



## wwPDB EM Validation Summary Report ⓘ

Nov 19, 2022 – 11:35 am GMT

PDB ID : 5M3M  
EMDB ID : EMD-4148  
Title : Free monomeric RNA polymerase I at 4.0A resolution  
Authors : Neyer, S.; Kunz, M.; Geiss, C.; Hantsche, M.; Hodirnau, V.-V.; Seybert, A.;  
Engel, C.; Scheffer, M.P.; Cramer, P.; Frangakis, A.S.  
Deposited on : 2016-10-15  
Resolution : 4.00 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>.

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The following versions of software and data (see [references ⓘ](#)) were used in the production of this report:

EMDB validation analysis : 0.0.1.dev43  
Mogul : 1.8.4, CSD as541be (2020)  
MolProbity : 4.02b-467  
Percentile statistics : 20191225.v01 (using entries in the PDB archive December 25th 2019)  
MapQ : 1.9.9  
Ideal geometry (proteins) : Engh & Huber (2001)  
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)  
Validation Pipeline (wwPDB-VP) : 2.31.2

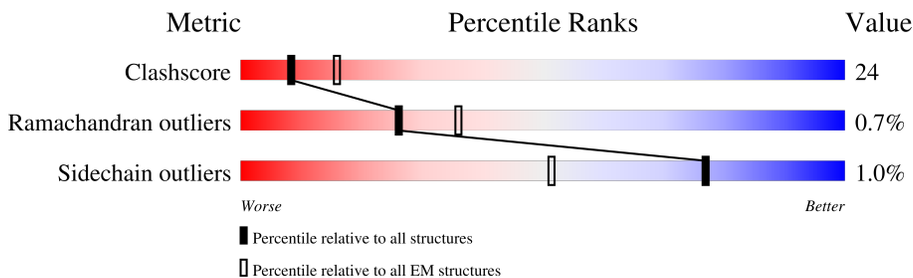
# 1 Overall quality at a glance i

The following experimental techniques were used to determine the structure:

*ELECTRON MICROSCOPY*

The reported resolution of this entry is 4.00 Å.

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



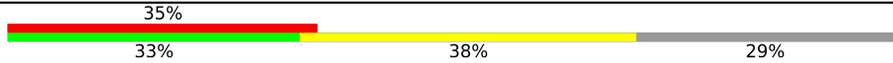
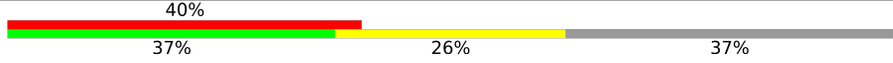
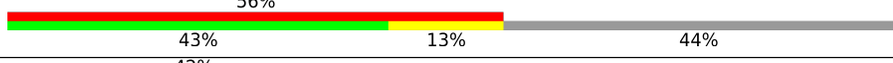
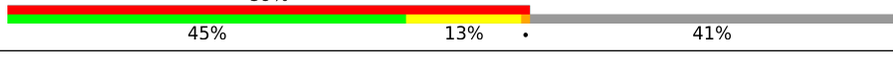
Metric	Whole archive (#Entries)	EM structures (#Entries)
Clashscore	158937	4297
Ramachandran outliers	154571	4023
Sidechain outliers	154315	3826

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	1664	61% 44% 43% 12%
2	B	1203	51% 49% 48%
3	C	335	52% 41% 49% 9%
4	E	215	87% 44% 54%
5	F	155	52% 32% 31% 37%
6	H	146	52% 48% 42% 10%
7	I	125	91% 46% 46% 7%
8	J	70	40% 44% 54%

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Mol	Chain	Length	Quality of chain
9	K	142	
10	L	70	
11	M	415	
12	N	233	
13	D	137	
14	G	326	

The following table lists non-polymeric compounds, carbohydrate monomers and non-standard residues in protein, DNA, RNA chains that are outliers for geometric or electron-density-fit criteria:

Mol	Type	Chain	Res	Chirality	Geometry	Clashes	Electron density
16	SO4	B	1301	-	-	X	-

## 2 Entry composition

There are 16 unique types of molecules in this entry. The entry contains 33233 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 DNA-directed RNA polymerase I subunit RPA190.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
1	A	1462	11558	7304	2006	2187	61	0	0

- Molecule 2 is a protein called DNA-directed RNA polymerase I subunit RPA135.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
2	B	1166	9266	5864	1617	1734	51	0	0

- Molecule 3 is a protein called DNA-directed RNA polymerases I and III subunit RPAC1.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
3	C	305	2423	1539	416	460	8	0	0

- Molecule 4 is a protein called DNA-directed RNA polymerases I, II, and III subunit RPABC1.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
4	E	212	1735	1102	306	316	11	0	0

- Molecule 5 is a protein called DNA-directed RNA polymerases I, II, and III subunit RPABC2.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
5	F	98	807	512	142	150	3	0	0

- Molecule 6 is a protein called DNA-directed RNA polymerases I, II, and III subunit RPABC3.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
6	H	131	1052	664	176	208	4	0	0

- Molecule 7 is a protein called DNA-directed RNA polymerase I subunit RPA12.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
7	I	116	883	550	148	176	9	0	0

- Molecule 8 is a protein called DNA-directed RNA polymerases I, II, and III subunit RPABC5.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
8	J	69	569	362	101	100	6	0	0

- Molecule 9 is a protein called DNA-directed RNA polymerases I and III subunit RPAC2.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
9	K	101	793	496	130	162	5	0	0

- Molecule 10 is a protein called DNA-directed RNA polymerases I, II, and III subunit RPABC4.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
10	L	44	352	217	70	61	4	0	0

- Molecule 11 is a protein called DNA-directed RNA polymerase I subunit RPA49.

Mol	Chain	Residues	Atoms				AltConf	Trace
			Total	C	N	O		
11	M	97	771	490	124	157	0	0

- Molecule 12 is a protein called DNA-directed RNA polymerase I subunit RPA34.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
12	N	131	1035	660	171	200	4	0	0

- Molecule 13 is a protein called DNA-directed RNA polymerase I subunit RPA14.

Mol	Chain	Residues	Atoms				AltConf	Trace
			Total	C	N	O		
13	D	58	459	289	78	92	0	0

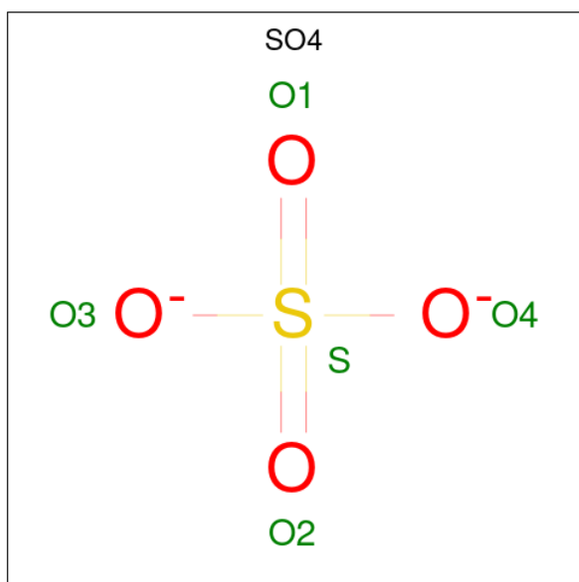
- Molecule 14 is a protein called DNA-directed RNA polymerase I subunit RPA43.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
14	G	192	1518	979	261	273	5	0	0

- Molecule 15 is ZINC ION (three-letter code: ZN) (formula: Zn).

Mol	Chain	Residues	Atoms		AltConf
15	A	2	Total	Zn	0
			2	2	
15	B	1	Total	Zn	0
			1	1	
15	I	2	Total	Zn	0
			2	2	
15	J	1	Total	Zn	0
			1	1	
15	L	1	Total	Zn	0
			1	1	

- Molecule 16 is SULFATE ION (three-letter code: SO4) (formula: O<sub>4</sub>S).

