



Full wwPDB EM Validation Report ⓘ

Nov 19, 2022 – 04:44 PM EST

PDB ID : 3J32
EMDB ID : EMD-5586
Title : An asymmetric unit map from electron cryo-microscopy of Haliotis diversicolor molluscan hemocyanin isoform 1 (HdH1)
Authors : Zhang, Q.; Dai, X.; Cong, Y.; Zhang, J.; Chen, D.-H.; Dougherty, M.; Wang, J.; Ludtke, S.; Schmid, M.F.; Chiu, W.
Deposited on : 2013-02-20
Resolution : 4.50 Å(reported)

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

We welcome your comments at validation@mail.wwpdb.org

A user guide is available at

<https://www.wwpdb.org/validation/2017/EMValidationReportHelp>
with specific help available everywhere you see the ⓘ symbol.

The types of validation reports are described at

<http://www.wwpdb.org/validation/2017/FAQs#types>.

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

EMDB validation analysis : 0.0.1.dev43
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) : 2.31.3

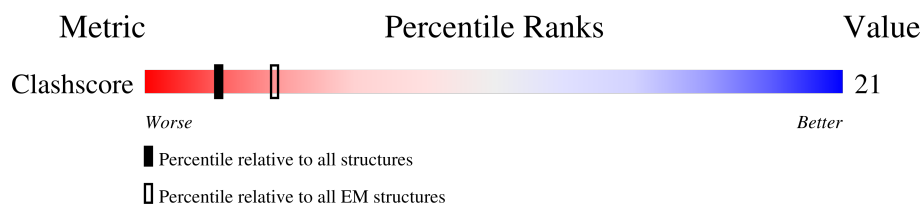
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

ELECTRON MICROSCOPY

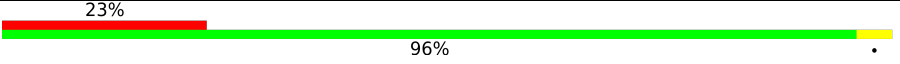
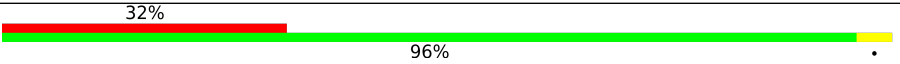
The reported resolution of this entry is 4.50 Å.

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

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	3303	
1	B	3303	

2 Entry composition

There is only 1 type of molecule in this entry. The entry contains 6606 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.

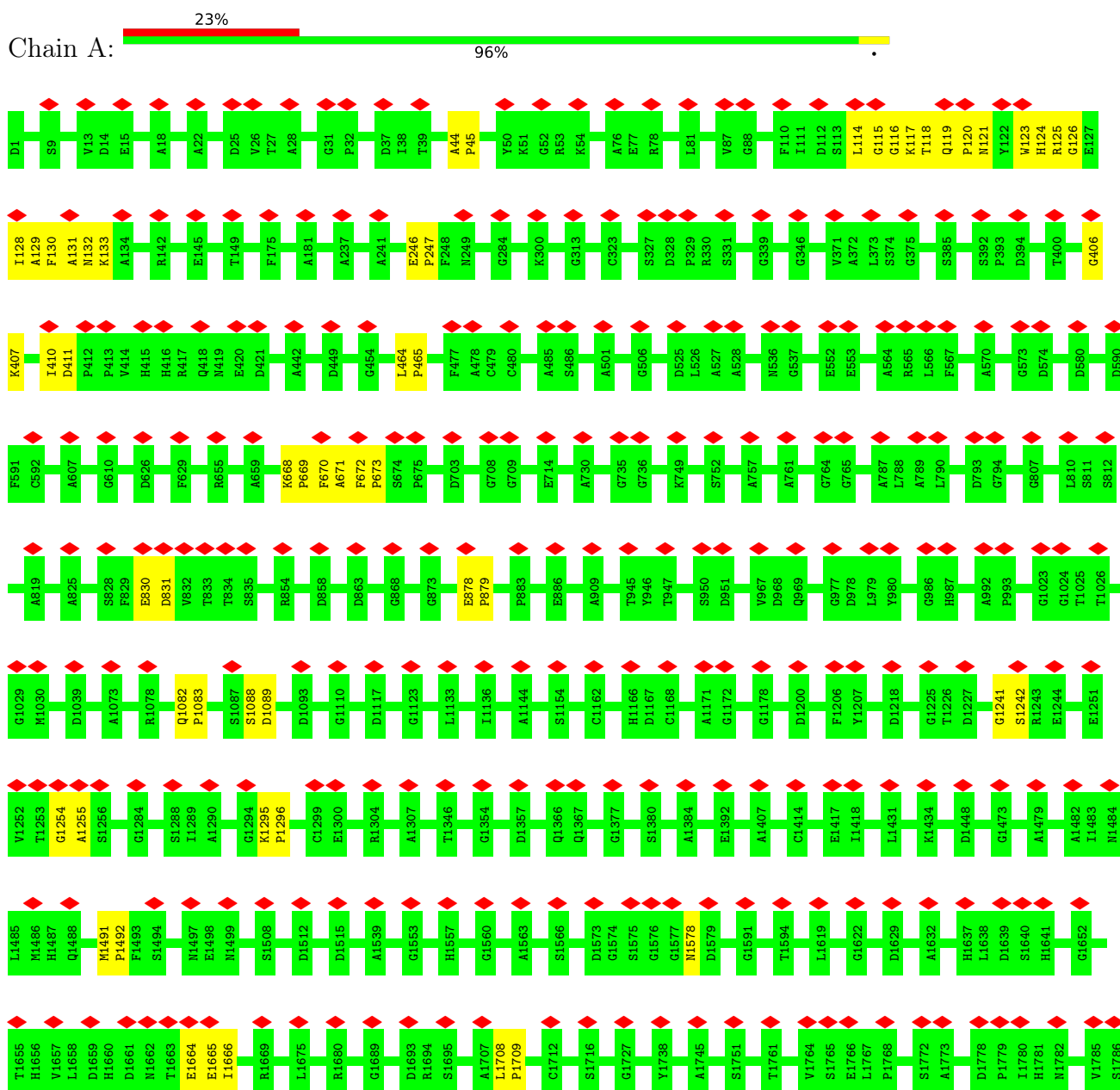
- Molecule 1 is a protein called Hemocyanin isoform 1.

