



wwPDB EM Validation Summary Report ⓘ

Oct 20, 2024 – 11:20 PM EDT

PDB ID : 8EYY
EMDB ID : EMD-28724
Title : Cryo-EM structure of 4 insulins bound full-length mouse IR mutant with physically decoupled alpha CTs (C684S/C685S/C687S, denoted as IR-3CS) Asymmetric conformation 2
Authors : Li, J.; Wu, J.Y.; Hall, C.; Bai, X.C.; Choi, E.
Deposited on : 2022-10-29
Resolution : 4.90 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

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.dev113
MolProbity : 4.02b-467
Percentile statistics : 20231227.v01 (using entries in the PDB archive December 27th 2023)
MapQ : 1.9.13
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP) : 2.39

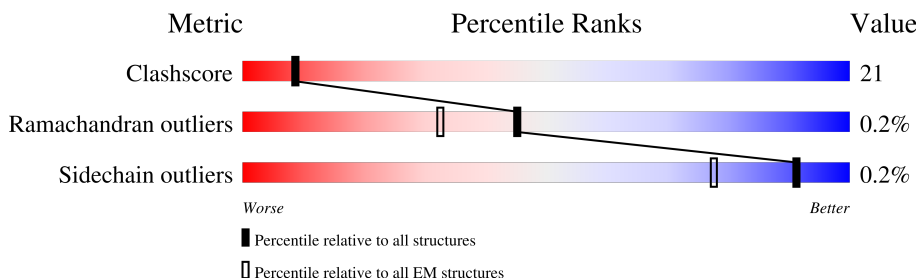
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

ELECTRON MICROSCOPY

The reported resolution of this entry is 4.90 Å.

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



Metric	Whole archive (#Entries)	EM structures (#Entries)
Clashscore	210492	15764
Ramachandran outliers	207382	16835
Sidechain outliers	206894	16415

The table below summarises the geometric issues observed across the polymeric chains and their fit to the map. The red, orange, yellow and green segments of the bar indicate the fraction of residues that contain outliers for ≥ 3 , 2, 1 and 0 types of geometric quality criteria respectively. A grey segment represents the fraction of residues that are not modelled. The numeric value for each fraction is indicated below the corresponding segment, with a dot representing fractions $\leq 5\%$. The upper red bar (where present) indicates the fraction of residues that have poor fit to the EM map (all-atom inclusion $< 40\%$). The numeric value is given above the bar.

Mol	Chain	Length	Quality of chain
1	A	1345	
1	B	1345	
2	C	110	
2	D	110	
2	E	110	
2	F	110	

2 Entry composition

There are 2 unique types of molecules in this entry. The entry contains 14301 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 Insulin receptor.

Mol	Chain	Residues	Atoms					AltConf	Trace
1	A	775	Total	C	N	O	S	0	0
			6284	3995	1080	1168	41		
1	B	810	Total	C	N	O	S	0	0
			6540	4151	1127	1213	49		

There are 6 discrepancies between the modelled and reference sequences:

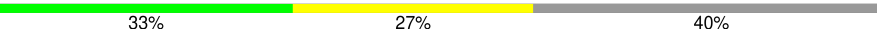
Chain	Residue	Modelled	Actual	Comment	Reference
A	684	SER	CYS	engineered mutation	UNP P15208
A	685	SER	CYS	engineered mutation	UNP P15208
A	687	SER	CYS	engineered mutation	UNP P15208
B	684	SER	CYS	engineered mutation	UNP P15208
B	685	SER	CYS	engineered mutation	UNP P15208
B	687	SER	CYS	engineered mutation	UNP P15208

- Molecule 2 is a protein called Insulin.

Mol	Chain	Residues	Atoms					AltConf	Trace
2	C	47	Total	C	N	O	S	0	0
			370	235	60	69	6		
2	D	46	Total	C	N	O	S	0	0
			361	229	58	68	6		
2	E	48	Total	C	N	O	S	0	0
			376	238	61	71	6		
2	F	47	Total	C	N	O	S	0	0
			370	235	60	69	6		

Y802	VAL	ASN	VAL	SER	PRO	GLY
R806	GLN	ALA	ALA	PHE	THR	GLY
T807	VAL	LYS	ALA	GLY	THR	GLY
M808	ASP	ASP	ALA	VAL	THR	GLY
D814	LEU	ILE	VAL	TRP	VAL	GLY
V815	GLY	LYS	VAL	GLU	ASP	GLY
I816	MET	GLY	ALA	GLY	ASP	GLY
V820	ASP	ALA	ASP	THR	THR	GLY
T821	LEU	THR	THR	LEU	THR	GLY
I824	TYR	ALA	VAL	ALA	ASP	GLY
F825	SER	VAL	LYS	GLU	ASP	GLY
N828	ARG	ASN	ILE	GLN	ASP	GLY
V829	PRO	VAL	ASP	PRO	ASP	GLY
V830	GLY	LYS	PHE	GLN	GLY	GLY
H831	TYR	VAL	GLY	GLY	GLY	GLY
L832	LEU	ASN	PRO	LEU	GLY	GLY
M833	SER	ALA	ASP	GLY	GLY	GLY
M834	ASP	ALA	ASP	GLY	GLY	GLY
P837	VAL	SER	ILE	VAL	VAL	GLY
K838	ASP	ARG	GLY	LEU	PRO	GLY
E839	TYR	GLU	THR	GLY	GLY	GLY
P840	LEU	ARG	ASP	PHE	GLY	GLY
N841	ASP	SER	VAL	VAL	GLY	GLY
G842	VAL	ILE	TYR	PRO	GLY	GLY
L843	VAL	GLU	TYR	PRO	GLY	GLY
I844	PRO	PHE	ASP	ASP	GLY	GLY
V845	SER	LEU	LYS	GLY	GLY	GLY
V849	ASN	PRO	GLY	GLY	GLY	GLY
S850	LYS	ALA	GLY	GLY	GLY	GLY
Y851	ILE	ALA	GLY	GLY	GLY	GLY
R852	ILE	VAL	ILE	PRO	PRO	GLY
R853	ILE	VAL	GLN	PRO	PRO	GLY
Y854	ILE	VAL	GLN	PRO	PRO	GLY
G855	PRO	GLY	GLY	GLY	GLY	GLY
D856	LEU	GLY	GLY	GLY	GLY	GLY
E857	ILE	ILE	ILE	GLY	GLY	GLY
E858	PHE	THR	THR	THR	THR	GLY
L859	VAL	VAL	VAL	VAL	VAL	GLY
V863	PHE	LEU	VAL	VAL	VAL	GLY
S864	PHE	GLY	VAL	VAL	VAL	GLY
R865	SER	LEU	VAL	VAL	VAL	GLY
K866	VAL	VAL	VAL	VAL	VAL	GLY
H867	VAL	VAL	VAL	VAL	VAL	GLY
E871	GLY	GLY	VAL	VAL	VAL	GLY
R872	LYS	SER	THR	THR	THR	GLY
G873	ILE	ILE	GLY	GLY	GLY	GLY
C874	TYR	VAL	VAL	VAL	VAL	GLY
R875	LEU	VAL	VAL	VAL	VAL	GLY
L876	PHE	TYR	VAL	VAL	VAL	GLY
L879	ARG	GLY	THR	THR	THR	GLY

