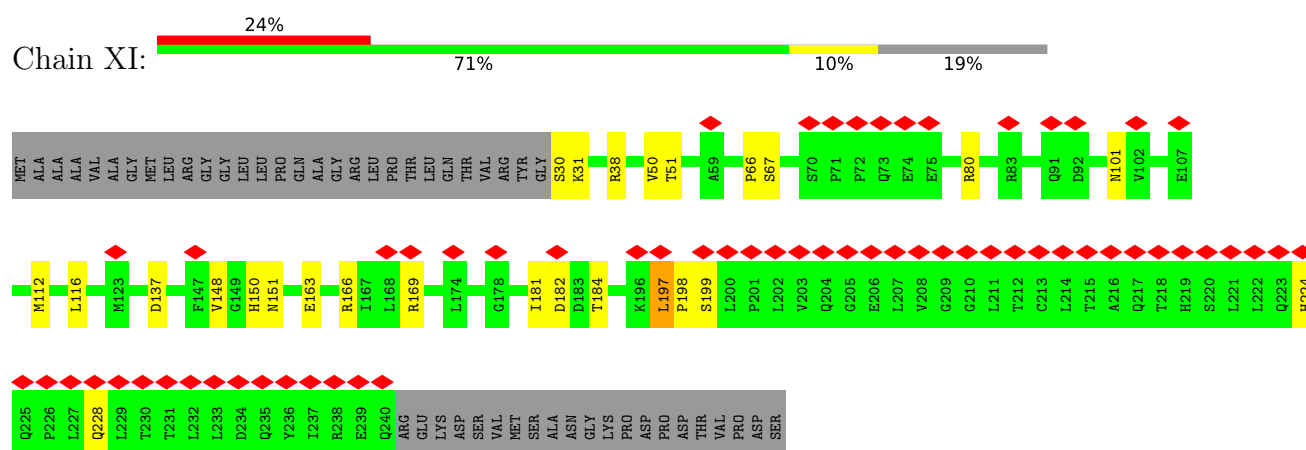
- Molecule 46: 39S ribosomal protein L4, mitochondrial



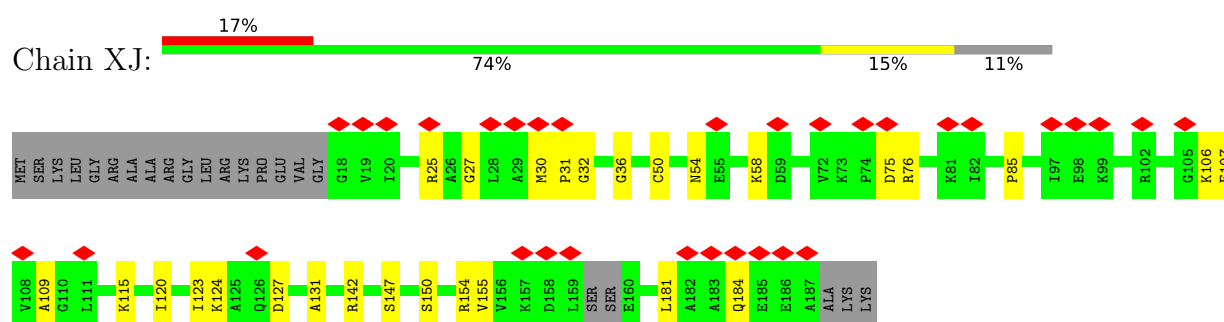
- Molecule 47: 39S ribosomal protein L9, mitochondrial



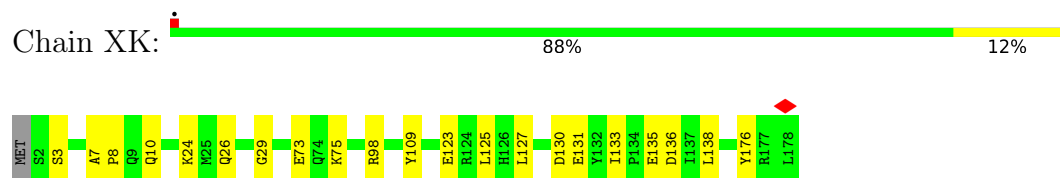
- Molecule 48: 39S ribosomal protein L10, mitochondrial



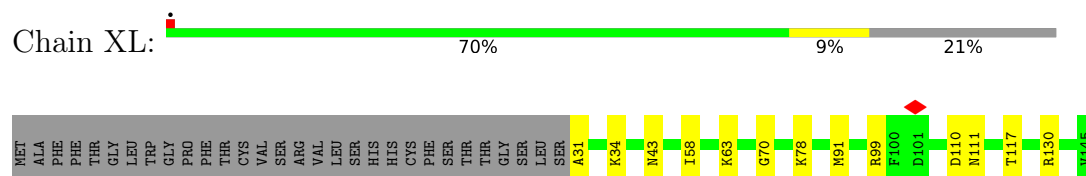
- Molecule 49: 39S ribosomal protein L11, mitochondrial



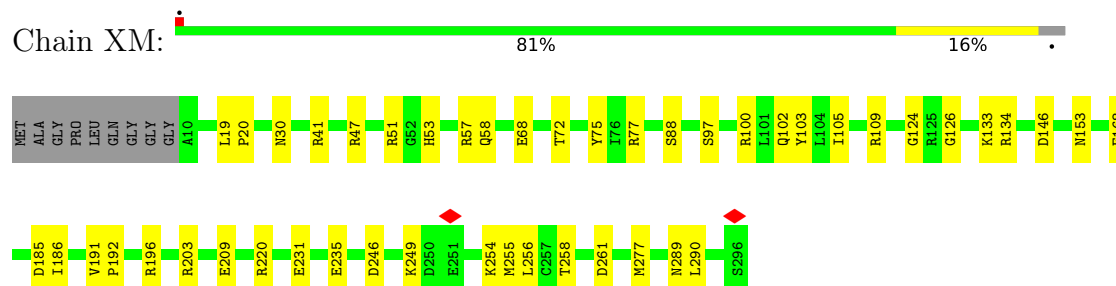
- Molecule 50: 39S ribosomal protein L13, mitochondrial



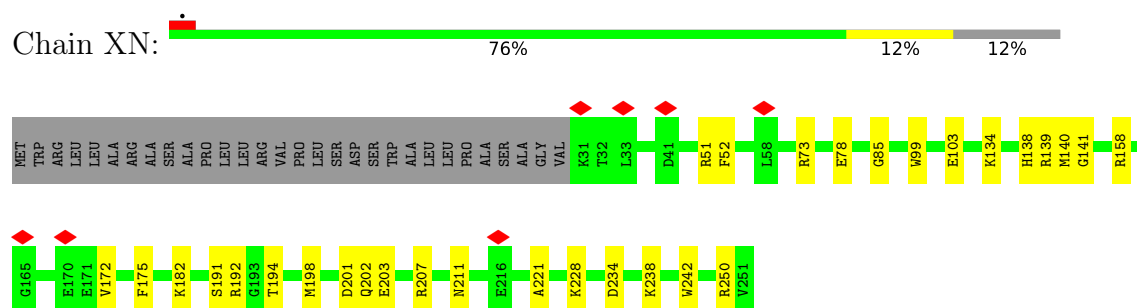
- Molecule 51: 39S ribosomal protein L14, mitochondrial



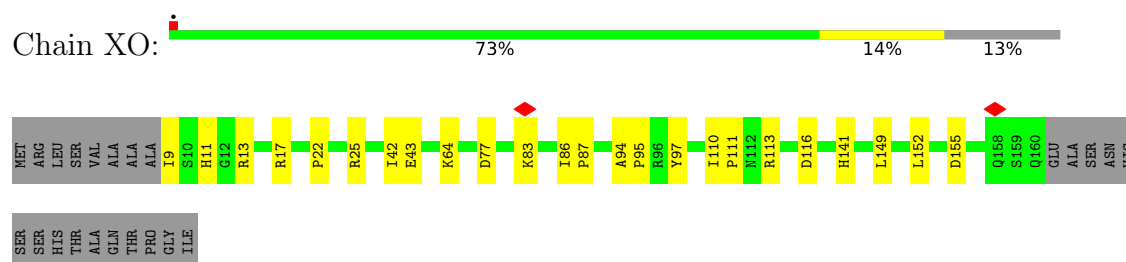
- Molecule 52: 39S ribosomal protein L15, mitochondrial



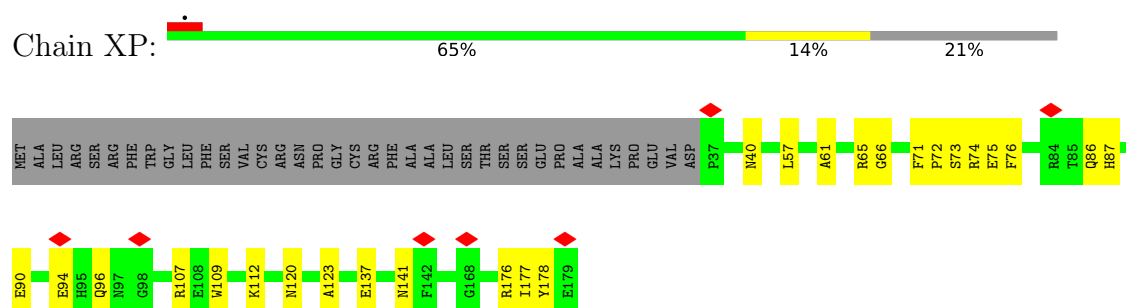
- Molecule 53: 39S ribosomal protein L16, mitochondrial



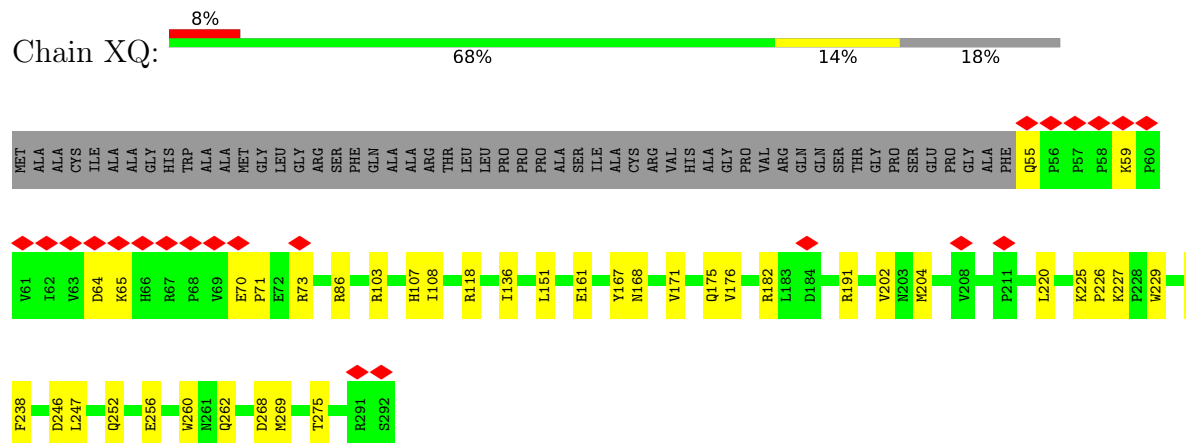
- Molecule 54: 39S ribosomal protein L17, mitochondrial



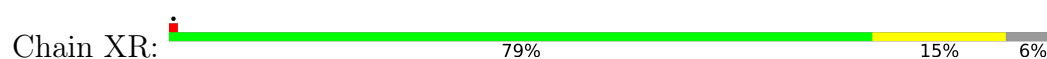
- Molecule 55: 39S ribosomal protein L18, mitochondrial



- Molecule 56: 39S ribosomal protein L19, mitochondrial

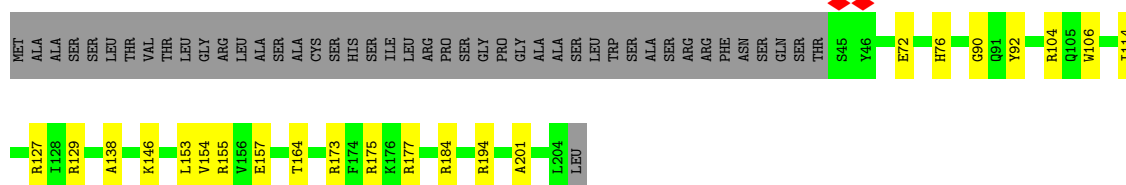


- Molecule 57: 39S ribosomal protein L20, mitochondrial

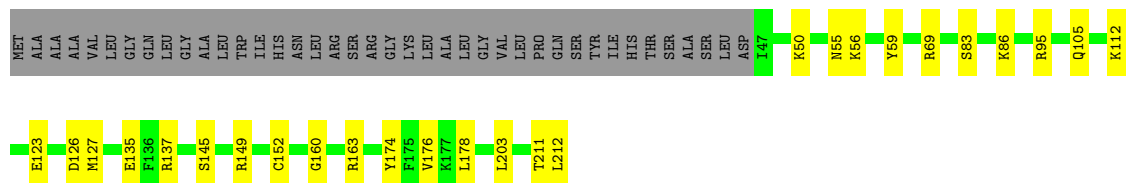




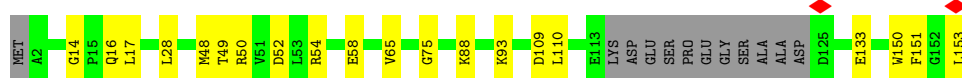
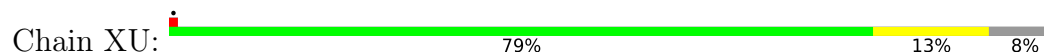
- Molecule 58: 39S ribosomal protein L21, mitochondrial



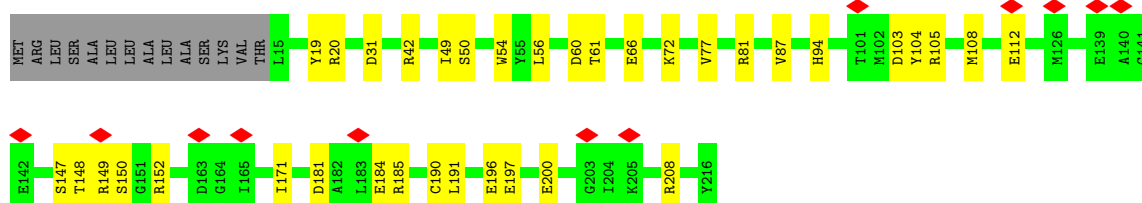
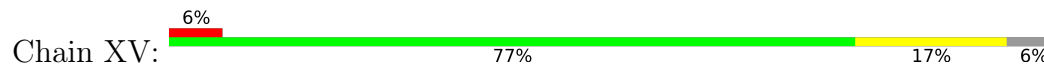
- Molecule 59: 39S ribosomal protein L22, mitochondrial



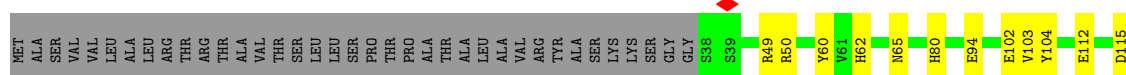
- Molecule 60: 39S ribosomal protein L23, mitochondrial

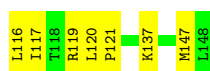


- Molecule 61: 39S ribosomal protein L24, mitochondrial



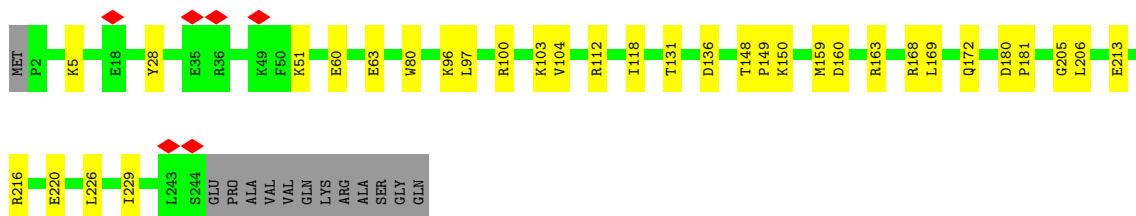
- Molecule 62: 39S ribosomal protein L27, mitochondrial





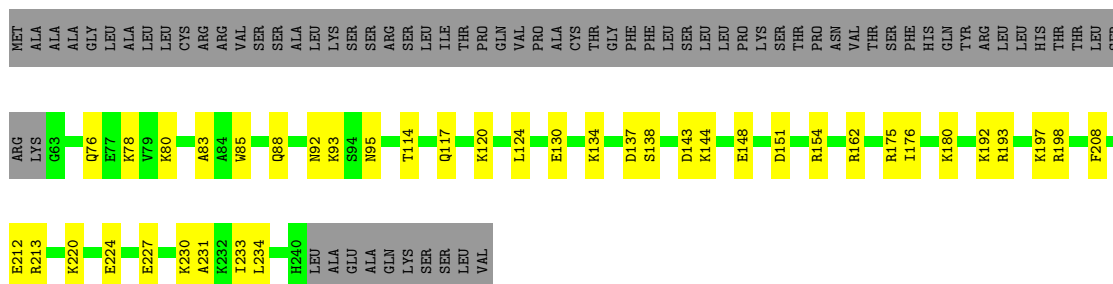
- Molecule 63: 39S ribosomal protein L28, mitochondrial

Chain XX: 82% 13% 5%



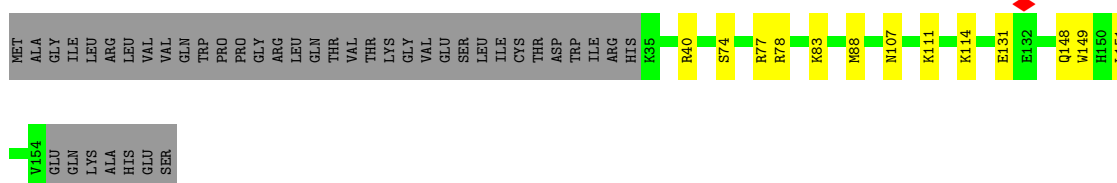
- Molecule 64: 39S ribosomal protein L47, mitochondrial

Chain XY: 55% 16% 29%



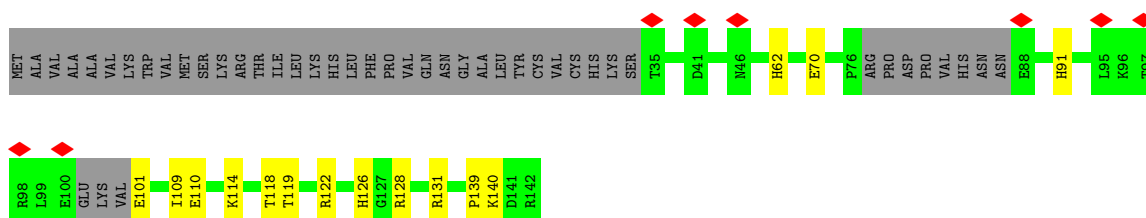
- Molecule 65: 39S ribosomal protein L30, mitochondrial

Chain XZ: 66% 8% 25%



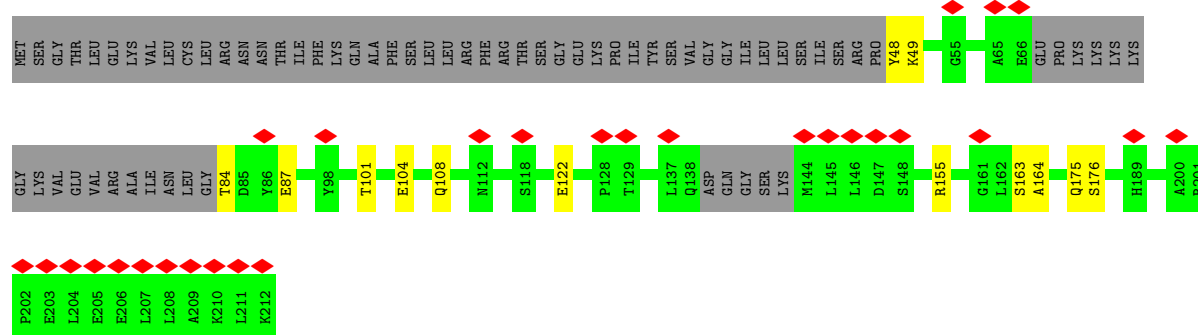
- Molecule 66: 39S ribosomal protein L42, mitochondrial

Chain a: 6% 58% 11% 32%

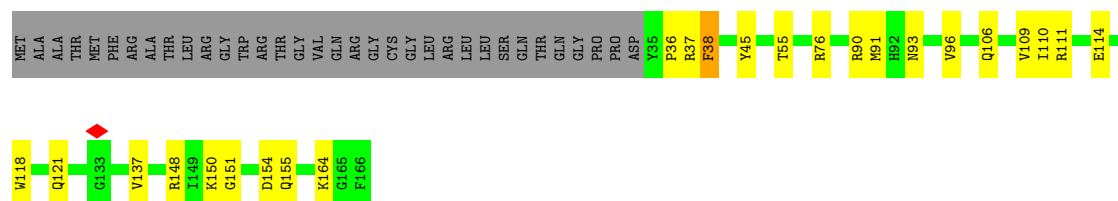


- Molecule 67: 39S ribosomal protein L43, mitochondrial

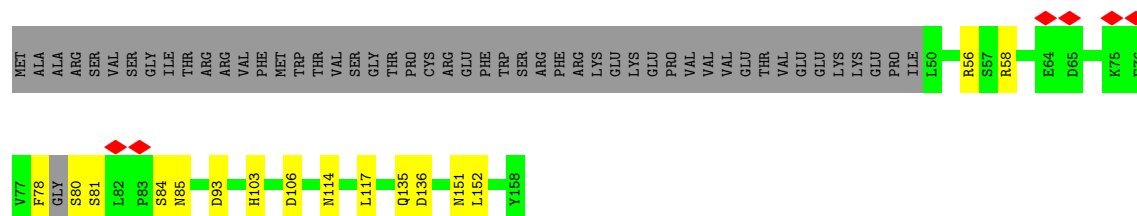
- Molecule 71: 39S ribosomal protein L48, mitochondrial



- Molecule 72: 39S ribosomal protein L49, mitochondrial

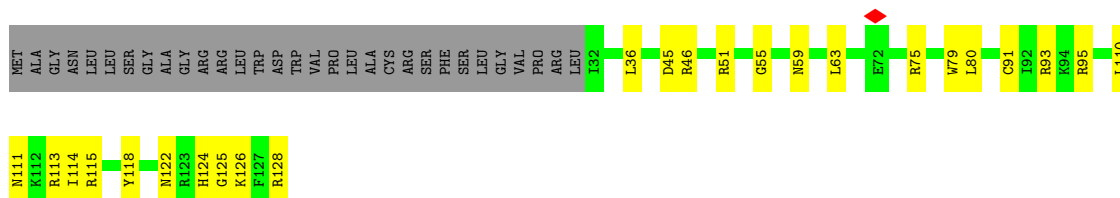


- Molecule 73: 39S ribosomal protein L50, mitochondrial

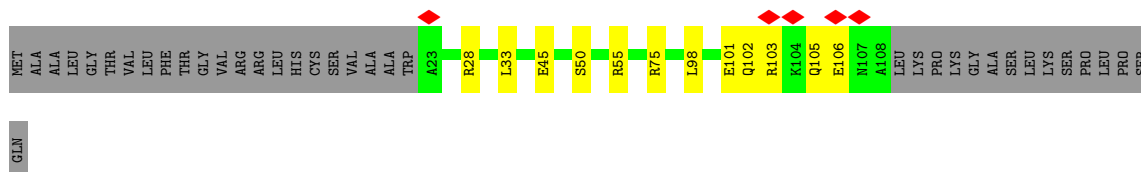


- Molecule 74: 39S ribosomal protein L51, mitochondrial

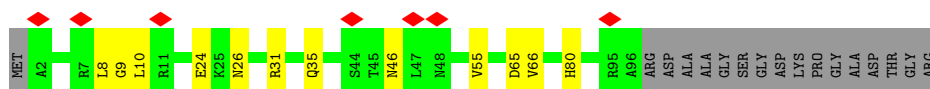




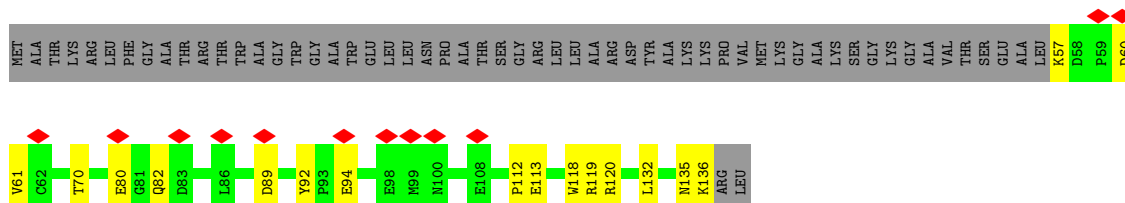
- Molecule 75: 39S ribosomal protein L52, mitochondrial



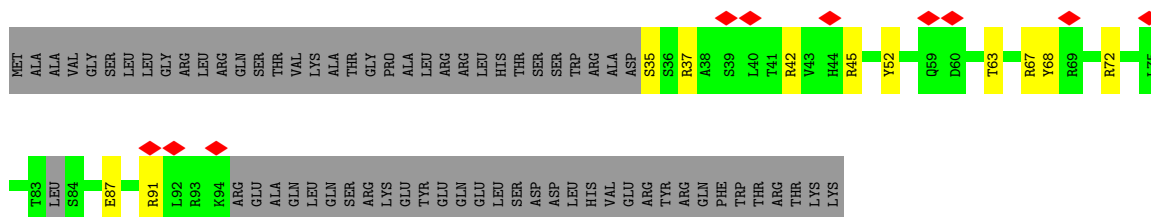
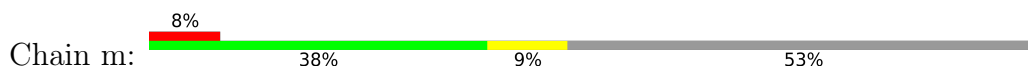
- Molecule 76: 39S ribosomal protein L53, mitochondrial



- Molecule 77: 39S ribosomal protein L54, mitochondrial



- Molecule 78: 39S ribosomal protein L55, mitochondrial

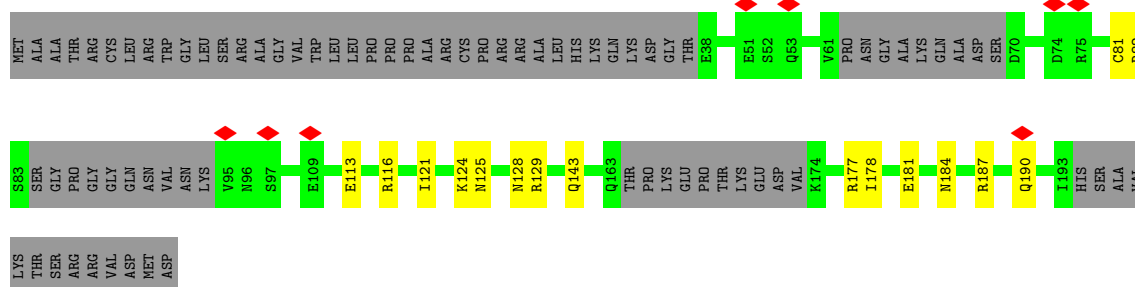


- Molecule 79: Ribosomal protein 63, mitochondrial

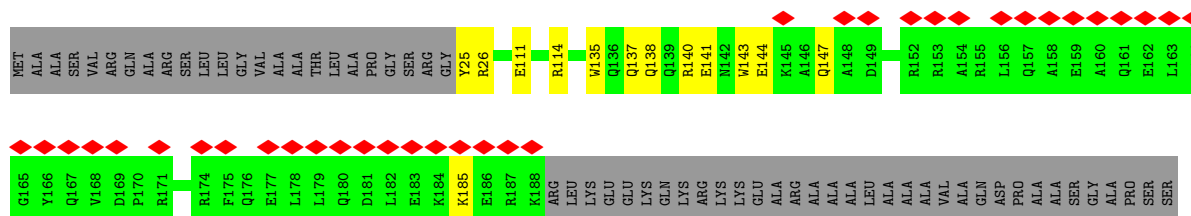




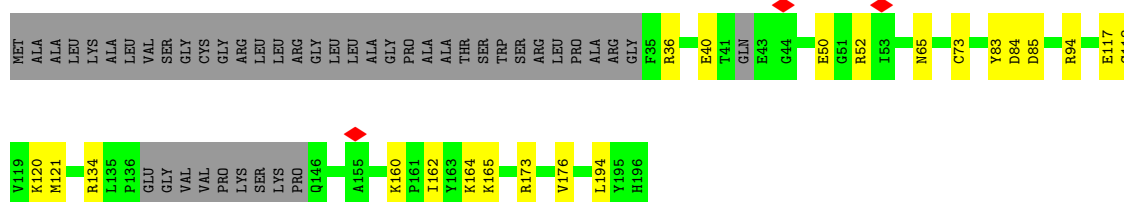
- Molecule 80: Peptidyl-tRNA hydrolase ICT1, mitochondrial



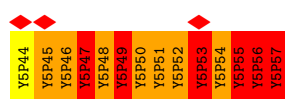
- Molecule 81: Growth arrest and DNA damage-inducible proteins-interacting protein 1



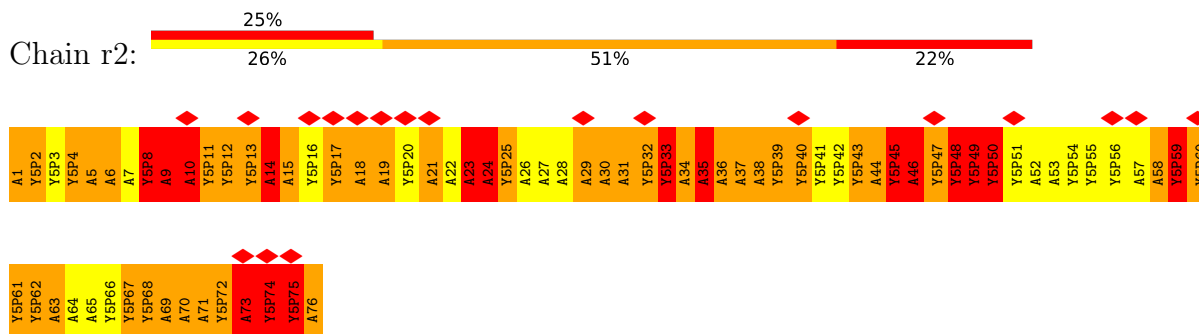
- Molecule 82: 39S ribosomal protein S18a, mitochondrial



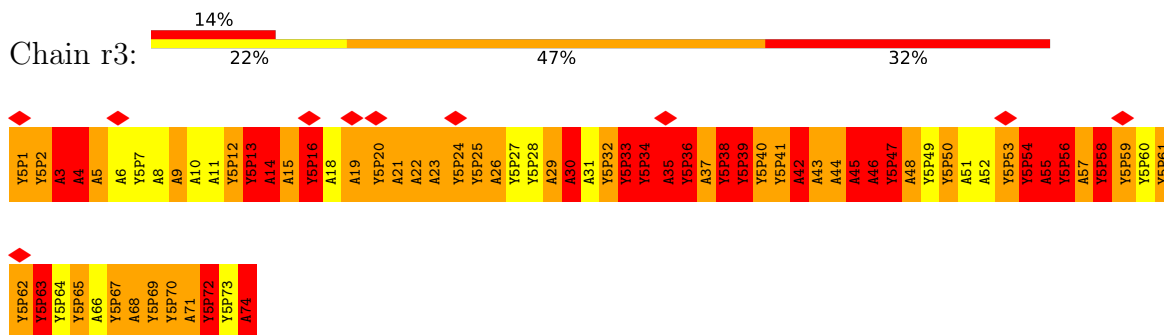
- Molecule 83: mRNA



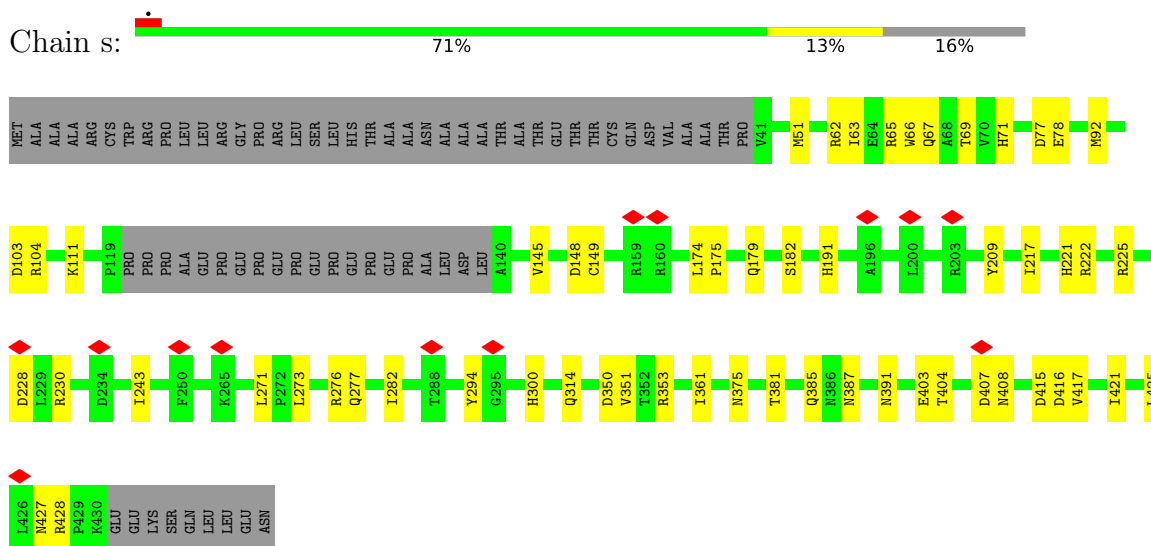
- Molecule 84: A/P-tRNA



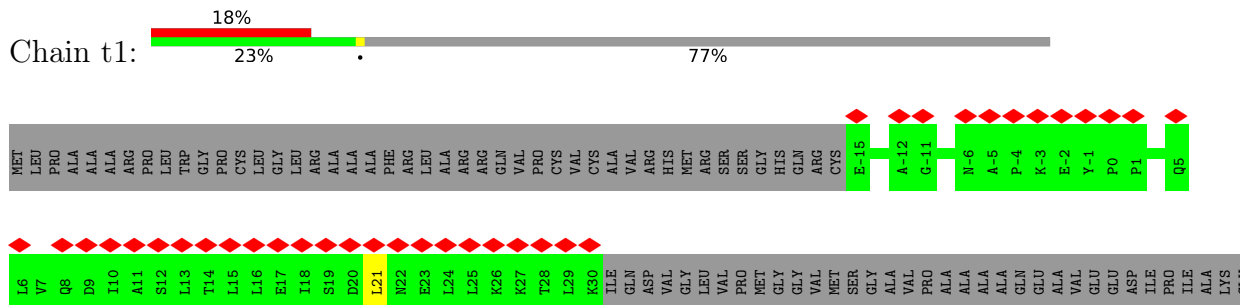
- Molecule 85: P/E-tRNA



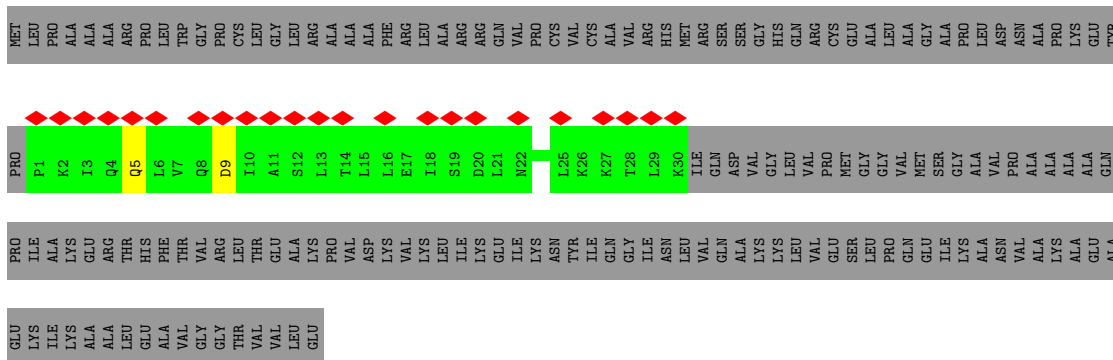
- Molecule 86: 39S ribosomal protein S30, mitochondrial



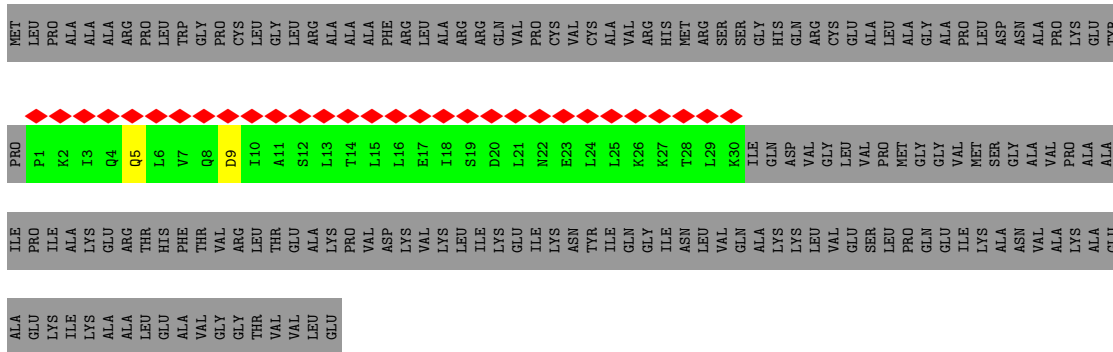
- Molecule 87: 39S ribosomal protein L12, mitochondrial



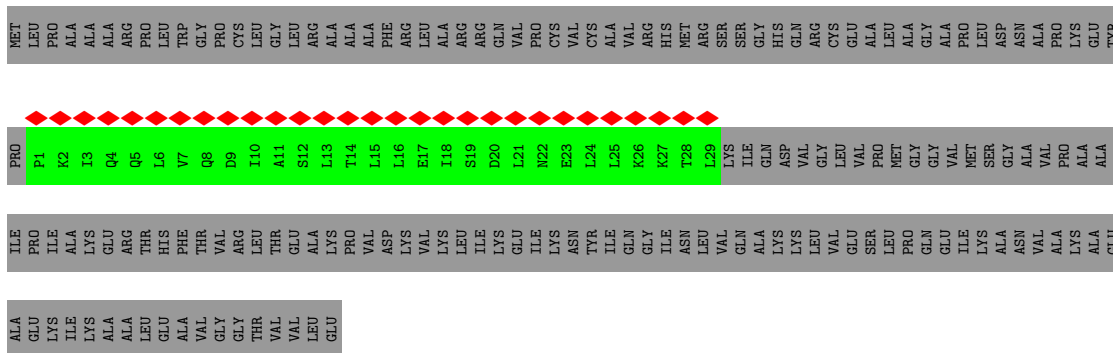
- Molecule 87: 39S ribosomal protein L12, mitochondrial



- Molecule 87: 39S ribosomal protein L12, mitochondrial



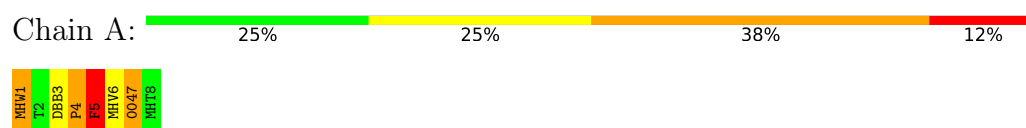
- Molecule 87: 39S ribosomal protein L12, mitochondrial



- Molecule 87: 39S ribosomal protein L12, mitochondrial

- Molecule 87: 39S ribosomal protein L12, mitochondrial

- Molecule 88: Quinupristin



4 Experimental information

Property	Value	Source
EM reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, Not provided	
Number of particles used	4966	Depositor
Resolution determination method	FSC 0.143 CUT-OFF	Depositor
CTF correction method	PHASE FLIPPING AND AMPLITUDE CORRECTION	Depositor
Microscope	FEI TITAN KRIOS	Depositor
Voltage (kV)	300	Depositor
Electron dose ($e^-/\text{\AA}^2$)	30	Depositor
Minimum defocus (nm)	Not provided	
Maximum defocus (nm)	Not provided	
Magnification	Not provided	
Image detector	GATAN K2 SUMMIT (4k x 4k)	Depositor
Maximum map value	0.208	Depositor
Minimum map value	-0.112	Depositor
Average map value	-0.000	Depositor
Map value standard deviation	0.004	Depositor
Recommended contour level	0.02	Depositor
Map size (Å)	546.0, 546.0, 546.0	wwPDB
Map dimensions	520, 520, 520	wwPDB
Map angles (°)	90.0, 90.0, 90.0	wwPDB
Pixel spacing (Å)	1.05, 1.05, 1.05	Depositor

5 Model quality

5.1 Standard geometry

Bond lengths and bond angles in the following residue types are not validated in this section: MHT, GTP, MHU, MHV, DOL, MG, 004, P5P, MHW, DBB, ZN, Y5P

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	0	0.17	0/895	0.30	0/1201
2	1	0.13	0/444	0.33	0/591
3	2	0.22	0/382	0.38	0/507
4	3	0.22	0/852	0.37	0/1136
5	4	0.14	0/349	0.31	0/461
6	5	0.13	0/3299	0.30	0/4495
7	6	0.13	0/3041	0.31	0/4137
8	7	0.13	0/2419	0.31	0/3267
9	8	0.11	0/1199	0.29	0/1612
10	9	0.13	0/1024	0.29	0/1379
11	XA	0.15	0/35615	0.27	0/55429
12	A0	0.11	0/1727	0.30	0/2338
13	A1	0.11	0/2276	0.28	0/3079
14	A2	0.13	0/939	0.30	0/1256
15	A3	0.15	0/621	0.35	0/820
16	A4	0.12	0/4559	0.32	0/6149
17	AA	0.10	0/21952	0.24	0/34164
18	AB	0.11	0/1819	0.28	0/2462
19	AC	0.10	0/1112	0.27	0/1505
20	AD	0.11	0/2768	0.29	0/3707
21	AE	0.11	0/989	0.31	0/1335
22	AF	0.10	0/1708	0.28	0/2291
23	AG	0.11	0/2559	0.29	0/3429
24	AH	0.12	0/1128	0.30	0/1529
25	AI	0.10	0/1031	0.26	0/1390
26	AJ	0.12	0/854	0.29	0/1148
27	AK	0.12	0/879	0.31	0/1182
28	AL	0.12	0/1406	0.30	0/1878
29	AM	0.11	0/941	0.29	0/1265
30	AN	0.11	0/864	0.30	0/1169
31	AO	0.10	0/1580	0.26	0/2150
32	AP	0.08	0/782	0.22	0/1050

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	# Z >5	RMSZ	# Z >5
33	AQ	0.12	0/746	0.27	0/993
34	AR	0.21	0/2103	0.44	0/2842
35	AS	0.12	0/1127	0.29	0/1518
36	AT	0.10	0/1361	0.30	0/1829
37	AU	0.13	0/1482	0.29	0/1987
38	AV	0.10	0/2925	0.29	0/3948
39	AW	0.10	0/778	0.25	0/1048
40	AX	0.11	0/2884	0.30	0/3903
41	AY	0.12	0/985	0.30	0/1329
42	AZ	0.11	0/748	0.27	0/1000
43	XB	0.09	0/1400	0.20	0/2168
44	XD	0.15	0/1879	0.34	0/2527
45	XE	0.15	0/2465	0.30	0/3344
46	XF	0.21	0/2071	0.34	0/2817
47	XH	0.13	0/798	0.31	0/1073
48	XI	0.14	0/1727	0.32	0/2340
49	XJ	0.10	0/1309	0.27	0/1764
50	XK	0.19	0/1495	0.30	0/2029
51	XL	0.14	0/904	0.27	0/1218
52	XM	0.18	0/2359	0.34	0/3185
53	XN	0.16	0/1825	0.32	0/2458
54	XO	0.16	0/1269	0.34	0/1708
55	XP	0.13	0/1190	0.30	0/1611
56	XQ	0.14	0/2026	0.32	0/2734
57	XR	0.23	0/1174	0.35	0/1572
58	XS	0.19	0/1311	0.33	0/1778
59	XT	0.19	0/1402	0.33	0/1886
60	XU	0.15	0/1200	0.30	0/1623
61	XV	0.13	0/1693	0.35	0/2297
62	XW	0.16	0/893	0.31	0/1204
63	XX	0.26	1/2090 (0.0%)	0.35	0/2825
64	XY	0.16	0/1571	0.34	0/2106
65	XZ	0.20	0/1003	0.36	0/1354
66	a	0.14	0/838	0.32	0/1138
67	b	0.18	0/1202	0.34	0/1626
68	c	0.15	0/2264	0.30	0/3059
69	d	0.12	0/1807	0.27	0/2450
70	e	0.11	0/1766	0.30	0/2382
71	f	0.10	0/1168	0.26	0/1573
72	g	0.27	0/1134	0.48	0/1547
73	h	0.13	0/905	0.35	0/1233
74	i	0.21	0/849	0.36	0/1135
75	j	0.17	0/703	0.35	0/947

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	# Z >5	RMSZ	# Z >5
76	k	0.12	0/743	0.28	0/1003
77	l	0.11	0/692	0.28	0/939
78	m	0.12	0/508	0.35	0/682
79	o	0.21	0/818	0.37	0/1097
80	p	0.12	0/1071	0.27	0/1433
81	q	0.14	0/1413	0.31	0/1906
82	r	0.15	0/1282	0.31	0/1734
86	s	0.14	0/3114	0.31	0/4225
87	t1	0.12	0/366	0.25	0/497
87	t2	0.08	0/238	0.20	0/319
87	t3	0.08	0/238	0.21	0/319
87	t4	0.07	0/229	0.20	0/308
87	t5	0.06	0/229	0.18	0/308
87	t6	0.10	0/213	0.23	0/286
88	A	0.65	0/13	0.82	0/15
All	All	0.14	1/176009 (0.0%)	0.29	0/249690

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
48	XI	0	1
70	e	0	1
88	A	2	3
All	All	2	5

All (1) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
63	XX	149	PRO	N-CD	7.78	1.58	1.47

There are no bond angle outliers.

All (2) chirality outliers are listed below:

Mol	Chain	Res	Type	Atom
88	A	2	THR	CB
88	A	4	PRO	CA

All (5) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
88	A	3	DBB	Peptide
88	A	4	PRO	Peptide
88	A	5	MHU	Peptide
48	XI	197	LEU	Peptide
70	e	265	LYS	Peptide

5.2 Too-close contacts

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	0	880	903	903	17	0
2	1	439	480	480	10	0
3	2	376	406	406	14	0
4	3	831	883	883	18	0
5	4	341	362	362	7	0
6	5	3204	3201	3201	33	0
7	6	2947	2840	2840	35	0
8	7	2365	2372	2371	36	0
9	8	1175	1202	1202	18	0
10	9	996	987	987	17	0
11	XA	31833	16159	16168	408	0
12	A0	1684	1685	1685	16	0
13	A1	2230	2261	2261	68	0
14	A2	925	964	964	15	0
15	A3	610	682	682	11	0
16	A4	4470	4485	4486	131	0
17	AA	19628	9964	9971	170	0
18	AB	1776	1769	1769	17	0
19	AC	1082	1088	1088	27	0
20	AD	2716	2783	2784	33	0
21	AE	972	1001	1001	18	0
22	AF	1668	1716	1716	22	0
23	AG	2505	2490	2490	32	0
24	AH	1105	1136	1136	38	0
25	AI	1011	1052	1052	6	0
26	AJ	838	887	887	14	0
27	AK	861	885	885	13	0
28	AL	1382	1473	1472	20	0
29	AM	920	951	951	11	0

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Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
30	AN	846	908	908	13	0
31	AO	1528	1490	1490	18	0
32	AP	765	796	796	8	0
33	AQ	734	749	749	6	0
34	AR	2060	2074	2074	25	0
35	AS	1100	1103	1103	13	0
36	AT	1330	1342	1342	11	0
37	AU	1461	1471	1471	17	0
38	AV	2867	2862	2862	25	0
39	AW	766	785	785	5	0
40	AX	2814	2803	2802	27	0
41	AY	956	912	911	23	0
42	AZ	731	734	734	8	0
43	XB	1255	635	640	11	0
44	XD	1842	1896	1896	40	0
45	XE	2396	2402	2402	29	0
46	XF	2013	2044	2044	33	0
47	XH	784	832	832	10	0
48	XI	1691	1783	1783	19	0
49	XJ	1291	1367	1364	20	0
50	XK	1451	1448	1448	16	0
51	XL	889	941	941	8	0
52	XM	2305	2376	2376	39	0
53	XN	1778	1808	1808	21	0
54	XO	1245	1283	1283	18	0
55	XP	1164	1162	1162	19	0
56	XQ	1978	2022	2022	30	0
57	XR	1153	1214	1214	20	0
58	XS	1284	1354	1354	17	0
59	XT	1368	1410	1410	23	0
60	XU	1171	1164	1164	18	0
61	XV	1648	1656	1654	43	0
62	XW	871	898	898	18	0
63	XX	2035	2054	2054	23	0
64	XY	1534	1575	1575	36	0
65	XZ	978	1030	1030	11	0
66	a	813	777	777	12	0
67	b	1178	1180	1180	16	0
68	c	2217	2220	2220	27	0
69	d	1758	1743	1742	61	0
70	e	1731	1732	1732	20	0
71	f	1149	1164	1164	13	0

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Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
72	g	1097	1086	1085	21	0
73	h	882	866	867	11	0
74	i	827	857	857	21	0
75	j	689	678	678	13	0
76	k	732	745	745	10	0
77	l	673	654	653	15	0
78	m	500	525	525	9	0
79	o	797	804	804	20	0
80	p	1058	1083	1083	13	0
81	q	1379	1359	1359	9	0
82	r	1247	1267	1267	17	0
83	r1	252	0	169	21	0
84	r2	1485	0	835	49	0
85	r3	1420	0	807	60	0
86	s	3036	3022	3022	37	0
87	t1	354	379	374	0	0
87	t2	238	268	270	1	0
87	t3	238	268	270	1	0
87	t4	229	255	257	0	0
87	t5	229	255	257	1	0
87	t6	214	236	236	1	0
88	A	73	67	64	7	0
89	0	1	0	0	0	0
89	4	1	0	0	0	0
89	AB	1	0	0	0	0
89	AO	1	0	0	0	0
89	AP	1	0	0	0	0
89	AT	1	0	0	0	0
89	r	1	0	0	0	0
90	AA	46	0	0	0	0
90	XA	141	0	0	0	0
90	XD	1	0	0	0	0
90	XE	1	0	0	0	0
90	XI	1	0	0	0	0
90	XM	2	0	0	0	0
90	XW	1	0	0	0	0
90	g	1	0	0	0	0
90	o	1	0	0	0	0
91	XA	48	50	50	1	0
92	AX	32	10	12	1	0
All	All	170629	143000	144825	1888	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 6.

