



wwPDB EM Map/Model Validation Report ⓘ

Apr 10, 2016 – 02:34 PM BST

PDB ID : 3J9W
EMDB ID: : EMD-6306
Title : Cryo-EM structure of the Bacillus subtilis MifM-stalled ribosome complex
Authors : Sohmen, D.; Chiba, S.; Shimokawa-Chiba, N.; Innis, C.A.; Berninghausen, O.; Beckmann, R.; Ito, K.; Wilson, D.N.
Deposited on : 2015-03-16
Resolution : 3.90 Å(reported)

This is a wwPDB EM Map/Model Validation Report for a publicly released PDB/EMDB entry.
For rigid body fitted models, validation errors reported here could stem from errors in the original structure(s) used in the fitting.
We welcome your comments at validation@mail.wwpdb.org
A user guide is available at
<http://wwpdb.org/validation/2016/EMValidationReportHelp>

MolProbity : 4.02b-467
Mogul : unknown
Percentile statistics : 20151230.v01 (using entries in the PDB archive December 30th 2015)
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et. al. (1996)
Validation Pipeline (wwPDB-VP) : trunk27241

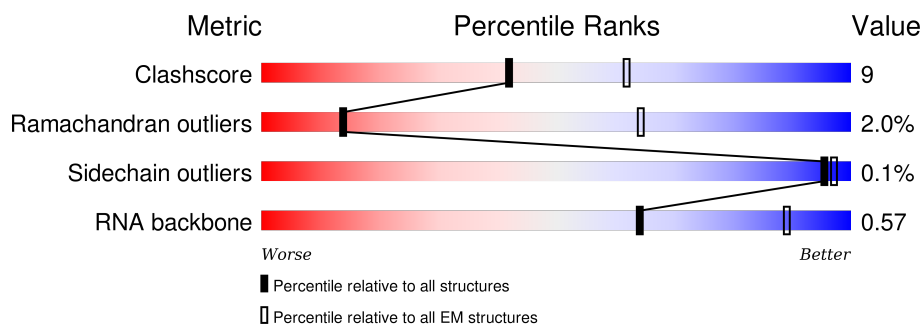
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

ELECTRON MICROSCOPY

The reported resolution of this entry is 3.90 Å.

Percentile scores (ranging between 0-100) for global validation metrics of the entry are shown in the following graphic. The table shows the number of entries on which the scores are based.




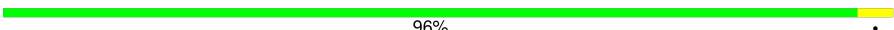








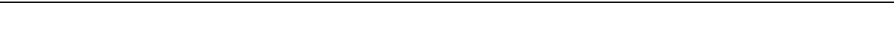

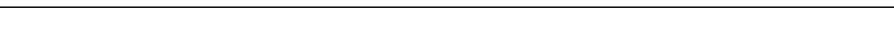
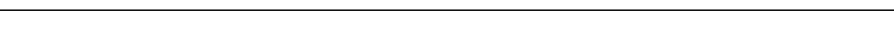











Metric	Whole archive (#Entries)	EM structures (#Entries)
Clashscore	114402	924
Ramachandran outliers	111179	726
Sidechain outliers	111093	686
RNA backbone	3027	244

The table below summarises the geometric issues observed across the polymeric chains. The red, orange, yellow and green segments on the bar indicate the fraction of residues that contain outliers for ≥ 3 , 2, 1 and 0 types of geometric quality criteria. A grey segment represents the fraction of residues that are not modelled. The numeric value for each fraction is indicated below the corresponding segment, with a dot representing fractions $\leq 5\%$.

Mol	Chain	Length	Quality of chain
1	AA	1555	51% 33% 12% ..
2	AB	246	88% . 9%
3	AC	218	94% . .
4	AD	200	97% ..
5	AE	166	96% . .
6	AF	95	98% .
7	AG	156	96% . .
8	AH	132	98% ..









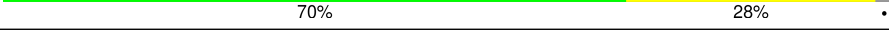
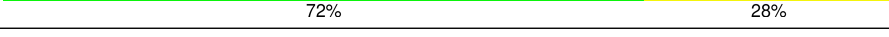
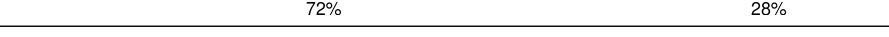

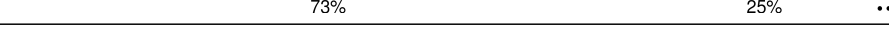
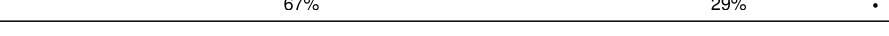
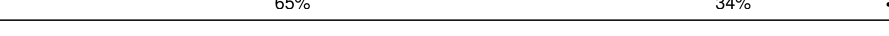


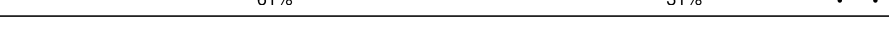
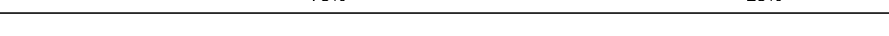


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Mol	Chain	Length	Quality of chain
9	AI	130	 93% 5% .
10	AJ	102	 96% .
11	AK	131	 85% 5% 10%
12	AL	138	 93% 7% .
13	AM	121	 89% 7% . .
14	AN	61	 90% 7% . .
15	AO	89	 97% . .
16	AP	90	 97% . .
17	AQ	87	 97% . .
18	AR	79	 85% . . 10%
19	AS	92	 87% . 9%
20	AT	88	 92% 6% .
21	AX	77	 57% 18% 22% .
22	AY	19	 11% 58% 32%
23	AZ	95	 15% 8% . 75%
24	B0	62	 56% 34% . 6%
25	B1	66	 71% 27% .
26	B2	59	 71% 27% .
27	B3	66	 70% 26% . .
28	B4	59	 66% 25% 8%
29	B5	49	 55% 43% .
30	B6	44	 80% 20%
31	B7	66	 47% 48% . .
32	B8	37	 84% 14% .
33	BA	2928	 44% 33% 18% 6%

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Mol	Chain	Length	Quality of chain
34	BB	119	
35	BD	277	
36	BE	209	
37	BF	207	
38	BG	179	
39	BH	179	
40	BJ	166	
41	BK	141	
42	BM	145	
43	BN	122	
44	BO	146	
45	BP	144	
46	BQ	120	
47	BR	120	
48	BS	115	
49	BT	119	
50	BU	102	
51	BV	113	
52	BW	95	
53	BX	103	
54	BZ	94	

2 Entry composition

There are 54 unique types of molecules in this entry. The entry contains 135425 atoms, of which 0 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 RNA chain called 16S ribosomal RNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
1	AA	1544	Total	C	N	O	P	0	0
			33115	14768	6067	10736	1544		

- Molecule 2 is a protein called 30S ribosomal protein uS2.

Mol	Chain	Residues	Atoms				AltConf	Trace
2	AB	224	Total	C	N	O	0	0
			896	448	224	224		

- Molecule 3 is a protein called 30S ribosomal protein uS3.

Mol	Chain	Residues	Atoms				AltConf	Trace
3	AC	210	Total	C	N	O	0	0
			840	420	210	210		

- Molecule 4 is a protein called 30S ribosomal protein uS4.

Mol	Chain	Residues	Atoms				AltConf	Trace
4	AD	199	Total	C	N	O	0	0
			797	398	199	200		

- Molecule 5 is a protein called 30S ribosomal protein uS5.

Mol	Chain	Residues	Atoms				AltConf	Trace
5	AE	165	Total	C	N	O	0	0
			661	330	165	166		

- Molecule 6 is a protein called 30S ribosomal protein bS6.

Mol	Chain	Residues	Atoms				AltConf	Trace
6	AF	95	Total	C	N	O	0	0
			381	190	95	96		

- Molecule 7 is a protein called 30S ribosomal protein uS7.

Mol	Chain	Residues	Atoms				AltConf	Trace
7	AG	153	Total	C	N	O	0	0
			613	306	153	154		

- Molecule 8 is a protein called 30S ribosomal protein uS8.

Mol	Chain	Residues	Atoms				AltConf	Trace
8	AH	131	Total	C	N	O	0	0
			525	262	131	132		

- Molecule 9 is a protein called 30S ribosomal protein uS9.

Mol	Chain	Residues	Atoms				AltConf	Trace
9	AI	130	Total	C	N	O	0	0
			521	260	130	131		

- Molecule 10 is a protein called 30S ribosomal protein uS10.

Mol	Chain	Residues	Atoms				AltConf	Trace
10	AJ	102	Total	C	N	O	0	0
			409	204	102	103		

- Molecule 11 is a protein called 30S ribosomal protein uS11.

Mol	Chain	Residues	Atoms				AltConf	Trace
11	AK	118	Total	C	N	O	0	0
			472	236	118	118		

- Molecule 12 is a protein called 30S ribosomal protein uS12.

Mol	Chain	Residues	Atoms				AltConf	Trace
12	AL	137	Total	C	N	O	0	0
			549	274	137	138		

- Molecule 13 is a protein called 30S ribosomal protein uS13.

