



## wwPDB EM Validation Summary Report ⓘ

Jul 14, 2024 – 01:39 pm BST

PDB ID : 8BJQ  
EMDB ID : EMD-16090  
Title : Structure of a yeast 80S ribosome-bound N-Acetyltransferase B complex  
Authors : Knorr, A.G.; Mackens-Kiani, T.; Musial, J.; Berninghausen, O.; Becker, T.;  
Beatrix, B.; Beckmann, R.  
Deposited on : 2022-11-05  
Resolution : 3.80 Å(reported)

This is a wwPDB EM Validation Summary Report for a publicly released PDB entry.

We welcome your comments at [validation@mail.wwpdb.org](mailto:validation@mail.wwpdb.org)

A user guide is available at

<https://www.wwpdb.org/validation/2017/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.dev92
MolProbity	:	4.02b-467
Percentile statistics	:	20191225.v01 (using entries in the PDB archive December 25th 2019)
MapQ	:	1.9.13
Ideal geometry (proteins)	:	Engh & Huber (2001)
Ideal geometry (DNA, RNA)	:	Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP)	:	2.37.1

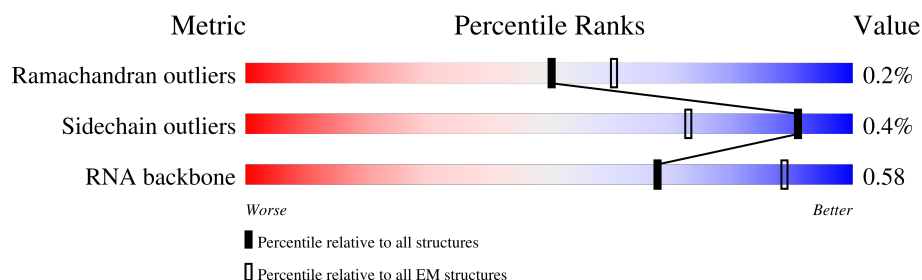
# 1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

*ELECTRON MICROSCOPY*

The reported resolution of this entry is 3.80 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.



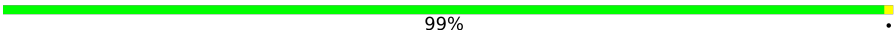
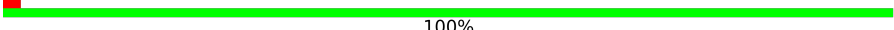
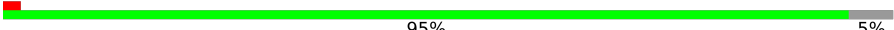
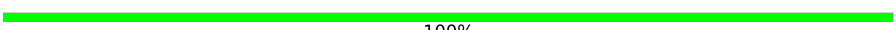
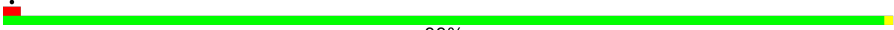





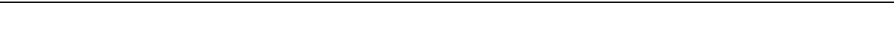

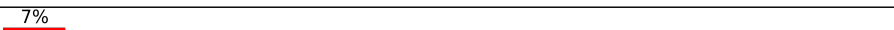
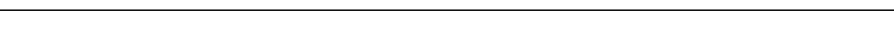
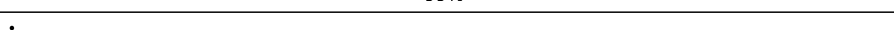

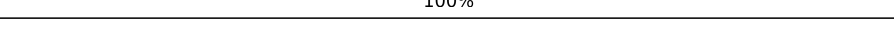
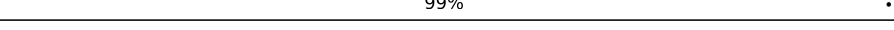
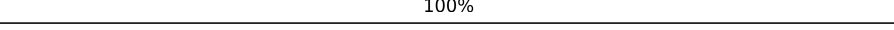
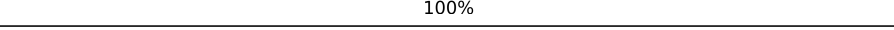

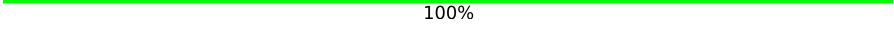
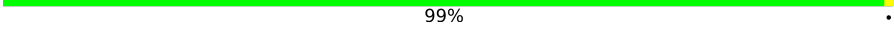
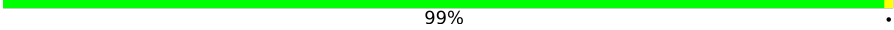
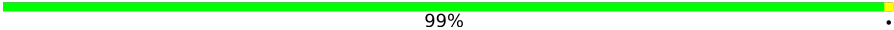
Metric	Whole archive (#Entries)	EM structures (#Entries)
Ramachandran outliers	154571	4023
Sidechain outliers	154315	3826
RNA backbone	4643	859

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  $\geq 3$ , 2, 1 and 0 types of geometric quality criteria respectively. A grey segment represents the fraction of residues that are not modelled. The numeric value for each fraction is indicated below the corresponding segment, with a dot representing fractions  $\leq 5\%$ . The upper red bar (where present) indicates the fraction of residues that have poor fit to the EM map (all-atom inclusion  $< 40\%$ ). The numeric value is given above the bar.

Mol	Chain	Length	Quality of chain
1	A	195	<div> <div>82%</div> <div>98%</div> </div>
1	C	195	<div> <div>29%</div> <div>34%</div> <div>65%</div> </div>
2	B	796	<div> <div>41%</div> <div>100%</div> </div>
2	D	796	<div> <div>56%</div> <div>98%</div> </div>
3	C4	121	<div> <div>90%</div> <div>10%</div> </div>
4	C3	158	<div> <div>84%</div> <div>16%</div> </div>
5	LA	251	<div> <div>100%</div> </div>
6	LB	386	<div> <div>100%</div> </div>

Continued on next page...

*Continued from previous page...*

Mol	Chain	Length	Quality of chain
7	LC	361	 99%
8	LD	294	 100%
9	LE	175	 95% 5%
10	LF	222	 100%
11	LG	233	 99%
12	LH	191	 99%
13	LI	218	 100%
14	LJ	169	 99%
15	LL	193	 98%
16	LM	136	 99%
17	LN	203	 100%
18	LO	197	 99%
19	LP	183	 7% 100%
20	LQ	185	 99%
21	LR	188	 90% 9%
22	LS	171	 100%
23	LT	159	 99%
24	LU	100	 100%
25	LV	136	 100%
26	LW	126	 53% 47%
27	LX	121	 100%
28	LY	125	 99%
29	LZ	135	 99%
30	La	148	 99%
31	Lb	58	 100%

*Continued on next page...*

*Continued from previous page...*

Mol	Chain	Length	Quality of chain
32	Lc	96	 99%
33	Ld	109	 99%
34	Le	127	 100%
35	Lf	106	 99%
36	Lg	112	 100%
37	Lh	119	 99%
38	Li	99	 100%
39	Lj	85	 100%
40	Lk	77	 100%
41	Ll	50	 100%
42	Lm	52	 98%
43	Ln	25	 96%
44	Lo	103	 100%
45	Lp	91	 100%
46	1	3395	 79% 18%

## 2 Entry composition [i](#)

There are 48 unique types of molecules in this entry. The entry contains 141384 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 protein called N-terminal acetyltransferase B complex catalytic subunit NAT3.

Mol	Chain	Residues	Atoms					AltConf	Trace
1	A	195	Total	C	N	O	S	0	0
			1612	1030	273	297	12		
1	C	68	Total	C	N	O	S	0	0
			571	378	81	107	5		

- Molecule 2 is a protein called N-terminal acetyltransferase B complex subunit MDM20.

Mol	Chain	Residues	Atoms					AltConf	Trace
2	B	796	Total	C	N	O	S	0	0
			6536	4195	1080	1231	30		
2	D	796	Total	C	N	O	S	0	0
			6536	4195	1080	1231	30		

- Molecule 3 is a RNA chain called 5S rRNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
3	C4	121	Total	C	N	O	P	0	0
			2579	1152	461	845	121		

- Molecule 4 is a RNA chain called 5.8S rRNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
4	C3	158	Total	C	N	O	P	0	0
			3353	1500	586	1109	158		

- Molecule 5 is a protein called 60S ribosomal protein L2-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
5	LA	251	Total	C	N	O	S	0	0
			1899	1182	385	331	1		

- Molecule 6 is a protein called 60S ribosomal protein L3.

Mol	Chain	Residues	Atoms					AltConf	Trace
6	LB	386	Total	C	N	O	S	0	0
			3075	1950	584	533	8		

- Molecule 7 is a protein called 60S ribosomal protein L4-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
7	LC	361	Total	C	N	O	S	0	0
			2748	1729	522	494	3		

- Molecule 8 is a protein called 60S ribosomal protein L5.

Mol	Chain	Residues	Atoms					AltConf	Trace
8	LD	294	Total	C	N	O	S	0	0
			2351	1484	410	455	2		

- Molecule 9 is a protein called 60S ribosomal protein L6-B.

Mol	Chain	Residues	Atoms					AltConf	Trace
9	LE	167	Total	C	N	O	S	0	0
			1305	841	234	229	1		

There are 3 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
LE	132	ALA	THR	conflict	UNP P05739
LE	146	ILE	LEU	conflict	UNP P05739
LE	173	MET	LEU	conflict	UNP P05739

- Molecule 10 is a protein called 60S ribosomal protein L7-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
10	LF	222	Total	C	N	O	S	0	0
			1784	1151	324	308	1		

- Molecule 11 is a protein called 60S ribosomal protein L8-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
11	LG	233	Total	C	N	O	S	0	0
			1804	1151	323	327	3		

- Molecule 12 is a protein called 60S ribosomal protein L9-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
12	LH	191	Total	C	N	O	S	0	0
			1508	957	274	273	4		

- Molecule 13 is a protein called 60S ribosomal protein L10.

Mol	Chain	Residues	Atoms					AltConf	Trace
13	LI	218	Total	C	N	O	S	0	0
			1764	1117	334	306	7		

- Molecule 14 is a protein called 60S ribosomal protein L11-B.

Mol	Chain	Residues	Atoms					AltConf	Trace
14	LJ	169	Total	C	N	O	S	0	0
			1350	846	253	247	4		

- Molecule 15 is a protein called 60S ribosomal protein L13-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
15	LL	193	Total	C	N	O		0	0
			1543	962	315	266			

- Molecule 16 is a protein called 60S ribosomal protein L14-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
16	LM	136	Total	C	N	O	S	0	0
			1053	675	199	177	2		

- Molecule 17 is a protein called 60S ribosomal protein L15-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
17	LN	203	Total	C	N	O	S	0	0
			1720	1077	361	281	1		

- Molecule 18 is a protein called 60S ribosomal protein L16-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
18	LO	197	Total	C	N	O	S	197	0
			1555	1003	289	262	1		

- Molecule 19 is a protein called 60S ribosomal protein L17-A.

Mol	Chain	Residues	Atoms				AltConf	Trace
19	LP	183	Total	C	N	O	0	0
			1416	879	284	253		

- Molecule 20 is a protein called 60S ribosomal protein L18-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
20	LQ	185	Total	C	N	O	S	0	0
			1441	908	290	241	2		

- Molecule 21 is a protein called 60S ribosomal protein L19-A.

Mol	Chain	Residues	Atoms				AltConf	Trace
21	LR	171	Total	C	N	O	0	0
			1378	850	294	234		

- Molecule 22 is a protein called 60S ribosomal protein L20-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
22	LS	171	Total	C	N	O	S	0	0
			1437	925	266	243	3		

- Molecule 23 is a protein called 60S ribosomal protein L21-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
23	LT	159	Total	C	N	O	S	0	0
			1276	805	246	221	4		

- Molecule 24 is a protein called 60S ribosomal protein L22-A.

Mol	Chain	Residues	Atoms				AltConf	Trace
24	LU	100	Total	C	N	O	0	0
			796	516	131	149		

- Molecule 25 is a protein called 60S ribosomal protein L23-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
25	LV	136	Total	C	N	O	S	0	0
			1003	628	189	179	7		

- Molecule 26 is a protein called 60S ribosomal protein L24-A.



Mol	Chain	Residues	Atoms					AltConf	Trace
26	LW	67	Total	C	N	O	S	0	0
			538	345	106	86	1		

There are 4 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
LW	104	GLN	ASN	conflict	UNP P04449
LW	109	GLN	LEU	conflict	UNP P04449
LW	112	ASP	ASN	conflict	UNP P04449
LW	119	ALA	GLU	conflict	UNP P04449

- Molecule 27 is a protein called 60S ribosomal protein L25.

Mol	Chain	Residues	Atoms					AltConf	Trace
27	LX	121	Total	C	N	O	S	0	0
			964	620	169	173	2		

- Molecule 28 is a protein called 60S ribosomal protein L26-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
28	LY	125	Total	C	N	O		0	0
			984	620	191	173			

- Molecule 29 is a protein called 60S ribosomal protein L27-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
29	LZ	135	Total	C	N	O		0	0
			1092	710	202	180			

- Molecule 30 is a protein called 60S ribosomal protein L28.