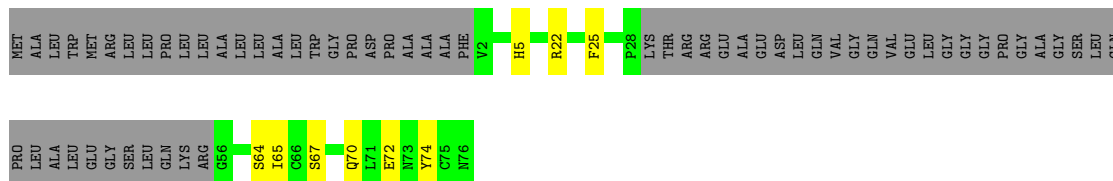
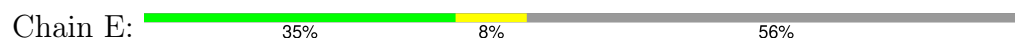
• Molecule 1: Insulin receptor

Chain B:  33% 27% 40%

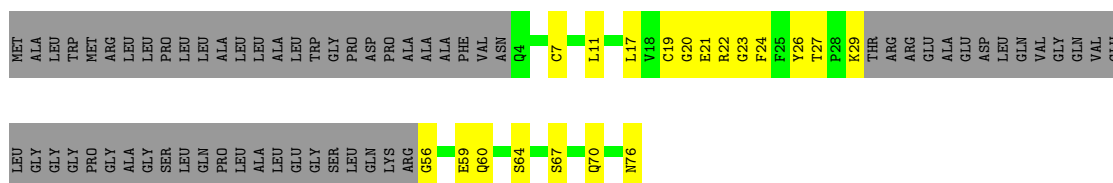
H1	Y91	P170	L262	I370	M462	P548
G5	A92	A171	H263	R371	G463	S549
R14	I93	I172	R267	S372	D464	H550
R19	Y94	V173	R275	S373	Q465	P551
S27	I95	I174	R276	L379	E469	G552
E30	F96	N175	H277	S380	M470	N553
R32	E97	R181	Q276	E471	M471	L554
L33	H98	C182	V277	L472	L472	M555
Q34	H100	W183	V278	L473	L473	R556
I35	L101	Q189	N281	K474	K474	G557
L36	K102	K190	N282	F475	F475	L558
L37	L104	L191	C283	S476	S476	N561
M38	L106	C192	C284	F477	F477	T562
L39	Y107	I195	I285	R478	R478	Q563
F39	N108	I199	E287	R479	R479	Y564
R40	L109	G200	C288	E394	E394	A565
T41	N110	C201	P289	Y398	Y398	I566
R42	N111	T202	S290	S399	S399	F567
P43	I112	A203	Y292	F400	F400	V568
E44	T113	C207	N294	N405	N405	R569
S50	R114	C208	G301	R409	R409	L571
F51	S116	H209	K310	Q410	Q410	F574
P52	V117	L210	L310	L411	L411	E577
K53	I119	E211	V311	W412	W412	R578
L54	E120	C212	G312	D413	D413	R579
L57	K121	L213	Q313	W414	W414	T580
T58	N122	D221	L314	T420	T420	Y581
L62	N123	P222	L315	L426	L426	G582
L63	E124	T223	G317	F428	F428	A583
F64	C126	K224	E318	H429	H429	K584
R65	R135	A227	T321	K433	K433	S585
V66	I136	C228	D322	L436	L436	I588
Y67	L137	R229	S323	ASP	ASP	T592
L72	V140	N230	V324	H440	H440	D593
K73	Y144	L233	T325	Y441	Y441	A594
D74	I145	Q236	Q328	M442	M442	P597
L75	N148	E239	V335	E443	E443	S598
P77	K149	N245	N337	E444	E444	V599
N78	D150	Y245	T341	V445	V445	P600
T80	P160	Y246	T342	T448	T448	L601
V81	G161	F248	R345	P537	P537	S609
R82	T162	Q249	R345	Q612	Q612	Q612
G84	ALA	D250	E353	T613	T613	I614
L87	LYS	R251	E353	L614	L614	L615
F88	GLY	R252	E362	K615	K615	K616
F89	LYS	C253	S365	W617	W617	K618
N90	THR	F258		K618	K618	D622
	C169			P623	P623	P623



- Molecule 2: Insulin



- Molecule 2: Insulin



4 Experimental information

Property	Value	Source
EM reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, C1	Depositor
Number of particles used	104347	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 ($e^-/\text{\AA}^2$)	60	Depositor
Minimum defocus (nm)	1600	Depositor
Maximum defocus (nm)	2600	Depositor
Magnification	Not provided	
Image detector	GATAN K3 BIOQUANTUM (6k x 4k)	Depositor
Maximum map value	0.110	Depositor
Minimum map value	-0.040	Depositor
Average map value	0.000	Depositor
Map value standard deviation	0.003	Depositor
Recommended contour level	0.018	Depositor
Map size (Å)	324.0, 324.0, 324.0	wwPDB
Map dimensions	300, 300, 300	wwPDB
Map angles (°)	90.0, 90.0, 90.0	wwPDB
Pixel spacing (Å)	1.08, 1.08, 1.08	Depositor

5 Model quality [i](#)

5.1 Standard geometry [i](#)

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	0.25	0/6437	0.50	0/8722
1	B	0.26	0/6703	0.51	0/9091
2	C	0.25	0/377	0.50	0/508
2	D	0.29	0/368	0.45	0/497
2	E	0.26	0/383	0.47	0/518
2	F	0.25	0/377	0.45	0/508
All	All	0.26	0/14645	0.50	0/19844

There are no bond length outliers.

There are no bond angle outliers.

There are no chirality outliers.

There are no planarity outliers.

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	6284	0	6118	264	0
1	B	6540	0	6344	280	0
2	C	370	0	344	23	0
2	D	361	0	331	18	0
2	E	376	0	346	7	0
2	F	370	0	344	14	0
All	All	14301	0	13827	585	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 21.

The worst 5 of 585 close contacts within the same asymmetric unit are listed below, sorted by their clash magnitude.

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:886:VAL:O	1:B:902:PRO:HA	1.72	0.88
1:A:354:LEU:O	1:A:358:LEU:HB2	1.83	0.78
1:B:853:ARG:NH1	1:B:881:PRO:O	2.18	0.77
1:A:245:TYR:HB3	1:A:253:CYS:HB3	1.68	0.76
1:B:547:THR:O	1:B:549:SER:N	2.19	0.75

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	761/1345 (57%)	710 (93%)	49 (6%)	2 (0%)	37	72
1	B	798/1345 (59%)	731 (92%)	66 (8%)	1 (0%)	48	83
2	C	43/110 (39%)	37 (86%)	6 (14%)	0	100	100
2	D	42/110 (38%)	42 (100%)	0	0	100	100
2	E	44/110 (40%)	44 (100%)	0	0	100	100
2	F	43/110 (39%)	39 (91%)	3 (7%)	1 (2%)	5	28
All	All	1731/3130 (55%)	1603 (93%)	124 (7%)	4 (0%)	45	78

All (4) Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	B	548	PRO
1	A	532	VAL
1	A	794	ASP
2	F	26	TYR

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	708/1211 (58%)	707 (100%)	1 (0%)	92	94
1	B	737/1211 (61%)	734 (100%)	3 (0%)	89	91
2	C	42/88 (48%)	42 (100%)	0	100	100
2	D	41/88 (47%)	41 (100%)	0	100	100
2	E	43/88 (49%)	43 (100%)	0	100	100
2	F	42/88 (48%)	42 (100%)	0	100	100
All	All	1613/2774 (58%)	1609 (100%)	4 (0%)	91	94

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

Mol	Chain	Res	Type
1	A	852	ARG
1	B	470	ASN
1	B	578	ARG
1	B	579	ARG

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. All (4) such sidechains are listed below:

Mol	Chain	Res	Type
1	A	34	GLN
1	A	629	HIS
1	B	230	ASN
2	E	76	ASN

5.3.3 RNA ⓘ

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 oligosaccharides in this entry.