Mol	Chain	Residues	Atoms				AltConf	Trace
13	AM	119	Total	C	N	O	0	0
			476	238	119	119		

- Molecule 14 is a protein called 30S ribosomal protein uS14.

Mol	Chain	Residues	Atoms				AltConf	Trace
14	AN	60	Total	C	N	O	0	0
			241	120	60	61		

- Molecule 15 is a protein called 30S ribosomal protein uS15.

Mol	Chain	Residues	Atoms				AltConf	Trace
15	AO	88	Total	C	N	O	0	0
			353	176	88	89		

- Molecule 16 is a protein called 30S ribosomal protein bS16.

Mol	Chain	Residues	Atoms				AltConf	Trace
16	AP	89	Total	C	N	O	0	0
			357	178	89	90		

- Molecule 17 is a protein called 30S ribosomal protein uS17.

Mol	Chain	Residues	Atoms				AltConf	Trace
17	AQ	86	Total	C	N	O	0	0
			345	172	86	87		

- Molecule 18 is a protein called 30S ribosomal protein bS18.

Mol	Chain	Residues	Atoms				AltConf	Trace
18	AR	71	Total	C	N	O	0	0
			285	142	71	72		

- Molecule 19 is a protein called 30S ribosomal protein uS19.

Mol	Chain	Residues	Atoms				AltConf	Trace
19	AS	84	Total	C	N	O	0	0
			336	168	84	84		

- Molecule 20 is a protein called 30S ribosomal protein bS20.

Mol	Chain	Residues	Atoms				AltConf	Trace
20	AT	86	Total	C	N	O	0	0
			345	172	86	87		

- Molecule 21 is a RNA chain called tRNA-Asp.

Mol	Chain	Residues	Atoms					AltConf	Trace
21	AX	77	Total	C	N	O	P	0	0
			1643	731	290	545	77		

- Molecule 22 is a RNA chain called mRNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
22	AY	19	Total	C	N	O	P	0	0
			415	185	82	129	19		

- Molecule 23 is a protein called MifM.

Mol	Chain	Residues	Atoms					AltConf	Trace
23	AZ	24	Total	C	N	O	S	0	0
			107	64	28	14	1		

- Molecule 24 is a protein called 50S ribosomal protein bL28.

Mol	Chain	Residues	Atoms					AltConf	Trace
24	B0	58	Total	C	N	O	S	0	0
			444	275	92	75	2		

- Molecule 25 is a protein called 50S ribosomal protein uL29.

Mol	Chain	Residues	Atoms					AltConf	Trace
25	B1	65	Total	C	N	O	S	0	0
			530	328	102	98	2		

- Molecule 26 is a protein called 50S ribosomal protein uL30.

Mol	Chain	Residues	Atoms					AltConf	Trace
26	B2	58	Total	C	N	O	S	0	0
			455	281	89	84	1		

- Molecule 27 is a protein called 50S ribosomal protein bL31.

Mol	Chain	Residues	Atoms					AltConf	Trace
27	B3	64	Total	C	N	O	S	0	0
			503	314	92	92	5		

- Molecule 28 is a protein called 50S ribosomal protein bL32.

Mol	Chain	Residues	Atoms					AltConf	Trace
28	B4	54	Total	C	N	O	S	0	0
			426	262	86	71	7		

- Molecule 29 is a protein called 50S ribosomal protein bL33.

Mol	Chain	Residues	Atoms					AltConf	Trace
29	B5	48	Total	C	N	O	S	0	0
			401	244	80	73	4		

- Molecule 30 is a protein called 50S ribosomal protein bL34.

Mol	Chain	Residues	Atoms					AltConf	Trace
30	B6	44	Total	C	N	O	S	0	0
			367	222	89	54	2		

- Molecule 31 is a protein called 50S ribosomal protein bL35.

Mol	Chain	Residues	Atoms					AltConf	Trace
31	B7	64	Total	C	N	O	S	0	0
			512	321	107	82	2		

- Molecule 32 is a protein called 50S ribosomal protein bL36.

Mol	Chain	Residues	Atoms					AltConf	Trace
32	B8	36	Total	C	N	O	S	0	0
			288	181	59	44	4		

- Molecule 33 is a RNA chain called 23S ribosomal RNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
33	BA	2923	Total	C	N	O	P	0	0
			62767	28002	11589	20253	2923		

- Molecule 34 is a RNA chain called 5S ribosomal RNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
34	BB	112	Total	C	N	O	P	0	0
			2395	1068	435	780	112		

- Molecule 35 is a protein called 50S ribosomal protein uL2.

Mol	Chain	Residues	Atoms					AltConf	Trace
35	BD	275	Total	C	N	O	S	0	0
			2111	1312	416	377	6		

- Molecule 36 is a protein called 50S ribosomal protein uL3.

Mol	Chain	Residues	Atoms					AltConf	Trace
36	BE	207	Total	C	N	O	S	0	0
			1575	988	290	292	5		

- Molecule 37 is a protein called 50S ribosomal protein uL4.

Mol	Chain	Residues	Atoms					AltConf	Trace
37	BF	205	Total	C	N	O	S	0	0
			1561	980	289	290	2		

- Molecule 38 is a protein called 50S ribosomal protein uL5.

Mol	Chain	Residues	Atoms					AltConf	Trace
38	BG	178	Total	C	N	O	S	0	0
			1404	893	245	259	7		

- Molecule 39 is a protein called 50S ribosomal protein uL6.

Mol	Chain	Residues	Atoms					AltConf	Trace
39	BH	175	Total	C	N	O	S	0	0
			1342	835	248	257	2		

- Molecule 40 is a protein called 50S ribosomal protein uL10.

Mol	Chain	Residues	Atoms					AltConf	Trace
40	BJ	123	Total	C	N	O	S	0	0
			955	602	163	189	1		

- Molecule 41 is a protein called 50S ribosomal protein uL11.

Mol	Chain	Residues	Atoms					AltConf	Trace
41	BK	133	Total	C	N	O	S	0	0
			981	617	173	185	6		

- Molecule 42 is a protein called 50S ribosomal protein uL13.

Mol	Chain	Residues	Atoms					AltConf	Trace
42	BM	142	Total	C	N	O	S	0	0
			1123	710	206	202	5		

- Molecule 43 is a protein called 50S ribosomal protein uL14.

Mol	Chain	Residues	Atoms					AltConf	Trace
43	BN	122	Total	C	N	O	S	0	0
			920	571	173	172	4		

- Molecule 44 is a protein called 50S ribosomal protein uL15.

Mol	Chain	Residues	Atoms					AltConf	Trace
44	BO	146	Total	C	N	O	S	0	0
			1081	671	207	201	2		

- Molecule 45 is a protein called 50S ribosomal protein uL16.

Mol	Chain	Residues	Atoms					AltConf	Trace
45	BP	138	Total	C	N	O	S	0	0
			1097	703	208	181	5		

- Molecule 46 is a protein called 50S ribosomal protein bL17.

Mol	Chain	Residues	Atoms					AltConf	Trace
46	BQ	119	Total	C	N	O	S	0	0
			953	583	186	180	4		

- Molecule 47 is a protein called 50S ribosomal protein uL18.

Mol	Chain	Residues	Atoms					AltConf	Trace
47	BR	120	Total	C	N	O	S	0	0
			912	564	176	171	1		

- Molecule 48 is a protein called 50S ribosomal protein bL19.

Mol	Chain	Residues	Atoms					AltConf	Trace
48	BS	114	Total	C	N	O	S	0	0
			936	595	184	157			

- Molecule 49 is a protein called 50S ribosomal protein bL20.

Mol	Chain	Residues	Atoms					AltConf	Trace
49	BT	117	Total	C	N	O	S	0	0
			940	591	189	156	4		

- Molecule 50 is a protein called 50S ribosomal protein bL21.

Mol	Chain	Residues	Atoms					AltConf	Trace
50	BU	101	Total	C	N	O	S	0	0
			786	501	139	146			

- Molecule 51 is a protein called 50S ribosomal protein uL22.

Mol	Chain	Residues	Atoms					AltConf	Trace
51	BV	109	Total	C	N	O	S	0	0
			842	525	164	150	3		

- Molecule 52 is a protein called 50S ribosomal protein uL23.

Mol	Chain	Residues	Atoms					AltConf	Trace
52	BW	93	Total	C	N	O	S	0	0
			752	472	137	139	4		

- Molecule 53 is a protein called 50S ribosomal protein uL24.