Mol	Chain	Residues	Atoms					AltConf	Trace
30	La	148	Total	C	N	O	S	0	0
			1173	749	231	190	3		

- Molecule 31 is a protein called 60S ribosomal protein L29.

Mol	Chain	Residues	Atoms					AltConf	Trace
31	Lb	58	Total	C	N	O		0	0
			462	289	100	73			

- Molecule 32 is a protein called 60S ribosomal protein L30.

Mol	Chain	Residues	Atoms					AltConf	Trace
32	Lc	96	Total	C	N	O	S	0	0
			737	476	123	137	1		

- Molecule 33 is a protein called 60S ribosomal protein L31-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
33	Ld	109	Total	C	N	O	S	0	0
			880	559	168	152	1		

- Molecule 34 is a protein called 60S ribosomal protein L32.

Mol	Chain	Residues	Atoms					AltConf	Trace
34	Le	127	Total	C	N	O	S	0	0
			1017	644	205	167	1		

- Molecule 35 is a protein called 60S ribosomal protein L33-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
35	Lf	106	Total	C	N	O	S	0	0
			850	540	165	144	1		

- Molecule 36 is a protein called 60S ribosomal protein L34-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
36	Lg	112	Total	C	N	O	S	0	0
			880	545	179	152	4		

- Molecule 37 is a protein called 60S ribosomal protein L35-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
37	Lh	119	Total	C	N	O	S	0	0
			969	615	186	167	1		

- Molecule 38 is a protein called 60S ribosomal protein L36-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
38	Li	99	Total	C	N	O	S	0	0
			766	478	154	132	2		

- Molecule 39 is a protein called 60S ribosomal protein L37-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
39	Lj	85	Total	C	N	O	S	0	0
			670	408	146	111	5		

- Molecule 40 is a protein called 60S ribosomal protein L38.

Mol	Chain	Residues	Atoms					AltConf	Trace
40	Lk	77	Total	C	N	O		0	0
			612	391	115	106			

- Molecule 41 is a protein called 60S ribosomal protein L39.

Mol	Chain	Residues	Atoms					AltConf	Trace
41	Ll	50	Total	C	N	O	S	0	0
			436	272	97	65	2		

- Molecule 42 is a protein called 60S ribosomal protein L40-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
42	Lm	52	Total	C	N	O	S	0	0
			417	259	86	67	5		

- Molecule 43 is a protein called 60S ribosomal protein L41-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
43	Ln	25	Total	C	N	O	S	0	0
			229	139	62	27	1		

- Molecule 44 is a protein called 60S ribosomal protein L42-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
44	Lo	103	Total	C	N	O	S	0	0
			824	517	167	135	5		

- Molecule 45 is a protein called 60S ribosomal protein L43-A.

Mol	Chain	Residues	Atoms					AltConf	Trace
45	Lp	91	Total	C	N	O	S	0	0
			694	429	138	121	6		

- Molecule 46 is a RNA chain called 25S rRNA.

Mol	Chain	Residues	Atoms					AltConf	Trace
46	1	3301	Total 70586	C 31525	N 12690	O 23070	P 3301	0	0

There are 2 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
1	?	-	G	deletion	GB 1262303
1	1962	A	G	conflict	GB 1262303

- Molecule 47 is MAGNESIUM ION (three-letter code: MG) (formula: Mg).

Mol	Chain	Residues	Atoms		AltConf
47	C4	1	Total 1	Mg 1	0
47	C3	1	Total 1	Mg 1	0
47	LA	2	Total 2	Mg 2	0
47	LB	1	Total 1	Mg 1	0
47	LN	1	Total 1	Mg 1	0
47	LP	1	Total 1	Mg 1	0
47	LR	1	Total 1	Mg 1	0
47	LV	1	Total 1	Mg 1	0
47	La	1	Total 1	Mg 1	0
47	Le	1	Total 1	Mg 1	0
47	1	195	Total 195	Mg 195	0

- Molecule 48 is ZINC ION (three-letter code: ZN) (formula: Zn).

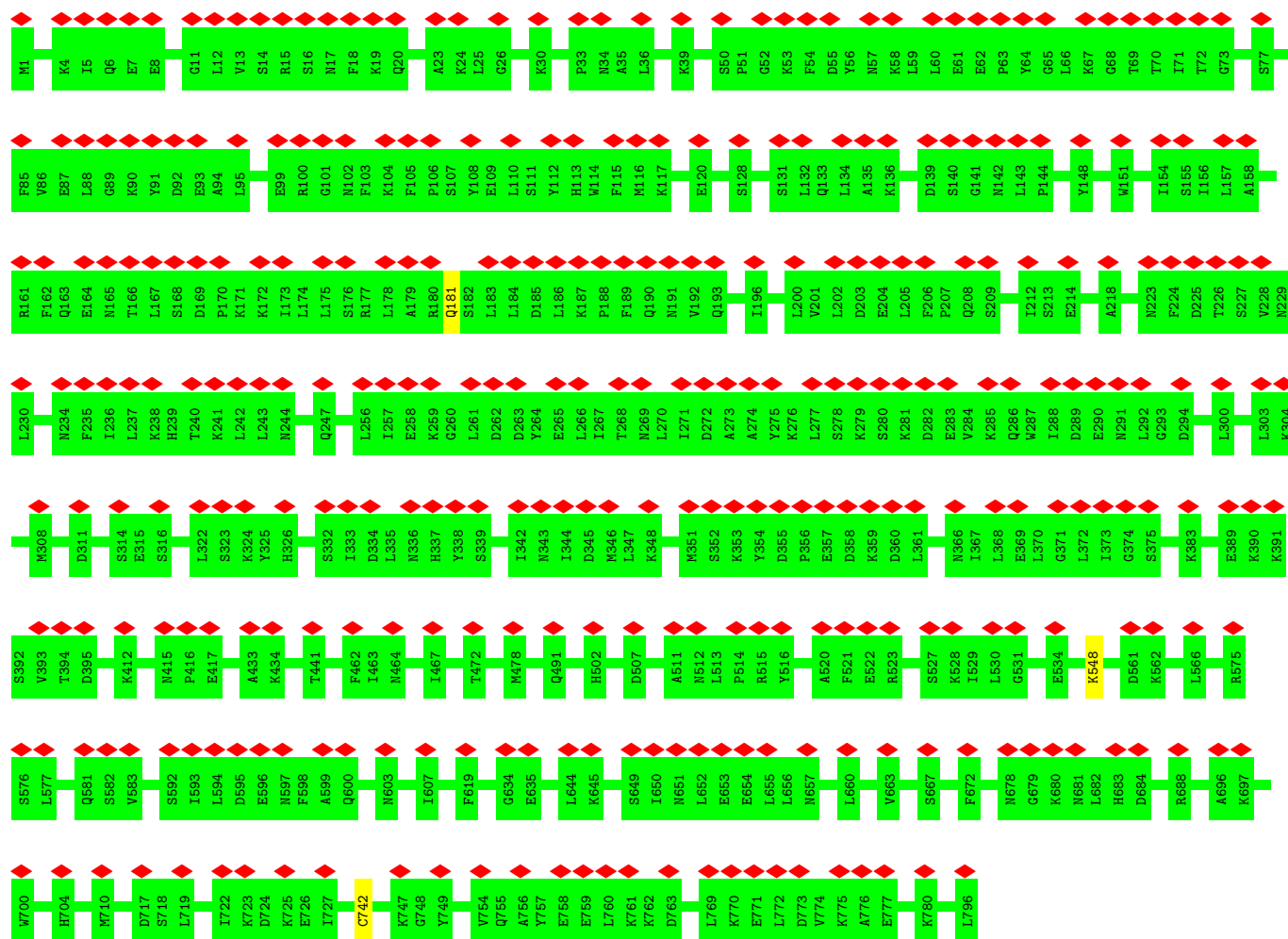
Mol	Chain	Residues	Atoms		AltConf
48	Lg	1	Total 1	Zn 1	0
48	Lj	1	Total 1	Zn 1	0

*Continued on next page...*

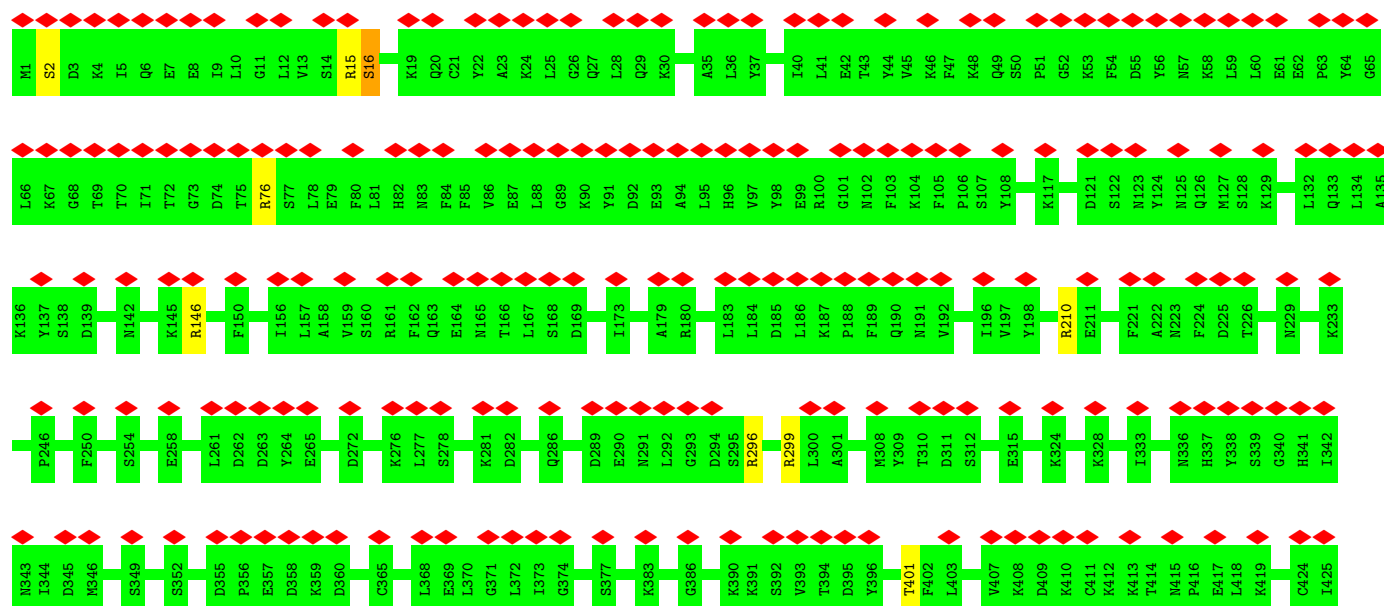
*Continued from previous page...*

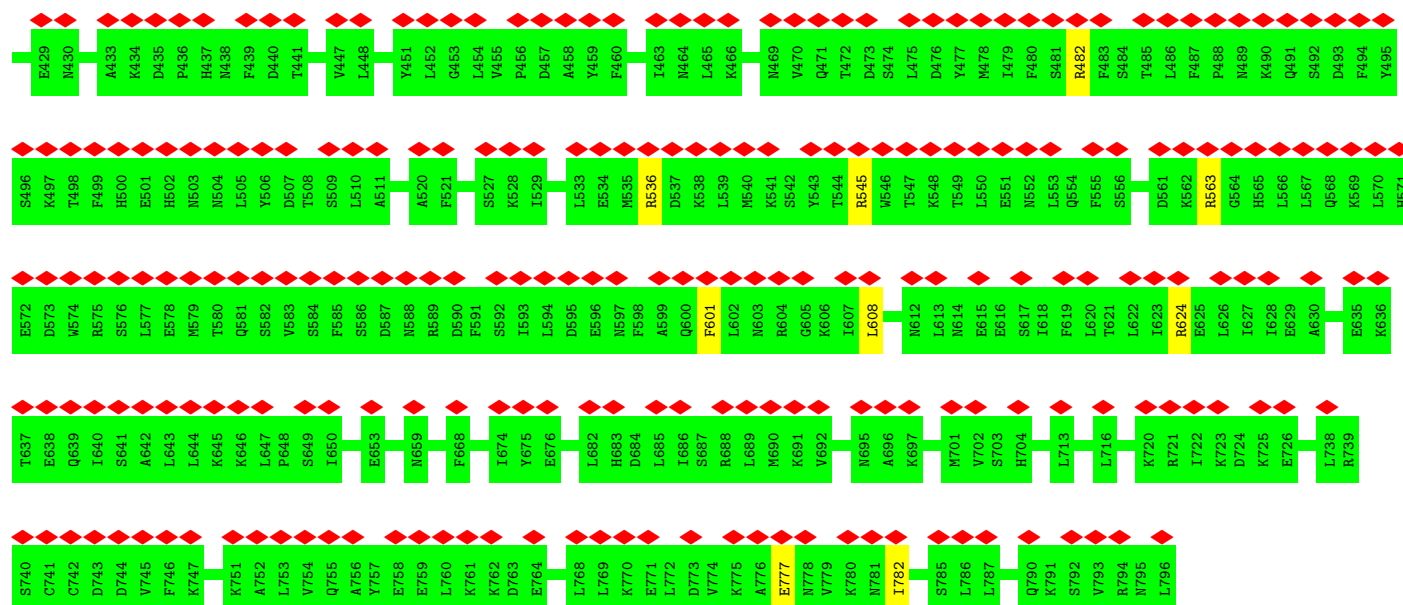
Mol	Chain	Residues	Atoms		AltConf
48	Lm	1	Total 1	Zn 1	0
48	Lo	1	Total 1	Zn 1	0
48	Lp	1	Total 1	Zn 1	0





• Molecule 2: N-terminal acetyltransferase B complex subunit MDM20





- Molecule 3: 5S rRNA

Chain C4: 90% 10%



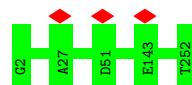
- Molecule 4: 5.8S rRNA

Chain C3: 84% 16%



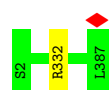
- Molecule 5: 60S ribosomal protein L2-A

Chain LA: 100%



- Molecule 6: 60S ribosomal protein L3

Chain LB: 100%



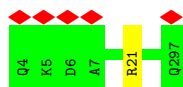
- Molecule 7: 60S ribosomal protein L4-A

Chain LC: 99%

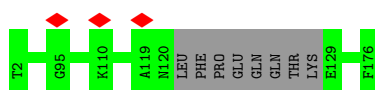




- Molecule 8: 60S ribosomal protein L5



- Molecule 9: 60S ribosomal protein L6-B



- Molecule 10: 60S ribosomal protein L7-A



There are no outlier residues recorded for this chain.