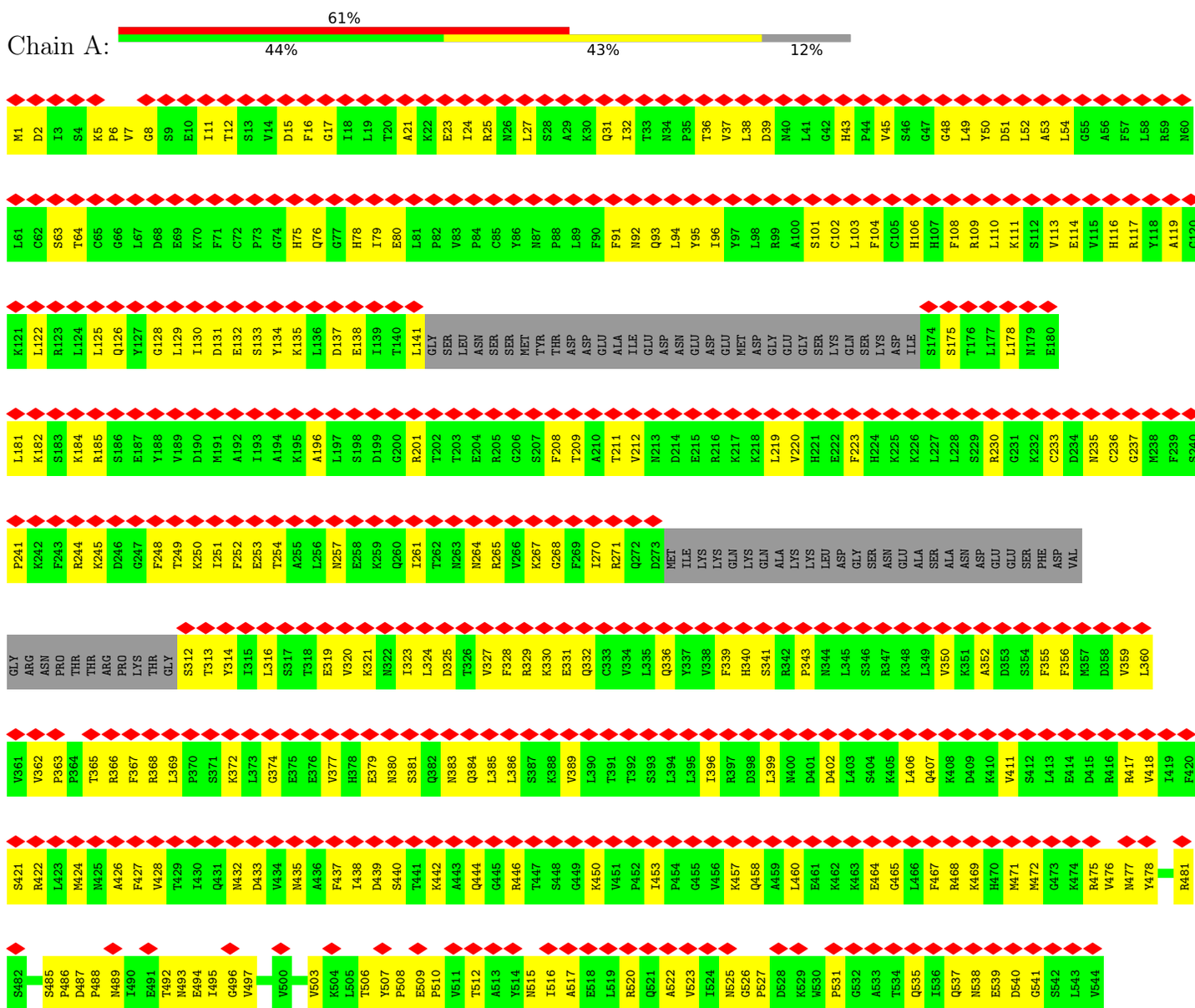


Mol	Chain	Residues	Atoms			AltConf
16	B	1	Total	O	S	0
			5	4	1	

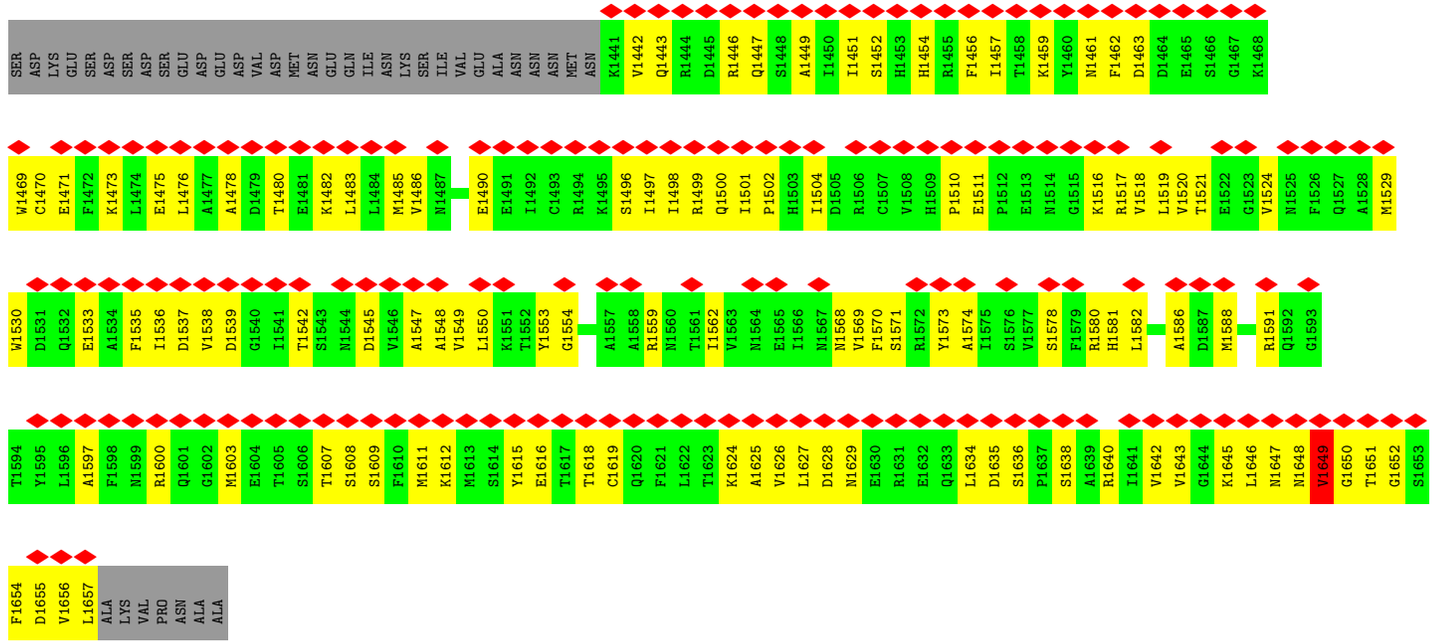
### 3 Residue-property plots

These plots are drawn for all protein, RNA, DNA and oligosaccharide chains in the entry. The first graphic for a chain summarises the proportions of the various outlier classes displayed in the second graphic. The second graphic shows the sequence view annotated by issues in geometry and atom inclusion in map density. Residues are color-coded according to the number of geometric quality criteria for which they contain at least one outlier: green = 0, yellow = 1, orange = 2 and red = 3 or more. A red diamond above a residue indicates a poor fit to the EM map for this residue (all-atom inclusion < 40%). Stretches of 2 or more consecutive residues without any outlier are shown as a green connector. Residues present in the sample, but not in the model, are shown in grey.

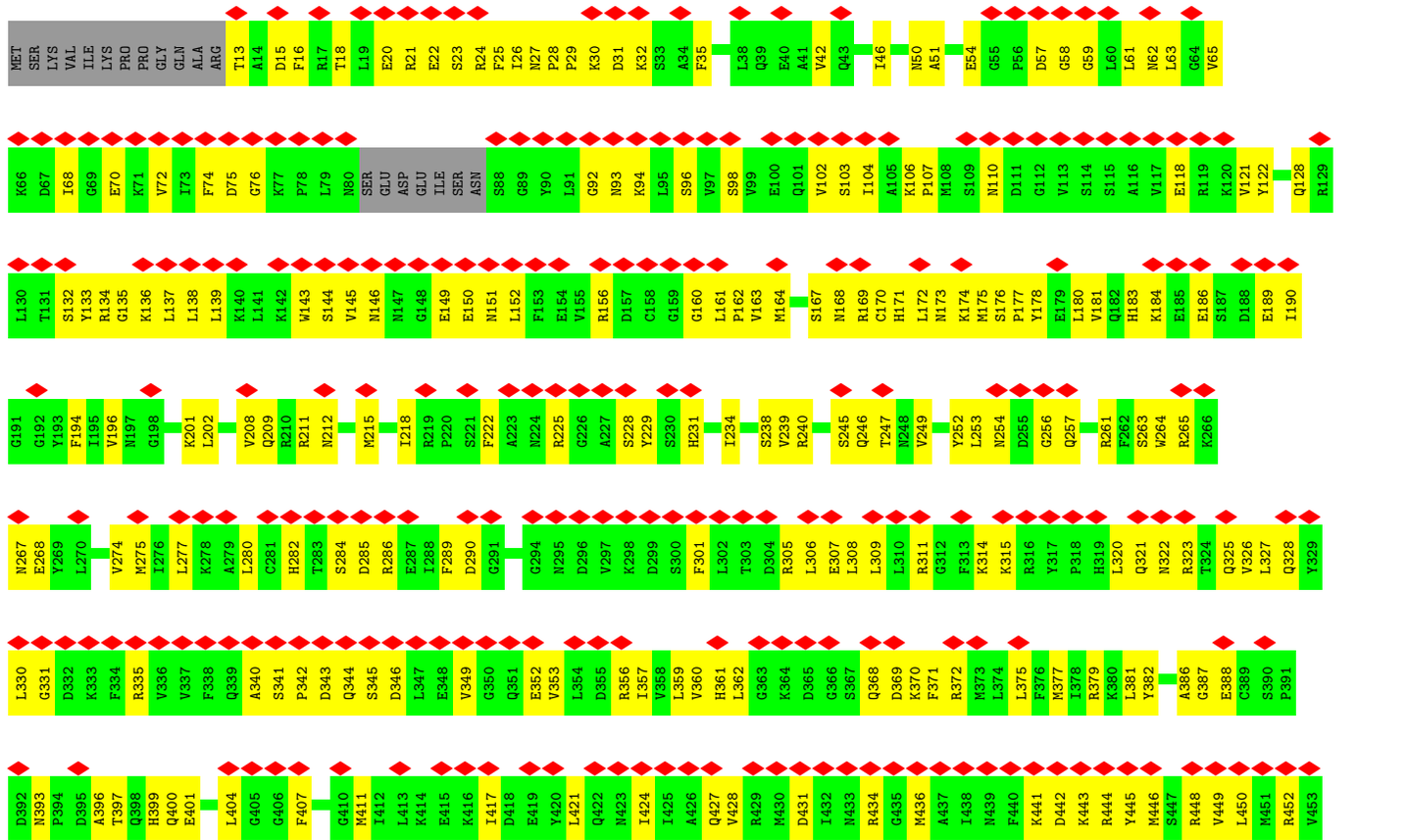
- Molecule 1: DNA-directed RNA polymerase I subunit RPA190