All (1888) 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)
61:XV:72:LYS:HE2	69:d:256:TYR:CE1	1.42	1.53
16:A4:449:GLY:HA2	41:AY:292:GLN:NE2	1.33	1.40
61:XV:72:LYS:CD	69:d:256:TYR:CZ	2.21	1.22
61:XV:72:LYS:CE	69:d:256:TYR:CE1	2.24	1.21
61:XV:72:LYS:HE2	69:d:256:TYR:CD1	1.79	1.15
61:XV:72:LYS:HD3	69:d:256:TYR:CZ	1.86	1.09
1:O:132:LYS:HE2	69:d:288:LYS:NZ	1.69	1.07
13:A1:162:SER:OG	16:A4:134:GLU:CG	2.03	1.05
16:A4:449:GLY:CA	41:AY:292:GLN:NE2	2.22	1.03
61:XV:105:ARG:HH12	69:d:171:ASP:HA	1.21	1.03
60:XU:151:PHE:CD2	69:d:222:LEU:HD11	1.94	1.03
85:r3:47:Y5P:H2'	85:r3:57:P5P:H1'	1.39	1.01
13:A1:162:SER:OG	16:A4:134:GLU:HG3	1.60	1.00
61:XV:72:LYS:HD2	69:d:256:TYR:CE2	1.97	0.98
11:XA:1881:A:OP2	72:g:111:ARG:NH2	1.97	0.97
17:AA:701:G:N2	17:AA:841:A:O2'	1.99	0.95
11:XA:3063:G:O2'	11:XA:3066:C:OP2	1.84	0.95
4:3:113:ARG:NH1	52:XM:75:TYR:O	1.99	0.95
11:XA:2724:G:OP1	46:XF:131:LYS:NZ	2.01	0.94
11:XA:2954:C:O2	53:XN:182:LYS:NZ	2.01	0.94
1:O:132:LYS:HE2	69:d:288:LYS:HZ3	1.28	0.93
11:XA:1828:A:N6	11:XA:2683:C:O2	2.02	0.92
13:A1:134:PRO:HB3	16:A4:58:VAL:HG23	1.53	0.91
61:XV:72:LYS:HD2	69:d:256:TYR:CZ	1.98	0.91
11:XA:1777:A:N6	11:XA:1780:U:OP2	2.04	0.90
74:i:95:ARG:NH1	74:i:111:ASN:OD1	2.04	0.90
16:A4:108:LEU:HD21	20:AD:154:VAL:HG21	1.50	0.90
13:A1:162:SER:OG	16:A4:134:GLU:HG2	1.71	0.90
85:r3:25:Y5P:H4A	85:r3:42:P5P:H6	1.51	0.90
51:XL:31:ALA:N	51:XL:91:MET:SD	2.46	0.89
51:XL:70:GLY:O	51:XL:130:ARG:NH1	2.04	0.89
61:XV:72:LYS:HD3	69:d:256:TYR:OH	1.71	0.89
10:9:26:GLY:O	10:9:31:ARG:NH1	2.05	0.89
70:e:264:LEU:O	70:e:273:ARG:NH2	2.07	0.88
13:A1:154:THR:OG1	24:AH:171:GLU:OE2	1.90	0.88
58:XS:138:ALA:O	67:b:127:GLN:NE2	2.07	0.88
22:AF:79:ALA:O	23:AG:312:GLN:NE2	2.06	0.88
48:XI:51:THR:O	53:XN:250:ARG:NH1	2.05	0.88

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
16:A4:449:GLY:HA2	41:AY:292:GLN:CD	2.00	0.88
11:XA:2145:G:O2'	11:XA:2147:G:OP1	1.91	0.87
2:1:34:ARG:NH2	2:1:38:ARG:O	2.08	0.87
11:XA:1912:A:OP1	11:XA:1986:A:N6	2.09	0.86
68:c:80:GLN:O	68:c:210:ARG:NH2	2.08	0.86
84:r2:45:Y5P:O2'	84:r2:46:P5P:OP1	1.94	0.86
6:5:30:ALA:N	44:XD:201:GLY:O	2.09	0.85
29:AM:93:LEU:O	34:AR:175:ARG:NH2	2.09	0.85
13:A1:52:ALA:HB2	16:A4:94:TYR:HB2	1.59	0.85
7:6:368:ARG:NH2	11:XA:2859:A:OP2	2.09	0.85
11:XA:2367:A:O3'	64:XY:120:LYS:NZ	2.09	0.85
18:AB:103:GLU:OE2	35:AS:52:ARG:NH2	2.10	0.85
31:AO:185:SER:O	34:AR:183:LYS:NZ	2.09	0.85
85:r3:54:Y5P:HC	85:r3:56:Y5P:H5	1.57	0.85
1:0:95:ARG:NH1	11:XA:1821:A:OP2	2.10	0.85
5:4:84:ARG:NE	11:XA:3188:U:OP2	2.10	0.84
83:r1:46:Y5P:H5	85:r3:35:P5P:H6	1.58	0.84
11:XA:2154:A:N7	79:o:30:ARG:NH2	2.25	0.84
11:XA:2191:A:N6	11:XA:2198:A:OP2	2.11	0.84
27:AK:90:ARG:NH2	27:AK:95:SER:O	2.11	0.84
52:XM:72:THR:OG1	52:XM:77:ARG:NH2	2.11	0.84
11:XA:2864:U:O5'	62:XW:50:ARG:NH1	2.10	0.83
9:8:87:ASP:O	78:m:45:ARG:NH1	2.11	0.83
16:A4:108:LEU:HD21	20:AD:154:VAL:CG2	2.08	0.83
17:AA:1557:A:O2'	26:AJ:72:LYS:NZ	2.11	0.83
11:XA:2726:C:N4	11:XA:2988:C:O2	2.11	0.83
25:AI:71:SER:O	25:AI:74:ARG:NH1	2.12	0.83
60:XU:16:GLN:NE2	60:XU:17:LEU:O	2.12	0.83
54:XO:77:ASP:OD1	54:XO:83:LYS:NZ	2.11	0.82
11:XA:2517:U:OP1	44:XD:287:ARG:NH2	2.13	0.82
61:XV:72:LYS:CD	69:d:256:TYR:CE1	2.58	0.82
37:AU:126:GLN:OE1	37:AU:129:ARG:NH2	2.12	0.82
46:XF:220:ASP:O	46:XF:245:ALA:N	2.12	0.82
13:A1:312:TYR:OH	40:AX:338:ASP:O	1.97	0.82
17:AA:1530:A:OP1	38:AV:64:LYS:NZ	2.13	0.81
67:b:132:PRO:O	67:b:136:LYS:NZ	2.13	0.81
2:1:53:ARG:NH2	11:XA:2879:A:O2'	2.13	0.81
4:3:104:ARG:NH1	4:3:160:LYS:O	2.14	0.81
11:XA:1765:C:OP2	74:i:75:ARG:NH2	2.13	0.81
11:XA:1985:G:OP1	44:XD:90:ARG:NH2	2.13	0.81
29:AM:55:ASP:OD2	36:AT:146:GLN:NE2	2.13	0.81

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
86:s:51:MET:O	86:s:62:ARG:NH2	2.12	0.81
17:AA:1017:A:OP2	17:AA:1018:G:O2'	1.99	0.81
16:A4:449:GLY:HA2	41:AY:292:GLN:HE21	1.41	0.80
17:AA:728:C:OP1	30:AN:5:ARG:NH2	2.14	0.80
11:XA:1680:A:OP1	64:XY:230:LYS:NZ	2.14	0.80
67:b:9:ARG:NH1	67:b:111:ASP:O	2.13	0.80
1:0:139:ARG:NH2	11:XA:2322:C:OP1	2.14	0.80
1:0:181:ARG:NH1	1:0:186:THR:O	2.15	0.80
11:XA:1773:A:OP2	68:c:31:VAL:N	2.13	0.80
11:XA:2347:C:O3'	69:d:69:LYS:NZ	2.14	0.80
13:A1:129:PHE:O	16:A4:63:LYS:NZ	2.15	0.80
52:XM:53:HIS:O	52:XM:58:GLN:NE2	2.14	0.80
14:A2:38:ARG:NH2	17:AA:1184:U:OP1	2.15	0.80
17:AA:826:A:OP1	26:AJ:55:ARG:NH1	2.13	0.80
38:AV:321:GLU:O	38:AV:326:LYS:NZ	2.15	0.80
16:A4:477:TRP:CZ3	24:AH:52:VAL:HG12	2.17	0.80
49:XJ:25:ARG:NH2	77:l:57:LYS:O	2.15	0.80
11:XA:2052:A:OP1	72:g:150:LYS:NZ	2.15	0.80
11:XA:2563:U:OP1	44:XD:284:ARG:NH1	2.13	0.79
8:7:322:LYS:NZ	54:XO:155:ASP:OD2	2.15	0.79
61:XV:105:ARG:NH1	69:d:171:ASP:HA	1.97	0.79
11:XA:3078:C:N4	11:XA:3079:G:O6	2.15	0.79
34:AR:176:GLU:OE2	34:AR:182:ARG:NE	2.15	0.79
9:8:164:ARG:NH1	71:f:84:THR:O	2.16	0.79
43:XB:1610:A:N6	43:XB:1623:G:O6	2.14	0.79
3:2:82:ARG:NH2	11:XA:1790:A:OP1	2.14	0.79
7:6:367:ASP:OD1	7:6:370:ARG:NH1	2.15	0.79
11:XA:2863:U:O2	11:XA:2869:A:N6	2.15	0.79
16:A4:62:LYS:HB2	24:AH:70:ASP:HA	1.64	0.79
52:XM:30:ASN:OD1	79:o:86:ARG:NH1	2.16	0.79
11:XA:1882:A:N6	11:XA:1893:A:O4'	2.15	0.79
17:AA:860:A:N7	17:AA:919:A:O2'	2.15	0.78
11:XA:3068:G:OP2	11:XA:3068:G:N2	2.17	0.78
11:XA:3173:G:OP2	82:r:160:LYS:NZ	2.17	0.78
23:AG:198:ARG:N	23:AG:246:ARG:O	2.17	0.78
26:AJ:84:ARG:NH1	26:AJ:85:LEU:O	2.16	0.78
11:XA:1832:A:N6	11:XA:1837:C:OP2	2.17	0.78
11:XA:2458:A:OP2	54:XO:9:ILE:N	2.17	0.78
17:AA:1508:C:N3	17:AA:1542:U:O4	2.16	0.78
69:d:288:LYS:O	69:d:290:GLU:N	2.15	0.78
58:XS:129:ARG:NH2	58:XS:154:VAL:O	2.16	0.78

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
60:XU:151:PHE:CE2	69:d:222:LEU:HD11	2.18	0.78
6:5:381:LEU:O	6:5:407:LYS:NZ	2.16	0.78
85:r3:13:Y5P:H2	85:r3:14:P5P:H6	1.65	0.78
11:XA:1939:G:O5'	44:XD:259:LYS:NZ	2.16	0.78
64:XY:76:GLN:NE2	64:XY:78:LYS:O	2.16	0.78
11:XA:1700:U:O4	64:XY:193:ARG:NH2	2.17	0.77
61:XV:72:LYS:CE	69:d:256:TYR:CD1	2.59	0.77
22:AF:126:TYR:O	22:AF:134:GLN:NE2	2.17	0.77
46:XF:277:ASP:O	72:g:90:ARG:NH2	2.16	0.77
45:XE:69:ASP:OD1	45:XE:154:ARG:NH1	2.17	0.77
84:r2:17:Y5P:O2'	84:r2:19:P5P:OP2	2.03	0.77
11:XA:1723:A:N6	11:XA:1726:C:OP2	2.17	0.77
14:A2:42:GLU:N	22:AF:241:TRP:O	2.17	0.77
16:A4:59:ILE:HG23	24:AH:72:LEU:HD11	1.67	0.77
23:AG:276:ARG:NH1	23:AG:373:ASP:OD2	2.18	0.77
46:XF:223:HIS:O	81:q:26:ARG:NH2	2.18	0.77
81:q:143:TRP:O	81:q:147:GLN:NE2	2.18	0.77
11:XA:1696:C:OP2	64:XY:180:LYS:NZ	2.18	0.77
17:AA:868:C:OP2	17:AA:870:C:N4	2.18	0.77
18:AB:180:ARG:NH1	18:AB:185:PRO:O	2.18	0.76
6:5:384:GLN:OE1	11:XA:2395:A:O2'	2.03	0.76
11:XA:1957:A:O4'	59:XT:163:ARG:NH1	2.18	0.76
17:AA:1429:C:O2	17:AA:1460:C:N4	2.19	0.76
21:AE:42:LEU:O	37:AU:184:ARG:NE	2.18	0.76
63:XX:118:ILE:O	63:XX:168:ARG:NH1	2.19	0.76
11:XA:3220:A:OP1	45:XE:260:LYS:NZ	2.18	0.76
77:l:135:ASN:OD1	77:l:136:LYS:N	2.18	0.76
57:XR:93:CYS:O	58:XS:146:LYS:NZ	2.18	0.76
44:XD:128:GLN:NE2	44:XD:129:VAL:O	2.19	0.76
4:3:124:ARG:NH2	11:XA:2868:C:OP1	2.19	0.76
9:8:93:LYS:O	9:8:99:ARG:NE	2.19	0.76
10:9:22:THR:OG1	10:9:36:ARG:NH1	2.19	0.76
11:XA:2537:G:O2'	11:XA:2634:U:OP2	2.03	0.76
50:XK:3:SER:O	68:c:302:ARG:NH2	2.19	0.76
52:XM:203:ARG:NH2	52:XM:261:ASP:O	2.19	0.76
64:XY:124:LEU:HD21	69:d:68:ARG:HH11	1.50	0.76
11:XA:2581:A:O2'	11:XA:2583:C:N4	2.19	0.76
13:A1:69:VAL:HA	16:A4:81:ASP:OD2	1.85	0.76
17:AA:906:C:OP1	20:AD:117:ARG:NE	2.19	0.76
59:XT:69:ARG:NH1	69:d:228:PHE:O	2.19	0.76
71:f:101:THR:OG1	78:m:67:ARG:NH1	2.18	0.76

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
13:A1:134:PRO:HG3	16:A4:60:PRO:HD3	1.68	0.76
43:XB:1635:C:OP1	71:f:108:GLN:NE2	2.18	0.76
64:XY:151:ASP:OD1	64:XY:154:ARG:NH2	2.19	0.76
10:9:127:LEU:O	10:9:134:ASN:ND2	2.18	0.75
16:A4:269:HIS:O	16:A4:270:ARG:NE	2.20	0.75
1:0:132:LYS:HE2	69:d:288:LYS:HZ1	1.51	0.75
12:A0:90:ASP:OD1	31:AO:215:ARG:NH1	2.18	0.75
53:XN:201:ASP:OD1	53:XN:202:GLN:N	2.19	0.75
22:AF:122:GLN:NE2	22:AF:138:GLU:O	2.19	0.75
20:AD:127:ASN:O	42:AZ:72:ARG:NH1	2.19	0.75
17:AA:1558:A:N6	84:r2:36:P5P:O2'	2.20	0.75
56:XQ:118:ARG:NH2	56:XQ:202:VAL:O	2.20	0.75
27:AK:116:ASP:OD1	27:AK:125:ARG:NH2	2.20	0.74
11:XA:1958:G:OP2	59:XT:160:GLY:N	2.18	0.74
40:AX:206:GLU:OE1	40:AX:249:ARG:NH1	2.20	0.74
67:b:50:GLU:OE2	67:b:57:ARG:NH2	2.19	0.74
11:XA:1689:C:OP2	63:XX:5:LYS:NZ	2.19	0.74
15:A3:175:ARG:NH1	17:AA:1541:U:O3'	2.20	0.74
23:AG:103:ASP:OD1	23:AG:106:ARG:NH2	2.19	0.74
37:AU:138:GLU:OE2	37:AU:141:ARG:NH2	2.21	0.74
9:8:173:LYS:NZ	71:f:175:GLN:OE1	2.21	0.74
24:AH:74:LYS:N	24:AH:175:THR:O	2.20	0.74
61:XV:150:SER:O	61:XV:152:ARG:NH1	2.20	0.74
11:XA:2457:A:O2'	54:XO:17:ARG:NH2	2.19	0.74
22:AF:176:ASP:OD1	22:AF:179:ARG:NH2	2.19	0.74
38:AV:34:TYR:OH	38:AV:371:ASP:OD2	2.06	0.74
16:A4:108:LEU:CD2	20:AD:154:VAL:HG21	2.18	0.74
4:3:113:ARG:NH2	11:XA:1750:G:OP2	2.20	0.74
40:AX:53:GLU:N	40:AX:67:HIS:O	2.21	0.74
9:8:110:GLU:OE2	9:8:114:ARG:NE	2.21	0.73
58:XS:194:ARG:NH2	67:b:13:SER:OG	2.21	0.73
23:AG:117:PHE:O	23:AG:122:ARG:NH1	2.21	0.73
11:XA:1787:G:N2	11:XA:1790:A:OP2	2.22	0.73
17:AA:869:C:OP2	31:AO:97:ARG:NH2	2.22	0.73
1:0:133:GLU:OE1	1:0:179:ARG:NH2	2.22	0.73
4:3:169:ARG:NH2	11:XA:1892:A:OP1	2.21	0.73
13:A1:53:LEU:O	16:A4:93:PRO:HB2	1.88	0.73
16:A4:477:TRP:HZ3	24:AH:52:VAL:HG12	1.50	0.73
31:AO:196:GLY:O	34:AR:146:LYS:NZ	2.22	0.73
58:XS:72:GLU:O	58:XS:76:HIS:ND1	2.20	0.73
11:XA:2524:A:OP1	44:XD:67:LYS:NZ	2.20	0.73

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:2510:U:OP2	11:XA:2539:A:N6	2.21	0.73
17:AA:949:U:O3'	30:AN:29:ARG:NH1	2.22	0.73
17:AA:1233:C:OP1	17:AA:1353:A:N6	2.22	0.73
11:XA:1962:A:OP2	11:XA:2501:C:N4	2.22	0.73
11:XA:3178:U:O2'	82:r:134:ARG:NH2	2.21	0.73
17:AA:1008:A:OP1	21:AE:50:ARG:NE	2.21	0.73
33:AQ:55:GLU:OE2	33:AQ:59:ARG:NE	2.21	0.73
16:A4:477:TRP:CE3	24:AH:52:VAL:CG1	2.71	0.72
17:AA:825:U:N3	17:AA:827:A:OP1	2.22	0.72
23:AG:312:GLN:OE1	23:AG:345:ARG:NH2	2.22	0.72
84:r2:49:Y5P:H4A	84:r2:50:Y5P:H4	1.70	0.72
60:XU:151:PHE:CD2	69:d:222:LEU:CD1	2.70	0.72
20:AD:178:GLU:OE2	20:AD:181:ARG:NH2	2.22	0.72
23:AG:272:SER:OG	23:AG:347:ALA:O	2.07	0.72
11:XA:1851:G:O4'	11:XA:2134:A:N6	2.23	0.72
71:f:176:SER:O	78:m:35:SER:N	2.22	0.72
16:A4:141:MET:HE1	19:AC:148:LYS:HE2	1.71	0.72
16:A4:479:GLU:HA	16:A4:482:ILE:HD12	1.71	0.72
11:XA:3050:U:O2'	51:XL:63:LYS:NZ	2.22	0.72
18:AB:230:CYS:O	35:AS:42:ARG:NH2	2.23	0.72
11:XA:1955:G:O2'	11:XA:1958:G:O2'	2.07	0.72
13:A1:163:VAL:O	41:AY:317:ASN:ND2	2.22	0.72
17:AA:1231:A:O2'	17:AA:1236:C:N4	2.23	0.72
21:AE:92:ASN:ND2	32:AP:117:MET:SD	2.63	0.72
9:8:134:ASP:OD1	9:8:135:THR:N	2.23	0.72
16:A4:82:THR:CG2	24:AH:51:HIS:CE1	2.73	0.72
17:AA:897:C:OP1	26:AJ:114:ARG:NH2	2.22	0.72
17:AA:1454:G:OP2	23:AG:377:ARG:NH2	2.22	0.71
11:XA:3012:U:O4'	11:XA:3173:G:N2	2.23	0.71
17:AA:1014:A:O2'	17:AA:1031:G:O4'	2.08	0.71
45:XE:221:ARG:NH1	45:XE:257:MET:O	2.22	0.71
57:XR:122:ARG:NH2	57:XR:126:GLU:OE2	2.23	0.71
75:j:50:SER:OG	75:j:55:ARG:O	2.07	0.71
25:AI:81:GLU:O	25:AI:148:ARG:NH1	2.23	0.71
11:XA:2499:U:OP2	11:XA:2504:A:N6	2.23	0.71
59:XT:126:ASP:OD1	59:XT:127:MET:N	2.23	0.71
11:XA:2082:G:N2	65:XZ:88:MET:SD	2.64	0.71
11:XA:2170:G:N2	49:XJ:147:SER:OG	2.23	0.71
16:A4:477:TRP:HE3	24:AH:52:VAL:HG11	1.56	0.71
40:AX:174:ASN:OD1	40:AX:177:ARG:NH1	2.23	0.71
37:AU:77:GLU:OE1	37:AU:81:LYS:NZ	2.23	0.71

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:2187:C:O3'	49:XJ:106:LYS:NZ	2.22	0.71
17:AA:703:A:OP2	37:AU:43:ASN:ND2	2.23	0.71
61:XV:184:GLU:O	64:XY:93:LYS:NZ	2.23	0.70
17:AA:780:C:N3	28:AL:197:ARG:NH2	2.40	0.70
10:9:74:VAL:O	64:XY:83:ALA:N	2.23	0.70
13:A1:169:ARG:O	13:A1:218:ASN:ND2	2.25	0.70
16:A4:470:GLN:OE1	16:A4:472:ASP:N	2.25	0.70
57:XR:31:PHE:O	57:XR:36:ASN:ND2	2.24	0.70
63:XX:220:GLU:N	63:XX:220:GLU:OE1	2.24	0.70
83:r1:50:Y5P:H4	84:r2:35:P5P:H2	1.71	0.70
85:r3:58:Y5P:H6	85:r3:58:Y5P:OP2	1.91	0.70
7:6:198:ALA:O	7:6:254:TYR:OH	2.09	0.70
11:XA:1749:C:OP2	11:XA:2899:C:O2'	2.10	0.70
13:A1:59:LYS:NZ	16:A4:83:THR:O	2.25	0.70
17:AA:1389:G:N1	17:AA:1416:A:OP2	2.23	0.70
45:XE:56:GLU:OE2	54:XO:141:HIS:ND1	2.24	0.70
58:XS:155:ARG:NE	58:XS:157:GLU:OE2	2.24	0.70
63:XX:63:GLU:OE1	63:XX:63:GLU:N	2.25	0.70
87:t2:5:GLN:NE2	87:t2:9:ASP:OD2	2.25	0.70
9:8:177:HIS:N	70:e:58:VAL:O	2.24	0.69
11:XA:1800:G:N1	11:XA:1803:A:OP2	2.24	0.69
15:A3:187:GLU:O	28:AL:212:ARG:NH2	2.24	0.69
17:AA:826:A:N7	26:AJ:55:ARG:NE	2.40	0.69
17:AA:1220:A:O2'	23:AG:395:LYS:O	2.10	0.69
49:XJ:27:GLY:O	49:XJ:58:LYS:NZ	2.24	0.69
8:7:192:TRP:O	8:7:295:ARG:NH1	2.25	0.69
11:XA:2579:C:N4	84:r2:13:Y5P:OP1	2.24	0.69
11:XA:2663:C:OP1	54:XO:13:ARG:NH1	2.25	0.69
16:A4:108:LEU:CD2	20:AD:154:VAL:CG2	2.71	0.69
61:XV:20:ARG:NH1	61:XV:42:ARG:O	2.25	0.69
85:r3:25:Y5P:H2'	85:r3:26:P5P:H8	1.74	0.69
40:AX:111:TYR:O	40:AX:115:THR:OG1	2.10	0.69
55:XP:61:ALA:O	55:XP:176:ARG:NE	2.24	0.69
85:r3:22:P5P:H2'	85:r3:23:P5P:H8	1.74	0.69
7:6:114:ARG:NH1	43:XB:1643:A:OP1	2.25	0.69
81:q:111:GLU:OE2	81:q:114:ARG:NH1	2.26	0.69
7:6:308:GLN:NE2	7:6:311:MET:SD	2.66	0.69
8:7:247:ASN:ND2	8:7:251:ILE:O	2.26	0.69
13:A1:52:ALA:HB2	16:A4:94:TYR:CB	2.23	0.68
11:XA:1864:A:OP1	57:XR:17:ARG:NH1	2.26	0.68
17:AA:1212:U:O2'	17:AA:1214:A:N6	2.26	0.68

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
40:AX:121:ALA:N	40:AX:299:ASN:OD1	2.26	0.68
41:AY:303:GLN:NE2	41:AY:307:GLU:OE1	2.26	0.68
17:AA:1429:C:OP1	23:AG:388:ARG:NH2	2.26	0.68
52:XM:168:GLU:OE2	52:XM:220:ARG:NH2	2.27	0.68
72:g:106:GLN:O	72:g:151:GLY:N	2.25	0.68
11:XA:2643:G:O2'	11:XA:2645:G:OP2	2.10	0.68
52:XM:185:ASP:OD1	52:XM:186:ILE:N	2.26	0.68
11:XA:2010:U:OP2	74:i:115:ARG:NH1	2.27	0.68
11:XA:2520:C:OP2	44:XD:295:TYR:OH	2.10	0.68
17:AA:752:C:O2'	17:AA:793:C:N4	2.27	0.68
49:XJ:154:ARG:NH1	49:XJ:155:VAL:O	2.27	0.68
11:XA:2531:U:O4	44:XD:246:ARG:NH2	2.26	0.68
18:AB:137:TYR:O	18:AB:264:ARG:NH2	2.26	0.68
68:c:58:VAL:O	68:c:63:LYS:NZ	2.27	0.68
23:AG:220:GLN:OE1	23:AG:223:ARG:NH1	2.27	0.67
6:5:343:GLN:NE2	6:5:417:LEU:O	2.27	0.67
11:XA:1864:A:O3'	57:XR:13:ARG:NH1	2.27	0.67
11:XA:3127:G:O2'	11:XA:3130:A:N6	2.28	0.67
60:XU:150:TRP:CH2	69:d:264:LYS:HG2	2.28	0.67
11:XA:3201:A:OP1	50:XK:98:ARG:NH1	2.26	0.67
16:A4:477:TRP:HE3	24:AH:52:VAL:CG1	2.07	0.67
61:XV:108:MET:HE1	69:d:169:PHE:CE2	2.30	0.67
11:XA:1725:C:O2	11:XA:2921:A:N6	2.28	0.67
10:9:28:ARG:NH1	11:XA:2376:A:N3	2.42	0.67
11:XA:2028:G:N1	11:XA:2264:A:OP2	2.27	0.67
16:A4:198:TYR:O	16:A4:239:ARG:NH1	2.28	0.67
11:XA:2822:C:O2'	11:XA:2915:C:OP2	2.13	0.67
13:A1:69:VAL:HG22	16:A4:81:ASP:OD2	1.94	0.67
11:XA:1917:A:O5'	11:XA:1984:A:N6	2.28	0.67
45:XE:103:LYS:NZ	45:XE:291:GLY:O	2.28	0.67
60:XU:153:LEU:HD22	69:d:240:LYS:HB2	1.76	0.67
11:XA:1708:A:O5'	64:XY:192:LYS:NZ	2.28	0.66
11:XA:1883:G:N7	46:XF:281:ARG:NH1	2.43	0.66
17:AA:1265:C:OP1	27:AK:112:ARG:NH1	2.28	0.66
66:a:101:GLU:N	66:a:101:GLU:OE1	2.28	0.66
3:2:54:GLN:NE2	11:XA:1918:G:O3'	2.24	0.66
12:A0:49:ARG:NH2	37:AU:41:ARG:O	2.27	0.66
40:AX:56:PRO:O	40:AX:59:HIS:NE2	2.28	0.66
48:XI:101:ASN:OD1	48:XI:151:ASN:N	2.28	0.66
9:8:178:TYR:N	78:m:63:THR:O	2.27	0.66
11:XA:1694:U:O4'	64:XY:162:ARG:NH2	2.28	0.66

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
13:A1:50:ARG:HG2	16:A4:94:TYR:CE1	2.30	0.66
66:a:126:HIS:O	66:a:131:ARG:NH2	2.27	0.66
72:g:37:ARG:O	72:g:38:PHE:CD2	2.49	0.66
19:AC:37:ASN:OD1	19:AC:41:ARG:NH1	2.28	0.66
41:AY:340:SER:OG	41:AY:377:ARG:NH2	2.28	0.66
9:8:164:ARG:NH1	71:f:87:GLU:O	2.26	0.66
7:6:124:ARG:NH2	9:8:112:GLU:OE1	2.28	0.66
7:6:261:ARG:NH2	80:p:125:ASN:OD1	2.28	0.66
11:XA:2021:U:O4	52:XM:41:ARG:NH2	2.28	0.66
48:XI:224:HIS:O	48:XI:228:GLN:N	2.24	0.66
84:r2:58:P5P:O2'	84:r2:60:Y5P:H5	1.96	0.66
11:XA:2529:U:OP2	44:XD:208:ARG:NH1	2.28	0.66
22:AF:129:ALA:O	22:AF:134:GLN:NE2	2.29	0.66
30:AN:30:VAL:N	30:AN:47:LYS:O	2.29	0.66
75:j:101:GLU:O	75:j:105:GLN:NE2	2.28	0.66
14:A2:55:CYS:SG	14:A2:59:ASN:ND2	2.69	0.66
11:XA:2011:G:O6	74:i:113:ARG:NH1	2.29	0.65
38:AV:131:ASN:ND2	38:AV:134:GLN:OE1	2.28	0.65
80:p:121:ILE:O	80:p:124:LYS:NZ	2.29	0.65
7:6:117:VAL:O	7:6:121:ARG:NH2	2.28	0.65
34:AR:279:LYS:NZ	34:AR:308:HIS:CE1	2.64	0.65
85:r3:9:P5P:H6	85:r3:12:Y5P:H4	1.78	0.65
44:XD:111:ARG:NE	44:XD:165:ASN:OD1	2.29	0.65
11:XA:1974:A:OP2	44:XD:265:TRP:NE1	2.29	0.65
11:XA:2511:C:O2'	44:XD:257:ILE:O	2.13	0.65
17:AA:1478:A:N6	17:AA:1565:A:O2'	2.30	0.65
11:XA:2900:C:O2	85:r3:74:P5P:H6	1.96	0.65
16:A4:478:TYR:CD2	16:A4:482:ILE:HD11	2.32	0.65
6:5:72:ARG:NH2	11:XA:1712:A:OP2	2.30	0.65
11:XA:2348:A:O4'	69:d:69:LYS:HB3	1.97	0.65
20:AD:227:ASN:HD22	83:r1:55:Y5P:H4	1.62	0.65
16:A4:62:LYS:HA	24:AH:70:ASP:HB2	1.76	0.65
46:XF:162:TYR:OH	46:XF:164:MET:SD	2.53	0.65
11:XA:1884:G:O2'	11:XA:1895:C:O2	2.15	0.65
16:A4:478:TYR:CE2	16:A4:482:ILE:HD11	2.31	0.65
11:XA:2167:A:N6	11:XA:2212:C:OP2	2.30	0.65
17:AA:1050:C:OP2	28:AL:198:ARG:NH1	2.29	0.65
49:XJ:115:LYS:N	77:l:92:TYR:OH	2.29	0.65
17:AA:1487:C:N4	83:r1:48:Y5P:OP1	2.30	0.64
60:XU:54:ARG:NH1	86:s:314:GLN:OE1	2.29	0.64
7:6:191:ASN:ND2	55:XP:137:GLU:O	2.30	0.64

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
84:r2:45:Y5P:HD	84:r2:46:P5P:P	2.20	0.64
6:5:80:ARG:NH2	6:5:82:TYR:OH	2.30	0.64
40:AX:266:ASN:ND2	40:AX:311:SER:O	2.29	0.64
75:j:103:ARG:NE	75:j:106:GLU:OE2	2.30	0.64
27:AK:89:ASN:OD1	27:AK:102:ARG:NH1	2.30	0.64
10:9:126:LYS:O	10:9:129:GLN:NE2	2.31	0.64
43:XB:1644:G:O6	55:XP:87:HIS:NE2	2.29	0.64
85:r3:38:Y5P:HB	85:r3:39:Y5P:H6	1.80	0.64
11:XA:2756:C:OP1	47:XH:121:ASN:ND2	2.29	0.64
13:A1:146:HIS:CD2	16:A4:57:VAL:HG22	2.33	0.64
11:XA:1889:C:OP1	52:XM:133:LYS:NZ	2.31	0.64
45:XE:54:SER:OG	45:XE:57:ASN:OD1	2.15	0.64
85:r3:61:Y5P:H2'	85:r3:62:Y5P:H6	1.80	0.64
19:AC:113:ARG:NE	24:AH:164:LEU:O	2.30	0.64
2:1:23:GLU:OE1	2:1:23:GLU:N	2.31	0.63
17:AA:1314:C:N3	22:AF:36:ARG:NH2	2.45	0.63
51:XL:34:LYS:NZ	51:XL:58:ILE:O	2.29	0.63
84:r2:34:P5P:O2'	84:r2:35:P5P:OP1	2.15	0.63
83:r1:49:Y5P:C6	83:r1:50:Y5P:H5	2.27	0.63
14:A2:9:ARG:NH2	17:AA:1021:U:OP2	2.32	0.63
18:AB:162:CYS:O	18:AB:261:LYS:NZ	2.30	0.63
80:p:113:GLU:OE1	80:p:116:ARG:NH2	2.31	0.63
3:2:70:LEU:O	64:XY:198:ARG:NH2	2.31	0.63
16:A4:62:LYS:HE3	24:AH:68:GLU:OE1	1.98	0.63
17:AA:700:A:OP2	37:AU:27:ARG:NH1	2.31	0.63
19:AC:89:ASP:OD1	19:AC:90:VAL:N	2.31	0.63
44:XD:132:ASP:OD2	44:XD:135:ARG:NH1	2.32	0.63
11:XA:1847:U:OP1	52:XM:47:ARG:NE	2.31	0.63
11:XA:2104:A:OP1	53:XN:73:ARG:NH1	2.32	0.63
17:AA:1149:G:OP1	28:AL:202:ARG:NH2	2.32	0.63
55:XP:141:ASN:OD1	80:p:184:ASN:ND2	2.31	0.63
11:XA:1953:A:O2'	11:XA:2463:A:OP1	2.17	0.63
11:XA:2529:U:O2'	44:XD:206:TYR:O	2.17	0.63
22:AF:119:LYS:NZ	40:AX:398:LEU:O	2.32	0.63
60:XU:75:GLY:O	60:XU:88:LYS:NZ	2.31	0.63
63:XX:51:LYS:O	63:XX:60:GLU:N	2.32	0.63
65:XZ:78:ARG:O	65:XZ:83:LYS:NZ	2.32	0.63
17:AA:1590:A:OP2	33:AQ:47:LYS:NZ	2.32	0.62
11:XA:2156:A:N7	82:r:173:ARG:NH1	2.47	0.62
83:r1:46:Y5P:H5	85:r3:35:P5P:C6	2.28	0.62
38:AV:173:PHE:O	38:AV:178:THR:OG1	2.15	0.62

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:2928:C:OP2	11:XA:3073:C:O2'	2.17	0.62
52:XM:246:ASP:OD1	52:XM:249:LYS:NZ	2.33	0.62
62:XW:62:HIS:N	62:XW:65:ASN:OD1	2.32	0.62
62:XW:115:ASP:C	62:XW:119:ARG:HE	2.06	0.62
62:XW:115:ASP:O	62:XW:119:ARG:NE	2.32	0.62
10:9:84:THR:N	64:XY:88:GLN:OE1	2.31	0.62
46:XF:58:HIS:NE2	72:g:114:GLU:OE1	2.33	0.62
16:A4:455:ASN:O	16:A4:486:TYR:OH	2.17	0.62
45:XE:136:GLU:N	45:XE:136:GLU:OE2	2.33	0.62
86:s:111:LYS:NZ	86:s:387:ASN:O	2.33	0.62
16:A4:477:TRP:CE3	24:AH:52:VAL:HG11	2.35	0.62
32:AP:140:TYR:O	32:AP:141:ARG:NE	2.29	0.62
44:XD:138:ASP:OD1	44:XD:249:ASN:ND2	2.32	0.62
17:AA:1322:C:OP1	19:AC:43:ARG:NH1	2.33	0.62
11:XA:1761:A:O2'	11:XA:1762:A:O5'	2.16	0.61
44:XD:126:VAL:O	44:XD:160:GLY:N	2.30	0.61
16:A4:99:SER:N	16:A4:102:GLU:OE2	2.33	0.61
56:XQ:103:ARG:NH2	56:XQ:167:TYR:OH	2.33	0.61
6:5:230:LEU:O	6:5:289:HIS:N	2.30	0.61
8:7:78:VAL:HG22	69:d:282:ILE:HD11	1.82	0.61
11:XA:2190:C:OP2	77:l:120:ARG:NH2	2.32	0.61
17:AA:722:C:N3	17:AA:798:C:O2'	2.32	0.61
1:0:138:ARG:NH1	11:XA:2320:A:O3'	2.33	0.61
39:AW:103:ARG:O	39:AW:115:ASP:N	2.32	0.61
2:1:23:GLU:OE2	2:1:57:VAL:N	2.33	0.61
21:AE:48:PRO:O	32:AP:124:TYR:OH	2.19	0.61
52:XM:153:ASN:ND2	52:XM:256:LEU:O	2.34	0.61
60:XU:48:MET:O	60:XU:93:LYS:NZ	2.33	0.61
13:A1:53:LEU:O	16:A4:93:PRO:CB	2.49	0.61
14:A2:12:ARG:NH2	17:AA:1125:A:O4'	2.34	0.61
17:AA:945:G:O2'	28:AL:154:ARG:NH2	2.34	0.61
29:AM:68:LEU:O	34:AR:161:ILE:N	2.32	0.61
44:XD:162:THR:O	44:XD:177:ARG:NH1	2.33	0.61
11:XA:2195:A:OP1	77:l:119:ARG:NE	2.34	0.60
13:A1:69:VAL:HG22	16:A4:81:ASP:CG	2.26	0.60
11:XA:2832:A:O4'	71:f:49:LYS:NZ	2.34	0.60
61:XV:66:GLU:OE1	61:XV:66:GLU:N	2.34	0.60
86:s:381:THR:O	86:s:385:GLN:NE2	2.34	0.60
11:XA:1689:C:O2	64:XY:213:ARG:NH2	2.34	0.60
11:XA:1837:C:OP1	67:b:4:ARG:NH2	2.32	0.60
11:XA:2065:A:OP1	71:f:48:TYR:N	2.34	0.60

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
17:AA:1021:U:O4	33:AQ:59:ARG:NH1	2.33	0.60
7:6:239:ASN:OD1	7:6:275:GLN:NE2	2.34	0.60
63:XX:80:TRP:O	63:XX:131:THR:OG1	2.18	0.60
7:6:283:GLU:OE1	7:6:283:GLU:N	2.34	0.60
78:m:52:TYR:N	78:m:68:TYR:O	2.30	0.60
8:7:116:TYR:OH	8:7:271:ARG:NE	2.33	0.60
16:A4:116:VAL:HG12	19:AC:138:TYR:CE1	2.36	0.60
61:XV:147:SER:OG	61:XV:152:ARG:N	2.35	0.60
70:e:203:LYS:N	70:e:237:LEU:O	2.35	0.60
85:r3:69:Y5P:H6	85:r3:69:Y5P:OP2	2.02	0.60
7:6:380:TYR:O	52:XM:97:SER:N	2.34	0.60
11:XA:1729:U:OP2	63:XX:100:ARG:NH1	2.35	0.60
61:XV:181:ASP:O	64:XY:93:LYS:NZ	2.30	0.60
55:XP:74:ARG:O	55:XP:96:GLN:NE2	2.34	0.60
67:b:27:GLN:NE2	68:c:177:GLN:OE1	2.33	0.60
6:5:221:GLN:NE2	6:5:272:ASP:O	2.34	0.59
11:XA:1672:C:OP1	59:XT:50:LYS:N	2.35	0.59
11:XA:2192:A:OP1	49:XJ:142:ARG:NE	2.35	0.59
86:s:148:ASP:OD1	86:s:149:CYS:N	2.34	0.59
11:XA:2074:A:N6	71:f:48:TYR:OH	2.35	0.59
29:AM:59:ASN:ND2	29:AM:63:GLU:OE2	2.35	0.59
84:r2:67:Y5P:H2'	84:r2:68:Y5P:H6	1.84	0.59
7:6:119:GLU:OE1	7:6:119:GLU:N	2.34	0.59
47:XH:136:ASN:OD1	47:XH:137:LYS:N	2.36	0.59
13:A1:50:ARG:HD3	16:A4:94:TYR:CZ	2.37	0.59
16:A4:339:LEU:O	16:A4:374:HIS:NE2	2.36	0.59
17:AA:702:C:O2'	17:AA:842:C:O2	2.16	0.59
48:XI:38:ARG:O	53:XN:242:TRP:NE1	2.34	0.59
85:r3:54:Y5P:O2'	85:r3:56:Y5P:OP2	2.18	0.59
11:XA:2832:A:O5'	71:f:49:LYS:NZ	2.36	0.59
46:XF:215:SER:OG	46:XF:257:GLN:N	2.31	0.59
52:XM:277:MET:SD	72:g:45:TYR:OH	2.61	0.59
84:r2:2:Y5P:H4	84:r2:72:Y5P:N3	2.16	0.59
3:2:56:SER:OG	11:XA:1980:A:OP1	2.21	0.59
57:XR:149:HIS:O	65:XZ:151:LEU:N	2.35	0.59
13:A1:162:SER:CB	16:A4:134:GLU:HG2	2.33	0.59
86:s:191:HIS:O	86:s:428:ARG:NH2	2.33	0.59
11:XA:2725:A:C2	88:A:1:MHW:CA	2.86	0.59
57:XR:96:GLU:N	57:XR:96:GLU:OE2	2.35	0.59
59:XT:95:ARG:NH2	59:XT:145:SER:O	2.35	0.59
17:AA:1259:U:O2	17:AA:1326:A:O2'	2.10	0.59

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
24:AH:70:ASP:OD2	24:AH:151:SER:N	2.36	0.59
46:XF:284:TYR:O	46:XF:290:TYR:OH	2.20	0.59
60:XU:133:GLU:HG3	69:d:89:ALA:HB1	1.84	0.59
11:XA:2618:U:O4	11:XA:3043:C:N4	2.33	0.59
17:AA:769:G:OP2	30:AN:73:ARG:NH2	2.35	0.59
17:AA:1164:U:OP1	26:AJ:122:LYS:NZ	2.36	0.59
74:i:45:ASP:O	74:i:51:ARG:NH1	2.36	0.59
11:XA:2151:A:OP2	11:XA:2249:G:N1	2.33	0.58
11:XA:2425:A:OP1	86:s:221:HIS:ND1	2.35	0.58
17:AA:1233:C:O2'	27:AK:86:ARG:NH1	2.36	0.58
36:AT:97:GLU:OE1	36:AT:97:GLU:N	2.34	0.58
40:AX:161:TRP:NE1	40:AX:183:GLU:OE2	2.36	0.58
49:XJ:115:LYS:NZ	77:l:70:THR:O	2.36	0.58
71:f:104:GLU:OE2	71:f:155:ARG:NH2	2.36	0.58
6:5:334:LYS:N	6:5:362:THR:OG1	2.36	0.58
11:XA:1699:C:OP1	64:XY:197:LYS:NZ	2.35	0.58
12:A0:96:ARG:N	12:A0:117:ILE:O	2.35	0.58
17:AA:732:A:O2'	29:AM:15:GLY:O	2.19	0.58
46:XF:75:GLU:OE2	46:XF:210:ARG:NE	2.34	0.58
61:XV:185:ARG:NE	64:XY:92:ASN:OD1	2.36	0.58
3:2:64:HIS:NE2	11:XA:1788:C:O2	2.36	0.58
17:AA:1526:U:O2'	17:AA:1527:A:O4'	2.18	0.58
8:7:190:ASP:O	8:7:295:ARG:NH1	2.36	0.58
11:XA:2575:U:O2	11:XA:2582:A:N6	2.36	0.58
11:XA:2826:G:OP1	62:XW:49:ARG:NH1	2.32	0.58
13:A1:134:PRO:HB3	16:A4:58:VAL:O	2.04	0.58
20:AD:94:THR:OG1	20:AD:97:GLU:OE1	2.21	0.58
77:l:94:GLU:OE1	77:l:94:GLU:N	2.33	0.58
11:XA:1737:A:O2'	74:i:93:ARG:NH2	2.36	0.58
13:A1:118:ALA:O	13:A1:122:HIS:ND1	2.35	0.58
17:AA:1150:C:OP1	28:AL:201:ARG:NH1	2.36	0.58
2:1:42:THR:O	81:q:135:TRP:NE1	2.33	0.58
11:XA:1885:A:OP2	46:XF:168:LYS:NZ	2.36	0.58
11:XA:3151:A:N6	11:XA:3163:G:O2'	2.36	0.58
26:AJ:72:LYS:HD3	83:r1:50:Y5P:HA	1.85	0.58
86:s:77:ASP:OD1	86:s:78:GLU:N	2.37	0.58
8:7:155:GLU:OE2	8:7:156:ARG:NH1	2.37	0.58
20:AD:136:ARG:N	42:AZ:67:PHE:O	2.37	0.58
39:AW:110:ASN:OD1	39:AW:125:ARG:NH1	2.37	0.58
43:XB:1625:A:N7	55:XP:86:GLN:NE2	2.52	0.58
50:XK:24:LYS:O	50:XK:26:GLN:NE2	2.37	0.58

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
75:j:101:GLU:OE1	75:j:105:GLN:NE2	2.37	0.58
84:r2:9:P5P:O2'	84:r2:45:Y5P:O2'	2.22	0.58
4:3:182:ASP:OD1	4:3:183:ARG:N	2.36	0.58
8:7:83:LYS:NZ	8:7:120:THR:O	2.36	0.58
91:XA:5142:DOL:H343	91:XA:5142:DOL:H311	1.84	0.58
12:A0:87:TRP:O	31:AO:215:ARG:NH2	2.36	0.58
13:A1:121:LYS:HG3	16:A4:77:THR:CG2	2.34	0.58
15:A3:143:TYR:OH	17:AA:1145:A:OP1	2.21	0.58
52:XM:146:ASP:OD1	80:p:143:GLN:NE2	2.36	0.58
56:XQ:71:PRO:O	56:XQ:73:ARG:NH1	2.36	0.58
56:XQ:118:ARG:NH2	56:XQ:204:MET:O	2.36	0.58
72:g:76:ARG:NH2	72:g:164:LYS:O	2.34	0.58
11:XA:2029:A:O2'	11:XA:2030:U:OP1	2.20	0.58
37:AU:58:GLU:OE2	37:AU:62:HIS:NE2	2.37	0.58
53:XN:228:LYS:NZ	65:XZ:77:ARG:O	2.35	0.58
36:AT:95:ASN:OD1	36:AT:96:LYS:N	2.36	0.58
46:XF:70:ARG:NH1	46:XF:194:GLU:OE2	2.37	0.58
17:AA:1048:C:O2'	17:AA:1049:A:OP1	2.20	0.57
34:AR:279:LYS:NZ	34:AR:308:HIS:NE2	2.51	0.57
85:r3:9:P5P:O3'	85:r3:45:P5P:O2'	2.16	0.57
1:0:104:LYS:O	59:XT:105:GLN:NE2	2.38	0.57
11:XA:2348:A:C4'	69:d:69:LYS:HB3	2.35	0.57
20:AD:192:GLY:O	31:AO:78:ARG:NH2	2.37	0.57
23:AG:382:PRO:O	24:AH:131:ARG:NH1	2.36	0.57
52:XM:289:ASN:OD1	52:XM:290:LEU:N	2.37	0.57
65:XZ:107:ASN:O	65:XZ:111:LYS:HG2	2.03	0.57
11:XA:2925:U:O2'	11:XA:2927:C:OP1	2.18	0.57
17:AA:767:C:O2'	30:AN:23:GLN:O	2.23	0.57
36:AT:9:ILE:O	36:AT:12:THR:OG1	2.21	0.57
47:XH:108:ARG:NH1	47:XH:143:GLU:OE2	2.35	0.57
84:r2:45:Y5P:HA	84:r2:46:P5P:OP2	2.03	0.57
17:AA:1282:G:N2	17:AA:1286:A:OP2	2.30	0.57
86:s:228:ASP:O	86:s:230:ARG:NH1	2.37	0.57
11:XA:2910:A:C2	85:r3:74:P5P:H8	2.39	0.57
17:AA:1293:C:N4	33:AQ:80:ARG:O	2.37	0.57
19:AC:115:ASN:ND2	41:AY:309:LYS:O	2.38	0.57
56:XQ:260:TRP:O	56:XQ:262:GLN:NE2	2.38	0.57
11:XA:2682:A:OP1	57:XR:34:ARG:NH2	2.37	0.57
17:AA:918:A:O2'	17:AA:919:A:O4'	2.22	0.57
27:AK:41:ARG:NH2	27:AK:88:ARG:O	2.37	0.57
38:AV:96:ARG:NH1	38:AV:101:CYS:SG	2.77	0.57

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
69:d:249:GLU:OE2	69:d:262:HIS:ND1	2.38	0.57
7:6:286:ARG:NE	7:6:295:GLN:O	2.34	0.57
17:AA:766:G:OP2	30:AN:76:HIS:NE2	2.33	0.57
9:8:132:GLU:OE2	78:m:37:ARG:NH2	2.37	0.57
17:AA:1320:G:OP1	19:AC:41:ARG:NH1	2.35	0.57
52:XM:109:ARG:NH2	72:g:91:MET:SD	2.74	0.57
56:XQ:227:LYS:O	56:XQ:229:TRP:N	2.38	0.57
83:r1:45:Y5P:O2'	83:r1:46:Y5P:O4'	2.23	0.57
6:5:200:ARG:NH1	6:5:234:ASP:OD2	2.38	0.57
10:9:134:ASN:OD1	10:9:135:PHE:N	2.38	0.57
12:A0:132:GLU:OE1	12:A0:205:ALA:N	2.36	0.57
24:AH:135:GLU:N	27:AK:124:GLN:O	2.36	0.57
55:XP:73:SER:OG	55:XP:75:GLU:OE1	2.23	0.57
72:g:36:PRO:O	72:g:38:PHE:HD2	1.87	0.57
85:r3:53:Y5P:OP2	85:r3:54:Y5P:H5	2.04	0.57
8:7:36:SER:N	8:7:39:GLU:OE2	2.37	0.56
8:7:159:LYS:NZ	66:a:91:HIS:O	2.37	0.56
11:XA:1750:G:O2'	11:XA:1751:A:O4'	2.23	0.56
29:AM:20:ARG:NH1	29:AM:42:PRO:O	2.37	0.56
45:XE:102:LEU:N	45:XE:123:GLN:O	2.37	0.56
68:c:315:ASN:OD1	68:c:316:TYR:N	2.38	0.56
11:XA:1730:U:OP1	47:XH:76:ARG:NH2	2.37	0.56
23:AG:129:GLU:OE1	23:AG:129:GLU:N	2.37	0.56
30:AN:53:ASP:OD2	30:AN:57:GLN:N	2.38	0.56
14:A2:24:ASN:OD1	14:A2:25:LYS:N	2.37	0.56
16:A4:117:ALA:HB2	19:AC:141:THR:OG1	2.05	0.56
31:AO:80:ASN:OD1	31:AO:103:ASN:ND2	2.38	0.56
43:XB:1639:U:O4	43:XB:1640:A:N6	2.37	0.56
59:XT:55:ASN:O	69:d:227:ARG:NH2	2.37	0.56
84:r2:5:P5P:H2'	84:r2:6:P5P:H8	1.87	0.56
11:XA:1673:U:O2	11:XA:1815:A:N6	2.38	0.56
21:AE:53:ALA:N	21:AE:56:GLN:O	2.36	0.56
21:AE:90:ARG:NH2	32:AP:117:MET:O	2.38	0.56
26:AJ:107:ILE:N	26:AJ:131:ASP:OD2	2.39	0.56
40:AX:170:GLN:NE2	40:AX:171:SER:O	2.38	0.56
66:a:70:GLU:O	67:b:141:ARG:N	2.37	0.56
11:XA:1820:A:N3	66:a:128:ARG:NH2	2.53	0.56
16:A4:141:MET:CE	19:AC:148:LYS:CE	2.84	0.56
17:AA:889:G:N1	17:AA:905:A:OP2	2.36	0.56
17:AA:1199:G:N1	17:AA:1424:U:O4	2.37	0.56
21:AE:47:LEU:N	21:AE:59:ASN:O	2.38	0.56

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
22:AF:116:GLU:OE2	40:AX:86:ARG:NH1	2.39	0.56
34:AR:89:LYS:O	34:AR:92:LYS:NZ	2.39	0.56
43:XB:1630:A:OP1	78:m:42:ARG:NE	2.38	0.56
61:XV:54:TRP:NE1	61:XV:56:LEU:O	2.38	0.56
11:XA:2195:A:O2'	11:XA:2196:A:O5'	2.21	0.56
12:A0:103:ASP:OD2	12:A0:105:THR:OG1	2.22	0.56
18:AB:164:GLU:OE1	23:AG:145:ARG:NH2	2.38	0.56
20:AD:191:ARG:NH1	31:AO:79:ARG:O	2.39	0.56
21:AE:5:GLU:OE2	21:AE:96:HIS:ND1	2.39	0.56
13:A1:81:VAL:O	13:A1:99:LYS:NZ	2.38	0.56
34:AR:279:LYS:HZ1	34:AR:308:HIS:CE1	2.24	0.56
11:XA:2248:U:OP1	57:XR:99:ARG:NH2	2.39	0.56
11:XA:2261:C:N3	58:XS:184:ARG:NH2	2.54	0.56
11:XA:2519:G:N2	11:XA:2530:A:OP2	2.36	0.56
85:r3:38:Y5P:HB	85:r3:39:Y5P:C6	2.36	0.56
2:l:14:LYS:NZ	11:XA:2848:A:O3'	2.39	0.56
16:A4:68:VAL:HG13	41:AY:302:ILE:HG12	1.88	0.56
11:XA:2239:A:OP2	50:XK:75:LYS:NZ	2.39	0.56
32:AP:49:ASP:OD2	39:AW:82:SER:N	2.39	0.56
53:XN:134:LYS:NZ	53:XN:141:GLY:O	2.37	0.56
69:d:200:SER:OG	69:d:204:ASN:OD1	2.24	0.56
87:t3:5:GLN:NE2	87:t3:9:ASP:OD2	2.39	0.56
17:AA:710:U:OP2	29:AM:13:ARG:NH1	2.39	0.55
18:AB:111:LEU:O	18:AB:113:HIS:ND1	2.39	0.55
23:AG:321:ASP:OD1	23:AG:322:ARG:NH1	2.36	0.55
13:A1:50:ARG:CG	16:A4:94:TYR:CE1	2.89	0.55
17:AA:819:A:O2'	17:AA:831:U:O2'	2.23	0.55
44:XD:232:ARG:NH1	44:XD:291:PRO:O	2.39	0.55
67:b:44:ARG:NH2	79:o:99:LYS:O	2.39	0.55
11:XA:1844:A:OP2	57:XR:48:ARG:NH2	2.37	0.55
16:A4:443:ASP:O	16:A4:446:LYS:NZ	2.35	0.55
11:XA:2139:U:OP2	65:XZ:74:SER:N	2.35	0.55
11:XA:2369:A:OP1	64:XY:117:GLN:NE2	2.40	0.55
77:l:112:PRO:O	77:l:118:TRP:NE1	2.39	0.55
85:r3:22:P5P:OP2	85:r3:46:P5P:H6	2.07	0.55
16:A4:70:VAL:HG21	24:AH:158:GLU:OE1	2.07	0.55
10:9:72:PRO:O	64:XY:85:TRP:NE1	2.39	0.55
13:A1:131:THR:OG1	24:AH:151:SER:OG	2.22	0.55
13:A1:156:TYR:O	13:A1:167:ARG:NH1	2.39	0.55
13:A1:256:SER:O	13:A1:260:ARG:NH1	2.39	0.55
16:A4:264:ARG:HE	16:A4:293:THR:HG22	1.71	0.55

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
17:AA:1106:C:O2'	17:AA:1108:C:OP2	2.24	0.55
46:XF:102:TRP:NE1	46:XF:163:TYR:O	2.34	0.55
48:XI:148:VAL:O	76:k:31:ARG:NH1	2.38	0.55
82:r:162:ILE:HG23	82:r:162:ILE:O	2.06	0.55
84:r2:70:P5P:H8	84:r2:70:P5P:O5'	2.06	0.55
11:XA:1905:C:OP1	46:XF:117:ARG:NH1	2.39	0.55
21:AE:38:ASP:N	21:AE:67:ASP:OD1	2.39	0.55
34:AR:70:PHE:O	34:AR:76:GLN:NE2	2.38	0.55
47:XH:95:GLU:OE2	47:XH:112:VAL:N	2.40	0.55
58:XS:153:LEU:O	58:XS:201:ALA:N	2.38	0.55
75:j:101:GLU:OE2	75:j:102:GLN:NE2	2.40	0.55
6:5:201:ARG:NH1	6:5:418:TYR:O	2.40	0.55
11:XA:2515:U:O2'	44:XD:282:ALA:O	2.25	0.55
11:XA:3190:A:O5'	82:r:120:LYS:NZ	2.36	0.55
17:AA:1430:A:OP1	23:AG:388:ARG:NH2	2.34	0.55
57:XR:23:GLU:N	57:XR:23:GLU:OE1	2.38	0.55
68:c:267:LEU:HD12	68:c:267:LEU:O	2.07	0.55
85:r3:70:Y5P:H4A	85:r3:71:P5P:C6	2.37	0.55
17:AA:710:U:O2'	37:AU:28:LYS:O	2.25	0.55
46:XF:191:ASP:OD1	46:XF:192:SER:N	2.40	0.55
6:5:114:LEU:N	6:5:264:ASP:OD2	2.40	0.55
6:5:160:HIS:HA	6:5:164:TRP:HB2	1.88	0.55
17:AA:1301:G:O3'	19:AC:165:LYS:NZ	2.40	0.55
35:AS:7:GLU:OE1	35:AS:7:GLU:N	2.39	0.55
16:A4:82:THR:HG23	24:AH:51:HIS:CE1	2.41	0.54
46:XF:292:ASP:OD1	46:XF:293:PHE:N	2.40	0.54
2:1:40:LYS:NZ	2:1:59:LYS:O	2.40	0.54
11:XA:2166:C:N4	11:XA:2212:C:OP2	2.41	0.54
11:XA:2275:U:H2'	11:XA:2276:C:C6	2.43	0.54
17:AA:1225:C:O2'	17:AA:1449:G:O2'	2.24	0.54
11:XA:2400:C:O2'	11:XA:2401:A:O5'	2.26	0.54
13:A1:83:LEU:O	13:A1:99:LYS:NZ	2.35	0.54
32:AP:65:CYS:SG	32:AP:68:CYS:N	2.81	0.54
84:r2:19:P5P:N7	84:r2:59:Y5P:H4	2.22	0.54
28:AL:112:MET:SD	28:AL:120:GLU:N	2.77	0.54
46:XF:86:VAL:O	46:XF:179:THR:OG1	2.26	0.54
63:XX:103:LYS:NZ	63:XX:104:VAL:O	2.40	0.54
69:d:158:ASP:OD1	69:d:159:ARG:N	2.40	0.54
84:r2:43:Y5P:H2'	84:r2:44:P5P:C8	2.38	0.54
11:XA:2990:A:O2'	11:XA:2992:G:OP2	2.22	0.54
13:A1:142:LYS:O	13:A1:146:HIS:ND1	2.40	0.54