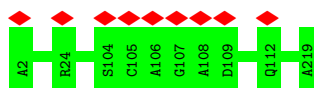
- Molecule 11: 60S ribosomal protein L8-A



- Molecule 12: 60S ribosomal protein L9-A

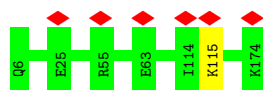


- Molecule 13: 60S ribosomal protein L10



- Molecule 14: 60S ribosomal protein L11-B





- Molecule 15: 60S ribosomal protein L13-A

Chain LL: 98%



- Molecule 16: 60S ribosomal protein L14-A

Chain LM: 99%



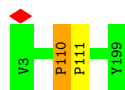
- Molecule 17: 60S ribosomal protein L15-A

Chain LN: 100%



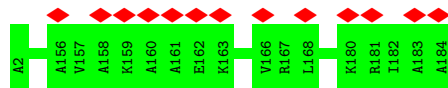
- Molecule 18: 60S ribosomal protein L16-A

Chain LO: 99%



- Molecule 19: 60S ribosomal protein L17-A

Chain LP: 7%

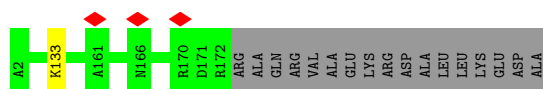


- Molecule 20: 60S ribosomal protein L18-A

Chain LQ: 99%



- Molecule 21: 60S ribosomal protein L19-A



- Molecule 22: 60S ribosomal protein L20-A



There are no outlier residues recorded for this chain.

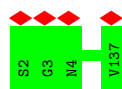
- Molecule 23: 60S ribosomal protein L21-A



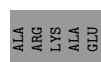
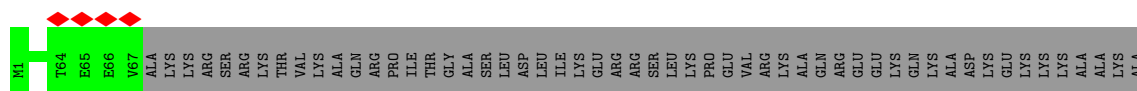
- Molecule 24: 60S ribosomal protein L22-A



- Molecule 25: 60S ribosomal protein L23-A



- Molecule 26: 60S ribosomal protein L24-A



- Molecule 27: 60S ribosomal protein L25





- Molecule 28: 60S ribosomal protein L26-A

Chain LY:  99%



- Molecule 29: 60S ribosomal protein L27-A

Chain LZ:  99%



- Molecule 30: 60S ribosomal protein L28

Chain La:  99%



- Molecule 31: 60S ribosomal protein L29

Chain Lb:  100%



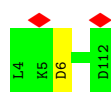
- Molecule 32: 60S ribosomal protein L30

Chain Lc:  99%



- Molecule 33: 60S ribosomal protein L31-A

Chain Ld:  99%



- Molecule 34: 60S ribosomal protein L32

Chain Le:  100%



- Molecule 35: 60S ribosomal protein L33-A

Chain Lf:  99%



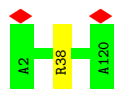
- Molecule 36: 60S ribosomal protein L34-A

Chain Lg:  100%

There are no outlier residues recorded for this chain.

- Molecule 37: 60S ribosomal protein L35-A

Chain Lh:  99%



- Molecule 38: 60S ribosomal protein L36-A

Chain Li:  100%



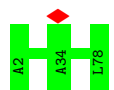
- Molecule 39: 60S ribosomal protein L37-A

Chain Lj:  100%

There are no outlier residues recorded for this chain.

- Molecule 40: 60S ribosomal protein L38

Chain Lk:  100%



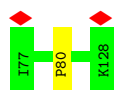
- Molecule 41: 60S ribosomal protein L39

Chain Ll:  100%

There are no outlier residues recorded for this chain.

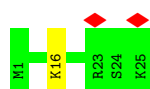
- Molecule 42: 60S ribosomal protein L40-A

Chain Lm:  98%



- Molecule 43: 60S ribosomal protein L41-A

Chain Ln:  96%



- Molecule 44: 60S ribosomal protein L42-A

Chain Lo:  100%




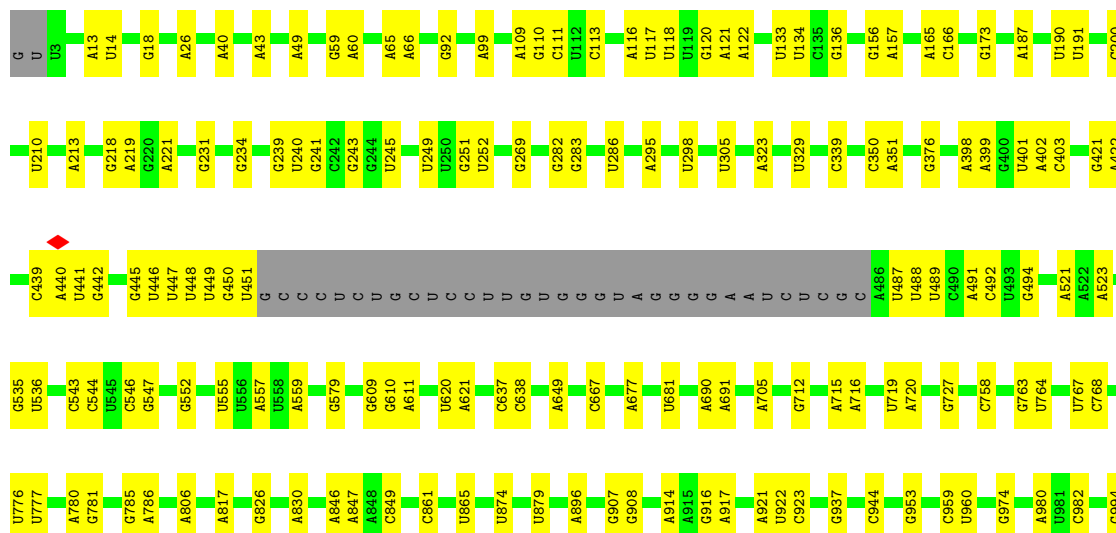
- Molecule 45: 60S ribosomal protein L43-A

Chain Lp:  100%



- Molecule 46: 25S rRNA

Chain 1:  79% 18%



G3101	C2821	G2614	A2402	A2188	U2056	G1979	C1788	U1572	U1325	C1192	A1002
U3104	U2843	C2638	G2403	C2192	U2059	C1980	A1797	G1573	A1330	A1193	G1010
A3113	C2844	U2652	A2404	U2193	A	G1981	G1808	C1574	U1348	C1196	U1015
A3114	A2845	A2656	U2411	G2194	G	G1982	U1814	G1575	G1349	C1201	G1016
C3115	G2856	A2666	A2419	G2206	U	G1983	U1815	A1580	A1350	C1201	C1017
U3119	C2867	G2672	G2435	A2207	U	G1984	U1816	C1581	U1351	U1208	G1018
C3120	U2867	A2673	U2436	A2208	C	G1985	A1817	A1582	A1352	G1217	G1019
U3121	G2871	A2674	G2437	U2209	C	U1986	U1818	C1583	U1353	G1217	G1020
A3122	U2872	G2677	A2438	G2210	U	U1989	U1819	A1587	G1354	G1222	G1024
A3129	U2873	U2681	A2439	A2222	U	U1990	U1820	A1588	U1355	G1222	A1025
A3130	G2874	U2682	G2440	A2223	G	U1991	U1821	A1589	U1356	C1227	A1026
U3131	U2875	A2683	A2441	A2224	A	G1992	U1822	U1599	G1357	C1232	A1027
A3139	A2887	G2689	C2442	A2244	C	G1993	U1823	U1606	A1386	C1232	U1028
A3142	U2888	U2690	U2443	G2249	A	G1994	U1824	U1607	U1389	U1235	C1032
C3143	C2889	A2691	A2444	G2250	C	U1619	U1825	U1607	G1400	G1236	G1035
U3151	A2911	A2694	G2445	G2255	C	U1620	U1826	U1607	G1417	G1236	A1036
U3152	U2923	A2695	G2446	A2255	U	U1629	U1827	U1607	G1418	U1241	A1047
U3153	G2923	A2696	G2447	A2270	C	U1639	U1828	U1607	A1419	G1242	A1048
C3154	C2929	A2697	A2448	G2271	G	U1642	U1829	U1607	G1434	A1245	C1049
U3155	U2935	A2698	G2449	G2272	C	A1643	U1830	U1607	C1437	G1246	G1063
U3156	A2936	A2699	G2450	G2273	C	U1644	U1831	U1607	U1446	U1253	A1064
U3157	G2947	A2700	G2451	G2274	C	U1645	U1832	U1607	G1450	C1254	A1065
G3158	U2971	A2701	G2452	G2275	C	U1657	U1833	U1607	U1481	C1255	G1072
A3165	C2971	U2737	U2537	G2288	A2088	C1687	U1834	U1607	A1482	U1258	A1080
A3170	C2983	U2752	U2538	G2307	A2089	A1683	U1835	U1607	G1483	G1262	U1081
G3173	G2990	U2753	U2539	U2310	A2093	U1717	U1836	U1607	U1484	A1263	A1093
A3174	A2991	G2753	U2540	A2313	C2094	U1724	U1837	U1607	G1488	G1264	U1094
U3175	U2992	C2772	U2541	G2314	C2101	C1725	U1838	U1607	U1495	U1269	U1095
G3176	U2996	C2773	U2542	G2315	U2102	G1736	U1839	U1607	C1496	A1270	U1096
U3179	U2997	G2777	U2543	U2318	A2107	U1741	U1840	U1607	U1508	A1271	G1097
A3180	C2997	G2778	U2544	G2334	G2111	A1750	U1841	U1607	U1523	A1274	A1098
C3181	G3012	C2788	U2545	G2335	G2112	G1751	U1842	U1607	U1556	A1278	A1103
A3187	G3030	G2789	U2546	U2336	A2113	G1761	U1843	U1607	A1557	C1279	G1104
U3196	U3056	G2796	U2547	G2337	G2114	U1762	U1844	U1607	G1560	C1280	G1117
U3207	G3059	G2800	U2548	G2373	G2121	C1763	U1845	U1607	G1561	A1279	G1131
A3210	U3078	A2801	U2549	G2375	G2122	U1764	U1846	U1607	C1562	G1283	U1144
G3217	G3080	A2802	U2550	G2385	A2126	U1765	U1847	U1607	U1563	G1285	U1159
A3218	U3079	A2803	U2551	G2386	A2131	G1766	U1848	U1607	U1564	C1286	G1178
G3219	A3086	A2804	U2552	U2388	A2144	U1767	U1849	U1607	A1565	A1287	A1179
A3227	C3092	C2810	U2553	G2393	C2039	G1770	U1850	U1607	A1566	G1307	A1180
C3228	C3093	A2816	U2554	G2394	U2040	U1771	U1851	U1607	A1567	U1309	U1181
G3229	G2807	A2817	U2555	A2397	U2043	G1775	U1852	U1607	U1568	U1315	
			U2556		U2044		U1853	U1607			
			U2557		U2045		U1854	U1607			
			U2558		U2046		U1855	U1607			
			U2559		U2047		U1856	U1607			
			U2560		U2048		U1857	U1607			
			U2561		U2049		U1858	U1607			
			U2562		U2050		U1859	U1607			
			U2563		U2051		U1860	U1607			
			U2564		U2052		U1861	U1607			
			U2565		U2053		U1862	U1607			
			U2566		U2054		U1863	U1607			
			U2567		U2055		U1864	U1607			
			U2568		U2056		U1865	U1607			
			U2569		U2057		U1866	U1607			
			U2570		U2058		U1867	U1607			
			U2571		U2059		U1868	U1607			
			U2572		U2060		U1869	U1607			
			U2573		U2061		U1870	U1607			
			U2574		U2062		U1871	U1607			
			U2575		U2063		U1872	U1607			
			U2576		U2064		U1873	U1607			
			U2577		U2065		U1874	U1607			
			U2578		U2066		U1875	U1607			
			U2579		U2067		U1876	U1607			
			U2580		U2068		U1877	U1607			
			U2581		U2069		U1878	U1607			
			U2582		U2070		U1879	U1607			
			U2583		U2071		U1880	U1607			
			U2584		U2072		U1881	U1607			
			U2585		U2073		U1882	U1607			
			U2586		U2074		U1883	U1607			
			U2587		U2075		U1884	U1607			
			U2588		U2076		U1885	U1607			
			U2589		U2077		U1886	U1607			
			U2590		U2078		U1887	U1607			
			U2591		U2079		U1888	U1607			
			U2592		U2080		U1889	U1607			
			U2593		U2081		U1890	U1607			
			U2594		U2082		U1891	U1607			
			U2595		U2083		U1892	U1607			
			U2596		U2084		U1893	U1607			
			U2597		U2085		U1894	U1607			
			U2598		U2086		U1895	U1607			
			U2599		U2087		U1896	U1607			
			U2600		U2088		U1897	U1607			
			U2601		U2089		U1898	U1607			
			U2602		U2090		U1899	U1607			
			U2603		U2091		U1900	U1607			
			U2604		U2092		U1901	U1607			
			U2605		U2093		U1902	U1607			
			U2606		U2094		U1903	U1607			
			U2607		U2095		U1904	U1607			
			U2608		U2096		U1905	U1607			
			U2609		U2097		U1906	U1607			
			U2610		U2098		U1907	U1607			
			U2611		U2099		U1908	U1607			
			U2612		U2100		U1909	U1607			
			U2613		U2101		U1910	U1607			
			U2614		U2102		U1911	U1607			
			U2615		U2103		U1912	U1607			
			U2616		U2104		U1913	U1607			
			U2617		U2105		U1914	U1607			
			U2618		U2106		U1915	U1607			
			U2619		U2107		U1916	U1607			
			U2620		U2108		U1917	U1607			
			U2621		U2109		U1918	U1607			
			U2622		U2110		U1919	U1607			
			U2623		U2111		U1920	U1607			
			U2624		U2112		U1921	U1607			
			U2625		U2113		U1922	U1607			
			U2626		U2114		U1923	U1607			
			U2627		U2115		U1924	U1607			
			U2628		U2116		U1925	U1607			
			U2629		U2117		U1926	U1607			
			U2630		U2118		U1927	U1607			
			U2631		U2119		U1928	U1607			
			U2632		U2120		U1929	U1607			
			U2633		U2121		U1930	U1607			
			U2634		U2122		U1931	U1607			
			U2635		U2123		U1932	U1607			
			U2636		U2124		U1933	U1607			
			U2637		U2125		U1934	U1607			
			U2638		U2126		U1935	U1607			
			U2639		U2127		U1936	U1607			
			U2640		U2128		U1937	U1607			
			U2641		U2129		U1938	U1607			
			U2642		U2130		U1939	U1607			
			U2643		U2131		U1940	U1607			
			U2644		U2132		U1941	U1607			
			U2645		U2133		U1942	U1607			
			U2646		U2134		U1943	U1607			
			U2647		U2135		U1944	U1607			
			U2648		U2136		U1945	U1607			
			U2649		U2137		U1946	U1607			