Mol	Chain	Residues	Atoms					AltConf	Trace
53	BX	100	Total	C	N	O	S	0	0
			754	473	141	137	3		

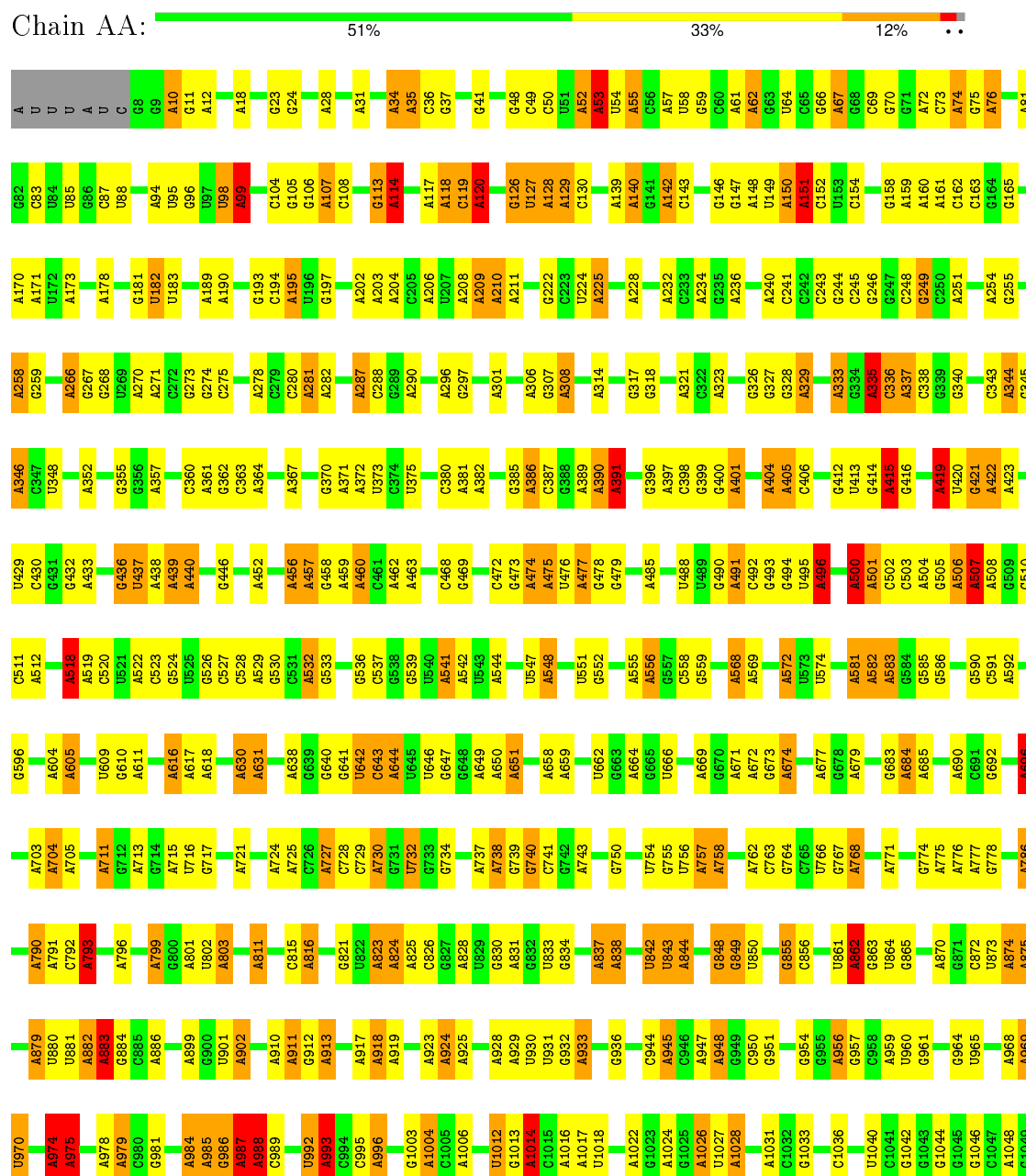
- Molecule 54 is a protein called 50S ribosomal protein bL27.

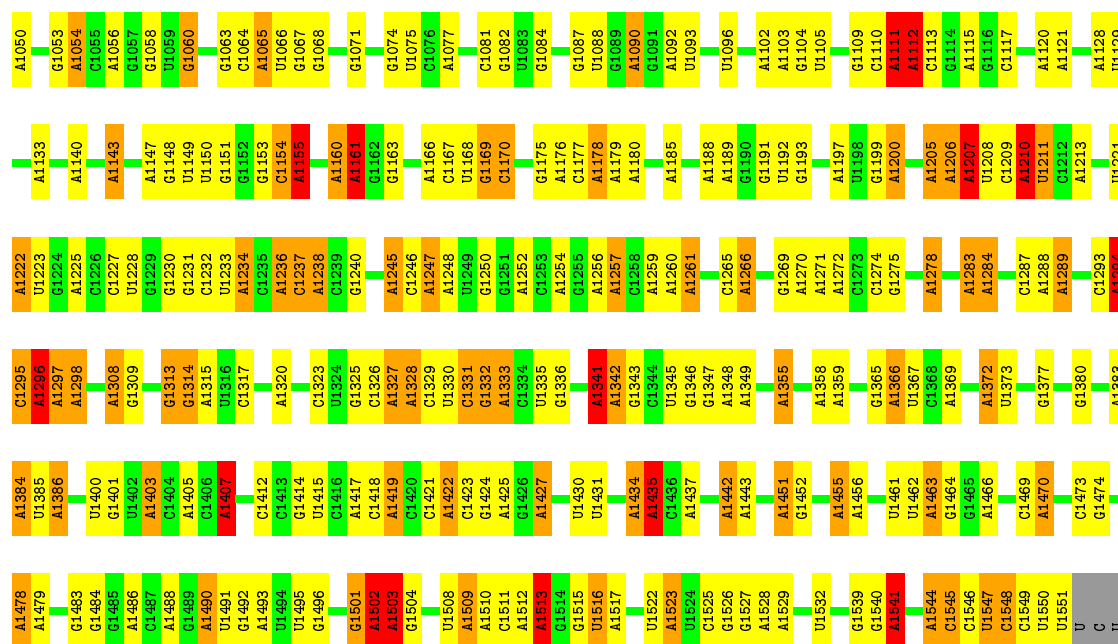
Mol	Chain	Residues	Atoms					AltConf	Trace
54	BZ	82	Total	C	N	O	S	0	0
			630	390	123	117			

3 Residue-property plots

These plots are drawn for all protein, RNA and DNA chains in the entry. The first graphic for a chain summarises the proportions of errors displayed in the second graphic. The second graphic shows the sequence view annotated by issues in geometry. Residues are color-coded according to the number of geometric quality criteria for which they contain at least one outlier: green = 0, yellow = 1, orange = 2 and red = 3 or more. Stretches of 2 or more consecutive residues without any outlier are shown as a green connector. Residues present in the sample, but not in the model, are shown in grey.

- Molecule 1: 16S ribosomal RNA





A

- Molecule 2: 30S ribosomal protein uS2

Chain AB: 88% 9%



- Molecule 3: 30S ribosomal protein uS3

Chain AC: 94%



- Molecule 4: 30S ribosomal protein uS4

Chain AD: 97%



- Molecule 5: 30S ribosomal protein uS5

Chain AE: 96%



- Molecule 6: 30S ribosomal protein bS6

Chain AF:  98% .



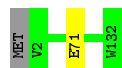
- Molecule 7: 30S ribosomal protein uS7

Chain AG:  96% ..



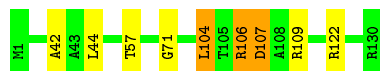
- Molecule 8: 30S ribosomal protein uS8

Chain AH:  98% ..



- Molecule 9: 30S ribosomal protein uS9

Chain AI:  93% 5% .




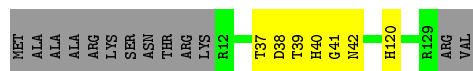
- Molecule 10: 30S ribosomal protein uS10

Chain AJ:  96% .



- Molecule 11: 30S ribosomal protein uS11

Chain AK:  85% 5% 10%



- Molecule 12: 30S ribosomal protein uS12

Chain AL:  93% 7% .



- Molecule 13: 30S ribosomal protein uS13

Chain AM:  89% 7% ..



- Molecule 14: 30S ribosomal protein uS14

Chain AN: 90% 7% ..



- Molecule 15: 30S ribosomal protein uS15

Chain AO: 97% ..



- Molecule 16: 30S ribosomal protein bS16

Chain AP: 97% ..



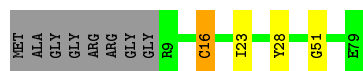
- Molecule 17: 30S ribosomal protein uS17

Chain AQ: 97% ..



- Molecule 18: 30S ribosomal protein bS18

Chain AR: 85% 10% ..



- Molecule 19: 30S ribosomal protein uS19

Chain AS: 87% 9% ..

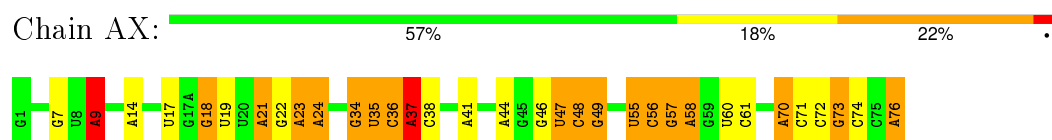


- Molecule 20: 30S ribosomal protein bS20

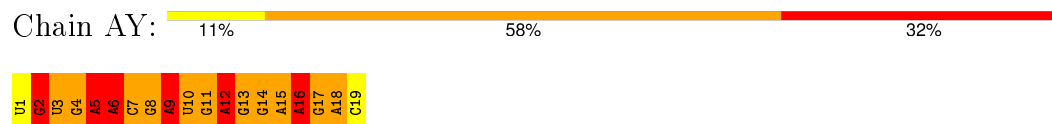
Chain AT: 92% 6% ..