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
28:AL:169:ASN:OD1	28:AL:170:LEU:N	2.40	0.54
40:AX:214:GLU:OE2	40:AX:232:ARG:NH2	2.41	0.54
38:AV:141:ASN:OD1	38:AV:142:PHE:N	2.39	0.54
44:XD:227:GLN:OE1	44:XD:232:ARG:N	2.41	0.54
34:AR:202:ARG:NE	34:AR:233:ALA:O	2.37	0.54
85:r3:37:P5P:H2'	85:r3:38:Y5P:O4'	2.07	0.54
20:AD:108:ALA:O	20:AD:114:ARG:NH1	2.41	0.54
25:AI:94:ASN:OD1	25:AI:95:THR:N	2.39	0.54
12:A0:30:ASP:OD1	12:A0:31:SER:N	2.42	0.53
35:AS:18:ASP:OD1	35:AS:19:LEU:N	2.41	0.53
1:0:163:GLU:OE1	1:0:181:ARG:NH2	2.41	0.53
11:XA:1769:C:O2'	57:XR:11:ARG:NH2	2.40	0.53
16:A4:477:TRP:CZ3	24:AH:52:VAL:CG1	2.87	0.53
85:r3:37:P5P:H8	85:r3:37:P5P:O5'	2.08	0.53
11:XA:2348:A:H5'	69:d:69:LYS:HE3	1.89	0.53
34:AR:279:LYS:HZ2	34:AR:308:HIS:CE1	2.26	0.53
61:XV:77:VAL:N	61:XV:87:VAL:O	2.40	0.53
86:s:103:ASP:OD1	86:s:104:ARG:N	2.40	0.53
15:A3:164:ARG:NH2	17:AA:1501:A:OP1	2.40	0.53
26:AJ:78:ARG:NH2	26:AJ:117:ASP:OD2	2.42	0.53
51:XL:110:ASP:OD1	51:XL:111:ASN:N	2.40	0.53
68:c:238:MET:N	68:c:238:MET:SD	2.80	0.53
11:XA:2348:A:H5'	69:d:69:LYS:CE	2.38	0.53
16:A4:175:GLN:O	16:A4:180:GLY:N	2.40	0.53
17:AA:1450:C:O2'	24:AH:131:ARG:NH2	2.42	0.53
50:XK:130:ASP:OD1	50:XK:131:GLU:N	2.40	0.53
68:c:182:GLU:OE1	68:c:182:GLU:N	2.36	0.53
84:r2:9:P5P:H2	84:r2:13:Y5P:H4	1.90	0.53
11:XA:1920:U:O2	11:XA:1996:C:O2'	2.25	0.53
22:AF:142:TYR:OH	40:AX:397:TYR:O	2.25	0.53
47:XH:131:TYR:O	47:XH:136:ASN:ND2	2.37	0.53
17:AA:1431:G:N1	17:AA:1458:A:OP2	2.37	0.53
23:AG:115:GLY:N	24:AH:84:ASP:OD2	2.42	0.53
60:XU:133:GLU:CG	69:d:89:ALA:HB1	2.39	0.53
72:g:37:ARG:O	72:g:38:PHE:CG	2.61	0.53
85:r3:16:Y5P:O4'	85:r3:57:P5P:H2	2.09	0.53
4:3:108:LYS:NZ	11:XA:1743:U:OP2	2.34	0.53
6:5:306:PRO:O	6:5:310:ARG:NE	2.41	0.53
10:9:82:GLU:OE2	10:9:84:THR:OG1	2.26	0.53
11:XA:1760:G:OP1	52:XM:196:ARG:NE	2.41	0.53
11:XA:3195:A:OP2	11:XA:3196:G:O2'	2.22	0.53

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
17:AA:1486:C:H2'	17:AA:1487:C:O4'	2.08	0.53
86:s:416:ASP:OD1	86:s:417:VAL:N	2.42	0.53
13:A1:53:LEU:HD12	16:A4:93:PRO:HB3	1.92	0.53
13:A1:54:PRO:HD2	16:A4:518:GLU:OE2	2.09	0.53
13:A1:134:PRO:HG3	16:A4:60:PRO:CD	2.39	0.53
17:AA:1261:C:OP1	20:AD:108:ALA:N	2.42	0.53
70:e:177:GLU:O	70:e:190:ARG:NH2	2.42	0.53
85:r3:32:Y5P:H2'	85:r3:33:Y5P:C2	2.39	0.53
11:XA:3082:G:N2	11:XA:3085:A:OP2	2.42	0.52
85:r3:72:Y5P:H6	85:r3:72:Y5P:HB	1.90	0.52
15:A3:190:GLN:OE1	15:A3:190:GLN:N	2.43	0.52
16:A4:141:MET:CE	19:AC:148:LYS:HE2	2.39	0.52
18:AB:153:TYR:O	18:AB:157:ASN:ND2	2.43	0.52
24:AH:161:GLN:HA	24:AH:164:LEU:CD1	2.39	0.52
33:AQ:67:GLU:OE1	33:AQ:70:ARG:NH1	2.42	0.52
52:XM:100:ARG:NE	52:XM:126:GLY:O	2.35	0.52
69:d:64:MET:N	69:d:64:MET:SD	2.83	0.52
6:5:182:ASP:O	6:5:185:ILE:HG22	2.10	0.52
11:XA:2681:G:O6	11:XA:2682:A:N6	2.42	0.52
76:k:24:GLU:OE2	76:k:26:ASN:N	2.42	0.52
6:5:337:GLU:OE1	6:5:337:GLU:N	2.43	0.52
54:XO:43:GLU:OE1	54:XO:43:GLU:N	2.42	0.52
45:XE:334:ASP:OD1	45:XE:335:GLU:N	2.42	0.52
8:7:109:ASP:N	8:7:124:GLU:OE2	2.43	0.52
11:XA:2457:A:N3	54:XO:17:ARG:NH2	2.57	0.52
55:XP:94:GLU:N	55:XP:94:GLU:OE1	2.42	0.52
12:A0:101:ARG:NH1	17:AA:1528:A:OP1	2.43	0.52
17:AA:668:U:O2'	31:AO:83:GLY:O	2.27	0.52
56:XQ:268:ASP:OD1	56:XQ:269:MET:N	2.43	0.52
69:d:197:VAL:O	69:d:198:ARG:NE	2.43	0.52
7:6:282:SER:OG	7:6:283:GLU:OE1	2.28	0.52
7:6:291:TYR:O	55:XP:40:ASN:ND2	2.43	0.52
86:s:62:ARG:O	86:s:65:ARG:HG2	2.10	0.52
13:A1:121:LYS:HG3	16:A4:77:THR:HG22	1.92	0.52
17:AA:659:U:OP1	20:AD:226:ARG:NH2	2.40	0.52
19:AC:45:SER:N	19:AC:167:LEU:O	2.42	0.52
23:AG:362:GLU:OE2	23:AG:365:ARG:NH1	2.42	0.52
75:j:103:ARG:NH1	75:j:106:GLU:OE2	2.43	0.52
16:A4:95:LEU:CD2	19:AC:157:THR:HG21	2.40	0.51
17:AA:948:U:OP2	17:AA:1045:G:N1	2.39	0.51
85:r3:1:Y5P:H6	85:r3:1:Y5P:HB	1.92	0.51

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
6:5:51:GLU:OE1	6:5:51:GLU:N	2.41	0.51
85:r3:54:Y5P:HC	85:r3:56:Y5P:C5	2.32	0.51
17:AA:1280:C:O3'	18:AB:210:ARG:NH2	2.44	0.51
25:AI:79:LYS:N	25:AI:82:GLU:OE2	2.38	0.51
11:XA:3175:A:OP2	11:XA:3187:C:N4	2.42	0.51
16:A4:478:TYR:O	16:A4:482:ILE:HG13	2.11	0.51
21:AE:37:ARG:NH2	37:AU:163:GLU:OE1	2.43	0.51
27:AK:49:ASP:OD1	27:AK:50:GLU:N	2.43	0.51
85:r3:26:P5P:H2	85:r3:44:P5P:N7	2.25	0.51
6:5:68:PRO:HB2	11:XA:1713:A:N6	2.25	0.51
18:AB:107:PHE:N	18:AB:115:ILE:O	2.42	0.51
59:XT:135:GLU:OE2	69:d:279:THR:OG1	2.28	0.51
68:c:94:ASN:OD1	68:c:122:ASN:ND2	2.40	0.51
85:r3:47:Y5P:H2'	85:r3:57:P5P:C1'	2.27	0.51
11:XA:2079:C:C4	11:XA:2080:U:C6	2.99	0.51
11:XA:2347:C:O3'	69:d:69:LYS:CE	2.58	0.51
11:XA:2462:A:OP1	45:XE:237:HIS:NE2	2.42	0.51
13:A1:53:LEU:HD21	16:A4:96:MET:HG2	1.92	0.51
16:A4:62:LYS:HB2	24:AH:70:ASP:CA	2.36	0.51
16:A4:442:GLY:HA3	24:AH:59:THR:CG2	2.40	0.51
28:AL:149:ASP:OD2	28:AL:152:HIS:ND1	2.43	0.51
56:XQ:252:GLN:NE2	56:XQ:256:GLU:OE2	2.43	0.51
84:r2:9:P5P:H8	84:r2:45:Y5P:H2'	1.92	0.51
11:XA:2714:A:OP1	45:XE:239:ARG:NH1	2.43	0.51
14:A2:102:ASN:OD1	14:A2:103:LYS:N	2.40	0.51
74:i:125:GLY:O	74:i:128:ARG:NH1	2.44	0.51
11:XA:1799:U:H2'	11:XA:1800:G:O4'	2.11	0.51
11:XA:2083:U:O4	79:o:67:ARG:NH2	2.42	0.51
11:XA:2449:G:OP2	86:s:225:ARG:NH2	2.38	0.51
38:AV:132:LYS:O	38:AV:136:GLY:N	2.32	0.51
41:AY:365:SER:N	41:AY:368:GLN:OE1	2.42	0.51
70:e:201:GLU:OE1	70:e:201:GLU:N	2.44	0.51
11:XA:1747:G:O3'	63:XX:96:LYS:NZ	2.44	0.51
11:XA:2040:G:H2'	11:XA:2041:U:O4'	2.11	0.51
11:XA:2384:A:N1	11:XA:2409:A:N6	2.58	0.51
16:A4:98:ALA:N	16:A4:102:GLU:OE2	2.44	0.51
16:A4:141:MET:HE2	19:AC:148:LYS:CE	2.40	0.51
85:r3:71:P5P:O3'	85:r3:72:Y5P:O4'	2.29	0.51
5:4:88:TRP:NE1	11:XA:2160:A:OP2	2.36	0.51
8:7:315:LYS:NZ	45:XE:68:GLU:OE1	2.43	0.51
11:XA:1902:C:O2	74:i:126:LYS:N	2.44	0.51

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:2937:A:OP1	11:XA:2984:A:O2'	2.27	0.51
23:AG:203:GLU:O	23:AG:207:GLU:OE1	2.29	0.51
39:AW:132:GLU:O	39:AW:135:GLN:NE2	2.40	0.51
84:r2:29:P5P:H2'	84:r2:30:P5P:H8	1.93	0.51
11:XA:2403:G:OP2	44:XD:105:ARG:NH2	2.43	0.50
13:A1:140:ASP:OD1	13:A1:141:GLU:N	2.44	0.50
16:A4:72:GLN:CD	24:AH:61:PRO:HG3	2.36	0.50
17:AA:723:A:OP1	17:AA:724:C:N4	2.37	0.50
17:AA:893:G:OP2	26:AJ:79:LYS:NZ	2.42	0.50
53:XN:203:GLU:OE1	53:XN:207:ARG:NE	2.39	0.50
11:XA:2723:A:O2'	11:XA:2724:G:OP1	2.26	0.50
12:A0:13:GLU:OE1	12:A0:16:ARG:NH1	2.43	0.50
37:AU:171:GLU:OE2	37:AU:171:GLU:N	2.42	0.50
50:XK:123:GLU:OE2	66:a:119:THR:N	2.44	0.50
80:p:128:ASN:OD1	80:p:129:ARG:N	2.44	0.50
17:AA:1433:A:N3	17:AA:1458:A:N6	2.60	0.50
44:XD:211:GLY:O	44:XD:246:ARG:NH2	2.44	0.50
49:XJ:107:GLU:OE1	49:XJ:109:ALA:N	2.43	0.50
7:6:27:ARG:N	11:XA:2073:A:OP2	2.45	0.50
8:7:203:THR:O	8:7:207:HIS:ND1	2.42	0.50
11:XA:2537:G:H2'	11:XA:2538:C:C6	2.45	0.50
11:XA:2939:C:H2'	11:XA:2940:A:O4'	2.11	0.50
13:A1:131:THR:CG2	16:A4:63:LYS:HE3	2.42	0.50
13:A1:146:HIS:HD2	16:A4:57:VAL:CG2	2.24	0.50
22:AF:70:LYS:O	23:AG:365:ARG:NH1	2.45	0.50
57:XR:17:ARG:HA	57:XR:20:ARG:HG2	1.92	0.50
68:c:135:THR:O	68:c:138:THR:OG1	2.27	0.50
85:r3:25:Y5P:O5'	85:r3:25:Y5P:H6	2.12	0.50
11:XA:2744:U:O2'	11:XA:2746:U:O4	2.26	0.50
16:A4:67:LYS:HB3	41:AY:312:GLU:OE2	2.11	0.50
7:6:376:THR:O	11:XA:1892:A:N6	2.45	0.50
63:XX:150:LYS:HG3	63:XX:159:MET:SD	2.51	0.50
68:c:42:GLN:O	68:c:46:GLU:OE1	2.30	0.50
3:2:87:ARG:NH2	11:XA:1792:G:N7	2.57	0.50
40:AX:272:THR:OG1	40:AX:282:ILE:O	2.25	0.50
64:XY:124:LEU:HD21	69:d:68:ARG:NH1	2.22	0.50
64:XY:208:PHE:O	64:XY:212:GLU:OE1	2.29	0.50
7:6:74:TYR:N	62:XW:102:GLU:OE2	2.44	0.50
11:XA:1820:A:O4'	66:a:128:ARG:NE	2.45	0.50
17:AA:1343:A:O2'	17:AA:1345:G:N7	2.43	0.50
55:XP:71:PHE:HB3	55:XP:72:PRO:HD3	1.92	0.50

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
9:8:140:LEU:O	9:8:143:GLN:NE2	2.43	0.50
14:A2:27:LEU:N	21:AE:118:TYR:OH	2.45	0.50
16:A4:62:LYS:CA	24:AH:70:ASP:HB2	2.41	0.50
16:A4:366:GLU:OE1	16:A4:366:GLU:N	2.42	0.50
17:AA:1310:C:O2'	27:AK:128:TRP:NE1	2.43	0.50
34:AR:140:ASP:OD1	34:AR:141:VAL:N	2.43	0.50
61:XV:103:ASP:OD1	61:XV:104:TYR:N	2.44	0.50
61:XV:105:ARG:HH12	69:d:171:ASP:CA	2.09	0.50
67:b:95:GLU:OE1	67:b:95:GLU:N	2.44	0.50
85:r3:70:Y5P:O5'	85:r3:70:Y5P:H6	2.11	0.50
29:AM:87:MET:SD	29:AM:88:GLU:N	2.85	0.49
48:XI:181:ILE:O	48:XI:184:THR:N	2.44	0.49
77:l:80:GLU:OE1	77:l:82:GLN:NE2	2.45	0.49
83:r1:46:Y5P:HB	83:r1:47:Y5P:H5	1.94	0.49
86:s:63:ILE:HA	86:s:66:TRP:NE1	2.27	0.49
12:A0:107:GLN:O	56:XQ:55:GLN:NE2	2.45	0.49
17:AA:969:A:N3	28:AL:146:HIS:NE2	2.59	0.49
53:XN:99:TRP:NE1	53:XN:103:GLU:OE2	2.45	0.49
70:e:65:PRO:O	70:e:69:GLU:OE1	2.30	0.49
7:6:282:SER:O	7:6:283:GLU:N	2.45	0.49
11:XA:2992:G:N2	88:A:5:MHU:HM1	2.28	0.49
45:XE:257:MET:HB2	45:XE:258:PRO:CD	2.42	0.49
55:XP:76:PHE:O	55:XP:96:GLN:NE2	2.40	0.49
84:r2:9:P5P:H6	84:r2:24:P5P:N7	2.28	0.49
11:XA:2135:A:N3	11:XA:2135:A:H2'	2.27	0.49
27:AK:28:HIS:NE2	42:AZ:60:GLU:OE2	2.41	0.49
6:5:203:CYS:O	86:s:179:GLN:NE2	2.46	0.49
6:5:350:ARG:NH1	6:5:384:GLN:OE1	2.45	0.49
11:XA:1774:U:O4'	46:XF:147:ARG:NH2	2.44	0.49
34:AR:279:LYS:HZ2	34:AR:308:HIS:CD2	2.30	0.49
10:9:69:LYS:O	10:9:71:LYS:NZ	2.44	0.49
17:AA:1234:C:O2'	17:AA:1235:U:OP1	2.28	0.49
77:l:135:ASN:OD1	77:l:135:ASN:C	2.55	0.49
84:r2:30:P5P:H2'	84:r2:31:P5P:H8	1.94	0.49
7:6:160:ASP:OD2	7:6:267:ARG:NH1	2.45	0.49
58:XS:127:ARG:NH2	58:XS:157:GLU:OE1	2.44	0.49
86:s:67:GLN:O	86:s:71:HIS:ND1	2.45	0.49
11:XA:2694:A:N3	11:XA:2942:C:O2'	2.41	0.49
17:AA:845:A:O2'	17:AA:846:A:O4'	2.30	0.49
54:XO:64:LYS:NZ	54:XO:97:TYR:O	2.43	0.49
62:XW:117:ILE:HA	62:XW:120:LEU:HD22	1.95	0.49

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
3:2:88:LYS:NZ	10:9:18:MET:SD	2.82	0.49
20:AD:124:LYS:NZ	20:AD:125:ASP:O	2.46	0.49
28:AL:66:ASP:N	28:AL:66:ASP:OD1	2.45	0.49
56:XQ:225:LYS:HG2	56:XQ:226:PRO:HD2	1.95	0.49
77:l:60:ASP:OD1	77:l:61:VAL:N	2.45	0.49
80:p:190:GLN:O	80:p:190:GLN:HG2	2.13	0.49
11:XA:2307:U:H2'	11:XA:2308:A:O4'	2.12	0.49
13:A1:236:THR:OG1	13:A1:240:GLU:OE1	2.30	0.49
16:A4:61:LYS:C	24:AH:70:ASP:HB2	2.38	0.49
17:AA:1078:A:H5'	85:r3:38:Y5P:O3'	2.13	0.49
63:XX:150:LYS:HG3	63:XX:159:MET:CE	2.42	0.49
69:d:116:MET:N	69:d:116:MET:SD	2.85	0.49
72:g:36:PRO:O	72:g:38:PHE:CD2	2.64	0.49
85:r3:26:P5P:H6	85:r3:42:P5P:C6	2.43	0.49
11:XA:1990:G:OP1	44:XD:269:ARG:NH2	2.38	0.48
11:XA:2279:U:OP2	74:i:46:ARG:NH2	2.44	0.48
16:A4:120:ILE:HG21	19:AC:142:LEU:HD21	1.95	0.48
69:d:235:GLN:OE1	69:d:235:GLN:N	2.45	0.48
85:r3:25:Y5P:C4	85:r3:42:P5P:H6	2.33	0.48
11:XA:2262:C:OP2	58:XS:173:ARG:NH1	2.46	0.48
11:XA:2685:U:O4'	11:XA:2697:G:N2	2.47	0.48
11:XA:2926:A:O2'	11:XA:3087:C:OP1	2.28	0.48
46:XF:167:MET:SD	46:XF:279:ARG:NH1	2.86	0.48
48:XI:163:GLU:O	48:XI:166:ARG:HG3	2.13	0.48
84:r2:2:Y5P:O5'	84:r2:2:Y5P:H6	2.12	0.48
10:9:54:LYS:NZ	11:XA:2415:C:O2'	2.45	0.48
11:XA:1678:C:O2	61:XV:42:ARG:NH2	2.46	0.48
44:XD:136:SER:O	44:XD:249:ASN:ND2	2.46	0.48
75:j:28:ARG:O	75:j:33:LEU:N	2.45	0.48
75:j:102:GLN:O	75:j:106:GLU:OE1	2.31	0.48
85:r3:33:Y5P:H6	85:r3:33:Y5P:HB	1.95	0.48
85:r3:42:P5P:O2'	85:r3:44:P5P:OP2	2.22	0.48
8:7:279:GLU:OE1	8:7:279:GLU:N	2.46	0.48
16:A4:68:VAL:CG1	41:AY:302:ILE:HG23	2.44	0.48
52:XM:88:SER:O	52:XM:134:ARG:NE	2.47	0.48
63:XX:169:LEU:O	63:XX:172:GLN:NE2	2.38	0.48
64:XY:175:ARG:NH2	64:XY:176:ILE:O	2.46	0.48
82:r:40:GLU:OE1	82:r:40:GLU:N	2.46	0.48
84:r2:34:P5P:H2'	84:r2:35:P5P:H8	1.95	0.48
11:XA:1904:C:C2	11:XA:2019:G:C2	3.02	0.48
11:XA:1911:C:C2	11:XA:2009:G:N2	2.81	0.48

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
13:A1:87:MET:O	13:A1:102:ASN:ND2	2.47	0.48
23:AG:265:GLN:NE2	23:AG:267:MET:SD	2.85	0.48
49:XJ:85:PRO:O	49:XJ:124:LYS:NZ	2.46	0.48
72:g:154:ASP:OD1	72:g:155:GLN:N	2.46	0.48
4:3:105:LYS:NZ	11:XA:1753:A:OP2	2.38	0.48
11:XA:2108:G:N2	11:XA:2112:A:OP2	2.39	0.48
13:A1:274:LYS:HD3	13:A1:274:LYS:N	2.29	0.48
38:AV:192:LYS:NZ	38:AV:194:THR:O	2.46	0.48
44:XD:194:ASN:ND2	44:XD:245:GLY:O	2.47	0.48
52:XM:68:GLU:OE1	52:XM:68:GLU:N	2.47	0.48
54:XO:113:ARG:NH1	54:XO:116:ASP:OD2	2.46	0.48
62:XW:112:GLU:O	62:XW:115:ASP:OD1	2.31	0.48
84:r2:62:Y5P:H2'	84:r2:63:P5P:H8	1.96	0.48
11:XA:2933:G:N2	11:XA:2936:U:O2	2.43	0.48
13:A1:50:ARG:HB3	16:A4:94:TYR:OH	2.13	0.48
17:AA:1059:U:O4	17:AA:1060:A:N6	2.44	0.48
20:AD:95:ALA:N	42:AZ:74:GLU:OE1	2.46	0.48
68:c:258:THR:N	68:c:271:PHE:O	2.40	0.48
10:9:52:GLN:NE2	11:XA:2416:U:O3'	2.45	0.48
11:XA:1870:A:N6	74:i:124:HIS:O	2.43	0.48
11:XA:2170:G:N2	49:XJ:147:SER:HG	2.11	0.48
16:A4:108:LEU:CD2	20:AD:154:VAL:HG22	2.42	0.48
20:AD:293:ASP:OD1	20:AD:307:LYS:N	2.43	0.48
61:XV:112:GLU:N	61:XV:112:GLU:OE1	2.42	0.48
85:r3:14:P5P:H2	85:r3:22:P5P:C6	2.43	0.48
11:XA:2074:A:OP1	65:XZ:40:ARG:N	2.41	0.48
11:XA:2173:G:N3	49:XJ:150:SER:OG	2.47	0.48
13:A1:77:LYS:CB	16:A4:91:ASP:OD2	2.62	0.48
41:AY:344:GLN:N	41:AY:344:GLN:OE1	2.46	0.48
53:XN:85:GLY:O	53:XN:192:ARG:NH2	2.41	0.48
63:XX:150:LYS:HE3	63:XX:159:MET:HE2	1.95	0.48
7:6:106:ARG:NH1	43:XB:1621:A:OP2	2.47	0.48
13:A1:68:SER:HA	16:A4:78:VAL:HG11	1.96	0.48
15:A3:142:LYS:NZ	17:AA:1490:U:OP1	2.39	0.48
21:AE:37:ARG:N	21:AE:67:ASP:O	2.41	0.48
25:AI:93:ASN:O	25:AI:123:GLY:N	2.47	0.48
28:AL:140:GLU:O	28:AL:144:GLU:OE1	2.31	0.48
40:AX:246:GLU:OE2	40:AX:250:GLN:NE2	2.41	0.48
61:XV:148:THR:HG22	61:XV:149:ARG:H	1.78	0.48
63:XX:160:ASP:OD1	63:XX:163:ARG:NH2	2.43	0.48
73:h:78:PHE:HB2	73:h:80:SER:C	2.39	0.48

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:2239:A:O2'	50:XK:29:GLY:HA3	2.14	0.47
11:XA:2714:A:N3	11:XA:2715:A:N6	2.62	0.47
13:A1:146:HIS:CD2	16:A4:57:VAL:CG2	2.97	0.47
17:AA:689:U:OP1	17:AA:827:A:O2'	2.29	0.47
21:AE:38:ASP:OD1	21:AE:39:LEU:N	2.44	0.47
23:AG:321:ASP:OD1	23:AG:321:ASP:C	2.56	0.47
34:AR:221:GLN:OE1	34:AR:223:ARG:NE	2.46	0.47
81:q:137:GLN:O	81:q:141:GLU:OE1	2.32	0.47
84:r2:1:P5P:H2	84:r2:73:P5P:O2'	2.14	0.47
4:3:140:ARG:NH2	11:XA:2870:G:OP1	2.39	0.47
9:8:178:TYR:O	70:e:136:ARG:NE	2.47	0.47
11:XA:1884:G:N3	11:XA:1895:C:O2'	2.47	0.47
11:XA:2230:A:N7	11:XA:2975:G:O2'	2.44	0.47
17:AA:1193:U:O3'	22:AF:178:ARG:NH2	2.45	0.47
78:m:87:GLU:OE1	78:m:91:ARG:NH2	2.47	0.47
84:r2:9:P5P:HO2'	84:r2:45:Y5P:HD	1.55	0.47
11:XA:1737:A:N6	11:XA:1760:G:O2'	2.47	0.47
11:XA:2829:C:OP2	11:XA:2830:A:O2'	2.23	0.47
20:AD:216:GLU:OE2	20:AD:216:GLU:N	2.45	0.47
62:XW:115:ASP:OD1	62:XW:116:LEU:N	2.47	0.47
70:e:185:ARG:O	70:e:189:GLU:OE1	2.31	0.47
79:o:98:THR:O	79:o:100:LYS:NZ	2.46	0.47
80:p:81:CYS:SG	80:p:82:ARG:N	2.85	0.47
80:p:187:ARG:O	80:p:190:GLN:NE2	2.48	0.47
3:2:54:GLN:OE1	3:2:54:GLN:N	2.42	0.47
16:A4:491:GLN:O	16:A4:495:HIS:ND1	2.45	0.47
36:AT:130:GLY:N	36:AT:135:CYS:SG	2.87	0.47
57:XR:14:VAL:O	68:c:32:LYS:NZ	2.37	0.47
61:XV:197:GLU:OE1	64:XY:95:ASN:ND2	2.45	0.47
62:XW:60:TYR:OH	62:XW:137:LYS:NZ	2.47	0.47
73:h:114:ASN:HA	73:h:117:LEU:HD22	1.96	0.47
83:r1:47:Y5P:C2	85:r3:35:P5P:H6	2.44	0.47
3:2:55:PRO:HB2	11:XA:2339:G:H4'	1.95	0.47
13:A1:77:LYS:HB3	16:A4:91:ASP:OD2	2.15	0.47
14:A2:59:ASN:OD1	14:A2:62:ARG:NH2	2.47	0.47
19:AC:77:ASP:OD2	24:AH:140:TYR:OH	2.31	0.47
50:XK:133:ILE:HB	50:XK:138:LEU:HB3	1.96	0.47
56:XQ:108:ILE:O	56:XQ:108:ILE:HG13	2.14	0.47
73:h:151:ASN:OD1	73:h:152:LEU:N	2.47	0.47
86:s:271:LEU:O	86:s:273:LEU:N	2.46	0.47
87:t5:4:GLN:OE1	87:t5:4:GLN:N	2.47	0.47

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
9:8:137:ARG:O	9:8:141:GLU:OE1	2.33	0.47
37:AU:52:GLU:OE1	37:AU:52:GLU:N	2.45	0.47
38:AV:271:GLU:OE1	38:AV:271:GLU:N	2.46	0.47
53:XN:221:ALA:O	65:XZ:114:LYS:NZ	2.45	0.47
55:XP:90:GLU:OE1	55:XP:107:ARG:NH1	2.48	0.47
69:d:157:HIS:O	69:d:161:HIS:ND1	2.47	0.47
70:e:269:LEU:O	70:e:273:ARG:HD3	2.13	0.47
84:r2:23:P5P:O2'	84:r2:24:P5P:O5'	2.33	0.47
1:0:86:THR:OG1	11:XA:2684:C:OP1	2.30	0.47
4:3:142:LYS:O	4:3:146:GLU:HG2	2.15	0.47
7:6:50:LYS:HA	62:XW:121:PRO:HA	1.97	0.47
11:XA:2460:A:OP1	45:XE:238:ARG:NE	2.44	0.47
11:XA:3061:G:H2'	11:XA:3062:U:O4'	2.14	0.47
16:A4:68:VAL:HG11	41:AY:302:ILE:HG23	1.96	0.47
16:A4:141:MET:HE2	19:AC:148:LYS:HE3	1.95	0.47
17:AA:1289:G:O2'	17:AA:1297:G:OP2	2.28	0.47
18:AB:202:ILE:O	18:AB:202:ILE:HG22	2.15	0.47
23:AG:302:LEU:O	40:AX:320:ARG:NH2	2.46	0.47
29:AM:111:ARG:NH2	31:AO:232:PRO:O	2.48	0.47
35:AS:92:PHE:O	39:AW:91:GLN:NE2	2.40	0.47
36:AT:116:GLU:HA	36:AT:119:GLU:CD	2.40	0.47
60:XU:14:GLY:O	61:XV:208:ARG:NE	2.44	0.47
77:l:89:ASP:N	77:l:89:ASP:OD1	2.48	0.47
11:XA:1671:G:C6	11:XA:1818:A:N1	2.83	0.47
11:XA:1787:G:N1	11:XA:1791:G:C6	2.82	0.47
11:XA:3097:U:C6	88:A:7:004:HD2	2.49	0.47
17:AA:894:C:N4	26:AJ:117:ASP:OD2	2.42	0.47
17:AA:1048:C:O2'	28:AL:196:TYR:O	2.32	0.47
17:AA:1187:U:OP2	17:AA:1189:U:N3	2.41	0.47
56:XQ:107:HIS:O	56:XQ:108:ILE:HG13	2.15	0.47
65:XZ:148:GLN:OE1	65:XZ:148:GLN:N	2.45	0.47
80:p:177:ARG:O	80:p:181:GLU:OE1	2.33	0.47
84:r2:60:Y5P:OP2	84:r2:61:Y5P:H5	2.15	0.47
11:XA:2693:A:OP1	79:o:15:ARG:NH2	2.46	0.47
17:AA:1234:C:O2	17:AA:1234:C:H2'	2.15	0.47
34:AR:114:ALA:O	34:AR:117:GLN:NE2	2.48	0.47
36:AT:115:GLU:O	36:AT:119:GLU:OE1	2.33	0.47
56:XQ:252:GLN:O	56:XQ:256:GLU:OE1	2.32	0.47
82:r:84:ASP:OD1	82:r:85:ASP:N	2.48	0.47
4:3:177:TYR:O	4:3:181:HIS:ND1	2.47	0.47
6:5:182:ASP:O	6:5:186:GLN:OE1	2.33	0.47

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:1681:G:OP2	64:XY:230:LYS:NZ	2.39	0.47
11:XA:2187:C:O2'	49:XJ:106:LYS:NZ	2.46	0.47
40:AX:100:MET:HB3	92:AX:500:GTP:HN1	1.79	0.47
11:XA:2520:C:O2	11:XA:2528:G:N2	2.43	0.46
17:AA:988:G:N2	17:AA:1002:C:O2'	2.45	0.46
62:XW:60:TYR:OH	62:XW:94:GLU:OE2	2.29	0.46
73:h:58:ARG:NH1	73:h:136:ASP:OD2	2.48	0.46
81:q:138:GLN:HA	81:q:141:GLU:OE2	2.15	0.46
16:A4:116:VAL:HG21	19:AC:134:PHE:HZ	1.80	0.46
17:AA:944:U:O2'	17:AA:1025:A:N3	2.34	0.46
22:AF:227:HIS:O	22:AF:231:GLU:OE1	2.33	0.46
38:AV:233:LYS:O	38:AV:288:LYS:NZ	2.46	0.46
48:XI:101:ASN:O	76:k:35:GLN:NE2	2.48	0.46
84:r2:9:P5P:O2'	84:r2:46:P5P:H5'2	2.15	0.46
5:4:99:LYS:NZ	11:XA:3013:G:O2'	2.44	0.46
8:7:290:GLN:N	8:7:291:PRO:HD2	2.30	0.46
11:XA:2574:G:O2'	11:XA:2575:U:P	2.74	0.46
11:XA:2803:A:H2'	11:XA:2804:A:O4'	2.16	0.46
11:XA:2870:G:C6	11:XA:2871:U:C5	3.03	0.46
12:A0:50:LEU:O	12:A0:55:TRP:NE1	2.48	0.46
17:AA:681:U:H2'	17:AA:682:A:C8	2.50	0.46
20:AD:254:ALA:O	20:AD:280:HIS:N	2.43	0.46
38:AV:106:ASN:OD1	38:AV:107:TRP:N	2.48	0.46
48:XI:198:PRO:O	48:XI:199:SER:C	2.58	0.46
11:XA:2151:A:O3'	79:o:38:ARG:NH1	2.45	0.46
16:A4:74:LEU:HB3	41:AY:298:PHE:CE1	2.50	0.46
17:AA:769:G:N2	17:AA:772:A:OP2	2.38	0.46
38:AV:222:SER:OG	38:AV:277:ARG:NH1	2.48	0.46
41:AY:367:LYS:O	41:AY:371:GLU:OE1	2.33	0.46
44:XD:172:MET:SD	44:XD:172:MET:N	2.87	0.46
51:XL:43:ASN:ND2	51:XL:117:THR:OG1	2.48	0.46
70:e:93:ASP:OD1	70:e:94:GLU:N	2.48	0.46
84:r2:5:P5P:H2'	84:r2:6:P5P:C8	2.46	0.46
86:s:174:LEU:HB3	86:s:175:PRO:HD3	1.97	0.46
11:XA:1837:C:O2	11:XA:1837:C:O4'	2.34	0.46
11:XA:2245:A:H1'	11:XA:2246:A:C8	2.50	0.46
11:XA:2934:G:C2'	11:XA:2987:U:OP2	2.63	0.46
13:A1:86:ARG:NH1	13:A1:96:PRO:O	2.49	0.46
17:AA:990:U:H2'	17:AA:991:G:O4'	2.15	0.46
18:AB:180:ARG:NH1	20:AD:211:CYS:SG	2.89	0.46
23:AG:75:LYS:O	23:AG:79:GLU:OE1	2.33	0.46

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
29:AM:18:THR:N	29:AM:37:ALA:O	2.38	0.46
44:XD:86:ASP:OD2	44:XD:87:HIS:ND1	2.49	0.46
45:XE:316:PHE:HB3	45:XE:317:PRO:HD3	1.97	0.46
45:XE:327:GLU:OE1	45:XE:327:GLU:N	2.48	0.46
51:XL:99:ARG:HH12	56:XQ:191:ARG:HE	1.64	0.46
70:e:158:VAL:O	70:e:255:VAL:N	2.42	0.46
76:k:9:GLY:O	76:k:46:ASN:ND2	2.49	0.46
79:o:33:GLN:HA	79:o:36:ILE:HG12	1.97	0.46
85:r3:67:Y5P:H2'	85:r3:68:P5P:H8	1.97	0.46
4:3:150:CYS:SG	4:3:154:GLN:HB2	2.56	0.46
11:XA:1939:G:O2'	11:XA:1973:G:H4'	2.16	0.46
16:A4:66:ASP:OD1	16:A4:67:LYS:N	2.49	0.46
47:XH:120:ARG:NH2	63:XX:136:ASP:OD2	2.43	0.46
61:XV:108:MET:HE1	69:d:169:PHE:CD2	2.51	0.46
85:r3:29:P5P:H2'	85:r3:30:P5P:O4'	2.15	0.46
11:XA:2835:C:H4'	62:XW:50:ARG:HE	1.81	0.46
17:AA:662:U:H2'	17:AA:663:A:O4'	2.16	0.46
17:AA:1024:G:C4	17:AA:1026:A:OP2	2.69	0.46
38:AV:108:THR:O	38:AV:111:THR:OG1	2.30	0.46
40:AX:297:MET:O	40:AX:297:MET:HG2	2.16	0.46
40:AX:393:ARG:O	40:AX:397:TYR:CD2	2.69	0.46
45:XE:310:LEU:O	45:XE:310:LEU:HG	2.16	0.46
55:XP:65:ARG:O	62:XW:80:HIS:N	2.41	0.46
83:r1:56:Y5P:O2'	83:r1:57:Y5P:O5'	2.34	0.46
7:6:283:GLU:OE2	7:6:306:LYS:NZ	2.46	0.46
8:7:314:ASP:O	8:7:318:GLU:OE1	2.34	0.46
10:9:137:ARG:NH2	64:XY:143:ASP:OD1	2.49	0.46
11:XA:2290:A:O2'	11:XA:2291:A:O4'	2.31	0.46
11:XA:2594:U:O4	11:XA:2632:A:N7	2.49	0.46
11:XA:3212:C:O2	11:XA:3212:C:O4'	2.33	0.46
17:AA:1287:A:OP2	20:AD:260:LYS:NZ	2.37	0.46
17:AA:1526:U:O2'	17:AA:1526:U:O2	2.34	0.46
22:AF:155:MET:SD	22:AF:179:ARG:NH1	2.89	0.46
22:AF:201:MET:N	22:AF:202:PRO:HD2	2.31	0.46
35:AS:116:LYS:O	35:AS:120:GLU:OE1	2.34	0.46
52:XM:255:MET:O	52:XM:258:THR:OG1	2.34	0.46
59:XT:149:ARG:NH1	59:XT:152:CYS:SG	2.89	0.46
74:i:59:ASN:O	74:i:63:LEU:HD23	2.16	0.46
74:i:110:LEU:O	74:i:114:ILE:HG23	2.16	0.46
3:2:49:ARG:NH1	11:XA:1919:C:O2	2.49	0.46
11:XA:3019:G:N2	11:XA:3131:G:O2'	2.44	0.46

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
13:A1:52:ALA:HB2	16:A4:94:TYR:CA	2.46	0.46
17:AA:1484:C:C5	85:r3:34:Y5P:H6	2.51	0.46
34:AR:260:ASP:HA	34:AR:263:ARG:HG2	1.98	0.46
45:XE:301:ASP:OD1	45:XE:302:SER:N	2.49	0.46
50:XK:125:LEU:HD23	50:XK:127:LEU:HD22	1.98	0.46
79:o:33:GLN:C	79:o:37:ARG:HE	2.23	0.46
84:r2:74:Y5P:O2'	84:r2:75:Y5P:OP1	2.25	0.46
85:r3:2:Y5P:H6	85:r3:2:Y5P:HB	1.98	0.46
8:7:47:LEU:O	8:7:50:LYS:HG2	2.16	0.45
11:XA:1984:A:OP1	44:XD:90:ARG:NE	2.47	0.45
11:XA:2295:C:O2'	11:XA:2297:A:OP1	2.32	0.45
11:XA:2945:A:N6	11:XA:2977:G:O2'	2.49	0.45
17:AA:1211:G:N1	17:AA:1354:A:C6	2.84	0.45
17:AA:1221:A:OP2	17:AA:1222:A:O2'	2.26	0.45
46:XF:209:TYR:OH	73:h:56:ARG:NH2	2.49	0.45
58:XS:106:TRP:CD2	58:XS:114:ILE:HD11	2.51	0.45
60:XU:49:THR:O	60:XU:52:ASP:OD1	2.35	0.45
64:XY:134:LYS:O	64:XY:137:ASP:OD1	2.34	0.45
65:XZ:149:TRP:NE1	75:j:45:GLU:OE1	2.45	0.45
72:g:118:TRP:O	72:g:121:GLN:HG3	2.15	0.45
74:i:118:TYR:O	74:i:122:ASN:ND2	2.48	0.45
84:r2:9:P5P:H4'	84:r2:46:P5P:C4'	2.47	0.45
85:r3:68:P5P:H2'	85:r3:69:Y5P:O4'	2.16	0.45
8:7:38:THR:O	8:7:42:GLU:OE1	2.34	0.45
11:XA:2381:A:N6	11:XA:2412:A:N1	2.65	0.45
17:AA:843:G:N1	17:AA:847:G:O6	2.49	0.45
17:AA:1124:A:OP2	21:AE:121:LYS:NZ	2.40	0.45
11:XA:2910:A:H2	85:r3:74:P5P:H8	1.80	0.45
16:A4:164:ARG:H	16:A4:167:LYS:HE3	1.82	0.45
79:o:33:GLN:O	79:o:37:ARG:NE	2.46	0.45
86:s:403:GLU:OE2	86:s:404:THR:OG1	2.33	0.45
5:4:67:LYS:NZ	11:XA:2966:U:OP1	2.46	0.45
5:4:99:LYS:NZ	11:XA:3013:G:O3'	2.50	0.45
8:7:78:VAL:CG2	69:d:282:ILE:HD11	2.45	0.45
11:XA:3160:A:O3'	45:XE:220:LYS:NZ	2.49	0.45
13:A1:52:ALA:CB	16:A4:94:TYR:CA	2.95	0.45
15:A3:161:ARG:NH2	17:AA:1146:C:OP1	2.44	0.45
17:AA:1235:U:H5''	17:AA:1236:C:OP2	2.16	0.45
41:AY:377:ARG:O	41:AY:381:ASN:ND2	2.49	0.45
44:XD:251:ASP:OD1	44:XD:251:ASP:C	2.59	0.45
50:XK:136:ASP:OD1	68:c:264:THR:OG1	2.30	0.45

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
56:XQ:226:PRO:O	56:XQ:229:TRP:NE1	2.49	0.45
59:XT:56:LYS:HG2	69:d:178:TYR:CE1	2.52	0.45
59:XT:123:GLU:O	59:XT:126:ASP:OD1	2.34	0.45
61:XV:19:TYR:OH	61:XV:31:ASP:OD2	2.35	0.45
81:q:140:ARG:O	81:q:144:GLU:OE1	2.34	0.45
5:4:87:ARG:NH2	5:4:102:GLN:O	2.41	0.45
11:XA:1759:U:H2'	11:XA:1760:G:O4'	2.16	0.45
11:XA:2796:G:O2'	47:XH:85:ASP:OD2	2.29	0.45
17:AA:1067:A:H2'	17:AA:1068:A:O4'	2.16	0.45
24:AH:67:ASP:OD1	24:AH:67:ASP:N	2.50	0.45
34:AR:259:TYR:O	34:AR:263:ARG:HG2	2.17	0.45
52:XM:231:GLU:O	52:XM:235:GLU:OE1	2.34	0.45
54:XO:149:LEU:HA	54:XO:152:LEU:CD2	2.45	0.45
84:r2:48:Y5P:HA	84:r2:49:Y5P:OP2	2.17	0.45
8:7:94:HIS:O	59:XT:137:ARG:HG2	2.17	0.45
11:XA:1917:A:N3	11:XA:1921:A:N6	2.64	0.45
11:XA:2475:U:N3	11:XA:2478:G:OP2	2.48	0.45
11:XA:2948:C:N3	11:XA:2977:G:N2	2.61	0.45
70:e:186:GLY:HA2	70:e:189:GLU:OE2	2.17	0.45
75:j:98:LEU:O	75:j:101:GLU:HG3	2.16	0.45
1:0:177:ARG:HG3	1:0:179:ARG:HG2	1.99	0.45
11:XA:2538:C:H2'	11:XA:2539:A:O4'	2.16	0.45
11:XA:2945:A:O2'	11:XA:2947:U:O4	2.34	0.45
14:A2:113:ASN:OD1	14:A2:114:LYS:N	2.49	0.45
24:AH:75:ARG:N	24:AH:175:THR:OG1	2.49	0.45
37:AU:178:GLU:OE1	37:AU:178:GLU:N	2.44	0.45
55:XP:120:ASN:OD1	55:XP:123:ALA:N	2.39	0.45
63:XX:226:LEU:HA	63:XX:229:ILE:HG12	1.99	0.45
82:r:176:VAL:N	82:r:194:LEU:O	2.49	0.45
88:A:1:MHW:O	88:A:1:MHW:OG1	2.33	0.45
4:3:118:HIS:NE2	11:XA:1891:A:OP1	2.50	0.45
6:5:250:ASN:OD1	6:5:251:HIS:N	2.50	0.45
11:XA:1805:A:OP2	61:XV:94:HIS:NE2	2.50	0.45
11:XA:2509:A:H5'	44:XD:273:GLY:HA2	1.99	0.45
11:XA:2642:C:O2'	11:XA:2643:G:H5'	2.17	0.45
17:AA:1421:G:H3'	17:AA:1421:G:OP2	2.16	0.45
59:XT:176:VAL:HG12	59:XT:178:LEU:HD12	1.99	0.45
73:h:78:PHE:CB	73:h:80:SER:C	2.90	0.45
85:r3:65:Y5P:O5'	85:r3:65:Y5P:H6	2.17	0.45
11:XA:2373:A:OP1	60:XU:50:ARG:NH2	2.44	0.45
13:A1:216:ARG:NH1	41:AY:326:SER:O	2.47	0.45

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
21:AE:96:HIS:O	21:AE:100:GLN:NE2	2.48	0.45
24:AH:77:SER:HB2	24:AH:173:THR:OG1	2.17	0.45
43:XB:1620:A:H2'	43:XB:1620:A:N3	2.31	0.45
48:XI:30:SER:OG	48:XI:31:LYS:N	2.49	0.45
70:e:175:GLN:OE1	70:e:175:GLN:N	2.49	0.45
79:o:9:ARG:CZ	82:r:164:LYS:HD3	2.46	0.45
84:r2:4:Y5P:O5'	84:r2:4:Y5P:H6	2.17	0.45
86:s:415:ASP:N	86:s:415:ASP:OD1	2.49	0.45
11:XA:1994:A:O2'	11:XA:2002:G:N7	2.46	0.45
11:XA:2262:C:O2'	58:XS:175:ARG:NH2	2.50	0.45
11:XA:2835:C:C5'	62:XW:50:ARG:HE	2.29	0.45
38:AV:235:GLU:O	38:AV:239:GLY:N	2.49	0.45
48:XI:80:ARG:NH1	77:l:113:GLU:OE2	2.48	0.45
50:XK:10:GLN:NE2	59:XT:203:LEU:O	2.42	0.45
53:XN:51:ARG:HE	53:XN:52:PHE:H	1.64	0.45
64:XY:137:ASP:OD1	64:XY:138:SER:N	2.50	0.45
70:e:120:LEU:HD21	70:e:124:TRP:CH2	2.52	0.45
70:e:265:LYS:N	70:e:266:PRO:HD3	2.32	0.45
81:q:185:LYS:HD3	81:q:185:LYS:N	2.30	0.45
11:XA:1747:G:OP2	11:XA:1749:C:N4	2.46	0.44
17:AA:682:A:N6	17:AA:865:A:H61	2.14	0.44
38:AV:47:HIS:N	38:AV:78:ASN:OD1	2.49	0.44
43:XB:1611:G:O6	43:XB:1644:G:N2	2.43	0.44
54:XO:94:ALA:HB3	54:XO:95:PRO:HD3	1.98	0.44
86:s:145:VAL:O	86:s:148:ASP:OD1	2.35	0.44
11:XA:2379:C:O2	11:XA:2379:C:O4'	2.35	0.44
11:XA:2407:U:H2'	11:XA:2408:U:O4'	2.17	0.44
17:AA:663:A:H2'	17:AA:664:G:C8	2.52	0.44
25:AI:115:GLU:OE2	25:AI:131:ALA:N	2.50	0.44
46:XF:249:ASN:ND2	74:i:55:GLY:O	2.50	0.44
56:XQ:175:GLN:NE2	56:XQ:176:VAL:O	2.51	0.44
61:XV:108:MET:CE	69:d:169:PHE:CD2	3.00	0.44
2:1:20:MET:N	2:1:20:MET:SD	2.90	0.44
7:6:288:SER:O	7:6:289:PRO:C	2.59	0.44
11:XA:2701:G:C6	11:XA:2702:G:C5	3.05	0.44
16:A4:449:GLY:C	41:AY:292:GLN:NE2	2.74	0.44
28:AL:141:GLU:HA	28:AL:144:GLU:OE2	2.16	0.44
41:AY:367:LYS:O	41:AY:370:VAL:HG22	2.18	0.44
49:XJ:181:LEU:O	49:XJ:184:GLN:HG3	2.17	0.44
61:XV:196:GLU:O	61:XV:200:GLU:OE1	2.35	0.44
66:a:139:PRO:O	66:a:140:LYS:HE2	2.17	0.44

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
85:r3:24:Y5P:O5'	85:r3:24:Y5P:H6	2.16	0.44
3:2:54:GLN:HG2	3:2:54:GLN:O	2.18	0.44
6:5:103:ALA:HB1	6:5:267:GLU:OE2	2.17	0.44
11:XA:1904:C:C2	11:XA:2019:G:N2	2.86	0.44
11:XA:2144:A:OP1	57:XR:57:ARG:NH1	2.51	0.44
11:XA:2511:C:N4	11:XA:2540:C:O4'	2.50	0.44
11:XA:2550:A:C5	11:XA:2590:A:C6	3.06	0.44
17:AA:1193:U:O2'	22:AF:178:ARG:NH1	2.51	0.44
17:AA:1517:A:O2'	17:AA:1518:C:O4'	2.33	0.44
38:AV:201:GLU:OE1	38:AV:233:LYS:NZ	2.49	0.44
61:XV:105:ARG:NH1	69:d:171:ASP:CA	2.76	0.44
83:r1:52:Y5P:C6	83:r1:53:Y5P:H5	2.48	0.44
6:5:50:TYR:OH	6:5:72:ARG:O	2.28	0.44
7:6:83:ASP:OD2	7:6:87:LYS:NZ	2.44	0.44
7:6:159:ARG:NH2	7:6:160:ASP:OD1	2.51	0.44
11:XA:3107:U:O2'	45:XE:258:PRO:O	2.35	0.44
16:A4:482:ILE:CG2	16:A4:519:TYR:HE2	2.30	0.44
46:XF:194:GLU:N	46:XF:194:GLU:OE1	2.51	0.44
53:XN:201:ASP:OD1	53:XN:201:ASP:C	2.60	0.44
83:r1:49:Y5P:H4A	83:r1:50:Y5P:C4	2.47	0.44
85:r3:53:Y5P:H5	85:r3:54:Y5P:H4	1.99	0.44
86:s:375:ASN:N	86:s:391:ASN:OD1	2.34	0.44
8:7:259:ASP:OD1	8:7:260:PHE:N	2.50	0.44
11:XA:2877:C:H2'	11:XA:2878:G:O4'	2.18	0.44
13:A1:255:ASN:OD1	13:A1:256:SER:N	2.51	0.44
16:A4:643:GLU:O	16:A4:646:THR:OG1	2.34	0.44
17:AA:1268:C:OP2	17:AA:1269:U:O2'	2.27	0.44
22:AF:38:SER:OG	22:AF:40:GLU:OE1	2.32	0.44
34:AR:247:HIS:O	34:AR:251:GLU:OE1	2.36	0.44
54:XO:86:ILE:HB	54:XO:87:PRO:HD3	1.99	0.44
64:XY:224:GLU:O	64:XY:227:GLU:HG3	2.18	0.44
70:e:124:TRP:CH2	78:m:72:ARG:HG2	2.52	0.44
84:r2:37:P5P:H3'	84:r2:38:P5P:H8	1.99	0.44
84:r2:39:Y5P:H4A	84:r2:40:Y5P:H4	1.98	0.44
7:6:27:ARG:N	11:XA:2832:A:N1	2.66	0.44
11:XA:2182:G:H2'	11:XA:2183:C:C6	2.53	0.44
17:AA:1443:U:OP2	27:AK:102:ARG:NH1	2.51	0.44
20:AD:112:LYS:O	20:AD:239:LYS:NZ	2.51	0.44
40:AX:157:ASP:OD1	40:AX:158:ALA:N	2.51	0.44
74:i:79:TRP:CD1	74:i:80:LEU:HD23	2.52	0.44
80:p:178:ILE:HA	80:p:181:GLU:CD	2.43	0.44