G3239	
G3242	
A3243	
A3244	
A3245	
G3246	
G3247	
U3259	
G3263	
U3269	
U3270	
A3273	
A3274	
U3275	
G3276	
U3281	
U3287	
G3288	
G3289	
A3294	
A3295	
U3304	
U3313	
A3316	
U3317	
G3318	
U3319	
A3320	
U3334	
U3341	
G3345	
C3350	
U3351	
U3352	
G3353	
U3354	
U3355	
G3369	
A3375	
C3378	
G3386	

U3389	
G3390	
U3396	



## 4 Experimental information

Property	Value	Source
EM reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, Not provided	
Number of particles used	9645	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 ( $e^-/\text{\AA}^2$ )	45.2	Depositor
Minimum defocus (nm)	500	Depositor
Maximum defocus (nm)	3500	Depositor
Magnification	Not provided	
Image detector	GATAN K2 SUMMIT (4k x 4k)	Depositor
Maximum map value	1.382	Depositor
Minimum map value	-0.498	Depositor
Average map value	0.011	Depositor
Map value standard deviation	0.093	Depositor
Recommended contour level	0.2	Depositor
Map size (Å)	417.99997, 417.99997, 417.99997	wwPDB
Map dimensions	400, 400, 400	wwPDB
Map angles (°)	90.0, 90.0, 90.0	wwPDB
Pixel spacing (Å)	1.045, 1.045, 1.045	Depositor

## 5 Model quality ⓘ

### 5.1 Standard geometry ⓘ

Bond lengths and bond angles in the following residue types are not validated in this section: MG, ZN

The Z score for a bond length (or angle) is the number of standard deviations the observed value is removed from the expected value. A bond length (or angle) with  $|Z| > 5$  is considered an outlier worth inspection. RMSZ is the root-mean-square of all Z scores of the bond lengths (or angles).

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	$\# Z  > 5$	RMSZ	$\# Z  > 5$
1	A	0.26	0/1654	0.56	1/2237 (0.0%)
1	C	0.67	0/589	0.82	0/800
2	B	0.25	0/6665	0.47	0/8979
2	D	0.64	0/6665	0.86	14/8979 (0.2%)
3	C4	0.37	0/2883	0.74	0/4491
4	C3	0.43	0/3746	0.77	0/5832
5	LA	0.31	0/1933	0.58	0/2598
6	LB	0.30	0/3146	0.53	0/4228
7	LC	0.30	0/2800	0.53	0/3790
8	LD	0.28	0/2400	0.52	0/3239
9	LE	0.29	0/1327	0.51	0/1790
10	LF	0.31	0/1821	0.50	0/2451
11	LG	0.29	0/1836	0.51	0/2481
12	LH	0.30	0/1529	0.55	0/2060
13	LI	0.29	0/1801	0.54	0/2416
14	LJ	0.28	0/1371	0.60	0/1838
15	LL	0.30	0/1568	0.62	0/2106
16	LM	0.29	0/1068	0.54	0/1438
17	LN	0.31	0/1757	0.59	0/2354
18	LO	0.31	0/1585	0.52	0/2128
19	LP	0.30	0/1439	0.56	0/1938
20	LQ	0.29	0/1465	0.56	0/1965
21	LR	0.29	0/1395	0.57	0/1861
22	LS	0.33	0/1473	0.60	0/1980
23	LT	0.31	0/1300	0.55	0/1743
24	LU	0.30	0/812	0.53	0/1099
25	LV	0.31	0/1018	0.57	0/1369
26	LW	0.31	0/550	0.58	0/731
27	LX	0.31	0/979	0.51	0/1321
28	LY	0.31	0/995	0.62	1/1329 (0.1%)
29	LZ	0.32	0/1118	0.55	0/1497
30	La	0.30	0/1204	0.54	0/1612

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	# Z  >5	RMSZ	# Z  >5
31	Lb	0.28	0/473	0.54	0/629
32	Lc	0.32	0/745	0.51	0/1001
33	Ld	0.31	0/894	0.58	0/1200
34	Le	0.29	0/1038	0.54	0/1390
35	Lf	0.33	0/868	0.57	0/1168
36	Lg	0.31	0/890	0.59	0/1189
37	Lh	0.29	0/978	0.54	0/1301
38	Li	0.28	0/772	0.56	0/1026
39	Lj	0.31	0/685	0.59	0/908
40	Lk	0.29	0/618	0.56	0/826
41	Ll	0.30	0/443	0.66	0/588
42	Lm	0.30	0/423	0.63	0/562
43	Ln	0.26	0/230	0.82	0/296
44	Lo	0.30	0/836	0.57	0/1104
45	Lp	0.33	0/701	0.64	0/934
46	1	0.42	2/79000 (0.0%)	0.80	26/123166 (0.0%)
All	All	0.39	2/151486 (0.0%)	0.72	42/221968 (0.0%)

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

Mol	Chain	#Chirality outliers	#Planarity outliers
7	LC	0	1
11	LG	0	2
15	LL	0	1
18	LO	0	1
All	All	0	5

All (2) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
46	1	2094	C	O3'-P	-16.91	1.40	1.61
46	1	1950	U	O3'-P	-14.84	1.43	1.61

The worst 5 of 42 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
46	1	1950	U	O3'-P-O5'	27.53	156.31	104.00
46	1	1950	U	OP2-P-O3'	-12.81	77.02	105.20
46	1	1950	U	P-O3'-C3'	-11.39	106.03	119.70

*Continued on next page...*

*Continued from previous page...*

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
46	1	1950	U	OP1-P-O3'	-7.96	87.69	105.20
28	LY	53	ASP	CB-CG-OD1	7.33	124.90	118.30

There are no chirality outliers.

All (5) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
7	LC	13	GLY	Peptide
11	LG	158	ASP	Peptide
11	LG	76	ALA	Peptide
15	LL	75	PHE	Peptide
18	LO	110[A]	PRO	Peptide

## 5.2 Too-close contacts [i](#)

Due to software issues we are unable to calculate clashes - this section is therefore empty.

## 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	A	193/195 (99%)	192 (100%)	1 (0%)	0	100	100
1	C	66/195 (34%)	66 (100%)	0	0	100	100
2	B	794/796 (100%)	790 (100%)	3 (0%)	1 (0%)	51	83
2	D	794/796 (100%)	770 (97%)	22 (3%)	2 (0%)	41	74
5	LA	249/251 (99%)	234 (94%)	15 (6%)	0	100	100
6	LB	384/386 (100%)	358 (93%)	26 (7%)	0	100	100
7	LC	359/361 (99%)	329 (92%)	28 (8%)	2 (1%)	25	62
8	LD	292/294 (99%)	274 (94%)	17 (6%)	1 (0%)	41	74
9	LE	163/175 (93%)	155 (95%)	8 (5%)	0	100	100

*Continued on next page...*

*Continued from previous page...*

Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
10	LF	220/222 (99%)	205 (93%)	15 (7%)	0	100	100
11	LG	231/233 (99%)	221 (96%)	10 (4%)	0	100	100
12	LH	189/191 (99%)	176 (93%)	13 (7%)	0	100	100
13	LI	216/218 (99%)	208 (96%)	8 (4%)	0	100	100
14	LJ	167/169 (99%)	152 (91%)	15 (9%)	0	100	100
15	LL	191/193 (99%)	174 (91%)	15 (8%)	2 (1%)	15	52
16	LM	134/136 (98%)	129 (96%)	5 (4%)	0	100	100
17	LN	201/203 (99%)	191 (95%)	10 (5%)	0	100	100
18	LO	195/197 (99%)	185 (95%)	8 (4%)	2 (1%)	15	52
19	LP	181/183 (99%)	171 (94%)	10 (6%)	0	100	100
20	LQ	183/185 (99%)	176 (96%)	7 (4%)	0	100	100
21	LR	169/188 (90%)	163 (96%)	6 (4%)	0	100	100
22	LS	169/171 (99%)	157 (93%)	12 (7%)	0	100	100
23	LT	157/159 (99%)	148 (94%)	9 (6%)	0	100	100
24	LU	98/100 (98%)	95 (97%)	3 (3%)	0	100	100
25	LV	134/136 (98%)	132 (98%)	2 (2%)	0	100	100
26	LW	65/126 (52%)	61 (94%)	4 (6%)	0	100	100
27	LX	119/121 (98%)	116 (98%)	3 (2%)	0	100	100
28	LY	123/125 (98%)	117 (95%)	6 (5%)	0	100	100
29	LZ	133/135 (98%)	123 (92%)	10 (8%)	0	100	100
30	La	146/148 (99%)	133 (91%)	13 (9%)	0	100	100
31	Lb	56/58 (97%)	52 (93%)	4 (7%)	0	100	100
32	Lc	94/96 (98%)	93 (99%)	1 (1%)	0	100	100
33	Ld	107/109 (98%)	99 (92%)	7 (6%)	1 (1%)	17	54
34	Le	125/127 (98%)	120 (96%)	5 (4%)	0	100	100
35	Lf	104/106 (98%)	100 (96%)	4 (4%)	0	100	100
36	Lg	110/112 (98%)	106 (96%)	4 (4%)	0	100	100
37	Lh	117/119 (98%)	110 (94%)	7 (6%)	0	100	100
38	Li	97/99 (98%)	92 (95%)	5 (5%)	0	100	100
39	Lj	83/85 (98%)	80 (96%)	3 (4%)	0	100	100
40	Lk	75/77 (97%)	74 (99%)	1 (1%)	0	100	100

*Continued on next page...*

Continued from previous page...

Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
41	Ll	48/50 (96%)	48 (100%)	0	0	100	100
42	Lm	50/52 (96%)	47 (94%)	2 (4%)	1 (2%)	7	41
43	Ln	23/25 (92%)	23 (100%)	0	0	100	100
44	Lo	101/103 (98%)	96 (95%)	5 (5%)	0	100	100
45	Lp	89/91 (98%)	88 (99%)	1 (1%)	0	100	100
All	All	7994/8297 (96%)	7629 (95%)	353 (4%)	12 (0%)	50	79

5 of 12 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
2	B	742	CYS
7	LC	293	SER
33	Ld	6	ASP
2	D	16	SER
2	D	777	GLU

### 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	A	180/180 (100%)	178 (99%)	2 (1%)	73	85
1	C	66/180 (37%)	64 (97%)	2 (3%)	41	66
2	B	743/743 (100%)	741 (100%)	2 (0%)	92	96
2	D	743/743 (100%)	738 (99%)	5 (1%)	84	91
5	LA	190/193 (98%)	190 (100%)	0	100	100
6	LB	318/322 (99%)	317 (100%)	1 (0%)	92	96
7	LC	288/288 (100%)	288 (100%)	0	100	100
8	LD	241/243 (99%)	241 (100%)	0	100	100
9	LE	138/153 (90%)	138 (100%)	0	100	100
10	LF	186/186 (100%)	186 (100%)	0	100	100
11	LG	187/191 (98%)	187 (100%)	0	100	100

Continued on next page...

*Continued from previous page...*

Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
12	LH	168/171 (98%)	167 (99%)	1 (1%)	86	92
13	LI	185/185 (100%)	185 (100%)	0	100	100
14	LJ	146/147 (99%)	145 (99%)	1 (1%)	84	91
15	LL	154/154 (100%)	153 (99%)	1 (1%)	86	92
16	LM	107/107 (100%)	106 (99%)	1 (1%)	78	88
17	LN	175/175 (100%)	175 (100%)	0	100	100
18	LO	160/160 (100%)	160 (100%)	0	100	100
19	LP	138/145 (95%)	138 (100%)	0	100	100
20	LQ	150/150 (100%)	149 (99%)	1 (1%)	84	91
21	LR	139/153 (91%)	138 (99%)	1 (1%)	84	91
22	LS	155/155 (100%)	155 (100%)	0	100	100
23	LT	136/136 (100%)	135 (99%)	1 (1%)	84	91
24	LU	87/87 (100%)	87 (100%)	0	100	100
25	LV	104/104 (100%)	104 (100%)	0	100	100
26	LW	54/107 (50%)	54 (100%)	0	100	100
27	LX	104/105 (99%)	104 (100%)	0	100	100
28	LY	108/108 (100%)	108 (100%)	0	100	100
29	LZ	115/115 (100%)	114 (99%)	1 (1%)	78	88
30	La	118/118 (100%)	117 (99%)	1 (1%)	81	89
31	Lb	46/46 (100%)	46 (100%)	0	100	100
32	Lc	81/81 (100%)	80 (99%)	1 (1%)	71	84
33	Ld	93/96 (97%)	93 (100%)	0	100	100
34	Le	108/109 (99%)	108 (100%)	0	100	100
35	Lf	90/90 (100%)	89 (99%)	1 (1%)	73	85
36	Lg	95/95 (100%)	95 (100%)	0	100	100
37	Lh	104/104 (100%)	103 (99%)	1 (1%)	76	86
38	Li	80/81 (99%)	80 (100%)	0	100	100
39	Lj	69/69 (100%)	69 (100%)	0	100	100
40	Lk	68/68 (100%)	68 (100%)	0	100	100
41	Ll	45/45 (100%)	45 (100%)	0	100	100
42	Lm	47/47 (100%)	47 (100%)	0	100	100

*Continued on next page...*

*Continued from previous page...*

Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
43	Ln	22/23 (96%)	21 (96%)	1 (4%)	27	57
44	Lo	87/88 (99%)	87 (100%)	0	100	100
45	Lp	71/71 (100%)	71 (100%)	0	100	100
All	All	6889/7117 (97%)	6864 (100%)	25 (0%)	91	95

5 of 25 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
32	Lc	22	LYS
43	Ln	16	LYS
2	D	782	ILE
37	Lh	38	ARG
1	C	43	PHE

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 23 such sidechains are listed below:

Mol	Chain	Res	Type
22	LS	8	GLN
30	La	62	HIS
25	LV	98	ASN
30	La	64	GLN
2	B	469	ASN

### 5.3.3 RNA ⓘ

Mol	Chain	Analysed	Backbone Outliers	Pucker Outliers
3	C4	120/121 (99%)	11 (9%)	1 (0%)
4	C3	157/158 (99%)	26 (16%)	1 (0%)
46	1	3297/3395 (97%)	609 (18%)	39 (1%)
All	All	3574/3674 (97%)	646 (18%)	41 (1%)

5 of 646 RNA backbone outliers are listed below:

Mol	Chain	Res	Type
3	C4	7	G
3	C4	11	A
3	C4	53	U
3	C4	54	U
3	C4	55	A



5 of 41 RNA pucker outliers are listed below:

Mol	Chain	Res	Type
46	1	2537	U
46	1	3228	C
46	1	2541	U
46	1	3154	C
46	1	3275	U

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

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

## 5.5 Carbohydrates [i](#)

There are no monosaccharides in this entry.

## 5.6 Ligand geometry [i](#)

Of 211 ligands modelled in this entry, 211 are monoatomic - leaving 0 for Mogul analysis.

There are no bond length outliers.

There are no bond angle outliers.

There are no chirality outliers.

There are no torsion outliers.

There are no ring outliers.

No monomer is involved in short contacts.

## 5.7 Other polymers [i](#)

There are no such residues in this entry.

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

There are no chain breaks in this entry.

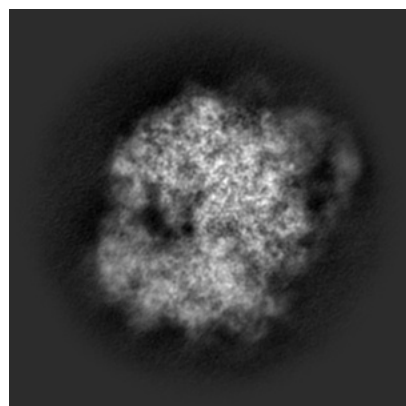
## 6 Map visualisation [i](#)

This section contains visualisations of the EMDB entry EMD-16090. These allow visual inspection of the internal detail of the map and identification of artifacts.

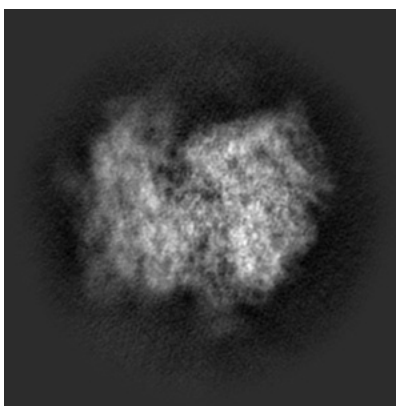
Images derived from a raw map, generated by summing the deposited half-maps, are presented below the corresponding image components of the primary map to allow further visual inspection and comparison with those of the primary map.

### 6.1 Orthogonal projections [i](#)

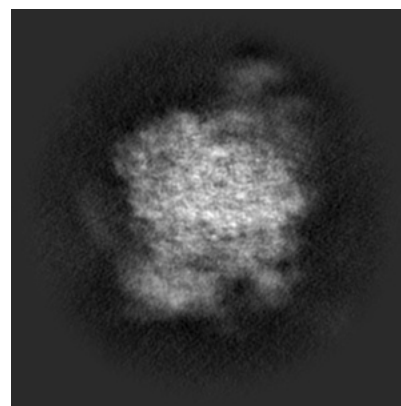
#### 6.1.1 Primary map



X

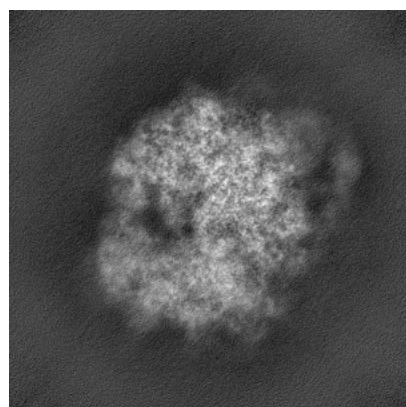


Y

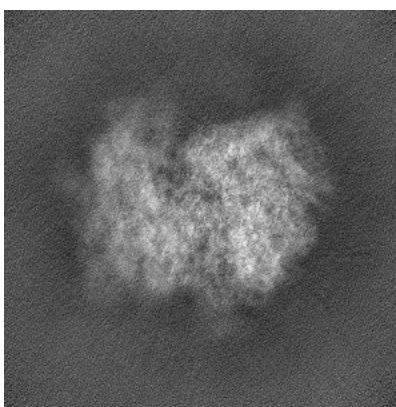


Z

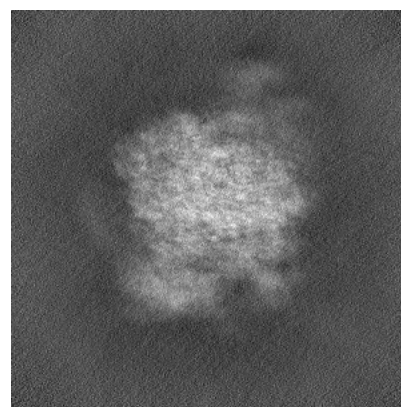
#### 6.1.2 Raw map



X



Y

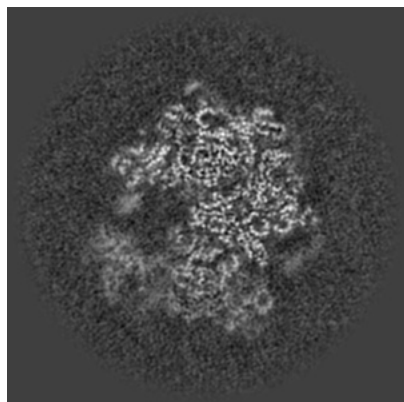


Z

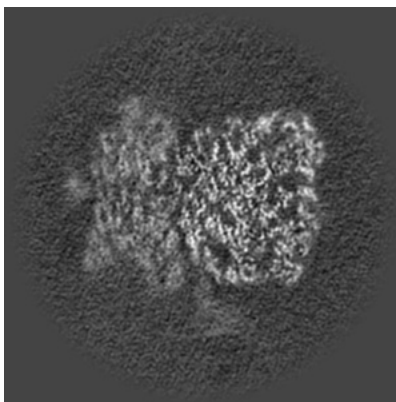
The images above show the map projected in three orthogonal directions.

## 6.2 Central slices [i](#)

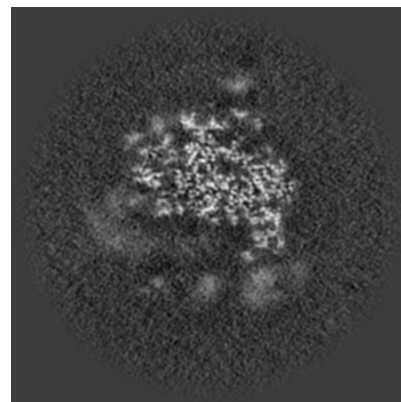
### 6.2.1 Primary map



X Index: 200

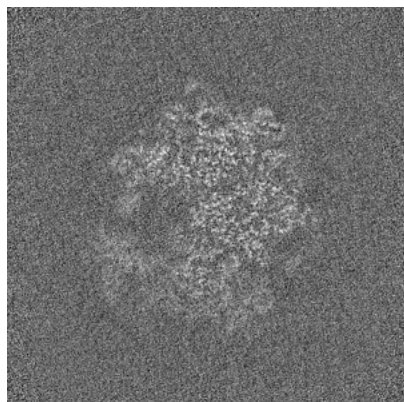


Y Index: 200

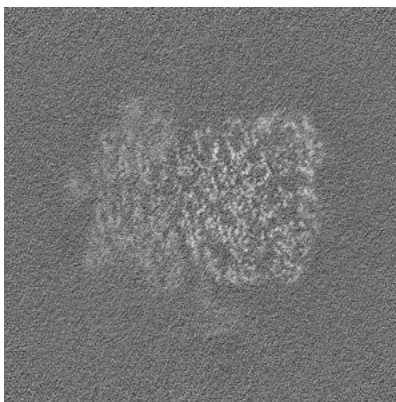


Z Index: 200

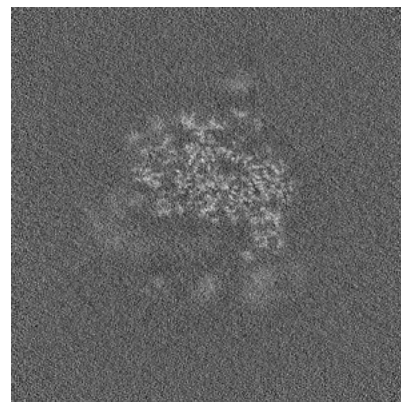
### 6.2.2 Raw map



X Index: 200



Y Index: 200

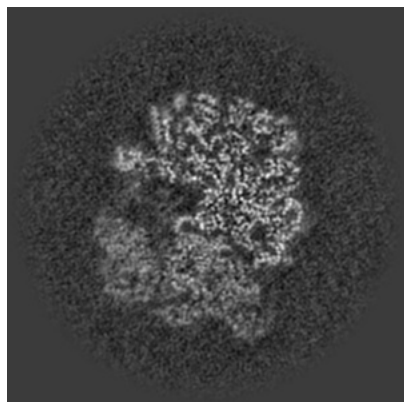


Z Index: 200

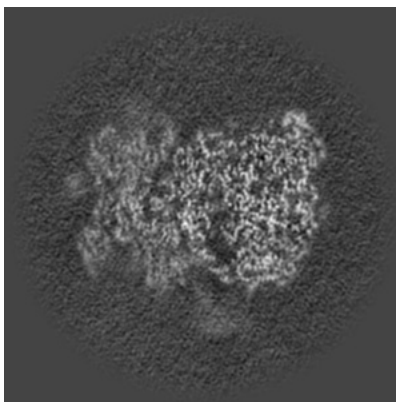
The images above show central slices of the map in three orthogonal directions.