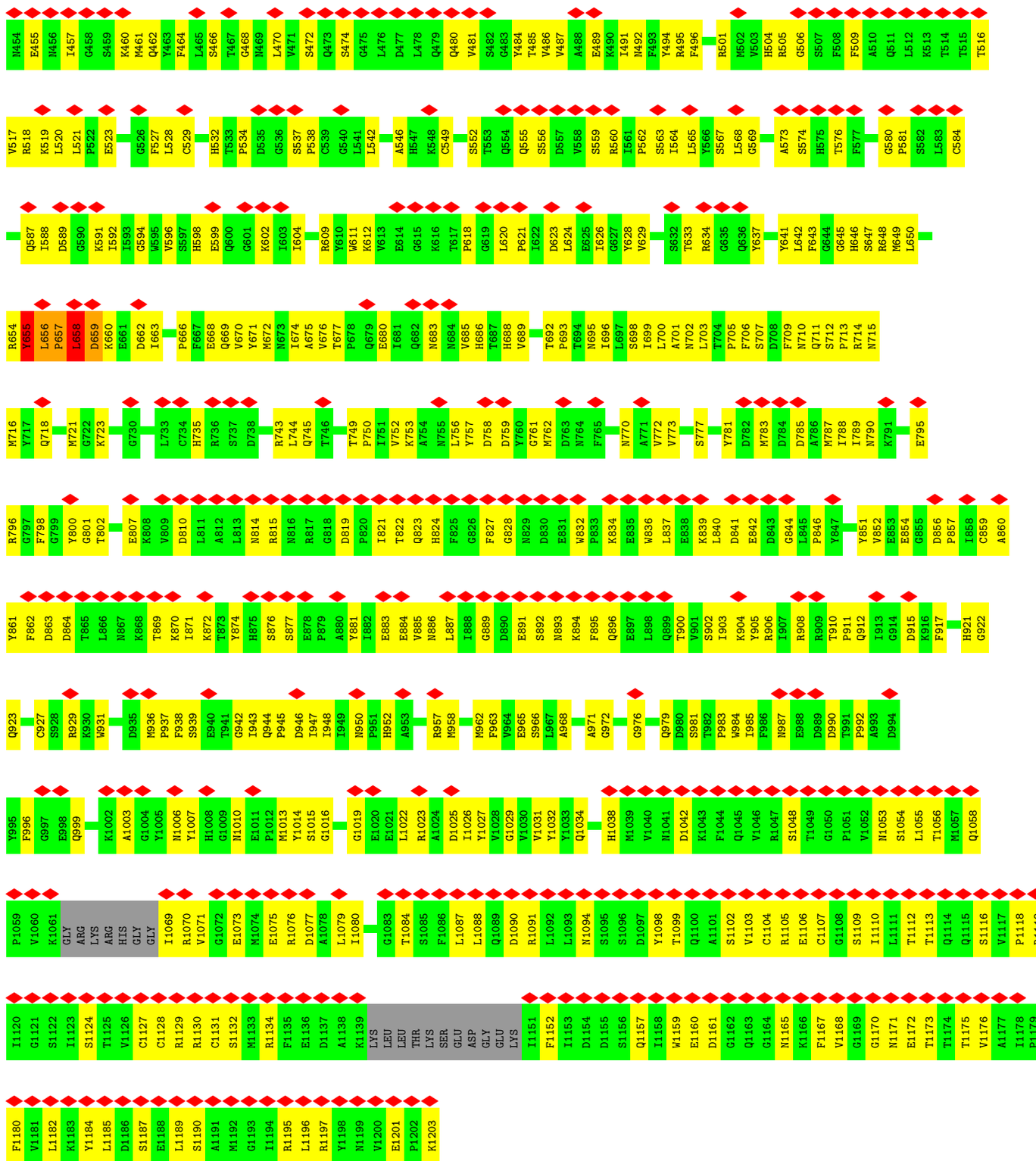


PRO	S1289	A1229	D1159	K1096	G1029	R956	R878	L810	T740	Q671	R606	S545
ARG	Y1290	S1230	G1160	Y1097	V1030	M960	L879	S811	P741	D672	V607	L646
LEU	V1291	A1231	V1161	S1098	H1031	V961	Q880	Y812	P742	D675	M610	I547
GLN	I1292	A1232	M1162	K1099	Y1034	S962	I882	R815	M743	A676	E811	G648
THR	H1293	I1233	K1163	K1100	D1035	G963	L883	T824	M744	T681	K612	M549
ASP	H1294	I1234	K1164	T1101	H1036	K964	R884	R825	M745	T682	L614	S550
VAL	M1295	T1235	K1165	L1102	R1039	T965	D885	A826	G746	S882	L615	V551
ALA	R1296	P1236	F1166	K1103	D1040	L966	P886	F826	N747	K883	R616	E552
ASN	F1297	Q1237	R1167	Y1104	A1041	P967	N887	T227	M748	D684	R615	Q553
SER	F1298	Q1238	M1168	Y1105	D1042	S968	N888	C828	L749	H617	Y618	Q554
SER	D1298	M1239	L1172	K1106	D1043	S969	K889	G829	I750	Y618	A619	K555
ASN	M1299	L1240	Y1174	K1107	G1043	Y972	S889	M830	S751	R689	M620	A556
LYS	F1241	L1241	M1175	K1108	T1044	E973	G890	D831	K752	E690	T621	L557
ARG	M1176	R1176	M1176	H1107	L1045	T974	I891	D832	N753	Q691	G622	L558
LEU	E1183	E1183	E1183	S1109	V1046	D975	A894	L833	N756	Y692	G623	M559
GLU	E1111	E1111	E1111	K1110	Q1047	A976	L894	R834	K757	Y692	A623	Q560
GLU	A1184	A1184	A1184	P1112	Q1048	M977	R899	L835	N757	Y692	Y624	L561
ASP	V1185	V1185	V1185	F1048	Y1050	A978	K899	T836	E758	Q694	M625	L562
ASP	V1186	V1186	V1186	L1049	Y1051	G979	R902	A837	Y759	Y697	A626	L563
GLU	I1187	I1187	I1187	Y1050	G1051	G980	A902	E838	K762	G998	D627	T563
GLN	I1188	I1188	I1188	G1051	Y981	I903	I903	G839	L765	F701	F628	P564
SER	A1189	A1189	A1189	G1052	V982	T904	T904	M940	L766	R702	D629	S665
HIS	S1190	S1190	S1190	D1053	K983	S905	S905	K941	E766	F703	G630	S666
LYS	Q1191	Q1191	Q1191	A1054	G984	Q906	Q906	W842	E767	E703	D631	M667
THR	S1192	S1192	S1192	A1054	R985	V907	V907	R843	E768	D704	D632	V568
LYS	V1193	V1193	V1193	D1055	F986	V908	V908	T944	G705	G705	E632	T570
LYS	G1194	G1194	G1194	D1056	Y987	S909	S909	D845	L770	H706	M633	H571
GLN	E1195	E1195	E1195	I1057	S988	K910	K910	I846	F771	T707	M634	T572
ALA	P1196	P1196	P1196	I1058	E994	P913	P913	K848	K772	T708	M635	L573
VAL	S1197	S1197	S1197	E1060	Y995	D914	D914	K949	K773	T708	H642	K575
SER	T1198	T1198	T1198	S1061	Y996	G915	G915	T849	D773	R709	E643	V577
TYR	Q1199	Q1199	Q1199	H1062	M1000	T916	T916	R852	G774	S710	E644	V578
GLU	M1200	M1200	M1200	M1063	G1005	R918	R918	T853	L776	E712	M640	H580
ASP	T1201	T1201	T1201	T1064	L1006	K919	K919	G854	L777	I713	E646	L581
GLU	L1202	L1202	L1202	Q1065	L1007	F920	F920	R855	L778	I713	L648	K582
ASP	M1203	M1203	M1203	E1067	I1008	P921	P921	S850	L779	I714	M649	M683
ILE	T1204	T1204	T1204	E1067	D1008	S924	S924	V851	L781	T717	L650	R584
GLU	F1205	F1205	F1205	L1070	T1009	Y930	Y930	R852	D782	P717	L651	D585
THR	HIS	HIS	HIS	Y1074	A1010	L930	L930	T853	K783	I719	M652	V587
ARG	PHE	PHE	PHE	Y1074	VAL	S936	S936	G854	Y786	I719	L648	H580
GLU	ALA	ALA	ALA	K1078	LYS	Y937	Y937	R855	G787	I719	M649	K582
GLY	HIS	GLY	GLY	K1079	THR	N937	N937	R855	A788	F722	L650	M683
GLY	HIS	GLY	GLY	Y1080	SER	Y938	Y938	S850	S789	Y723	L651	R584
ALA	ALA	ALA	ALA	M1081	ARG	V938	V938	S850	K790	P724	M652	D585
A1213	ALA	ALA	ALA	M1081	SER	S941	S941	R855	K791	L725	T653	V587
M1214	ALA	ALA	ALA	P1082	G1017	Q942	Q942	L865	G792	W726	D654	L588
V1215	ALA	ALA	ALA	S1083	Y1018	T943	T943	K866	I793	T727	S655	L588
T1216	ALA	ALA	ALA	A1084	L1019	M944	M944	R866	G799	K729	Q856	M589
L1217	ALA	ALA	ALA	Q1020	Q1020	C945	C945	T867	L797	Q730	Q857	M590
A1213	ALA	ALA	ALA	R1021	R1021	L946	L946	S868	V800	I731	L658	Q591
M1214	ALA	ALA	ALA	C1022	C1022	Q949	Q949	P869	P803	I732	T659	R593
V1215	ALA	ALA	ALA	L1023	L1023	Q950	Q950	A870	E804	T733	T659	L594
T1216	ALA	ALA	ALA	T1024	T1024	D871	D871	D871	V805	V735	G663	T594
L1217	ALA	ALA	ALA	K1025	K1025	D872	D872	D872	V806	L736	G664	L595
GLY	S1146	S1146	S1146	E1028	E1028	P873	P873	P873	V807	L737	P665	H596
THR	F1147	F1147	F1147	E1028	E1028	E874	E874	E874	K808	N738	V666	K597
ASN	L1148	L1148	L1148	E1028	E1028	L876	L876	L876	V809	V739	G668	S599
THR	D1149	D1149	D1149	E1028	E1028	K877	K877	K877			L669	M601
ALA	K1150	K1150	K1150	E1028	E1028						L670	K604
GLY	N1151	N1151	N1151	E1028	E1028							V605
GLY	S1152	S1152	S1152	E1028	E1028							
ASN	K1153	K1153	K1153	E1028	E1028							
ILE	L1154	L1154	L1154	E1028	E1028							
VAL	F1155	F1155	F1155	E1028	E1028							
ALA	K1156	K1156	K1156	E1028	E1028							
VAL	S1157	S1157	S1157	E1028	E1028							
VAL	S1158	S1158	S1158	E1028	E1028							



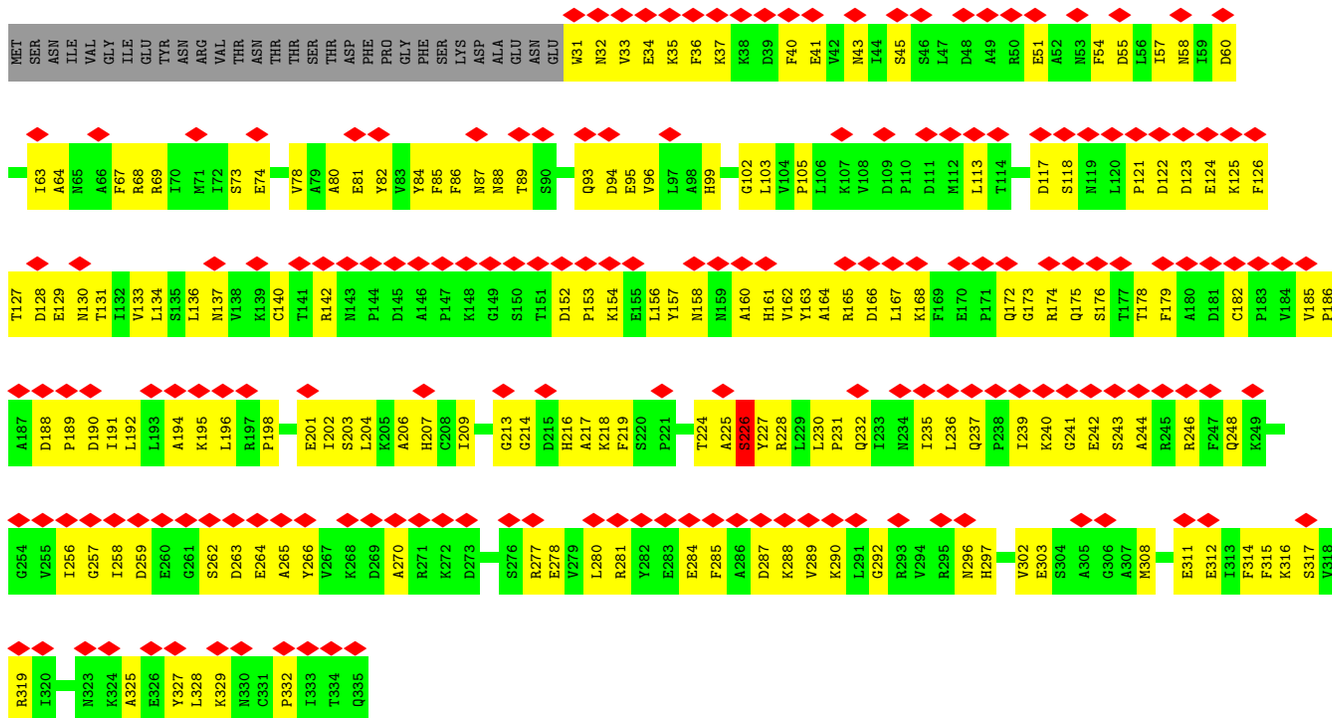
● Molecule 2: DNA-directed RNA polymerase I subunit RPA135



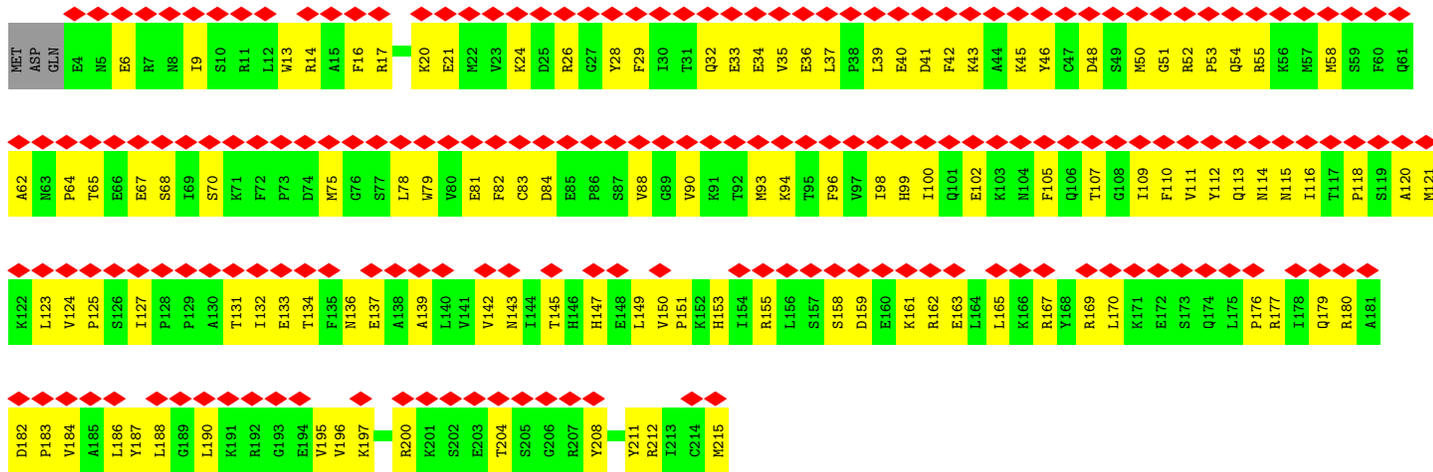
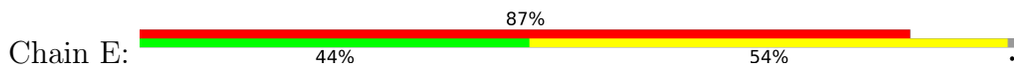