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
82:r:117:GLU:O	82:r:121:MET:HG2	2.18	0.44
86:s:63:ILE:HA	86:s:66:TRP:CD1	2.52	0.44
11:XA:2939:C:O2'	11:XA:2940:A:H5'	2.18	0.44
17:AA:1108:C:N4	17:AA:1125:A:N7	2.65	0.44
29:AM:71:ASP:N	29:AM:71:ASP:OD1	2.50	0.44
30:AN:93:ASP:OD1	30:AN:96:THR:OG1	2.35	0.44
34:AR:295:ASP:OD1	34:AR:296:ASP:N	2.50	0.44
35:AS:36:ASP:OD1	35:AS:37:ALA:N	2.51	0.44
76:k:55:VAL:O	76:k:55:VAL:HG12	2.18	0.44
84:r2:59:Y5P:H4A	84:r2:60:Y5P:C2	2.47	0.44
6:5:242:ARG:HA	6:5:245:ILE:HG12	1.99	0.44
8:7:262:ASP:OD1	8:7:263:VAL:N	2.50	0.44
11:XA:2066:C:H2'	11:XA:2067:C:O5'	2.18	0.44
11:XA:2145:G:H1'	58:XS:104:ARG:NH2	2.32	0.44
11:XA:2704:A:OP2	50:XK:109:TYR:OH	2.24	0.44
13:A1:68:SER:HA	16:A4:78:VAL:CG1	2.48	0.44
17:AA:1052:C:OP1	28:AL:150:LYS:NZ	2.46	0.44
21:AE:105:CYS:O	32:AP:64:LYS:NZ	2.51	0.44
22:AF:77:ALA:N	23:AG:368:GLY:O	2.49	0.44
35:AS:90:PRO:O	35:AS:91:ASN:OD1	2.36	0.44
58:XS:92:TYR:O	68:c:312:ARG:NE	2.50	0.44
68:c:195:PHE:O	68:c:198:VAL:HG12	2.17	0.44
8:7:109:ASP:O	8:7:111:GLN:NE2	2.51	0.43
11:XA:1733:C:O4'	74:i:95:ARG:NH2	2.51	0.43
11:XA:2995:G:C2	11:XA:3069:A:N6	2.85	0.43
35:AS:61:GLN:NE2	35:AS:62:ASP:O	2.51	0.43
52:XM:191:VAL:HB	52:XM:192:PRO:HD3	2.00	0.43
53:XN:191:SER:H	53:XN:194:THR:HG1	1.62	0.43
64:XY:220:LYS:O	64:XY:224:GLU:OE1	2.35	0.43
84:r2:11:Y5P:H2'	84:r2:12:Y5P:C6	2.48	0.43
86:s:300:HIS:HB2	86:s:361:ILE:HG22	2.00	0.43
7:6:338:ARG:NH1	62:XW:147:MET:O	2.50	0.43
11:XA:2276:C:N3	11:XA:2290:A:C2	2.86	0.43
11:XA:2314:C:H2'	11:XA:2315:A:C8	2.53	0.43
11:XA:3180:A:H2'	11:XA:3181:U:C6	2.53	0.43
16:A4:634:ALA:HB3	16:A4:641:ILE:HG21	2.01	0.43
17:AA:805:C:O2	17:AA:805:C:O4'	2.36	0.43
38:AV:236:LEU:CD2	38:AV:327:LEU:HD11	2.49	0.43
46:XF:141:ILE:O	46:XF:142:ARG:HB2	2.18	0.43
55:XP:109:TRP:HA	55:XP:112:LYS:HG2	2.00	0.43
72:g:150:LYS:HB2	79:o:82:PHE:HZ	1.83	0.43

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
76:k:80:HIS:O	82:r:36:ARG:NH1	2.49	0.43
85:r3:15:P5P:H2	85:r3:57:P5P:C5	2.48	0.43
11:XA:2081:U:H2'	11:XA:2082:G:C8	2.53	0.43
11:XA:2508:C:H2'	11:XA:2509:A:C8	2.53	0.43
11:XA:2605:C:OP2	11:XA:2606:U:O2'	2.29	0.43
11:XA:2755:A:O2'	63:XX:112:ARG:NH2	2.50	0.43
13:A1:134:PRO:CG	16:A4:60:PRO:HD3	2.42	0.43
22:AF:35:SER:OG	22:AF:36:ARG:N	2.50	0.43
22:AF:228:LYS:HA	22:AF:231:GLU:OE2	2.18	0.43
30:AN:39:LEU:O	36:AT:11:ARG:NH1	2.51	0.43
56:XQ:64:ASP:OD1	56:XQ:64:ASP:N	2.51	0.43
75:j:98:LEU:O	75:j:102:GLN:OE1	2.36	0.43
7:6:228:PRO:O	7:6:229:ASP:OD1	2.36	0.43
11:XA:1877:U:O3'	52:XM:30:ASN:ND2	2.52	0.43
11:XA:1965:A:H2'	11:XA:1966:G:O4'	2.18	0.43
11:XA:2143:G:C2	11:XA:2258:A:C2	3.06	0.43
11:XA:2146:A:N6	67:b:116:ARG:O	2.51	0.43
35:AS:36:ASP:OD1	35:AS:36:ASP:C	2.60	0.43
49:XJ:75:ASP:O	49:XJ:76:ARG:HB3	2.19	0.43
61:XV:197:GLU:HA	61:XV:200:GLU:OE2	2.18	0.43
84:r2:32:Y5P:O5'	84:r2:32:Y5P:H6	2.19	0.43
4:3:183:ARG:NH1	7:6:355:LYS:O	2.43	0.43
5:4:70:THR:O	5:4:102:GLN:NE2	2.51	0.43
8:7:147:ALA:O	8:7:150:MET:HG2	2.17	0.43
11:XA:1788:C:N4	11:XA:1998:U:C4	2.86	0.43
17:AA:748:G:N2	30:AN:7:SER:O	2.46	0.43
20:AD:147:PRO:O	20:AD:155:GLN:NE2	2.52	0.43
21:AE:41:ASN:OD1	21:AE:43:GLY:N	2.51	0.43
23:AG:214:SER:O	23:AG:217:ASP:OD1	2.37	0.43
31:AO:163:LEU:HD23	31:AO:163:LEU:H	1.84	0.43
45:XE:227:GLN:NE2	45:XE:239:ARG:O	2.46	0.43
48:XI:50:VAL:O	53:XN:211:ASN:ND2	2.51	0.43
4:3:107:VAL:HG13	4:3:161:MET:CE	2.49	0.43
8:7:312:ILE:O	8:7:316:LEU:HD23	2.19	0.43
11:XA:1852:C:O4'	11:XA:2693:A:N7	2.52	0.43
11:XA:2182:G:N2	11:XA:2199:A:N3	2.66	0.43
17:AA:864:U:O4	17:AA:865:A:N6	2.51	0.43
19:AC:98:GLY:HA3	42:AZ:71:TYR:CE2	2.54	0.43
26:AJ:49:LEU:HD23	26:AJ:50:GLY:H	1.83	0.43
40:AX:350:PRO:O	40:AX:354:GLU:OE1	2.36	0.43
62:XW:115:ASP:OD1	62:XW:119:ARG:NH2	2.52	0.43

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
16:A4:273:GLU:OE2	16:A4:274:GLN:NE2	2.51	0.43
17:AA:661:C:OP1	20:AD:339:SER:OG	2.33	0.43
17:AA:1401:G:H3'	17:AA:1402:A:H5''	1.99	0.43
36:AT:7:PHE:HB2	36:AT:10:ARG:HE	1.82	0.43
36:AT:36:THR:O	36:AT:45:ARG:NE	2.52	0.43
42:AZ:65:LEU:HD22	42:AZ:71:TYR:HB2	2.01	0.43
48:XI:66:PRO:O	48:XI:67:SER:OG	2.34	0.43
69:d:131:ASN:OD1	69:d:131:ASN:N	2.47	0.43
6:5:189:LYS:NZ	86:s:182:SER:O	2.45	0.43
9:8:165:ASP:OD1	9:8:165:ASP:N	2.51	0.43
11:XA:2216:A:N3	48:XI:150:HIS:NE2	2.63	0.43
17:AA:701:G:O5'	37:AU:38:LYS:NZ	2.51	0.43
17:AA:865:A:H2'	17:AA:866:A:N9	2.33	0.43
18:AB:230:CYS:SG	18:AB:231:LEU:N	2.92	0.43
31:AO:65:GLN:O	31:AO:69:GLY:N	2.51	0.43
50:XK:73:GLU:OE1	50:XK:73:GLU:N	2.52	0.43
52:XM:254:LYS:O	52:XM:258:THR:HG23	2.19	0.43
55:XP:57:LEU:O	80:p:177:ARG:NH2	2.52	0.43
57:XR:65:ARG:O	57:XR:69:ILE:HG12	2.19	0.43
65:XZ:131:GLU:OE2	75:j:75:ARG:NH2	2.52	0.43
69:d:288:LYS:HG2	69:d:289:PRO:HD2	2.01	0.43
73:h:103:HIS:O	73:h:106:ASP:OD1	2.37	0.43
4:3:135:LYS:NZ	11:XA:2895:U:O5'	2.52	0.43
11:XA:2042:U:OP2	52:XM:57:ARG:NH1	2.52	0.43
11:XA:2347:C:O2'	69:d:69:LYS:NZ	2.49	0.43
11:XA:2373:A:C4	11:XA:2375:C:C5	3.07	0.43
11:XA:2529:U:H2'	44:XD:206:TYR:O	2.19	0.43
11:XA:3122:U:O2	11:XA:3122:U:O4'	2.36	0.43
12:A0:63:ARG:NH1	12:A0:110:ASP:OD2	2.47	0.43
16:A4:335:PHE:HA	16:A4:338:ILE:HG22	2.01	0.43
17:AA:1259:U:C5	84:r2:34:P5P:H1'	2.54	0.43
22:AF:116:GLU:O	22:AF:120:ARG:HG2	2.19	0.43
44:XD:204:ALA:HB1	44:XD:208:ARG:HE	1.83	0.43
46:XF:234:THR:O	81:q:25:TYR:N	2.52	0.43
48:XI:181:ILE:O	48:XI:182:ASP:OD1	2.37	0.43
49:XJ:127:ASP:O	49:XJ:131:ALA:N	2.41	0.43
53:XN:138:HIS:ND1	53:XN:139:ARG:O	2.52	0.43
56:XQ:64:ASP:O	56:XQ:65:LYS:HE2	2.19	0.43
61:XV:50:SER:O	61:XV:81:ARG:NH1	2.51	0.43
74:i:80:LEU:O	74:i:80:LEU:HD12	2.18	0.43
6:5:393:LYS:O	6:5:396:VAL:HG12	2.19	0.43

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:1864:A:C6	11:XA:1865:C:C4	3.07	0.43
11:XA:2094:G:OP2	52:XM:57:ARG:NH2	2.52	0.43
11:XA:2096:U:O4	52:XM:57:ARG:NH1	2.39	0.43
11:XA:2672:A:C2	11:XA:2673:G:C8	3.07	0.43
11:XA:2725:A:C5	88:A:1:MHW:CE	3.01	0.43
13:A1:52:ALA:HB1	16:A4:94:TYR:N	2.34	0.43
17:AA:1486:C:O2	17:AA:1486:C:O5'	2.37	0.43
17:AA:1563:U:H2'	83:r1:47:Y5P:OP1	2.19	0.43
28:AL:169:ASN:OD1	28:AL:169:ASN:C	2.62	0.43
34:AR:176:GLU:OE1	34:AR:176:GLU:N	2.52	0.43
44:XD:204:ALA:CB	44:XD:208:ARG:HE	2.32	0.43
53:XN:198:MET:O	53:XN:201:ASP:OD1	2.37	0.43
57:XR:134:ASP:OD1	57:XR:134:ASP:N	2.51	0.43
67:b:28:ARG:NH2	67:b:78:GLU:OE1	2.52	0.43
86:s:350:ASP:OD1	86:s:351:VAL:N	2.51	0.43
3:2:66:TRP:CG	11:XA:1702:A:C8	3.07	0.42
11:XA:1827:C:C5	11:XA:2698:G:C2	3.07	0.42
11:XA:2041:U:OP1	52:XM:57:ARG:NE	2.52	0.42
11:XA:2814:G:C5	53:XN:140:MET:HE3	2.54	0.42
17:AA:1554:G:H2'	17:AA:1555:A:O4'	2.18	0.42
44:XD:216:LEU:HD23	44:XD:216:LEU:H	1.84	0.42
56:XQ:182:ARG:HE	56:XQ:220:LEU:HD21	1.83	0.42
61:XV:147:SER:OG	61:XV:150:SER:O	2.31	0.42
68:c:42:GLN:NE2	74:i:36:LEU:O	2.52	0.42
68:c:128:GLN:O	68:c:131:SER:OG	2.37	0.42
8:7:306:LEU:O	8:7:306:LEU:HG	2.19	0.42
11:XA:2151:A:H2'	11:XA:2152:A:C8	2.54	0.42
11:XA:2952:U:C2	11:XA:2953:U:C5	3.07	0.42
11:XA:3096:U:C5	88:A:7:004:O	2.72	0.42
11:XA:3189:C:C2'	11:XA:3190:A:OP2	2.67	0.42
15:A3:161:ARG:NH1	17:AA:1147:G:OP2	2.53	0.42
20:AD:97:GLU:OE1	20:AD:97:GLU:N	2.51	0.42
54:XO:110:ILE:HB	54:XO:111:PRO:CD	2.50	0.42
63:XX:163:ARG:NH2	63:XX:205:GLY:O	2.53	0.42
66:a:110:GLU:O	66:a:114:LYS:HG2	2.19	0.42
67:b:14:VAL:C	67:b:15:LEU:HD23	2.44	0.42
83:r1:56:Y5P:HD	83:r1:57:Y5P:P	2.41	0.42
84:r2:8:Y5P:N3	84:r2:14:P5P:H8	2.34	0.42
84:r2:10:P5P:C6	84:r2:11:Y5P:H4	2.48	0.42
84:r2:33:Y5P:H2'	84:r2:33:Y5P:H6	1.77	0.42
6:5:311:ALA:O	6:5:315:LEU:HD23	2.18	0.42

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:1883:G:OP2	52:XM:100:ARG:NH2	2.43	0.42
11:XA:1903:C:O2'	11:XA:1904:C:H5'	2.19	0.42
11:XA:2672:A:C2	11:XA:2673:G:N7	2.86	0.42
11:XA:3068:G:C2	11:XA:3097:U:C4	3.06	0.42
11:XA:3181:U:OP2	11:XA:3182:A:O2'	2.29	0.42
11:XA:3188:U:H4'	11:XA:3189:C:O5'	2.20	0.42
15:A3:171:LYS:O	15:A3:175:ARG:HG2	2.18	0.42
23:AG:276:ARG:HG3	23:AG:277:LYS:H	1.84	0.42
28:AL:175:TYR:O	28:AL:179:GLU:OE1	2.37	0.42
31:AO:81:HIS:ND1	31:AO:82:LYS:O	2.52	0.42
56:XQ:161:GLU:OE1	56:XQ:161:GLU:N	2.53	0.42
60:XU:58:GLU:OE2	60:XU:65:VAL:N	2.52	0.42
66:a:118:THR:HG21	66:a:122:ARG:HD3	2.01	0.42
68:c:42:GLN:O	68:c:45:LEU:HB3	2.20	0.42
79:o:47:TYR:O	79:o:50:SER:OG	2.36	0.42
82:r:65:ASN:ND2	82:r:73:CYS:O	2.52	0.42
83:r1:46:Y5P:H4A	85:r3:36:Y5P:N1	2.34	0.42
86:s:427:ASN:OD1	86:s:428:ARG:N	2.52	0.42
9:8:131:MET:O	9:8:134:ASP:OD1	2.37	0.42
11:XA:2235:C:O2'	82:r:165:LYS:N	2.48	0.42
11:XA:2376:A:C6	11:XA:2421:G:O6	2.72	0.42
11:XA:2546:G:C2	11:XA:2547:C:C5	3.08	0.42
11:XA:2942:C:C2	11:XA:2943:G:N7	2.87	0.42
11:XA:3148:C:OP1	45:XE:211:ILE:HG12	2.19	0.42
13:A1:52:ALA:CB	16:A4:94:TYR:HA	2.49	0.42
13:A1:53:LEU:CD1	16:A4:93:PRO:HB3	2.50	0.42
13:A1:211:ARG:NH2	41:AY:359:SER:OG	2.52	0.42
17:AA:920:G:C2	17:AA:921:U:C4	3.07	0.42
17:AA:1251:A:N6	17:AA:1341:C:O2'	2.52	0.42
17:AA:1347:G:OP1	27:AK:36:ARG:NH1	2.45	0.42
28:AL:118:ASN:ND2	28:AL:121:ASP:OD2	2.52	0.42
45:XE:121:LEU:C	45:XE:122:LEU:HD22	2.45	0.42
63:XX:28:TYR:OH	63:XX:206:LEU:O	2.34	0.42
77:l:132:LEU:HA	77:l:135:ASN:ND2	2.34	0.42
82:r:83:TYR:HA	82:r:118:CYS:SG	2.59	0.42
86:s:209:TYR:HB3	86:s:282:ILE:HD11	2.01	0.42
86:s:407:ASP:O	86:s:408:ASN:OD1	2.38	0.42
6:5:83:LYS:NZ	11:XA:2412:A:O2'	2.50	0.42
14:A2:10:LEU:O	14:A2:15:ASN:ND2	2.41	0.42
16:A4:256:GLU:O	16:A4:257:HIS:C	2.61	0.42
16:A4:372:TYR:O	16:A4:376:ILE:HG12	2.19	0.42

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
17:AA:1132:U:H2'	17:AA:1133:C:C6	2.54	0.42
21:AE:15:ARG:HA	21:AE:18:THR:OG1	2.20	0.42
32:AP:134:LYS:O	37:AU:193:ARG:NE	2.38	0.42
60:XU:28:LEU:N	64:XY:114:THR:HG22	2.34	0.42
66:a:62:HIS:ND1	66:a:62:HIS:O	2.52	0.42
73:h:135:GLN:N	73:h:135:GLN:OE1	2.52	0.42
85:r3:2:Y5P:O3'	85:r3:3:P5P:O4'	2.37	0.42
86:s:92:MET:SD	86:s:276:ARG:NE	2.88	0.42
11:XA:2558:A:C4'	11:XA:2559:U:OP2	2.68	0.42
16:A4:62:LYS:N	24:AH:70:ASP:N	2.67	0.42
42:AZ:80:ASP:OD1	42:AZ:81:GLU:N	2.53	0.42
44:XD:204:ALA:O	44:XD:208:ARG:NH2	2.49	0.42
52:XM:51:ARG:HD2	74:i:126:LYS:HD2	2.02	0.42
55:XP:177:ILE:O	55:XP:178:TYR:C	2.63	0.42
76:k:65:ASP:OD1	76:k:66:VAL:N	2.52	0.42
83:r1:49:Y5P:H6	83:r1:49:Y5P:HB	2.00	0.42
84:r2:4:Y5P:C2	84:r2:70:P5P:H2	2.50	0.42
86:s:66:TRP:O	86:s:69:THR:OG1	2.28	0.42
11:XA:2334:C:O2	86:s:277:GLN:NE2	2.53	0.42
11:XA:2726:C:O2	11:XA:2726:C:H2'	2.19	0.42
12:A0:61:GLU:OE2	12:A0:139:TRP:N	2.52	0.42
17:AA:1343:A:H2'	17:AA:1343:A:N3	2.35	0.42
23:AG:200:LEU:O	23:AG:218:TYR:OH	2.32	0.42
30:AN:66:LEU:HD13	30:AN:79:HIS:HB3	2.01	0.42
30:AN:98:LYS:NZ	30:AN:99:PRO:O	2.50	0.42
40:AX:337:LEU:O	40:AX:337:LEU:HG	2.19	0.42
48:XI:112:MET:O	48:XI:116:LEU:HD23	2.20	0.42
58:XS:164:THR:CG2	67:b:107:GLN:HA	2.49	0.42
76:k:10:LEU:O	76:k:46:ASN:ND2	2.53	0.42
85:r3:3:P5P:H2	85:r3:70:Y5P:C2	2.50	0.42
1:0:132:LYS:CE	69:d:288:LYS:HZ1	2.26	0.42
1:0:138:ARG:HA	1:0:141:ILE:HG12	2.01	0.42
3:2:54:GLN:OE1	11:XA:1918:G:O2'	2.38	0.42
8:7:38:THR:O	8:7:41:THR:OG1	2.32	0.42
9:8:159:ALA:O	9:8:163:LYS:HG2	2.20	0.42
11:XA:1764:C:H3'	11:XA:1765:C:C5'	2.50	0.42
11:XA:2044:A:C4	11:XA:2045:A:C8	3.08	0.42
11:XA:2154:A:OP2	79:o:28:SER:OG	2.31	0.42
11:XA:2337:A:C4	11:XA:2338:A:C8	3.08	0.42
16:A4:116:VAL:HG12	19:AC:138:TYR:CD1	2.54	0.42
16:A4:616:ASP:HA	16:A4:619:LYS:HG2	2.01	0.42

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
17:AA:922:C:H1'	17:AA:923:A:N7	2.35	0.42
31:AO:137:ALA:HB3	31:AO:138:PRO:HD3	2.01	0.42
49:XJ:32:GLY:O	49:XJ:36:GLY:N	2.41	0.42
56:XQ:225:LYS:CG	56:XQ:226:PRO:HD2	2.50	0.42
70:e:124:TRP:O	70:e:127:LYS:HG3	2.20	0.42
72:g:137:VAL:O	72:g:137:VAL:HG12	2.20	0.42
79:o:45:ASN:O	79:o:49:LEU:HD23	2.19	0.42
82:r:50:GLU:O	82:r:52:ARG:NH1	2.53	0.42
86:s:243:ILE:N	86:s:294:TYR:O	2.44	0.42
6:5:270:ILE:HG22	6:5:270:ILE:O	2.19	0.42
7:6:379:ILE:O	7:6:380:TYR:CG	2.73	0.42
8:7:315:LYS:HA	8:7:318:GLU:CD	2.45	0.42
11:XA:1748:G:C5	11:XA:1750:G:N2	2.88	0.42
11:XA:2349:G:H2'	11:XA:2350:A:C8	2.54	0.42
13:A1:100:GLU:O	19:AC:156:GLN:NE2	2.48	0.42
17:AA:991:G:N1	17:AA:992:U:O4	2.53	0.42
17:AA:1462:G:C2	17:AA:1463:G:C5	3.08	0.42
23:AG:140:TRP:HA	23:AG:146:PRO:HA	2.01	0.42
35:AS:15:ARG:O	35:AS:18:ASP:OD1	2.38	0.42
52:XM:19:LEU:HB3	52:XM:20:PRO:HD2	2.01	0.42
52:XM:185:ASP:OD1	52:XM:185:ASP:C	2.61	0.42
52:XM:209:GLU:OE1	52:XM:209:GLU:N	2.53	0.42
53:XN:172:VAL:HG13	53:XN:175:PHE:CZ	2.55	0.42
64:XY:230:LYS:HA	64:XY:233:ILE:HG12	2.02	0.42
71:f:122:GLU:N	71:f:122:GLU:OE1	2.52	0.42
73:h:84:SER:OG	73:h:85:ASN:N	2.52	0.42
85:r3:63:Y5P:OP2	85:r3:63:Y5P:H6	2.20	0.42
4:3:150:CYS:SG	4:3:154:GLN:NE2	2.84	0.42
8:7:237:VAL:O	8:7:241:GLU:OE1	2.38	0.42
17:AA:1516:G:C6	17:AA:1517:A:N6	2.88	0.42
24:AH:54:VAL:O	24:AH:54:VAL:HG23	2.20	0.42
34:AR:106:MET:HB2	34:AR:110:GLN:HB2	2.02	0.42
38:AV:144:PHE:CZ	38:AV:167:VAL:HG21	2.55	0.42
46:XF:228:GLN:HA	46:XF:231:VAL:HG12	2.02	0.42
48:XI:166:ARG:HA	48:XI:169:ARG:HG2	2.01	0.42
59:XT:211:THR:OG1	59:XT:212:LEU:N	2.47	0.42
61:XV:190:CYS:O	61:XV:191:LEU:HB3	2.20	0.42
64:XY:144:LYS:O	64:XY:148:GLU:OE1	2.36	0.42
72:g:55:THR:HG22	72:g:55:THR:O	2.20	0.42
74:i:91:CYS:SG	74:i:114:ILE:HG22	2.60	0.42
86:s:243:ILE:O	86:s:294:TYR:N	2.53	0.42

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
86:s:294:TYR:OH	86:s:353:ARG:NH2	2.52	0.42
10:9:17:ARG:NH2	11:XA:1789:A:OP1	2.51	0.41
11:XA:1845:C:HO2'	11:XA:2297:A:H61	1.68	0.41
11:XA:2146:A:C4	57:XR:64:MET:HE1	2.55	0.41
11:XA:2390:A:C8	11:XA:2403:G:C5	3.08	0.41
11:XA:2574:G:O2'	11:XA:2575:U:OP1	2.35	0.41
11:XA:2871:U:C2	11:XA:2872:C:C5	3.08	0.41
16:A4:638:SER:OG	16:A4:640:PRO:HD2	2.20	0.41
17:AA:1161:A:C2	17:AA:1162:A:C8	3.08	0.41
17:AA:1559:G:H2'	17:AA:1560:U:C6	2.55	0.41
23:AG:263:ASP:OD1	23:AG:267:MET:HB2	2.20	0.41
31:AO:106:PRO:HA	31:AO:109:ARG:HG2	2.01	0.41
42:AZ:77:ASP:O	42:AZ:80:ASP:OD1	2.38	0.41
45:XE:58:VAL:HB	45:XE:59:PRO:HD3	2.02	0.41
46:XF:87:PHE:CZ	46:XF:178:LEU:HD21	2.55	0.41
47:XH:70:LYS:HD2	47:XH:71:PRO:O	2.19	0.41
51:XL:78:LYS:HG3	51:XL:78:LYS:O	2.20	0.41
52:XM:103:TYR:OH	72:g:93:ASN:ND2	2.47	0.41
68:c:216:ARG:O	68:c:217:ASP:C	2.62	0.41
72:g:150:LYS:HB2	79:o:82:PHE:CZ	2.55	0.41
83:r1:50:Y5P:C4	84:r2:35:P5P:H2	2.44	0.41
85:r3:25:Y5P:H2'	85:r3:26:P5P:C8	2.48	0.41
9:8:134:ASP:OD1	9:8:134:ASP:C	2.63	0.41
11:XA:2029:A:O2'	11:XA:2124:A:N1	2.54	0.41
11:XA:3217:A:H4'	54:XO:11:HIS:ND1	2.35	0.41
16:A4:243:ASN:O	16:A4:247:ILE:HG12	2.19	0.41
17:AA:917:C:OP2	31:AO:91:ARG:NH2	2.53	0.41
17:AA:1053:A:N1	17:AA:1100:C:O2'	2.52	0.41
17:AA:1390:A:H1'	17:AA:1392:A:N7	2.35	0.41
40:AX:374:GLU:HG2	40:AX:375:GLU:N	2.35	0.41
55:XP:75:GLU:OE1	55:XP:75:GLU:N	2.53	0.41
69:d:244:GLU:OE1	69:d:244:GLU:N	2.52	0.41
72:g:109:VAL:HG12	72:g:148:ARG:HA	2.02	0.41
83:r1:50:Y5P:H2'	83:r1:51:Y5P:O4'	2.20	0.41
1:0:85:ARG:NH2	11:XA:3102:U:O3'	2.41	0.41
2:1:34:ARG:NH2	2:1:35:ASN:O	2.54	0.41
2:1:35:ASN:OD1	2:1:36:ARG:N	2.53	0.41
6:5:180:ILE:O	6:5:184:LEU:HD23	2.20	0.41
11:XA:1713:A:H2'	11:XA:1714:C:C6	2.55	0.41
11:XA:2319:A:O3'	54:XO:42:ILE:HD11	2.19	0.41
13:A1:291:GLU:O	13:A1:294:LYS:HG3	2.20	0.41

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
16:A4:62:LYS:HA	24:AH:70:ASP:CB	2.45	0.41
16:A4:319:LEU:HA	16:A4:322:HIS:CD2	2.55	0.41
17:AA:896:A:H2'	17:AA:897:C:C6	2.55	0.41
38:AV:79:ILE:CD1	38:AV:88:ALA:HB2	2.50	0.41
83:r1:49:Y5P:C4	83:r1:50:Y5P:H5	2.50	0.41
7:6:360:ARG:NH1	11:XA:2870:G:O6	2.53	0.41
8:7:82:ASN:O	8:7:86:SER:OG	2.36	0.41
8:7:315:LYS:HA	8:7:318:GLU:OE2	2.21	0.41
11:XA:1770:G:OP2	57:XR:11:ARG:NH1	2.50	0.41
11:XA:1853:A:OP2	58:XS:177:ARG:NH2	2.43	0.41
11:XA:2051:A:OP2	79:o:81:LYS:NZ	2.45	0.41
11:XA:2153:A:C5	79:o:30:ARG:HD3	2.56	0.41
11:XA:2841:U:C4	11:XA:2842:C:C5	3.08	0.41
11:XA:3127:G:C2	11:XA:3129:A:OP2	2.74	0.41
11:XA:3215:C:N3	11:XA:3226:G:N1	2.67	0.41
14:A2:44:THR:HG22	14:A2:45:CYS:N	2.35	0.41
14:A2:66:CYS:O	14:A2:67:ARG:C	2.61	0.41
23:AG:167:GLY:O	23:AG:171:ASN:ND2	2.53	0.41
38:AV:57:MET:O	38:AV:61:PHE:CD2	2.73	0.41
54:XO:113:ARG:HB3	54:XO:116:ASP:OD1	2.21	0.41
56:XQ:168:ASN:O	56:XQ:171:VAL:HG22	2.20	0.41
61:XV:148:THR:O	61:XV:149:ARG:C	2.64	0.41
64:XY:233:ILE:HG13	64:XY:234:LEU:N	2.35	0.41
66:a:109:ILE:HG13	66:a:110:GLU:N	2.35	0.41
70:e:258:ASP:OD1	70:e:259:GLU:N	2.52	0.41
11:XA:2058:C:N3	11:XA:2081:U:O4	2.53	0.41
11:XA:2065:A:C4	11:XA:2066:C:C5	3.09	0.41
11:XA:2192:A:OP2	77:l:119:ARG:NH1	2.53	0.41
11:XA:2976:G:H2'	11:XA:2977:G:O4'	2.20	0.41
11:XA:3125:A:H2'	11:XA:3126:C:O4'	2.21	0.41
16:A4:455:ASN:HA	16:A4:486:TYR:CE1	2.55	0.41
16:A4:561:SER:O	16:A4:563:PRO:HD3	2.21	0.41
19:AC:88:GLU:O	19:AC:92:LEU:HD23	2.21	0.41
38:AV:275:LEU:HD13	38:AV:353:LEU:HB3	2.02	0.41
48:XI:197:LEU:O	48:XI:198:PRO:C	2.63	0.41
60:XU:150:TRP:NE1	69:d:172:MET:HE2	2.34	0.41
62:XW:103:VAL:HG12	62:XW:104:TYR:N	2.35	0.41
64:XY:231:ALA:HA	64:XY:234:LEU:CD2	2.51	0.41
68:c:164:GLU:HG2	68:c:165:VAL:N	2.35	0.41
76:k:8:LEU:HD12	76:k:10:LEU:H	1.85	0.41
85:r3:2:Y5P:H4A	85:r3:70:Y5P:N3	2.35	0.41

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
85:r3:36:Y5P:H2'	85:r3:37:P5P:C8	2.50	0.41
87:t6:9:ASP:O	87:t6:13:LEU:HG	2.20	0.41
1:0:113:CYS:SG	1:0:114:GLY:N	2.94	0.41
11:XA:2279:U:OP1	46:XF:255:LYS:NZ	2.41	0.41
11:XA:2674:U:OP1	59:XT:112:LYS:NZ	2.46	0.41
14:A2:18:LYS:NZ	17:AA:1028:G:O6	2.45	0.41
16:A4:601:LYS:HA	16:A4:601:LYS:HE3	2.03	0.41
17:AA:777:G:C2	17:AA:778:C:C6	3.09	0.41
19:AC:52:THR:OG1	19:AC:55:GLU:OE1	2.39	0.41
37:AU:102:HIS:O	37:AU:106:MET:SD	2.79	0.41
43:XB:1607:U:O2'	43:XB:1608:G:H5'	2.20	0.41
54:XO:22:PRO:O	54:XO:25:ARG:HG2	2.20	0.41
59:XT:59:TYR:N	69:d:227:ARG:O	2.52	0.41
59:XT:83:SER:HB3	59:XT:86:LYS:HD3	2.03	0.41
70:e:261:GLY:HA2	70:e:273:ARG:NH2	2.35	0.41
72:g:96:VAL:HG12	72:g:110:ILE:CD1	2.51	0.41
73:h:93:ASP:OD1	73:h:93:ASP:N	2.53	0.41
76:k:9:GLY:C	76:k:10:LEU:HD23	2.46	0.41
86:s:421:ILE:O	86:s:425:LEU:HD23	2.20	0.41
1:0:113:CYS:SG	1:0:115:HIS:N	2.83	0.41
6:5:143:PRO:HA	6:5:146:HIS:ND1	2.35	0.41
11:XA:2045:A:C5	11:XA:2046:C:C5	3.09	0.41
11:XA:2674:U:C4	11:XA:2675:G:C5	3.09	0.41
11:XA:2943:G:C6	11:XA:2944:C:N4	2.89	0.41
11:XA:3217:A:O4'	56:XQ:86:ARG:NH2	2.52	0.41
16:A4:640:PRO:O	16:A4:643:GLU:HG2	2.20	0.41
17:AA:1146:C:C2	17:AA:1147:G:C8	3.09	0.41
17:AA:1428:G:H4'	23:AG:388:ARG:HG3	2.03	0.41
20:AD:273:ASN:OD1	20:AD:277:HIS:NE2	2.54	0.41
26:AJ:49:LEU:HD23	26:AJ:50:GLY:N	2.35	0.41
28:AL:137:ARG:O	28:AL:140:GLU:HG2	2.21	0.41
46:XF:77:VAL:HG13	46:XF:77:VAL:O	2.21	0.41
50:XK:7:ALA:HB3	50:XK:8:PRO:HD3	2.02	0.41
55:XP:66:GLY:N	55:XP:75:GLU:OE2	2.51	0.41
56:XQ:70:GLU:HG3	56:XQ:71:PRO:HD3	2.02	0.41
59:XT:69:ARG:HD2	69:d:228:PHE:HB3	2.03	0.41
4:3:152:LYS:O	4:3:156:LYS:HG2	2.20	0.41
11:XA:2156:A:N7	82:r:173:ARG:CZ	2.83	0.41
11:XA:2459:A:C4	11:XA:2460:A:C8	3.08	0.41
11:XA:2589:A:C4	11:XA:2591:A:OP2	2.74	0.41
11:XA:2867:C:H2'	11:XA:2868:C:O4'	2.20	0.41

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
11:XA:3058:U:O3'	45:XE:247:ASP:OD2	2.39	0.41
12:A0:135:MET:SD	12:A0:135:MET:N	2.89	0.41
16:A4:80:ARG:NH1	24:AH:53:ASP:OD2	2.53	0.41
20:AD:287:ASP:OD2	20:AD:330:LYS:NZ	2.54	0.41
35:AS:33:ASP:O	35:AS:36:ASP:OD1	2.39	0.41
49:XJ:30:MET:HB2	49:XJ:31:PRO:HD3	2.03	0.41
59:XT:174:TYR:CE2	59:XT:176:VAL:CG2	3.03	0.41
60:XU:109:ASP:OD1	60:XU:110:LEU:N	2.54	0.41
68:c:195:PHE:O	68:c:199:ILE:HG12	2.21	0.41
68:c:259:ARG:HB2	68:c:271:PHE:HB2	2.03	0.41
80:p:178:ILE:HA	80:p:181:GLU:OE2	2.20	0.41
85:r3:4:P5P:O2'	85:r3:5:P5P:H8	2.20	0.41
10:9:123:GLN:O	10:9:129:GLN:NE2	2.47	0.41
11:XA:1936:A:O5'	11:XA:1937:A:N6	2.54	0.41
11:XA:1962:A:C5	11:XA:2501:C:O4'	2.73	0.41
11:XA:1983:U:H2'	11:XA:1988:G:O2'	2.21	0.41
11:XA:2034:A:H2'	11:XA:2035:U:O4'	2.21	0.41
11:XA:2080:U:O2'	79:o:60:ARG:O	2.36	0.41
11:XA:2188:A:N1	11:XA:2198:A:H2'	2.36	0.41
11:XA:2339:G:C2	11:XA:2340:C:C5	3.09	0.41
11:XA:2548:C:C2	11:XA:2568:G:N2	2.89	0.41
11:XA:2553:G:N2	44:XD:280:GLY:O	2.49	0.41
11:XA:2869:A:C5	11:XA:2870:G:C8	3.09	0.41
11:XA:3180:A:C4	11:XA:3190:A:C6	3.09	0.41
13:A1:131:THR:HG23	16:A4:63:LYS:HE3	2.01	0.41
16:A4:104:ARG:HA	16:A4:107:LEU:CD2	2.51	0.41
16:A4:449:GLY:CA	41:AY:292:GLN:CD	2.83	0.41
16:A4:639:LEU:N	16:A4:640:PRO:CD	2.83	0.41
17:AA:663:A:H2'	17:AA:664:G:H8	1.86	0.41
17:AA:1054:A:C2	17:AA:1055:U:H1'	2.55	0.41
17:AA:1231:A:OP1	27:AK:98:ARG:NE	2.54	0.41
17:AA:1323:G:N7	19:AC:36:LYS:NZ	2.61	0.41
17:AA:1367:A:N1	17:AA:1388:C:O4'	2.54	0.41
31:AO:105:CYS:HB2	31:AO:106:PRO:HD2	2.01	0.41
33:AQ:70:ARG:O	33:AQ:73:ASN:OD1	2.39	0.41
35:AS:69:ARG:HG3	35:AS:70:ILE:HD12	2.03	0.41
40:AX:371:ALA:N	40:AX:372:PRO:CD	2.84	0.41
44:XD:234:MET:HA	44:XD:293:LYS:O	2.20	0.41
45:XE:145:LEU:HD13	45:XE:181:ILE:HG21	2.03	0.41
46:XF:109:ILE:O	46:XF:109:ILE:HG13	2.20	0.41
46:XF:141:ILE:HD12	46:XF:146:TRP:CZ3	2.56	0.41

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
50:XK:135:GLU:HA	50:XK:138:LEU:CD2	2.51	0.41
56:XQ:246:ASP:OD1	56:XQ:247:LEU:N	2.54	0.41
58:XS:90:GLY:O	68:c:312:ARG:NH2	2.53	0.41
59:XT:212:LEU:HA	67:b:119:PHE:HE1	1.85	0.41
61:XV:171:ILE:O	64:XY:80:LYS:NZ	2.53	0.41
64:XY:130:GLU:OE1	64:XY:130:GLU:N	2.41	0.41
67:b:80:LEU:HB3	68:c:172:ASN:O	2.21	0.41
69:d:218:THR:OG1	69:d:220:GLN:NE2	2.53	0.41
73:h:78:PHE:HB2	73:h:81:SER:N	2.35	0.41
79:o:71:PHE:CE2	79:o:75:LYS:HD2	2.56	0.41
11:XA:1795:A:O2'	11:XA:1796:A:H5'	2.21	0.41
11:XA:2719:G:C2	11:XA:2720:A:N7	2.89	0.41
11:XA:2919:A:C6	63:XX:97:LEU:HD12	2.56	0.41
11:XA:3075:G:C6	11:XA:3094:G:N1	2.88	0.41
12:A0:71:LEU:HB2	12:A0:72:PRO:HD2	2.03	0.41
17:AA:905:A:H4'	20:AD:117:ARG:HG2	2.02	0.41
18:AB:135:MET:O	18:AB:140:GLY:N	2.46	0.41
19:AC:86:THR:O	19:AC:89:ASP:OD1	2.39	0.41
30:AN:30:VAL:O	30:AN:47:LYS:N	2.38	0.41
34:AR:135:ARG:NH1	34:AR:236:GLU:OE2	2.53	0.41
37:AU:39:ILE:HG13	37:AU:40:GLU:N	2.36	0.41
38:AV:90:TYR:CZ	56:XQ:59:LYS:HB3	2.55	0.41
40:AX:108:LEU:HD23	40:AX:112:LEU:HD23	2.02	0.41
44:XD:250:VAL:O	44:XD:251:ASP:CG	2.64	0.41
45:XE:209:LYS:HA	45:XE:264:ILE:O	2.20	0.41
46:XF:220:ASP:OD1	46:XF:221:LEU:N	2.50	0.41
46:XF:280:TYR:OH	52:XM:124:GLY:C	2.64	0.41
61:XV:49:ILE:HG21	61:XV:54:TRP:HE3	1.86	0.41
63:XX:148:THR:O	63:XX:148:THR:HG23	2.20	0.41
71:f:163:SER:OG	71:f:164:ALA:N	2.53	0.41
85:r3:54:Y5P:H6	85:r3:54:Y5P:O5'	2.20	0.41
86:s:217:ILE:CD1	86:s:222:ARG:HE	2.34	0.41
6:5:267:GLU:OE2	6:5:268:CYS:N	2.53	0.40
8:7:276:PHE:H	8:7:303:PRO:HA	1.86	0.40
8:7:313:TRP:O	8:7:317:LEU:HD23	2.21	0.40
11:XA:2072:A:H2'	11:XA:2073:A:C8	2.55	0.40
11:XA:2727:C:O2'	11:XA:2815:G:N2	2.46	0.40
11:XA:3064:A:O4'	11:XA:3099:C:N4	2.54	0.40
13:A1:254:GLU:OE2	13:A1:255:ASN:ND2	2.53	0.40
15:A3:171:LYS:O	15:A3:174:ARG:HG2	2.21	0.40
15:A3:174:ARG:HA	15:A3:177:TRP:CE2	2.55	0.40

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
17:AA:765:C:O2	17:AA:765:C:C2'	2.69	0.40
17:AA:873:G:O2'	17:AA:921:U:O2	2.32	0.40
17:AA:1192:C:C5	17:AA:1193:U:C6	3.09	0.40
19:AC:84:GLU:OE1	19:AC:84:GLU:N	2.52	0.40
20:AD:228:VAL:N	20:AD:240:SER:O	2.54	0.40
34:AR:161:ILE:O	34:AR:162:SER:C	2.64	0.40
40:AX:69:ASN:OD1	40:AX:70:ILE:N	2.54	0.40
47:XH:70:LYS:HD2	47:XH:70:LYS:C	2.46	0.40
48:XI:137:ASP:OD1	48:XI:137:ASP:N	2.54	0.40
49:XJ:120:ILE:HA	49:XJ:123:ILE:HG22	2.02	0.40
50:XK:176:TYR:OH	82:r:94:ARG:NH2	2.46	0.40
68:c:150:THR:HA	68:c:153:ILE:HG22	2.03	0.40
84:r2:70:P5P:H2'	84:r2:71:P5P:H8	2.03	0.40
3:2:49:ARG:NH2	11:XA:2500:A:N1	2.70	0.40
7:6:144:GLY:N	7:6:145:PRO:CD	2.84	0.40
8:7:225:VAL:O	8:7:229:ILE:HG12	2.21	0.40
11:XA:1687:A:H2'	11:XA:1688:A:O4'	2.21	0.40
11:XA:2065:A:C2	11:XA:2066:C:C4	3.09	0.40
11:XA:2091:A:H2'	11:XA:2092:C:O4'	2.21	0.40
11:XA:2292:G:C6	57:XR:10:LEU:N	2.89	0.40
11:XA:2349:G:N2	11:XA:2365:U:C2	2.89	0.40
11:XA:2453:G:O6	11:XA:2672:A:N6	2.54	0.40
11:XA:2671:C:H2'	11:XA:2672:A:H8	1.86	0.40
11:XA:2992:G:C2	88:A:5:MHU:HM1	2.55	0.40
13:A1:208:LYS:NZ	13:A1:210:ASP:OD2	2.49	0.40
14:A2:39:GLU:OE2	14:A2:39:GLU:N	2.54	0.40
16:A4:62:LYS:CE	24:AH:68:GLU:OE1	2.67	0.40
17:AA:718:A:H2'	17:AA:719:G:O4'	2.21	0.40
17:AA:825:U:C2	17:AA:827:A:OP1	2.74	0.40
17:AA:1456:U:O2'	22:AF:102:GLY:O	2.34	0.40
17:AA:1464:G:O2'	17:AA:1465:C:O4'	2.37	0.40
18:AB:162:CYS:SG	18:AB:254:GLN:HA	2.61	0.40
18:AB:222:ILE:O	18:AB:222:ILE:HG13	2.22	0.40
20:AD:422:TRP:HA	20:AD:425:LEU:HD21	2.03	0.40
31:AO:54:GLU:N	31:AO:55:PRO:CD	2.85	0.40
44:XD:163:ILE:HG22	44:XD:164:LEU:N	2.36	0.40
56:XQ:237:ASN:OD1	56:XQ:238:PHE:N	2.54	0.40
7:6:228:PRO:O	7:6:229:ASP:C	2.65	0.40
8:7:189:LEU:HD23	8:7:189:LEU:H	1.87	0.40
11:XA:2475:U:C6	11:XA:2477:G:OP2	2.75	0.40
11:XA:3151:A:C5	11:XA:3152:C:C5	3.09	0.40

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
13:A1:216:ARG:NH2	41:AY:326:SER:O	2.54	0.40
17:AA:807:A:O2'	17:AA:809:G:O6	2.33	0.40
46:XF:223:HIS:HA	46:XF:226:MET:HE1	2.04	0.40
49:XJ:50:CYS:O	49:XJ:54:ASN:ND2	2.54	0.40
52:XM:102:GLN:HA	52:XM:105:ILE:HG12	2.03	0.40
53:XN:234:ASP:O	53:XN:238:LYS:HA	2.21	0.40
56:XQ:136:ILE:O	56:XQ:151:LEU:HA	2.21	0.40
59:XT:126:ASP:OD1	59:XT:126:ASP:C	2.64	0.40
63:XX:180:ASP:N	63:XX:181:PRO:HD3	2.37	0.40
83:r1:46:Y5P:HB	83:r1:47:Y5P:C5	2.51	0.40
85:r3:54:Y5P:O2'	85:r3:55:P5P:H3'	2.21	0.40
8:7:66:GLU:O	8:7:122:SER:HA	2.21	0.40
11:XA:1804:A:C2	69:d:83:ILE:HG13	2.55	0.40
13:A1:69:VAL:HG22	16:A4:81:ASP:OD1	2.22	0.40
13:A1:165:ASN:HB3	13:A1:167:ARG:CZ	2.52	0.40
17:AA:970:A:H1'	28:AL:146:HIS:HE1	1.86	0.40
20:AD:232:THR:HG22	20:AD:233:ALA:N	2.36	0.40
36:AT:149:CYS:SG	36:AT:150:PRO:HD2	2.61	0.40
38:AV:63:ARG:O	38:AV:64:LYS:HG3	2.22	0.40
45:XE:257:MET:HB2	45:XE:258:PRO:HD2	2.02	0.40
52:XM:133:LYS:C	52:XM:134:ARG:HG2	2.47	0.40
63:XX:213:GLU:OE2	63:XX:216:ARG:NH1	2.53	0.40
69:d:204:ASN:O	69:d:207:ASN:N	2.51	0.40
84:r2:23:P5P:O2'	84:r2:24:P5P:H8	2.21	0.40
84:r2:23:P5P:H8	84:r2:23:P5P:OP1	2.21	0.40
11:XA:1913:G:C2	11:XA:1914:A:C8	3.09	0.40
11:XA:2233:U:C2	45:XE:248:ILE:CD1	3.05	0.40
11:XA:2279:U:OP2	74:i:46:ARG:HD2	2.21	0.40
11:XA:2552:U:C2	11:XA:2553:G:C8	3.09	0.40
11:XA:3055:U:H2'	11:XA:3056:C:O4'	2.21	0.40
11:XA:3056:C:H2'	11:XA:3057:C:C6	2.57	0.40
12:A0:44:PRO:O	12:A0:45:PHE:HB3	2.21	0.40
13:A1:122:HIS:CD2	16:A4:74:LEU:CD1	3.04	0.40
16:A4:158:LYS:HD3	16:A4:162:GLU:HG3	2.04	0.40
16:A4:409:ASP:O	16:A4:412:ASP:OD2	2.40	0.40
16:A4:465:ILE:HG22	16:A4:474:THR:HB	2.04	0.40
18:AB:194:ILE:HA	18:AB:220:VAL:O	2.22	0.40
26:AJ:49:LEU:HD23	26:AJ:51:PRO:HD2	2.03	0.40
34:AR:243:ILE:HG22	34:AR:247:HIS:CD2	2.57	0.40
38:AV:218:SER:HA	38:AV:221:PHE:CE2	2.56	0.40
53:XN:78:GLU:OE2	53:XN:158:ARG:NH2	2.51	0.40

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
56:XQ:70:GLU:N	56:XQ:71:PRO:CD	2.84	0.40
56:XQ:275:THR:HG22	56:XQ:275:THR:O	2.21	0.40
61:XV:60:ASP:OD1	61:XV:61:THR:N	2.54	0.40
70:e:66:LEU:HA	70:e:69:GLU:OE2	2.22	0.40
75:j:102:GLN:HA	75:j:105:GLN:HE21	1.86	0.40

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 PDB entries followed by that with respect to all EM entries.

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	0	106/188 (56%)	102 (96%)	4 (4%)	0	100	100
2	1	51/65 (78%)	47 (92%)	4 (8%)	0	100	100
3	2	44/92 (48%)	43 (98%)	1 (2%)	0	100	100
4	3	93/188 (50%)	91 (98%)	2 (2%)	0	100	100
5	4	36/103 (35%)	35 (97%)	1 (3%)	0	100	100
6	5	391/423 (92%)	365 (93%)	26 (7%)	0	100	100
7	6	350/380 (92%)	325 (93%)	25 (7%)	0	100	100
8	7	283/338 (84%)	266 (94%)	17 (6%)	0	100	100
9	8	137/206 (66%)	129 (94%)	8 (6%)	0	100	100
10	9	122/137 (89%)	116 (95%)	6 (5%)	0	100	100
12	A0	197/218 (90%)	187 (95%)	10 (5%)	0	100	100
13	A1	273/323 (84%)	258 (94%)	15 (6%)	0	100	100
14	A2	114/118 (97%)	111 (97%)	3 (3%)	0	100	100
15	A3	67/199 (34%)	67 (100%)	0	0	100	100
16	A4	526/689 (76%)	493 (94%)	33 (6%)	0	100	100
18	AB	216/296 (73%)	210 (97%)	6 (3%)	0	100	100

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Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
19	AC	130/167 (78%)	128 (98%)	2 (2%)	0	100	100
20	AD	341/430 (79%)	327 (96%)	14 (4%)	0	100	100
21	AE	120/125 (96%)	116 (97%)	4 (3%)	0	100	100
22	AF	197/242 (81%)	193 (98%)	4 (2%)	0	100	100
23	AG	300/396 (76%)	290 (97%)	10 (3%)	0	100	100
24	AH	133/201 (66%)	127 (96%)	6 (4%)	0	100	100
25	AI	134/194 (69%)	129 (96%)	5 (4%)	0	100	100
26	AJ	106/138 (77%)	97 (92%)	9 (8%)	0	100	100
27	AK	99/128 (77%)	97 (98%)	2 (2%)	0	100	100
28	AL	162/257 (63%)	160 (99%)	2 (1%)	0	100	100
29	AM	114/137 (83%)	112 (98%)	2 (2%)	0	100	100
30	AN	105/130 (81%)	101 (96%)	4 (4%)	0	100	100
31	AO	183/258 (71%)	179 (98%)	4 (2%)	0	100	100
32	AP	93/142 (66%)	87 (94%)	6 (6%)	0	100	100
33	AQ	83/87 (95%)	79 (95%)	4 (5%)	0	100	100
34	AR	248/360 (69%)	238 (96%)	10 (4%)	0	100	100
35	AS	131/190 (69%)	123 (94%)	8 (6%)	0	100	100
36	AT	160/173 (92%)	150 (94%)	10 (6%)	0	100	100
37	AU	171/205 (83%)	169 (99%)	2 (1%)	0	100	100
38	AV	341/414 (82%)	322 (94%)	19 (6%)	0	100	100
39	AW	95/187 (51%)	91 (96%)	4 (4%)	0	100	100
40	AX	342/398 (86%)	327 (96%)	15 (4%)	0	100	100
41	AY	111/395 (28%)	102 (92%)	9 (8%)	0	100	100
42	AZ	84/106 (79%)	82 (98%)	2 (2%)	0	100	100
44	XD	234/305 (77%)	214 (92%)	18 (8%)	2 (1%)	14	50
45	XE	302/348 (87%)	287 (95%)	15 (5%)	0	100	100
46	XF	248/311 (80%)	240 (97%)	8 (3%)	0	100	100
47	XH	93/267 (35%)	90 (97%)	3 (3%)	0	100	100
48	XI	209/261 (80%)	192 (92%)	17 (8%)	0	100	100
49	XJ	168/192 (88%)	157 (94%)	11 (6%)	0	100	100
50	XK	175/178 (98%)	169 (97%)	6 (3%)	0	100	100

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Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
51	XL	113/145 (78%)	109 (96%)	4 (4%)	0	100	100
52	XM	285/296 (96%)	272 (95%)	13 (5%)	0	100	100
53	XN	219/251 (87%)	210 (96%)	9 (4%)	0	100	100
54	XO	150/175 (86%)	144 (96%)	6 (4%)	0	100	100
55	XP	141/180 (78%)	131 (93%)	10 (7%)	0	100	100
56	XQ	236/292 (81%)	223 (94%)	13 (6%)	0	100	100
57	XR	138/149 (93%)	132 (96%)	6 (4%)	0	100	100
58	XS	158/205 (77%)	153 (97%)	5 (3%)	0	100	100
59	XT	164/206 (80%)	156 (95%)	8 (5%)	0	100	100
60	XU	137/153 (90%)	132 (96%)	5 (4%)	0	100	100
61	XV	200/216 (93%)	191 (96%)	9 (4%)	0	100	100
62	XW	109/148 (74%)	105 (96%)	4 (4%)	0	100	100
63	XX	241/256 (94%)	231 (96%)	10 (4%)	0	100	100
64	XY	176/250 (70%)	171 (97%)	5 (3%)	0	100	100
65	XZ	118/161 (73%)	116 (98%)	2 (2%)	0	100	100
66	a	93/142 (66%)	86 (92%)	7 (8%)	0	100	100
67	b	146/215 (68%)	137 (94%)	9 (6%)	0	100	100
68	c	271/332 (82%)	259 (96%)	12 (4%)	0	100	100
69	d	212/306 (69%)	200 (94%)	11 (5%)	1 (0%)	24	63
70	e	207/279 (74%)	200 (97%)	7 (3%)	0	100	100
71	f	137/212 (65%)	131 (96%)	6 (4%)	0	100	100
72	g	130/166 (78%)	121 (93%)	8 (6%)	1 (1%)	16	53
73	h	106/158 (67%)	100 (94%)	6 (6%)	0	100	100
74	i	95/128 (74%)	91 (96%)	4 (4%)	0	100	100
75	j	84/123 (68%)	84 (100%)	0	0	100	100
76	k	93/112 (83%)	86 (92%)	7 (8%)	0	100	100
77	l	78/138 (56%)	73 (94%)	5 (6%)	0	100	100
78	m	58/128 (45%)	54 (93%)	4 (7%)	0	100	100
79	o	92/102 (90%)	87 (95%)	5 (5%)	0	100	100
80	p	119/206 (58%)	113 (95%)	6 (5%)	0	100	100
81	q	162/222 (73%)	159 (98%)	3 (2%)	0	100	100

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Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
82	r	144/196 (74%)	136 (94%)	8 (6%)	0	100	100
86	s	366/439 (83%)	349 (95%)	17 (5%)	0	100	100
87	t1	45/198 (23%)	41 (91%)	4 (9%)	0	100	100
87	t2	28/198 (14%)	28 (100%)	0	0	100	100
87	t3	28/198 (14%)	28 (100%)	0	0	100	100
87	t4	27/198 (14%)	26 (96%)	1 (4%)	0	100	100
87	t5	27/198 (14%)	26 (96%)	1 (4%)	0	100	100
87	t6	25/198 (13%)	25 (100%)	0	0	100	100
88	A	2/8 (25%)	0	1 (50%)	1 (50%)	0	0
All	All	13778/19168 (72%)	13136 (95%)	637 (5%)	5 (0%)	100	100

All (5) Ramachandran outliers are listed below:

Mol	Chain	Res	Type
88	A	4	PRO
72	g	38	PHE
44	XD	207	ILE
44	XD	208	ARG
69	d	289	PRO

5.3.2 Protein sidechains ⓘ

In the following table, the Percentiles column shows the percent sidechain outliers of the chain as a percentile score with respect to all PDB entries followed by that with respect to all EM entries.