## 6.3 Largest variance slices [i](#)

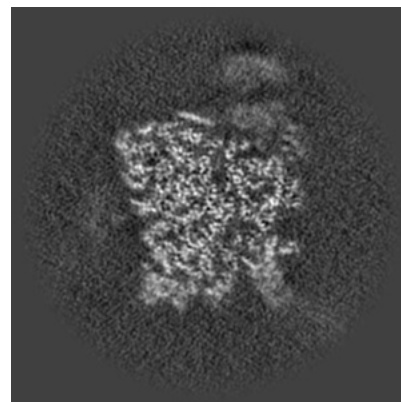
### 6.3.1 Primary map



X Index: 182

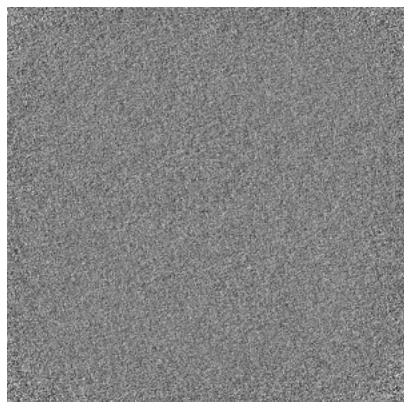


Y Index: 194

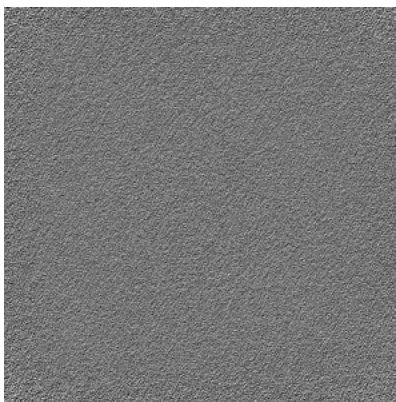


Z Index: 237

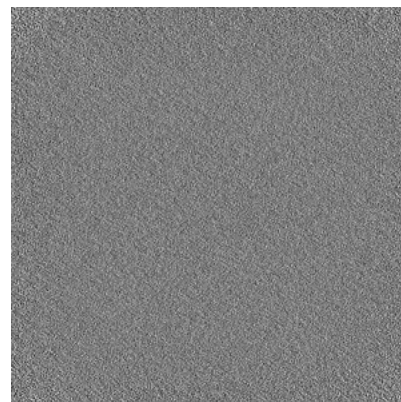
### 6.3.2 Raw map



X Index: 0



Y Index: 0



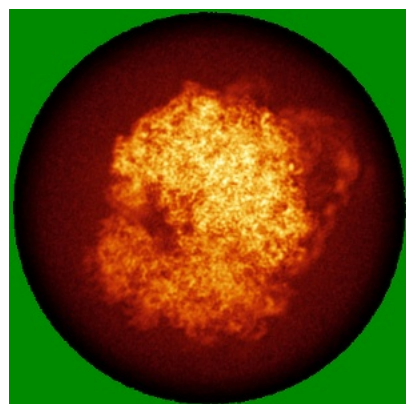
Z Index: 0

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

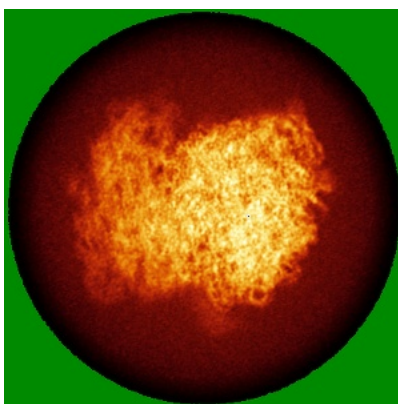


## 6.4 Orthogonal standard-deviation projections (False-color) [i](#)

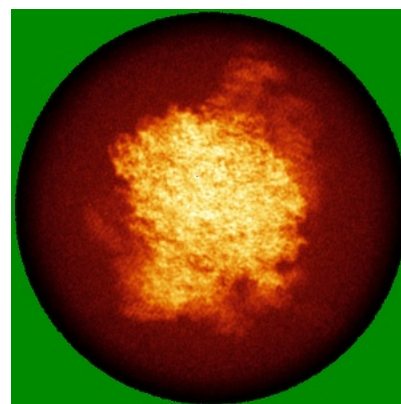
### 6.4.1 Primary map



X

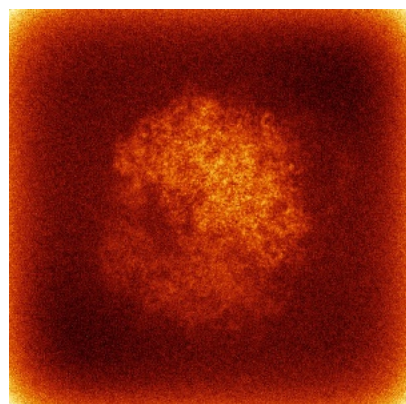


Y

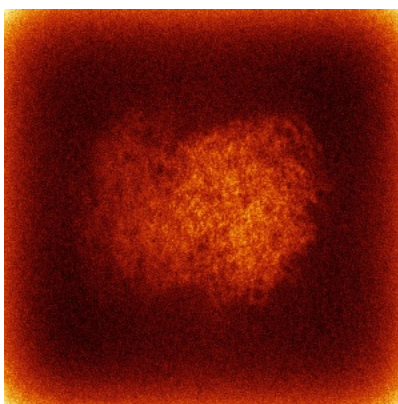


Z

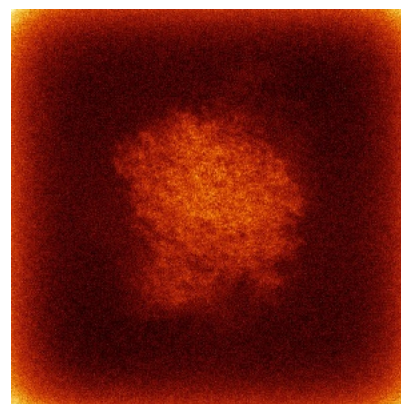
### 6.4.2 Raw map



X



Y

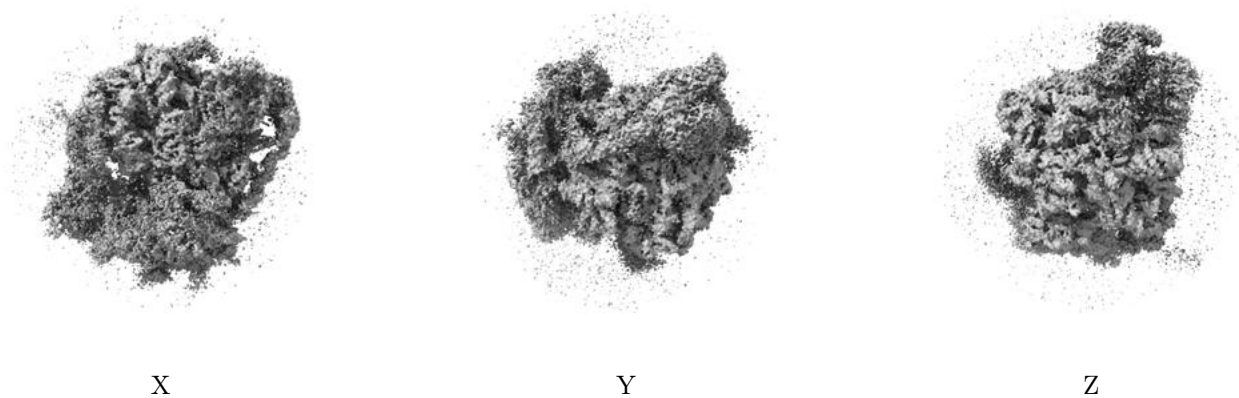


Z

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.

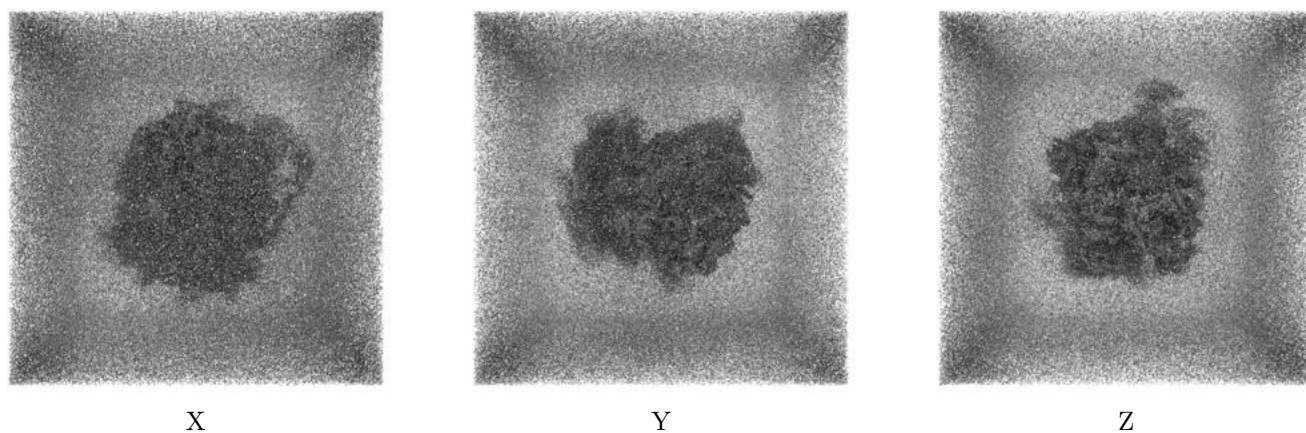
## 6.5 Orthogonal surface views [i](#)

### 6.5.1 Primary map



The images above show the 3D surface view of the map at the recommended contour level 0.2. These images, in conjunction with the slice images, may facilitate assessment of whether an appropriate contour level has been provided.

### 6.5.2 Raw map



These images show the 3D surface of the raw map. The raw map's contour level was selected so that its surface encloses the same volume as the primary map does at its recommended contour level.

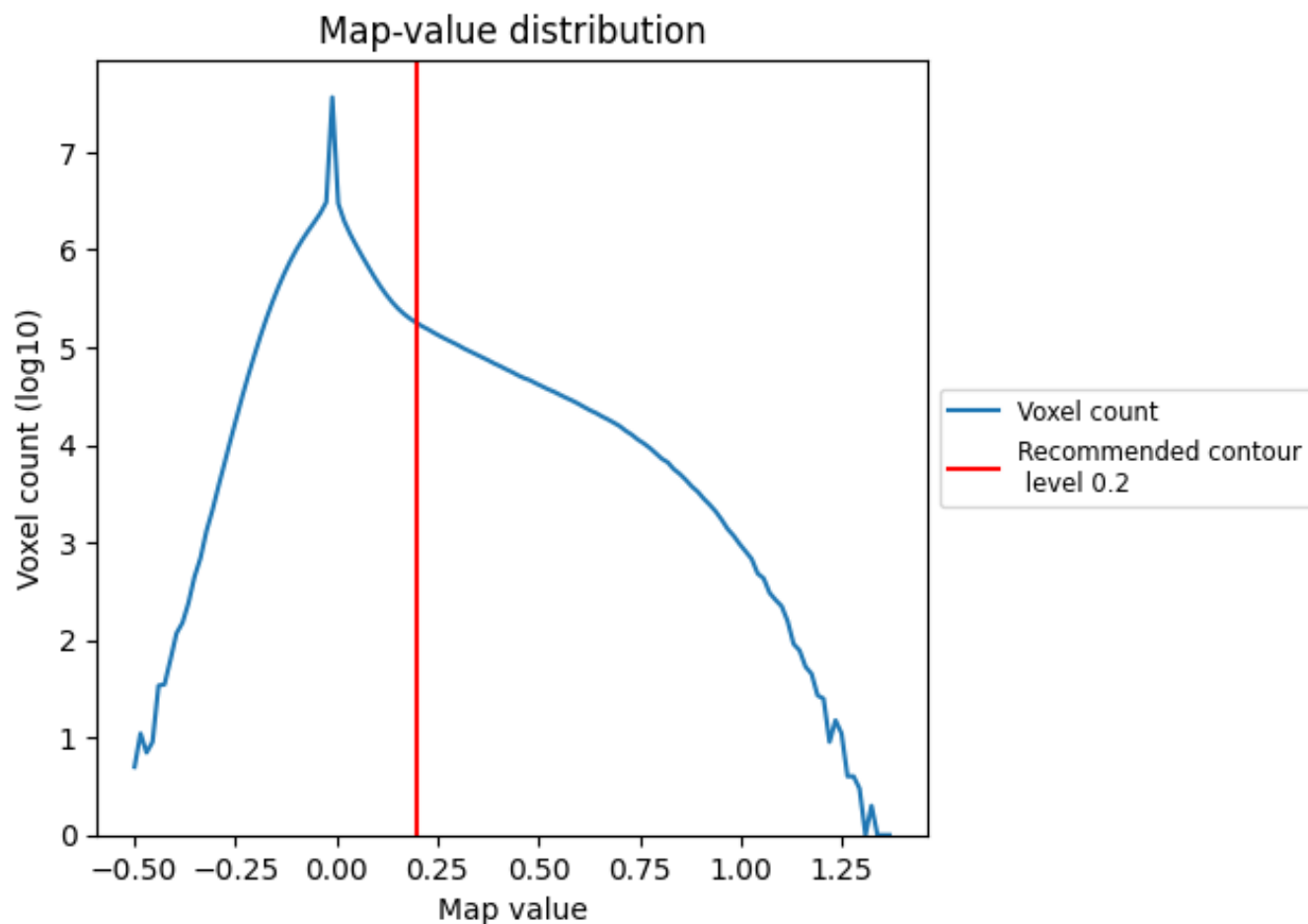
## 6.6 Mask visualisation [i](#)