● Molecule 3: DNA-directed RNA polymerases I and III subunit RPAC1

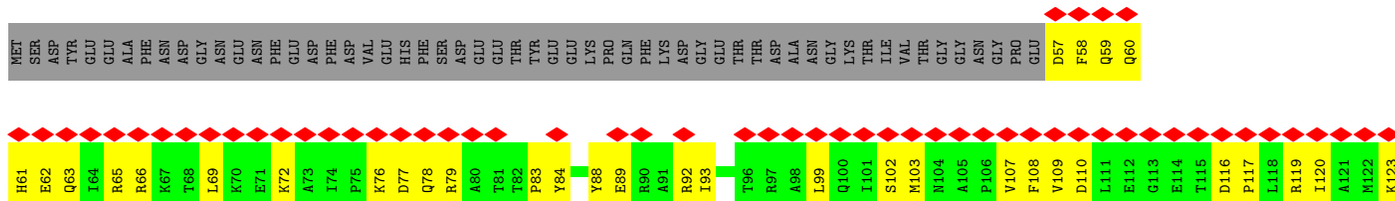




• Molecule 4: DNA-directed RNA polymerases I, II, and III subunit RPABC1

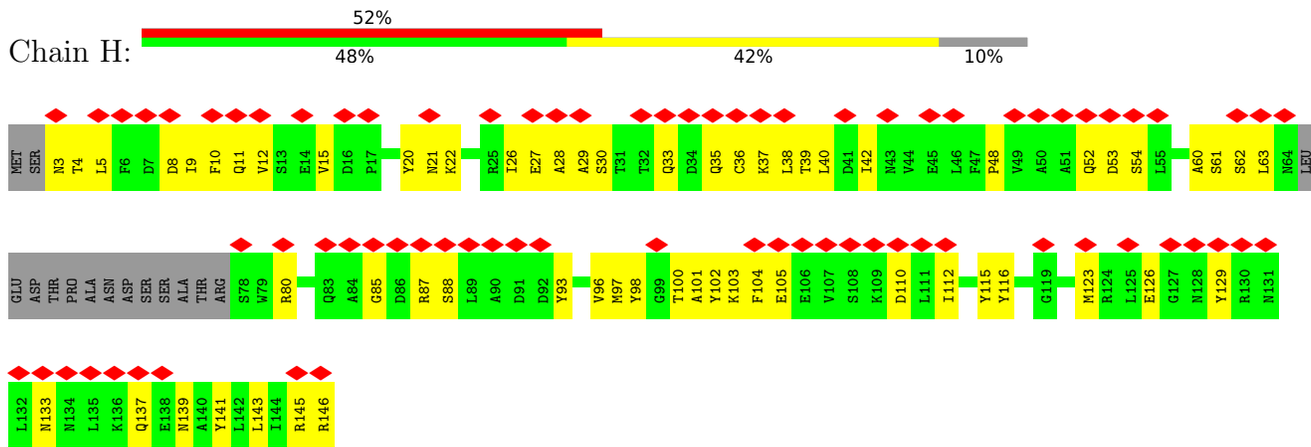


• Molecule 5: DNA-directed RNA polymerases I, II, and III subunit RPABC2

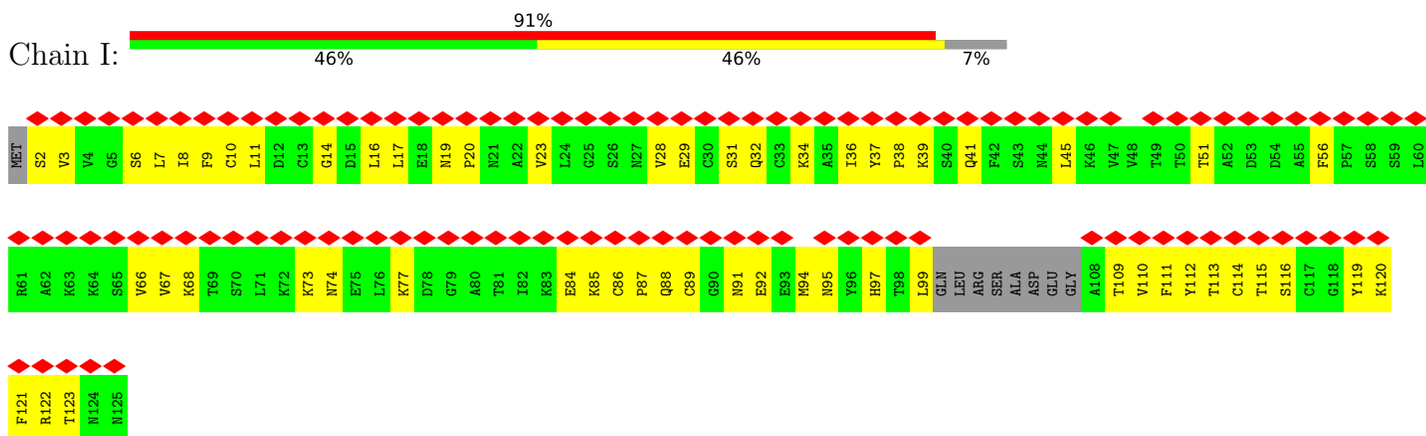




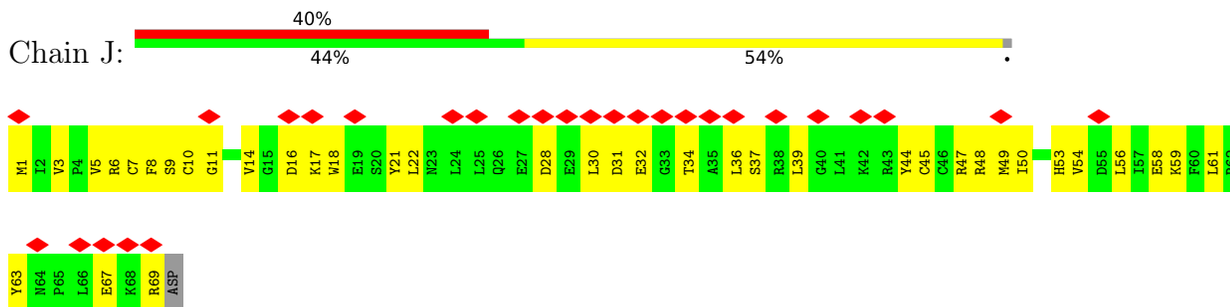
- Molecule 6: DNA-directed RNA polymerases I, II, and III subunit RPABC3



- Molecule 7: DNA-directed RNA polymerase I subunit RPA12

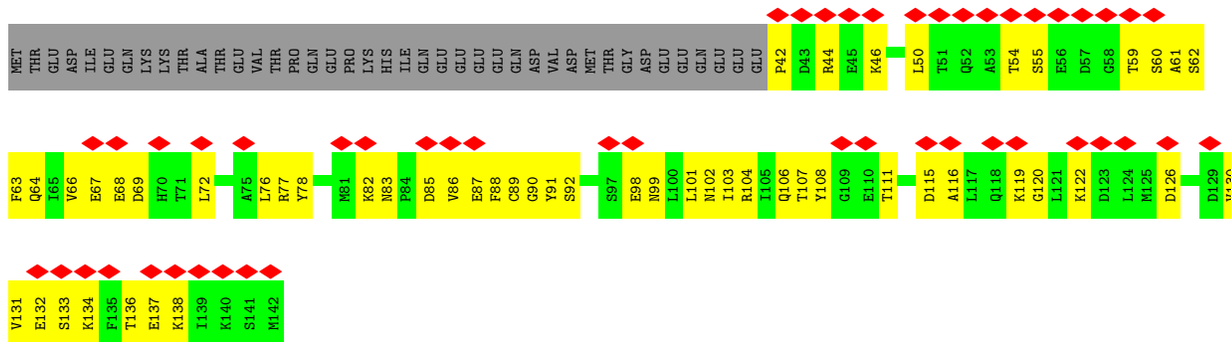


- Molecule 8: DNA-directed RNA polymerases I, II, and III subunit RPABC5

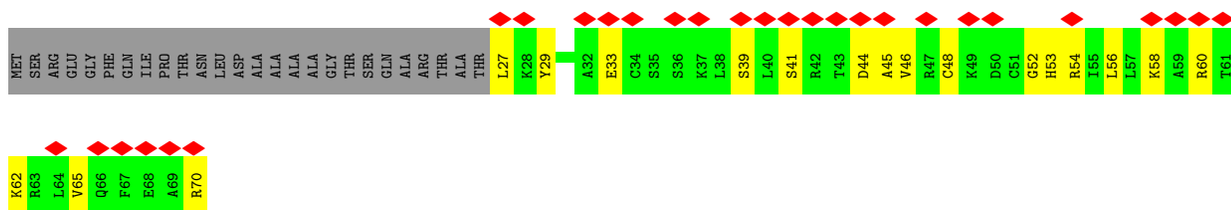


- Molecule 9: DNA-directed RNA polymerases I and III subunit RPAC2

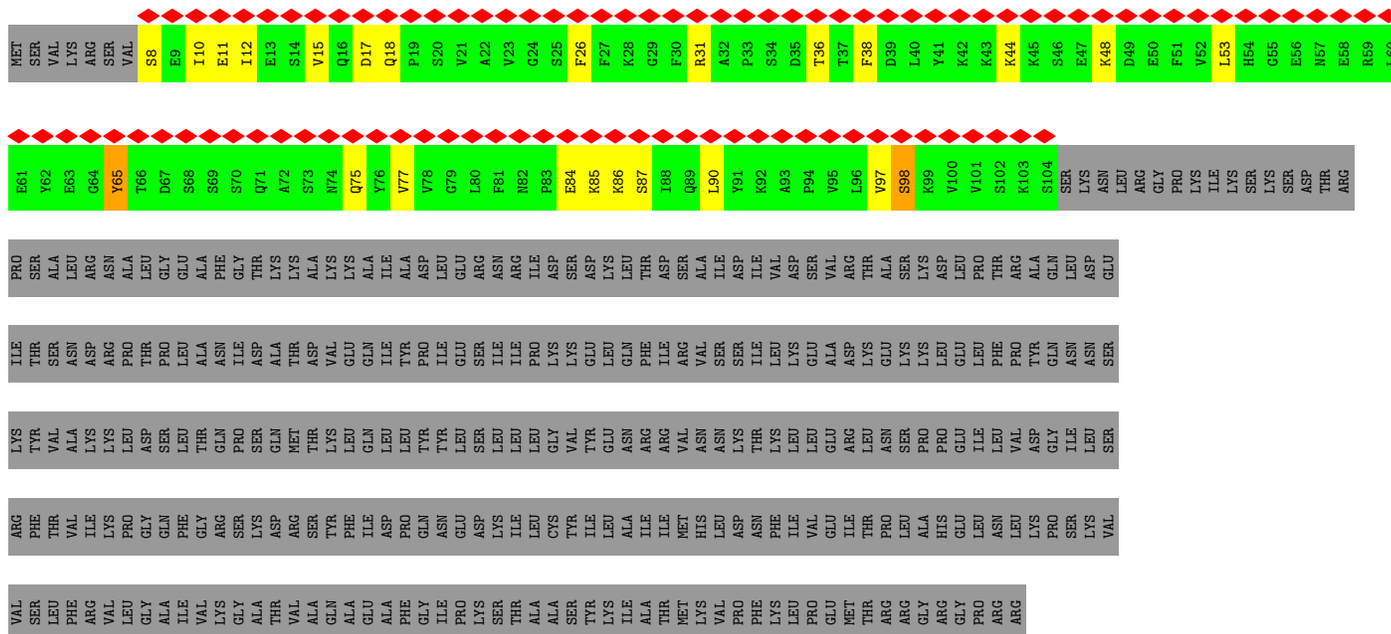




• Molecule 10: DNA-directed RNA polymerases I, II, and III subunit RPABC4



• Molecule 11: DNA-directed RNA polymerase I subunit RPA49



• Molecule 12: DNA-directed RNA polymerase I subunit RPA34





## 4 Experimental information

Property	Value	Source
EM reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, C1	Depositor
Number of particles used	94000	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 ( $e^-/\text{\AA}^2$ )	56	Depositor
Minimum defocus (nm)	Not provided	
Maximum defocus (nm)	Not provided	
Magnification	Not provided	
Image detector	GATAN K2 QUANTUM (4k x 4k)	Depositor
Maximum map value	0.152	Depositor
Minimum map value	-0.094	Depositor
Average map value	-0.000	Depositor
Map value standard deviation	0.007	Depositor
Recommended contour level	0.04	Depositor
Map size (Å)	241.49998, 241.49998, 241.49998	wwPDB
Map dimensions	230, 230, 230	wwPDB
Map angles (°)	90.0, 90.0, 90.0	wwPDB
Pixel spacing (Å)	1.05, 1.05, 1.05	Depositor

## 5 Model quality [i](#)

### 5.1 Standard geometry [i](#)

Bond lengths and bond angles in the following residue types are not validated in this section: ZN, SO4

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	0.40	0/11770	0.51	0/15895
2	B	0.45	0/9471	0.53	0/12805
3	C	0.45	0/2475	0.51	0/3354
4	E	0.40	0/1771	0.50	0/2383
5	F	0.37	0/821	0.48	0/1106
6	H	0.46	0/1070	0.54	0/1449
7	I	0.38	0/895	0.49	0/1205
8	J	0.50	0/578	0.53	0/775
9	K	0.45	0/804	0.55	0/1083
10	L	0.40	0/354	0.53	0/468
11	M	0.40	0/786	0.55	0/1057
12	N	0.39	0/1052	0.55	0/1418
13	D	0.40	0/465	0.58	0/630
14	G	0.37	0/1555	0.66	3/2113 (0.1%)
All	All	0.42	0/33867	0.53	3/45741 (0.0%)

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

Mol	Chain	#Chirality outliers	#Planarity outliers
1	A	0	1

There are no bond length outliers.