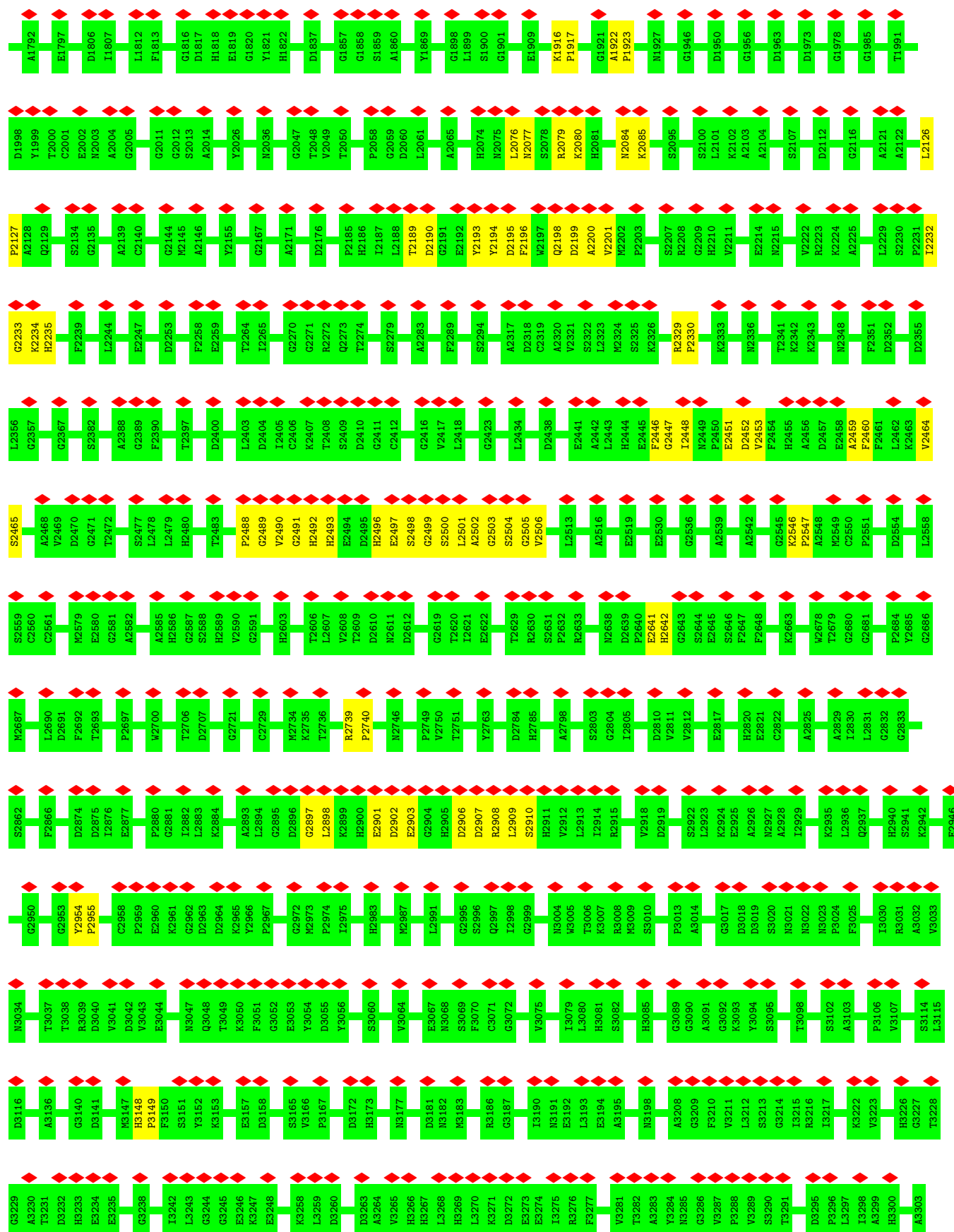
Mol	Chain	Residues	Atoms		AltConf	Trace
1	A	3303	Total 3303	C 3303	0	3303
1	B	3303	Total 3303	C 3303	0	3303

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.