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

## 7 Map analysis [i](#)

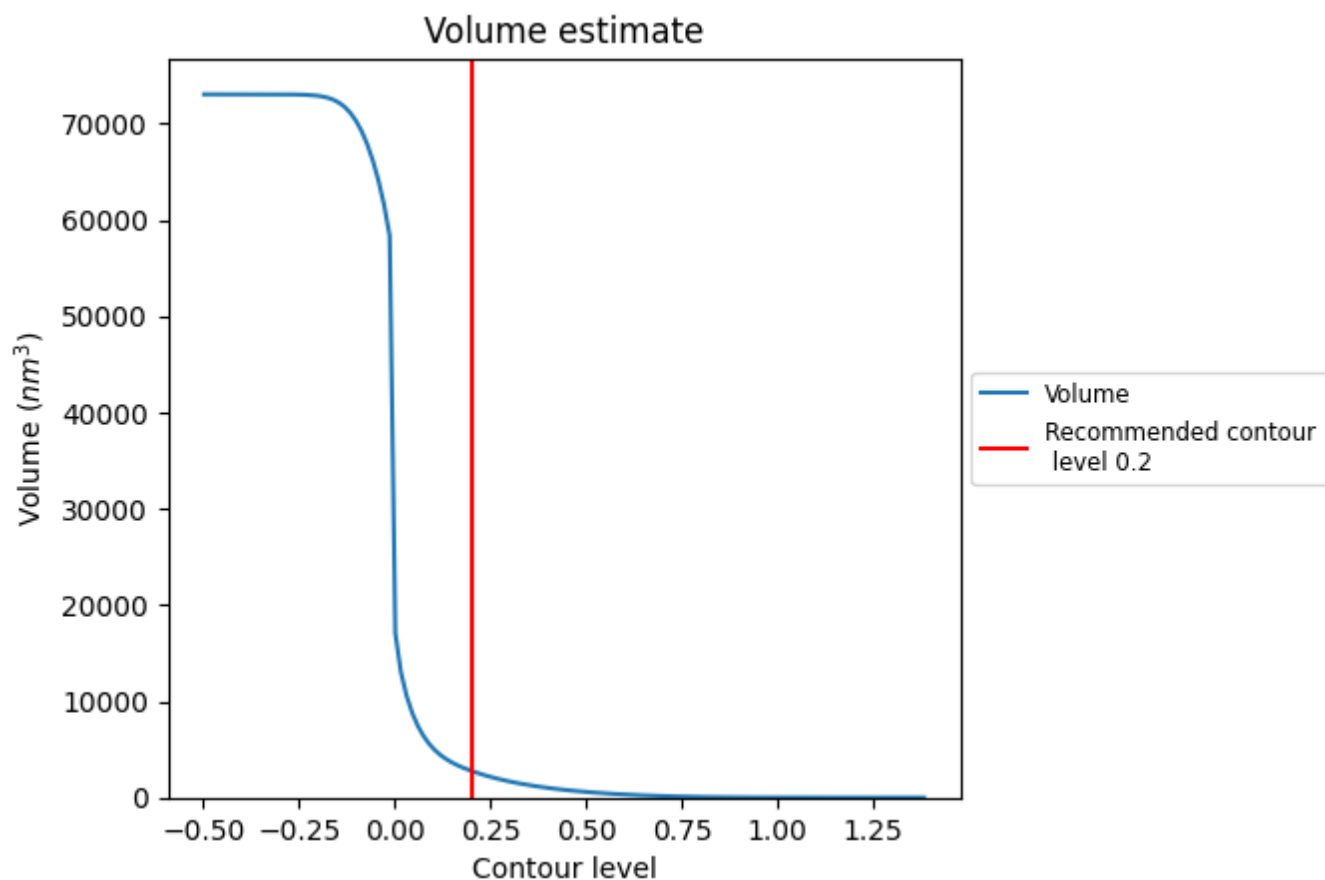
This section contains the results of statistical analysis of the map.

### 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](#)

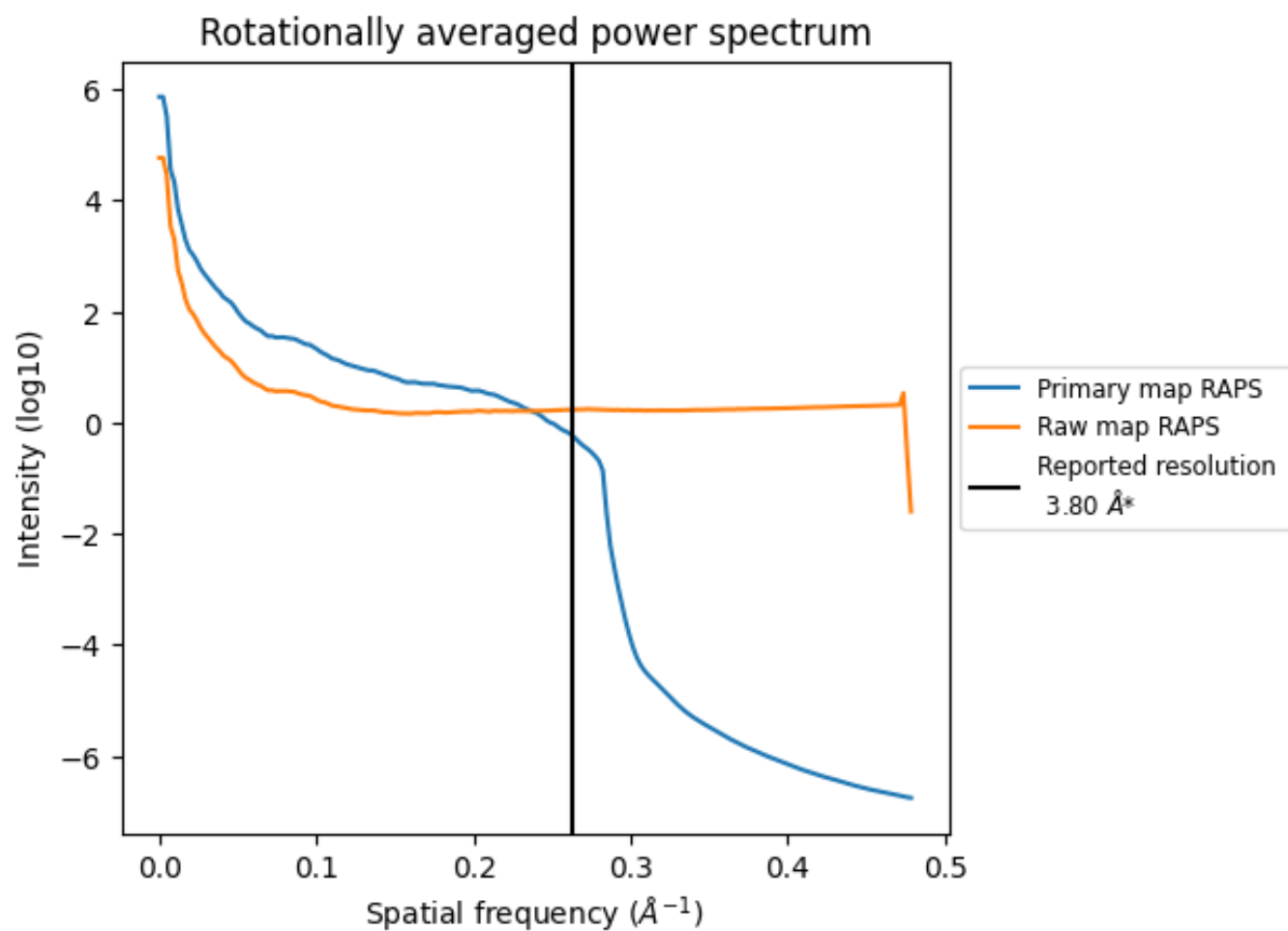


The volume at the recommended contour level is 2769 nm<sup>3</sup>; this corresponds to an approximate mass of 2501 kDa.

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 ⓘ

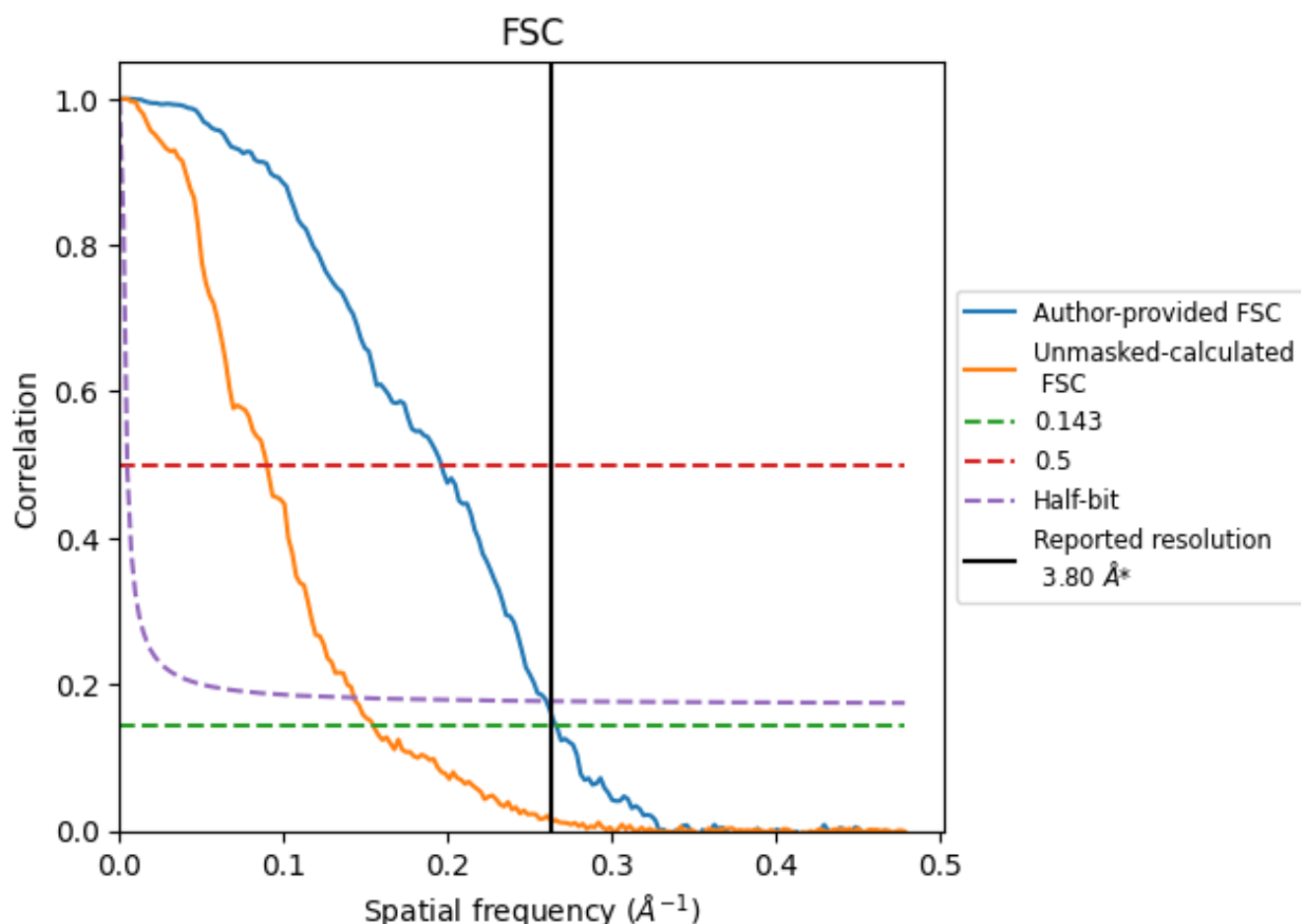


\*Reported resolution corresponds to spatial frequency of 0.263 Å<sup>-1</sup>

## 8 Fourier-Shell correlation [i](#)

Fourier-Shell Correlation (FSC) is the most commonly used method to estimate the resolution of single-particle and subtomogram-averaged maps. The shape of the curve depends on the imposed symmetry, mask and whether or not the two 3D reconstructions used were processed from a common reference. The reported resolution is shown as a black line. A curve is displayed for the half-bit criterion in addition to lines showing the 0.143 gold standard cut-off and 0.5 cut-off.