5.6 Ligand geometry [i](#)

There are no ligands in this entry.

5.7 Other polymers [i](#)

There are no such residues in this entry.

5.8 Polymer linkage issues [i](#)

There are no chain breaks in this entry.

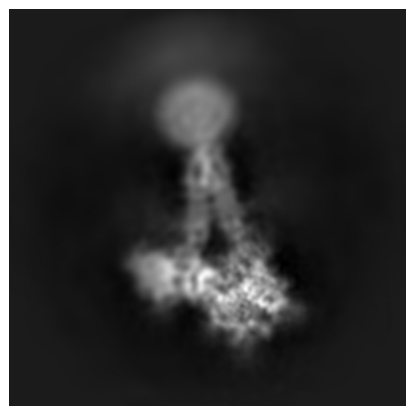
6 Map visualisation [i](#)

This section contains visualisations of the EMDB entry EMD-28724. These allow visual inspection of the internal detail of the map and identification of artifacts.

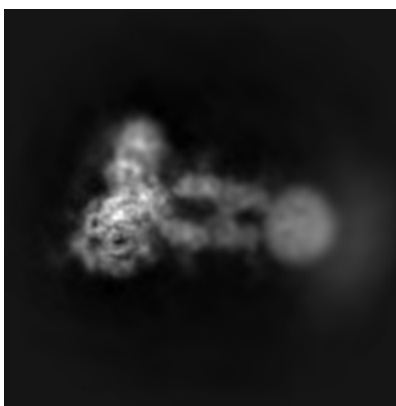
Images derived from a raw map, generated by summing the deposited half-maps, are presented below the corresponding image components of the primary map to allow further visual inspection and comparison with those of the primary map.

6.1 Orthogonal projections [i](#)

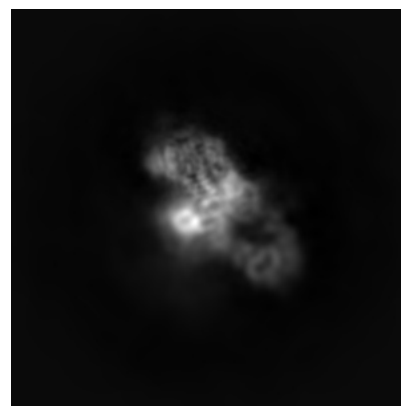
6.1.1 Primary map



X

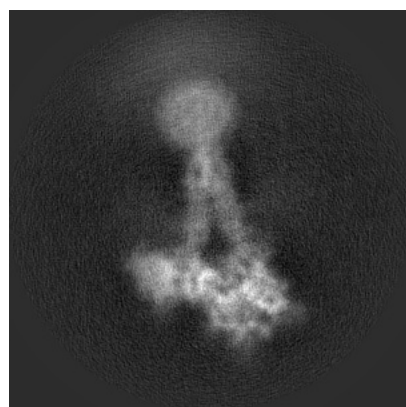


Y

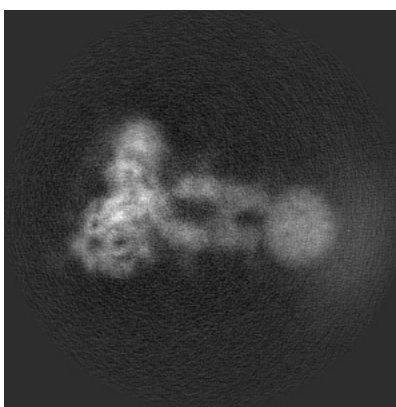


Z

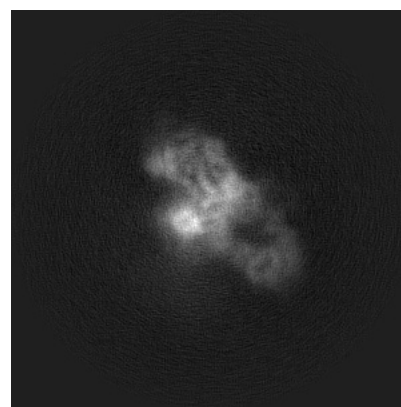
6.1.2 Raw map



X



Y

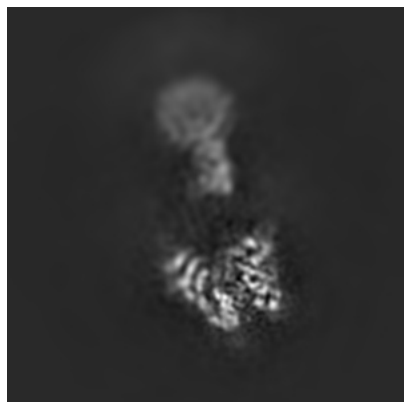


Z

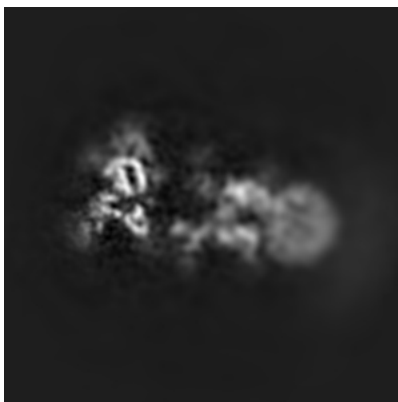
The images above show the map projected in three orthogonal directions.