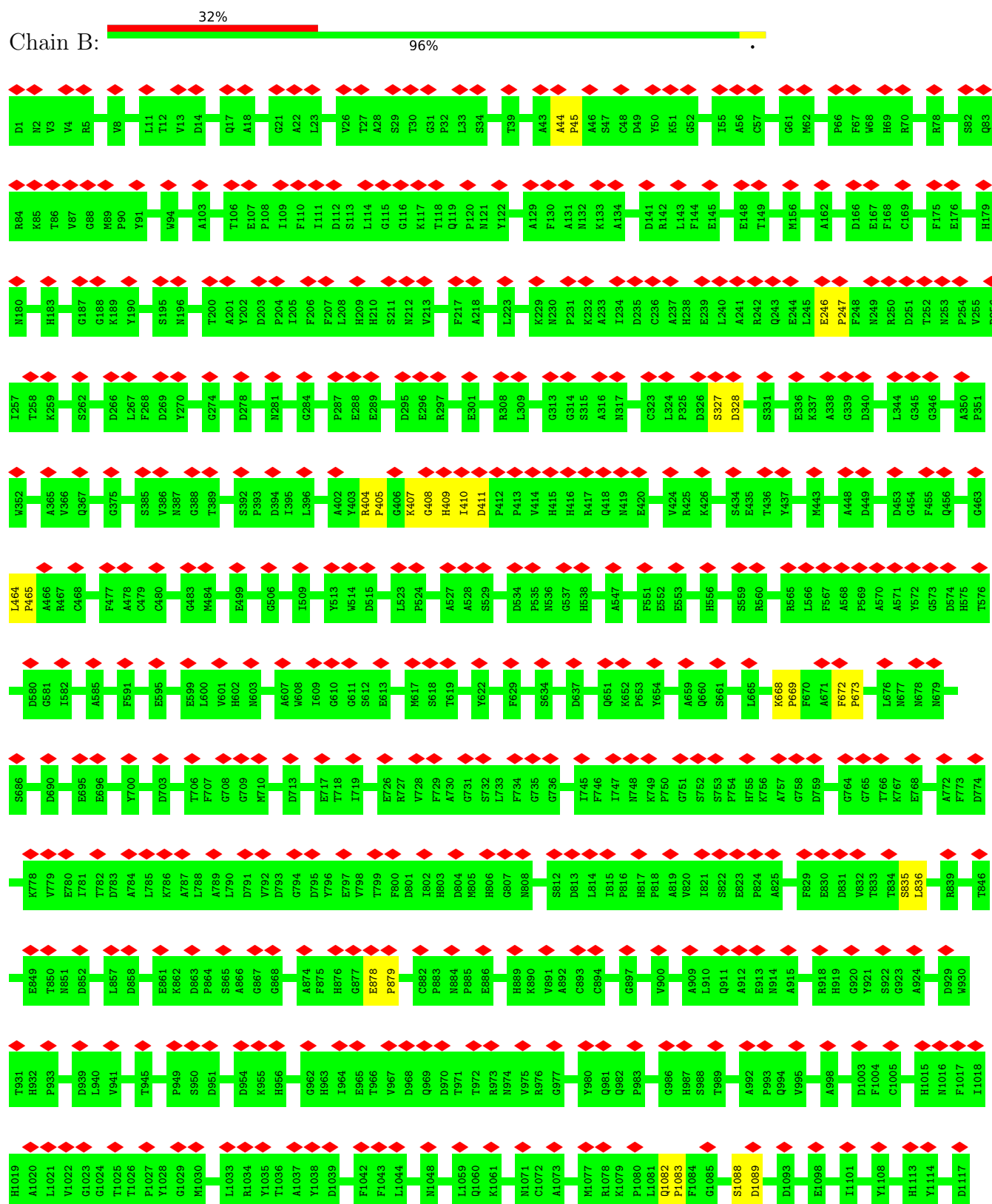
• Molecule 1: Hemocyanin isoform 1





• Molecule 1: Hemocyanin isoform 1

Chain B:



T1118	L1119	G1123	Q1128	L1133	F1143	A1144	G1145	F1146	L1147	L1148	K1153	S1154	V1155	V1161	C1162	A1163	P1164	P1165	H1166	D1167	A1171	G1172	E1173	L1177	G1178	D1179	E1180	M1181	E1182	M1183	A1184	Y1187	D1188	R1189	V1190	F1191	D1194	T1195	T1196	N1197	V1198	E1201	Y1207	F1217	G1221				
T1222	S1223	L1224	G1225	D1226	D1227	I1228	F1229	A1232	N1233	D1237	G1241	S1242	R1243	E1244	H1245	D1246	R1247	Y1248	E1251	V1252	T1253	G1254	A1255	S1256	D1264	E1269	D1283	A1290	G1294	K1295	P1296	G1297	E1300	L1301	N1302	Q1318	A1325	L1332	A1339	D1350	D1358	A1359	T1360						
Q1366	Q1367	R1368	Y1369	D1370	P1371	N1372	R1376	G1377	S1380	F1381	D1389	E1392	F1395	A1407	G1414	G1432	G1433	K1434	G1435	Q1436	Y1437	D1443	Y1444	S1445	F1451	H1454	M1457	E1478	I1483	M1486	H1487	Q1488	M1491	P1492	E1498	M1499	H1500	G1501	N1502	V1503									
K1509	G1521	D1525	S1533	T1534	P1535	S1536	K1542	A1543	G1553	G1560	C1570	G1574	D1579	C1580	S1581	H1582	T1583	A1584	G1591	G1592	E1593	G1621	G1622	L1623	F1624	E1627	A1632	S1636	S1640	G1652	I1653	D1654	D1659	H1660	T1661	N1662	T1663	E1664	E1665	T1666	L1667								
L1675	R1680	D1693	R1694	D1697	A1701	S1704	L1708	P1709	P1710	P1713	S1716	A1717	A1718	N1719	R1720	F1721	V1725	A1745	S1751	L1752	V1753	G1621	G1622	L1623	F1624	E1627	A1632	S1636	S1640	G1652	I1653	D1654	D1659	H1660	T1661	N1662	T1663	E1664	E1665	T1666	L1667								
G1816	D1817	H1818	E1819	G1820	Y1821	H1822	C1839	E1846	G1851	I1856	G1857	G1858	S1859	A1860	D1884	A1888	G1898	L1899	S1900	G1901	N1905	C1906	E1909	K1916	P1917	F1918	S1919	F1920	G1921	A1922	P1923	Y1924	S1928	S1934	K1935	D1938	G1946	D1950	L1954	E1955	G1956	D1973							
A1977	G1983	A1988	D1997	C2001	G2011	G2012	S2013	A2014	A2019	H2039	E2040	D2041	T2048	V2049	T2050	A2051	Y2052	D2053	G2054	T2055	D2060	A2065	H2074	N2075	L2076	N2077	S2078	R2079	N2084	K2085	L2090	T2091	S2092	A2099	S2100	A2104	G2116	A2119	I2120	A2121	A2122								
G2125	L2126	P2127	A2128	Q2129	D2132	T2133	S2134	G2135	A2139	G2144	Y2155	A2162	G2167	A2171	G2191	E2192	Y2193	Y2194	D2195	F2196	W2197	Q2198	D2199	A2200	V2201	N2202	P2203	S2207	R2208	N2221	V2222	K2224	A2225	L2226	P2231	I2232	G2233	K2234	H2235	D2240	Q2241	A2242	E2247						
C2252	G2271	A2317	A2320	R2329	P2330	A2345	D2352	N2369	E2374	A2378	S2382	G2389	D2400	K2407	T2408	S2409	D2410	G2411	H2414	A2415	G2416	G2423	D2438	D2452	A2456	D2457	E2458	A2459	F2460	F2461	L2462	A2468	L2474	S2477	T2483	G2489													
D2495	H2496	E2497	S2498	G2499	S2500	L2501	A2502	G2503	S2504	V2506	R2507	L2513	T2514	T2515	A2524	G2527	H2532	G2536	A2542	G2545	K2546	P2547	D2548	M2549	C2550	P2551	M2552	A2553	D2554	L2558	S2559	C2560	G2564	M2565	A2566	M2579	E2580	G2581	G2587	G2591	L2592	P2593	A2599						
S2602	T2609	D2610	N2611	D2612	N2613	N2614	P2615	D2618	V2625	T2626	T2627	T2628	S2631	P2632	R2633	F2637	N2638	D2639	P2640	E2641	G2642	G2643	S2644	E2645	S2646	V2652	E2657	E2665	E2669	N2673	V2678	T2679	G2680	G2681	G2686	N2687	A2694	P2697	L2698	F2699	N2700	N2705	A2711						
Y2719	R2720	G2721	A2727	N2728	C2729	T2736	R2739	P2740	D2743	N2744	V2750	A2753	D2759	R2767	D2794	A2799	F2800	L2801	L2802	S2803	G2804	S2808	V2811	I2815	C2816	N2819	C2822	A2825	G2826	T2827	F2828	A2829	G2832	G2833	S2840	D2848	Q2854												
S2865	F2866	K2867	D2874	F2880	G2881	I2882	L2883	T2887	L2894	G2895	D2896	G2897	L2898	K2899	H2900	E2901	D2902	E2903	G2904	H2905	D2906	D2907	L2908	L2909	S2910	H2911	V2912	L2913	I2914	E2917	V2918	S2922	E2925	A2926	I2929	K2930	T2931	A2932	L2933	L2936	Q2937	N2938	D2939	K2942	G2943	G2944	F2945	E2946	E2947