### 8.1 FSC [i](#)



\*Reported resolution corresponds to spatial frequency of 0.263 Å<sup>-1</sup>

## 8.2 Resolution estimates [i](#)

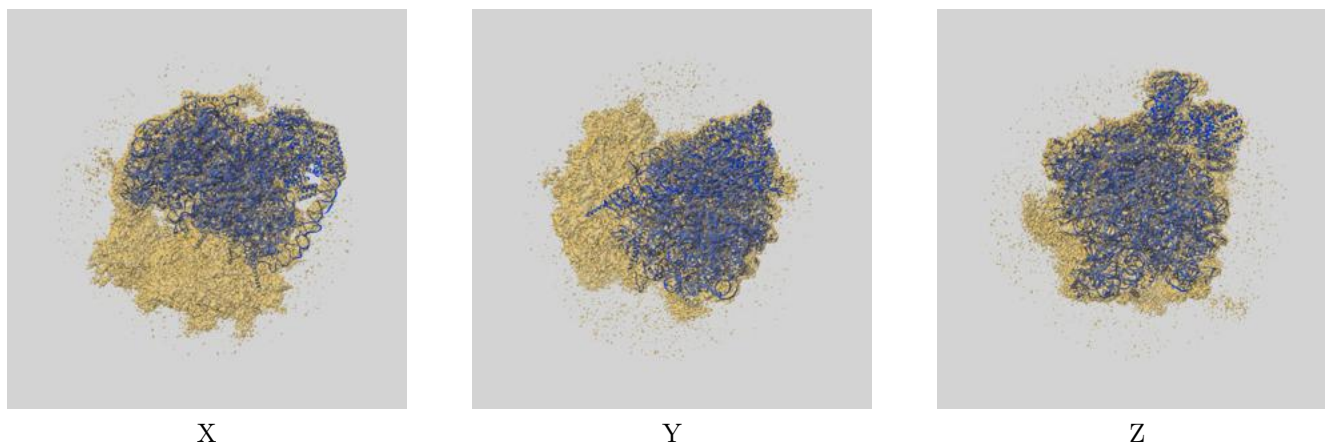
Resolution estimate (Å)	Estimation criterion (FSC cut-off)		
	0.143	0.5	Half-bit
Reported by author	3.80	-	-
Author-provided FSC curve	3.76	5.10	3.85
Unmasked-calculated*	6.46	11.15	6.99

\*Resolution estimate based on FSC curve calculated by comparison of deposited half-maps. The value from deposited half-maps intersecting FSC 0.143 CUT-OFF 6.46 differs from the reported value 3.8 by more than 10 %

## 9 Map-model fit [i](#)

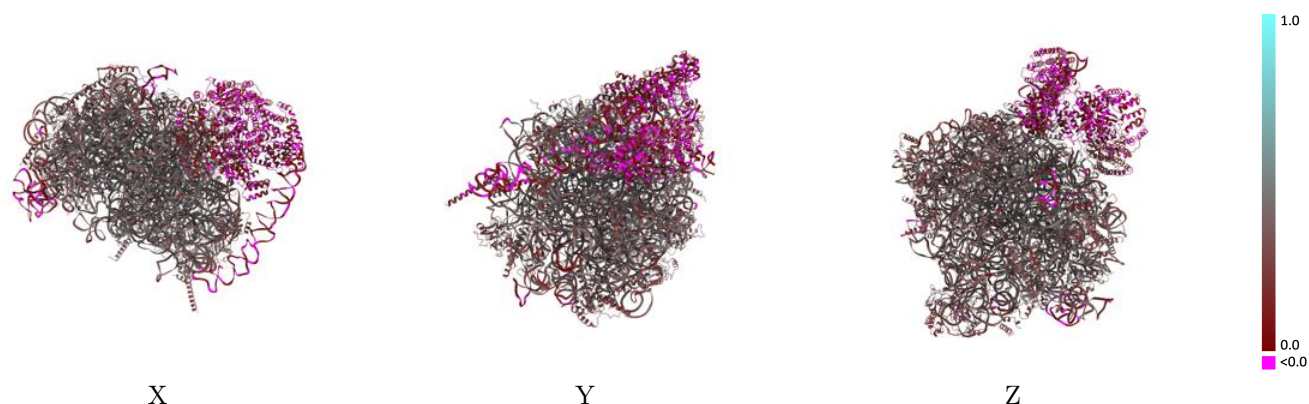
This section contains information regarding the fit between EMDB map EMD-16090 and PDB model 8BJQ. Per-residue inclusion information can be found in section [3](#) on page [14](#).

### 9.1 Map-model overlay [i](#)



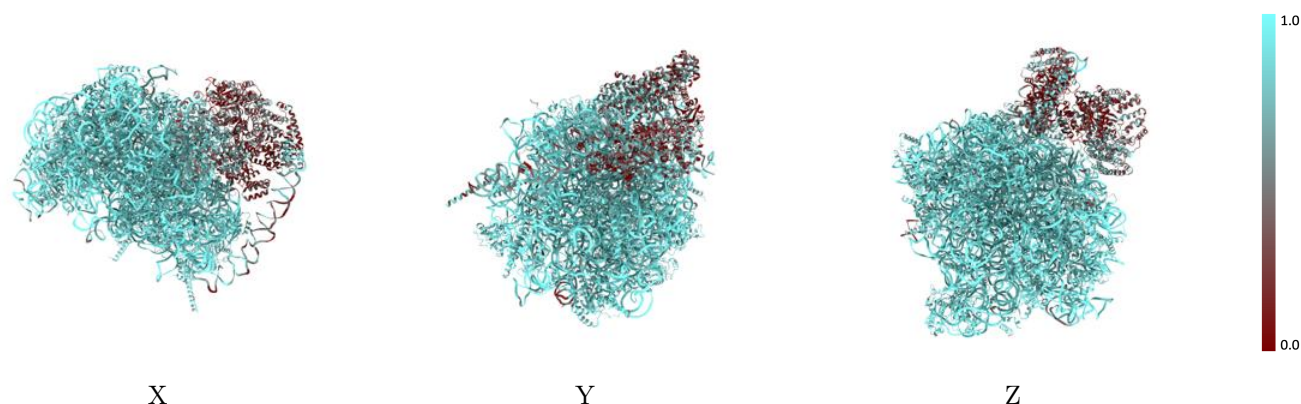
The images above show the 3D surface view of the map at the recommended contour level 0.2 at 50% transparency in yellow overlaid with a ribbon representation of the model coloured in blue. These images allow for the visual assessment of the quality of fit between the atomic model and the map.

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



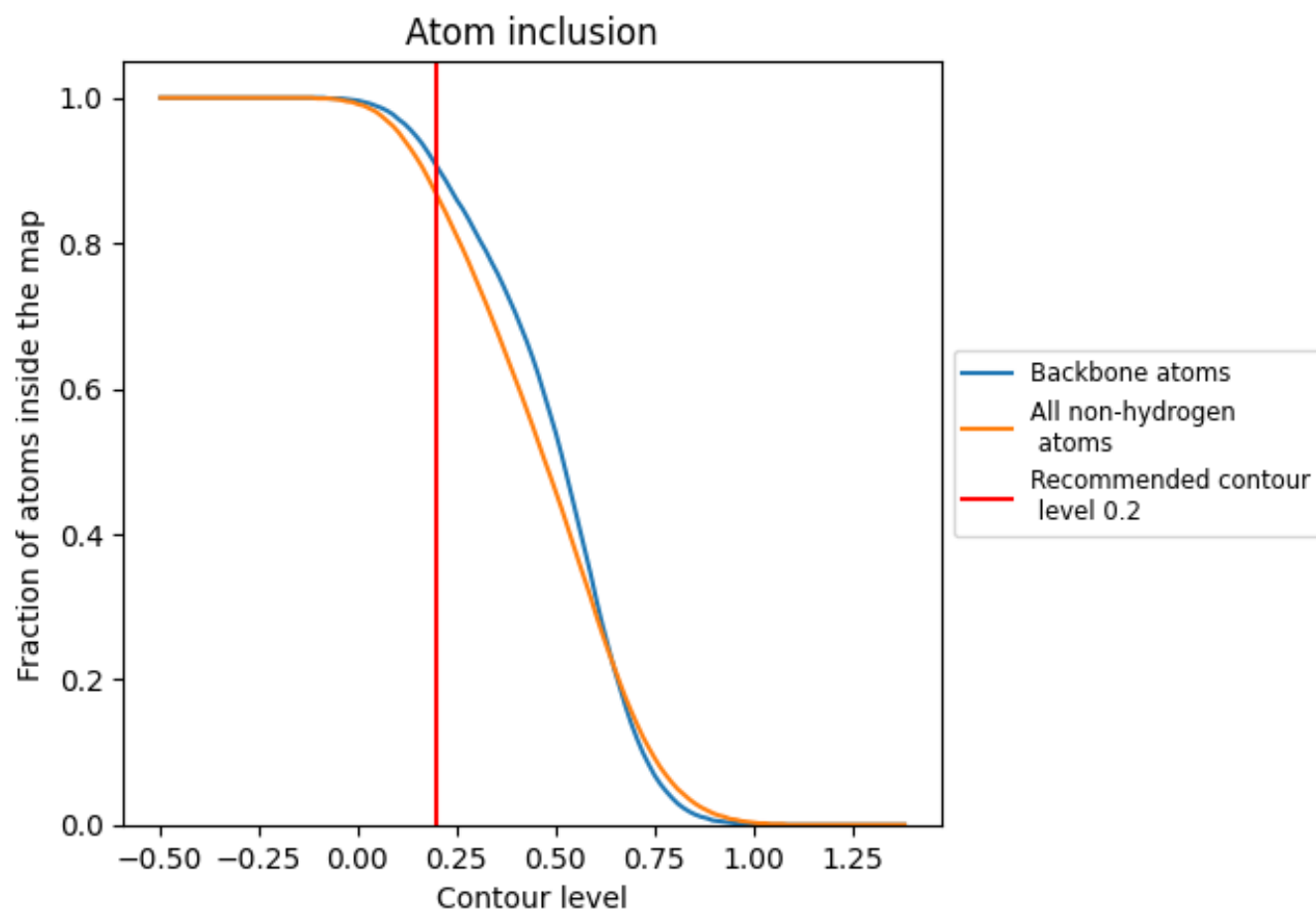
The images above show the model with each residue coloured according its Q-score. This shows their resolvability in the map with higher Q-score values reflecting better resolvability. Please note: Q-score is calculating the resolvability of atoms, and thus high values are only expected at resolutions 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 (0.2).







































































## 9.4 Atom inclusion [i](#)



At the recommended contour level, 91% of all backbone atoms, 87% 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 (0.2) and Q-score for the entire model and for each chain.

Chain	Atom inclusion	Q-score
All	 0.8660	 0.3370
1	 0.9560	 0.3660
A	 0.1720	 0.0460
B	 0.4710	 0.1120
C	 0.1710	 0.0510
C3	 0.9890	 0.3920
C4	 0.9940	 0.3720
D	 0.3710	 0.0800
LA	 0.8480	 0.4090
LB	 0.9000	 0.4110
LC	 0.8880	 0.3830
LD	 0.8750	 0.2840
LE	 0.8540	 0.2940
LF	 0.8850	 0.3770
LG	 0.8540	 0.3350
LH	 0.8950	 0.3730
LI	 0.8570	 0.3560
LJ	 0.8060	 0.2790
LL	 0.8630	 0.3360
LM	 0.8920	 0.3610
LN	 0.8650	 0.3780
LO	 0.8940	 0.4090
LP	 0.8370	 0.3760
LQ	 0.8890	 0.3760
LR	 0.8480	 0.3690
LS	 0.8840	 0.3990
LT	 0.8800	 0.3910
LU	 0.8340	 0.3000
LV	 0.8140	 0.4190
LW	 0.8200	 0.3950
LX	 0.8710	 0.3860
LY	 0.8640	 0.3510
LZ	 0.8950	 0.3510
La	 0.8760	 0.3470
Lb	 0.8430	 0.3460



*Continued on next page...*

*Continued from previous page...*

Chain	Atom inclusion	Q-score
Lc	 0.8880	 0.3450
Ld	 0.8760	 0.4000
Le	 0.8780	 0.4100
Lf	 0.8990	 0.4200
Lg	 0.8770	 0.3860
Lh	 0.8810	 0.3370
Li	 0.8260	 0.3310
Lj	 0.9180	 0.4140
Lk	 0.8330	 0.3420
Ll	 0.8580	 0.3570
Lm	 0.8740	 0.3840
Ln	 0.8120	 0.3120
Lo	 0.8610	 0.3750
Lp	 0.8390	 0.3730