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	0	97/164 (59%)	97 (100%)	0	100	100
2	1	50/60 (83%)	50 (100%)	0	100	100
3	2	40/72 (56%)	40 (100%)	0	100	100
4	3	88/166 (53%)	88 (100%)	0	100	100
5	4	37/89 (42%)	37 (100%)	0	100	100
6	5	353/368 (96%)	353 (100%)	0	100	100

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Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
7	6	313/332 (94%)	313 (100%)	0	100	100
8	7	267/303 (88%)	267 (100%)	0	100	100
9	8	128/190 (67%)	128 (100%)	0	100	100
10	9	104/112 (93%)	104 (100%)	0	100	100
12	A0	176/190 (93%)	176 (100%)	0	100	100
13	A1	253/291 (87%)	253 (100%)	0	100	100
14	A2	99/101 (98%)	99 (100%)	0	100	100
15	A3	63/166 (38%)	63 (100%)	0	100	100
16	A4	494/609 (81%)	494 (100%)	0	100	100
18	AB	192/249 (77%)	192 (100%)	0	100	100
19	AC	115/143 (80%)	115 (100%)	0	100	100
20	AD	283/357 (79%)	283 (100%)	0	100	100
21	AE	104/107 (97%)	104 (100%)	0	100	100
22	AF	178/209 (85%)	178 (100%)	0	100	100
23	AG	264/342 (77%)	264 (100%)	0	100	100
24	AH	125/180 (69%)	125 (100%)	0	100	100
25	AI	104/147 (71%)	104 (100%)	0	100	100
26	AJ	93/118 (79%)	93 (100%)	0	100	100
27	AK	91/113 (80%)	91 (100%)	0	100	100
28	AL	152/226 (67%)	152 (100%)	0	100	100
29	AM	95/113 (84%)	95 (100%)	0	100	100
30	AN	93/115 (81%)	93 (100%)	0	100	100
31	AO	166/230 (72%)	166 (100%)	0	100	100
32	AP	86/123 (70%)	86 (100%)	0	100	100
33	AQ	77/79 (98%)	77 (100%)	0	100	100
34	AR	229/318 (72%)	229 (100%)	0	100	100
35	AS	115/164 (70%)	115 (100%)	0	100	100
36	AT	150/157 (96%)	150 (100%)	0	100	100
37	AU	149/174 (86%)	149 (100%)	0	100	100
38	AV	315/364 (86%)	315 (100%)	0	100	100
39	AW	84/158 (53%)	84 (100%)	0	100	100

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Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
40	AX	307/351 (88%)	307 (100%)	0	100	100
41	AY	104/357 (29%)	104 (100%)	0	100	100
42	AZ	79/95 (83%)	79 (100%)	0	100	100
44	XD	190/245 (78%)	190 (100%)	0	100	100
45	XE	259/290 (89%)	259 (100%)	0	100	100
46	XF	217/262 (83%)	217 (100%)	0	100	100
47	XH	86/228 (38%)	86 (100%)	0	100	100
48	XI	194/232 (84%)	194 (100%)	0	100	100
49	XJ	133/150 (89%)	133 (100%)	0	100	100
50	XK	155/156 (99%)	155 (100%)	0	100	100
51	XL	98/124 (79%)	98 (100%)	0	100	100
52	XM	245/249 (98%)	245 (100%)	0	100	100
53	XN	188/211 (89%)	188 (100%)	0	100	100
54	XO	133/150 (89%)	133 (100%)	0	100	100
55	XP	125/155 (81%)	125 (100%)	0	100	100
56	XQ	220/256 (86%)	220 (100%)	0	100	100
57	XR	118/126 (94%)	118 (100%)	0	100	100
58	XS	145/180 (81%)	145 (100%)	0	100	100
59	XT	146/176 (83%)	146 (100%)	0	100	100
60	XU	126/135 (93%)	126 (100%)	0	100	100
61	XV	179/191 (94%)	179 (100%)	0	100	100
62	XW	91/119 (76%)	91 (100%)	0	100	100
63	XX	219/229 (96%)	219 (100%)	0	100	100
64	XY	161/223 (72%)	161 (100%)	0	100	100
65	XZ	111/147 (76%)	111 (100%)	0	100	100
66	a	93/133 (70%)	93 (100%)	0	100	100
67	b	130/186 (70%)	130 (100%)	0	100	100
68	c	241/288 (84%)	241 (100%)	0	100	100
69	d	196/274 (72%)	196 (100%)	0	100	100
70	e	185/236 (78%)	185 (100%)	0	100	100
71	f	128/188 (68%)	128 (100%)	0	100	100

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Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
72	g	122/148 (82%)	122 (100%)	0	100	100
73	h	103/148 (70%)	103 (100%)	0	100	100
74	i	86/110 (78%)	86 (100%)	0	100	100
75	j	68/97 (70%)	68 (100%)	0	100	100
76	k	80/90 (89%)	80 (100%)	0	100	100
77	l	74/116 (64%)	74 (100%)	0	100	100
78	m	54/113 (48%)	54 (100%)	0	100	100
79	o	80/87 (92%)	80 (100%)	0	100	100
80	p	117/181 (65%)	117 (100%)	0	100	100
81	q	141/178 (79%)	141 (100%)	0	100	100
82	r	138/169 (82%)	138 (100%)	0	100	100
86	s	326/381 (86%)	326 (100%)	0	100	100
87	t1	41/158 (26%)	40 (98%)	1 (2%)	43	63
87	t2	29/158 (18%)	29 (100%)	0	100	100
87	t3	29/158 (18%)	29 (100%)	0	100	100
87	t4	28/158 (18%)	28 (100%)	0	100	100
87	t5	28/158 (18%)	28 (100%)	0	100	100
87	t6	26/158 (16%)	26 (100%)	0	100	100
88	A	2/2 (100%)	2 (100%)	0	100	100
All	All	12396/16509 (75%)	12395 (100%)	1 (0%)	100	100

All (1) residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
87	t1	21[A]	LEU

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. All (87) such sidechains are listed below:

Mol	Chain	Res	Type
1	0	106	ASN
4	3	154	GLN
5	4	102	GLN
7	6	108	GLN
7	6	307	HIS

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Mol	Chain	Res	Type
7	6	354	GLN
8	7	284	HIS
9	8	126	GLN
9	8	144	GLN
9	8	158	HIS
12	A0	88	GLN
13	A1	275	ASN
15	A3	139	ASN
16	A4	72	GLN
16	A4	129	GLN
16	A4	284	ASN
16	A4	577	ASN
16	A4	590	GLN
18	AB	266	GLN
20	AD	292	HIS
21	AE	59	ASN
22	AF	147	GLN
23	AG	296	ASN
23	AG	340	GLN
24	AH	83	HIS
24	AH	147	HIS
25	AI	92	HIS
25	AI	146	HIS
27	AK	60	ASN
27	AK	117	HIS
29	AM	61	HIS
31	AO	119	ASN
31	AO	204	ASN
34	AR	108	GLN
38	AV	47	HIS
38	AV	329	GLN
38	AV	381	GLN
38	AV	388	GLN
39	AW	86	HIS
40	AX	66	GLN
40	AX	67	HIS
40	AX	190	ASN
40	AX	364	ASN
40	AX	394	HIS
41	AY	285	GLN
41	AY	292	GLN
45	XE	216	GLN

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Mol	Chain	Res	Type
46	XF	103	GLN
46	XF	130	GLN
46	XF	276	GLN
47	XH	88	HIS
48	XI	189	GLN
50	XK	9	GLN
50	XK	64	HIS
50	XK	126	HIS
51	XL	113	ASN
53	XN	101	HIS
53	XN	237	HIS
57	XR	94	GLN
59	XT	75	HIS
61	XV	18	HIS
62	XW	86	ASN
63	XX	27	HIS
63	XX	237	GLN
63	XX	240	GLN
63	XX	241	GLN
68	c	168	HIS
68	c	172	ASN
69	d	157	HIS
69	d	211	GLN
70	e	75	GLN
71	f	158	GLN
72	g	92	HIS
72	g	93	ASN
74	i	122	ASN
76	k	15	GLN
76	k	48	ASN
76	k	80	HIS
79	o	21	HIS
79	o	33	GLN
80	p	53	GLN
82	r	123	HIS
82	r	148	ASN
86	s	191	HIS
86	s	246	GLN
86	s	296	HIS
86	s	298	GLN

5.3.3 RNA ⓘ

Mol	Chain	Analysed	Backbone Outliers	Pucker Outliers
11	XA	1490/1561 (95%)	270 (18%)	7 (0%)
17	AA	916/924 (99%)	167 (18%)	6 (0%)
43	XB	55/72 (76%)	11 (20%)	0
83	r1	13/14 (92%)	7 (53%)	0
84	r2	75/76 (98%)	24 (32%)	0
85	r3	71/73 (97%)	32 (45%)	0
All	All	2620/2720 (96%)	511 (19%)	13 (0%)

All (511) RNA backbone outliers are listed below:

Mol	Chain	Res	Type
11	XA	1681	G
11	XA	1685	C
11	XA	1689	C
11	XA	1692	A
11	XA	1693	C
11	XA	1695	C
11	XA	1699	C
11	XA	1700	U
11	XA	1704	U
11	XA	1707	C
11	XA	1708	A
11	XA	1709	G
11	XA	1710	A
11	XA	1711	C
11	XA	1712	A
11	XA	1715	C
11	XA	1724	A
11	XA	1727	A
11	XA	1733	C
11	XA	1734	C
11	XA	1736	A
11	XA	1737	A
11	XA	1741	A
11	XA	1745	U
11	XA	1748	G
11	XA	1762	A
11	XA	1763	A
11	XA	1764	C
11	XA	1765	C
11	XA	1770	G
11	XA	1804	A
11	XA	1805	A

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Mol	Chain	Res	Type
11	XA	1807	U
11	XA	1809	U
11	XA	1810	A
11	XA	1811	A
11	XA	1821	A
11	XA	1823	A
11	XA	1827	C
11	XA	1828	A
11	XA	1832	A
11	XA	1836	A
11	XA	1844	A
11	XA	1853	A
11	XA	1854	U
11	XA	1856	A
11	XA	1869	A
11	XA	1872	U
11	XA	1878	U
11	XA	1882	A
11	XA	1887	A
11	XA	1893	A
11	XA	1902	C
11	XA	1903	C
11	XA	1907	A
11	XA	1909	A
11	XA	1918	G
11	XA	1919	C
11	XA	1937	A
11	XA	1940	A
11	XA	1944	C
11	XA	1950	U
11	XA	1958	G
11	XA	1974	A
11	XA	1975	U
11	XA	1985	G
11	XA	1986	A
11	XA	1992	C
11	XA	1993	A
11	XA	1994	A
11	XA	2000	C
11	XA	2001	C
11	XA	2002	G
11	XA	2003	A

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Mol	Chain	Res	Type
11	XA	2010	U
11	XA	2015	G
11	XA	2022	G
11	XA	2030	U
11	XA	2036	C
11	XA	2037	U
11	XA	2039	A
11	XA	2055	U
11	XA	2067	C
11	XA	2079	C
11	XA	2099	U
11	XA	2111	C
11	XA	2113	G
11	XA	2125	C
11	XA	2126	U
11	XA	2135	A
11	XA	2138	U
11	XA	2147	G
11	XA	2159	U
11	XA	2168	U
11	XA	2169	A
11	XA	2176	C
11	XA	2177	U
11	XA	2178	A
11	XA	2179	A
11	XA	2180	A
11	XA	2181	A
11	XA	2182	G
11	XA	2188	A
11	XA	2195	A
11	XA	2196	A
11	XA	2198	A
11	XA	2200	A
11	XA	2236	C
11	XA	2237	A
11	XA	2241	A
11	XA	2243	A
11	XA	2244	U
11	XA	2245	A
11	XA	2251	A
11	XA	2260	A
11	XA	2262	C

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Mol	Chain	Res	Type
11	XA	2263	C
11	XA	2283	C
11	XA	2284	C
11	XA	2285	U
11	XA	2297	A
11	XA	2299	U
11	XA	2300	G
11	XA	2316	U
11	XA	2322	C
11	XA	2332	C
11	XA	2345	G
11	XA	2357	C
11	XA	2374	A
11	XA	2375	C
11	XA	2381	A
11	XA	2407	U
11	XA	2414	C
11	XA	2415	C
11	XA	2418	A
11	XA	2432	A
11	XA	2446	A
11	XA	2451	A
11	XA	2458	A
11	XA	2476	C
11	XA	2478	G
11	XA	2485	U
11	XA	2493	C
11	XA	2520	C
11	XA	2523	C
11	XA	2527	A
11	XA	2530	A
11	XA	2540	C
11	XA	2557	C
11	XA	2558	A
11	XA	2559	U
11	XA	2564	A
11	XA	2570	C
11	XA	2575	U
11	XA	2576	A
11	XA	2577	C
11	XA	2578	C
11	XA	2579	C

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Mol	Chain	Res	Type
11	XA	2581	A
11	XA	2592	G
11	XA	2594	U
11	XA	2618	U
11	XA	2626	U
11	XA	2627	G
11	XA	2628	U
11	XA	2630	U
11	XA	2633	A
11	XA	2635	G
11	XA	2654	U
11	XA	2656	U
11	XA	2659	C
11	XA	2683	C
11	XA	2686	G
11	XA	2694	A
11	XA	2695	G
11	XA	2696	A
11	XA	2706	A
11	XA	2715	A
11	XA	2718	C
11	XA	2719	G
11	XA	2722	A
11	XA	2723	A
11	XA	2724	G
11	XA	2725	A
11	XA	2732	G
11	XA	2733	G
11	XA	2740	A
11	XA	2745	A
11	XA	2758	G
11	XA	2788	C
11	XA	2789	C
11	XA	2791	A
11	XA	2810	G
11	XA	2832	A
11	XA	2833	A
11	XA	2847	C
11	XA	2854	U
11	XA	2859	A
11	XA	2864	U
11	XA	2865	C

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Mol	Chain	Res	Type
11	XA	2869	A
11	XA	2871	U
11	XA	2879	A
11	XA	2892	A
11	XA	2906	C
11	XA	2910	A
11	XA	2913	A
11	XA	2916	G
11	XA	2917	G
11	XA	2918	A
11	XA	2921	A
11	XA	2928	C
11	XA	2934	G
11	XA	2935	A
11	XA	2939	C
11	XA	2952	U
11	XA	2956	A
11	XA	2962	C
11	XA	2963	A
11	XA	2978	U
11	XA	2985	C
11	XA	2989	G
11	XA	2990	A
11	XA	2992	G
11	XA	3000	A
11	XA	3005	A
11	XA	3007	C
11	XA	3016	G
11	XA	3021	C
11	XA	3041	U
11	XA	3049	U
11	XA	3053	A
11	XA	3054	G
11	XA	3059	A
11	XA	3061	G
11	XA	3065	U
11	XA	3067	U
11	XA	3069	A
11	XA	3073	C
11	XA	3086	U
11	XA	3089	A
11	XA	3090	G

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Mol	Chain	Res	Type
11	XA	3096	U
11	XA	3100	U
11	XA	3122	U
11	XA	3124	U
11	XA	3129	A
11	XA	3150	U
11	XA	3151	A
11	XA	3154	U
11	XA	3157	C
11	XA	3158	A
11	XA	3160	A
11	XA	3162	C
11	XA	3169	C
11	XA	3172	C
11	XA	3177	A
11	XA	3182	A
11	XA	3184	C
11	XA	3189	C
11	XA	3190	A
11	XA	3194	U
11	XA	3208	C
11	XA	3209	A
11	XA	3210	C
11	XA	3212	C
11	XA	3217	A
11	XA	3218	A
11	XA	3219	G
11	XA	3228	U
17	AA	651	A
17	AA	680	U
17	AA	688	A
17	AA	694	C
17	AA	704	U
17	AA	721	U
17	AA	722	C
17	AA	730	A
17	AA	753	A
17	AA	757	A
17	AA	761	A
17	AA	766	G
17	AA	771	A
17	AA	791	G

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Mol	Chain	Res	Type
17	AA	792	C
17	AA	794	U
17	AA	796	G
17	AA	811	G
17	AA	814	A
17	AA	825	U
17	AA	829	C
17	AA	830	U
17	AA	832	U
17	AA	835	C
17	AA	836	A
17	AA	844	A
17	AA	851	A
17	AA	856	A
17	AA	860	A
17	AA	861	U
17	AA	865	A
17	AA	866	A
17	AA	868	C
17	AA	869	C
17	AA	880	C
17	AA	881	A
17	AA	890	C
17	AA	893	G
17	AA	897	C
17	AA	899	G
17	AA	903	U
17	AA	905	A
17	AA	909	G
17	AA	919	A
17	AA	923	A
17	AA	932	C
17	AA	933	G
17	AA	938	A
17	AA	939	A
17	AA	941	G
17	AA	942	A
17	AA	950	A
17	AA	967	A
17	AA	975	A
17	AA	992	U
17	AA	993	A

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Mol	Chain	Res	Type
17	AA	994	A
17	AA	1001	C
17	AA	1002	C
17	AA	1009	C
17	AA	1015	A
17	AA	1031	G
17	AA	1042	U
17	AA	1049	A
17	AA	1062	G
17	AA	1069	A
17	AA	1081	U
17	AA	1082	A
17	AA	1103	A
17	AA	1105	C
17	AA	1106	C
17	AA	1109	A
17	AA	1121	A
17	AA	1128	C
17	AA	1142	A
17	AA	1143	C
17	AA	1144	U
17	AA	1151	C
17	AA	1167	A
17	AA	1185	C
17	AA	1188	A
17	AA	1189	U
17	AA	1190	C
17	AA	1193	U
17	AA	1194	C
17	AA	1213	A
17	AA	1214	A
17	AA	1215	U
17	AA	1220	A
17	AA	1223	C
17	AA	1225	C
17	AA	1226	C
17	AA	1227	G
17	AA	1228	A
17	AA	1229	U
17	AA	1230	C
17	AA	1235	U
17	AA	1236	C

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Mol	Chain	Res	Type
17	AA	1237	A
17	AA	1248	C
17	AA	1250	C
17	AA	1251	A
17	AA	1261	C
17	AA	1268	C
17	AA	1271	C
17	AA	1284	U
17	AA	1290	C
17	AA	1293	C
17	AA	1294	A
17	AA	1295	A
17	AA	1296	A
17	AA	1297	G
17	AA	1307	G
17	AA	1326	A
17	AA	1327	G
17	AA	1330	C
17	AA	1331	A
17	AA	1341	C
17	AA	1342	C
17	AA	1343	A
17	AA	1344	U
17	AA	1349	U
17	AA	1353	A
17	AA	1354	A
17	AA	1356	A
17	AA	1365	A
17	AA	1369	U
17	AA	1378	C
17	AA	1390	A
17	AA	1391	U
17	AA	1402	A
17	AA	1416	A
17	AA	1419	G
17	AA	1420	U
17	AA	1421	G
17	AA	1422	G
17	AA	1423	A
17	AA	1424	U
17	AA	1430	A
17	AA	1432	U

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Mol	Chain	Res	Type
17	AA	1448	U
17	AA	1459	A
17	AA	1461	A
17	AA	1463	G
17	AA	1478	A
17	AA	1482	A
17	AA	1488	C
17	AA	1489	G
17	AA	1503	G
17	AA	1525	C
17	AA	1526	U
17	AA	1527	A
17	AA	1528	A
17	AA	1531	C
17	AA	1537	C
17	AA	1539	C
17	AA	1551	G
17	AA	1557	A
17	AA	1558	A
17	AA	1559	G
17	AA	1568	U
17	AA	1571	U
17	AA	1582	G
17	AA	1594	G
17	AA	1595	G
17	AA	1598	G
17	AA	1599	A
43	XB	1608	G
43	XB	1609	U
43	XB	1611	G
43	XB	1615	A
43	XB	1619	C
43	XB	1620	A
43	XB	1621	A
43	XB	1646	U
43	XB	1649	C
43	XB	1659	U
43	XB	1663	C
83	r1	47	Y5P
83	r1	49	Y5P
83	r1	53	Y5P
83	r1	54	Y5P

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Mol	Chain	Res	Type
83	r1	55	Y5P
83	r1	56	Y5P
83	r1	57	Y5P
84	r2	8	Y5P
84	r2	9	P5P
84	r2	10	P5P
84	r2	14	P5P
84	r2	15	P5P
84	r2	18	P5P
84	r2	21	P5P
84	r2	23	P5P
84	r2	24	P5P
84	r2	25	Y5P
84	r2	33	Y5P
84	r2	35	P5P
84	r2	45	Y5P
84	r2	46	P5P
84	r2	47	Y5P
84	r2	48	Y5P
84	r2	49	Y5P
84	r2	50	Y5P
84	r2	59	Y5P
84	r2	69	P5P
84	r2	73	P5P
84	r2	74	Y5P
84	r2	75	Y5P
84	r2	76	P5P
85	r3	3	P5P
85	r3	4	P5P
85	r3	13	Y5P
85	r3	14	P5P
85	r3	16	Y5P
85	r3	19	P5P
85	r3	20	Y5P
85	r3	21	P5P
85	r3	30	P5P
85	r3	33	Y5P
85	r3	34	Y5P
85	r3	35	P5P
85	r3	36	Y5P
85	r3	38	Y5P
85	r3	39	Y5P

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Mol	Chain	Res	Type
85	r3	40	Y5P
85	r3	41	Y5P
85	r3	42	P5P
85	r3	43	P5P
85	r3	45	P5P
85	r3	46	P5P
85	r3	47	Y5P
85	r3	48	P5P
85	r3	50	Y5P
85	r3	54	Y5P
85	r3	55	P5P
85	r3	56	Y5P
85	r3	58	Y5P
85	r3	59	Y5P
85	r3	63	Y5P
85	r3	72	Y5P
85	r3	74	P5P

All (13) RNA pucker outliers are listed below:

Mol	Chain	Res	Type
11	XA	2066	C
11	XA	2195	A
11	XA	2417	C
11	XA	2558	A
11	XA	2574	G
11	XA	2961	C
11	XA	2962	C
17	AA	687	G
17	AA	770	C
17	AA	1048	C
17	AA	1234	C
17	AA	1419	G
17	AA	1420	U

5.4 Non-standard residues in protein, DNA, RNA chains ⓘ

168 non-standard protein/DNA/RNA residues 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
84	P5P	r2	24	84	20,23,24	1.29	3 (15%)	27,33,36	2.49	11 (40%)
84	P5P	r2	71	84	20,23,24	1.30	3 (15%)	27,33,36	2.46	10 (37%)
85	P5P	r3	4	85	20,23,24	1.30	3 (15%)	27,33,36	2.49	10 (37%)
85	P5P	r3	22	85	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
84	Y5P	r2	4	84	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
84	Y5P	r2	2	84	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
84	P5P	r2	38	84	20,23,24	1.29	3 (15%)	27,33,36	2.46	11 (40%)
84	P5P	r2	69	84	20,23,24	1.28	3 (15%)	27,33,36	2.48	11 (40%)
84	Y5P	r2	72	84	14,19,20	7.38	3 (21%)	18,26,29	0.56	0
84	Y5P	r2	66	84	14,19,20	7.42	3 (21%)	18,26,29	0.57	0
84	P5P	r2	76	84	20,23,24	1.28	3 (15%)	27,33,36	2.46	11 (40%)
85	P5P	r3	23	85	20,23,24	1.29	3 (15%)	27,33,36	2.47	11 (40%)
84	P5P	r2	6	84	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
84	P5P	r2	35	84	20,23,24	1.28	3 (15%)	27,33,36	2.51	11 (40%)
84	Y5P	r2	61	84	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
84	P5P	r2	5	84	20,23,24	1.29	3 (15%)	27,33,36	2.47	11 (40%)
83	Y5P	r1	51	83	14,19,20	7.35	3 (21%)	18,26,29	0.57	0
84	P5P	r2	19	84	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
85	Y5P	r3	34	85	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
85	Y5P	r3	56	85	14,19,20	7.42	3 (21%)	18,26,29	0.59	0
84	P5P	r2	22	84	20,23,24	1.26	3 (15%)	27,33,36	2.50	11 (40%)
85	Y5P	r3	73	85	14,19,20	7.37	3 (21%)	18,26,29	0.55	0
85	P5P	r3	44	85	20,23,24	1.28	3 (15%)	27,33,36	2.52	11 (40%)
85	P5P	r3	18	85	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
84	Y5P	r2	41	84	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
83	Y5P	r1	57	83	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
85	P5P	r3	30	85	20,23,24	1.28	3 (15%)	27,33,36	2.49	11 (40%)
85	Y5P	r3	72	85	14,19,20	7.41	3 (21%)	18,26,29	0.57	0
85	P5P	r3	29	85	20,23,24	1.28	3 (15%)	27,33,36	2.48	11 (40%)
85	P5P	r3	43	85	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
83	Y5P	r1	50	83	14,19,20	7.36	3 (21%)	18,26,29	0.59	0

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	# Z > 2	Counts	RMSZ	# Z > 2
85	Y5P	r3	70	85	14,19,20	7.42	3 (21%)	18,26,29	0.61	0
84	Y5P	r2	49	84	14,19,20	7.41	3 (21%)	18,26,29	0.61	0
85	P5P	r3	3	85	20,23,24	1.29	3 (15%)	27,33,36	2.47	11 (40%)
83	Y5P	r1	53	83	14,19,20	7.43	3 (21%)	18,26,29	0.57	0
84	Y5P	r2	25	84	14,19,20	7.38	3 (21%)	18,26,29	0.54	0
84	P5P	r2	34	84	20,23,24	1.28	3 (15%)	27,33,36	2.48	10 (37%)
83	Y5P	r1	47	83	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
85	P5P	r3	15	85	20,23,24	1.29	3 (15%)	27,33,36	2.52	11 (40%)
83	Y5P	r1	52	83	14,19,20	7.35	3 (21%)	18,26,29	0.54	0
84	P5P	r2	65	84	20,23,24	1.29	3 (15%)	27,33,36	2.48	10 (37%)
85	Y5P	r3	62	85	14,19,20	7.44	3 (21%)	18,26,29	0.61	0
84	Y5P	r2	68	84	14,19,20	7.36	3 (21%)	18,26,29	0.55	0
83	Y5P	r1	46	83	14,19,20	7.40	3 (21%)	18,26,29	0.57	0
85	P5P	r3	74	85	20,23,24	1.31	3 (15%)	27,33,36	2.43	10 (37%)
85	Y5P	r3	69	85	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
85	Y5P	r3	58	85	14,19,20	7.42	3 (21%)	18,26,29	0.64	0
84	Y5P	r2	51	84	14,19,20	7.37	3 (21%)	18,26,29	0.54	0
85	Y5P	r3	61	85	14,19,20	7.38	3 (21%)	18,26,29	0.54	0
85	P5P	r3	19	85	20,23,24	1.28	3 (15%)	27,33,36	2.48	10 (37%)
84	P5P	r2	9	84	20,23,24	1.28	3 (15%)	27,33,36	2.51	11 (40%)
85	Y5P	r3	60	85	14,19,20	7.42	3 (21%)	18,26,29	0.55	0
88	MHU	A	5	88	14,15,16	0.44	0	18,19,21	1.25	3 (16%)
85	P5P	r3	8	85	20,23,24	1.29	3 (15%)	27,33,36	2.49	11 (40%)
84	Y5P	r2	47	84	14,19,20	7.40	3 (21%)	18,26,29	0.56	0
84	Y5P	r2	16	84	14,19,20	7.37	3 (21%)	18,26,29	0.53	0
85	Y5P	r3	53	85	14,19,20	7.40	3 (21%)	18,26,29	0.56	0
85	Y5P	r3	38	85	14,19,20	7.36	3 (21%)	18,26,29	0.62	0
84	Y5P	r2	17	84	14,19,20	7.40	3 (21%)	18,26,29	0.54	0
85	Y5P	r3	32	85	14,19,20	7.40	3 (21%)	18,26,29	0.59	0
85	Y5P	r3	49	85	14,19,20	7.39	3 (21%)	18,26,29	0.56	0
85	Y5P	r3	2	85	14,19,20	7.42	3 (21%)	18,26,29	0.59	0
83	Y5P	r1	45	83	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
84	Y5P	r2	60	84	14,19,20	7.40	3 (21%)	18,26,29	0.58	0
85	Y5P	r3	25	85	14,19,20	7.37	3 (21%)	18,26,29	0.56	0
84	P5P	r2	10	84	20,23,24	1.27	3 (15%)	27,33,36	2.49	11 (40%)

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	# Z > 2	Counts	RMSZ	# Z > 2
84	P5P	r2	57	84	20,23,24	1.28	3 (15%)	27,33,36	2.46	10 (37%)
85	Y5P	r3	33	85	14,19,20	7.39	3 (21%)	18,26,29	0.54	0
84	P5P	r2	37	84	20,23,24	1.29	3 (15%)	27,33,36	2.46	10 (37%)
84	Y5P	r2	62	84	14,19,20	7.40	3 (21%)	18,26,29	0.58	0
85	P5P	r3	45	85	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
84	P5P	r2	23	84	20,23,24	1.29	3 (15%)	27,33,36	2.43	10 (37%)
84	P5P	r2	63	84	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
85	P5P	r3	48	85	20,23,24	1.27	3 (15%)	27,33,36	2.46	10 (37%)
83	Y5P	r1	48	83	14,19,20	7.38	3 (21%)	18,26,29	0.56	0
85	P5P	r3	31	85	20,23,24	1.28	3 (15%)	27,33,36	2.46	11 (40%)
84	P5P	r2	36	83,84	20,23,24	1.28	3 (15%)	27,33,36	2.50	11 (40%)
84	P5P	r2	52	84	20,23,24	1.30	3 (15%)	27,33,36	2.49	11 (40%)
84	P5P	r2	58	84	20,23,24	1.28	3 (15%)	27,33,36	2.46	10 (37%)
85	P5P	r3	6	85	20,23,24	1.28	3 (15%)	27,33,36	2.48	10 (37%)
85	P5P	r3	42	85	20,23,24	1.30	3 (15%)	27,33,36	2.45	10 (37%)
84	P5P	r2	27	84	20,23,24	1.28	3 (15%)	27,33,36	2.48	11 (40%)
84	P5P	r2	18	84	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
85	P5P	r3	21	85	20,23,24	1.28	3 (15%)	27,33,36	2.50	11 (40%)
84	P5P	r2	46	84	20,23,24	1.27	3 (15%)	27,33,36	2.49	11 (40%)
84	Y5P	r2	3	84	14,19,20	7.39	3 (21%)	18,26,29	0.57	0
85	P5P	r3	37	85	20,23,24	1.27	3 (15%)	27,33,36	2.47	11 (40%)
85	Y5P	r3	65	85	14,19,20	7.43	3 (21%)	18,26,29	0.63	0
84	P5P	r2	28	84	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
84	P5P	r2	1	84	20,23,24	1.27	3 (15%)	27,33,36	2.46	11 (40%)
84	Y5P	r2	12	84	14,19,20	7.38	3 (21%)	18,26,29	0.58	0
85	Y5P	r3	7	85	14,19,20	7.39	3 (21%)	18,26,29	0.56	0
84	P5P	r2	29	84	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
84	P5P	r2	73	84	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
84	Y5P	r2	33	84	14,19,20	7.44	3 (21%)	18,26,29	0.60	0
85	Y5P	r3	50	85	14,19,20	7.40	3 (21%)	18,26,29	0.56	0
84	P5P	r2	15	84	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
83	Y5P	r1	44	83	14,19,20	7.38	3 (21%)	18,26,29	0.56	0
84	P5P	r2	53	84	20,23,24	1.28	3 (15%)	27,33,36	2.47	10 (37%)
85	P5P	r3	9	85	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
85	Y5P	r3	39	85	14,19,20	7.39	3 (21%)	18,26,29	0.55	0

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	# Z > 2	Counts	RMSZ	# Z > 2
84	P5P	r2	21	84	20,23,24	1.29	3 (15%)	27,33,36	2.47	10 (37%)
84	Y5P	r2	59	84	14,19,20	7.36	3 (21%)	18,26,29	0.56	0
84	Y5P	r2	45	84	14,19,20	7.36	3 (21%)	18,26,29	0.54	0
85	P5P	r3	71	85	20,23,24	1.28	3 (15%)	27,33,36	2.48	11 (40%)
85	Y5P	r3	13	85	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
84	Y5P	r2	48	84	14,19,20	7.40	3 (21%)	18,26,29	0.61	0
85	P5P	r3	51	85	20,23,24	1.27	3 (15%)	27,33,36	2.47	11 (40%)
84	Y5P	r2	74	84	14,19,20	7.40	3 (21%)	18,26,29	0.54	0
84	P5P	r2	31	84	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
85	P5P	r3	10	85	20,23,24	1.28	3 (15%)	27,33,36	2.47	10 (37%)
85	P5P	r3	46	85	20,23,24	1.28	3 (15%)	27,33,36	2.46	10 (37%)
85	Y5P	r3	1	85	18,20,20	6.54	3 (16%)	25,29,29	0.67	0
83	Y5P	r1	55	83	14,19,20	7.40	3 (21%)	18,26,29	0.64	0
88	DBB	A	3	88	4,5,6	0.58	0	1,5,7	0.28	0
85	Y5P	r3	40	85	14,19,20	7.41	3 (21%)	18,26,29	0.55	0
85	P5P	r3	11	85	20,23,24	1.29	3 (15%)	27,33,36	2.47	11 (40%)
84	P5P	r2	7	84	20,23,24	1.26	3 (15%)	27,33,36	2.48	11 (40%)
85	Y5P	r3	27	85	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
83	Y5P	r1	56	20,83	14,19,20	7.40	3 (21%)	18,26,29	0.59	0
84	P5P	r2	26	84	20,23,24	1.30	3 (15%)	27,33,36	2.47	10 (37%)
85	P5P	r3	68	85	20,23,24	1.29	3 (15%)	27,33,36	2.47	11 (40%)
84	Y5P	r2	8	84	14,19,20	7.37	3 (21%)	18,26,29	0.55	0
85	Y5P	r3	64	85	14,19,20	7.38	3 (21%)	18,26,29	0.56	0
85	Y5P	r3	63	85	14,19,20	7.39	3 (21%)	18,26,29	0.58	0
85	Y5P	r3	41	85	14,19,20	7.42	3 (21%)	18,26,29	0.62	0
84	Y5P	r2	20	84	14,19,20	7.42	3 (21%)	18,26,29	0.56	0
84	Y5P	r2	67	84	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
84	P5P	r2	44	84	20,23,24	1.29	3 (15%)	27,33,36	2.47	11 (40%)
85	Y5P	r3	16	85	14,19,20	7.42	3 (21%)	18,26,29	0.55	0
85	P5P	r3	52	85	20,23,24	1.28	3 (15%)	27,33,36	2.48	11 (40%)
85	Y5P	r3	24	85	14,19,20	7.43	3 (21%)	18,26,29	0.59	0
85	Y5P	r3	36	85	14,19,20	7.36	3 (21%)	18,26,29	0.54	0
85	Y5P	r3	59	85	14,19,20	7.36	3 (21%)	18,26,29	0.53	0
84	P5P	r2	64	84	20,23,24	1.29	3 (15%)	27,33,36	2.48	11 (40%)
85	Y5P	r3	54	85	14,19,20	7.40	3 (21%)	18,26,29	0.55	0

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	# Z > 2	Counts	RMSZ	# Z > 2
84	Y5P	r2	32	84	14,19,20	7.39	3 (21%)	18,26,29	0.54	0
84	Y5P	r2	43	84	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
84	Y5P	r2	55	84	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
83	Y5P	r1	49	83,84	14,19,20	7.44	3 (21%)	18,26,29	0.59	0
85	Y5P	r3	28	85	14,19,20	7.40	3 (21%)	18,26,29	0.56	0
85	Y5P	r3	12	85	14,19,20	7.39	3 (21%)	18,26,29	0.55	0
84	P5P	r2	14	84	20,23,24	1.28	3 (15%)	27,33,36	2.47	10 (37%)
88	004	A	7	88	9,10,11	1.14	1 (11%)	9,12,14	1.60	2 (22%)
85	P5P	r3	14	85	20,23,24	1.27	3 (15%)	27,33,36	2.47	11 (40%)
88	MHV	A	6	88	7,9,10	0.34	0	8,11,13	1.75	3 (37%)
85	P5P	r3	35	85	20,23,24	1.30	3 (15%)	27,33,36	2.45	10 (37%)
85	Y5P	r3	47	85	14,19,20	7.39	3 (21%)	18,26,29	0.61	0
85	P5P	r3	26	85	20,23,24	1.31	3 (15%)	27,33,36	2.45	10 (37%)
84	P5P	r2	70	84	20,23,24	1.27	3 (15%)	27,33,36	2.49	11 (40%)
85	P5P	r3	55	85	20,23,24	1.27	3 (15%)	27,33,36	2.48	11 (40%)
84	Y5P	r2	13	84	14,19,20	7.42	3 (21%)	18,26,29	0.57	0
85	Y5P	r3	20	85	14,19,20	7.43	3 (21%)	18,26,29	0.56	0
84	P5P	r2	30	84	20,23,24	1.27	3 (15%)	27,33,36	2.47	11 (40%)
84	Y5P	r2	39	84	14,19,20	7.37	3 (21%)	18,26,29	0.55	0
85	P5P	r3	5	85	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
88	MHW	A	1	88	9,9,10	0.79	0	10,11,13	2.49	3 (30%)
85	Y5P	r3	67	85	14,19,20	7.41	3 (21%)	18,26,29	0.56	0
85	P5P	r3	66	85	20,23,24	1.29	3 (15%)	27,33,36	2.48	10 (37%)
85	P5P	r3	57	85	20,23,24	1.28	3 (15%)	27,33,36	2.47	11 (40%)
84	Y5P	r2	50	84	14,19,20	7.39	3 (21%)	18,26,29	0.56	0
84	Y5P	r2	40	84	14,19,20	7.38	3 (21%)	18,26,29	0.55	0
83	Y5P	r1	54	83	14,19,20	7.41	3 (21%)	18,26,29	0.57	0
84	Y5P	r2	42	84	14,19,20	7.37	3 (21%)	18,26,29	0.55	0
84	Y5P	r2	56	84	14,19,20	7.40	3 (21%)	18,26,29	0.55	0
84	Y5P	r2	11	84	14,19,20	7.39	3 (21%)	18,26,29	0.57	0
84	Y5P	r2	75	84	14,19,20	7.44	3 (21%)	18,26,29	0.62	0
84	Y5P	r2	54	84	14,19,20	7.37	3 (21%)	18,26,29	0.53	0

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
84	P5P	r2	24	84	-	2/7/25/26	0/3/3/3
84	P5P	r2	71	84	-	0/7/25/26	0/3/3/3
85	P5P	r3	4	85	-	3/7/25/26	0/3/3/3
85	P5P	r3	22	85	-	0/7/25/26	0/3/3/3
84	Y5P	r2	4	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	2	84	-	1/7/33/34	0/2/2/2
84	P5P	r2	38	84	-	0/7/25/26	0/3/3/3
84	P5P	r2	69	84	-	3/7/25/26	0/3/3/3
84	Y5P	r2	72	84	-	3/7/33/34	0/2/2/2
84	Y5P	r2	66	84	-	3/7/33/34	0/2/2/2
84	P5P	r2	76	84	-	1/7/25/26	0/3/3/3
85	P5P	r3	23	85	-	0/7/25/26	0/3/3/3
84	P5P	r2	6	84	-	0/7/25/26	0/3/3/3
84	P5P	r2	35	84	-	2/7/25/26	0/3/3/3
84	Y5P	r2	61	84	-	2/7/33/34	0/2/2/2
84	P5P	r2	5	84	-	0/7/25/26	0/3/3/3
83	Y5P	r1	51	83	-	2/7/33/34	0/2/2/2
84	P5P	r2	19	84	-	3/7/25/26	0/3/3/3
85	Y5P	r3	34	85	-	5/7/33/34	0/2/2/2
85	Y5P	r3	56	85	-	7/7/33/34	0/2/2/2
84	P5P	r2	22	84	-	1/7/25/26	0/3/3/3
85	Y5P	r3	73	85	-	1/7/33/34	0/2/2/2
85	P5P	r3	44	85	-	3/7/25/26	0/3/3/3
85	P5P	r3	18	85	-	3/7/25/26	0/3/3/3
84	Y5P	r2	41	84	-	1/7/33/34	0/2/2/2
83	Y5P	r1	57	83	-	1/7/33/34	0/2/2/2
85	P5P	r3	30	85	-	2/7/25/26	0/3/3/3
85	Y5P	r3	72	85	-	2/7/33/34	0/2/2/2
85	P5P	r3	29	85	-	3/7/25/26	0/3/3/3
85	P5P	r3	43	85	-	2/7/25/26	0/3/3/3
83	Y5P	r1	50	83	-	2/7/33/34	0/2/2/2
85	Y5P	r3	70	85	-	1/7/33/34	0/2/2/2
84	Y5P	r2	49	84	-	3/7/33/34	0/2/2/2
85	P5P	r3	3	85	-	3/7/25/26	0/3/3/3
83	Y5P	r1	53	83	-	4/7/33/34	0/2/2/2
84	Y5P	r2	25	84	-	1/7/33/34	0/2/2/2

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Mol	Type	Chain	Res	Link	Chirals	Torsions	Rings
84	P5P	r2	34	84	-	0/7/25/26	0/3/3/3
83	Y5P	r1	47	83	-	3/7/33/34	0/2/2/2
85	P5P	r3	15	85	-	0/7/25/26	0/3/3/3
83	Y5P	r1	52	83	-	1/7/33/34	0/2/2/2
84	P5P	r2	65	84	-	0/7/25/26	0/3/3/3
85	Y5P	r3	62	85	-	1/7/33/34	0/2/2/2
84	Y5P	r2	68	84	-	1/7/33/34	0/2/2/2
83	Y5P	r1	46	83	-	4/7/33/34	0/2/2/2
85	P5P	r3	74	85	-	4/7/25/26	0/3/3/3
85	Y5P	r3	69	85	-	3/7/33/34	0/2/2/2
85	Y5P	r3	58	85	-	4/7/33/34	0/2/2/2
84	Y5P	r2	51	84	-	1/7/33/34	0/2/2/2
85	Y5P	r3	61	85	-	2/7/33/34	0/2/2/2
85	P5P	r3	19	85	-	2/7/25/26	0/3/3/3
84	P5P	r2	9	84	-	5/7/25/26	0/3/3/3
85	Y5P	r3	60	85	-	2/7/33/34	0/2/2/2
88	MHU	A	5	88	-	5/9/12/14	0/1/1/1
85	P5P	r3	8	85	-	0/7/25/26	0/3/3/3
84	Y5P	r2	47	84	-	3/7/33/34	0/2/2/2
84	Y5P	r2	16	84	-	1/7/33/34	0/2/2/2
85	Y5P	r3	53	85	-	3/7/33/34	0/2/2/2
85	Y5P	r3	38	85	-	4/7/33/34	0/2/2/2
84	Y5P	r2	17	84	-	3/7/33/34	0/2/2/2
85	Y5P	r3	32	85	-	1/7/33/34	0/2/2/2
85	Y5P	r3	49	85	-	1/7/33/34	0/2/2/2
85	Y5P	r3	2	85	-	3/7/33/34	0/2/2/2
83	Y5P	r1	45	83	-	1/7/33/34	0/2/2/2
84	Y5P	r2	60	84	-	1/7/33/34	0/2/2/2
85	Y5P	r3	25	85	-	1/7/33/34	0/2/2/2
84	P5P	r2	10	84	-	0/7/25/26	0/3/3/3
84	P5P	r2	57	84	-	0/7/25/26	0/3/3/3
85	Y5P	r3	33	85	-	2/7/33/34	0/2/2/2
84	P5P	r2	37	84	-	0/7/25/26	0/3/3/3
84	Y5P	r2	62	84	-	1/7/33/34	0/2/2/2
85	P5P	r3	45	85	-	3/7/25/26	0/3/3/3
84	P5P	r2	23	84	-	1/7/25/26	0/3/3/3
84	P5P	r2	63	84	-	0/7/25/26	0/3/3/3
85	P5P	r3	48	85	-	2/7/25/26	0/3/3/3

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Mol	Type	Chain	Res	Link	Chirals	Torsions	Rings
83	Y5P	r1	48	83	-	1/7/33/34	0/2/2/2
85	P5P	r3	31	85	-	0/7/25/26	0/3/3/3
84	P5P	r2	36	83,84	-	0/7/25/26	0/3/3/3
84	P5P	r2	52	84	-	0/7/25/26	0/3/3/3
84	P5P	r2	58	84	-	0/7/25/26	0/3/3/3
85	P5P	r3	6	85	-	0/7/25/26	0/3/3/3
85	P5P	r3	42	85	-	2/7/25/26	0/3/3/3
84	P5P	r2	27	84	-	0/7/25/26	0/3/3/3
84	P5P	r2	18	84	-	0/7/25/26	0/3/3/3
85	P5P	r3	21	85	-	3/7/25/26	0/3/3/3
84	P5P	r2	46	84	-	5/7/25/26	0/3/3/3
84	Y5P	r2	3	84	-	2/7/33/34	0/2/2/2
85	P5P	r3	37	85	-	0/7/25/26	0/3/3/3
85	Y5P	r3	65	85	-	1/7/33/34	0/2/2/2
84	P5P	r2	28	84	-	0/7/25/26	0/3/3/3
84	P5P	r2	1	84	-	3/7/25/26	0/3/3/3
84	Y5P	r2	12	84	-	1/7/33/34	0/2/2/2
85	Y5P	r3	7	85	-	3/7/33/34	0/2/2/2
84	P5P	r2	29	84	-	0/7/25/26	0/3/3/3
84	P5P	r2	73	84	-	2/7/25/26	0/3/3/3
84	Y5P	r2	33	84	-	6/7/33/34	0/2/2/2
85	Y5P	r3	50	85	-	3/7/33/34	0/2/2/2
84	P5P	r2	15	84	-	4/7/25/26	0/3/3/3
83	Y5P	r1	44	83	-	2/7/33/34	0/2/2/2
84	P5P	r2	53	84	-	0/7/25/26	0/3/3/3
85	P5P	r3	9	85	-	5/7/25/26	0/3/3/3
85	Y5P	r3	39	85	-	3/7/33/34	0/2/2/2
84	P5P	r2	21	84	-	2/7/25/26	0/3/3/3
84	Y5P	r2	59	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	45	84	-	3/7/33/34	0/2/2/2
85	P5P	r3	71	85	-	0/7/25/26	0/3/3/3
85	Y5P	r3	13	85	-	6/7/33/34	0/2/2/2
84	Y5P	r2	48	84	-	2/7/33/34	0/2/2/2
85	P5P	r3	51	85	-	0/7/25/26	0/3/3/3
84	Y5P	r2	74	84	-	3/7/33/34	0/2/2/2
84	P5P	r2	31	84	-	0/7/25/26	0/3/3/3
85	P5P	r3	10	85	-	0/7/25/26	0/3/3/3
85	P5P	r3	46	85	-	2/7/25/26	0/3/3/3

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Mol	Type	Chain	Res	Link	Chirals	Torsions	Rings
85	Y5P	r3	1	85	-	3/10/34/34	0/2/2/2
83	Y5P	r1	55	83	-	3/7/33/34	0/2/2/2
88	DBB	A	3	88	-	0/3/4/6	-
85	Y5P	r3	40	85	-	3/7/33/34	0/2/2/2
85	P5P	r3	11	85	-	0/7/25/26	0/3/3/3
84	P5P	r2	7	84	-	2/7/25/26	0/3/3/3
85	Y5P	r3	27	85	-	1/7/33/34	0/2/2/2
83	Y5P	r1	56	20,83	-	1/7/33/34	0/2/2/2
84	P5P	r2	26	84	-	0/7/25/26	0/3/3/3
85	P5P	r3	68	85	-	0/7/25/26	0/3/3/3
84	Y5P	r2	8	84	-	1/7/33/34	0/2/2/2
85	Y5P	r3	64	85	-	1/7/33/34	0/2/2/2
85	Y5P	r3	63	85	-	3/7/33/34	0/2/2/2
85	Y5P	r3	41	85	-	1/7/33/34	0/2/2/2
84	Y5P	r2	20	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	67	84	-	1/7/33/34	0/2/2/2
84	P5P	r2	44	84	-	0/7/25/26	0/3/3/3
85	Y5P	r3	16	85	-	4/7/33/34	0/2/2/2
85	P5P	r3	52	85	-	0/7/25/26	0/3/3/3
85	Y5P	r3	24	85	-	1/7/33/34	0/2/2/2
85	Y5P	r3	36	85	-	2/7/33/34	0/2/2/2
85	Y5P	r3	59	85	-	2/7/33/34	0/2/2/2
84	P5P	r2	64	84	-	0/7/25/26	0/3/3/3
85	Y5P	r3	54	85	-	3/7/33/34	0/2/2/2
84	Y5P	r2	32	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	43	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	55	84	-	2/7/33/34	0/2/2/2
83	Y5P	r1	49	83,84	-	3/7/33/34	0/2/2/2
85	Y5P	r3	28	85	-	1/7/33/34	0/2/2/2
85	Y5P	r3	12	85	-	1/7/33/34	0/2/2/2
84	P5P	r2	14	84	-	2/7/25/26	0/3/3/3
88	004	A	7	88	-	0/4/6/8	0/1/1/1
85	P5P	r3	14	85	-	3/7/25/26	0/3/3/3
88	MHV	A	6	88	-	0/1/12/14	0/1/1/1
85	P5P	r3	35	85	-	3/7/25/26	0/3/3/3
85	Y5P	r3	47	85	-	4/7/33/34	0/2/2/2
85	P5P	r3	26	85	-	0/7/25/26	0/3/3/3
84	P5P	r2	70	84	-	0/7/25/26	0/3/3/3

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Mol	Type	Chain	Res	Link	Chirals	Torsions	Rings
85	P5P	r3	55	85	-	2/7/25/26	0/3/3/3
84	Y5P	r2	13	84	-	3/7/33/34	0/2/2/2
85	Y5P	r3	20	85	-	1/7/33/34	0/2/2/2
84	P5P	r2	30	84	-	0/7/25/26	0/3/3/3
84	Y5P	r2	39	84	-	1/7/33/34	0/2/2/2
85	P5P	r3	5	85	-	1/7/25/26	0/3/3/3
88	MHW	A	1	88	-	0/2/2/4	0/1/1/1
85	Y5P	r3	67	85	-	1/7/33/34	0/2/2/2
85	P5P	r3	66	85	-	1/7/25/26	0/3/3/3
85	P5P	r3	57	85	-	3/7/25/26	0/3/3/3
84	Y5P	r2	50	84	-	3/7/33/34	0/2/2/2
84	Y5P	r2	40	84	-	1/7/33/34	0/2/2/2
83	Y5P	r1	54	83	-	3/7/33/34	0/2/2/2
84	Y5P	r2	42	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	56	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	11	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	75	84	-	1/7/33/34	0/2/2/2
84	Y5P	r2	54	84	-	1/7/33/34	0/2/2/2

All (490) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
85	r3	62	Y5P	C2-N3	26.95	1.48	1.27
84	r2	75	Y5P	C2-N3	26.94	1.48	1.27
84	r2	33	Y5P	C2-N3	26.94	1.48	1.27
83	r1	49	Y5P	C2-N3	26.94	1.48	1.27
83	r1	53	Y5P	C2-N3	26.89	1.48	1.27
85	r3	65	Y5P	C2-N3	26.88	1.48	1.27
85	r3	24	Y5P	C2-N3	26.88	1.48	1.27
85	r3	20	Y5P	C2-N3	26.88	1.48	1.27
85	r3	41	Y5P	C2-N3	26.88	1.48	1.27
85	r3	56	Y5P	C2-N3	26.87	1.48	1.27
85	r3	2	Y5P	C2-N3	26.87	1.48	1.27
85	r3	16	Y5P	C2-N3	26.86	1.48	1.27
85	r3	58	Y5P	C2-N3	26.86	1.48	1.27
84	r2	20	Y5P	C2-N3	26.86	1.48	1.27
85	r3	70	Y5P	C2-N3	26.86	1.48	1.27
84	r2	13	Y5P	C2-N3	26.85	1.48	1.27
84	r2	66	Y5P	C2-N3	26.85	1.48	1.27
85	r3	60	Y5P	C2-N3	26.84	1.48	1.27
84	r2	49	Y5P	C2-N3	26.82	1.48	1.27

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
85	r3	1	Y5P	C2-N3	26.82	1.48	1.27
83	r1	54	Y5P	C2-N3	26.81	1.48	1.27
85	r3	72	Y5P	C2-N3	26.81	1.48	1.27
83	r1	46	Y5P	C2-N3	26.81	1.48	1.27
84	r2	17	Y5P	C2-N3	26.80	1.48	1.27
83	r1	55	Y5P	C2-N3	26.79	1.48	1.27
84	r2	60	Y5P	C2-N3	26.79	1.48	1.27
85	r3	67	Y5P	C2-N3	26.79	1.48	1.27
85	r3	40	Y5P	C2-N3	26.79	1.48	1.27
84	r2	62	Y5P	C2-N3	26.78	1.48	1.27
84	r2	47	Y5P	C2-N3	26.78	1.48	1.27
84	r2	74	Y5P	C2-N3	26.77	1.48	1.27
85	r3	50	Y5P	C2-N3	26.77	1.48	1.27
83	r1	56	Y5P	C2-N3	26.76	1.48	1.27
84	r2	48	Y5P	C2-N3	26.76	1.48	1.27
84	r2	56	Y5P	C2-N3	26.76	1.48	1.27
85	r3	32	Y5P	C2-N3	26.76	1.48	1.27
84	r2	55	Y5P	C2-N3	26.75	1.48	1.27
85	r3	53	Y5P	C2-N3	26.75	1.48	1.27
85	r3	28	Y5P	C2-N3	26.75	1.48	1.27
85	r3	54	Y5P	C2-N3	26.75	1.48	1.27
83	r1	57	Y5P	C2-N3	26.75	1.48	1.27
85	r3	13	Y5P	C2-N3	26.75	1.48	1.27
85	r3	39	Y5P	C2-N3	26.74	1.48	1.27
84	r2	32	Y5P	C2-N3	26.74	1.48	1.27
84	r2	3	Y5P	C2-N3	26.74	1.48	1.27
85	r3	12	Y5P	C2-N3	26.73	1.48	1.27
85	r3	47	Y5P	C2-N3	26.73	1.48	1.27
85	r3	63	Y5P	C2-N3	26.73	1.48	1.27
85	r3	49	Y5P	C2-N3	26.73	1.48	1.27
85	r3	7	Y5P	C2-N3	26.73	1.48	1.27
85	r3	27	Y5P	C2-N3	26.72	1.48	1.27
83	r1	45	Y5P	C2-N3	26.72	1.48	1.27
84	r2	41	Y5P	C2-N3	26.72	1.48	1.27
84	r2	50	Y5P	C2-N3	26.72	1.48	1.27
84	r2	4	Y5P	C2-N3	26.72	1.48	1.27
85	r3	33	Y5P	C2-N3	26.72	1.48	1.27
84	r2	12	Y5P	C2-N3	26.71	1.48	1.27
84	r2	61	Y5P	C2-N3	26.71	1.48	1.27
84	r2	11	Y5P	C2-N3	26.71	1.48	1.27
84	r2	43	Y5P	C2-N3	26.71	1.48	1.27
83	r1	48	Y5P	C2-N3	26.71	1.48	1.27