4 Experimental information

Property	Value	Source
EM reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, D5	Depositor
Number of particles used	28641	Depositor
Resolution determination method	FSC 0.5 CUT-OFF	Depositor
CTF correction method	each micrograph	Depositor
Microscope	JEOL 3200FSC	Depositor
Voltage (kV)	300	Depositor
Electron dose ($e^-/\text{\AA}^2$)	20	Depositor
Minimum defocus (nm)	500	Depositor
Maximum defocus (nm)	3600	Depositor
Magnification	60000	Depositor
Image detector	KODAK SO-163 FILM	Depositor
Maximum map value	11.359	Depositor
Minimum map value	-0.071	Depositor
Average map value	0.006	Depositor
Map value standard deviation	0.111	Depositor
Recommended contour level	1.5	Depositor
Map size (\AA)	612.0, 612.0, 612.0	wwPDB
Map dimensions	600, 600, 600	wwPDB
Map angles ($^\circ$)	90.0, 90.0, 90.0	wwPDB
Pixel spacing (\AA)	1.02, 1.02, 1.02	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).

There are no protein, RNA or DNA chains available to summarize Z scores of covalent bonds and angles.

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	3303	0	0	77	0
1	B	3303	0	0	67	0
All	All	6606	0	0	139	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.

All (139) 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:A:2641:GLU:CA	1:B:1536:SER:CA	2.39	0.99
1:A:2642:HIS:CA	1:B:1533:SER:CA	2.47	0.92
1:B:2075:ASN:CA	1:B:2076:LEU:CA	2.68	0.72
1:A:2641:GLU:CA	1:B:1535:PRO:CA	2.69	0.71
1:A:116:GLY:CA	1:A:117:LYS:CA	2.70	0.70
1:A:2489:GLY:CA	1:A:2490:VAL:CA	2.69	0.70
1:A:128:ILE:CA	1:A:129:ALA:CA	2.70	0.70

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:A:2198:GLN:CA	1:A:2199:ASP:CA	2.71	0.68
1:A:2232:ILE:CA	1:A:2233:GLY:CA	2.71	0.68
1:B:1241:GLY:CA	1:B:1242:SER:CA	2.72	0.68
1:B:1580:CYS:CA	1:B:1581:SER:CA	2.73	0.67
1:A:2234:LYS:CA	1:A:2235:HIS:CA	2.72	0.67
1:A:2195:ASP:CA	1:A:2196:PHE:CA	2.72	0.67
1:A:2499:GLY:CA	1:A:2500:SER:CA	2.74	0.66
1:A:2505:GLY:CA	1:A:2506:VAL:CA	2.74	0.66
1:A:132:ASN:CA	1:A:133:LYS:CA	2.74	0.65
1:A:2084:ASN:CA	1:A:2085:LYS:CA	2.75	0.65
1:A:1254:GLY:CA	1:A:1255:ALA:CA	2.75	0.65
1:A:2504:SER:CA	1:A:2505:GLY:CA	2.75	0.65
1:A:2451:GLU:CA	1:A:2452:ASP:CA	2.76	0.64
1:A:668:LYS:CA	1:A:669:PRO:CA	2.75	0.64
1:B:672:PHE:CA	1:B:673:PRO:CA	2.76	0.64
1:A:44:ALA:CA	1:A:45:PRO:CA	2.76	0.64
1:A:1241:GLY:CA	1:A:1242:SER:CA	2.76	0.63
1:A:2502:ALA:CA	1:A:2503:GLY:CA	2.76	0.63
1:B:3148:HIS:CA	1:B:3149:PRO:CA	2.77	0.63
1:A:1088:SER:CA	1:A:1089:ASP:CA	2.77	0.63
1:A:2447:GLY:CA	1:A:2448:ILE:CA	2.76	0.63
1:B:2546:LYS:CA	1:B:2547:PRO:CA	2.77	0.63
1:B:2894:LEU:CA	1:B:2895:GLY:CA	2.77	0.63
1:A:2546:LYS:CA	1:A:2547:PRO:CA	2.77	0.63
1:B:1778:ASP:CA	1:B:1779:PRO:CA	2.77	0.63
1:B:2126:LEU:CA	1:B:2127:PRO:CA	2.77	0.63
1:A:2200:ALA:CA	1:A:2201:VAL:CA	2.77	0.63
1:B:1782:ASN:CA	1:B:1783:VAL:CA	2.77	0.63
1:A:1664:GLU:CA	1:A:1665:GLU:CA	2.77	0.62
1:B:1819:GLU:CA	1:B:1820:GLY:CA	2.77	0.62
1:B:246:GLU:CA	1:B:247:PRO:CA	2.77	0.62
1:B:1922:ALA:CA	1:B:1923:PRO:CA	2.77	0.62
1:B:2231:PRO:CA	1:B:2232:ILE:CA	2.77	0.62
1:A:1922:ALA:CA	1:A:1923:PRO:CA	2.77	0.62
1:B:1708:LEU:CA	1:B:1709:PRO:CA	2.77	0.62
1:B:2233:GLY:CA	1:B:2234:LYS:CA	2.77	0.62
1:A:1295:LYS:CA	1:A:1296:PRO:CA	2.77	0.62
1:B:44:ALA:CA	1:B:45:PRO:CA	2.77	0.62
1:B:464:LEU:CA	1:B:465:PRO:CA	2.77	0.62
1:B:1082:GLN:CA	1:B:1083:PRO:CA	2.78	0.62
1:B:2234:LYS:CA	1:B:2235:HIS:CA	2.77	0.62