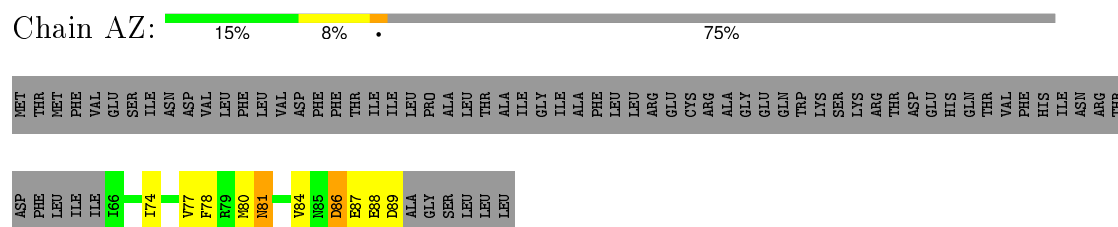
- Molecule 21: tRNA-Asp



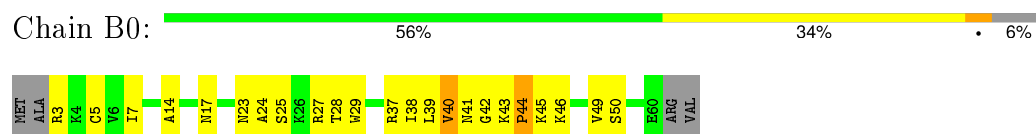
- Molecule 22: mRNA



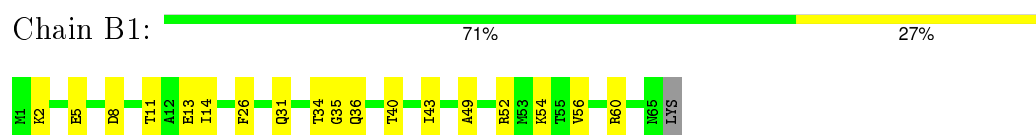
- Molecule 23: MifM



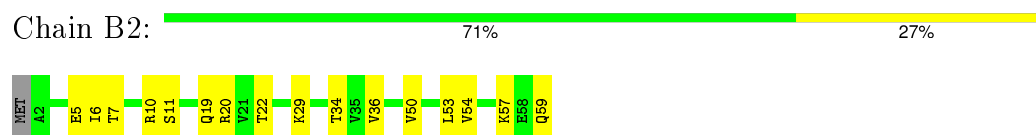
- Molecule 24: 50S ribosomal protein bL28



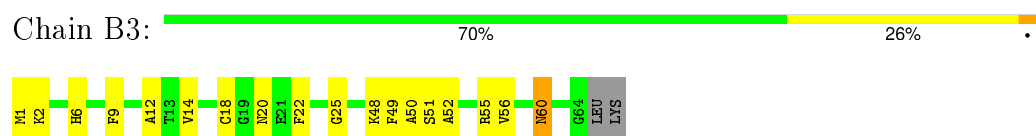
- Molecule 25: 50S ribosomal protein uL29



- Molecule 26: 50S ribosomal protein uL30



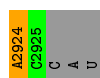
- Molecule 27: 50S ribosomal protein bL31



- Molecule 28: 50S ribosomal protein bL32

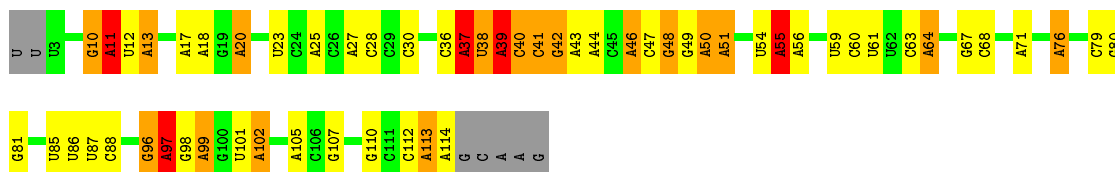
A1555	C1474	G1309	G1284	G1145	A1078	A1005	C930	C959	C785	A702	C480	C387
A1556	A1392	A1312	A1235	A1148	U1079	A1006	C931	U960	A786	G703	U481	A388
C1558	G1394	A1313	G1236	A1149	A1084	A1008	C932	C961	C787	U704	U485	A389
G1561	G1395	A1314	U1242	C1150	U1087	U1013	U934	G865	C789	A705	A486	A390
A1562	C1396	G1315	A1243	U1151	G1088	A1014	C936	A867	C791	G709	G489	A391
G1563	G1397	A1316	A1244	G1152	G1089	G1015	C937	A868	G792	G710	A490	C392
U1483	A1398	G1317	G1245	G1153	A1092	U1016	C938	U869	U793	U711	U494	C393
U1484	G1318	G1246	G1247	G1157	A1093	C1017	C939	U870	U794	C712	U495	U394
A1485	G1319	G1248	C1249	G1158	A1094	G1018	C940	A871	U795	U713	A496	U405
A1490	G1322	U1249	U1250	A1161	G1095	A1019	U942	C572	A796	A715	A497	G406
A1491	A1323	A1405	G1251	A1172	A1096	A1020	U943	G873	A797	A716	A498	A407
C1573	G1407	A1406	U1252	A1173	A1097	A1021	C944	U874	A798	C718	U499	G411
A1574	G1408	G1408	G1253	A1174	A1098	G1022	C945	U875	A799	C719	G499	G412
U1575	U1411	A1326	A1254	A1175	A1099	G1023	C946	U876	G800	C720	A500	A412
G1578	A1412	U1327	G1255	U1176	G1101	A1024	A947	G877	G804	A721	A501	A418
A1579	A1413	A1335	C1256	U1177	G1102	A1025	U948	G878	G805	A722	C502	A419
A1580	A1414	A1336	C1257	G1178	A1103	A1026	A952	G879	A808	A723	C503	U420
U1581	U1417	A1339	A1258	U1179	U1104	G1027	C953	C880	A809	A724	A504	A421
A1583	U1418	A1340	G1259	U1180	G1105	C1028	A956	A882	A811	C725	A507	G426
U1584	A1421	U1341	A1260	C1181	U1106	G1030	A957	A888	G812	A726	C508	G427
A1585	G1422	A1342	C1261	U1182	U1107	A1031	A958	A889	G821	G728	C509	A428
U1586	A1423	C1344	G1262	U1183	U1108	G1032	C959	U892	A821	U729	U510	A429
U1587	A1424	U1345	G1263	U1184	G1109	A1033	U966	G895	G822	G731	G512	C430
U1588	C1425	A1346	G1264	U1185	G1110	G1034	A970	G897	G823	U732	G513	A431
U1513	A1426	G1348	A1265	U1186	C1111	G1035	A971	U900	A828	C733	G514	G432
A1516	G1431	U1351	G1266	A1190	U1112	C1037	C975	G903	G824	U734	G515	U433
A1517	A1432	U1352	G1267	A1191	A1113	G1038	C976	G904	G825	G735	G516	U434
G1593	A1433	C1353	A1268	A1192	U1114	G1039	A977	G897	G827	U736	A517	G435
U1594	U1434	G1354	A1269	A1193	A1115	G1040	A978	U907	A829	A737	A518	A436
U1595	A1435	U1355	A1277	U1194	U1116	A1042	A979	U908	A830	C738	A519	A437
U1596	U1436	U1356	G1278	U1201	A1117	A1043	C977	U909	G831	A740	A524	C442
C1597	A1437	G1357	G1285	A1202	U1118	A1044	U976	U907	G832	A743	A525	U448
U1598	G1438	U1358	A1286	A1203	A1119	A1045	U977	U907	G833	A752	A526	A449
U1599	U1439	A1360	A1287	A1204	C1120	A1046	U978	A908	A834	A753	A527	U453
A1600	A1440	A1361	G1288	A1205	G1121	A1047	C978	U909	G835	A754	C528	U454
A1601	U1441	U1362	G1289	U1206	C1122	A1048	U979	U909	G836	A755	A530	G453
A1606	A1442	G1363	G1290	G1207	A1123	A1049	C980	A910	U837	U758	A537	A456
A1607	U1443	U1364	G1291	G1208	U1124	A1050	C981	A911	U838	A759	A538	A459
A1608	A1444	U1365	A1292	G1209	U1125	A1051	U982	G911	G839	A762	A543	C460
G1613	G1450	G1371	G1293	U1218	A1130	A1052	U983	C912	A840	A763	U544	A462
A1614	U1451	A1294	A1294	C1219	A1131	A1053	U984	A913	A841	C764	G546	U474
A1615	A1452	C1374	U1295	G1220	A1132	U1064	U985	C914	C842	U765	G547	A475
A1616	A1453	A1454	G1296	A1221	G1133	U1065	G986	U915	C843	A766	G548	A476
A1617	G1454	A1375	G1297	A1222	A1134	A1066	A987	U917	A847	A769	G549	U477
A1618	C1455	G1376	C1298	A1223	G1135	A1067	A991	U918	G848	A770	A549	A478
U1619	A1456	G1377	G1299	G1224	U1136	A1068	A992	U919	A849	A771	A550	U479
A1620	U1459	U1378	G1300	U1225	G1137	U1069	A993	C921	G850	A772	G551	G468
U1626	A1500	U1380	A1302	U1226	C1138	G1070	A994	A922	A851	G775	A552	A469
A1627	A1501	A1381	U1303	G1227	G1139	G1071	C997	U923	G852	G776	G553	C461
G1628	A1502	G1382	G1304	U1228	U1140	A1072	C998	U924	C853	C777	A554	A462
C1629	U1543	U1383	A1305	A1230	A1141	A1073	A999	A925	U854	C778	A555	U475
U1630	A1465	C1384	G1306	G1231	U1142	A1074	A1003	G928	G855	A781	U556	A476
A1553	U1554	A1473	A1308	A1233	A1144	A1075	U1004	G929	U858	A782	C556	A477