All (3) bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
14	G	241	ARG	NE-CZ-NH1	11.80	126.20	120.30
14	G	241	ARG	NE-CZ-NH2	-11.62	114.49	120.30

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
14	G	241	ARG	CD-NE-CZ	6.08	132.12	123.60

There are no chirality outliers.

All (1) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	A	1649	VAL	Peptide

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

In the following table, the Non-H and H(model) columns list the number of non-hydrogen atoms and hydrogen atoms in the chain respectively. The H(added) column lists the number of hydrogen atoms added and optimized by MolProbity. The Clashes column lists the number of clashes within the asymmetric unit, whereas Symm-Clashes lists symmetry-related clashes.

Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
1	A	11558	0	11642	651	0
2	B	9266	0	9151	558	0
3	C	2423	0	2412	156	0
4	E	1735	0	1764	98	0
5	F	807	0	827	45	0
6	H	1052	0	1021	47	0
7	I	883	0	879	60	0
8	J	569	0	585	36	0
9	K	793	0	790	56	0
10	L	352	0	374	20	0
11	M	771	0	755	11	0
12	N	1035	0	1069	29	0
13	D	459	0	462	8	0
14	G	1518	0	1528	31	0
15	A	2	0	0	0	0
15	B	1	0	0	0	0
15	I	2	0	0	0	0
15	J	1	0	0	0	0
15	L	1	0	0	0	0
16	B	5	0	0	8	0
All	All	33233	0	33259	1613	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 24.

The worst 5 of 1613 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)
2:B:656:LEU:O	12:N:153:VAL:HG11	1.62	0.97
3:C:225:ALA:O	3:C:226:SER:HB2	1.63	0.95
1:A:1501:ILE:HG22	1:A:1502:PRO:HD2	1.49	0.94
2:B:894:LYS:HG2	10:L:54:ARG:HH21	1.33	0.94
5:F:66:ARG:HA	5:F:69:LEU:HD12	1.50	0.93

There are no symmetry-related clashes.

## 5.3 Torsion angles [i](#)

### 5.3.1 Protein backbone [i](#)

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

The Analysed column shows the number of residues for which the backbone conformation was analysed, and the total number of residues.

Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
1	A	1448/1664 (87%)	1364 (94%)	75 (5%)	9 (1%)	25	63
2	B	1158/1203 (96%)	1110 (96%)	41 (4%)	7 (1%)	25	63
3	C	303/335 (90%)	283 (93%)	19 (6%)	1 (0%)	41	75
4	E	210/215 (98%)	202 (96%)	8 (4%)	0	100	100
5	F	96/155 (62%)	94 (98%)	2 (2%)	0	100	100
6	H	127/146 (87%)	124 (98%)	3 (2%)	0	100	100
7	I	112/125 (90%)	108 (96%)	4 (4%)	0	100	100
8	J	67/70 (96%)	61 (91%)	6 (9%)	0	100	100
9	K	99/142 (70%)	95 (96%)	4 (4%)	0	100	100
10	L	42/70 (60%)	38 (90%)	4 (10%)	0	100	100
11	M	95/415 (23%)	86 (90%)	7 (7%)	2 (2%)	7	39
12	N	125/233 (54%)	108 (86%)	13 (10%)	4 (3%)	4	31
13	D	54/137 (39%)	50 (93%)	2 (4%)	2 (4%)	3	28
14	G	186/326 (57%)	171 (92%)	13 (7%)	2 (1%)	14	51
All	All	4122/5236 (79%)	3894 (94%)	201 (5%)	27 (1%)	26	61

5 of 27 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	A	946	LEU
2	B	657	PRO
2	B	658	LEU
2	B	659	ASP
3	C	226	SER

### 5.3.2 Protein sidechains [i](#)

In the following table, the Percentiles column shows the percent sidechain outliers of the chain as a percentile score with respect to all 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	1292/1465 (88%)	1288 (100%)	4 (0%)	92	95
2	B	1022/1053 (97%)	1020 (100%)	2 (0%)	93	96
3	C	269/296 (91%)	268 (100%)	1 (0%)	91	94
4	E	194/197 (98%)	194 (100%)	0	100	100
5	F	88/137 (64%)	88 (100%)	0	100	100
6	H	115/128 (90%)	115 (100%)	0	100	100
7	I	103/110 (94%)	103 (100%)	0	100	100
8	J	64/65 (98%)	64 (100%)	0	100	100
9	K	91/130 (70%)	91 (100%)	0	100	100
10	L	39/57 (68%)	39 (100%)	0	100	100
11	M	88/371 (24%)	79 (90%)	9 (10%)	7	28
12	N	124/220 (56%)	120 (97%)	4 (3%)	39	62
13	D	55/116 (47%)	49 (89%)	6 (11%)	6	26
14	G	170/291 (58%)	158 (93%)	12 (7%)	14	42
All	All	3714/4636 (80%)	3676 (99%)	38 (1%)	77	86

5 of 38 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
14	G	35	SER
14	G	230	ARG

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Mol	Chain	Res	Type
14	G	39	VAL
14	G	167	THR
14	G	243	VAL

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 50 such sidechains are listed below:

Mol	Chain	Res	Type
2	B	587	GLN
3	C	58	ASN
9	K	99	ASN
2	B	686	HIS
2	B	1008	HIS

### 5.3.3 RNA [i](#)

There are no RNA molecules in this entry.

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

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

## 5.5 Carbohydrates [i](#)

There are no monosaccharides in this entry.

## 5.6 Ligand geometry [i](#)

Of 8 ligands modelled in this entry, 7 are monoatomic - leaving 1 for Mogul analysis.

In the following table, the Counts columns list the number of bonds (or angles) for which Mogul statistics could be retrieved, the number of bonds (or angles) that are observed in the model and the number of bonds (or angles) that are defined in the Chemical Component Dictionary. The Link column lists molecule types, if any, to which the group is linked. The Z score for a bond length (or angle) is the number of standard deviations the observed value is removed from the expected value. A bond length (or angle) with  $|Z| > 2$  is considered an outlier worth inspection. RMSZ is the root-mean-square of all Z scores of the bond lengths (or angles).

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	$\# Z  > 2$	Counts	RMSZ	$\# Z  > 2$
16	SO4	B	1301	-	4,4,4	0.20	0	6,6,6	0.35	0

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.

1 monomer is involved in 8 short contacts:

Mol	Chain	Res	Type	Clashes	Symm-Clashes
16	B	1301	SO4	8	0

## 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.

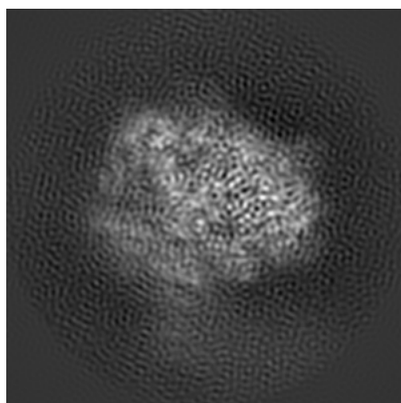
## 6 Map visualisation [i](#)

This section contains visualisations of the EMDB entry EMD-4148. These allow visual inspection of the internal detail of the map and identification of artifacts.

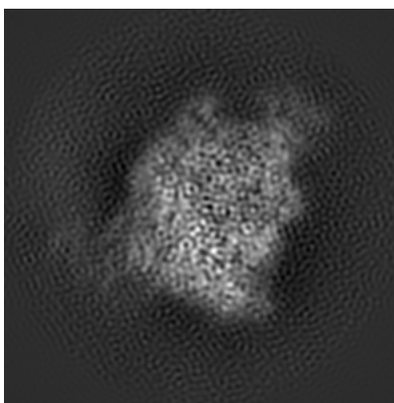
Images derived from a raw map, generated by summing the deposited half-maps, are presented below the corresponding image components of the primary map to allow further visual inspection and comparison with those of the primary map.

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

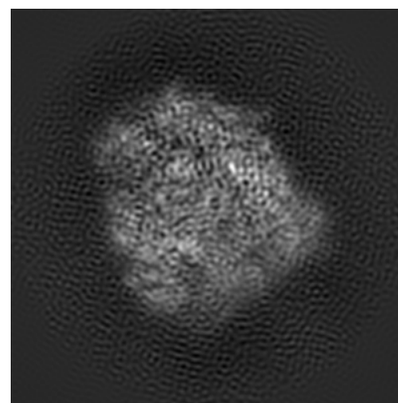
#### 6.1.1 Primary map



X

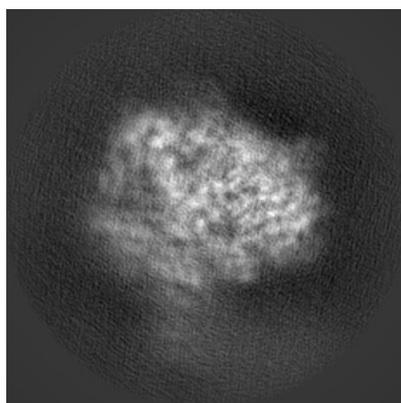


Y

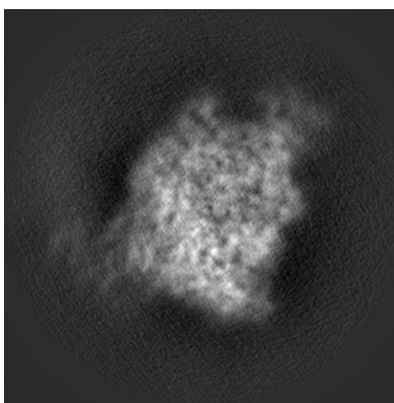


Z

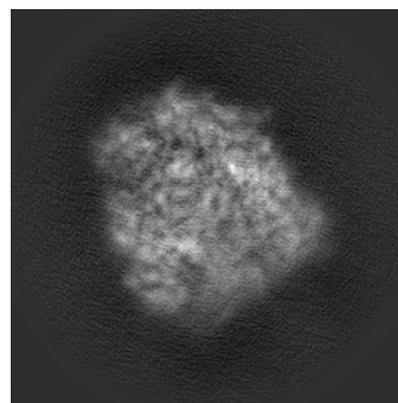
#### 6.1.2 Raw map



X



Y

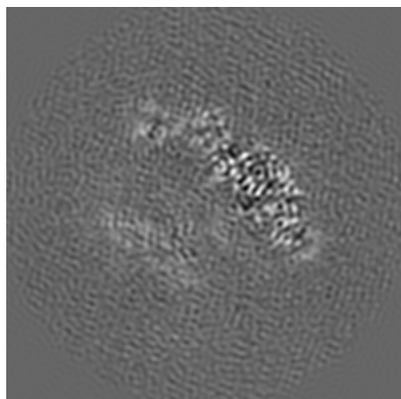


Z

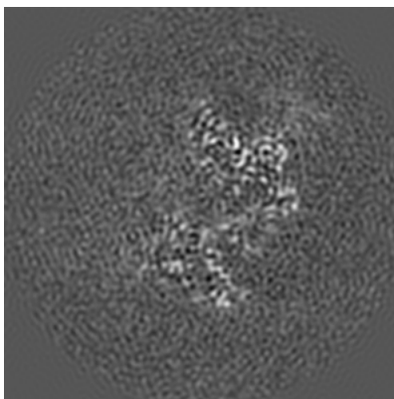
The images above show the map projected in three orthogonal directions.