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1295:LYS:CA	1:B:1296:PRO:CA	2.77	0.62
1:A:1665:GLU:CA	1:A:1666:ILE:CA	2.77	0.62
1:A:1708:LEU:CA	1:A:1709:PRO:CA	2.77	0.62
1:A:1916:LYS:CA	1:A:1917:PRO:CA	2.78	0.62
1:B:327:SER:CA	1:B:328:ASP:CA	2.78	0.62
1:B:1253:THR:CA	1:B:1254:GLY:CA	2.77	0.62
1:B:2461:PHE:CA	1:B:2462:LEU:CA	2.78	0.62
1:B:2954:TYR:CA	1:B:2955:PRO:CA	2.77	0.62
1:A:2739:ARG:CA	1:A:2740:PRO:CA	2.78	0.62
1:B:1916:LYS:CA	1:B:1917:PRO:CA	2.78	0.62
1:A:878:GLU:CA	1:A:879:PRO:CA	2.77	0.62
1:A:2954:TYR:CA	1:A:2955:PRO:CA	2.77	0.62
1:B:878:GLU:CA	1:B:879:PRO:CA	2.77	0.62
1:B:1491:MET:CA	1:B:1492:PRO:CA	2.78	0.62
1:B:2202:MET:CA	1:B:2203:PRO:CA	2.77	0.62
1:A:1491:MET:CA	1:A:1492:PRO:CA	2.78	0.62
1:B:1817:ASP:CA	1:B:1818:HIS:CA	2.77	0.62
1:B:2200:ALA:CA	1:B:2201:VAL:CA	2.77	0.62
1:A:246:GLU:CA	1:A:247:PRO:CA	2.78	0.61
1:A:830:GLU:CA	1:A:831:ASP:CA	2.77	0.61
1:A:672:PHE:CA	1:A:673:PRO:CA	2.78	0.61
1:B:1088:SER:CA	1:B:1089:ASP:CA	2.77	0.61
1:B:2198:GLN:CA	1:B:2199:ASP:CA	2.78	0.61
1:B:2329:ARG:CA	1:B:2330:PRO:CA	2.78	0.61
1:A:2488:PRO:CA	1:A:2489:GLY:CA	2.78	0.61
1:A:3148:HIS:CA	1:A:3149:PRO:CA	2.78	0.61
1:A:118:THR:CA	1:A:119:GLN:CA	2.78	0.61
1:A:464:LEU:CA	1:A:465:PRO:CA	2.78	0.61
1:A:1082:GLN:CA	1:A:1083:PRO:CA	2.77	0.61
1:B:2739:ARG:CA	1:B:2740:PRO:CA	2.78	0.61
1:A:2079:ARG:CA	1:A:2080:LYS:CA	2.79	0.61
1:B:668:LYS:CA	1:B:669:PRO:CA	2.78	0.61
1:B:2084:ASN:CA	1:B:2085:LYS:CA	2.77	0.61
1:B:2502:ALA:CA	1:B:2503:GLY:CA	2.78	0.61
1:A:2459:ALA:CA	1:A:2460:PHE:CA	2.79	0.61
1:A:2906:ASP:CA	1:A:2907:ASP:CA	2.80	0.60
1:B:835:SER:CA	1:B:836:LEU:CA	2.79	0.60
1:B:2221:ASN:CA	1:B:2222:VAL:CA	2.79	0.60
1:A:2897:GLY:CA	1:A:2898:LEU:CA	2.80	0.60
1:A:2076:LEU:CA	1:A:2077:ASN:CA	2.80	0.60
1:B:2901:GLU:CA	1:B:2902:ASP:CA	2.80	0.60

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1773:ALA:CA	1:B:1774:GLU:CA	2.80	0.60
1:A:2452:ASP:CA	1:A:2453:VAL:CA	2.80	0.60
1:A:2464:VAL:CA	1:A:2465:SER:CA	2.80	0.59
1:A:2497:GLU:CA	1:A:2498:SER:CA	2.80	0.59
1:A:2496:HIS:CA	1:A:2497:GLU:CA	2.80	0.59
1:A:669:PRO:CA	1:A:670:PHE:CA	2.80	0.59
1:A:120:PRO:CA	1:A:121:ASN:CA	2.81	0.58
1:B:3040:ASP:CA	1:B:3041:VAL:CA	2.81	0.58
1:A:410:ILE:CA	1:A:411:ASP:CA	2.81	0.58
1:A:2492:HIS:CA	1:A:2493:HIS:CA	2.80	0.58
1:A:2193:TYR:CA	1:A:2194:TYR:CA	2.82	0.58
1:B:3042:ASP:CA	1:B:3043:VAL:CA	2.82	0.57
1:B:409:HIS:CA	1:B:410:ILE:CA	2.82	0.57
1:B:2207:SER:CA	1:B:2208:ARG:CA	2.82	0.57
1:B:2641:GLU:CA	1:B:2642:HIS:CA	2.82	0.57
1:B:3045:LEU:CA	1:B:3046:PHE:CA	2.83	0.57
1:B:1581:SER:CA	1:B:1582:HIS:CA	2.83	0.57
1:A:130:PHE:CA	1:A:131:ALA:CA	2.83	0.57
1:B:407:LYS:CA	1:B:408:GLY:CA	2.82	0.57
1:A:131:ALA:CA	1:A:132:ASN:CA	2.83	0.57
1:B:410:ILE:CA	1:B:411:ASP:CA	2.82	0.57
1:B:1371:PRO:CA	1:B:1372:ASN:CA	2.83	0.56
1:B:2644:SER:CA	1:B:2645:GLU:CA	2.84	0.56
1:A:406:GLY:CA	1:A:407:LYS:CA	2.84	0.56
1:A:2126:LEU:CA	1:A:2127:PRO:CA	2.84	0.55
1:A:670:PHE:CA	1:A:671:ALA:CA	2.85	0.54
1:B:2191:GLY:CA	1:B:2192:GLU:CA	2.85	0.54
1:A:2901:GLU:CA	1:A:2902:ASP:CA	2.86	0.54
1:B:1813:PHE:CA	1:B:1814:HIS:CA	2.85	0.54
1:A:2446:PHE:CA	1:A:2447:GLY:CA	2.86	0.54
1:A:2909:LEU:CA	1:A:2910:SER:CA	2.85	0.54
1:B:404:ARG:CA	1:B:405:PRO:CA	2.86	0.54
1:A:114:LEU:CA	1:A:115:GLY:CA	2.85	0.53
1:B:2193:TYR:CA	1:B:2194:TYR:CA	2.87	0.53
1:A:2902:ASP:CA	1:A:2903:GLU:CA	2.85	0.53
1:A:2498:SER:CA	1:A:2499:GLY:CA	2.86	0.53
1:B:2913:LEU:CA	1:B:2914:ILE:CA	2.85	0.53
1:A:2500:SER:CA	1:A:2501:LEU:CA	2.87	0.52
1:B:3043:VAL:CA	1:B:3044:GLU:CA	2.88	0.52
1:A:2907:ASP:CA	1:A:2908:ARG:CA	2.88	0.51
1:A:2491:GLY:CA	1:A:2492:HIS:CA	2.88	0.51