A2851	A2852	A2853	A2854	A2855	A2856	A2857	A2858	A2859	A2860	A2861	A2862	A2863	A2864	A2865	A2866	A2867	A2868	A2869	A2870	A2871	A2872	A2873	A2874	A2875	A2876	A2877	A2878	A2879	A2880	A2881	A2882	A2883	A2884	A2885	A2886	A2887	A2888	A2889	A2890	A2891	A2892	A2893	A2897	A2898	A2899	A2900	A2901	A2902	A2903	A2904	A2905	A2906	A2907	A2908	A2909	A2912	A2916	A2917	A2918	A2919	A2923																																																																																																																																																																																																																																																																																																																																																																																																																																					
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A2302	A2303	A2307	G2308	C2310	C2312	C2313	C2314	C2315	C2316	C2317	C2324	C2327	C2328	C2329	C2330	C2331	C2332	C2333	C2334	C2335	C2336	C2337	C2338	C2339	C2340	C2341	C2342	C2343	C2344	C2345	C2346	C2347	C2348	C2349	C2350	C2351	C2352	C2355	C2356	C2357	C2358	C2359	C2362	C2363	C2364	C2365	C2366	C2369	C2370	C2371	C2372	C2373	C2374	C2375	C2376	C2377	C2378	C2379	C2380	C2381	C2382	C2383	C2386	C2387	C2388	C2389	C2390	C2393	C2394	C2395	C2398	C2399	C2400	C2401	C2402	C2403	C2404	C2405	C2406	C2407	C2411	C2412	C2413	C2414	C2417	C2418	C2419	C2420	C2421	C2422	C2423	C2427	C2431	C2434	C2435	C2436	C2439	C2440	C2441	C2447	C2448	C2453	C2454																																																																																																																																																																																																																																																																																																																																																																																											
G2195	C2196	C2197	A2200	C2201	A2202	A2205	A2206	A2214	C2215	A2216	A2220	C2221	C2222	C2223	C2224	A2227	A2228	C2232	C2233	C2236	C2237	C2238	C2239	C2240	C2241	C2242	C2243	C2246	C2252	C2253	C2254	C2255	C2256	A2262	C2263	C2267	C2268	C2269	C2270	A2276	C2277	C2278	C2279	A2285	A2286	A2298	C2299	C2300	C2301	C2302	C2303	C2304	C2305	C2306	C2307	C2308	C2309	C2310	C2311	C2312	C2313	C2314	C2315	C2316	C2317	C2318	C2319	C2320	C2321	C2322	C2323	C2324	C2325	C2326	C2327	C2328	C2329	C2330	C2331	C2332	C2333	C2334	C2335	C2336	C2337	C2338	C2339	C2340	C2341	C2342	C2343	C2344	C2345	C2346	C2347	C2348	C2349	C2350	C2351	C2352	C2353	C2354	C2355	C2356	C2357	C2358	C2359	C2360	C2361	C2362	C2363	C2364	C2365	C2366	C2367	C2368	C2369	C2370	C2371	C2372	C2373	C2374	C2375	C2376	C2377	C2378	C2379	C2380	C2381	C2382	C2383	C2384	C2385	C2386	C2387	C2388	C2389	C2390	C2391	C2392	C2393	C2394	C2395	C2396	C2397	C2398	C2399	C2400	C2401	C2402	C2403	C2404	C2405	C2406	C2407	C2408	C2409	C2410	C2411	C2412	C2413	C2414	C2415	C2416	C2417	C2418	C2419	C2420	C2421	C2422	C2423	C2424	C2425	C2426	C2427	C2428	C2429	C2430	C2431	C2432	C2433	C2434	C2435	C2436	C2437	C2438	C2439	C2440	C2441	C2442	C2443	C2444	C2445	C2446	C2447	C2448	C2449	C2450	C2451	C2452	C2453	C2454	C2455	C2456	C2457	C2458	C2459	C2460	C2461	C2462	C2463	C2464	C2465	C2466	C2467	C2468	C2469	C2470	C2471	C2472	C2473	C2474	C2475	C2476	C2477	C2478	C2479	C2480	C2481	C2482	C2483	C2484	C2485	C2486	C2487	C2488	C2489	C2490	C2491	C2492	C2493	C2494	C2495	C2496	C2497	C2498	C2499	C2500	C2501	C2502	C2503	C2504	C2505	C2506	C2507	C2508	C2509	C2510	C2511	C2512	C2513	C2514	C2515	C2516	C2517	C2518	C2519	C2520	C2521	C2522	C2523	C2524	C2525	C2526	C2527	C2528	C2529	C2530	C2531	C2532	C2533	C2534	C2535	C2536	C2537	C2538	C2539	C2540	C2541	C2542	C2543	C2544	C2545	C2546	C2547	C2548	C2549	C2550	C2551	C2552	C2553	C2554	C2555	C2556	C2557	C2558	C2559	C2560	C2561	C2562	C2563	C2564	C2565	C2566	C2567	C2568	C2569	C2570	C2571	C2572	C2573	C2574	C2575	C2576	C2577	C2578	C2579	C2580	C2581	C2582	C2583	C2584	C2585	C2586	C2587	C2588	C2589	C2590	C2591	C2592	C2593	C2594	C2595	C2596	C2597	C2598	C2599	C2600	C2601	C2602	C2603	C2604	C2605	C2606	C2607	C2608	C2609	C2610	C2611	C2612	C2613	C2614	C2615	C2616	C2617	C2618	C2619	C2620	C2621	C2622	C2623	C2624	C2625	C2626	C2627	C2628	C2629	C2630	C2631	C2632	C2633	C2634	C2635	C2636	C2637	C2638	C2639	C2640	C2641	C2642	C2643	C2644	C2645	C2646	C2647	C2648	C2649	C2650	C2651	C2652	C2653	C2654	C2655	C2656	C2657	C2658	C2659	C2660	C2661	C2662	C2663	C2664	C2665	C2666	C2667	C2668	C2669	C2670	C2671	C2672	C2673	C2674	C2675	C2676	C2677	C2678	C2679	C2680	C2681	C2682	C2683	C2684	C2685	C2686	C2687	C2688	C2689	C2690	C2691	C2692	C2693	C2694	C2695	C2696	C2697	C2698	C2699	C2700	C2701	C2702	C2703	C2704	C2705	C2706	C2707	C2708	C2709	C2710	C2711	C2712	C2713	C2714	C2715	C2716	C2717	C2718	C2719	C2720	C2721	C2722	C2723	C2724	C2725	C2726	C2727	C2728	C2729	C2730	C2731	C2732	C2733	C2



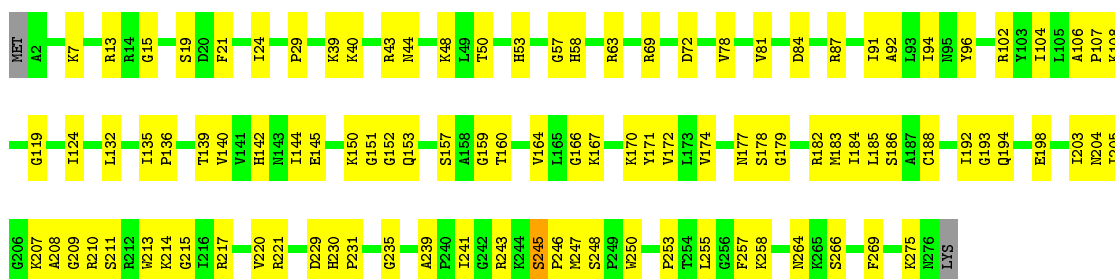
- Molecule 34: 5S ribosomal RNA

Chain BB:



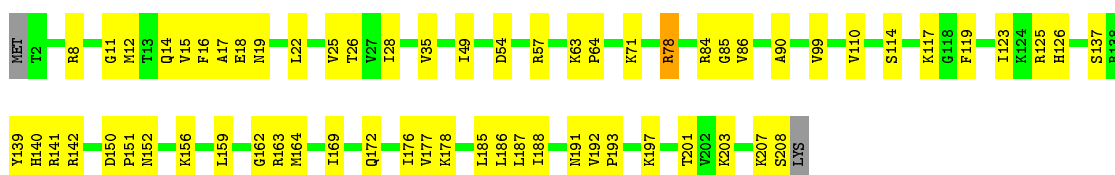
- Molecule 35: 50S ribosomal protein uL2

Chain BD:



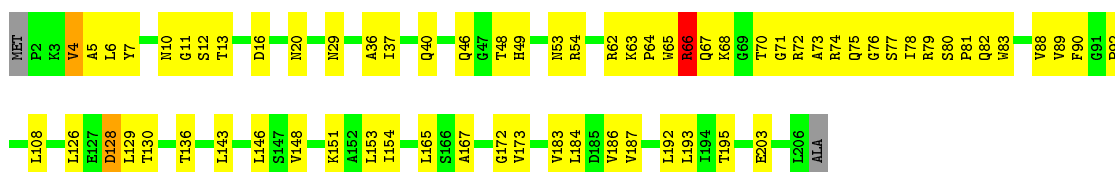
- Molecule 36: 50S ribosomal protein uL3

Chain BE:



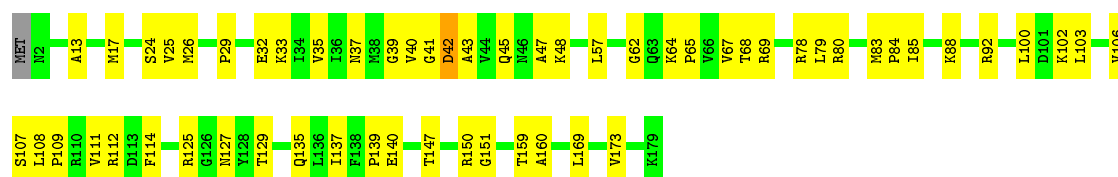
- Molecule 37: 50S ribosomal protein uL4

Chain BF:



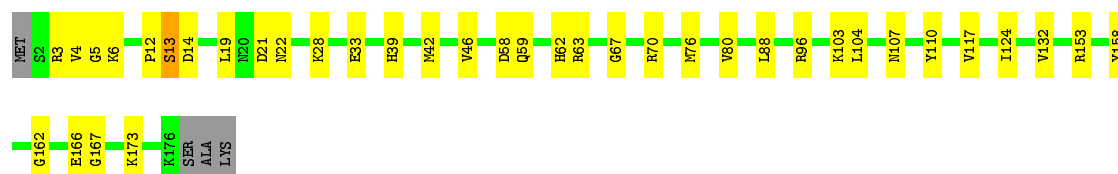
- Molecule 38: 50S ribosomal protein uL5

Chain BG:



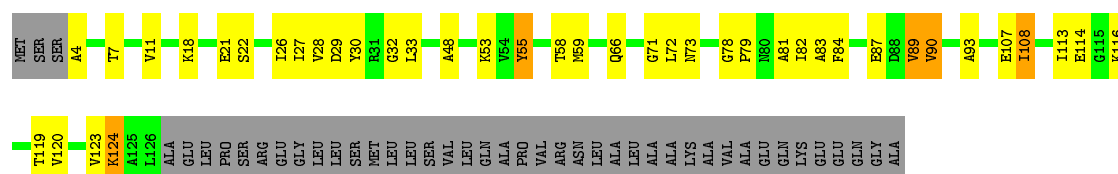
- Molecule 39: 50S ribosomal protein uL6

Chain BH: 77% 21% ..



- Molecule 40: 50S ribosomal protein uL10

Chain BJ: 49% 22% 26%



- Molecule 41: 50S ribosomal protein uL11

Chain BK: 77% 16% 6%



- Molecule 42: 50S ribosomal protein uL13

Chain BM: 70% 28%



- Molecule 43: 50S ribosomal protein uL14

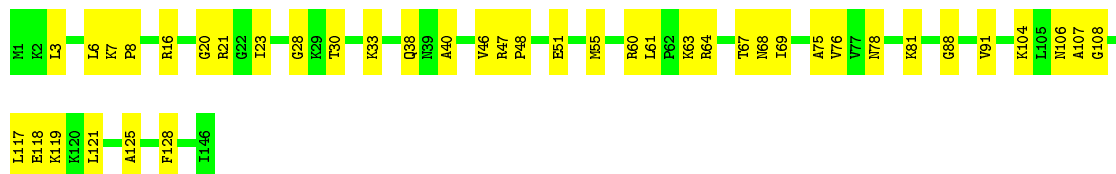
Chain BN: 72% 28%





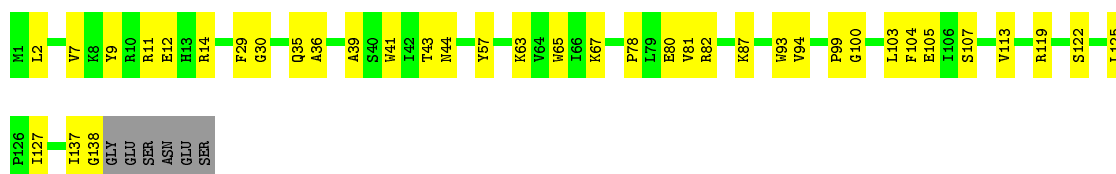
- Molecule 44: 50S ribosomal protein uL15

Chain BO: 72% 28%



- Molecule 45: 50S ribosomal protein uL16

Chain BP: 69% 26%



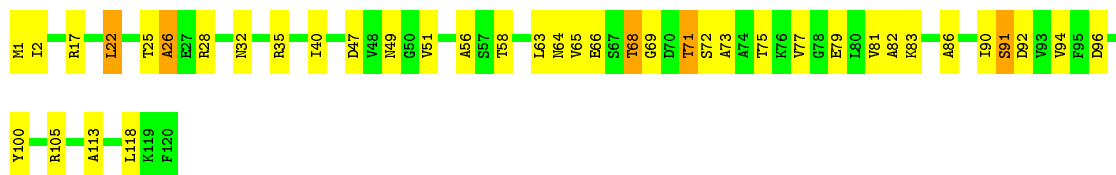
- Molecule 46: 50S ribosomal protein bL17

Chain BQ: 73% 25%



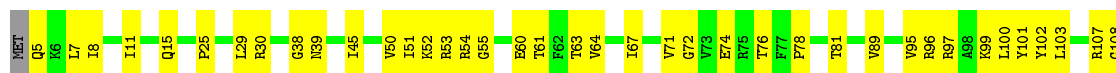
- Molecule 47: 50S ribosomal protein uL18

Chain BR: 67% 29%

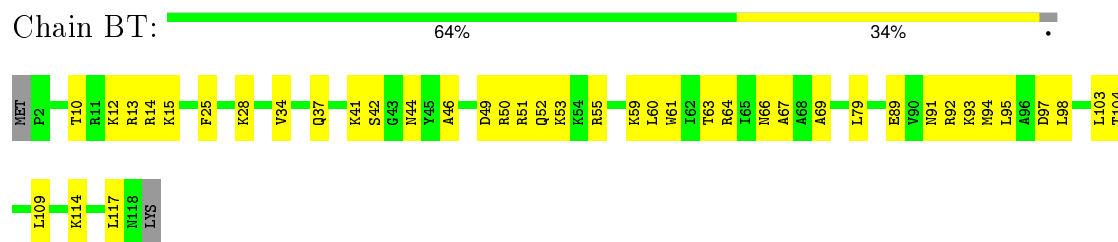


- Molecule 48: 50S ribosomal protein bL19

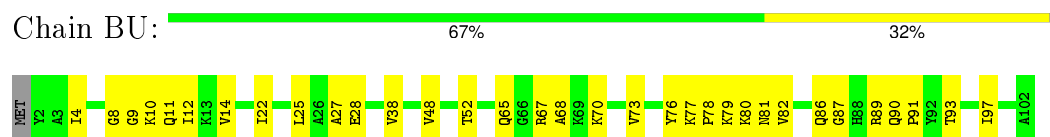
Chain BS: 65% 34%



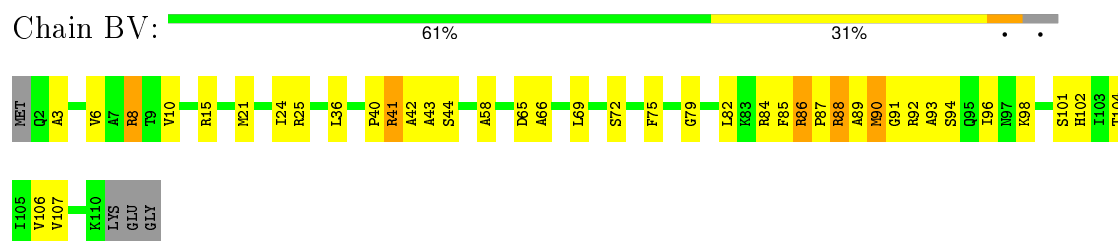
- Molecule 49: 50S ribosomal protein bL20



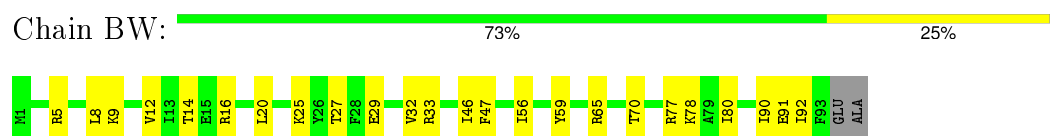
- Molecule 50: 50S ribosomal protein bL21