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
84	r2	40	Y5P	C2-N3	26.70	1.48	1.27
84	r2	25	Y5P	C2-N3	26.70	1.48	1.27
84	r2	72	Y5P	C2-N3	26.70	1.48	1.27
83	r1	44	Y5P	C2-N3	26.69	1.48	1.27
83	r1	47	Y5P	C2-N3	26.69	1.48	1.27
85	r3	61	Y5P	C2-N3	26.69	1.48	1.27
85	r3	34	Y5P	C2-N3	26.69	1.48	1.27
84	r2	2	Y5P	C2-N3	26.69	1.48	1.27
84	r2	67	Y5P	C2-N3	26.68	1.48	1.27
85	r3	69	Y5P	C2-N3	26.68	1.48	1.27
84	r2	51	Y5P	C2-N3	26.68	1.48	1.27
85	r3	64	Y5P	C2-N3	26.67	1.48	1.27
85	r3	73	Y5P	C2-N3	26.66	1.48	1.27
84	r2	54	Y5P	C2-N3	26.66	1.48	1.27
84	r2	42	Y5P	C2-N3	26.66	1.48	1.27
85	r3	25	Y5P	C2-N3	26.65	1.48	1.27
84	r2	8	Y5P	C2-N3	26.65	1.48	1.27
84	r2	39	Y5P	C2-N3	26.65	1.48	1.27
83	r1	50	Y5P	C2-N3	26.64	1.48	1.27
85	r3	36	Y5P	C2-N3	26.63	1.48	1.27
84	r2	16	Y5P	C2-N3	26.63	1.48	1.27
84	r2	45	Y5P	C2-N3	26.61	1.47	1.27
85	r3	59	Y5P	C2-N3	26.61	1.47	1.27
84	r2	59	Y5P	C2-N3	26.60	1.47	1.27
85	r3	38	Y5P	C2-N3	26.60	1.47	1.27
84	r2	68	Y5P	C2-N3	26.60	1.47	1.27
83	r1	52	Y5P	C2-N3	26.59	1.47	1.27
83	r1	51	Y5P	C2-N3	26.58	1.47	1.27
84	r2	59	Y5P	C6-C5	6.24	1.52	1.33
84	r2	25	Y5P	C6-C5	6.22	1.52	1.33
84	r2	16	Y5P	C6-C5	6.22	1.52	1.33
85	r3	59	Y5P	C6-C5	6.22	1.52	1.33
84	r2	41	Y5P	C6-C5	6.21	1.52	1.33
84	r2	48	Y5P	C6-C5	6.21	1.52	1.33
85	r3	16	Y5P	C6-C5	6.21	1.52	1.33
85	r3	13	Y5P	C6-C5	6.21	1.52	1.33
84	r2	68	Y5P	C6-C5	6.20	1.52	1.33
85	r3	69	Y5P	C6-C5	6.20	1.52	1.33
85	r3	53	Y5P	C6-C5	6.20	1.52	1.33
84	r2	32	Y5P	C6-C5	6.20	1.52	1.33
83	r1	44	Y5P	C6-C5	6.20	1.52	1.33
85	r3	38	Y5P	C6-C5	6.20	1.52	1.33

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
84	r2	8	Y5P	C6-C5	6.20	1.52	1.33
84	r2	45	Y5P	C6-C5	6.20	1.52	1.33
85	r3	61	Y5P	C6-C5	6.20	1.52	1.33
83	r1	52	Y5P	C6-C5	6.20	1.52	1.33
84	r2	2	Y5P	C6-C5	6.20	1.52	1.33
85	r3	64	Y5P	C6-C5	6.20	1.52	1.33
85	r3	27	Y5P	C6-C5	6.19	1.52	1.33
85	r3	12	Y5P	C6-C5	6.19	1.52	1.33
83	r1	53	Y5P	C6-C5	6.19	1.52	1.33
84	r2	12	Y5P	C6-C5	6.19	1.52	1.33
84	r2	11	Y5P	C6-C5	6.19	1.52	1.33
85	r3	7	Y5P	C6-C5	6.19	1.52	1.33
84	r2	40	Y5P	C6-C5	6.19	1.52	1.33
85	r3	20	Y5P	C6-C5	6.19	1.52	1.33
84	r2	43	Y5P	C6-C5	6.19	1.52	1.33
84	r2	17	Y5P	C6-C5	6.19	1.52	1.33
84	r2	54	Y5P	C6-C5	6.18	1.52	1.33
84	r2	20	Y5P	C6-C5	6.18	1.52	1.33
83	r1	49	Y5P	C6-C5	6.18	1.52	1.33
84	r2	51	Y5P	C6-C5	6.18	1.52	1.33
84	r2	4	Y5P	C6-C5	6.18	1.52	1.33
83	r1	57	Y5P	C6-C5	6.18	1.52	1.33
84	r2	62	Y5P	C6-C5	6.18	1.52	1.33
84	r2	72	Y5P	C6-C5	6.18	1.52	1.33
85	r3	54	Y5P	C6-C5	6.18	1.52	1.33
85	r3	33	Y5P	C6-C5	6.18	1.52	1.33
85	r3	47	Y5P	C6-C5	6.18	1.52	1.33
83	r1	48	Y5P	C6-C5	6.18	1.52	1.33
85	r3	34	Y5P	C6-C5	6.18	1.52	1.33
85	r3	40	Y5P	C6-C5	6.18	1.52	1.33
85	r3	50	Y5P	C6-C5	6.18	1.52	1.33
85	r3	28	Y5P	C6-C5	6.18	1.52	1.33
83	r1	47	Y5P	C6-C5	6.18	1.52	1.33
85	r3	67	Y5P	C6-C5	6.18	1.52	1.33
84	r2	74	Y5P	C6-C5	6.18	1.52	1.33
83	r1	56	Y5P	C6-C5	6.18	1.52	1.33
85	r3	60	Y5P	C6-C5	6.17	1.52	1.33
84	r2	3	Y5P	C6-C5	6.17	1.52	1.33
84	r2	42	Y5P	C6-C5	6.17	1.52	1.33
85	r3	63	Y5P	C6-C5	6.17	1.52	1.33
85	r3	36	Y5P	C6-C5	6.17	1.52	1.33
84	r2	39	Y5P	C6-C5	6.17	1.52	1.33

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
85	r3	49	Y5P	C6-C5	6.17	1.52	1.33
83	r1	54	Y5P	C6-C5	6.17	1.52	1.33
84	r2	66	Y5P	C6-C5	6.17	1.52	1.33
84	r2	55	Y5P	C6-C5	6.17	1.52	1.33
84	r2	60	Y5P	C6-C5	6.17	1.52	1.33
83	r1	50	Y5P	C6-C5	6.17	1.52	1.33
84	r2	47	Y5P	C6-C5	6.17	1.52	1.33
84	r2	49	Y5P	C6-C5	6.17	1.52	1.33
84	r2	61	Y5P	C6-C5	6.17	1.52	1.33
85	r3	58	Y5P	C6-C5	6.17	1.52	1.33
84	r2	50	Y5P	C6-C5	6.16	1.52	1.33
84	r2	56	Y5P	C6-C5	6.16	1.52	1.33
84	r2	67	Y5P	C6-C5	6.16	1.52	1.33
84	r2	13	Y5P	C6-C5	6.16	1.52	1.33
85	r3	56	Y5P	C6-C5	6.16	1.52	1.33
85	r3	72	Y5P	C6-C5	6.16	1.52	1.33
84	r2	75	Y5P	C6-C5	6.16	1.52	1.33
85	r3	39	Y5P	C6-C5	6.16	1.52	1.33
83	r1	45	Y5P	C6-C5	6.16	1.52	1.33
85	r3	25	Y5P	C6-C5	6.16	1.52	1.33
83	r1	51	Y5P	C6-C5	6.16	1.52	1.33
85	r3	32	Y5P	C6-C5	6.15	1.52	1.33
85	r3	1	Y5P	C6-C5	6.15	1.52	1.33
85	r3	24	Y5P	C6-C5	6.15	1.52	1.33
85	r3	2	Y5P	C6-C5	6.15	1.52	1.33
85	r3	73	Y5P	C6-C5	6.14	1.52	1.33
85	r3	62	Y5P	C6-C5	6.14	1.52	1.33
85	r3	70	Y5P	C6-C5	6.13	1.52	1.33
85	r3	41	Y5P	C6-C5	6.12	1.51	1.33
83	r1	46	Y5P	C6-C5	6.12	1.51	1.33
84	r2	33	Y5P	C6-C5	6.12	1.51	1.33
85	r3	65	Y5P	C6-C5	6.12	1.51	1.33
83	r1	55	Y5P	C6-C5	6.11	1.51	1.33
85	r3	74	P5P	C5-C4	-3.37	1.34	1.40
85	r3	26	P5P	C5-C4	-3.32	1.35	1.40
85	r3	35	P5P	C5-C4	-3.31	1.35	1.40
85	r3	42	P5P	C5-C4	-3.31	1.35	1.40
85	r3	4	P5P	C5-C4	-3.30	1.35	1.40
84	r2	52	P5P	C5-C4	-3.30	1.35	1.40
84	r2	65	P5P	C5-C4	-3.28	1.35	1.40
84	r2	37	P5P	C5-C4	-3.27	1.35	1.40
85	r3	46	P5P	C5-C4	-3.27	1.35	1.40

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
84	r2	38	P5P	C5-C4	-3.27	1.35	1.40
85	r3	23	P5P	C5-C4	-3.27	1.35	1.40
84	r2	5	P5P	C5-C4	-3.26	1.35	1.40
84	r2	34	P5P	C5-C4	-3.26	1.35	1.40
84	r2	24	P5P	C5-C4	-3.26	1.35	1.40
84	r2	71	P5P	C5-C4	-3.26	1.35	1.40
84	r2	69	P5P	C5-C4	-3.26	1.35	1.40
84	r2	31	P5P	C5-C4	-3.26	1.35	1.40
85	r3	68	P5P	C5-C4	-3.25	1.35	1.40
84	r2	26	P5P	C5-C4	-3.25	1.35	1.40
84	r2	53	P5P	C5-C4	-3.25	1.35	1.40
85	r3	21	P5P	C5-C4	-3.25	1.35	1.40
85	r3	3	P5P	C5-C4	-3.25	1.35	1.40
84	r2	76	P5P	C5-C4	-3.25	1.35	1.40
85	r3	71	P5P	C5-C4	-3.25	1.35	1.40
85	r3	5	P5P	C5-C4	-3.25	1.35	1.40
85	r3	10	P5P	C5-C4	-3.25	1.35	1.40
85	r3	48	P5P	C5-C4	-3.24	1.35	1.40
84	r2	6	P5P	C5-C4	-3.24	1.35	1.40
85	r3	11	P5P	C5-C4	-3.24	1.35	1.40
85	r3	31	P5P	C5-C4	-3.24	1.35	1.40
84	r2	57	P5P	C5-C4	-3.24	1.35	1.40
84	r2	27	P5P	C5-C4	-3.24	1.35	1.40
84	r2	15	P5P	C5-C4	-3.24	1.35	1.40
84	r2	64	P5P	C5-C4	-3.24	1.35	1.40
85	r3	29	P5P	C5-C4	-3.24	1.35	1.40
85	r3	66	P5P	C5-C4	-3.23	1.35	1.40
85	r3	9	P5P	C5-C4	-3.23	1.35	1.40
84	r2	70	P5P	C5-C4	-3.23	1.35	1.40
84	r2	30	P5P	C5-C4	-3.23	1.35	1.40
84	r2	35	P5P	C5-C4	-3.23	1.35	1.40
84	r2	23	P5P	C5-C4	-3.23	1.35	1.40
84	r2	14	P5P	C5-C4	-3.23	1.35	1.40
85	r3	51	P5P	C5-C4	-3.23	1.35	1.40
84	r2	58	P5P	C5-C4	-3.23	1.35	1.40
85	r3	19	P5P	C5-C4	-3.22	1.35	1.40
85	r3	43	P5P	C5-C4	-3.22	1.35	1.40
84	r2	63	P5P	C5-C4	-3.22	1.35	1.40
84	r2	18	P5P	C5-C4	-3.22	1.35	1.40
85	r3	22	P5P	C5-C4	-3.22	1.35	1.40
84	r2	21	P5P	C5-C4	-3.22	1.35	1.40
84	r2	19	P5P	C5-C4	-3.21	1.35	1.40

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
84	r2	44	P5P	C5-C4	-3.21	1.35	1.40
85	r3	15	P5P	C5-C4	-3.21	1.35	1.40
84	r2	36	P5P	C5-C4	-3.21	1.35	1.40
84	r2	46	P5P	C5-C4	-3.21	1.35	1.40
85	r3	6	P5P	C5-C4	-3.20	1.35	1.40
85	r3	8	P5P	C5-C4	-3.20	1.35	1.40
84	r2	29	P5P	C5-C4	-3.20	1.35	1.40
85	r3	52	P5P	C5-C4	-3.20	1.35	1.40
85	r3	18	P5P	C5-C4	-3.20	1.35	1.40
84	r2	7	P5P	C5-C4	-3.19	1.35	1.40
84	r2	9	P5P	C5-C4	-3.19	1.35	1.40
84	r2	28	P5P	C5-C4	-3.19	1.35	1.40
88	A	7	004	CB-CA	-3.19	1.49	1.52
85	r3	57	P5P	C5-C4	-3.19	1.35	1.40
85	r3	30	P5P	C5-C4	-3.19	1.35	1.40
85	r3	45	P5P	C5-C4	-3.18	1.35	1.40
85	r3	37	P5P	C5-C4	-3.18	1.35	1.40
84	r2	1	P5P	C5-C4	-3.18	1.35	1.40
85	r3	55	P5P	C5-C4	-3.17	1.35	1.40
85	r3	44	P5P	C5-C4	-3.16	1.35	1.40
84	r2	22	P5P	C5-C4	-3.16	1.35	1.40
85	r3	14	P5P	C5-C4	-3.16	1.35	1.40
84	r2	10	P5P	C5-C4	-3.15	1.35	1.40
84	r2	73	P5P	C5-C4	-3.14	1.35	1.40
85	r3	24	Y5P	C6-N1	2.80	1.44	1.37
85	r3	64	Y5P	C6-N1	2.79	1.44	1.37
85	r3	2	Y5P	C6-N1	2.79	1.44	1.37
84	r2	11	Y5P	C6-N1	2.78	1.44	1.37
85	r3	72	Y5P	C6-N1	2.78	1.44	1.37
85	r3	33	Y5P	C6-N1	2.78	1.44	1.37
85	r3	13	Y5P	C6-N1	2.78	1.44	1.37
85	r3	27	Y5P	C6-N1	2.78	1.44	1.37
84	r2	62	Y5P	C6-N1	2.77	1.44	1.37
85	r3	56	Y5P	C6-N1	2.77	1.44	1.37
83	r1	45	Y5P	C6-N1	2.77	1.44	1.37
85	r3	41	Y5P	C6-N1	2.77	1.44	1.37
85	r3	12	Y5P	C6-N1	2.77	1.44	1.37
84	r2	2	Y5P	C6-N1	2.77	1.44	1.37
85	r3	16	Y5P	C6-N1	2.77	1.44	1.37
83	r1	49	Y5P	C6-N1	2.77	1.44	1.37
85	r3	54	Y5P	C6-N1	2.77	1.44	1.37
84	r2	4	Y5P	C6-N1	2.77	1.44	1.37

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
84	r2	48	Y5P	C6-N1	2.77	1.44	1.37
83	r1	52	Y5P	C6-N1	2.77	1.44	1.37
83	r1	48	Y5P	C6-N1	2.77	1.44	1.37
85	r3	40	Y5P	C6-N1	2.77	1.44	1.37
85	r3	69	Y5P	C6-N1	2.77	1.44	1.37
84	r2	47	Y5P	C6-N1	2.76	1.44	1.37
85	r3	32	Y5P	C6-N1	2.76	1.44	1.37
85	r3	60	Y5P	C6-N1	2.76	1.44	1.37
83	r1	53	Y5P	C6-N1	2.76	1.44	1.37
84	r2	49	Y5P	C6-N1	2.76	1.44	1.37
85	r3	59	Y5P	C6-N1	2.76	1.44	1.37
84	r2	17	Y5P	C6-N1	2.76	1.43	1.37
84	r2	41	Y5P	C6-N1	2.76	1.43	1.37
84	r2	74	Y5P	C6-N1	2.76	1.43	1.37
85	r3	62	Y5P	C6-N1	2.76	1.43	1.37
85	r3	70	Y5P	C6-N1	2.76	1.43	1.37
84	r2	8	Y5P	C6-N1	2.76	1.43	1.37
85	r3	50	Y5P	C6-N1	2.76	1.43	1.37
84	r2	39	Y5P	C6-N1	2.76	1.43	1.37
83	r1	56	Y5P	C6-N1	2.75	1.43	1.37
84	r2	16	Y5P	C6-N1	2.75	1.43	1.37
85	r3	47	Y5P	C6-N1	2.75	1.43	1.37
83	r1	57	Y5P	C6-N1	2.75	1.43	1.37
84	r2	75	Y5P	C6-N1	2.75	1.43	1.37
84	r2	3	Y5P	C6-N1	2.75	1.43	1.37
84	r2	43	Y5P	C6-N1	2.75	1.43	1.37
84	r2	40	Y5P	C6-N1	2.75	1.43	1.37
84	r2	32	Y5P	C6-N1	2.75	1.43	1.37
84	r2	33	Y5P	C6-N1	2.75	1.43	1.37
85	r3	53	Y5P	C6-N1	2.75	1.43	1.37
85	r3	61	Y5P	C6-N1	2.75	1.43	1.37
85	r3	1	Y5P	C6-N1	2.74	1.43	1.37
84	r2	56	Y5P	C6-N1	2.74	1.43	1.37
84	r2	42	Y5P	C6-N1	2.74	1.43	1.37
85	r3	65	Y5P	C6-N1	2.74	1.43	1.37
84	r2	25	Y5P	C6-N1	2.74	1.43	1.37
84	r2	50	Y5P	C6-N1	2.74	1.43	1.37
83	r1	47	Y5P	C6-N1	2.74	1.43	1.37
85	r3	7	Y5P	C6-N1	2.74	1.43	1.37
85	r3	28	Y5P	C6-N1	2.74	1.43	1.37
85	r3	39	Y5P	C6-N1	2.74	1.43	1.37
85	r3	20	Y5P	C6-N1	2.74	1.43	1.37

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
85	r3	49	Y5P	C6-N1	2.74	1.43	1.37
84	r2	66	Y5P	C6-N1	2.74	1.43	1.37
85	r3	25	Y5P	C6-N1	2.74	1.43	1.37
85	r3	67	Y5P	C6-N1	2.74	1.43	1.37
83	r1	44	Y5P	C6-N1	2.73	1.43	1.37
84	r2	13	Y5P	C6-N1	2.73	1.43	1.37
83	r1	55	Y5P	C6-N1	2.73	1.43	1.37
84	r2	68	Y5P	C6-N1	2.73	1.43	1.37
84	r2	72	Y5P	C6-N1	2.73	1.43	1.37
84	r2	59	Y5P	C6-N1	2.73	1.43	1.37
84	r2	61	Y5P	C6-N1	2.73	1.43	1.37
85	r3	58	Y5P	C6-N1	2.73	1.43	1.37
85	r3	63	Y5P	C6-N1	2.73	1.43	1.37
83	r1	54	Y5P	C6-N1	2.73	1.43	1.37
84	r2	54	Y5P	C6-N1	2.72	1.43	1.37
85	r3	34	Y5P	C6-N1	2.72	1.43	1.37
85	r3	38	Y5P	C6-N1	2.72	1.43	1.37
84	r2	20	Y5P	C6-N1	2.72	1.43	1.37
84	r2	51	Y5P	C6-N1	2.72	1.43	1.37
85	r3	36	Y5P	C6-N1	2.72	1.43	1.37
84	r2	60	Y5P	C6-N1	2.72	1.43	1.37
84	r2	55	Y5P	C6-N1	2.72	1.43	1.37
84	r2	12	Y5P	C6-N1	2.72	1.43	1.37
84	r2	67	Y5P	C6-N1	2.72	1.43	1.37
85	r3	73	Y5P	C6-N1	2.72	1.43	1.37
83	r1	51	Y5P	C6-N1	2.71	1.43	1.37
84	r2	45	Y5P	C6-N1	2.71	1.43	1.37
83	r1	46	Y5P	C6-N1	2.71	1.43	1.37
83	r1	50	Y5P	C6-N1	2.70	1.43	1.37
85	r3	3	P5P	C5-N7	-2.70	1.34	1.39
84	r2	1	P5P	C5-N7	-2.70	1.34	1.39
85	r3	57	P5P	C5-N7	-2.70	1.34	1.39
85	r3	15	P5P	C5-N7	-2.70	1.34	1.39
85	r3	30	P5P	C5-N7	-2.69	1.34	1.39
84	r2	22	P5P	C5-N7	-2.69	1.34	1.39
85	r3	44	P5P	C5-N7	-2.68	1.34	1.39
84	r2	23	P5P	C5-N7	-2.68	1.34	1.39
85	r3	8	P5P	C5-N7	-2.67	1.34	1.39
85	r3	52	P5P	C5-N7	-2.67	1.34	1.39
84	r2	73	P5P	C5-N7	-2.67	1.34	1.39
85	r3	66	P5P	C5-N7	-2.67	1.34	1.39
85	r3	18	P5P	C5-N7	-2.67	1.34	1.39

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
84	r2	44	P5P	C5-N7	-2.67	1.34	1.39
85	r3	45	P5P	C5-N7	-2.67	1.34	1.39
84	r2	21	P5P	C5-N7	-2.66	1.34	1.39
84	r2	10	P5P	C5-N7	-2.66	1.34	1.39
84	r2	69	P5P	C5-N7	-2.66	1.34	1.39
85	r3	29	P5P	C5-N7	-2.66	1.34	1.39
84	r2	28	P5P	C5-N7	-2.66	1.34	1.39
84	r2	24	P5P	C5-N7	-2.66	1.34	1.39
84	r2	70	P5P	C5-N7	-2.66	1.34	1.39
85	r3	19	P5P	C5-N7	-2.66	1.34	1.39
85	r3	14	P5P	C5-N7	-2.66	1.34	1.39
84	r2	38	P5P	C5-N7	-2.65	1.34	1.39
84	r2	58	P5P	C5-N7	-2.65	1.34	1.39
84	r2	64	P5P	C5-N7	-2.65	1.34	1.39
85	r3	42	P5P	C5-N7	-2.65	1.34	1.39
84	r2	9	P5P	C5-N7	-2.65	1.34	1.39
85	r3	74	P5P	C5-N7	-2.65	1.34	1.39
85	r3	23	P5P	C5-N7	-2.65	1.34	1.39
84	r2	26	P5P	C5-N7	-2.65	1.34	1.39
85	r3	43	P5P	C5-N7	-2.65	1.34	1.39
84	r2	18	P5P	C5-N7	-2.65	1.34	1.39
84	r2	63	P5P	C5-N7	-2.65	1.34	1.39
85	r3	4	P5P	C5-N7	-2.65	1.34	1.39
85	r3	11	P5P	C5-N7	-2.65	1.34	1.39
84	r2	52	P5P	C5-N7	-2.64	1.34	1.39
85	r3	31	P5P	C5-N7	-2.64	1.34	1.39
84	r2	30	P5P	C5-N7	-2.64	1.34	1.39
84	r2	5	P5P	C5-N7	-2.64	1.34	1.39
84	r2	76	P5P	C5-N7	-2.64	1.34	1.39
85	r3	26	P5P	C5-N7	-2.64	1.34	1.39
84	r2	7	P5P	C5-N7	-2.64	1.34	1.39
85	r3	68	P5P	C5-N7	-2.64	1.34	1.39
84	r2	19	P5P	C5-N7	-2.64	1.34	1.39
85	r3	55	P5P	C5-N7	-2.64	1.34	1.39
84	r2	36	P5P	C5-N7	-2.63	1.34	1.39
84	r2	27	P5P	C5-N7	-2.63	1.34	1.39
84	r2	71	P5P	C5-N7	-2.63	1.34	1.39
85	r3	35	P5P	C5-N7	-2.63	1.34	1.39
84	r2	34	P5P	C5-N7	-2.63	1.34	1.39
85	r3	9	P5P	C5-N7	-2.63	1.34	1.39
85	r3	37	P5P	C5-N7	-2.63	1.34	1.39
85	r3	21	P5P	C5-N7	-2.63	1.34	1.39

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
85	r3	5	P5P	C5-N7	-2.62	1.34	1.39
85	r3	71	P5P	C5-N7	-2.62	1.34	1.39
84	r2	31	P5P	C5-N7	-2.62	1.34	1.39
84	r2	29	P5P	C5-N7	-2.62	1.34	1.39
84	r2	35	P5P	C5-N7	-2.62	1.34	1.39
84	r2	65	P5P	C5-N7	-2.62	1.34	1.39
84	r2	46	P5P	C5-N7	-2.62	1.34	1.39
84	r2	57	P5P	C5-N7	-2.61	1.34	1.39
85	r3	22	P5P	C5-N7	-2.61	1.34	1.39
84	r2	15	P5P	C5-N7	-2.61	1.34	1.39
85	r3	48	P5P	C5-N7	-2.61	1.34	1.39
85	r3	46	P5P	C5-N7	-2.61	1.34	1.39
84	r2	53	P5P	C5-N7	-2.60	1.34	1.39
85	r3	6	P5P	C5-N7	-2.60	1.34	1.39
85	r3	10	P5P	C5-N7	-2.60	1.34	1.39
84	r2	14	P5P	C5-N7	-2.58	1.35	1.39
84	r2	37	P5P	C5-N7	-2.58	1.35	1.39
84	r2	6	P5P	C5-N7	-2.57	1.35	1.39
85	r3	51	P5P	C5-N7	-2.57	1.35	1.39
85	r3	74	P5P	C4-N9	-2.38	1.32	1.37
84	r2	26	P5P	C4-N9	-2.37	1.32	1.37
84	r2	71	P5P	C4-N9	-2.36	1.32	1.37
85	r3	26	P5P	C4-N9	-2.35	1.32	1.37
84	r2	44	P5P	C4-N9	-2.31	1.32	1.37
85	r3	42	P5P	C4-N9	-2.31	1.32	1.37
85	r3	3	P5P	C4-N9	-2.30	1.32	1.37
84	r2	21	P5P	C4-N9	-2.29	1.32	1.37
84	r2	38	P5P	C4-N9	-2.29	1.32	1.37
84	r2	24	P5P	C4-N9	-2.29	1.32	1.37
84	r2	35	P5P	C4-N9	-2.29	1.32	1.37
84	r2	9	P5P	C4-N9	-2.28	1.32	1.37
84	r2	37	P5P	C4-N9	-2.28	1.32	1.37
85	r3	5	P5P	C4-N9	-2.28	1.32	1.37
84	r2	52	P5P	C4-N9	-2.28	1.32	1.37
84	r2	10	P5P	C4-N9	-2.28	1.32	1.37
85	r3	46	P5P	C4-N9	-2.28	1.33	1.37
85	r3	10	P5P	C4-N9	-2.28	1.33	1.37
84	r2	64	P5P	C4-N9	-2.28	1.33	1.37
84	r2	76	P5P	C4-N9	-2.27	1.33	1.37
85	r3	23	P5P	C4-N9	-2.27	1.33	1.37
84	r2	46	P5P	C4-N9	-2.27	1.33	1.37
84	r2	5	P5P	C4-N9	-2.27	1.33	1.37

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
84	r2	65	P5P	C4-N9	-2.27	1.33	1.37
85	r3	31	P5P	C4-N9	-2.26	1.33	1.37
85	r3	35	P5P	C4-N9	-2.26	1.33	1.37
84	r2	15	P5P	C4-N9	-2.26	1.33	1.37
85	r3	66	P5P	C4-N9	-2.26	1.33	1.37
84	r2	36	P5P	C4-N9	-2.26	1.33	1.37
85	r3	11	P5P	C4-N9	-2.26	1.33	1.37
84	r2	34	P5P	C4-N9	-2.26	1.33	1.37
85	r3	71	P5P	C4-N9	-2.26	1.33	1.37
84	r2	23	P5P	C4-N9	-2.26	1.33	1.37
85	r3	44	P5P	C4-N9	-2.26	1.33	1.37
84	r2	53	P5P	C4-N9	-2.26	1.33	1.37
85	r3	9	P5P	C4-N9	-2.26	1.33	1.37
85	r3	19	P5P	C4-N9	-2.25	1.33	1.37
85	r3	15	P5P	C4-N9	-2.25	1.33	1.37
84	r2	57	P5P	C4-N9	-2.25	1.33	1.37
84	r2	28	P5P	C4-N9	-2.25	1.33	1.37
85	r3	68	P5P	C4-N9	-2.25	1.33	1.37
85	r3	14	P5P	C4-N9	-2.24	1.33	1.37
84	r2	14	P5P	C4-N9	-2.24	1.33	1.37
84	r2	1	P5P	C4-N9	-2.24	1.33	1.37
85	r3	55	P5P	C4-N9	-2.24	1.33	1.37
85	r3	8	P5P	C4-N9	-2.24	1.33	1.37
84	r2	27	P5P	C4-N9	-2.24	1.33	1.37
84	r2	6	P5P	C4-N9	-2.24	1.33	1.37
84	r2	31	P5P	C4-N9	-2.24	1.33	1.37
85	r3	6	P5P	C4-N9	-2.24	1.33	1.37
84	r2	63	P5P	C4-N9	-2.24	1.33	1.37
84	r2	19	P5P	C4-N9	-2.24	1.33	1.37
85	r3	21	P5P	C4-N9	-2.24	1.33	1.37
85	r3	4	P5P	C4-N9	-2.23	1.33	1.37
85	r3	22	P5P	C4-N9	-2.23	1.33	1.37
84	r2	58	P5P	C4-N9	-2.23	1.33	1.37
85	r3	45	P5P	C4-N9	-2.23	1.33	1.37
85	r3	51	P5P	C4-N9	-2.22	1.33	1.37
84	r2	73	P5P	C4-N9	-2.22	1.33	1.37
85	r3	29	P5P	C4-N9	-2.22	1.33	1.37
84	r2	18	P5P	C4-N9	-2.21	1.33	1.37
85	r3	48	P5P	C4-N9	-2.21	1.33	1.37
84	r2	30	P5P	C4-N9	-2.21	1.33	1.37
84	r2	22	P5P	C4-N9	-2.21	1.33	1.37
84	r2	29	P5P	C4-N9	-2.21	1.33	1.37

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
85	r3	37	P5P	C4-N9	-2.21	1.33	1.37
85	r3	52	P5P	C4-N9	-2.20	1.33	1.37
85	r3	57	P5P	C4-N9	-2.20	1.33	1.37
85	r3	30	P5P	C4-N9	-2.20	1.33	1.37
84	r2	7	P5P	C4-N9	-2.20	1.33	1.37
84	r2	69	P5P	C4-N9	-2.20	1.33	1.37
85	r3	18	P5P	C4-N9	-2.19	1.33	1.37
85	r3	43	P5P	C4-N9	-2.18	1.33	1.37
84	r2	70	P5P	C4-N9	-2.16	1.33	1.37

All (803) bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
88	A	1	MHW	O-C-CA	-6.19	116.60	124.37
84	r2	35	P5P	N9-C8-N7	-5.39	106.28	113.94
85	r3	42	P5P	N9-C8-N7	-5.38	106.31	113.94
85	r3	26	P5P	N9-C8-N7	-5.37	106.32	113.94
84	r2	71	P5P	N9-C8-N7	-5.36	106.33	113.94
85	r3	35	P5P	N9-C8-N7	-5.35	106.35	113.94
85	r3	4	P5P	N9-C8-N7	-5.34	106.36	113.94
84	r2	52	P5P	N9-C8-N7	-5.32	106.39	113.94
84	r2	24	P5P	N9-C8-N7	-5.31	106.40	113.94
85	r3	71	P5P	N9-C8-N7	-5.31	106.40	113.94
85	r3	15	P5P	N9-C8-N7	-5.30	106.42	113.94
84	r2	34	P5P	N9-C8-N7	-5.30	106.42	113.94
85	r3	74	P5P	N9-C8-N7	-5.30	106.42	113.94
85	r3	21	P5P	N9-C8-N7	-5.30	106.42	113.94
84	r2	21	P5P	N9-C8-N7	-5.29	106.43	113.94
84	r2	26	P5P	N9-C8-N7	-5.29	106.44	113.94
84	r2	46	P5P	N9-C8-N7	-5.29	106.44	113.94
85	r3	46	P5P	N9-C8-N7	-5.28	106.45	113.94
84	r2	37	P5P	N9-C8-N7	-5.28	106.45	113.94
85	r3	3	P5P	N9-C8-N7	-5.28	106.45	113.94
84	r2	57	P5P	N9-C8-N7	-5.28	106.45	113.94
84	r2	65	P5P	N9-C8-N7	-5.27	106.45	113.94
85	r3	19	P5P	N9-C8-N7	-5.27	106.45	113.94
84	r2	53	P5P	N9-C8-N7	-5.27	106.46	113.94
85	r3	44	P5P	N9-C8-N7	-5.27	106.46	113.94
84	r2	19	P5P	N9-C8-N7	-5.27	106.46	113.94
84	r2	58	P5P	N9-C8-N7	-5.27	106.46	113.94
85	r3	66	P5P	N9-C8-N7	-5.27	106.46	113.94
85	r3	11	P5P	N9-C8-N7	-5.27	106.47	113.94

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	23	P5P	N9-C8-N7	-5.27	106.47	113.94
85	r3	51	P5P	N9-C8-N7	-5.27	106.47	113.94
84	r2	38	P5P	N9-C8-N7	-5.26	106.47	113.94
85	r3	52	P5P	N9-C8-N7	-5.26	106.47	113.94
84	r2	69	P5P	N9-C8-N7	-5.26	106.48	113.94
84	r2	63	P5P	N9-C8-N7	-5.26	106.48	113.94
84	r2	27	P5P	N9-C8-N7	-5.25	106.48	113.94
85	r3	6	P5P	N9-C8-N7	-5.25	106.48	113.94
85	r3	68	P5P	N9-C8-N7	-5.25	106.48	113.94
84	r2	30	P5P	N9-C8-N7	-5.25	106.48	113.94
84	r2	36	P5P	N9-C8-N7	-5.25	106.49	113.94
85	r3	10	P5P	N9-C8-N7	-5.25	106.49	113.94
85	r3	31	P5P	N9-C8-N7	-5.24	106.50	113.94
84	r2	14	P5P	N9-C8-N7	-5.24	106.50	113.94
85	r3	8	P5P	N9-C8-N7	-5.24	106.50	113.94
84	r2	18	P5P	N9-C8-N7	-5.24	106.50	113.94
84	r2	44	P5P	N9-C8-N7	-5.24	106.50	113.94
85	r3	22	P5P	N9-C8-N7	-5.24	106.50	113.94
84	r2	10	P5P	N9-C8-N7	-5.24	106.51	113.94
84	r2	31	P5P	N9-C8-N7	-5.23	106.52	113.94
85	r3	5	P5P	N9-C8-N7	-5.23	106.52	113.94
85	r3	29	P5P	N9-C8-N7	-5.23	106.52	113.94
84	r2	64	P5P	N9-C8-N7	-5.23	106.52	113.94
84	r2	5	P5P	N9-C8-N7	-5.23	106.52	113.94
84	r2	15	P5P	N9-C8-N7	-5.22	106.52	113.94
84	r2	73	P5P	N9-C8-N7	-5.22	106.53	113.94
84	r2	28	P5P	N9-C8-N7	-5.22	106.53	113.94
85	r3	55	P5P	N9-C8-N7	-5.22	106.53	113.94
85	r3	48	P5P	N9-C8-N7	-5.22	106.54	113.94
85	r3	18	P5P	N9-C8-N7	-5.21	106.54	113.94
84	r2	9	P5P	N9-C8-N7	-5.21	106.55	113.94
85	r3	30	P5P	N9-C8-N7	-5.21	106.55	113.94
84	r2	22	P5P	N9-C8-N7	-5.20	106.55	113.94
84	r2	6	P5P	N9-C8-N7	-5.20	106.56	113.94
85	r3	45	P5P	N9-C8-N7	-5.20	106.56	113.94
85	r3	43	P5P	N9-C8-N7	-5.18	106.59	113.94
85	r3	9	P5P	N9-C8-N7	-5.17	106.60	113.94
84	r2	23	P5P	N9-C8-N7	-5.17	106.60	113.94
85	r3	57	P5P	N9-C8-N7	-5.17	106.61	113.94
84	r2	29	P5P	N9-C8-N7	-5.16	106.61	113.94
84	r2	7	P5P	N9-C8-N7	-5.16	106.62	113.94
84	r2	70	P5P	N9-C8-N7	-5.16	106.62	113.94

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	76	P5P	N9-C8-N7	-5.16	106.62	113.94
85	r3	37	P5P	N9-C8-N7	-5.14	106.65	113.94
85	r3	14	P5P	N9-C8-N7	-5.14	106.65	113.94
84	r2	1	P5P	N9-C8-N7	-5.13	106.65	113.94
85	r3	37	P5P	C5-C4-N3	-5.07	120.22	127.18
84	r2	70	P5P	C5-C4-N3	-5.05	120.25	127.18
84	r2	1	P5P	C5-C4-N3	-5.03	120.27	127.18
85	r3	30	P5P	C5-C4-N3	-5.03	120.27	127.18
84	r2	22	P5P	C5-C4-N3	-5.03	120.27	127.18
84	r2	7	P5P	C5-C4-N3	-5.02	120.28	127.18
85	r3	15	P5P	C5-C4-N3	-5.02	120.28	127.18
84	r2	9	P5P	C5-C4-N3	-5.02	120.28	127.18
84	r2	69	P5P	C5-C4-N3	-5.02	120.29	127.18
85	r3	44	P5P	C5-C4-N3	-5.02	120.29	127.18
84	r2	29	P5P	C5-C4-N3	-5.01	120.30	127.18
84	r2	76	P5P	C5-C4-N3	-5.00	120.31	127.18
85	r3	9	P5P	C5-C4-N3	-5.00	120.31	127.18
85	r3	43	P5P	C5-C4-N3	-5.00	120.32	127.18
85	r3	45	P5P	C5-C4-N3	-5.00	120.32	127.18
85	r3	68	P5P	C5-C4-N3	-5.00	120.32	127.18
85	r3	21	P5P	C5-C4-N3	-5.00	120.32	127.18
85	r3	29	P5P	C5-C4-N3	-5.00	120.32	127.18
84	r2	36	P5P	C5-C4-N3	-4.99	120.32	127.18
85	r3	18	P5P	C5-C4-N3	-4.99	120.33	127.18
85	r3	57	P5P	C5-C4-N3	-4.99	120.33	127.18
85	r3	14	P5P	C5-C4-N3	-4.99	120.33	127.18
84	r2	28	P5P	C5-C4-N3	-4.97	120.35	127.18
85	r3	23	P5P	C5-C4-N3	-4.97	120.35	127.18
85	r3	55	P5P	C5-C4-N3	-4.97	120.35	127.18
84	r2	73	P5P	C5-C4-N3	-4.97	120.36	127.18
85	r3	31	P5P	C5-C4-N3	-4.96	120.36	127.18
84	r2	6	P5P	C5-C4-N3	-4.96	120.37	127.18
84	r2	18	P5P	C5-C4-N3	-4.96	120.37	127.18
84	r2	27	P5P	C5-C4-N3	-4.96	120.37	127.18
85	r3	71	P5P	C5-C4-N3	-4.96	120.37	127.18
85	r3	52	P5P	C5-C4-N3	-4.96	120.37	127.18
85	r3	8	P5P	C5-C4-N3	-4.96	120.37	127.18
85	r3	19	P5P	C5-C4-N3	-4.96	120.37	127.18
84	r2	30	P5P	C5-C4-N3	-4.95	120.38	127.18
85	r3	6	P5P	C5-C4-N3	-4.95	120.38	127.18
84	r2	46	P5P	C5-C4-N3	-4.95	120.38	127.18
85	r3	66	P5P	C5-C4-N3	-4.95	120.38	127.18

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	10	P5P	C5-C4-N3	-4.95	120.39	127.18
84	r2	64	P5P	C5-C4-N3	-4.95	120.39	127.18
84	r2	31	P5P	C5-C4-N3	-4.95	120.39	127.18
85	r3	11	P5P	C5-C4-N3	-4.95	120.39	127.18
84	r2	35	P5P	C5-C4-N3	-4.95	120.39	127.18
85	r3	5	P5P	C5-C4-N3	-4.94	120.39	127.18
85	r3	51	P5P	C5-C4-N3	-4.93	120.41	127.18
85	r3	4	P5P	C5-C4-N3	-4.93	120.41	127.18
84	r2	15	P5P	C5-C4-N3	-4.93	120.41	127.18
84	r2	58	P5P	C5-C4-N3	-4.93	120.41	127.18
84	r2	19	P5P	C5-C4-N3	-4.93	120.41	127.18
84	r2	65	P5P	C5-C4-N3	-4.93	120.42	127.18
84	r2	63	P5P	C5-C4-N3	-4.93	120.42	127.18
84	r2	37	P5P	C5-C4-N3	-4.92	120.42	127.18
84	r2	23	P5P	C5-C4-N3	-4.92	120.42	127.18
85	r3	22	P5P	C5-C4-N3	-4.92	120.42	127.18
84	r2	14	P5P	C5-C4-N3	-4.92	120.43	127.18
84	r2	52	P5P	C5-C4-N3	-4.92	120.43	127.18
84	r2	24	P5P	C5-C4-N3	-4.91	120.43	127.18
85	r3	46	P5P	C5-C4-N3	-4.91	120.44	127.18
84	r2	53	P5P	C5-C4-N3	-4.90	120.45	127.18
84	r2	5	P5P	C5-C4-N3	-4.90	120.46	127.18
85	r3	10	P5P	C5-C4-N3	-4.90	120.46	127.18
84	r2	34	P5P	C5-C4-N3	-4.89	120.47	127.18
85	r3	48	P5P	C5-C4-N3	-4.89	120.47	127.18
84	r2	57	P5P	C5-C4-N3	-4.88	120.47	127.18
84	r2	38	P5P	C5-C4-N3	-4.87	120.49	127.18
84	r2	44	P5P	C5-C4-N3	-4.87	120.49	127.18
85	r3	3	P5P	C5-C4-N3	-4.86	120.50	127.18
85	r3	35	P5P	C5-C4-N3	-4.85	120.52	127.18
84	r2	26	P5P	C5-C4-N3	-4.84	120.53	127.18
84	r2	21	P5P	C5-C4-N3	-4.84	120.54	127.18
85	r3	26	P5P	C5-C4-N3	-4.83	120.55	127.18
85	r3	74	P5P	C5-C4-N3	-4.81	120.57	127.18
85	r3	42	P5P	C5-C4-N3	-4.79	120.60	127.18
84	r2	71	P5P	C5-C4-N3	-4.75	120.65	127.18
84	r2	21	P5P	N1-C2-N3	-4.38	119.89	127.33
85	r3	35	P5P	N1-C2-N3	-4.36	119.92	127.33
85	r3	42	P5P	N1-C2-N3	-4.35	119.93	127.33
84	r2	26	P5P	N1-C2-N3	-4.34	119.96	127.33
85	r3	74	P5P	N1-C2-N3	-4.34	119.96	127.33
85	r3	10	P5P	N1-C2-N3	-4.32	119.99	127.33

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	6	P5P	N1-C2-N3	-4.32	120.00	127.33
85	r3	19	P5P	N1-C2-N3	-4.31	120.00	127.33
84	r2	34	P5P	N1-C2-N3	-4.31	120.00	127.33
84	r2	71	P5P	N1-C2-N3	-4.30	120.02	127.33
85	r3	66	P5P	N1-C2-N3	-4.30	120.02	127.33
84	r2	52	P5P	N1-C2-N3	-4.30	120.02	127.33
85	r3	26	P5P	N1-C2-N3	-4.30	120.03	127.33
84	r2	37	P5P	N1-C2-N3	-4.30	120.03	127.33
85	r3	44	P5P	N1-C2-N3	-4.30	120.03	127.33
85	r3	6	P5P	N1-C2-N3	-4.30	120.03	127.33
84	r2	10	P5P	N1-C2-N3	-4.30	120.03	127.33
84	r2	46	P5P	N1-C2-N3	-4.29	120.03	127.33
85	r3	48	P5P	N1-C2-N3	-4.29	120.03	127.33
84	r2	28	P5P	N1-C2-N3	-4.29	120.03	127.33
84	r2	53	P5P	N1-C2-N3	-4.29	120.05	127.33
84	r2	24	P5P	N1-C2-N3	-4.28	120.05	127.33
84	r2	64	P5P	N1-C2-N3	-4.28	120.05	127.33
85	r3	4	P5P	N1-C2-N3	-4.28	120.05	127.33
84	r2	19	P5P	N1-C2-N3	-4.28	120.05	127.33
84	r2	15	P5P	N1-C2-N3	-4.28	120.06	127.33
85	r3	52	P5P	N1-C2-N3	-4.28	120.06	127.33
85	r3	8	P5P	N1-C2-N3	-4.28	120.06	127.33
85	r3	51	P5P	N1-C2-N3	-4.28	120.06	127.33
85	r3	22	P5P	N1-C2-N3	-4.28	120.06	127.33
84	r2	65	P5P	N1-C2-N3	-4.28	120.06	127.33
84	r2	44	P5P	N1-C2-N3	-4.28	120.07	127.33
84	r2	5	P5P	N1-C2-N3	-4.27	120.07	127.33
84	r2	27	P5P	N1-C2-N3	-4.27	120.07	127.33
84	r2	29	P5P	N1-C2-N3	-4.27	120.08	127.33
84	r2	14	P5P	N1-C2-N3	-4.27	120.08	127.33
84	r2	58	P5P	N1-C2-N3	-4.27	120.08	127.33
84	r2	57	P5P	N1-C2-N3	-4.27	120.08	127.33
85	r3	71	P5P	N1-C2-N3	-4.27	120.08	127.33
85	r3	29	P5P	N1-C2-N3	-4.27	120.08	127.33
85	r3	30	P5P	N1-C2-N3	-4.27	120.08	127.33
85	r3	11	P5P	N1-C2-N3	-4.27	120.08	127.33
84	r2	63	P5P	N1-C2-N3	-4.26	120.09	127.33
85	r3	23	P5P	N1-C2-N3	-4.26	120.09	127.33
84	r2	38	P5P	N1-C2-N3	-4.26	120.09	127.33
85	r3	5	P5P	N1-C2-N3	-4.26	120.09	127.33
85	r3	9	P5P	N1-C2-N3	-4.26	120.09	127.33
84	r2	70	P5P	N1-C2-N3	-4.26	120.09	127.33

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	68	P5P	N1-C2-N3	-4.26	120.10	127.33
84	r2	7	P5P	N1-C2-N3	-4.26	120.10	127.33
85	r3	21	P5P	N1-C2-N3	-4.26	120.10	127.33
85	r3	43	P5P	N1-C2-N3	-4.26	120.10	127.33
84	r2	18	P5P	N1-C2-N3	-4.26	120.10	127.33
84	r2	31	P5P	N1-C2-N3	-4.26	120.10	127.33
85	r3	3	P5P	N1-C2-N3	-4.25	120.10	127.33
85	r3	37	P5P	N1-C2-N3	-4.25	120.11	127.33
84	r2	76	P5P	N1-C2-N3	-4.25	120.11	127.33
85	r3	31	P5P	N1-C2-N3	-4.25	120.11	127.33
85	r3	46	P5P	N1-C2-N3	-4.25	120.11	127.33
84	r2	23	P5P	N1-C2-N3	-4.24	120.12	127.33
85	r3	55	P5P	N1-C2-N3	-4.24	120.12	127.33
85	r3	18	P5P	N1-C2-N3	-4.24	120.12	127.33
85	r3	45	P5P	N1-C2-N3	-4.24	120.12	127.33
84	r2	69	P5P	N1-C2-N3	-4.24	120.13	127.33
84	r2	9	P5P	N1-C2-N3	-4.23	120.14	127.33
84	r2	30	P5P	N1-C2-N3	-4.23	120.14	127.33
84	r2	35	P5P	N1-C2-N3	-4.23	120.15	127.33
84	r2	35	P5P	C4-N9-C8	4.23	110.18	105.74
84	r2	36	P5P	N1-C2-N3	-4.21	120.17	127.33
85	r3	14	P5P	N1-C2-N3	-4.21	120.18	127.33
85	r3	57	P5P	N1-C2-N3	-4.21	120.19	127.33
85	r3	42	P5P	C4-N9-C8	4.20	110.15	105.74
84	r2	22	P5P	N1-C2-N3	-4.20	120.20	127.33
84	r2	71	P5P	C4-N9-C8	4.18	110.13	105.74
84	r2	1	P5P	N1-C2-N3	-4.17	120.25	127.33
85	r3	15	P5P	N1-C2-N3	-4.17	120.25	127.33
85	r3	26	P5P	C4-N9-C8	4.16	110.10	105.74
84	r2	34	P5P	C4-N9-C8	4.14	110.09	105.74
84	r2	73	P5P	N1-C2-N3	-4.14	120.29	127.33
84	r2	52	P5P	C4-N9-C8	4.12	110.06	105.74
85	r3	74	P5P	C4-N9-C8	4.11	110.06	105.74
85	r3	4	P5P	C4-N9-C8	4.11	110.05	105.74
84	r2	9	P5P	C1'-N9-C8	-4.11	117.98	127.09
84	r2	14	P5P	C4-N9-C8	4.10	110.05	105.74
84	r2	22	P5P	C1'-N9-C8	-4.10	117.99	127.09
85	r3	71	P5P	C4-N9-C8	4.10	110.04	105.74
84	r2	24	P5P	C4-N9-C8	4.10	110.04	105.74
84	r2	26	P5P	C4-N9-C8	4.10	110.04	105.74
84	r2	21	P5P	C4-N9-C8	4.09	110.03	105.74
84	r2	36	P5P	C4-N9-C8	4.09	110.03	105.74

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	35	P5P	C4-N9-C8	4.09	110.03	105.74
84	r2	46	P5P	C4-N9-C8	4.08	110.02	105.74
85	r3	15	P5P	C4-N9-C8	4.08	110.02	105.74
85	r3	15	P5P	C1'-N9-C8	-4.08	118.04	127.09
85	r3	44	P5P	C4-N9-C8	4.08	110.02	105.74
85	r3	44	P5P	C1'-N9-C8	-4.07	118.05	127.09
84	r2	57	P5P	C4-N9-C8	4.07	110.01	105.74
84	r2	53	P5P	C4-N9-C8	4.07	110.01	105.74
84	r2	37	P5P	C4-N9-C8	4.06	110.00	105.74
85	r3	3	P5P	C4-N9-C8	4.06	110.00	105.74
85	r3	51	P5P	C4-N9-C8	4.06	110.00	105.74
85	r3	6	P5P	C4-N9-C8	4.05	109.99	105.74
85	r3	11	P5P	C4-N9-C8	4.05	109.99	105.74
84	r2	19	P5P	C4-N9-C8	4.05	109.99	105.74
85	r3	46	P5P	C4-N9-C8	4.05	109.99	105.74
84	r2	38	P5P	C4-N9-C8	4.04	109.98	105.74
84	r2	44	P5P	C4-N9-C8	4.04	109.98	105.74
84	r2	36	P5P	C1'-N9-C8	-4.04	118.14	127.09
84	r2	64	P5P	C4-N9-C8	4.03	109.97	105.74
85	r3	8	P5P	C4-N9-C8	4.03	109.97	105.74
85	r3	52	P5P	C4-N9-C8	4.03	109.97	105.74
84	r2	65	P5P	C4-N9-C8	4.03	109.97	105.74
85	r3	21	P5P	C4-N9-C8	4.03	109.97	105.74
85	r3	14	P5P	C1'-N9-C8	-4.03	118.16	127.09
84	r2	73	P5P	C4-N9-C8	4.02	109.96	105.74
84	r2	9	P5P	C4-N9-C8	4.02	109.96	105.74
84	r2	15	P5P	C4-N9-C8	4.01	109.95	105.74
85	r3	22	P5P	C4-N9-C8	4.01	109.95	105.74
85	r3	10	P5P	C4-N9-C8	4.01	109.95	105.74
84	r2	10	P5P	C4-N9-C8	4.01	109.95	105.74
85	r3	66	P5P	C4-N9-C8	4.01	109.95	105.74
84	r2	18	P5P	C4-N9-C8	4.01	109.94	105.74
84	r2	63	P5P	C4-N9-C8	4.01	109.94	105.74
85	r3	19	P5P	C4-N9-C8	4.00	109.94	105.74
84	r2	27	P5P	C4-N9-C8	4.00	109.94	105.74
85	r3	57	P5P	C1'-N9-C8	-4.00	118.22	127.09
84	r2	58	P5P	C4-N9-C8	4.00	109.94	105.74
85	r3	29	P5P	C4-N9-C8	3.99	109.93	105.74
84	r2	69	P5P	C4-N9-C8	3.99	109.93	105.74
85	r3	68	P5P	C4-N9-C8	3.99	109.92	105.74
84	r2	30	P5P	C4-N9-C8	3.99	109.92	105.74
85	r3	23	P5P	C4-N9-C8	3.99	109.92	105.74