6.2 Central slices [i](#)

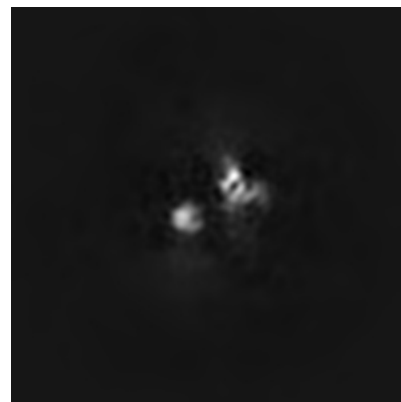
6.2.1 Primary map



X Index: 150

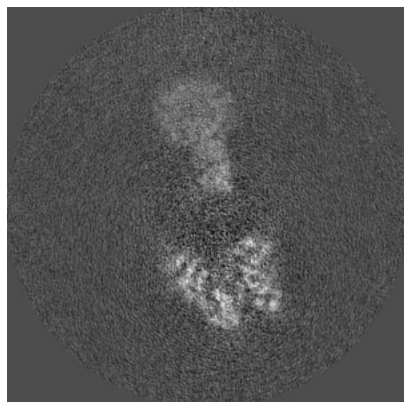


Y Index: 150

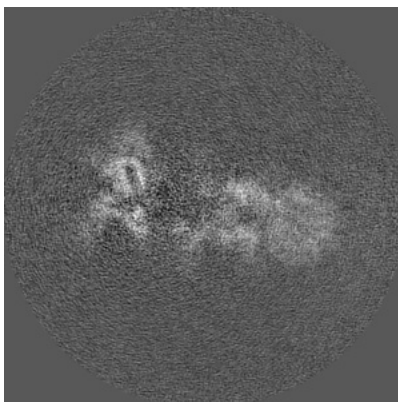


Z Index: 150

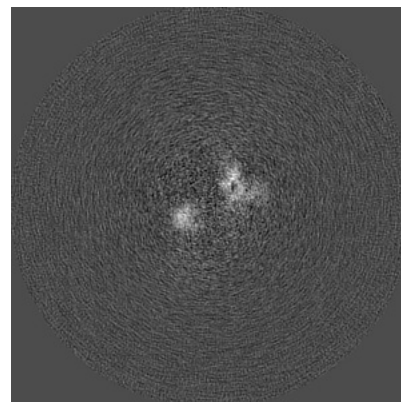
6.2.2 Raw map



X Index: 150



Y Index: 150

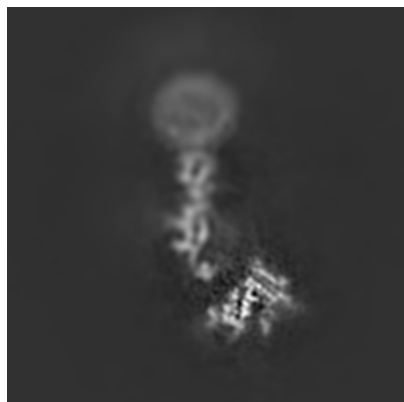


Z Index: 150

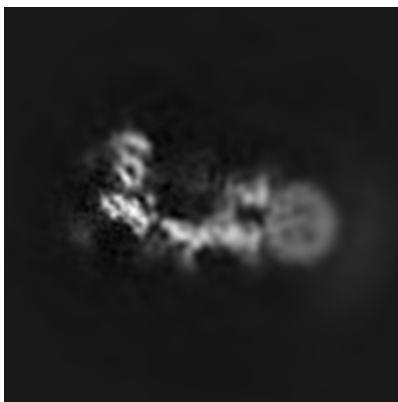
The images above show central slices of the map in three orthogonal directions.

6.3 Largest variance slices [i](#)

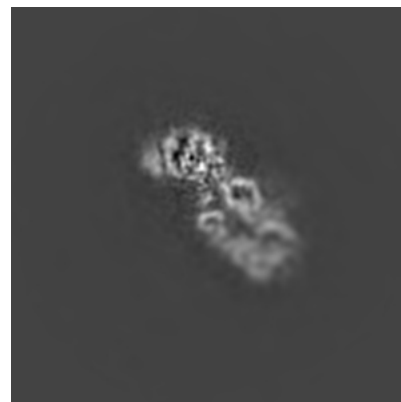
6.3.1 Primary map



X Index: 133

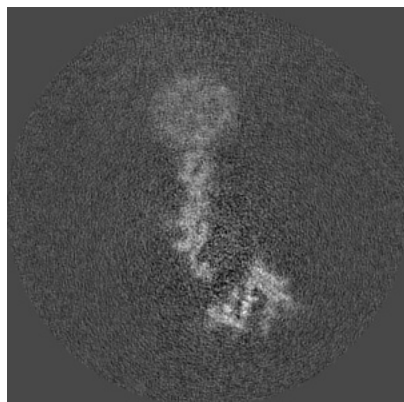


Y Index: 145

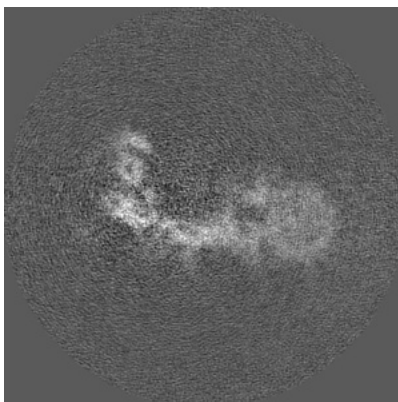


Z Index: 88

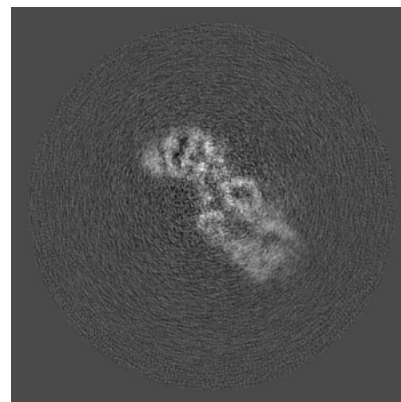
6.3.2 Raw map



X Index: 134



Y Index: 143



Z Index: 88

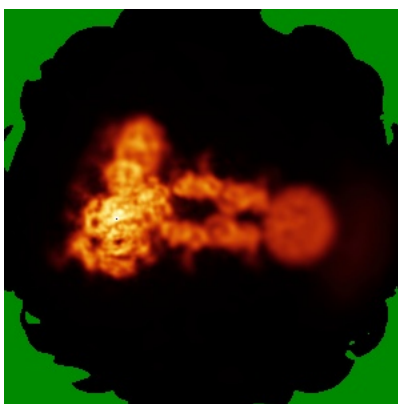
The images above show the largest variance slices of the map in three orthogonal directions.