## 6.2 Central slices [i](#)

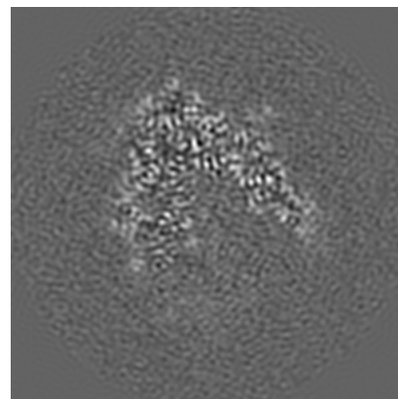
### 6.2.1 Primary map



X Index: 115

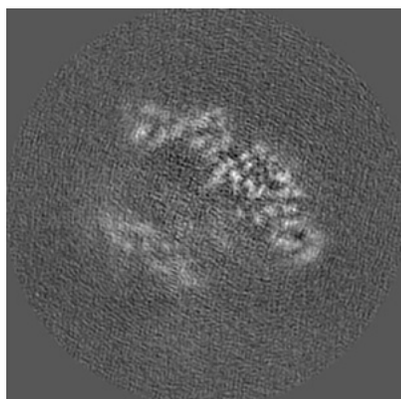


Y Index: 115

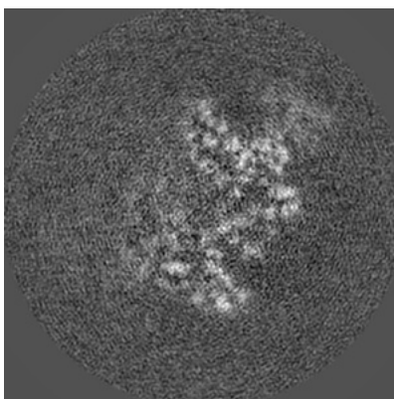


Z Index: 115

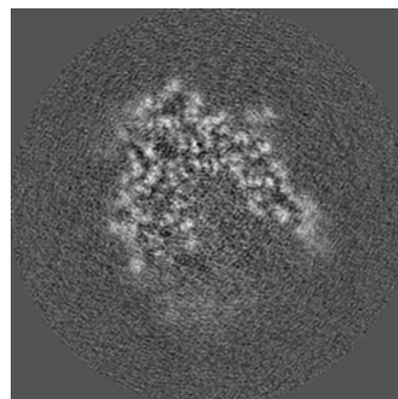
### 6.2.2 Raw map



X Index: 115



Y Index: 115

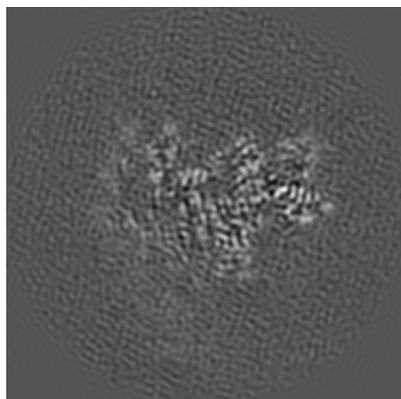


Z Index: 115

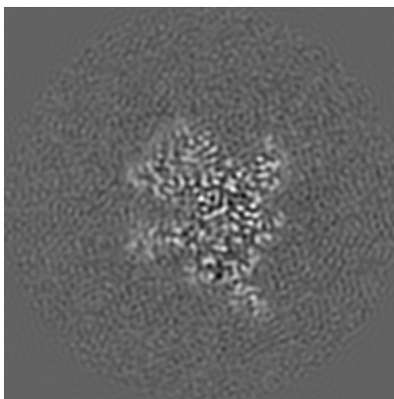
The images above show central slices of the map in three orthogonal directions.

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

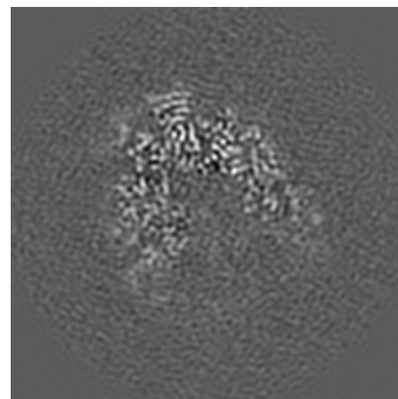
### 6.3.1 Primary map



X Index: 95

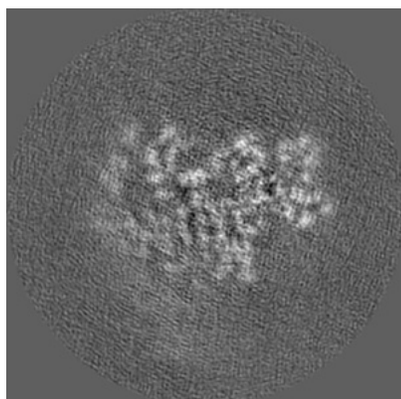


Y Index: 137

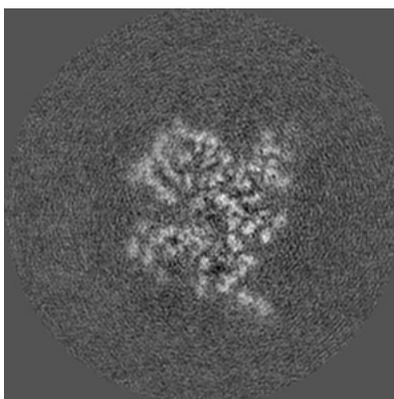


Z Index: 120

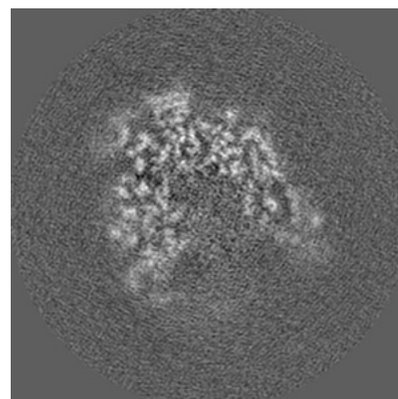
### 6.3.2 Raw map



X Index: 96



Y Index: 134

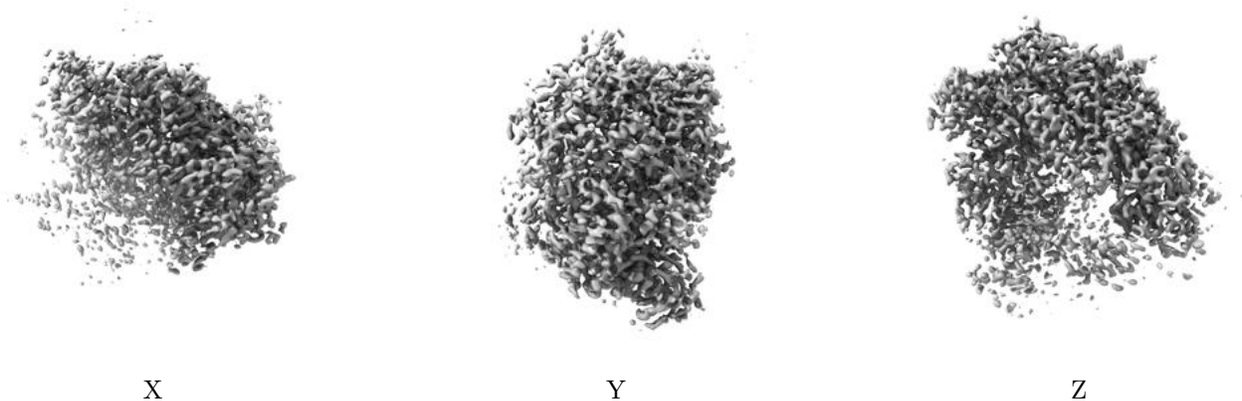


Z Index: 120

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

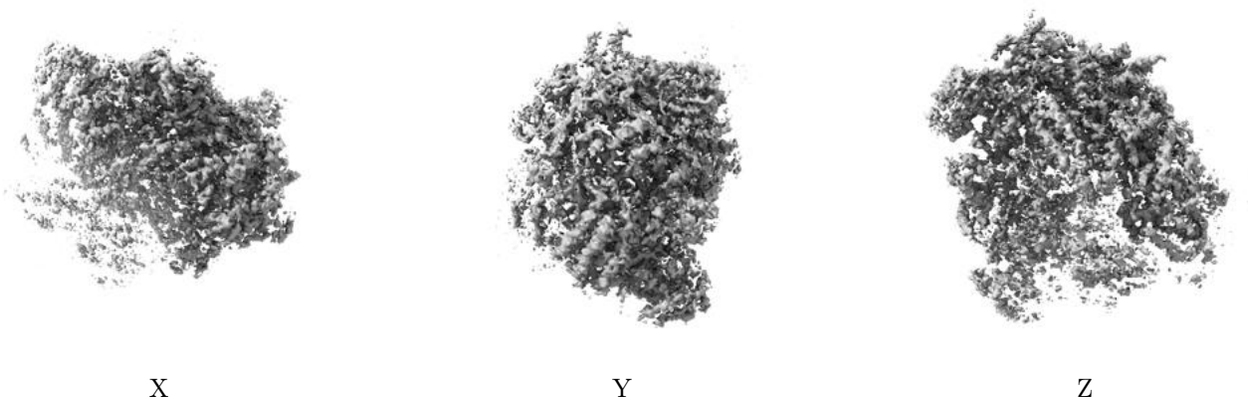
## 6.4 Orthogonal surface views [i](#)

### 6.4.1 Primary map



The images above show the 3D surface view of the map at the recommended contour level 0.04. These images, in conjunction with the slice images, may facilitate assessment of whether an appropriate contour level has been provided.

### 6.4.2 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.

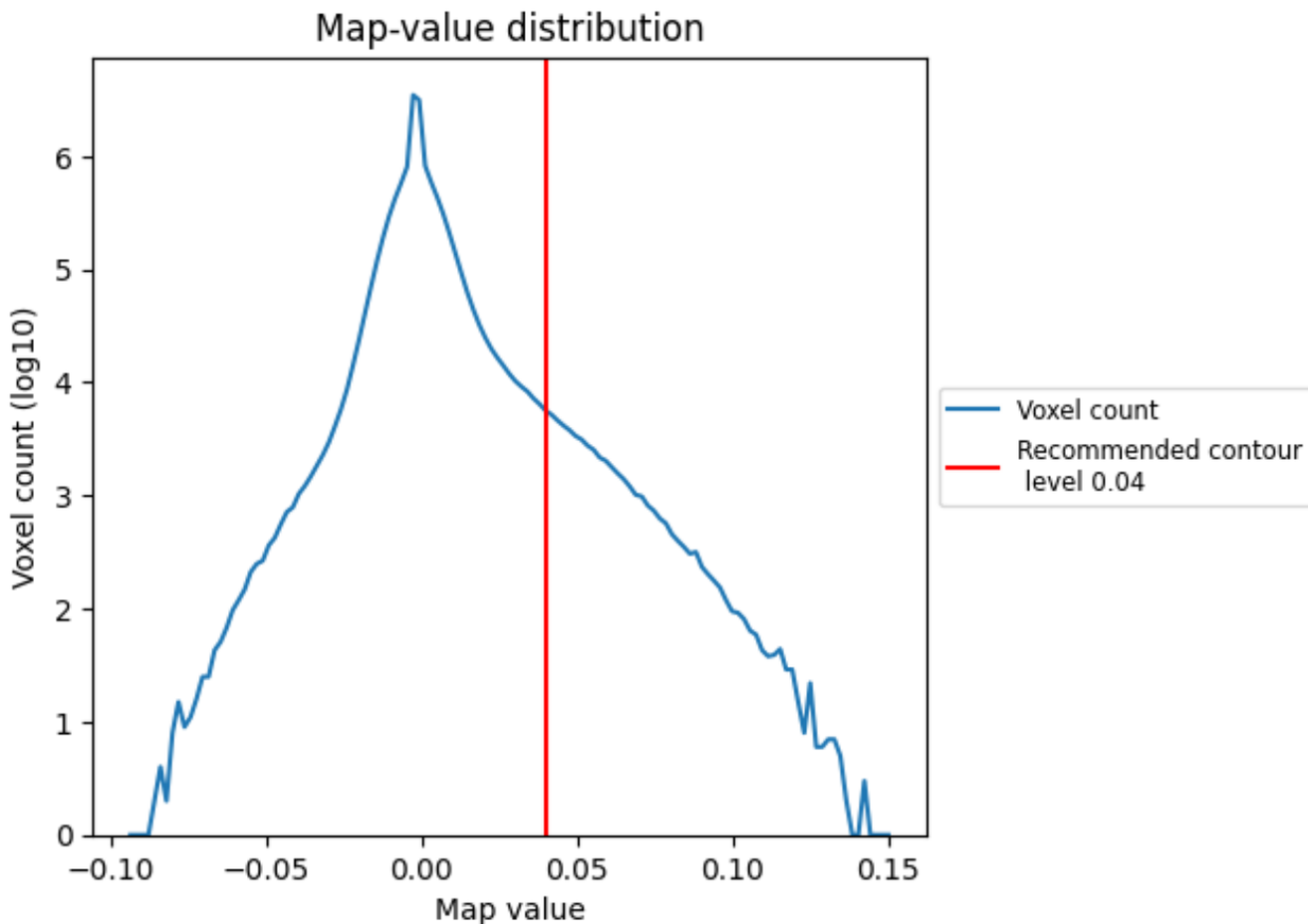
## 6.5 Mask visualisation [i](#)