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	73	P5P	C1'-N9-C8	-3.98	118.25	127.09
84	r2	5	P5P	C4-N9-C8	3.98	109.92	105.74
85	r3	5	P5P	C4-N9-C8	3.98	109.92	105.74
84	r2	22	P5P	C4-N9-C8	3.98	109.92	105.74
85	r3	18	P5P	C4-N9-C8	3.98	109.91	105.74
85	r3	48	P5P	C4-N9-C8	3.97	109.91	105.74
85	r3	45	P5P	C4-N9-C8	3.97	109.91	105.74
85	r3	18	P5P	C1'-N9-C8	-3.96	118.30	127.09
85	r3	31	P5P	C4-N9-C8	3.96	109.90	105.74
84	r2	28	P5P	C4-N9-C8	3.96	109.90	105.74
85	r3	55	P5P	C4-N9-C8	3.96	109.90	105.74
84	r2	70	P5P	C1'-N9-C8	-3.96	118.31	127.09
84	r2	31	P5P	C4-N9-C8	3.96	109.89	105.74
85	r3	9	P5P	C1'-N9-C8	-3.96	118.31	127.09
84	r2	6	P5P	C4-N9-C8	3.95	109.89	105.74
84	r2	21	P5P	C6-N1-C2	3.95	120.46	115.80
84	r2	64	P5P	C1'-N9-C8	-3.95	118.34	127.09
84	r2	76	P5P	C4-N9-C8	3.94	109.88	105.74
85	r3	14	P5P	C4-N9-C8	3.93	109.86	105.74
85	r3	8	P5P	C1'-N9-C8	-3.93	118.37	127.09
85	r3	9	P5P	C4-N9-C8	3.93	109.86	105.74
84	r2	10	P5P	C1'-N9-C8	-3.93	118.38	127.09
85	r3	30	P5P	C4-N9-C8	3.93	109.86	105.74
84	r2	35	P5P	C1'-N9-C8	-3.93	118.38	127.09
85	r3	30	P5P	C1'-N9-C8	-3.92	118.39	127.09
84	r2	29	P5P	C4-N9-C8	3.92	109.86	105.74
85	r3	57	P5P	C4-N9-C8	3.92	109.85	105.74
84	r2	1	P5P	C1'-N9-C8	-3.92	118.40	127.09
84	r2	18	P5P	C1'-N9-C8	-3.91	118.41	127.09
85	r3	43	P5P	C4-N9-C8	3.91	109.84	105.74
84	r2	52	P5P	C1'-N9-C8	-3.90	118.43	127.09
84	r2	69	P5P	C1'-N9-C8	-3.90	118.44	127.09
84	r2	5	P5P	C1'-N9-C8	-3.89	118.45	127.09
84	r2	70	P5P	C4-N9-C8	3.89	109.83	105.74
84	r2	23	P5P	C4-N9-C8	3.89	109.82	105.74
84	r2	27	P5P	C1'-N9-C8	-3.89	118.47	127.09
84	r2	7	P5P	C1'-N9-C8	-3.89	118.47	127.09
85	r3	21	P5P	C1'-N9-C8	-3.89	118.47	127.09
85	r3	29	P5P	C1'-N9-C8	-3.89	118.47	127.09
85	r3	55	P5P	C1'-N9-C8	-3.88	118.47	127.09
84	r2	7	P5P	C4-N9-C8	3.88	109.82	105.74
85	r3	37	P5P	C4-N9-C8	3.88	109.81	105.74

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	26	P5P	C6-N1-C2	3.88	120.38	115.80
85	r3	3	P5P	C1'-N9-C8	-3.88	118.49	127.09
85	r3	45	P5P	C1'-N9-C8	-3.87	118.50	127.09
84	r2	15	P5P	C1'-N9-C8	-3.87	118.51	127.09
84	r2	44	P5P	C1'-N9-C8	-3.86	118.52	127.09
84	r2	46	P5P	C1'-N9-C8	-3.86	118.53	127.09
85	r3	68	P5P	C1'-N9-C8	-3.86	118.53	127.09
84	r2	31	P5P	C1'-N9-C8	-3.86	118.54	127.09
85	r3	71	P5P	C1'-N9-C8	-3.85	118.55	127.09
84	r2	1	P5P	C4-N9-C8	3.85	109.78	105.74
85	r3	37	P5P	C1'-N9-C8	-3.85	118.56	127.09
84	r2	28	P5P	C1'-N9-C8	-3.84	118.56	127.09
84	r2	10	P5P	C6-N1-C2	3.83	120.33	115.80
85	r3	10	P5P	C6-N1-C2	3.83	120.32	115.80
84	r2	76	P5P	C1'-N9-C8	-3.83	118.60	127.09
85	r3	48	P5P	C6-N1-C2	3.83	120.32	115.80
84	r2	19	P5P	C1'-N9-C8	-3.83	118.60	127.09
85	r3	22	P5P	C1'-N9-C8	-3.83	118.60	127.09
84	r2	24	P5P	C1'-N9-C8	-3.82	118.61	127.09
84	r2	34	P5P	C1'-N9-C8	-3.82	118.61	127.09
84	r2	15	P5P	C6-N1-C2	3.82	120.31	115.80
85	r3	52	P5P	C1'-N9-C8	-3.81	118.63	127.09
85	r3	43	P5P	C1'-N9-C8	-3.81	118.64	127.09
85	r3	6	P5P	C6-N1-C2	3.81	120.29	115.80
84	r2	30	P5P	C1'-N9-C8	-3.80	118.65	127.09
85	r3	11	P5P	C1'-N9-C8	-3.80	118.66	127.09
84	r2	5	P5P	C6-N1-C2	3.80	120.29	115.80
84	r2	52	P5P	C6-N1-C2	3.80	120.28	115.80
84	r2	29	P5P	C1'-N9-C8	-3.80	118.67	127.09
84	r2	71	P5P	C6-N1-C2	3.79	120.28	115.80
84	r2	34	P5P	C6-N1-C2	3.79	120.28	115.80
84	r2	14	P5P	C1'-N9-C8	-3.79	118.68	127.09
84	r2	53	P5P	C6-N1-C2	3.79	120.28	115.80
84	r2	63	P5P	C1'-N9-C8	-3.79	118.68	127.09
84	r2	44	P5P	C6-N1-C2	3.79	120.28	115.80
85	r3	51	P5P	C1'-N9-C8	-3.79	118.68	127.09
84	r2	24	P5P	C6-N1-C2	3.79	120.27	115.80
84	r2	26	P5P	C1'-N9-C8	-3.79	118.69	127.09
84	r2	38	P5P	C1'-N9-C8	-3.79	118.69	127.09
85	r3	23	P5P	C1'-N9-C8	-3.79	118.69	127.09
85	r3	3	P5P	C6-N1-C2	3.79	120.27	115.80
85	r3	5	P5P	C1'-N9-C8	-3.78	118.69	127.09

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	6	P5P	C6-N1-C2	3.78	120.27	115.80
85	r3	51	P5P	C6-N1-C2	3.78	120.27	115.80
84	r2	65	P5P	C6-N1-C2	3.78	120.26	115.80
84	r2	64	P5P	C6-N1-C2	3.77	120.25	115.80
85	r3	42	P5P	C6-N1-C2	3.77	120.25	115.80
84	r2	6	P5P	C1'-N9-C8	-3.77	118.73	127.09
85	r3	52	P5P	C6-N1-C2	3.77	120.25	115.80
85	r3	66	P5P	C6-N1-C2	3.77	120.25	115.80
85	r3	8	P5P	C6-N1-C2	3.76	120.24	115.80
85	r3	22	P5P	C6-N1-C2	3.76	120.24	115.80
84	r2	46	P5P	C6-N1-C2	3.76	120.24	115.80
85	r3	4	P5P	C1'-N9-C8	-3.76	118.75	127.09
84	r2	65	P5P	C1'-N9-C8	-3.76	118.75	127.09
85	r3	19	P5P	C6-N1-C2	3.75	120.23	115.80
85	r3	66	P5P	C1'-N9-C8	-3.75	118.77	127.09
84	r2	19	P5P	C6-N1-C2	3.75	120.23	115.80
84	r2	38	P5P	C6-N1-C2	3.75	120.23	115.80
85	r3	31	P5P	C1'-N9-C8	-3.75	118.77	127.09
85	r3	9	P5P	C6-N1-C2	3.75	120.23	115.80
84	r2	18	P5P	C6-N1-C2	3.74	120.22	115.80
84	r2	28	P5P	C6-N1-C2	3.74	120.22	115.80
85	r3	19	P5P	C1'-N9-C8	-3.74	118.79	127.09
84	r2	57	P5P	C6-N1-C2	3.74	120.22	115.80
85	r3	55	P5P	C6-N1-C2	3.74	120.22	115.80
85	r3	4	P5P	C6-N1-C2	3.74	120.22	115.80
84	r2	58	P5P	C1'-N9-C8	-3.73	118.81	127.09
84	r2	63	P5P	C6-N1-C2	3.73	120.20	115.80
85	r3	6	P5P	C1'-N9-C8	-3.73	118.81	127.09
84	r2	23	P5P	C6-N1-C2	3.73	120.20	115.80
84	r2	70	P5P	C6-N1-C2	3.73	120.20	115.80
84	r2	27	P5P	C6-N1-C2	3.73	120.20	115.80
85	r3	5	P5P	C6-N1-C2	3.73	120.20	115.80
85	r3	10	P5P	C1'-N9-C8	-3.73	118.83	127.09
85	r3	46	P5P	C6-N1-C2	3.73	120.20	115.80
85	r3	21	P5P	C6-N1-C2	3.72	120.20	115.80
85	r3	23	P5P	C6-N1-C2	3.72	120.19	115.80
84	r2	37	P5P	C6-N1-C2	3.72	120.19	115.80
84	r2	58	P5P	C6-N1-C2	3.72	120.19	115.80
85	r3	44	P5P	C6-N1-C2	3.72	120.19	115.80
84	r2	29	P5P	C6-N1-C2	3.72	120.19	115.80
85	r3	48	P5P	C1'-N9-C8	-3.72	118.85	127.09
84	r2	53	P5P	C1'-N9-C8	-3.72	118.85	127.09

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	31	P5P	C6-N1-C2	3.71	120.19	115.80
85	r3	45	P5P	C6-N1-C2	3.71	120.18	115.80
85	r3	30	P5P	C6-N1-C2	3.71	120.18	115.80
84	r2	7	P5P	C6-N1-C2	3.71	120.18	115.80
85	r3	43	P5P	C6-N1-C2	3.71	120.17	115.80
84	r2	30	P5P	C6-N1-C2	3.70	120.17	115.80
85	r3	18	P5P	C6-N1-C2	3.70	120.17	115.80
84	r2	9	P5P	C6-N1-C2	3.70	120.17	115.80
84	r2	36	P5P	C6-N1-C2	3.70	120.17	115.80
85	r3	29	P5P	C6-N1-C2	3.70	120.17	115.80
84	r2	14	P5P	C6-N1-C2	3.70	120.17	115.80
84	r2	35	P5P	C6-N1-C2	3.69	120.16	115.80
84	r2	21	P5P	C1'-N9-C8	-3.69	118.90	127.09
84	r2	71	P5P	C1'-N9-C8	-3.69	118.91	127.09
85	r3	31	P5P	C6-N1-C2	3.69	120.15	115.80
85	r3	26	P5P	C1'-N9-C8	-3.68	118.92	127.09
85	r3	11	P5P	C6-N1-C2	3.68	120.15	115.80
84	r2	37	P5P	C1'-N9-C8	-3.68	118.92	127.09
85	r3	15	P5P	C6-N1-C2	3.68	120.15	115.80
85	r3	26	P5P	C6-N1-C2	3.68	120.14	115.80
85	r3	35	P5P	C6-N1-C2	3.68	120.14	115.80
85	r3	37	P5P	C6-N1-C2	3.68	120.14	115.80
84	r2	57	P5P	C1'-N9-C8	-3.68	118.94	127.09
85	r3	14	P5P	C6-N1-C2	3.67	120.14	115.80
85	r3	57	P5P	C6-N1-C2	3.67	120.13	115.80
85	r3	71	P5P	C6-N1-C2	3.67	120.13	115.80
85	r3	46	P5P	C1'-N9-C8	-3.67	118.96	127.09
84	r2	1	P5P	C6-N1-C2	3.66	120.12	115.80
84	r2	22	P5P	C6-N1-C2	3.66	120.12	115.80
84	r2	73	P5P	C6-N1-C2	3.65	120.11	115.80
84	r2	69	P5P	C6-N1-C2	3.64	120.10	115.80
85	r3	74	P5P	C6-N1-C2	3.63	120.09	115.80
85	r3	68	P5P	C6-N1-C2	3.63	120.09	115.80
85	r3	74	P5P	C1'-N9-C8	-3.63	119.04	127.09
84	r2	76	P5P	C6-N1-C2	3.61	120.06	115.80
84	r2	23	P5P	C1'-N9-C8	-3.52	119.29	127.09
85	r3	35	P5P	C1'-N9-C8	-3.51	119.31	127.09
84	r2	35	P5P	C5-N7-C8	3.49	108.68	104.40
85	r3	42	P5P	C1'-N9-C8	-3.46	119.41	127.09
85	r3	44	P5P	N3-C4-N9	3.44	133.01	127.17
84	r2	22	P5P	N3-C4-N9	3.43	133.00	127.17
84	r2	9	P5P	N3-C4-N9	3.41	132.97	127.17

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
84	r2	36	P5P	N3-C4-N9	3.41	132.97	127.17
85	r3	15	P5P	N3-C4-N9	3.41	132.96	127.17
85	r3	21	P5P	C5-N7-C8	3.41	108.58	104.40
84	r2	70	P5P	N3-C4-N9	3.41	132.96	127.17
85	r3	15	P5P	C5-N7-C8	3.40	108.58	104.40
85	r3	37	P5P	N3-C4-N9	3.39	132.93	127.17
84	r2	73	P5P	N3-C4-N9	3.39	132.93	127.17
84	r2	69	P5P	C5-N7-C8	3.38	108.55	104.40
85	r3	14	P5P	N3-C4-N9	3.38	132.91	127.17
85	r3	30	P5P	N3-C4-N9	3.38	132.91	127.17
85	r3	4	P5P	C5-N7-C8	3.38	108.54	104.40
84	r2	35	P5P	N3-C4-N9	3.38	132.91	127.17
85	r3	18	P5P	N3-C4-N9	3.38	132.91	127.17
85	r3	57	P5P	N3-C4-N9	3.38	132.91	127.17
85	r3	45	P5P	N3-C4-N9	3.37	132.90	127.17
84	r2	69	P5P	N3-C4-N9	3.37	132.90	127.17
84	r2	76	P5P	N3-C4-N9	3.37	132.90	127.17
85	r3	19	P5P	C5-N7-C8	3.36	108.53	104.40
85	r3	29	P5P	N3-C4-N9	3.36	132.88	127.17
84	r2	7	P5P	N3-C4-N9	3.36	132.88	127.17
85	r3	8	P5P	N3-C4-N9	3.36	132.88	127.17
84	r2	65	P5P	C5-N7-C8	3.36	108.52	104.40
84	r2	71	P5P	C5-N7-C8	3.35	108.52	104.40
84	r2	30	P5P	C5-N7-C8	3.35	108.51	104.40
85	r3	43	P5P	N3-C4-N9	3.35	132.87	127.17
85	r3	71	P5P	N3-C4-N9	3.35	132.86	127.17
84	r2	58	P5P	C5-N7-C8	3.35	108.51	104.40
85	r3	9	P5P	N3-C4-N9	3.35	132.86	127.17
85	r3	52	P5P	N3-C4-N9	3.35	132.86	127.17
84	r2	1	P5P	N3-C4-N9	3.34	132.85	127.17
84	r2	18	P5P	N3-C4-N9	3.34	132.85	127.17
85	r3	44	P5P	C5-N7-C8	3.34	108.50	104.40
84	r2	29	P5P	N3-C4-N9	3.34	132.85	127.17
84	r2	46	P5P	N3-C4-N9	3.34	132.84	127.17
84	r2	14	P5P	N3-C4-N9	3.34	132.84	127.17
84	r2	46	P5P	C5-N7-C8	3.34	108.50	104.40
84	r2	24	P5P	C5-N7-C8	3.34	108.50	104.40
85	r3	21	P5P	N3-C4-N9	3.34	132.84	127.17
84	r2	64	P5P	N3-C4-N9	3.34	132.84	127.17
85	r3	35	P5P	C5-N7-C8	3.34	108.50	104.40
85	r3	66	P5P	C5-N7-C8	3.34	108.50	104.40
84	r2	27	P5P	C5-N7-C8	3.34	108.49	104.40

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	71	P5P	C5-N7-C8	3.34	108.49	104.40
85	r3	52	P5P	C5-N7-C8	3.33	108.49	104.40
84	r2	34	P5P	N3-C4-N9	3.33	132.84	127.17
85	r3	55	P5P	N3-C4-N9	3.33	132.84	127.17
85	r3	31	P5P	C5-N7-C8	3.33	108.49	104.40
85	r3	68	P5P	N3-C4-N9	3.33	132.84	127.17
85	r3	23	P5P	C5-N7-C8	3.33	108.49	104.40
85	r3	46	P5P	C5-N7-C8	3.33	108.49	104.40
84	r2	10	P5P	N3-C4-N9	3.33	132.83	127.17
84	r2	19	P5P	C5-N7-C8	3.33	108.48	104.40
84	r2	52	P5P	N3-C4-N9	3.32	132.82	127.17
84	r2	10	P5P	C5-N7-C8	3.32	108.48	104.40
84	r2	53	P5P	C5-N7-C8	3.32	108.48	104.40
85	r3	6	P5P	N3-C4-N9	3.32	132.81	127.17
88	A	7	004	CG2-CB-CA	-3.32	115.48	120.64
84	r2	57	P5P	C5-N7-C8	3.32	108.47	104.40
84	r2	28	P5P	C5-N7-C8	3.32	108.47	104.40
84	r2	19	P5P	N3-C4-N9	3.32	132.81	127.17
85	r3	11	P5P	N3-C4-N9	3.32	132.81	127.17
85	r3	30	P5P	C5-N7-C8	3.32	108.47	104.40
84	r2	63	P5P	C5-N7-C8	3.32	108.47	104.40
84	r2	52	P5P	C5-N7-C8	3.32	108.47	104.40
85	r3	3	P5P	C5-N7-C8	3.32	108.47	104.40
84	r2	27	P5P	N3-C4-N9	3.31	132.80	127.17
84	r2	28	P5P	N3-C4-N9	3.31	132.80	127.17
85	r3	4	P5P	N3-C4-N9	3.31	132.80	127.17
84	r2	73	P5P	C5-N7-C8	3.31	108.47	104.40
85	r3	55	P5P	C5-N7-C8	3.31	108.47	104.40
84	r2	30	P5P	N3-C4-N9	3.31	132.80	127.17
85	r3	51	P5P	N3-C4-N9	3.31	132.80	127.17
85	r3	66	P5P	N3-C4-N9	3.31	132.80	127.17
85	r3	29	P5P	C5-N7-C8	3.31	108.46	104.40
84	r2	70	P5P	C5-N7-C8	3.31	108.46	104.40
84	r2	24	P5P	N3-C4-N9	3.31	132.79	127.17
85	r3	23	P5P	N3-C4-N9	3.31	132.79	127.17
84	r2	63	P5P	N3-C4-N9	3.31	132.79	127.17
85	r3	22	P5P	N3-C4-N9	3.31	132.79	127.17
84	r2	21	P5P	C5-N7-C8	3.31	108.46	104.40
84	r2	18	P5P	C5-N7-C8	3.30	108.46	104.40
85	r3	5	P5P	C5-N7-C8	3.30	108.46	104.40
85	r3	68	P5P	C5-N7-C8	3.30	108.46	104.40
84	r2	37	P5P	C5-N7-C8	3.30	108.45	104.40

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	6	P5P	C5-N7-C8	3.30	108.45	104.40
84	r2	34	P5P	C5-N7-C8	3.30	108.45	104.40
84	r2	15	P5P	N3-C4-N9	3.30	132.78	127.17
85	r3	10	P5P	C5-N7-C8	3.30	108.45	104.40
85	r3	26	P5P	C5-N7-C8	3.30	108.45	104.40
85	r3	42	P5P	C5-N7-C8	3.30	108.45	104.40
85	r3	48	P5P	C5-N7-C8	3.30	108.45	104.40
84	r2	29	P5P	C5-N7-C8	3.30	108.45	104.40
84	r2	31	P5P	C5-N7-C8	3.30	108.45	104.40
84	r2	22	P5P	C5-N7-C8	3.30	108.45	104.40
85	r3	19	P5P	N3-C4-N9	3.30	132.77	127.17
84	r2	36	P5P	C5-N7-C8	3.29	108.44	104.40
85	r3	8	P5P	C5-N7-C8	3.29	108.44	104.40
85	r3	11	P5P	C5-N7-C8	3.29	108.44	104.40
85	r3	51	P5P	C5-N7-C8	3.29	108.44	104.40
84	r2	31	P5P	N3-C4-N9	3.29	132.76	127.17
85	r3	5	P5P	N3-C4-N9	3.29	132.76	127.17
85	r3	3	P5P	N3-C4-N9	3.29	132.76	127.17
85	r3	45	P5P	C5-N7-C8	3.29	108.44	104.40
85	r3	18	P5P	C5-N7-C8	3.29	108.43	104.40
85	r3	31	P5P	N3-C4-N9	3.29	132.76	127.17
84	r2	26	P5P	C5-N7-C8	3.28	108.43	104.40
84	r2	37	P5P	N3-C4-N9	3.28	132.75	127.17
85	r3	37	P5P	C2-N3-C4	3.28	119.85	111.83
84	r2	6	P5P	N3-C4-N9	3.28	132.75	127.17
84	r2	53	P5P	N3-C4-N9	3.28	132.75	127.17
84	r2	58	P5P	N3-C4-N9	3.28	132.75	127.17
85	r3	35	P5P	C2-N3-C4	3.28	119.84	111.83
84	r2	23	P5P	C5-N7-C8	3.28	108.42	104.40
85	r3	48	P5P	N3-C4-N9	3.28	132.74	127.17
84	r2	5	P5P	C5-N7-C8	3.27	108.42	104.40
84	r2	57	P5P	N3-C4-N9	3.27	132.74	127.17
84	r2	7	P5P	C5-N7-C8	3.27	108.42	104.40
84	r2	64	P5P	C5-N7-C8	3.27	108.42	104.40
85	r3	43	P5P	C5-N7-C8	3.27	108.42	104.40
85	r3	57	P5P	C5-N7-C8	3.27	108.41	104.40
85	r3	22	P5P	C5-N7-C8	3.27	108.41	104.40
84	r2	38	P5P	C5-N7-C8	3.27	108.41	104.40
84	r2	76	P5P	C2-N3-C4	3.27	119.81	111.83
84	r2	65	P5P	N3-C4-N9	3.27	132.72	127.17
85	r3	46	P5P	N3-C4-N9	3.27	132.72	127.17
85	r3	30	P5P	C2-N3-C4	3.27	119.81	111.83

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	44	P5P	C2-N3-C4	3.27	119.81	111.83
84	r2	70	P5P	C2-N3-C4	3.27	119.81	111.83
84	r2	5	P5P	N3-C4-N9	3.27	132.72	127.17
85	r3	71	P5P	C2-N3-C4	3.27	119.81	111.83
85	r3	29	P5P	C2-N3-C4	3.26	119.80	111.83
84	r2	1	P5P	C5-N7-C8	3.26	108.41	104.40
85	r3	19	P5P	C2-N3-C4	3.26	119.80	111.83
85	r3	68	P5P	C2-N3-C4	3.26	119.80	111.83
84	r2	7	P5P	C2-N3-C4	3.26	119.80	111.83
84	r2	44	P5P	N3-C4-N9	3.26	132.71	127.17
85	r3	43	P5P	C2-N3-C4	3.26	119.80	111.83
85	r3	66	P5P	C2-N3-C4	3.26	119.80	111.83
84	r2	38	P5P	N3-C4-N9	3.26	132.71	127.17
85	r3	37	P5P	C5-N7-C8	3.26	108.40	104.40
84	r2	69	P5P	C2-N3-C4	3.26	119.78	111.83
85	r3	74	P5P	C2-N3-C4	3.26	119.78	111.83
84	r2	15	P5P	C5-N7-C8	3.26	108.40	104.40
84	r2	44	P5P	C5-N7-C8	3.26	108.40	104.40
84	r2	6	P5P	C2-N3-C4	3.25	119.78	111.83
85	r3	10	P5P	N3-C4-N9	3.25	132.70	127.17
84	r2	14	P5P	C5-N7-C8	3.25	108.39	104.40
84	r2	28	P5P	C2-N3-C4	3.25	119.77	111.83
84	r2	29	P5P	C2-N3-C4	3.25	119.77	111.83
84	r2	6	P5P	C5-N7-C8	3.25	108.39	104.40
84	r2	37	P5P	C2-N3-C4	3.25	119.77	111.83
84	r2	46	P5P	C2-N3-C4	3.25	119.77	111.83
84	r2	14	P5P	C2-N3-C4	3.25	119.76	111.83
84	r2	9	P5P	C5-N7-C8	3.25	108.39	104.40
85	r3	18	P5P	C2-N3-C4	3.25	119.76	111.83
85	r3	45	P5P	C2-N3-C4	3.25	119.76	111.83
84	r2	9	P5P	C2-N3-C4	3.25	119.76	111.83
85	r3	9	P5P	C5-N7-C8	3.25	108.39	104.40
85	r3	9	P5P	C2-N3-C4	3.25	119.76	111.83
85	r3	52	P5P	C2-N3-C4	3.24	119.76	111.83
84	r2	34	P5P	C2-N3-C4	3.24	119.75	111.83
84	r2	21	P5P	N3-C4-N9	3.24	132.69	127.17
85	r3	11	P5P	C2-N3-C4	3.24	119.75	111.83
85	r3	10	P5P	C2-N3-C4	3.24	119.75	111.83
84	r2	22	P5P	C2-N3-C4	3.24	119.75	111.83
85	r3	21	P5P	C2-N3-C4	3.24	119.75	111.83
85	r3	8	P5P	C2-N3-C4	3.24	119.75	111.83
84	r2	10	P5P	C2-N3-C4	3.24	119.75	111.83

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
88	A	1	MHW	CB-CA-C	-3.24	117.68	120.09
84	r2	21	P5P	C2-N3-C4	3.24	119.74	111.83
84	r2	23	P5P	N3-C4-N9	3.24	132.68	127.17
84	r2	19	P5P	C2-N3-C4	3.24	119.74	111.83
84	r2	58	P5P	C2-N3-C4	3.24	119.74	111.83
85	r3	4	P5P	C2-N3-C4	3.24	119.74	111.83
85	r3	42	P5P	N3-C4-N9	3.24	132.67	127.17
85	r3	74	P5P	N3-C4-N9	3.24	132.67	127.17
85	r3	6	P5P	C2-N3-C4	3.24	119.74	111.83
84	r2	27	P5P	C2-N3-C4	3.24	119.74	111.83
84	r2	30	P5P	C2-N3-C4	3.24	119.74	111.83
85	r3	14	P5P	C5-N7-C8	3.24	108.37	104.40
84	r2	31	P5P	C2-N3-C4	3.24	119.73	111.83
85	r3	42	P5P	C2-N3-C4	3.24	119.73	111.83
85	r3	51	P5P	C2-N3-C4	3.24	119.73	111.83
85	r3	31	P5P	C2-N3-C4	3.24	119.73	111.83
85	r3	55	P5P	C2-N3-C4	3.23	119.73	111.83
84	r2	63	P5P	C2-N3-C4	3.23	119.73	111.83
84	r2	18	P5P	C2-N3-C4	3.23	119.73	111.83
84	r2	1	P5P	C2-N3-C4	3.23	119.72	111.83
84	r2	26	P5P	N3-C4-N9	3.23	132.66	127.17
85	r3	22	P5P	C2-N3-C4	3.23	119.72	111.83
85	r3	23	P5P	C2-N3-C4	3.23	119.72	111.83
85	r3	26	P5P	N3-C4-N9	3.23	132.66	127.17
84	r2	57	P5P	C2-N3-C4	3.23	119.72	111.83
84	r2	64	P5P	C2-N3-C4	3.23	119.71	111.83
85	r3	5	P5P	C2-N3-C4	3.23	119.71	111.83
85	r3	46	P5P	C2-N3-C4	3.23	119.71	111.83
84	r2	24	P5P	C2-N3-C4	3.23	119.71	111.83
84	r2	36	P5P	C2-N3-C4	3.23	119.71	111.83
84	r2	52	P5P	C2-N3-C4	3.22	119.71	111.83
85	r3	57	P5P	C2-N3-C4	3.22	119.71	111.83
84	r2	53	P5P	C2-N3-C4	3.22	119.70	111.83
84	r2	23	P5P	C2-N3-C4	3.22	119.70	111.83
85	r3	26	P5P	C2-N3-C4	3.22	119.70	111.83
85	r3	35	P5P	N3-C4-N9	3.22	132.64	127.17
84	r2	15	P5P	C2-N3-C4	3.22	119.69	111.83
85	r3	48	P5P	C2-N3-C4	3.22	119.69	111.83
84	r2	65	P5P	C2-N3-C4	3.22	119.69	111.83
84	r2	26	P5P	C2-N3-C4	3.22	119.69	111.83
84	r2	35	P5P	C2-N3-C4	3.22	119.69	111.83
85	r3	15	P5P	C6-C5-C4	3.21	119.75	116.24

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	74	P5P	C5-N7-C8	3.21	108.34	104.40
84	r2	44	P5P	C2-N3-C4	3.21	119.67	111.83
84	r2	38	P5P	C2-N3-C4	3.21	119.66	111.83
85	r3	15	P5P	C2-N3-C4	3.20	119.66	111.83
85	r3	14	P5P	C2-N3-C4	3.20	119.66	111.83
88	A	7	004	CG1-CB-CA	3.20	125.62	120.64
84	r2	76	P5P	C5-N7-C8	3.20	108.33	104.40
84	r2	5	P5P	C2-N3-C4	3.20	119.64	111.83
85	r3	3	P5P	C2-N3-C4	3.20	119.64	111.83
84	r2	71	P5P	C2-N3-C4	3.20	119.64	111.83
84	r2	71	P5P	N3-C4-N9	3.17	132.57	127.17
84	r2	73	P5P	C2-N3-C4	3.17	119.58	111.83
84	r2	70	P5P	C6-C5-C4	3.17	119.70	116.24
84	r2	1	P5P	C6-C5-C4	3.16	119.69	116.24
84	r2	22	P5P	C6-C5-C4	3.16	119.69	116.24
84	r2	73	P5P	C6-C5-C4	3.15	119.69	116.24
85	r3	21	P5P	C6-C5-C4	3.12	119.66	116.24
84	r2	36	P5P	C6-C5-C4	3.12	119.65	116.24
84	r2	7	P5P	C6-C5-C4	3.12	119.65	116.24
85	r3	37	P5P	C6-C5-C4	3.12	119.65	116.24
85	r3	30	P5P	C6-C5-C4	3.11	119.64	116.24
84	r2	35	P5P	C6-C5-C4	3.11	119.64	116.24
85	r3	9	P5P	C6-C5-C4	3.10	119.64	116.24
84	r2	9	P5P	C6-C5-C4	3.10	119.63	116.24
84	r2	27	P5P	C6-C5-C4	3.10	119.63	116.24
84	r2	29	P5P	C6-C5-C4	3.10	119.63	116.24
84	r2	5	P5P	C6-C5-C4	3.10	119.63	116.24
84	r2	15	P5P	C6-C5-C4	3.09	119.62	116.24
85	r3	45	P5P	C6-C5-C4	3.09	119.62	116.24
85	r3	44	P5P	C6-C5-C4	3.09	119.62	116.24
85	r3	4	P5P	C6-C5-C4	3.09	119.62	116.24
84	r2	65	P5P	C6-C5-C4	3.09	119.62	116.24
85	r3	14	P5P	C6-C5-C4	3.09	119.61	116.24
85	r3	55	P5P	C6-C5-C4	3.09	119.61	116.24
85	r3	57	P5P	C6-C5-C4	3.08	119.61	116.24
84	r2	52	P5P	C6-C5-C4	3.08	119.61	116.24
84	r2	64	P5P	C6-C5-C4	3.08	119.60	116.24
85	r3	51	P5P	C6-C5-C4	3.08	119.60	116.24
85	r3	43	P5P	C6-C5-C4	3.08	119.60	116.24
84	r2	6	P5P	C6-C5-C4	3.07	119.60	116.24
84	r2	18	P5P	C6-C5-C4	3.07	119.60	116.24
85	r3	23	P5P	C6-C5-C4	3.07	119.60	116.24

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	18	P5P	C6-C5-C4	3.07	119.60	116.24
84	r2	46	P5P	C6-C5-C4	3.07	119.59	116.24
85	r3	29	P5P	C6-C5-C4	3.07	119.59	116.24
85	r3	31	P5P	C6-C5-C4	3.07	119.59	116.24
84	r2	69	P5P	C6-C5-C4	3.07	119.59	116.24
85	r3	5	P5P	C6-C5-C4	3.06	119.59	116.24
85	r3	8	P5P	C6-C5-C4	3.06	119.59	116.24
85	r3	3	P5P	C6-C5-C4	3.06	119.59	116.24
85	r3	68	P5P	C6-C5-C4	3.06	119.59	116.24
84	r2	31	P5P	C6-C5-C4	3.06	119.58	116.24
84	r2	30	P5P	C6-C5-C4	3.05	119.58	116.24
84	r2	53	P5P	C6-C5-C4	3.05	119.57	116.24
84	r2	10	P5P	C6-C5-C4	3.05	119.57	116.24
85	r3	6	P5P	C6-C5-C4	3.05	119.57	116.24
85	r3	22	P5P	C6-C5-C4	3.05	119.57	116.24
84	r2	58	P5P	C6-C5-C4	3.04	119.57	116.24
84	r2	23	P5P	C6-C5-C4	3.04	119.57	116.24
84	r2	24	P5P	C6-C5-C4	3.04	119.57	116.24
85	r3	52	P5P	C6-C5-C4	3.04	119.57	116.24
84	r2	37	P5P	C6-C5-C4	3.03	119.56	116.24
85	r3	11	P5P	C6-C5-C4	3.03	119.56	116.24
84	r2	28	P5P	C6-C5-C4	3.03	119.56	116.24
84	r2	19	P5P	C6-C5-C4	3.03	119.55	116.24
85	r3	48	P5P	C6-C5-C4	3.02	119.55	116.24
85	r3	46	P5P	C6-C5-C4	3.02	119.55	116.24
84	r2	38	P5P	C6-C5-C4	3.02	119.54	116.24
84	r2	76	P5P	C6-C5-C4	3.02	119.54	116.24
85	r3	19	P5P	C6-C5-C4	3.02	119.54	116.24
85	r3	10	P5P	C6-C5-C4	3.01	119.54	116.24
84	r2	57	P5P	C6-C5-C4	3.01	119.54	116.24
85	r3	66	P5P	C6-C5-C4	3.01	119.53	116.24
84	r2	14	P5P	C6-C5-C4	3.01	119.53	116.24
85	r3	71	P5P	C6-C5-C4	3.01	119.53	116.24
84	r2	26	P5P	C6-C5-C4	3.00	119.53	116.24
84	r2	63	P5P	C6-C5-C4	3.00	119.53	116.24
84	r2	44	P5P	C6-C5-C4	3.00	119.52	116.24
84	r2	34	P5P	C6-C5-C4	2.97	119.49	116.24
84	r2	21	P5P	C6-C5-C4	2.94	119.45	116.24
85	r3	26	P5P	C6-C5-C4	2.88	119.39	116.24
84	r2	71	P5P	C6-C5-C4	2.87	119.38	116.24
88	A	5	MHU	O-C-CA	-2.79	117.60	124.77
85	r3	42	P5P	C6-C5-C4	2.79	119.29	116.24

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
85	r3	35	P5P	C6-C5-C4	2.78	119.28	116.24
85	r3	74	P5P	C6-C5-C4	2.71	119.21	116.24
88	A	1	MHW	CE-N-CA	2.66	121.26	116.83
88	A	6	MHV	CD2-CG-CB	2.62	119.86	115.89
88	A	5	MHU	CM-N-CA	2.44	121.00	113.70
88	A	6	MHV	CB-CA-N	-2.42	107.50	112.50
84	r2	22	P5P	C4-N9-C1'	2.33	132.09	126.63
84	r2	9	P5P	C4-N9-C1'	2.32	132.06	126.63
85	r3	14	P5P	C4-N9-C1'	2.28	131.97	126.63
85	r3	15	P5P	C4-N9-C1'	2.27	131.93	126.63
85	r3	44	P5P	C4-N9-C1'	2.26	131.93	126.63
85	r3	57	P5P	C4-N9-C1'	2.26	131.93	126.63
84	r2	70	P5P	C4-N9-C1'	2.24	131.86	126.63
84	r2	36	P5P	C4-N9-C1'	2.22	131.84	126.63
84	r2	1	P5P	C4-N9-C1'	2.22	131.82	126.63
85	r3	9	P5P	C4-N9-C1'	2.22	131.82	126.63
85	r3	18	P5P	C4-N9-C1'	2.21	131.79	126.63
84	r2	73	P5P	C4-N9-C1'	2.20	131.78	126.63
85	r3	30	P5P	C4-N9-C1'	2.19	131.75	126.63
84	r2	7	P5P	C4-N9-C1'	2.17	131.71	126.63
84	r2	64	P5P	C4-N9-C1'	2.16	131.69	126.63
84	r2	10	P5P	C4-N9-C1'	2.16	131.68	126.63
85	r3	8	P5P	C4-N9-C1'	2.15	131.65	126.63
84	r2	18	P5P	C4-N9-C1'	2.14	131.64	126.63
84	r2	69	P5P	C4-N9-C1'	2.14	131.64	126.63
85	r3	55	P5P	C4-N9-C1'	2.14	131.63	126.63
85	r3	37	P5P	C4-N9-C1'	2.14	131.63	126.63
84	r2	5	P5P	C4-N9-C1'	2.13	131.62	126.63
85	r3	29	P5P	C4-N9-C1'	2.12	131.60	126.63
84	r2	27	P5P	C4-N9-C1'	2.12	131.60	126.63
85	r3	45	P5P	C4-N9-C1'	2.12	131.59	126.63
84	r2	31	P5P	C4-N9-C1'	2.11	131.57	126.63
85	r3	21	P5P	C4-N9-C1'	2.11	131.56	126.63
88	A	5	MHU	CB-CA-N	-2.10	107.38	110.48
85	r3	68	P5P	C4-N9-C1'	2.10	131.54	126.63
84	r2	28	P5P	C4-N9-C1'	2.10	131.54	126.63
84	r2	15	P5P	C4-N9-C1'	2.10	131.54	126.63
84	r2	76	P5P	C4-N9-C1'	2.09	131.53	126.63
85	r3	43	P5P	C4-N9-C1'	2.09	131.52	126.63
84	r2	52	P5P	C4-N9-C1'	2.08	131.51	126.63
85	r3	3	P5P	C4-N9-C1'	2.08	131.50	126.63
84	r2	44	P5P	C4-N9-C1'	2.08	131.50	126.63

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
88	A	6	MHV	CE-N-CA	2.07	116.43	111.75
84	r2	29	P5P	C4-N9-C1'	2.07	131.47	126.63
85	r3	22	P5P	C4-N9-C1'	2.06	131.44	126.63
84	r2	46	P5P	C4-N9-C1'	2.06	131.44	126.63
84	r2	35	P5P	C4-N9-C1'	2.06	131.44	126.63
84	r2	30	P5P	C4-N9-C1'	2.05	131.42	126.63
84	r2	19	P5P	C4-N9-C1'	2.04	131.41	126.63
85	r3	71	P5P	C4-N9-C1'	2.04	131.41	126.63
85	r3	52	P5P	C4-N9-C1'	2.04	131.40	126.63
85	r3	23	P5P	C4-N9-C1'	2.03	131.39	126.63
84	r2	6	P5P	C4-N9-C1'	2.03	131.39	126.63
85	r3	5	P5P	C4-N9-C1'	2.03	131.38	126.63
84	r2	63	P5P	C4-N9-C1'	2.03	131.37	126.63
85	r3	11	P5P	C4-N9-C1'	2.02	131.35	126.63
84	r2	24	P5P	C4-N9-C1'	2.02	131.35	126.63
84	r2	38	P5P	C4-N9-C1'	2.01	131.33	126.63
85	r3	31	P5P	C4-N9-C1'	2.01	131.33	126.63
85	r3	51	P5P	C4-N9-C1'	2.00	131.32	126.63

There are no chirality outliers.

All (284) torsion outliers are listed below:

Mol	Chain	Res	Type	Atoms
83	r1	46	Y5P	C4'-C5'-O5'-P
83	r1	46	Y5P	O4'-C1'-N1-C6
83	r1	47	Y5P	O4'-C4'-C5'-O5'
83	r1	55	Y5P	O4'-C4'-C5'-O5'
83	r1	57	Y5P	O4'-C1'-N1-C2
84	r2	4	Y5P	O4'-C1'-N1-C2
84	r2	11	Y5P	O4'-C1'-N1-C2
84	r2	33	Y5P	C2'-C1'-N1-C6
84	r2	35	P5P	C3'-C4'-C5'-O5'
84	r2	35	P5P	O4'-C4'-C5'-O5'
84	r2	39	Y5P	O4'-C1'-N1-C2
84	r2	41	Y5P	O4'-C1'-N1-C2
84	r2	45	Y5P	O4'-C4'-C5'-O5'
84	r2	46	P5P	C2'-C1'-N9-C8
84	r2	47	Y5P	O4'-C1'-N1-C2
84	r2	50	Y5P	O4'-C4'-C5'-O5'
84	r2	54	Y5P	O4'-C1'-N1-C2
84	r2	75	Y5P	O4'-C1'-N1-C2
85	r3	3	P5P	O4'-C4'-C5'-O5'

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Mol	Chain	Res	Type	Atoms
85	r3	13	Y5P	C4'-C5'-O5'-P
85	r3	16	Y5P	C4'-C5'-O5'-P
85	r3	21	P5P	C3'-C4'-C5'-O5'
85	r3	21	P5P	C4'-C5'-O5'-P
85	r3	24	Y5P	O4'-C1'-N1-C2
85	r3	25	Y5P	O4'-C1'-N1-C2
85	r3	27	Y5P	O4'-C1'-N1-C2
85	r3	30	P5P	O4'-C4'-C5'-O5'
85	r3	34	Y5P	O4'-C4'-C5'-O5'
85	r3	34	Y5P	C2'-C1'-N1-C2
85	r3	38	Y5P	O4'-C4'-C5'-O5'
85	r3	47	Y5P	O4'-C4'-C5'-O5'
85	r3	47	Y5P	O4'-C1'-N1-C2
85	r3	59	Y5P	O4'-C1'-N1-C2
85	r3	61	Y5P	O4'-C1'-N1-C2
85	r3	63	Y5P	O4'-C1'-N1-C2
85	r3	70	Y5P	O4'-C1'-N1-C2
85	r3	74	P5P	O4'-C1'-N9-C4
85	r3	74	P5P	O4'-C1'-N9-C8
88	A	5	MHU	O-C-CA-CB
83	r1	44	Y5P	O4'-C1'-N1-C2
83	r1	47	Y5P	O4'-C1'-N1-C2
83	r1	48	Y5P	O4'-C1'-N1-C2
83	r1	50	Y5P	O4'-C1'-N1-C2
83	r1	51	Y5P	O4'-C1'-N1-C2
83	r1	52	Y5P	O4'-C1'-N1-C2
83	r1	54	Y5P	O4'-C1'-N1-C2
84	r2	2	Y5P	O4'-C1'-N1-C2
84	r2	3	Y5P	O4'-C1'-N1-C2
84	r2	8	Y5P	O4'-C1'-N1-C2
84	r2	12	Y5P	O4'-C1'-N1-C2
84	r2	13	Y5P	O4'-C1'-N1-C2
84	r2	16	Y5P	O4'-C1'-N1-C2
84	r2	17	Y5P	O4'-C1'-N1-C2
84	r2	20	Y5P	O4'-C1'-N1-C2
84	r2	25	Y5P	O4'-C1'-N1-C2
84	r2	32	Y5P	O4'-C1'-N1-C2
84	r2	40	Y5P	O4'-C1'-N1-C2
84	r2	42	Y5P	O4'-C1'-N1-C2
84	r2	43	Y5P	O4'-C1'-N1-C2
84	r2	45	Y5P	O4'-C1'-N1-C2
84	r2	48	Y5P	O4'-C1'-N1-C2

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Mol	Chain	Res	Type	Atoms
84	r2	49	Y5P	O4'-C1'-N1-C2
84	r2	50	Y5P	O4'-C1'-N1-C2
84	r2	51	Y5P	O4'-C1'-N1-C2
84	r2	56	Y5P	O4'-C1'-N1-C2
84	r2	59	Y5P	O4'-C1'-N1-C2
84	r2	60	Y5P	O4'-C1'-N1-C2
84	r2	61	Y5P	O4'-C1'-N1-C2
84	r2	62	Y5P	O4'-C1'-N1-C2
84	r2	66	Y5P	O4'-C1'-N1-C2
84	r2	67	Y5P	O4'-C1'-N1-C2
84	r2	68	Y5P	O4'-C1'-N1-C2
84	r2	72	Y5P	O4'-C1'-N1-C2
84	r2	74	Y5P	O4'-C1'-N1-C2
85	r3	12	Y5P	O4'-C1'-N1-C2
85	r3	16	Y5P	O4'-C1'-N1-C2
85	r3	28	Y5P	O4'-C1'-N1-C2
85	r3	36	Y5P	O4'-C1'-N1-C2
85	r3	38	Y5P	O4'-C1'-N1-C2
85	r3	39	Y5P	O4'-C1'-N1-C2
85	r3	49	Y5P	O4'-C1'-N1-C2
85	r3	53	Y5P	O4'-C1'-N1-C2
85	r3	54	Y5P	O4'-C1'-N1-C2
85	r3	60	Y5P	O4'-C1'-N1-C2
85	r3	62	Y5P	O4'-C1'-N1-C2
85	r3	64	Y5P	O4'-C1'-N1-C2
85	r3	65	Y5P	O4'-C1'-N1-C2
85	r3	67	Y5P	O4'-C1'-N1-C2
85	r3	69	Y5P	O4'-C1'-N1-C2
85	r3	73	Y5P	O4'-C1'-N1-C2
85	r3	41	Y5P	O4'-C1'-N1-C6
84	r2	33	Y5P	C2'-C1'-N1-C2
88	A	5	MHU	CE1-CZ-NZ-CZ2
83	r1	47	Y5P	C3'-C4'-C5'-O5'
84	r2	7	P5P	O4'-C4'-C5'-O5'
84	r2	13	Y5P	O4'-C4'-C5'-O5'
84	r2	13	Y5P	C3'-C4'-C5'-O5'
84	r2	45	Y5P	C3'-C4'-C5'-O5'
84	r2	50	Y5P	C3'-C4'-C5'-O5'
84	r2	74	Y5P	O4'-C4'-C5'-O5'
84	r2	74	Y5P	C3'-C4'-C5'-O5'
85	r3	19	P5P	C3'-C4'-C5'-O5'
85	r3	30	P5P	C3'-C4'-C5'-O5'

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Mol	Chain	Res	Type	Atoms
85	r3	38	Y5P	C3'-C4'-C5'-O5'
85	r3	39	Y5P	O4'-C4'-C5'-O5'
85	r3	42	P5P	C3'-C4'-C5'-O5'
85	r3	45	P5P	O4'-C4'-C5'-O5'
85	r3	47	Y5P	C3'-C4'-C5'-O5'
85	r3	54	Y5P	O4'-C4'-C5'-O5'
85	r3	55	P5P	C3'-C4'-C5'-O5'
85	r3	55	P5P	O4'-C4'-C5'-O5'
85	r3	57	P5P	O4'-C4'-C5'-O5'
85	r3	63	Y5P	O4'-C4'-C5'-O5'
85	r3	63	Y5P	C3'-C4'-C5'-O5'
85	r3	74	P5P	C3'-C4'-C5'-O5'
88	A	5	MHU	CE2-CZ-NZ-CZ1
84	r2	47	Y5P	C4'-C5'-O5'-P
88	A	5	MHU	CE1-CZ-NZ-CZ1
88	A	5	MHU	CE2-CZ-NZ-CZ2
85	r3	34	Y5P	C2'-C1'-N1-C6
83	r1	53	Y5P	O4'-C4'-C5'-O5'
84	r2	1	P5P	O4'-C4'-C5'-O5'
84	r2	9	P5P	O4'-C4'-C5'-O5'
84	r2	14	P5P	O4'-C4'-C5'-O5'
84	r2	21	P5P	C3'-C4'-C5'-O5'
84	r2	21	P5P	O4'-C4'-C5'-O5'
84	r2	66	Y5P	O4'-C4'-C5'-O5'
84	r2	66	Y5P	C3'-C4'-C5'-O5'
84	r2	72	Y5P	C3'-C4'-C5'-O5'
84	r2	73	P5P	O4'-C4'-C5'-O5'
85	r3	7	Y5P	O4'-C4'-C5'-O5'
85	r3	16	Y5P	O4'-C4'-C5'-O5'
85	r3	18	P5P	O4'-C4'-C5'-O5'
85	r3	21	P5P	O4'-C4'-C5'-O5'
85	r3	29	P5P	O4'-C4'-C5'-O5'
85	r3	45	P5P	C3'-C4'-C5'-O5'
85	r3	50	Y5P	O4'-C4'-C5'-O5'
85	r3	50	Y5P	C3'-C4'-C5'-O5'
85	r3	74	P5P	O4'-C4'-C5'-O5'
85	r3	35	P5P	C4'-C5'-O5'-P
84	r2	46	P5P	C2'-C1'-N9-C4
85	r3	50	Y5P	O4'-C1'-N1-C2
83	r1	55	Y5P	C3'-C4'-C5'-O5'
84	r2	69	P5P	C3'-C4'-C5'-O5'
85	r3	18	P5P	C3'-C4'-C5'-O5'

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Mol	Chain	Res	Type	Atoms
85	r3	34	Y5P	C3'-C4'-C5'-O5'
85	r3	48	P5P	O4'-C4'-C5'-O5'
83	r1	53	Y5P	O4'-C1'-N1-C2
83	r1	55	Y5P	O4'-C1'-N1-C2
85	r3	7	Y5P	O4'-C1'-N1-C2
85	r3	20	Y5P	O4'-C1'-N1-C2
85	r3	34	Y5P	O4'-C1'-N1-C2
85	r3	47	Y5P	C4'-C5'-O5'-P
83	r1	46	Y5P	C2'-C1'-N1-C2
83	r1	46	Y5P	C2'-C1'-N1-C6
83	r1	54	Y5P	C3'-C4'-C5'-O5'
84	r2	7	P5P	C3'-C4'-C5'-O5'
84	r2	14	P5P	C3'-C4'-C5'-O5'
85	r3	3	P5P	C3'-C4'-C5'-O5'
85	r3	7	Y5P	C3'-C4'-C5'-O5'
85	r3	16	Y5P	C3'-C4'-C5'-O5'
85	r3	54	Y5P	C3'-C4'-C5'-O5'
83	r1	53	Y5P	C3'-C4'-C5'-O5'
84	r2	33	Y5P	O4'-C4'-C5'-O5'
84	r2	72	Y5P	O4'-C4'-C5'-O5'
85	r3	19	P5P	O4'-C4'-C5'-O5'
85	r3	42	P5P	O4'-C4'-C5'-O5'
85	r3	53	Y5P	O4'-C4'-C5'-O5'
85	r3	56	Y5P	O4'-C4'-C5'-O5'
85	r3	59	Y5P	O4'-C4'-C5'-O5'
83	r1	49	Y5P	O4'-C4'-C5'-O5'
84	r2	46	P5P	O4'-C4'-C5'-O5'
85	r3	43	P5P	O4'-C4'-C5'-O5'
85	r3	56	Y5P	C3'-C4'-C5'-O5'
83	r1	45	Y5P	O4'-C1'-N1-C2
84	r2	55	Y5P	O4'-C1'-N1-C2
84	r2	1	P5P	C3'-C4'-C5'-O5'
84	r2	33	Y5P	C3'-C4'-C5'-O5'
85	r3	35	P5P	C3'-C4'-C5'-O5'
85	r3	53	Y5P	C3'-C4'-C5'-O5'
85	r3	57	P5P	C3'-C4'-C5'-O5'
84	r2	33	Y5P	C4'-C5'-O5'-P
85	r3	46	P5P	C4'-C5'-O5'-P
83	r1	54	Y5P	O4'-C4'-C5'-O5'
84	r2	69	P5P	O4'-C4'-C5'-O5'
85	r3	39	Y5P	C3'-C4'-C5'-O5'
85	r3	66	P5P	O4'-C4'-C5'-O5'

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Mol	Chain	Res	Type	Atoms
84	r2	73	P5P	C3'-C4'-C5'-O5'
85	r3	29	P5P	C3'-C4'-C5'-O5'
84	r2	15	P5P	C2'-C1'-N9-C8
84	r2	9	P5P	C3'-C4'-C5'-O5'
84	r2	46	P5P	C3'-C4'-C5'-O5'
85	r3	36	Y5P	O4'-C4'-C5'-O5'
85	r3	3	P5P	C4'-C5'-O5'-P
84	r2	33	Y5P	O4'-C1'-N1-C6
84	r2	19	P5P	C2'-C1'-N9-C8
85	r3	9	P5P	C2'-C1'-N9-C8
83	r1	49	Y5P	O4'-C1'-N1-C2
85	r3	33	Y5P	O4'-C1'-N1-C2
85	r3	72	Y5P	O4'-C1'-N1-C2
85	r3	32	Y5P	O4'-C1'-N1-C2
84	r2	19	P5P	O4'-C1'-N9-C4
83	r1	56	Y5P	O4'-C1'-N1-C6
85	r3	44	P5P	C2'-C1'-N9-C4
85	r3	1	Y5P	O4'-C4'-C5'-O5'
85	r3	4	P5P	O4'-C4'-C5'-O5'
85	r3	58	Y5P	O4'-C4'-C5'-O5'
84	r2	19	P5P	O4'-C1'-N9-C8
84	r2	48	Y5P	C4'-C5'-O5'-P
84	r2	69	P5P	C4'-C5'-O5'-P
84	r2	76	P5P	C4'-C5'-O5'-P
85	r3	44	P5P	C2'-C1'-N9-C8
84	r2	9	P5P	C4'-C5'-O5'-P
84	r2	23	P5P	C4'-C5'-O5'-P
85	r3	18	P5P	C4'-C5'-O5'-P
85	r3	33	Y5P	C4'-C5'-O5'-P
85	r3	45	P5P	C4'-C5'-O5'-P
85	r3	58	Y5P	C4'-C5'-O5'-P
85	r3	72	Y5P	C4'-C5'-O5'-P
84	r2	15	P5P	C2'-C1'-N9-C4
84	r2	24	P5P	C2'-C1'-N9-C4
85	r3	14	P5P	C4'-C5'-O5'-P
85	r3	29	P5P	C4'-C5'-O5'-P
84	r2	49	Y5P	O4'-C4'-C5'-O5'
84	r2	9	P5P	O4'-C1'-N9-C8
84	r2	9	P5P	O4'-C1'-N9-C4
84	r2	24	P5P	C2'-C1'-N9-C8
85	r3	2	Y5P	O4'-C1'-N1-C2
85	r3	13	Y5P	O4'-C1'-N1-C2

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Mol	Chain	Res	Type	Atoms
85	r3	56	Y5P	O4'-C1'-N1-C2
85	r3	58	Y5P	O4'-C1'-N1-C2
85	r3	40	Y5P	C4'-C5'-O5'-P
85	r3	56	Y5P	C4'-C5'-O5'-P
85	r3	9	P5P	O4'-C1'-N9-C8
85	r3	1	Y5P	O4'-C1'-N1-C2
85	r3	40	Y5P	O4'-C1'-N1-C2
85	r3	13	Y5P	C2'-C1'-N1-C2
85	r3	56	Y5P	C2'-C1'-N1-C2
84	r2	55	Y5P	C4'-C5'-O5'-P
83	r1	51	Y5P	O4'-C4'-C5'-O5'
85	r3	35	P5P	O4'-C4'-C5'-O5'
85	r3	48	P5P	C3'-C4'-C5'-O5'
85	r3	61	Y5P	C3'-C4'-C5'-O5'
85	r3	9	P5P	O4'-C1'-N9-C4
85	r3	2	Y5P	C2'-C1'-N1-C2
84	r2	17	Y5P	C4'-C5'-O5'-P
85	r3	4	P5P	C4'-C5'-O5'-P
85	r3	69	Y5P	C4'-C5'-O5'-P
84	r2	1	P5P	C4'-C5'-O5'-P
83	r1	50	Y5P	C2'-C1'-N1-C6
84	r2	17	Y5P	C2'-C1'-N1-C6
85	r3	60	Y5P	C2'-C1'-N1-C6
85	r3	69	Y5P	C2'-C1'-N1-C6
85	r3	14	P5P	C2'-C1'-N9-C8
85	r3	38	Y5P	C2'-C1'-N1-C6
85	r3	56	Y5P	C2'-C1'-N1-C6
84	r2	46	P5P	C4'-C5'-O5'-P
85	r3	58	Y5P	C3'-C4'-C5'-O5'
85	r3	1	Y5P	C2'-C1'-N1-C2
85	r3	44	P5P	O4'-C1'-N9-C8
84	r2	3	Y5P	C2'-C1'-N1-C6
84	r2	47	Y5P	C3'-C4'-C5'-O5'
84	r2	61	Y5P	C4'-C5'-O5'-P
85	r3	46	P5P	C2'-C1'-N9-C8
85	r3	13	Y5P	C2'-C1'-N1-C6
85	r3	40	Y5P	C2'-C1'-N1-C2
83	r1	49	Y5P	C3'-C4'-C5'-O5'
84	r2	49	Y5P	C3'-C4'-C5'-O5'
85	r3	4	P5P	C3'-C4'-C5'-O5'
85	r3	5	P5P	C3'-C4'-C5'-O5'
84	r2	22	P5P	C4'-C5'-O5'-P

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Mol	Chain	Res	Type	Atoms
85	r3	9	P5P	C4'-C5'-O5'-P
85	r3	43	P5P	C2'-C1'-N9-C8
85	r3	57	P5P	C2'-C1'-N9-C8
85	r3	9	P5P	C2'-C1'-N9-C4
85	r3	14	P5P	C2'-C1'-N9-C4
85	r3	13	Y5P	C3'-C4'-C5'-O5'
84	r2	15	P5P	O4'-C1'-N9-C8
85	r3	2	Y5P	O4'-C1'-N1-C6
85	r3	13	Y5P	O4'-C1'-N1-C6
85	r3	56	Y5P	O4'-C1'-N1-C6
84	r2	15	P5P	O4'-C1'-N9-C4
83	r1	44	Y5P	C4'-C5'-O5'-P
83	r1	53	Y5P	C4'-C5'-O5'-P

There are no ring outliers.