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

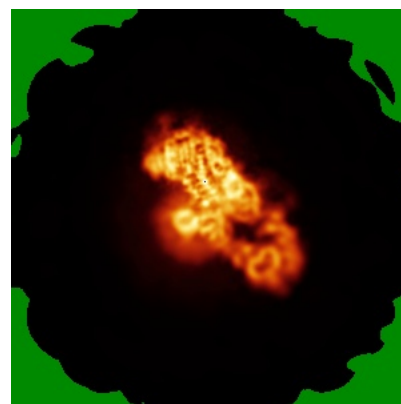
6.4.1 Primary map



X

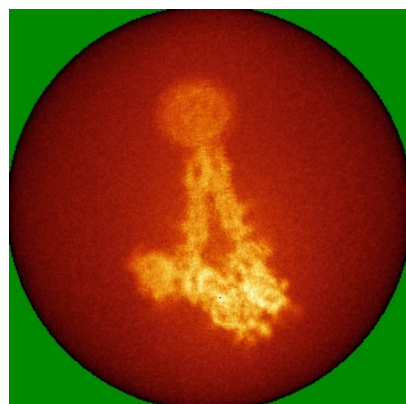


Y

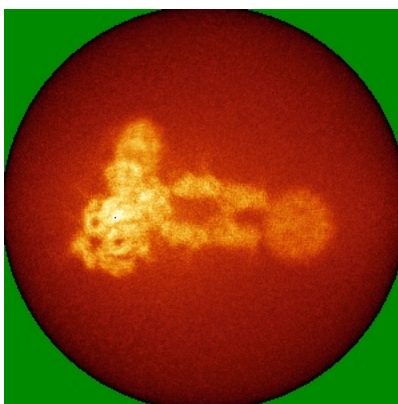


Z

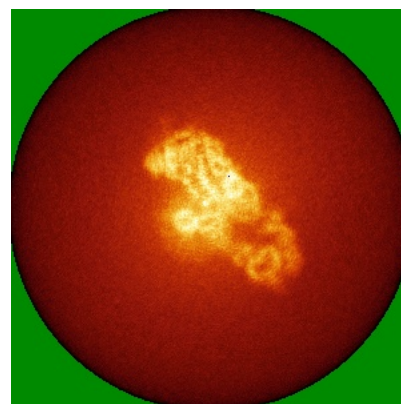
6.4.2 Raw map



X



Y

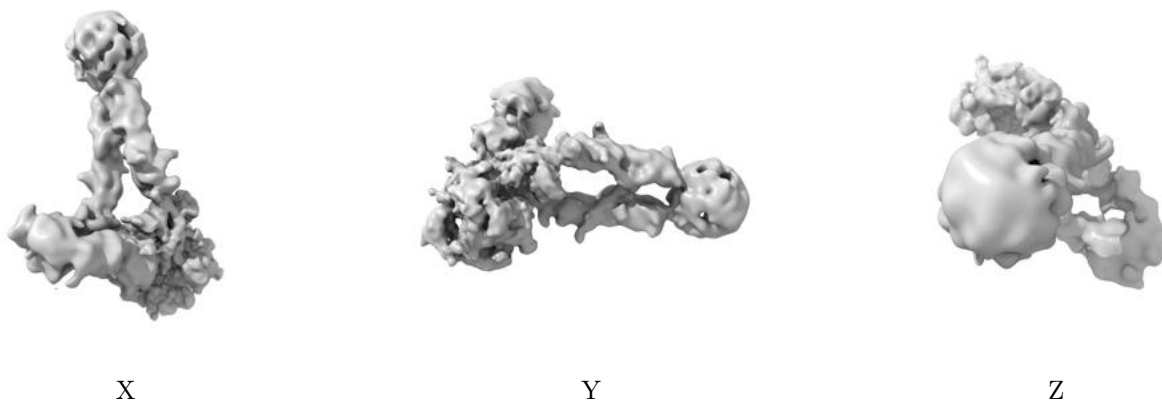


Z

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.

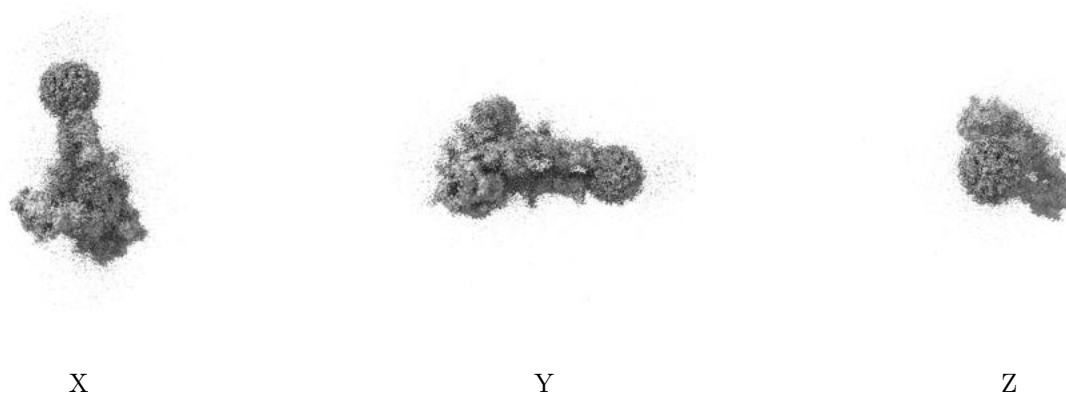
6.5 Orthogonal surface views [i](#)

6.5.1 Primary map



The images above show the 3D surface view of the map at the recommended contour level 0.018. 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.

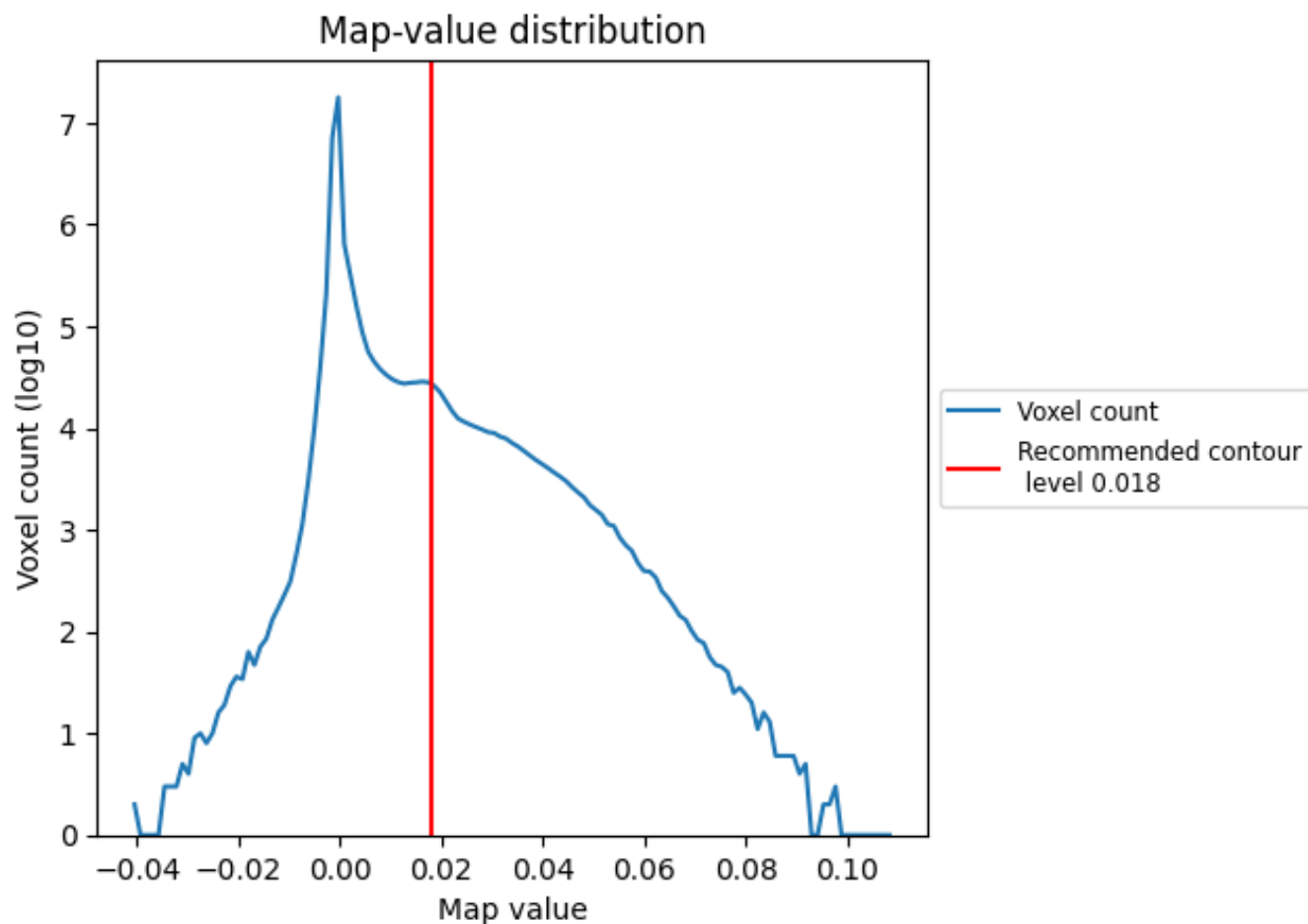
6.6 Mask visualisation [i](#)