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

## 7 Map analysis [i](#)

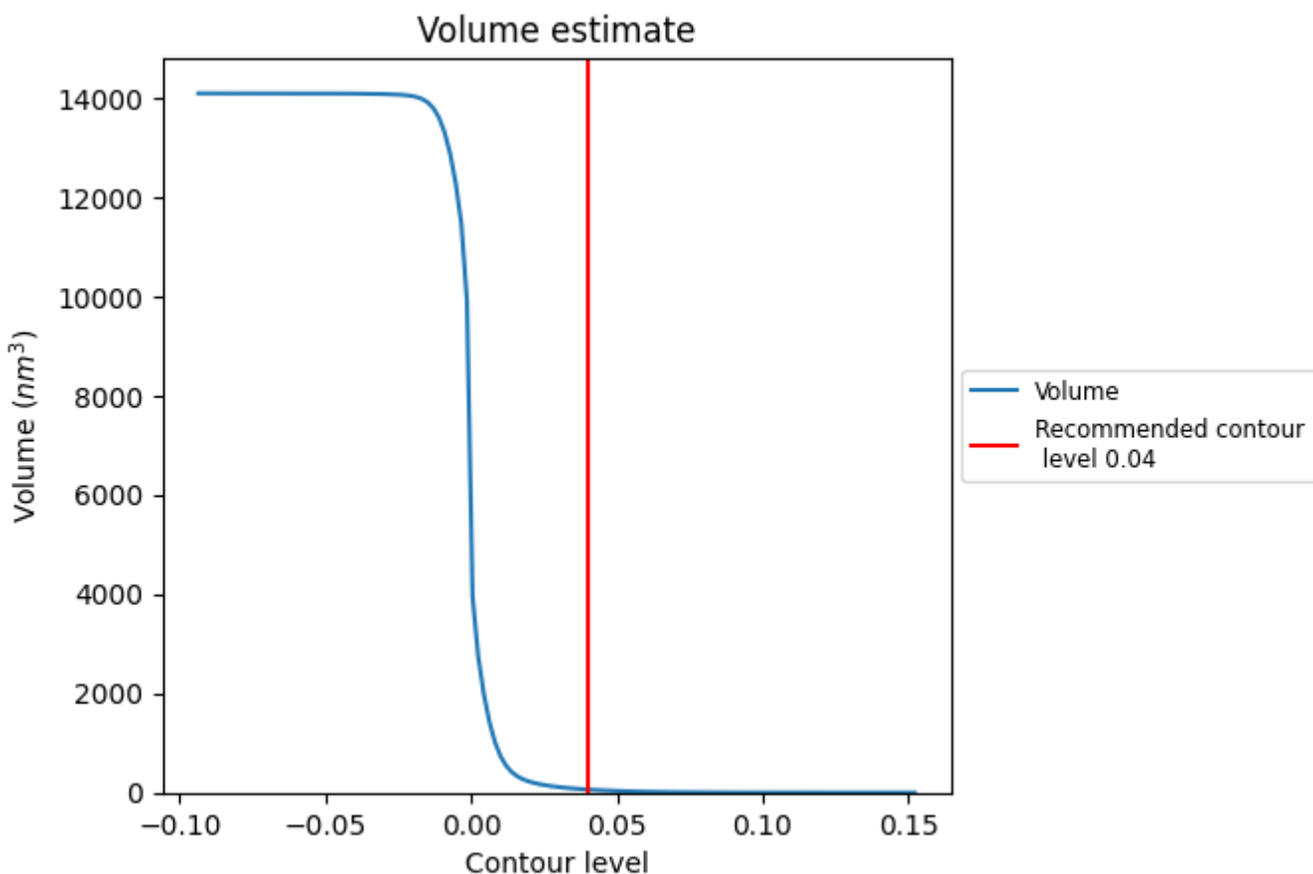
This section contains the results of statistical analysis of the map.

### 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.

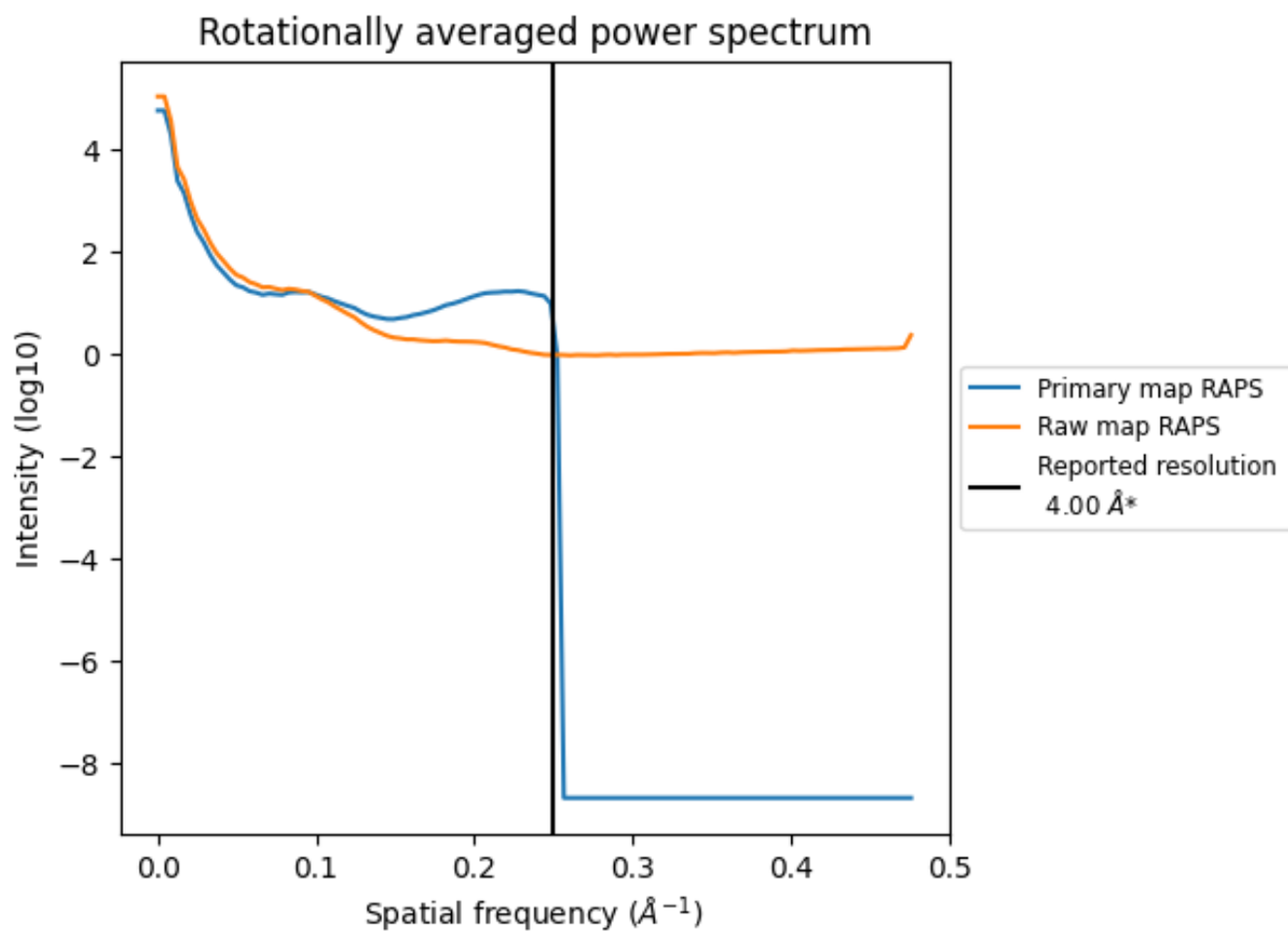
## 7.2 Volume estimate [\(i\)](#)



The volume at the recommended contour level is 62 nm<sup>3</sup>; this corresponds to an approximate mass of 56 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 i