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1665:GLU:CA	1:B:1666:ILE:CA	2.89	0.50
1:B:3039:ARG:CA	1:B:3040:ASP:CA	2.90	0.49
1:A:124:HIS:CA	1:A:125:ARG:CA	2.90	0.49
1:B:3041:VAL:CA	1:B:3042:ASP:CA	2.91	0.48
1:A:123:TRP:CA	1:A:126:GLY:CA	2.95	0.45
1:A:1578:ASN:CA	1:B:2719:TYR:CA	2.95	0.45
1:A:2189:THR:CA	1:A:2190:ASP:CA	2.95	0.45
1:A:1578:ASN:CA	1:B:2720:ARG:CA	2.96	0.44
1:A:2329:ARG:CA	1:A:2330:PRO:CA	2.97	0.43

There are no symmetry-related clashes.

5.3 Torsion angles [i](#)

5.3.1 Protein backbone [i](#)

There are no protein backbone outliers to report in this entry.

5.3.2 Protein sidechains [i](#)

There are no protein residues with a non-rotameric sidechain to report in this entry.

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](#)

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.

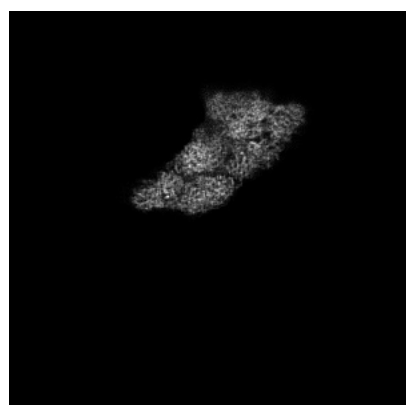
6 Map visualisation [i](#)

This section contains visualisations of the EMDB entry EMD-5586. These allow visual inspection of the internal detail of the map and identification of artifacts.

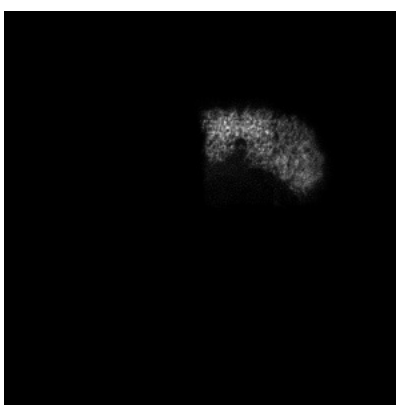
No raw map or half-maps were deposited for this entry and therefore no images, graphs, etc. pertaining to the raw map can be shown.

6.1 Orthogonal projections [i](#)

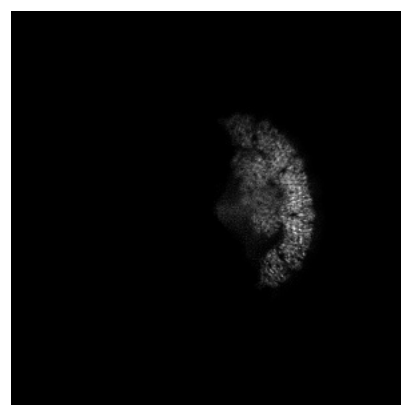
6.1.1 Primary map



X



Y

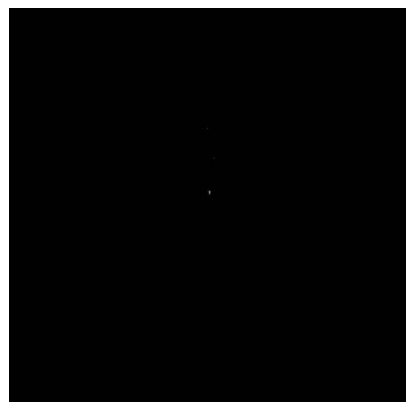


Z

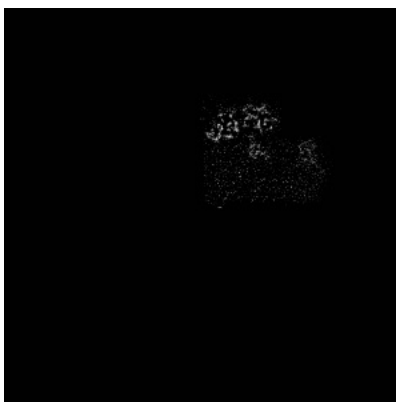
The images above show the map projected in three orthogonal directions.