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

7 Map analysis [i](#)

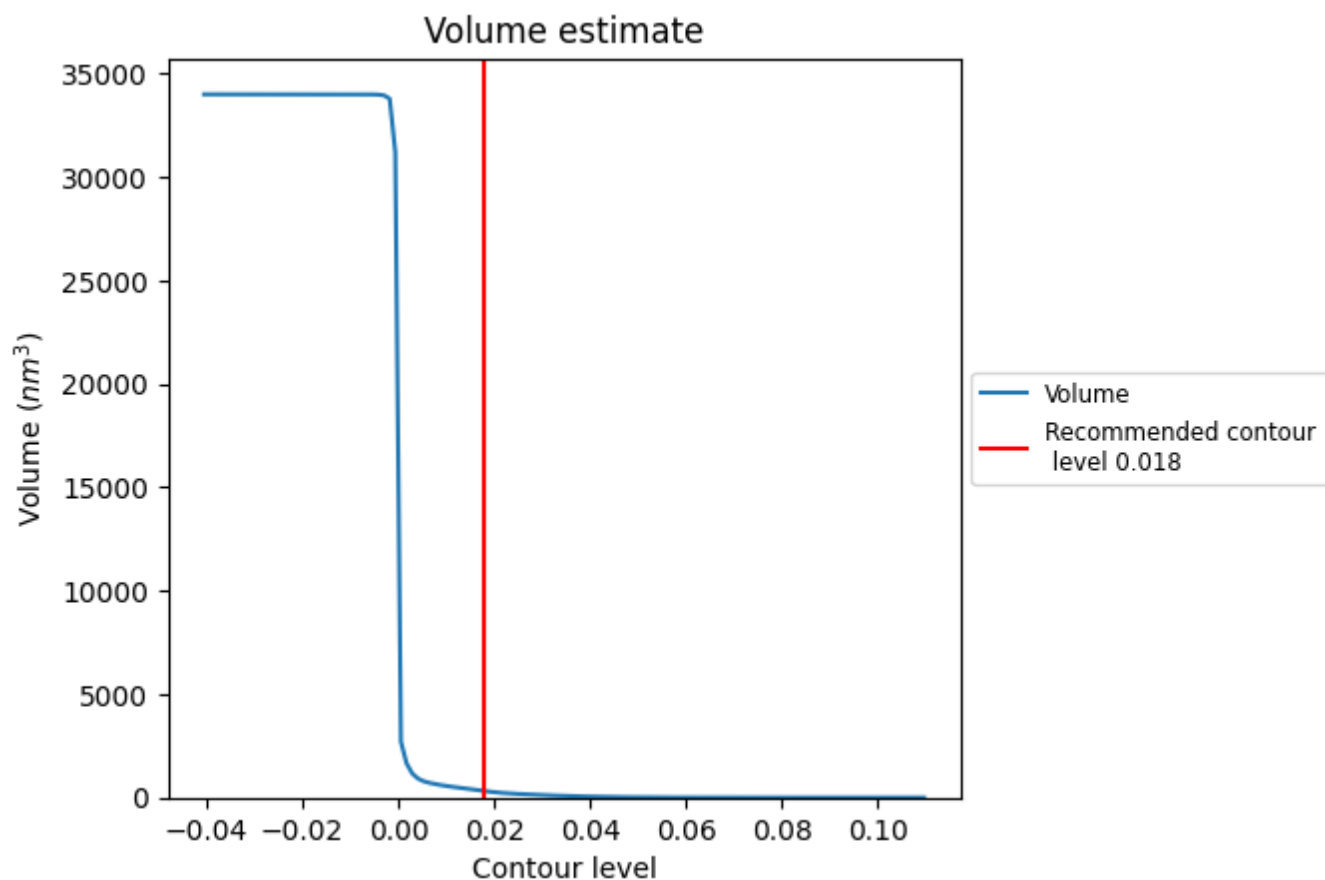
This section contains the results of statistical analysis of the map.

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.

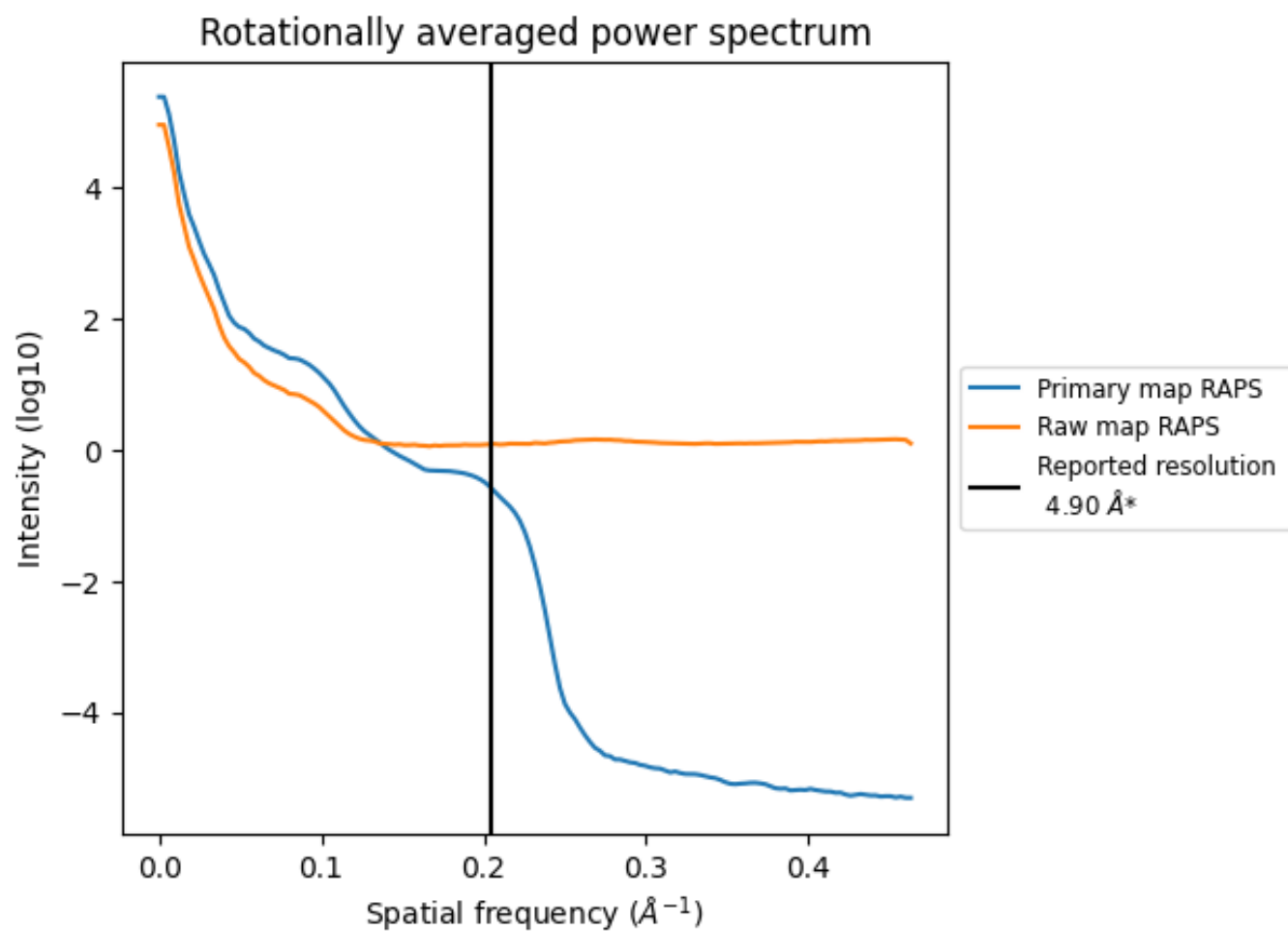
7.2 Volume estimate [i](#)