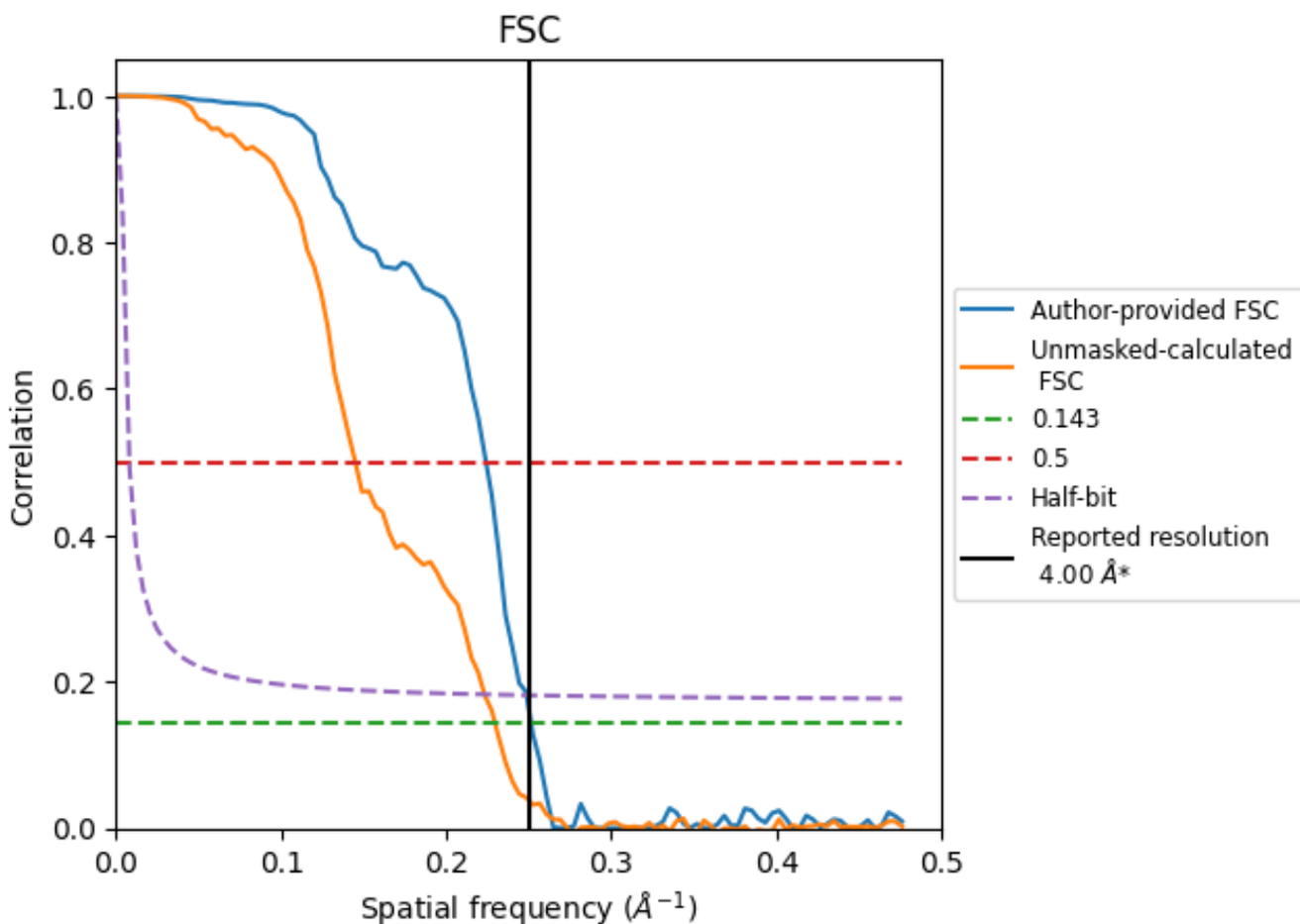


\*Reported resolution corresponds to spatial frequency of 0.250 Å<sup>-1</sup>

## 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 0.250 Å<sup>-1</sup>

## 8.2 Resolution estimates [i](#)

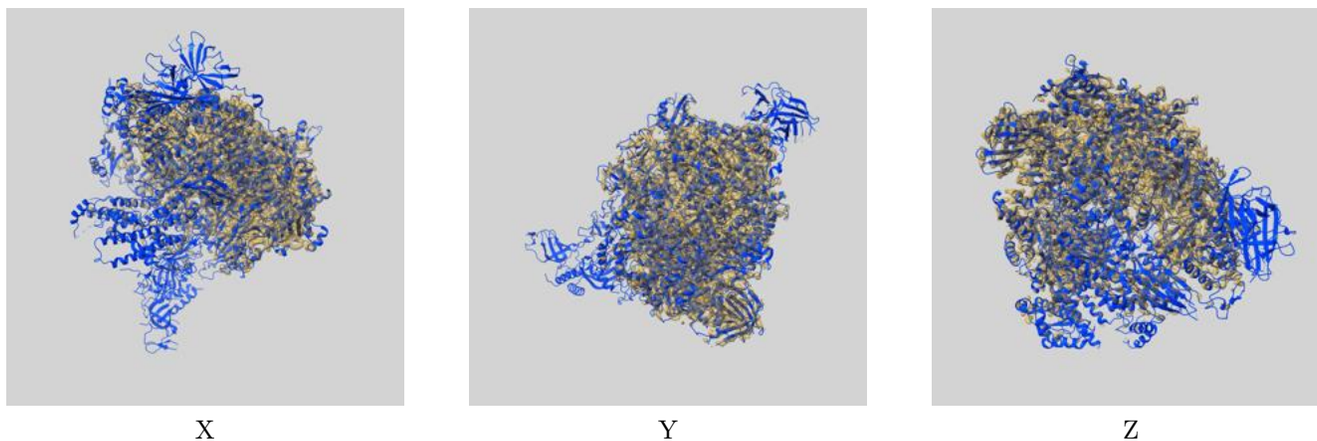
Resolution estimate (Å)	Estimation criterion (FSC cut-off)		
	0.143	0.5	Half-bit
Reported by author	4.00	-	-
Author-provided FSC curve	3.98	4.46	4.02
Unmasked-calculated*	4.35	6.89	4.48

\*Resolution estimate based on FSC curve calculated by comparison of deposited half-maps.

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

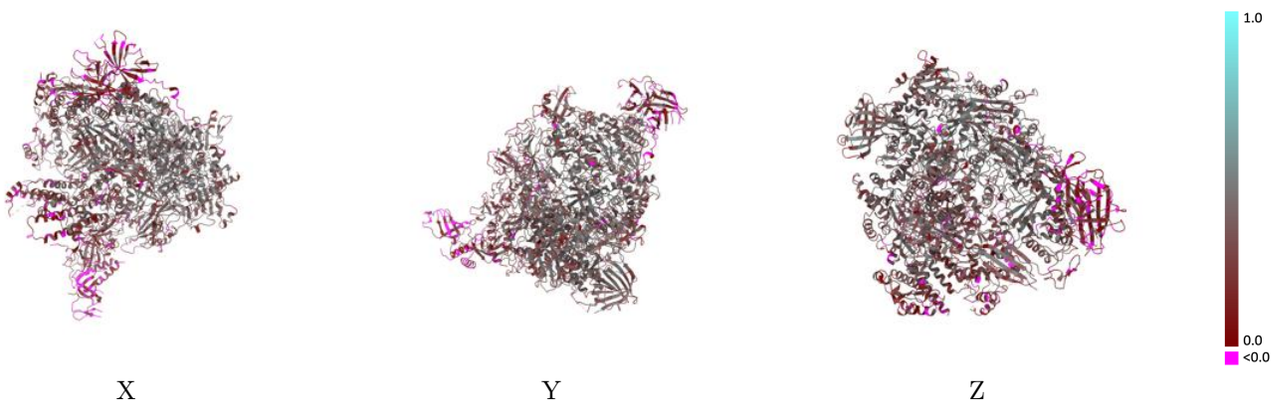
This section contains information regarding the fit between EMDB map EMD-4148 and PDB model 5M3M. Per-residue inclusion information can be found in section [3](#) on page [7](#).

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



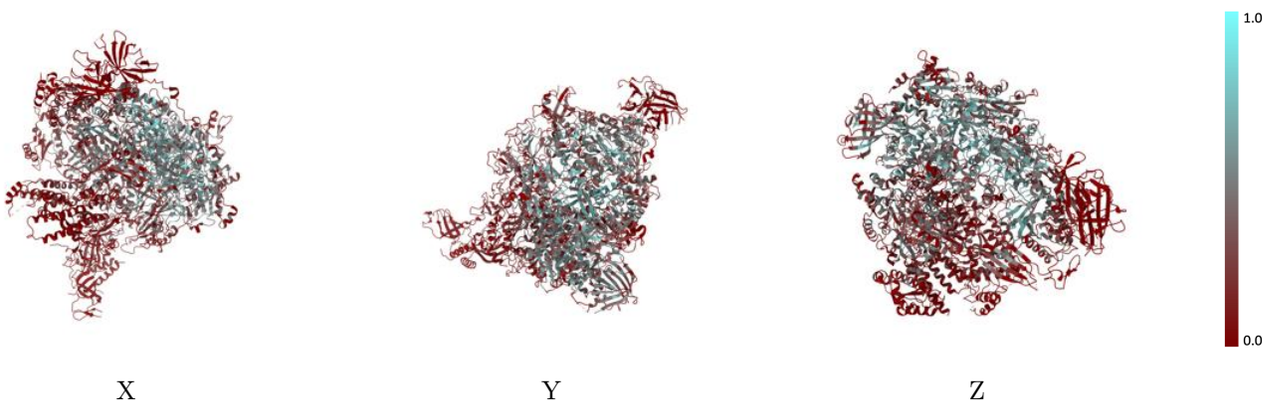
The images above show the 3D surface view of the map at the recommended contour level 0.04 at 50% transparency in yellow overlaid with a ribbon representation of the model coloured in blue. These images allow for the visual assessment of the quality of fit between the atomic model and the map.

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



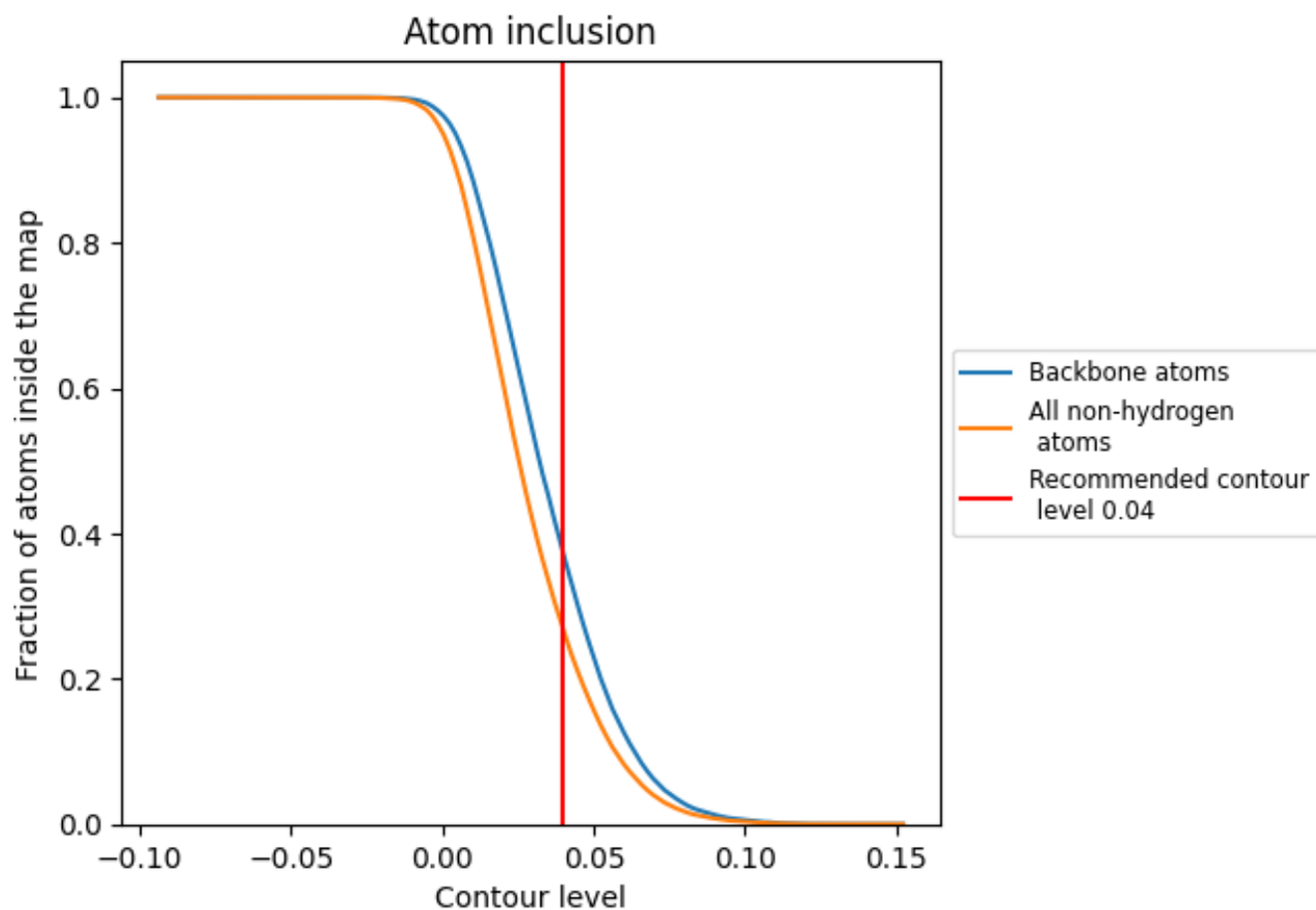
The images above show the model with each residue coloured according to its Q-score. This shows their resolvability in the map with higher Q-score values reflecting better resolvability. Please note: Q-score is calculating the resolvability of atoms, and thus high values are only expected at resolutions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

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



The images above show the model with each residue coloured according to its atom inclusion. This shows to what extent they are inside the map at the recommended contour level (0.04).

## 9.4 Atom inclusion [i](#)



At the recommended contour level, 37% of all backbone atoms, 27% 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 (0.04) and Q-score for the entire model and for each chain.

Chain	Atom inclusion	Q-score
All	 0.2669	 0.3090
A	 0.2526	 0.3180
B	 0.3859	 0.3650
C	 0.3501	 0.3600
D	 0.0044	 0.0810
E	 0.1454	 0.2700
F	 0.1924	 0.2950
G	 0.0080	 0.1250
H	 0.3411	 0.3680
I	 0.0627	 0.2470
J	 0.4728	 0.3920
K	 0.3929	 0.3660
L	 0.3353	 0.3610
M	 0.0026	 0.1250
N	 0.0000	 0.0710