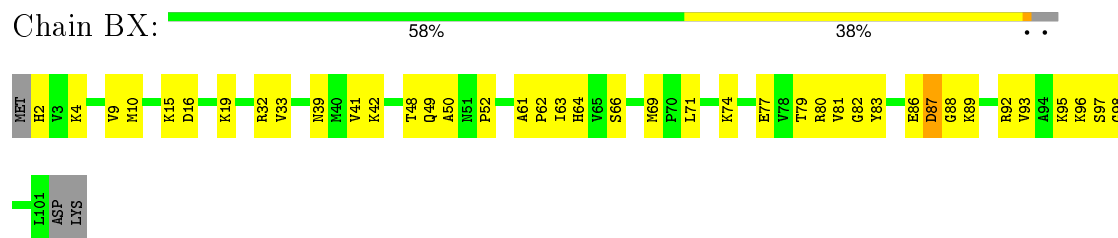
- Molecule 51: 50S ribosomal protein uL22



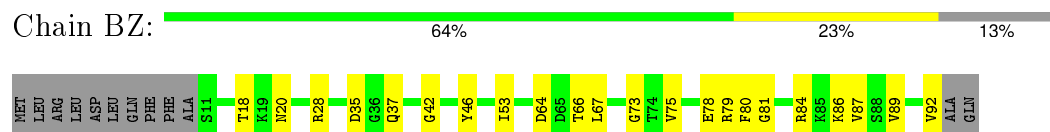
- Molecule 52: 50S ribosomal protein uL23



- Molecule 53: 50S ribosomal protein uL24



- Molecule 54: 50S ribosomal protein bL27



4 Experimental information

Property	Value	Source
Reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, Not provided	Depositor
Number of images	305045	Depositor
Resolution determination method	FSC 0.143	Depositor
CTF correction method	defocus groups	Depositor
Microscope	FEI TITAN KRIOS	Depositor
Voltage (kV)	300	Depositor
Electron dose ($e^-/\text{\AA}^2$)	28	Depositor
Minimum defocus (nm)	1000	Depositor
Maximum defocus (nm)	2500	Depositor
Magnification	125085	Depositor
Image detector	FEI FALCON II (4k x 4k)	Depositor

5 Model quality ⓘ

5.1 Standard geometry ⓘ

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 >2	RMSZ	# Z >2
1	AA	1.36	984/37074 (2.7%)	2.90	3806/57834 (6.6%)
10	AJ	0.28	0/408	0.39	0/507
11	AK	0.22	0/471	0.42	0/587
12	AL	0.24	0/548	0.50	0/682
13	AM	0.30	0/475	0.52	0/592
14	AN	0.21	0/240	0.48	0/297
15	AO	0.27	0/352	0.41	0/437
16	AP	0.27	0/356	0.41	0/442
17	AQ	0.27	0/344	0.44	0/427
18	AR	0.31	0/284	0.43	0/352
19	AS	0.33	0/335	0.46	0/417
2	AB	0.31	0/895	0.40	0/1117
20	AT	0.26	0/344	0.40	0/427
21	AX	1.03	30/1834 (1.6%)	2.18	104/2858 (3.6%)
22	AY	0.38	0/466	0.93	2/726 (0.3%)
23	AZ	0.50	0/106	1.02	0/122
24	B0	0.30	0/448	0.58	0/596
25	B1	0.24	0/531	0.47	0/707
26	B2	0.24	0/457	0.44	0/613
27	B3	0.27	0/513	0.43	0/683
28	B4	0.23	0/433	0.48	0/574
29	B5	0.25	0/406	0.44	0/540
3	AC	0.29	0/839	0.38	0/1047
30	B6	0.20	0/370	0.44	0/483
31	B7	0.23	0/519	0.48	0/680
32	B8	0.19	0/291	0.36	0/383
33	BA	1.40	1948/70307 (2.8%)	2.98	7636/109687 (7.0%)
34	BB	1.30	66/2678 (2.5%)	2.78	248/4174 (5.9%)
35	BD	0.27	0/2148	0.48	0/2881
36	BE	0.27	0/1597	0.47	0/2140
37	BF	0.26	0/1580	0.50	0/2132
38	BG	0.29	0/1423	0.50	0/1910
39	BH	0.23	0/1360	0.43	0/1832
4	AD	0.25	0/796	0.41	0/992

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	# Z >2	RMSZ	# Z >2
40	BJ	0.26	0/963	0.49	0/1298
41	BK	0.26	0/995	0.48	0/1346
42	BM	0.25	0/1146	0.49	0/1542
43	BN	0.28	0/927	0.47	0/1245
44	BO	0.23	0/1093	0.44	0/1457
45	BP	0.20	0/1120	0.38	0/1496
46	BQ	0.26	0/960	0.50	0/1284
47	BR	0.30	0/921	0.54	1/1236 (0.1%)
48	BS	0.24	0/949	0.44	0/1269
49	BT	0.26	0/952	0.45	0/1266
5	AE	0.26	0/660	0.46	0/822
50	BU	0.28	0/797	0.53	0/1070
51	BV	0.34	0/851	0.59	0/1146
52	BW	0.29	0/759	0.47	0/1011
53	BX	0.26	0/764	0.52	0/1022
54	BZ	0.30	0/638	0.49	0/847
6	AF	0.31	0/380	0.41	0/472
7	AG	0.26	0/612	0.39	0/762
8	AH	0.24	0/524	0.43	0/652
9	AI	0.26	0/520	0.51	0/647
All	All	1.20	3028/147759 (2.0%)	2.62	11797/221768 (5.3%)

Chiral center outliers are detected by calculating the chiral volume of a chiral center and verifying if the center is modelled as a planar moiety or with the opposite hand. A planarity outlier is detected by checking planarity of atoms in a peptide group, atoms in a mainchain group or atoms of a sidechain that are expected to be planar.

Mol	Chain	#Chirality outliers	#Planarity outliers
1	AA	0	36
21	AX	0	2
22	AY	0	7
33	BA	0	84
34	BB	0	2
37	BF	0	1
51	BV	0	2
All	All	0	134

The worst 5 of 3028 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
1	AA	508	A	C8-N7	8.50	1.37	1.31
1	AA	439	A	C8-N7	8.28	1.37	1.31

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Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
33	BA	1188	A	C8-N7	8.21	1.37	1.31
1	AA	1372	A	C8-N7	8.18	1.37	1.31
33	BA	526	A	C8-N7	8.14	1.37	1.31

The worst 5 of 11797 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
33	BA	1134	A	C2-N3-C4	20.56	120.88	110.60
33	BA	226	A	C2-N3-C4	20.47	120.84	110.60
33	BA	1691	A	C2-N3-C4	20.46	120.83	110.60
1	AA	1308	A	C2-N3-C4	20.36	120.78	110.60
1	AA	195	A	C2-N3-C4	20.24	120.72	110.60

There are no chirality outliers.

5 of 134 planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	AA	209	A	Sidechain
1	AA	308	A	Sidechain
1	AA	335	A	Sidechain
1	AA	53	A	Sidechain
1	AA	76	A	Sidechain

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	AA	33115	0	16676	250	0
2	AB	896	0	244	2	0
3	AC	840	0	241	3	0
4	AD	797	0	224	3	0
5	AE	661	0	197	1	0
6	AF	381	0	104	0	0
7	AG	613	0	164	1	0
8	AH	525	0	146	0	0
9	AI	521	0	155	4	0
10	AJ	409	0	104	0	0

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Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
11	AK	472	0	135	7	0
12	AL	549	0	157	1	0
13	AM	476	0	137	2	0
14	AN	241	0	62	2	0
15	AO	353	0	94	0	0
16	AP	357	0	95	0	0
17	AQ	345	0	90	1	0
18	AR	285	0	76	1	0
19	AS	336	0	93	2	0
20	AT	345	0	89	2	0
21	AX	1643	0	830	30	0
22	AY	415	0	207	80	0
23	AZ	107	0	58	9	0
24	B0	444	0	486	61	0
25	B1	530	0	568	12	0
26	B2	455	0	491	12	0
27	B3	503	0	494	12	0
28	B4	426	0	445	14	0
29	B5	401	0	413	18	0
30	B6	367	0	410	20	0
31	B7	512	0	564	31	0
32	B8	288	0	330	3	0
33	BA	62767	0	31584	776	0
34	BB	2395	0	1212	21	0
35	BD	2111	0	2200	85	0
36	BE	1575	0	1642	48	0
37	BF	1561	0	1647	92	0
38	BG	1404	0	1467	44	0
39	BH	1342	0	1388	26	0
40	BJ	955	0	990	24	0
41	BK	981	0	1020	27	0
42	BM	1123	0	1162	31	0
43	BN	920	0	977	21	0
44	BO	1081	0	1132	40	0
45	BP	1097	0	1165	25	0
46	BQ	953	0	983	28	0
47	BR	912	0	947	36	0
48	BS	936	0	1008	31	0
49	BT	940	0	1005	41	0
50	BU	786	0	826	40	0
51	BV	842	0	899	46	0
52	BW	752	0	802	19	0

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Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
53	BX	754	0	809	31	0
54	BZ	630	0	644	16	0
All	All	135425	0	80088	1864	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 9.