The volume at the recommended contour level is 317 nm³; this corresponds to an approximate mass of 287 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 ⓘ

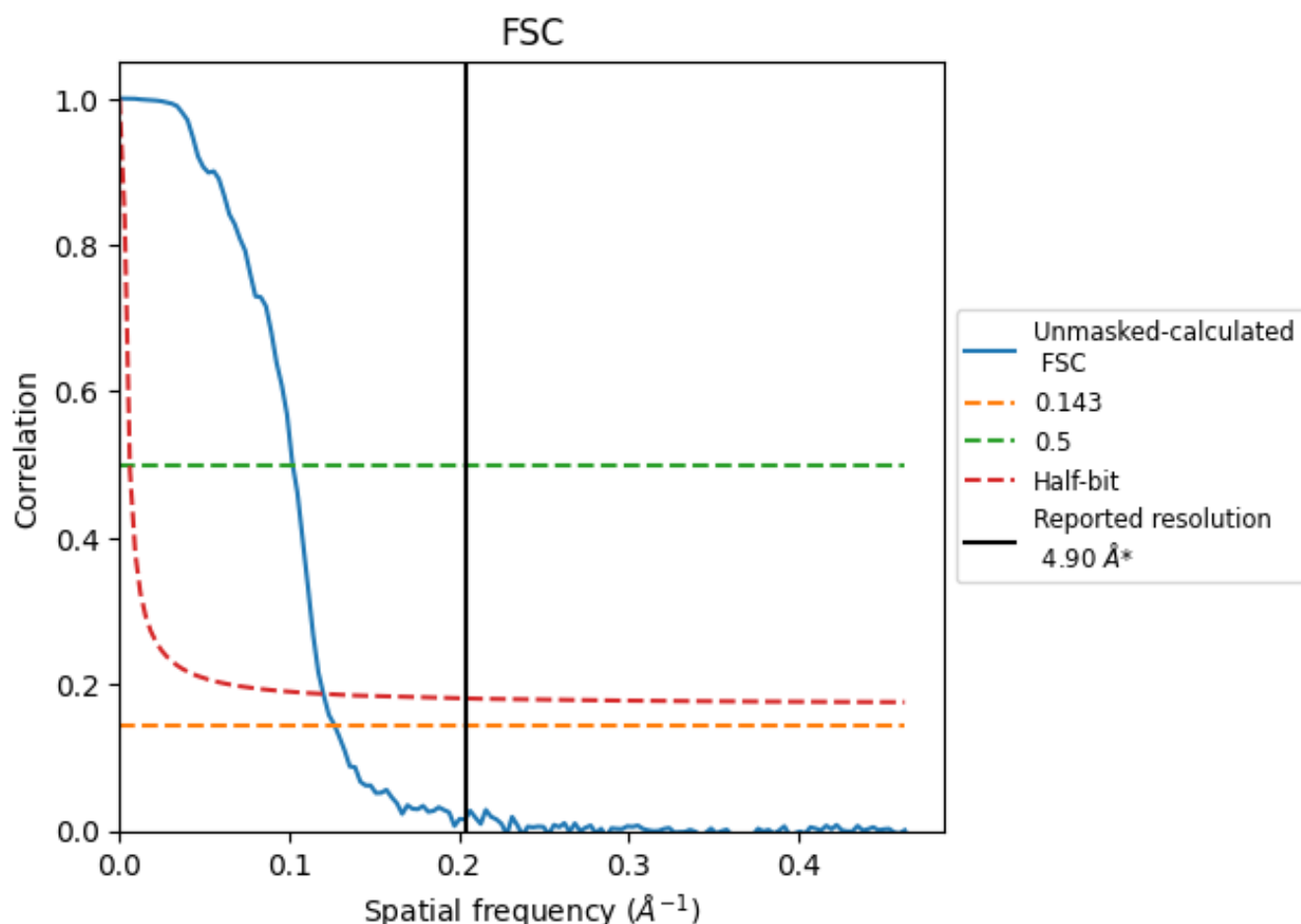


*Reported resolution corresponds to spatial frequency of 0.204 Å⁻¹

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 0.204 \AA^{-1}

8.2 Resolution estimates [i](#)

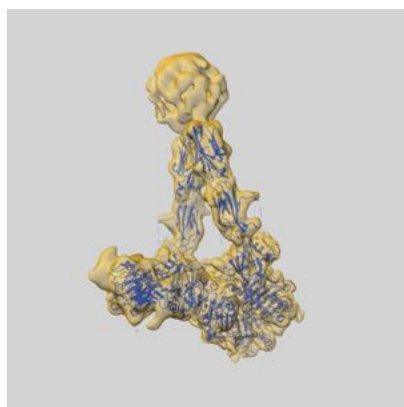
Resolution estimate (Å)	Estimation criterion (FSC cut-off)		
	0.143	0.5	Half-bit
Reported by author	4.90	-	-
Author-provided FSC curve	-	-	-
Unmasked-calculated*	7.87	9.79	8.32

*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 7.87 differs from the reported value 4.9 by more than 10 %

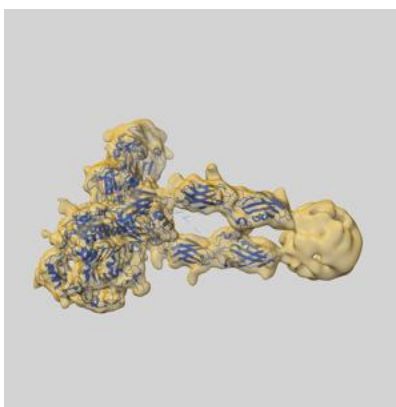
9 Map-model fit [i](#)

This section contains information regarding the fit between EMDB map EMD-28724 and PDB model 8EYY. Per-residue inclusion information can be found in [section 3](#) on [page 4](#).

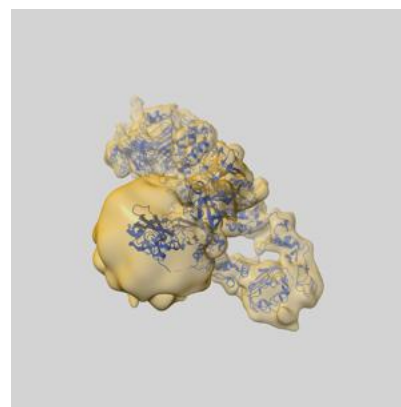
9.1 Map-model overlay [i](#)