112 monomers are involved in 131 short contacts:

Mol	Chain	Res	Type	Clashes	Symm-Clashes
84	r2	24	P5P	3	0
84	r2	71	P5P	1	0
85	r3	4	P5P	1	0
85	r3	22	P5P	3	0
84	r2	4	Y5P	2	0
84	r2	2	Y5P	2	0
84	r2	38	P5P	1	0
84	r2	72	Y5P	1	0
85	r3	23	P5P	1	0
84	r2	6	P5P	2	0
84	r2	35	P5P	4	0
84	r2	61	Y5P	1	0
84	r2	5	P5P	2	0
83	r1	51	Y5P	1	0
84	r2	19	P5P	2	0
85	r3	34	Y5P	1	0
85	r3	56	Y5P	3	0
85	r3	44	P5P	2	0
83	r1	57	Y5P	2	0
85	r3	30	P5P	1	0
85	r3	72	Y5P	2	0
85	r3	29	P5P	1	0
83	r1	50	Y5P	7	0
85	r3	70	Y5P	4	0

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Mol	Chain	Res	Type	Clashes	Symm-Clashes
84	r2	49	Y5P	2	0
85	r3	3	P5P	2	0
83	r1	53	Y5P	1	0
84	r2	34	P5P	3	0
83	r1	47	Y5P	4	0
85	r3	15	P5P	1	0
83	r1	52	Y5P	1	0
85	r3	62	Y5P	1	0
84	r2	68	Y5P	1	0
83	r1	46	Y5P	6	0
85	r3	74	P5P	3	0
85	r3	69	Y5P	2	0
85	r3	58	Y5P	1	0
85	r3	61	Y5P	1	0
84	r2	9	P5P	7	0
88	A	5	MHU	2	0
85	r3	53	Y5P	2	0
85	r3	38	Y5P	4	0
84	r2	17	Y5P	1	0
85	r3	32	Y5P	1	0
85	r3	2	Y5P	3	0
83	r1	45	Y5P	1	0
84	r2	60	Y5P	3	0
85	r3	25	Y5P	5	0
84	r2	10	P5P	1	0
85	r3	33	Y5P	2	0
84	r2	37	P5P	1	0
84	r2	62	Y5P	1	0
85	r3	45	P5P	1	0
84	r2	23	P5P	3	0
84	r2	63	P5P	1	0
83	r1	48	Y5P	1	0
84	r2	36	P5P	1	0
84	r2	58	P5P	1	0
85	r3	42	P5P	4	0
84	r2	46	P5P	5	0
85	r3	37	P5P	3	0
85	r3	65	Y5P	1	0
84	r2	1	P5P	1	0
84	r2	12	Y5P	1	0
84	r2	29	P5P	1	0
84	r2	73	P5P	1	0

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Mol	Chain	Res	Type	Clashes	Symm-Clashes
84	r2	33	Y5P	1	0
85	r3	9	P5P	2	0
85	r3	39	Y5P	2	0
84	r2	59	Y5P	2	0
84	r2	45	Y5P	6	0
85	r3	71	P5P	2	0
85	r3	13	Y5P	1	0
84	r2	48	Y5P	1	0
84	r2	74	Y5P	1	0
84	r2	31	P5P	1	0
85	r3	46	P5P	1	0
85	r3	1	Y5P	1	0
83	r1	55	Y5P	1	0
83	r1	56	Y5P	2	0
85	r3	68	P5P	2	0
84	r2	8	Y5P	1	0
85	r3	63	Y5P	1	0
84	r2	67	Y5P	1	0
84	r2	44	P5P	1	0
85	r3	16	Y5P	1	0
85	r3	24	Y5P	1	0
85	r3	36	Y5P	2	0
85	r3	54	Y5P	7	0
84	r2	32	Y5P	1	0
84	r2	43	Y5P	1	0
83	r1	49	Y5P	4	0
85	r3	12	Y5P	1	0
84	r2	14	P5P	1	0
88	A	7	004	2	0
85	r3	14	P5P	2	0
85	r3	35	P5P	3	0
85	r3	47	Y5P	2	0
85	r3	26	P5P	4	0
84	r2	70	P5P	3	0
85	r3	55	P5P	1	0
84	r2	13	Y5P	2	0
84	r2	30	P5P	2	0
84	r2	39	Y5P	1	0
85	r3	5	P5P	1	0
88	A	1	MHW	3	0
85	r3	67	Y5P	1	0
85	r3	57	P5P	4	0

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Mol	Chain	Res	Type	Clashes	Symm-Clashes
84	r2	50	Y5P	1	0
84	r2	40	Y5P	1	0
84	r2	11	Y5P	2	0
84	r2	75	Y5P	1	0

5.5 Carbohydrates [i](#)

There are no oligosaccharides in this entry.

5.6 Ligand geometry [i](#)

Of 204 ligands modelled in this entry, 202 are monoatomic - leaving 2 for Mogul analysis.

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$
92	GTP	AX	500	-	33,34,34	0.95	2 (6%)	50,54,54	1.54	8 (16%)
91	DOL	XA	5142	-	47,50,50	3.97	20 (42%)	57,70,70	2.99	15 (26%)

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
92	GTP	AX	500	-	-	6/22/38/38	0/3/3/3
91	DOL	XA	5142	-	-	22/64/77/77	0/2/3/3

All (22) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
91	XA	5142	DOL	C28-C29	10.70	1.56	1.32
91	XA	5142	DOL	C6-N5	10.01	1.49	1.34

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
91	XA	5142	DOL	C22-C23	9.30	1.57	1.32
91	XA	5142	DOL	C10-N9	9.26	1.39	1.29
91	XA	5142	DOL	C19-C20	8.81	1.57	1.34
91	XA	5142	DOL	C26-N25	6.96	1.48	1.34
91	XA	5142	DOL	C22-C20	5.94	1.58	1.46
91	XA	5142	DOL	C13-C10	5.78	1.57	1.50
91	XA	5142	DOL	O36-C37	5.24	1.46	1.34
91	XA	5142	DOL	C8-C6	4.28	1.56	1.49
91	XA	5142	DOL	C16-C14	4.17	1.57	1.51
91	XA	5142	DOL	C28-C26	3.91	1.56	1.48
91	XA	5142	DOL	C12-C8	3.80	1.38	1.34
91	XA	5142	DOL	O18-C17	-2.87	1.38	1.43
91	XA	5142	DOL	O27-C26	-2.78	1.19	1.24
91	XA	5142	DOL	C3-C2	-2.35	1.50	1.54
91	XA	5142	DOL	C24-C23	2.34	1.58	1.49
92	AX	500	GTP	C2-N3	2.16	1.38	1.33
91	XA	5142	DOL	O36-C32	-2.12	1.41	1.44
91	XA	5142	DOL	C17-C19	2.05	1.53	1.50
91	XA	5142	DOL	O7-C6	-2.04	1.18	1.23
92	AX	500	GTP	PB-O3B	2.00	1.61	1.59

All (23) bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
91	XA	5142	DOL	O40-S39-O41	-14.90	101.23	118.13
91	XA	5142	DOL	O11-C10-C13	9.88	126.83	117.70
91	XA	5142	DOL	C24-N25-C26	-5.26	113.64	122.17
91	XA	5142	DOL	O11-C10-N9	-5.26	108.41	114.09
92	AX	500	GTP	C5-C4-N3	-5.01	120.42	128.39
92	AX	500	GTP	C2-N3-C4	4.58	120.19	112.30
91	XA	5142	DOL	C23-C22-C20	-4.13	119.66	125.89
91	XA	5142	DOL	C32-O36-C37	-3.96	111.19	117.76
91	XA	5142	DOL	C4-N5-C1	-3.71	108.16	112.51
91	XA	5142	DOL	O40-S39-C2	3.33	111.72	108.50
92	AX	500	GTP	N9-C4-N3	3.18	132.32	125.95
92	AX	500	GTP	C2-N1-C6	-3.06	119.56	125.11
91	XA	5142	DOL	C17-C16-C14	-2.75	108.14	113.90
92	AX	500	GTP	N9-C8-N7	-2.65	108.49	113.40
91	XA	5142	DOL	C3-C4-N5	2.63	106.11	103.22
92	AX	500	GTP	C5-C6-N1	2.55	119.74	113.25
92	AX	500	GTP	C8-N7-C5	2.51	108.73	104.26
92	AX	500	GTP	O6-C6-C5	-2.42	120.14	126.53

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
91	XA	5142	DOL	O41-S39-C2	2.40	110.81	108.50
91	XA	5142	DOL	C14-C13-C10	-2.28	107.84	112.73
91	XA	5142	DOL	C30-C29-C28	-2.27	120.30	126.30
91	XA	5142	DOL	C30-C32-C33	-2.24	110.48	115.98
91	XA	5142	DOL	O36-C32-C30	2.10	110.59	107.12

There are no chirality outliers.

All (28) torsion outliers are listed below:

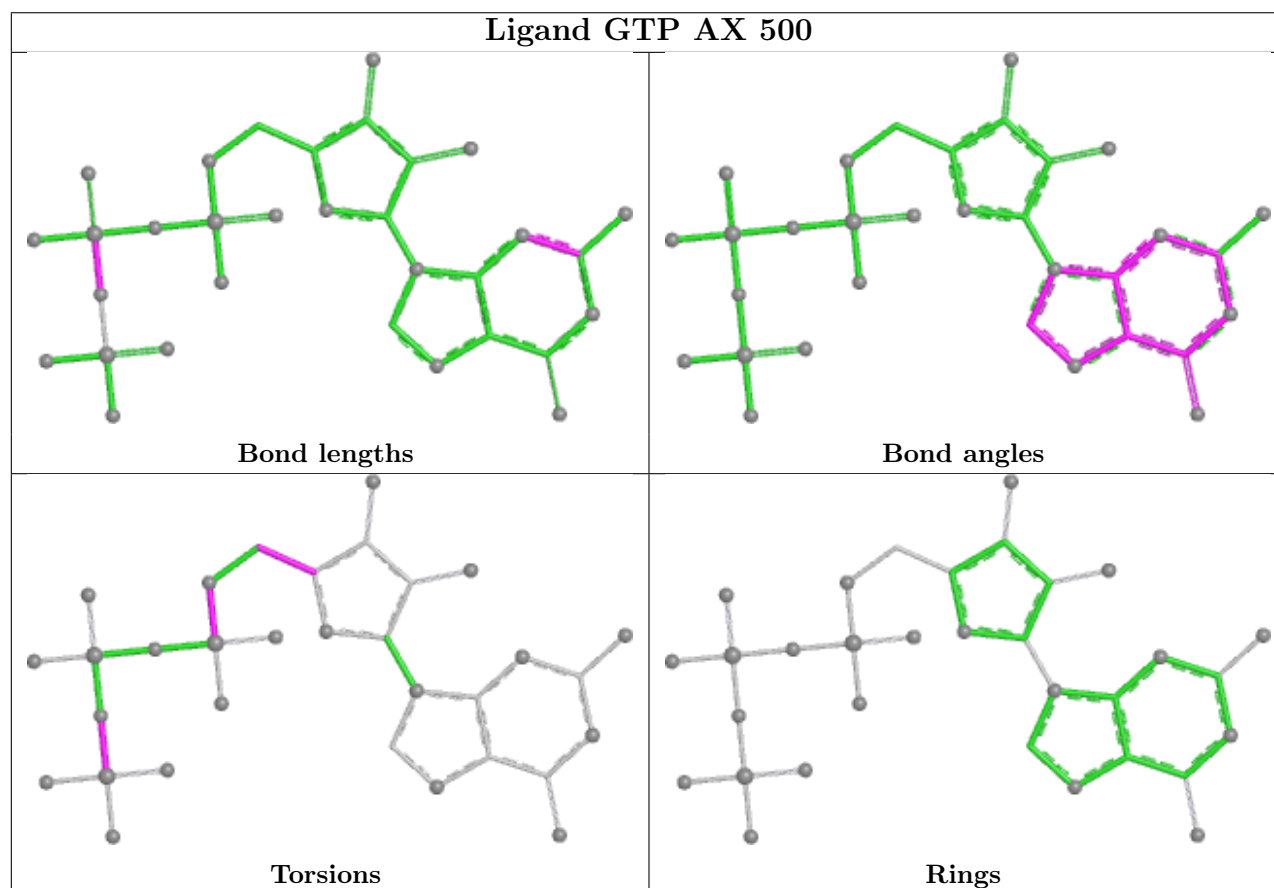
Mol	Chain	Res	Type	Atoms
91	XA	5142	DOL	C3-C2-S39-O41
91	XA	5142	DOL	C3-C2-S39-C42
91	XA	5142	DOL	C1-C2-S39-O41
91	XA	5142	DOL	C1-C2-S39-O40
91	XA	5142	DOL	C1-C2-S39-C42
91	XA	5142	DOL	C43-C42-S39-C2
91	XA	5142	DOL	C43-C42-S39-O41
91	XA	5142	DOL	O11-C10-C13-C14
91	XA	5142	DOL	C16-C17-C19-C20
91	XA	5142	DOL	O18-C17-C19-C20
91	XA	5142	DOL	C29-C30-C32-C33
91	XA	5142	DOL	C31-C30-C32-C33
92	AX	500	GTP	PB-O3B-PG-O3G
92	AX	500	GTP	C5'-O5'-PA-O3A
92	AX	500	GTP	C5'-O5'-PA-O2A
91	XA	5142	DOL	C1-C37-O36-C32
91	XA	5142	DOL	O38-C37-O36-C32
91	XA	5142	DOL	C28-C29-C30-C31
91	XA	5142	DOL	N9-C10-C13-C14
92	AX	500	GTP	O4'-C4'-C5'-O5'
92	AX	500	GTP	C3'-C4'-C5'-O5'
91	XA	5142	DOL	C29-C30-C32-O36
91	XA	5142	DOL	C31-C30-C32-O36
91	XA	5142	DOL	C28-C29-C30-C32
92	AX	500	GTP	PB-O3B-PG-O1G
91	XA	5142	DOL	S39-C42-C43-N44
91	XA	5142	DOL	C42-C43-N44-C47
91	XA	5142	DOL	C42-C43-N44-C45

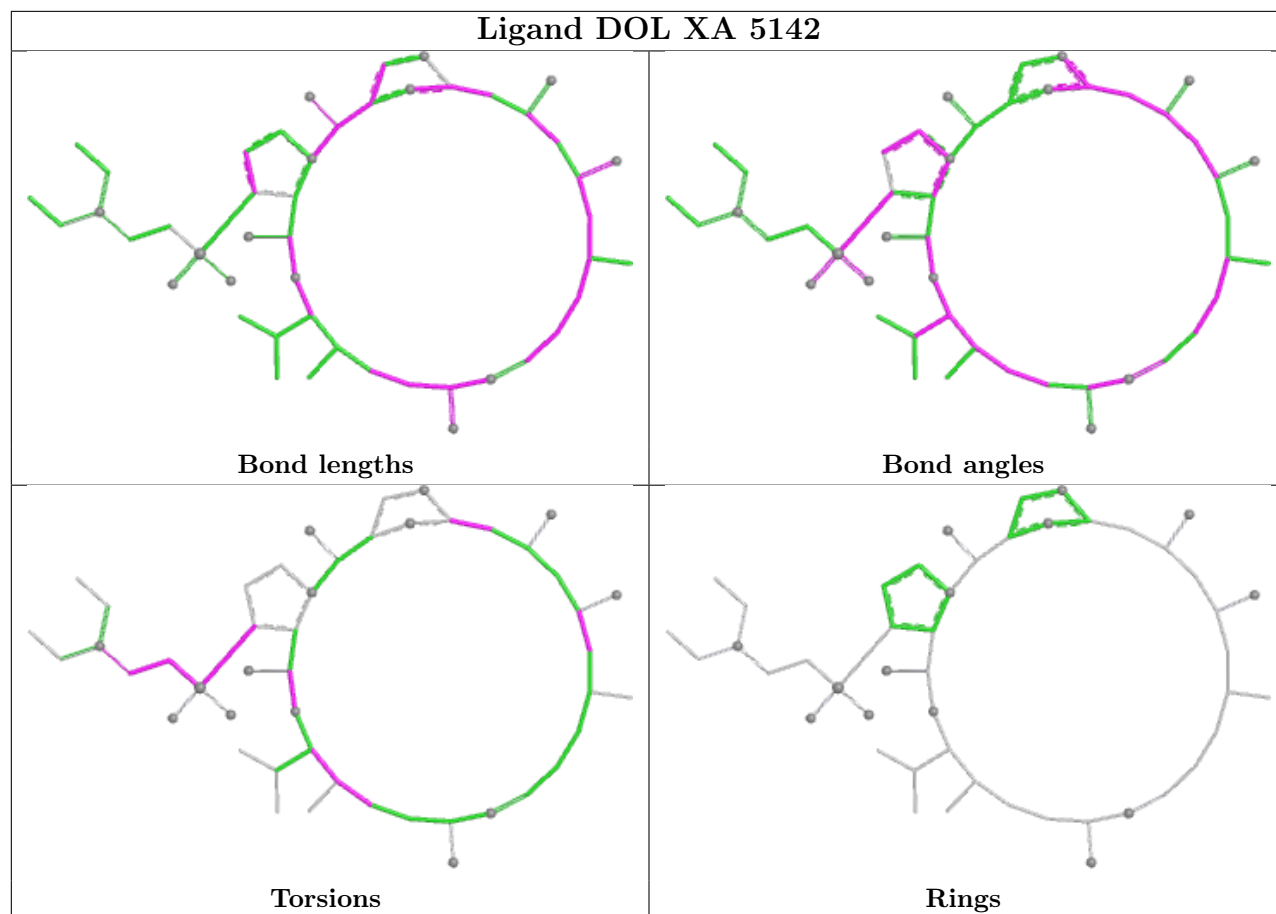
There are no ring outliers.

2 monomers are involved in 2 short contacts:

Mol	Chain	Res	Type	Clashes	Symm-Clashes
92	AX	500	GTP	1	0
91	XA	5142	DOL	1	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
17	AA	7
8	7	3
16	A4	2
40	AX	2
82	r	1
85	r3	1
38	AV	1
7	6	1

All chain breaks are listed below:

Model	Chain	Residue-1	Atom-1	Residue-2	Atom-2	Distance (Å)
1	AA	955:A	O3'	965:C	P	31.06
1	AA	734:C	O3'	740:G	P	17.04
1	AA	1403:A	O3'	1407:U	P	14.09
1	AA	1115:U	O3'	1120:C	P	10.73
1	AA	928:A	O3'	931:C	P	9.83
1	AA	1518:C	O3'	1521:U	P	8.42
1	AA	1383:A	O3'	1388:C	P	7.91
1	7	158:PHE	C	159:LYS	N	7.12
1	A4	537:ARG	C	538:ASP	N	6.17
1	7	285:ASN	C	286:LEU	N	6.07
1	r	134:ARG	C	135:LEU	N	5.77
1	r3	16:Y5P	O3'	18:P5P	P	5.69
1	AV	269:SER	C	270:PRO	N	4.66
1	7	185:LEU	C	186:ASP	N	3.47
1	AX	168:LEU	C	169:LEU	N	3.10
1	AX	195:ASN	C	196:GLU	N	3.08
1	A4	143:GLU	C	144:TYR	N	3.07
1	6	282:SER	C	283:GLU	N	3.05

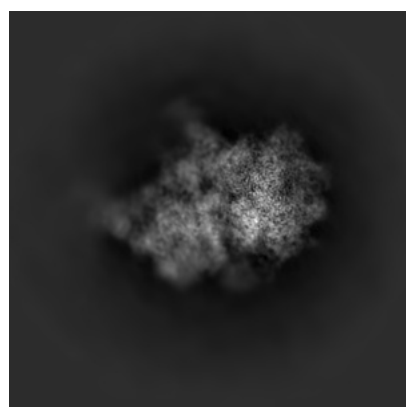
6 Map visualisation [i](#)

This section contains visualisations of the EMDB entry EMD-11395. These allow visual inspection of the internal detail of the map and identification of artifacts.

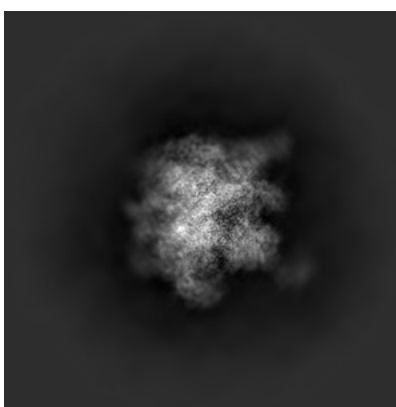
No raw map or half-maps were deposited for this entry and therefore no images, graphs, etc. pertaining to the raw map can be shown.

6.1 Orthogonal projections [i](#)

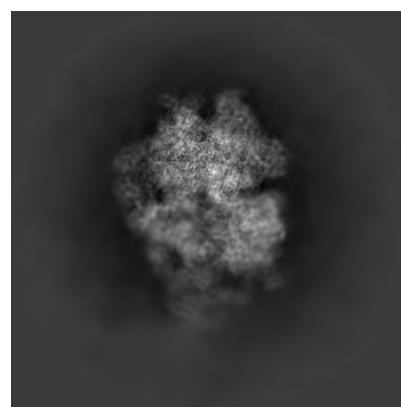
6.1.1 Primary map



X



Y

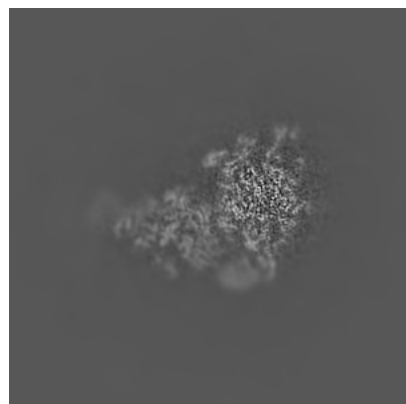


Z

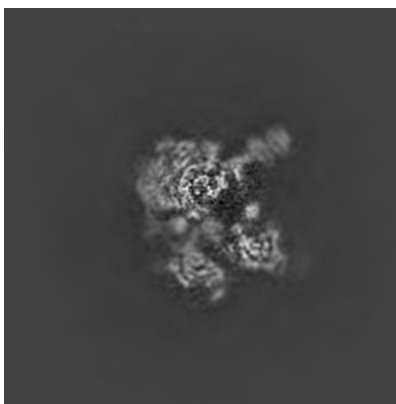
The images above show the map projected in three orthogonal directions.

6.2 Central slices [i](#)

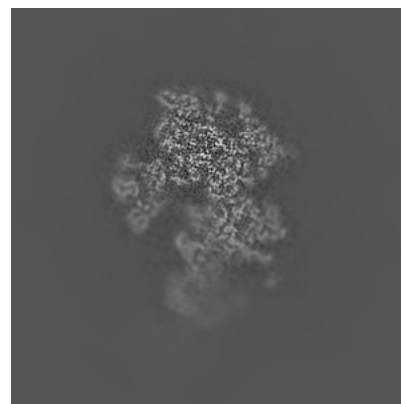
6.2.1 Primary map



X Index: 260



Y Index: 260

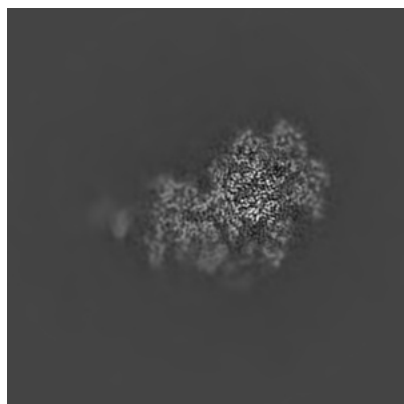


Z Index: 260

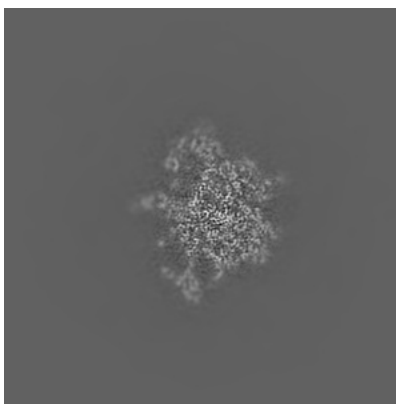
The images above show central slices of the map in three orthogonal directions.

6.3 Largest variance slices [i](#)

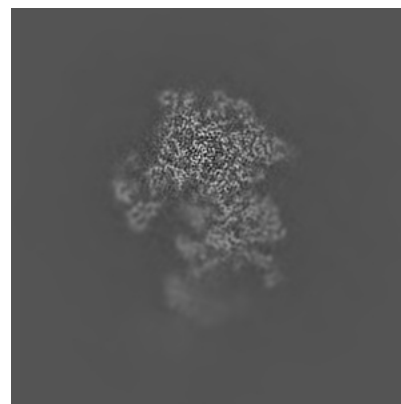
6.3.1 Primary map



X Index: 277



Y Index: 333

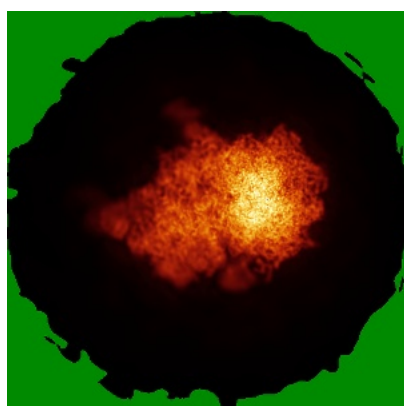


Z Index: 265

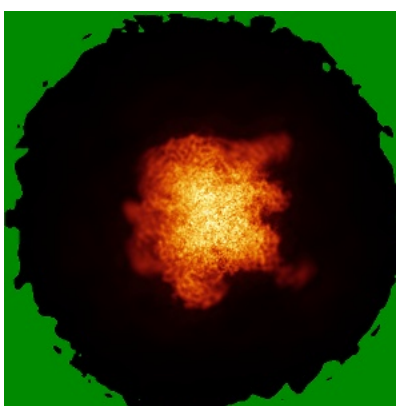
The images above show the largest variance slices of the map in three orthogonal directions.

6.4 Orthogonal standard-deviation projections (False-color) [i](#)

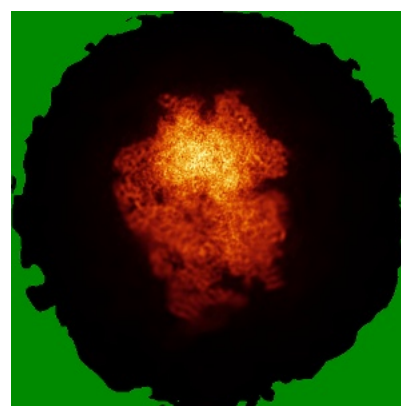
6.4.1 Primary map



X



Y



Z

The images above show the map standard deviation projections with false color in three orthogonal directions. Minimum values are shown in green, max in blue, and dark to light orange shades represent small to large values respectively.

6.5 Orthogonal surface views [i](#)

6.5.1 Primary map



The images above show the 3D surface view of the map at the recommended contour level 0.02. These images, in conjunction with the slice images, may facilitate assessment of whether an appropriate contour level has been provided.

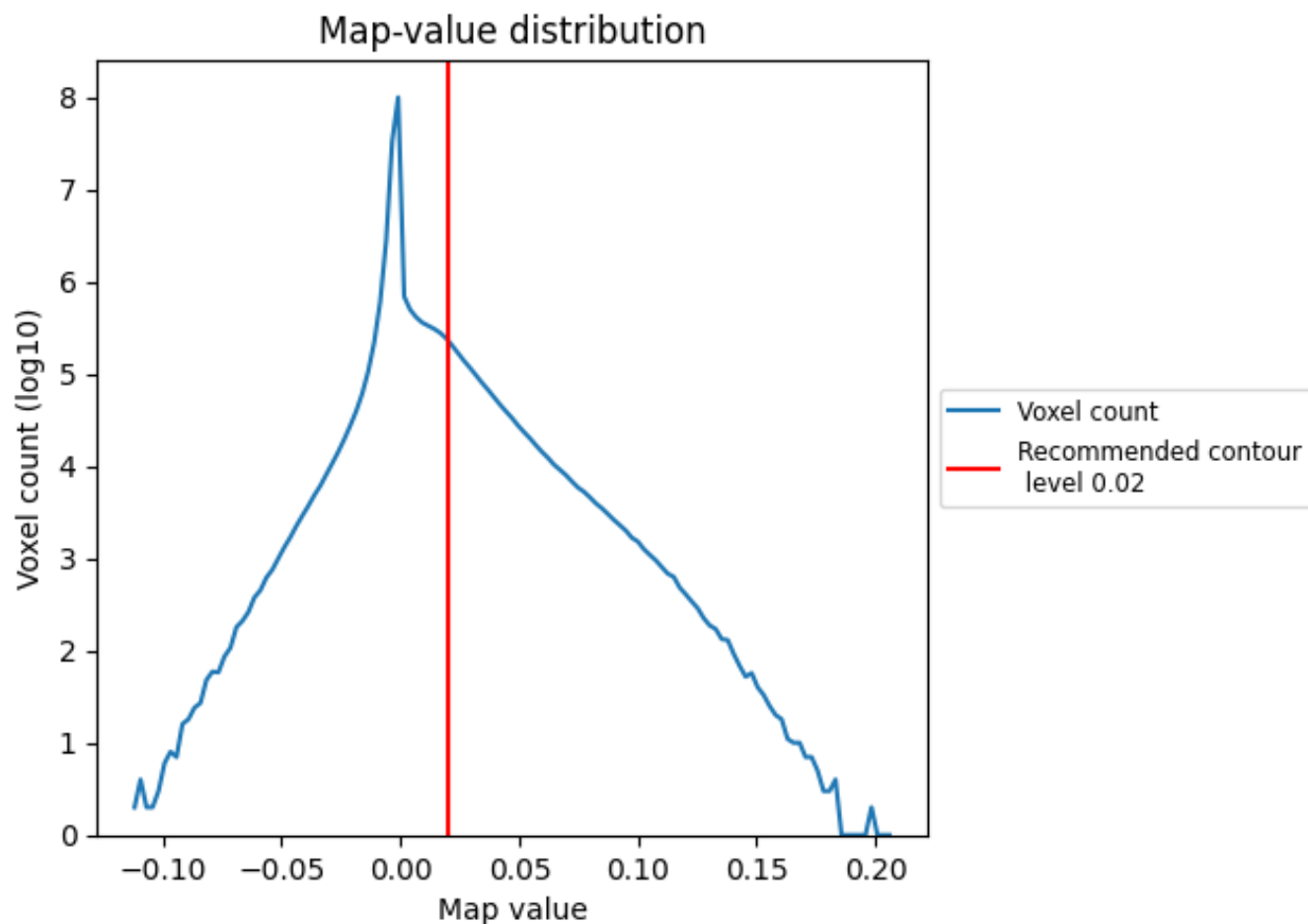
6.6 Mask visualisation [i](#)

This section was not generated. No masks/segmentation were deposited.

7 Map analysis [i](#)

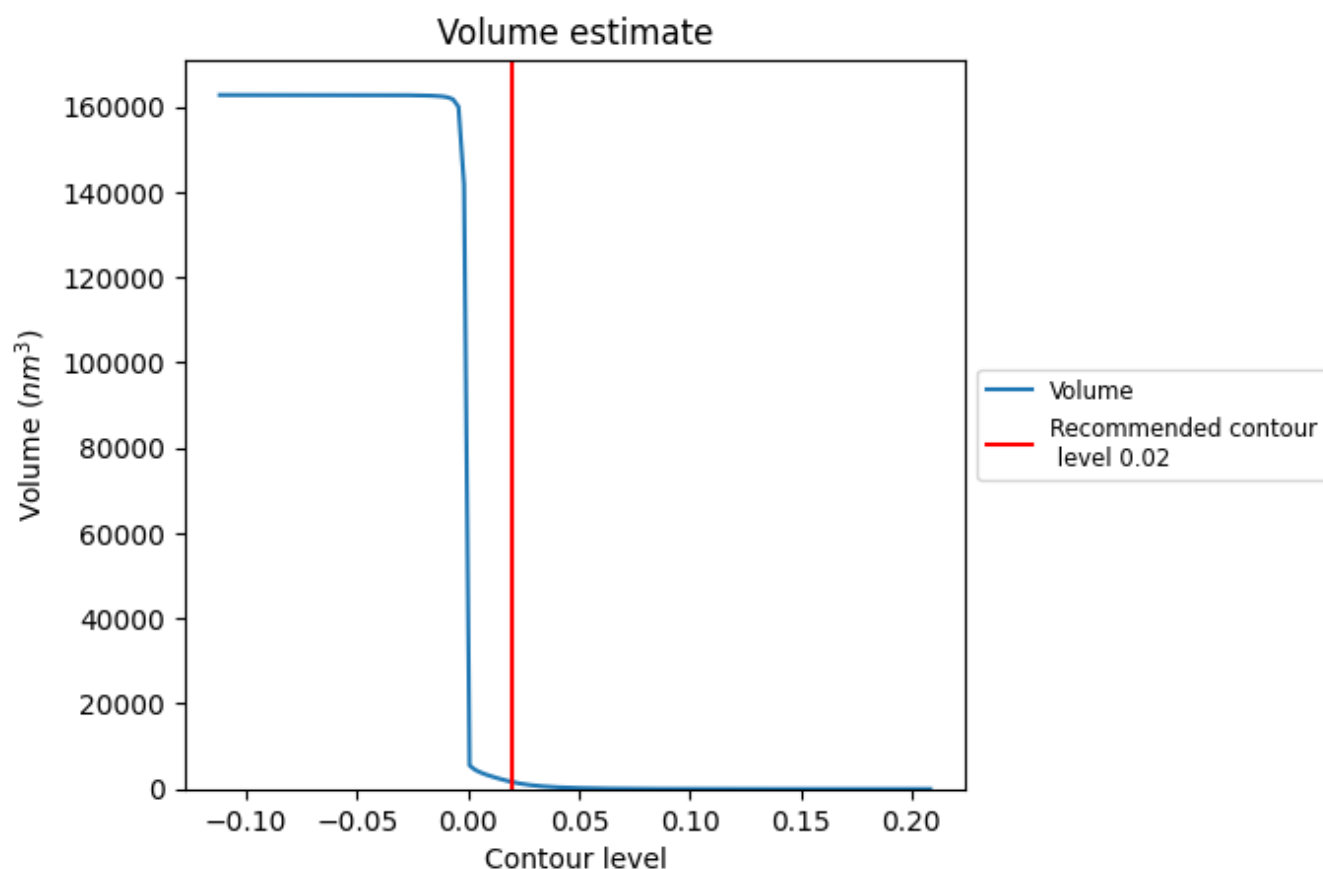
This section contains the results of statistical analysis of the map.

7.1 Map-value distribution [i](#)



The map-value distribution is plotted in 128 intervals along the x-axis. The y-axis is logarithmic. A spike in this graph at zero usually indicates that the volume has been masked.

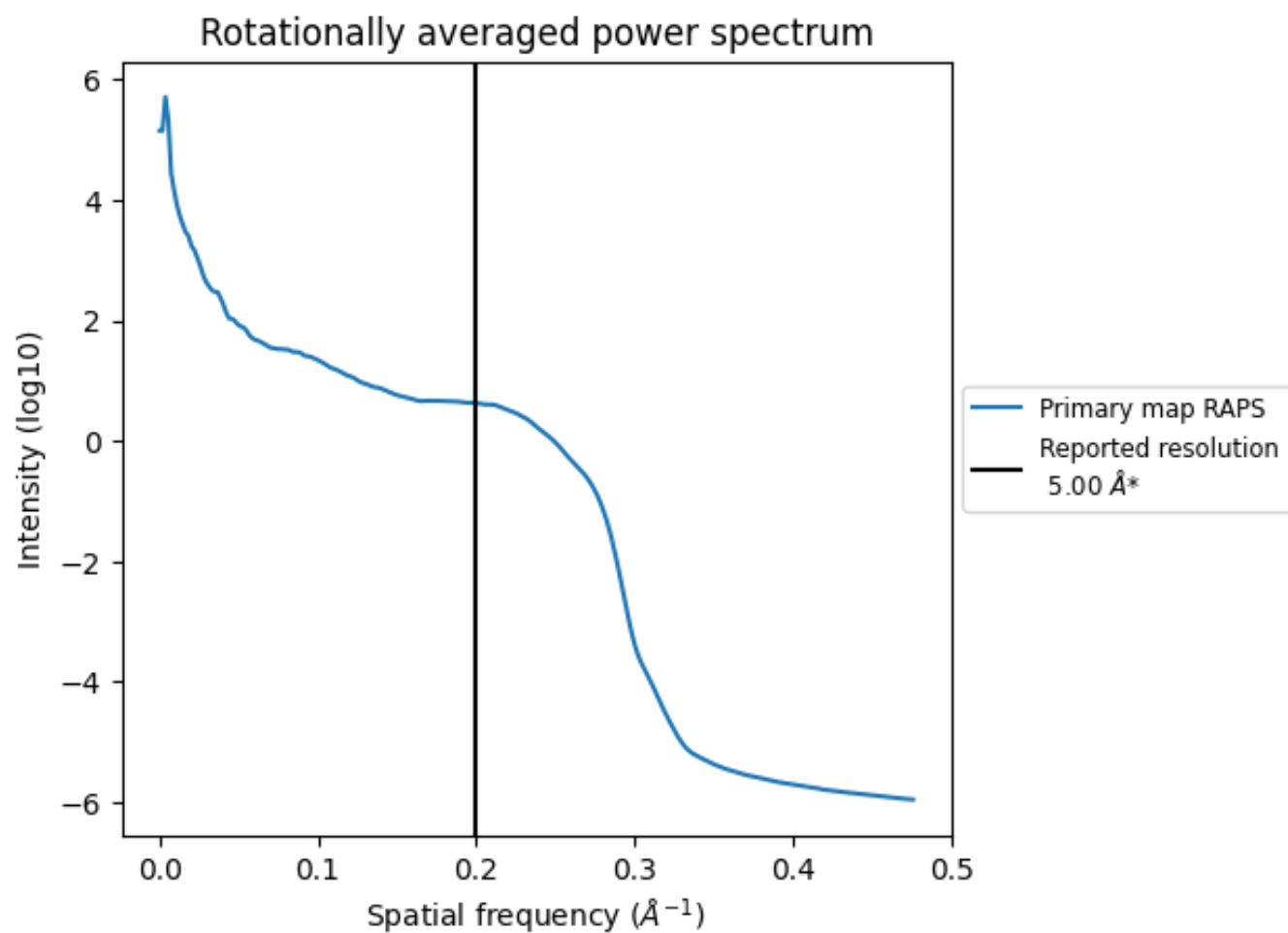
7.2 Volume estimate [i](#)



The volume at the recommended contour level is 1624 nm^3 ; this corresponds to an approximate mass of 1467 kDa.

The volume estimate graph shows how the enclosed volume varies with the contour level. The recommended contour level is shown as a vertical line and the intersection between the line and the curve gives the volume of the enclosed surface at the given level.

7.3 Rotationally averaged power spectrum ⓘ



*Reported resolution corresponds to spatial frequency of 0.200 Å⁻¹

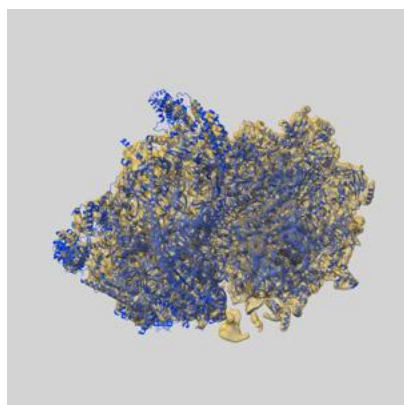
8 Fourier-Shell correlation

This section was not generated. No FSC curve or half-maps provided.

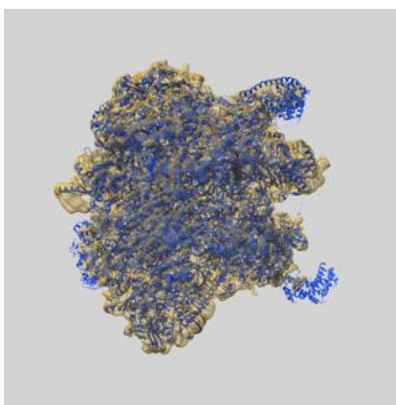
9 Map-model fit [i](#)

This section contains information regarding the fit between EMDB map EMD-11395 and PDB model 6ZSE. Per-residue inclusion information can be found in section [3](#) on page [22](#).

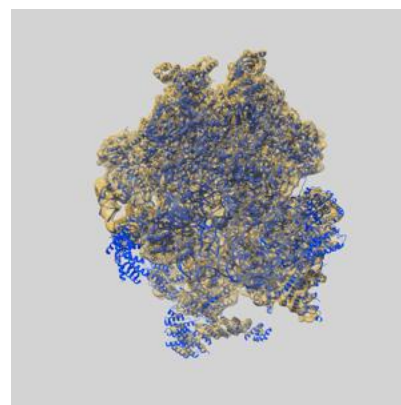
9.1 Map-model overlay [i](#)



X



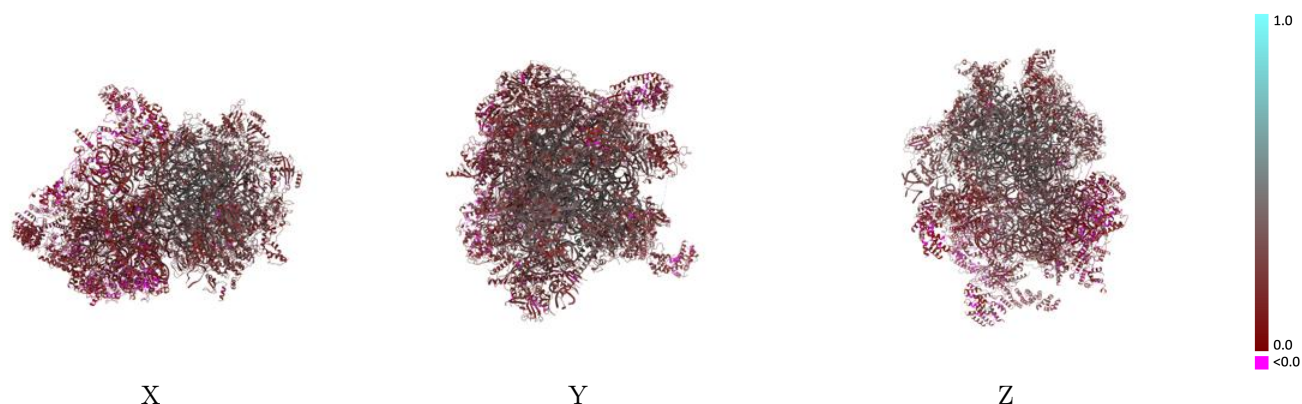
Y



Z

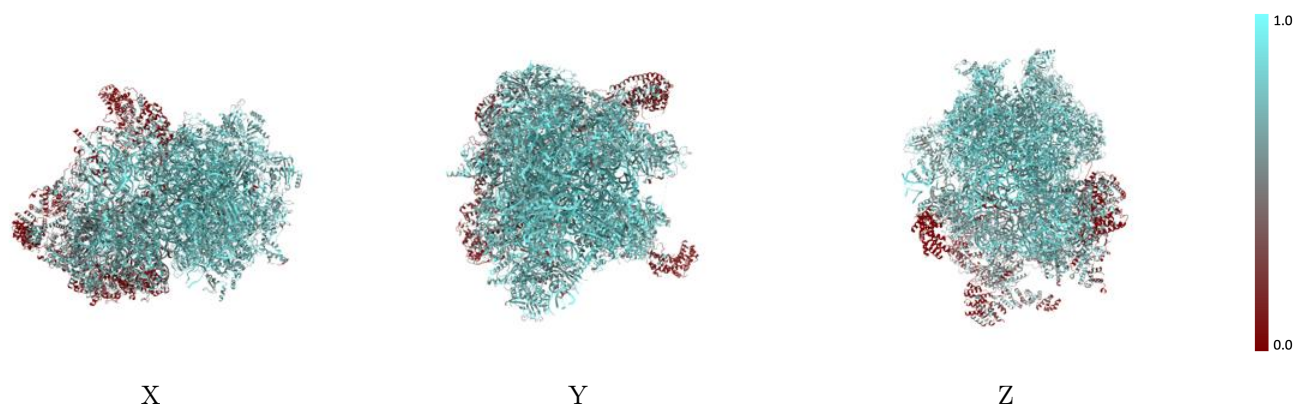
The images above show the 3D surface view of the map at the recommended contour level 0.02 at 50% transparency in yellow overlaid with a ribbon representation of the model coloured in blue. These images allow for the visual assessment of the quality of fit between the atomic model and the map.

9.2 Q-score mapped to coordinate model [i](#)



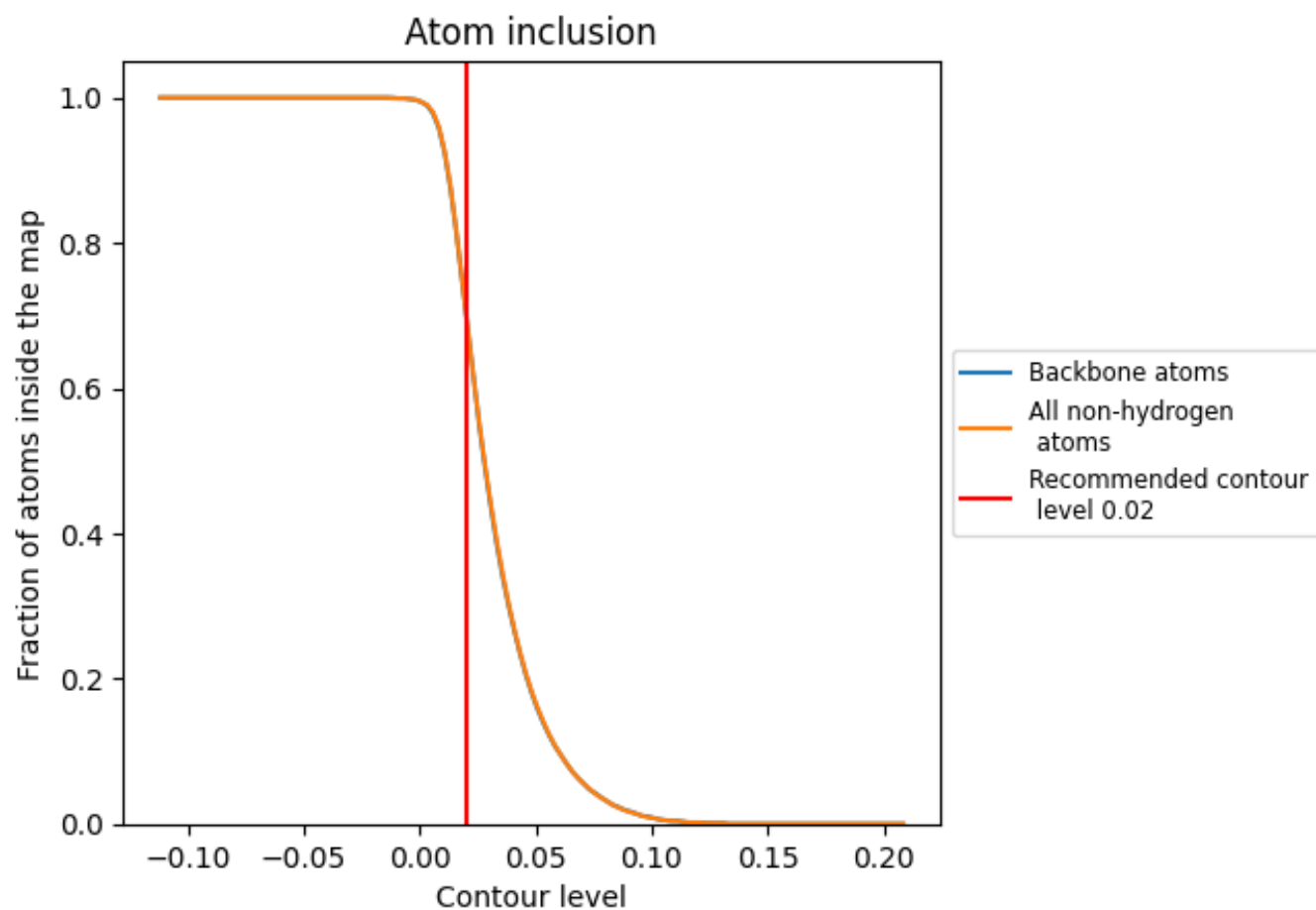
The images above show the model with each residue coloured according to its Q-score. This shows their resolvability in the map with higher Q-score values reflecting better resolvability. Please note: Q-score is calculating the resolvability of atoms, and thus high values are only expected at resolutions at which atoms can be resolved. Low Q-score values may therefore be expected for many entries.

9.3 Atom inclusion mapped to coordinate model [i](#)



The images above show the model with each residue coloured according to its atom inclusion. This shows to what extent they are inside the map at the recommended contour level (0.02).

9.4 Atom inclusion [i](#)



At the recommended contour level, 70% of all backbone atoms, 70% of all non-hydrogen atoms, are inside the map.

9.5 Map-model fit summary ⓘ

The table lists the average atom inclusion at the recommended contour level (0.02) and Q-score for the entire model and for each chain.

Chain	Atom inclusion	Q-score
All	 0.7000	 0.2530
0	 0.7850	 0.3180
1	 0.7180	 0.2780
2	 0.8440	 0.3950
3	 0.8410	 0.3950
4	 0.7940	 0.3320
5	 0.7500	 0.2530
6	 0.7330	 0.2450
7	 0.7290	 0.2560
8	 0.5410	 0.1860
9	 0.7510	 0.2860
A	 0.8080	 0.3540
A0	 0.4340	 0.1130
A1	 0.2850	 0.1000
A2	 0.3800	 0.1620
A3	 0.7050	 0.2760
A4	 0.2690	 0.1270
AA	 0.8690	 0.2190
AB	 0.5830	 0.1560
AC	 0.5160	 0.1450
AD	 0.5530	 0.1840
AE	 0.4320	 0.1380
AF	 0.3730	 0.1010
AG	 0.4930	 0.1390
AH	 0.2820	 0.0970
AI	 0.4310	 0.1040
AJ	 0.6710	 0.2200
AK	 0.5100	 0.0690
AL	 0.5170	 0.1400
AM	 0.5000	 0.1140
AN	 0.5760	 0.1320
AO	 0.5710	 0.1650
AP	 0.4550	 0.1270
AQ	 0.5140	 0.1640
AR	 0.4700	 0.1250



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Chain	Atom inclusion	Q-score
AS	 0.5010	 0.1550
AT	 0.5910	 0.1590
AU	 0.5200	 0.1510
AV	 0.2660	 0.1200
AW	 0.4760	 0.1330
AX	 0.1910	 0.0640
AY	 0.2400	 0.0880
AZ	 0.4090	 0.1160
XA	 0.9340	 0.3770
XB	 0.9410	 0.2600
XD	 0.8010	 0.3150
XE	 0.8000	 0.3390
XF	 0.8350	 0.3660
XH	 0.7270	 0.2790
XI	 0.5390	 0.1960
XJ	 0.6010	 0.1650
XK	 0.8350	 0.3660
XL	 0.8080	 0.3560
XM	 0.8020	 0.3440
XN	 0.7780	 0.3330
XO	 0.7900	 0.3170
XP	 0.7410	 0.2520
XQ	 0.7130	 0.2890
XR	 0.8340	 0.3650
XS	 0.7860	 0.3580
XT	 0.8290	 0.3730
XU	 0.7950	 0.3260
XV	 0.7630	 0.2730
XW	 0.8170	 0.3620
XX	 0.7610	 0.2840
XY	 0.7940	 0.2940
XZ	 0.8260	 0.3640
a	 0.7450	 0.3170
b	 0.8410	 0.3550
c	 0.7630	 0.2800
d	 0.6810	 0.2310
e	 0.5970	 0.1610
f	 0.6050	 0.1920
g	 0.8090	 0.3390
h	 0.7230	 0.2650
i	 0.8450	 0.3800
j	 0.8030	 0.3350

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Chain	Atom inclusion	Q-score
k	 0.7000	 0.2050
l	 0.6420	 0.1990
m	 0.6930	 0.1780
o	 0.8530	 0.3760
p	 0.7210	 0.2470
q	 0.6190	 0.2310
r	 0.7920	 0.2920
r1	 0.6270	 0.2510
r2	 0.5370	 0.1930
r3	 0.6720	 0.1550
s	 0.7650	 0.2790
t1	 0.1730	 0.1940
t2	 0.2230	 0.1700
t3	 0.0000	 0.1470
t4	 0.0000	 0.1300
t5	 0.0000	 0.1430
t6	 0.0000	 0.0830