The worst 5 of 1864 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)
23:AZ:74:ILE:CA	51:BV:85:PHE:HE2	1.11	1.60
23:AZ:74:ILE:CA	51:BV:85:PHE:CE2	1.99	1.44
24:B0:37:ARG:NH1	24:B0:44:PRO:HG3	1.37	1.40
37:BF:80:SER:OG	37:BF:81:PRO:HD2	1.30	1.29
41:BK:15:ALA:HB1	41:BK:46:THR:CG2	1.62	1.28

There are no symmetry-related clashes.

5.3 Torsion angles [i](#)

5.3.1 Protein backbone [i](#)

In the following table, the Percentiles column shows the percent Ramachandran outliers of the chain as a percentile score with respect to all 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	
2	AB	222/246 (90%)	204 (92%)	13 (6%)	5 (2%)	8	50
3	AC	208/218 (95%)	193 (93%)	14 (7%)	1 (0%)	34	76
4	AD	197/200 (98%)	191 (97%)	4 (2%)	2 (1%)	19	64
5	AE	163/166 (98%)	150 (92%)	9 (6%)	4 (2%)	7	48
6	AF	93/95 (98%)	88 (95%)	3 (3%)	2 (2%)	8	50
7	AG	151/156 (97%)	144 (95%)	6 (4%)	1 (1%)	26	70
8	AH	129/132 (98%)	123 (95%)	5 (4%)	1 (1%)	24	68
9	AI	128/130 (98%)	113 (88%)	10 (8%)	5 (4%)	4	37

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Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
10	AJ	100/102 (98%)	88 (88%)	8 (8%)	4 (4%)	4	37
11	AK	116/131 (88%)	106 (91%)	9 (8%)	1 (1%)	21	65
12	AL	135/138 (98%)	119 (88%)	9 (7%)	7 (5%)	2	31
13	AM	117/121 (97%)	94 (80%)	13 (11%)	10 (8%)	1	17
14	AN	58/61 (95%)	51 (88%)	4 (7%)	3 (5%)	2	31
15	AO	86/89 (97%)	82 (95%)	2 (2%)	2 (2%)	8	50
16	AP	87/90 (97%)	82 (94%)	3 (3%)	2 (2%)	8	50
17	AQ	84/87 (97%)	78 (93%)	6 (7%)	0	100	100
18	AR	69/79 (87%)	64 (93%)	2 (3%)	3 (4%)	3	35
19	AS	82/92 (89%)	75 (92%)	5 (6%)	2 (2%)	7	49
20	AT	84/88 (96%)	77 (92%)	6 (7%)	1 (1%)	16	61
23	AZ	22/95 (23%)	17 (77%)	2 (9%)	3 (14%)	0	6
24	B0	56/62 (90%)	53 (95%)	1 (2%)	2 (4%)	4	40
25	B1	63/66 (96%)	60 (95%)	3 (5%)	0	100	100
26	B2	56/59 (95%)	54 (96%)	1 (2%)	1 (2%)	11	54
27	B3	62/66 (94%)	56 (90%)	4 (6%)	2 (3%)	5	43
28	B4	52/59 (88%)	47 (90%)	4 (8%)	1 (2%)	10	53
29	B5	46/49 (94%)	44 (96%)	2 (4%)	0	100	100
30	B6	42/44 (96%)	41 (98%)	1 (2%)	0	100	100
31	B7	62/66 (94%)	56 (90%)	5 (8%)	1 (2%)	12	56
32	B8	34/37 (92%)	33 (97%)	1 (3%)	0	100	100
35	BD	273/277 (99%)	264 (97%)	8 (3%)	1 (0%)	39	79
36	BE	205/209 (98%)	189 (92%)	11 (5%)	5 (2%)	7	49
37	BF	203/207 (98%)	184 (91%)	16 (8%)	3 (2%)	13	57
38	BG	176/179 (98%)	154 (88%)	18 (10%)	4 (2%)	8	50
39	BH	173/179 (97%)	165 (95%)	7 (4%)	1 (1%)	30	73
40	BJ	121/166 (73%)	97 (80%)	14 (12%)	10 (8%)	1	18
41	BK	131/141 (93%)	122 (93%)	7 (5%)	2 (2%)	13	57
42	BM	140/145 (97%)	130 (93%)	9 (6%)	1 (1%)	26	70
43	BN	120/122 (98%)	112 (93%)	6 (5%)	2 (2%)	11	55
44	BO	144/146 (99%)	132 (92%)	10 (7%)	2 (1%)	14	58

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Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
45	BP	136/144 (94%)	129 (95%)	7 (5%)	0	100	100
46	BQ	117/120 (98%)	109 (93%)	7 (6%)	1 (1%)	21	65
47	BR	118/120 (98%)	106 (90%)	7 (6%)	5 (4%)	3	35
48	BS	112/115 (97%)	100 (89%)	12 (11%)	0	100	100
49	BT	115/119 (97%)	112 (97%)	3 (3%)	0	100	100
50	BU	99/102 (97%)	82 (83%)	15 (15%)	2 (2%)	9	52
51	BV	107/113 (95%)	96 (90%)	8 (8%)	3 (3%)	6	46
52	BW	91/95 (96%)	86 (94%)	5 (6%)	0	100	100
53	BX	98/103 (95%)	87 (89%)	8 (8%)	3 (3%)	5	44
54	BZ	80/94 (85%)	77 (96%)	3 (4%)	0	100	100
All	All	5563/5920 (94%)	5116 (92%)	336 (6%)	111 (2%)	14	52

5 of 111 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
5	AE	4	ILE
6	AF	70	ALA
12	AL	127	ARG
13	AM	101	ASN
23	AZ	78	PHE

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	
23	AZ	6/87 (7%)	4 (67%)	2 (33%)	0	3
24	B0	47/50 (94%)	47 (100%)	0	100	100
25	B1	56/57 (98%)	56 (100%)	0	100	100
26	B2	52/53 (98%)	52 (100%)	0	100	100
27	B3	53/55 (96%)	53 (100%)	0	100	100
28	B4	48/53 (91%)	48 (100%)	0	100	100

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Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
29	B5	46/47 (98%)	46 (100%)	0	100	100
30	B6	39/39 (100%)	39 (100%)	0	100	100
31	B7	54/56 (96%)	54 (100%)	0	100	100
32	B8	34/35 (97%)	34 (100%)	0	100	100
35	BD	223/225 (99%)	223 (100%)	0	100	100
36	BE	168/170 (99%)	168 (100%)	0	100	100
37	BF	169/170 (99%)	168 (99%)	1 (1%)	90	95
38	BG	153/154 (99%)	153 (100%)	0	100	100
39	BH	148/151 (98%)	148 (100%)	0	100	100
40	BJ	105/138 (76%)	105 (100%)	0	100	100
41	BK	103/110 (94%)	103 (100%)	0	100	100
42	BM	120/123 (98%)	120 (100%)	0	100	100
43	BN	101/101 (100%)	101 (100%)	0	100	100
44	BO	110/110 (100%)	110 (100%)	0	100	100
45	BP	111/116 (96%)	111 (100%)	0	100	100
46	BQ	99/100 (99%)	99 (100%)	0	100	100
47	BR	93/93 (100%)	93 (100%)	0	100	100
48	BS	99/100 (99%)	99 (100%)	0	100	100
49	BT	96/98 (98%)	96 (100%)	0	100	100
50	BU	83/84 (99%)	83 (100%)	0	100	100
51	BV	90/93 (97%)	89 (99%)	1 (1%)	80	90
52	BW	84/85 (99%)	84 (100%)	0	100	100
53	BX	84/87 (97%)	84 (100%)	0	100	100
54	BZ	64/74 (86%)	64 (100%)	0	100	100
All	All	2738/2914 (94%)	2734 (100%)	4 (0%)	95	97

All (4) residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
23	AZ	81	ASN
23	AZ	86	ASP
37	BF	66	ARG
51	BV	90	MET

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

Mol	Chain	Res	Type
37	BF	67	GLN
39	BH	62	HIS
53	BX	39	ASN
37	BF	75	GLN
37	BF	82	GLN

5.3.3 RNA ⓘ

Mol	Chain	Analysed	Backbone Outliers	Pucker Outliers
1	AA	1543/1555 (99%)	235 (15%)	17 (1%)
21	AX	76/77 (98%)	15 (19%)	1 (1%)
22	AY	18/19 (94%)	14 (77%)	3 (16%)
33	BA	2922/2928 (99%)	791 (27%)	80 (2%)
34	BB	111/119 (93%)	32 (28%)	4 (3%)
All	All	4670/4698 (99%)	1087 (23%)	105 (2%)

5 of 1087 RNA backbone outliers are listed below:

Mol	Chain	Res	Type
1	AA	10	A
1	AA	11	G
1	AA	34	A
1	AA	41	G
1	AA	49	C

5 of 105 RNA pucker outliers are listed below:

Mol	Chain	Res	Type
33	BA	837	U
33	BA	1245	G
33	BA	2784	C
33	BA	847	A
33	BA	1092	A

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

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

5.5 Carbohydrates [i](#)

There are no carbohydrates in this entry.

5.6 Ligand geometry [i](#)

There are no ligands in this entry.

5.7 Other polymers [i](#)

There are no such residues in this entry.

5.8 Polymer linkage issues [i](#)

There are no chain breaks in this entry.