X



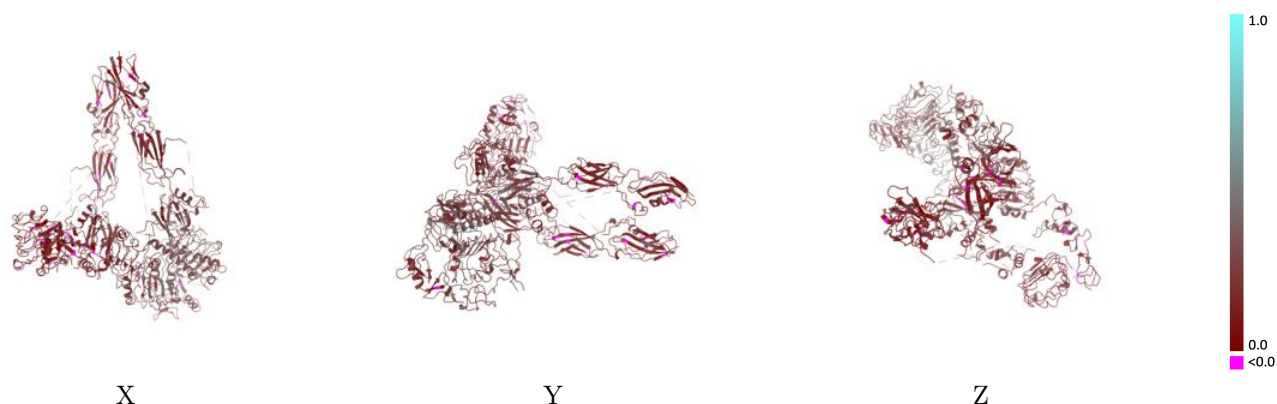
Y



Z

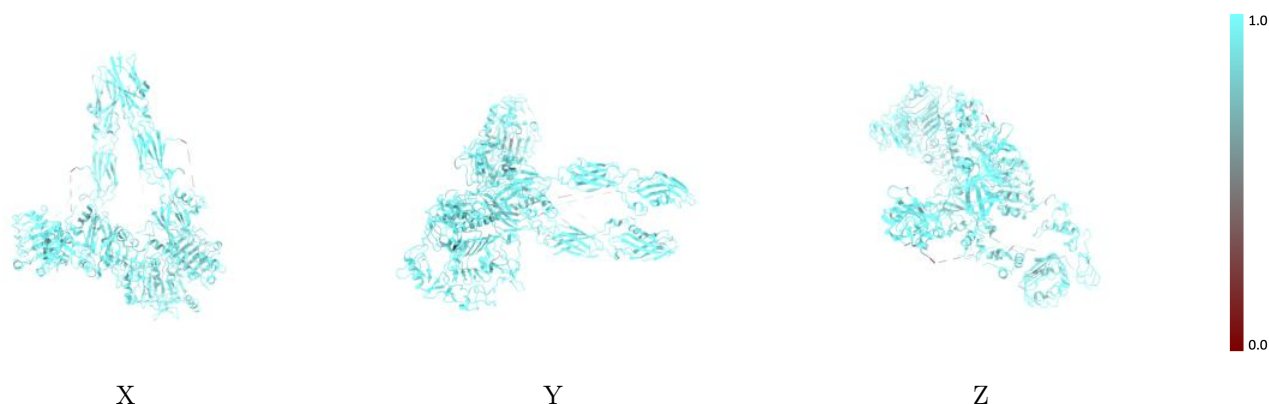
The images above show the 3D surface view of the map at the recommended contour level 0.018 at 50% transparency in yellow overlaid with a ribbon representation of the model coloured in blue. These images allow for the visual assessment of the quality of fit between the atomic model and the map.

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



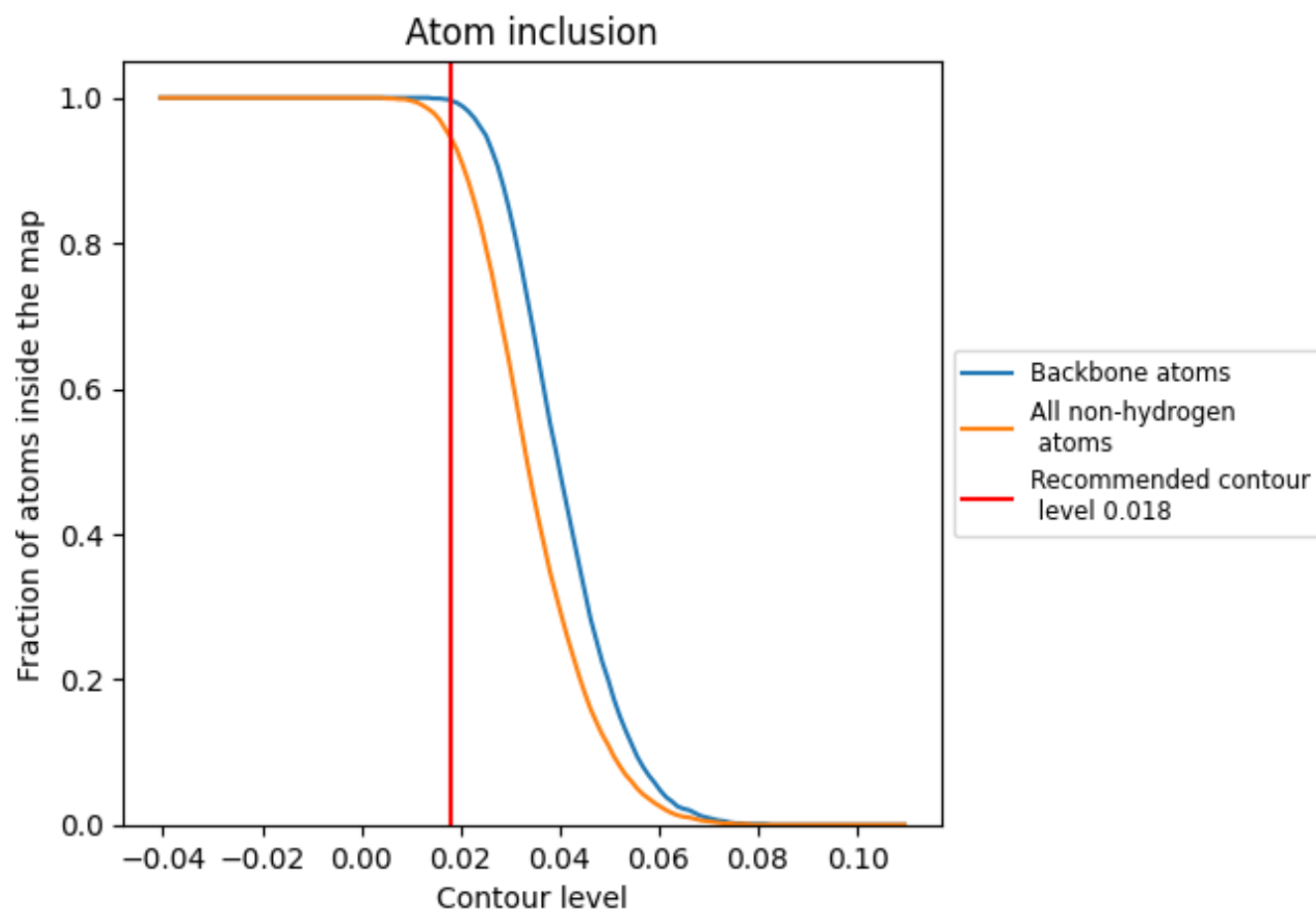
The images above show the model with each residue coloured according to its Q-score. This shows their resolvability in the map with higher Q-score values reflecting better resolvability. Please note: Q-score is calculating the resolvability of atoms, and thus high values are only expected at resolutions 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 (0.018).

9.4 Atom inclusion [i](#)



At the recommended contour level, 100% of all backbone atoms, 95% 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 (0.018) and Q-score for the entire model and for each chain.

Chain	Atom inclusion	Q-score
All	<div><div></div>0.9460</div>	<div><div></div>0.2200</div>
A	<div><div></div>0.9460</div>	<div><div></div>0.1990</div>
B	<div><div></div>0.9450</div>	<div><div></div>0.2390</div>
C	<div><div></div>0.9370</div>	<div><div></div>0.2060</div>
D	<div><div></div>0.9470</div>	<div><div></div>0.1800</div>
E	<div><div></div>0.9650</div>	<div><div></div>0.2850</div>
F	<div><div></div>0.9560</div>	<div><div></div>0.2300</div>

1.0

0.0

<0.0