6.2 Central slices [i](#)

6.2.1 Primary map



X Index: 300



Y Index: 300



Z Index: 300

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: 425



Y Index: 284



Z Index: 331

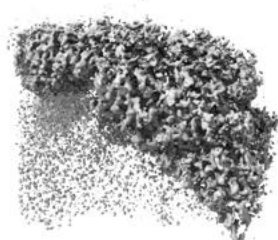
The images above show the largest variance slices of the map in three orthogonal directions.

6.4 Orthogonal surface views [i](#)

6.4.1 Primary map



X



Y



Z

The images above show the 3D surface view of the map at the recommended contour level 1.5. These images, in conjunction with the slice images, may facilitate assessment of whether an appropriate contour level has been provided.

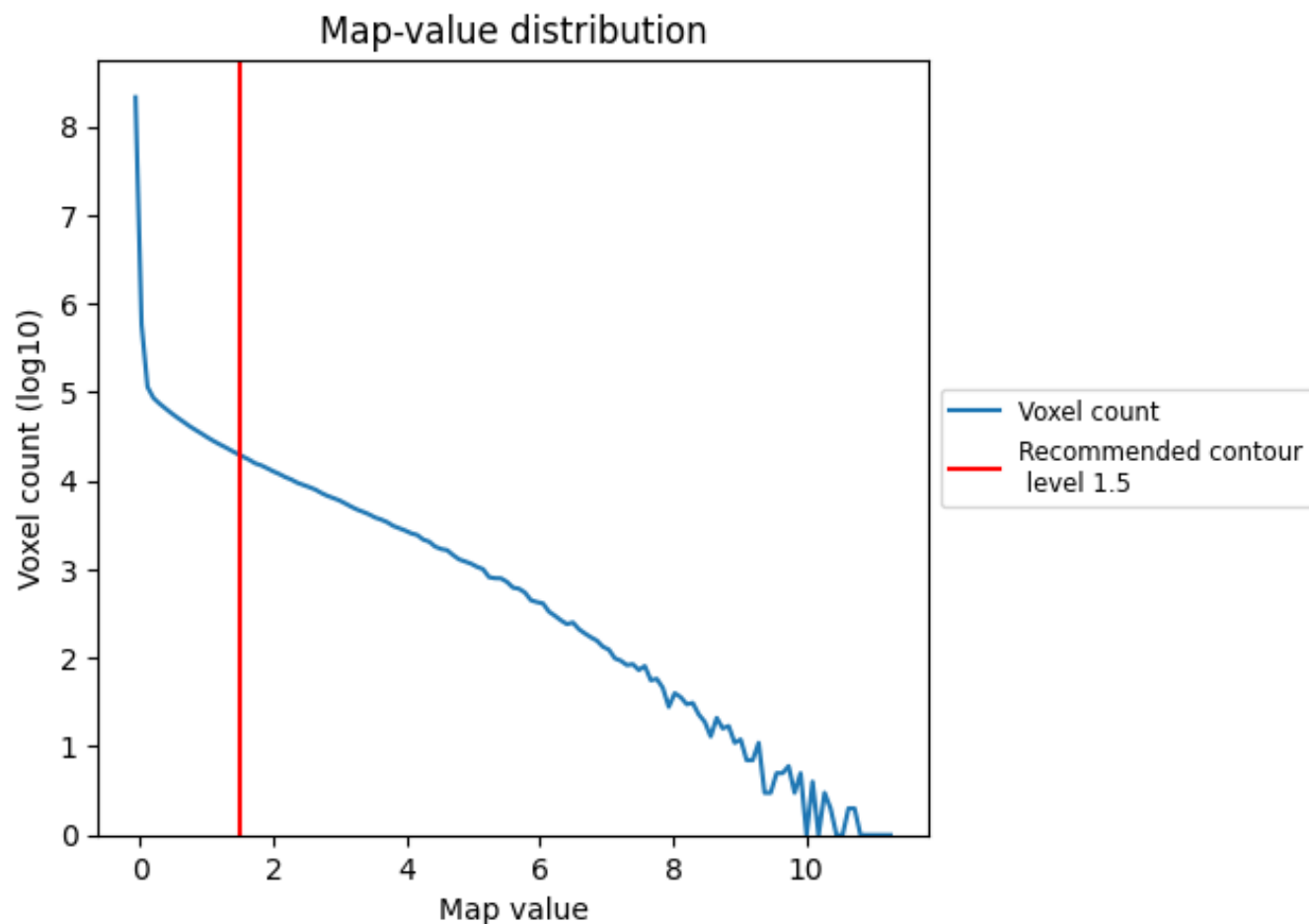
6.5 Mask visualisation

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

7 Map analysis [i](#)

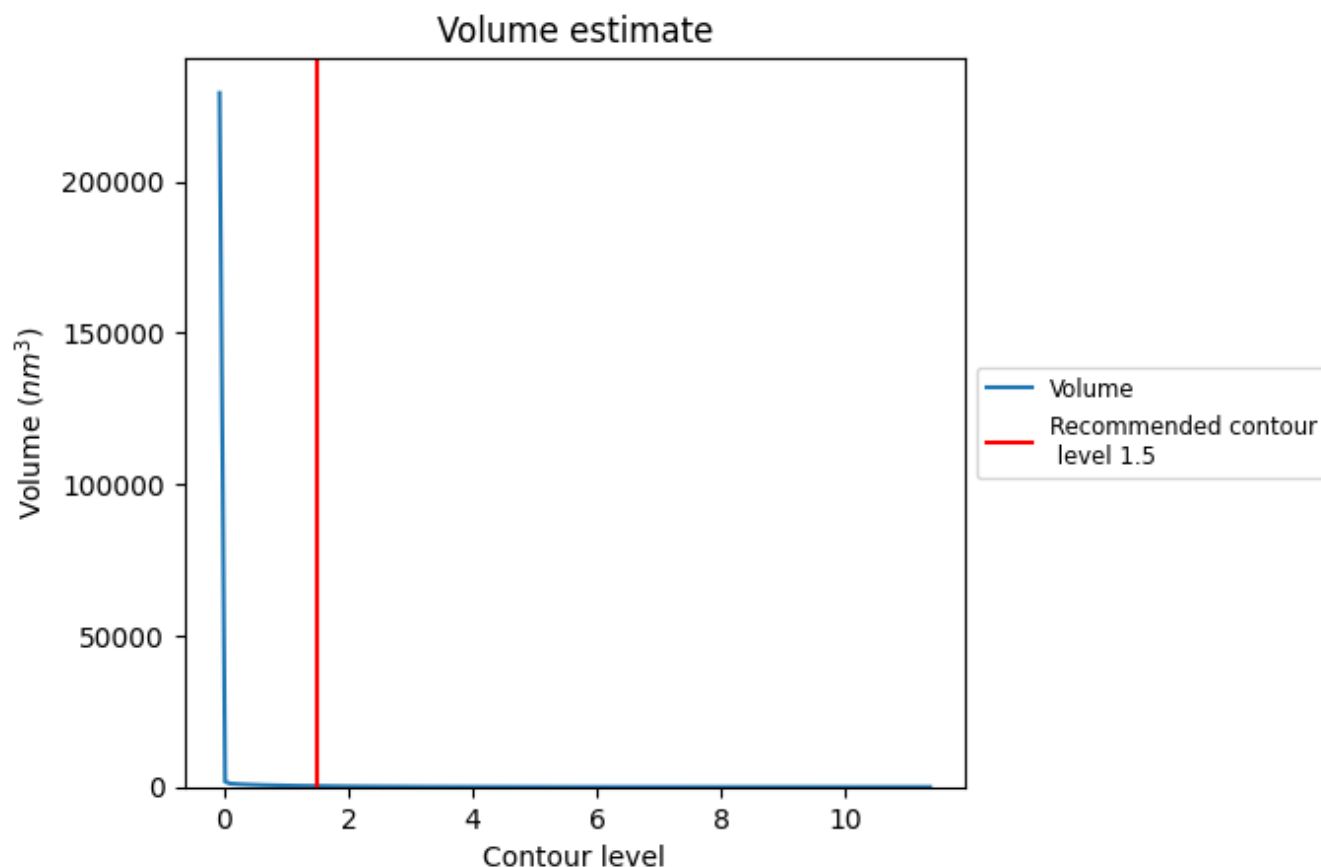
This section contains the results of statistical analysis of the map.

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.

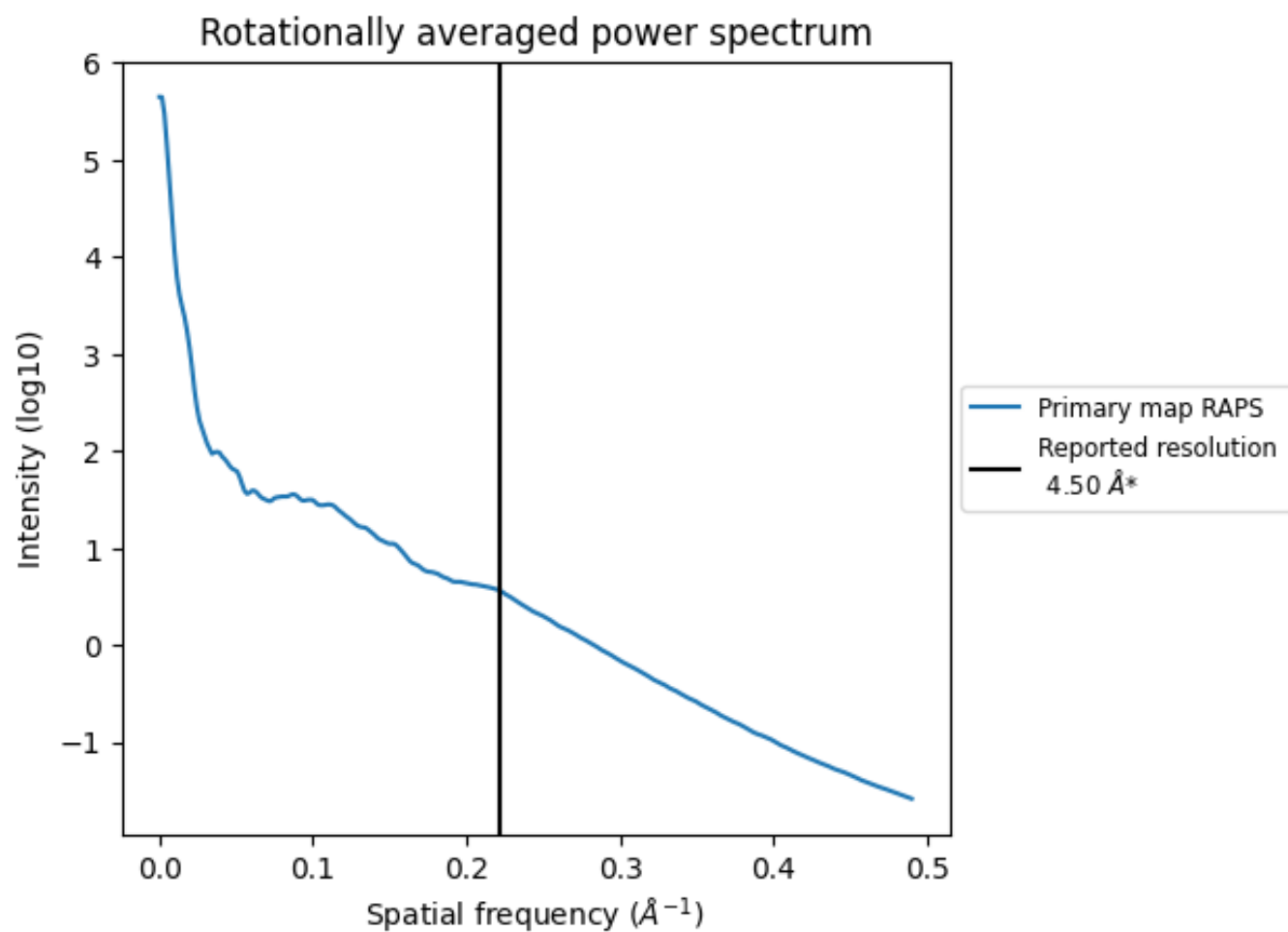
7.2 Volume estimate [i](#)



The volume at the recommended contour level is 290 nm³; this corresponds to an approximate mass of 262 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 0.222 \AA^{-1}

8 Fourier-Shell correlation

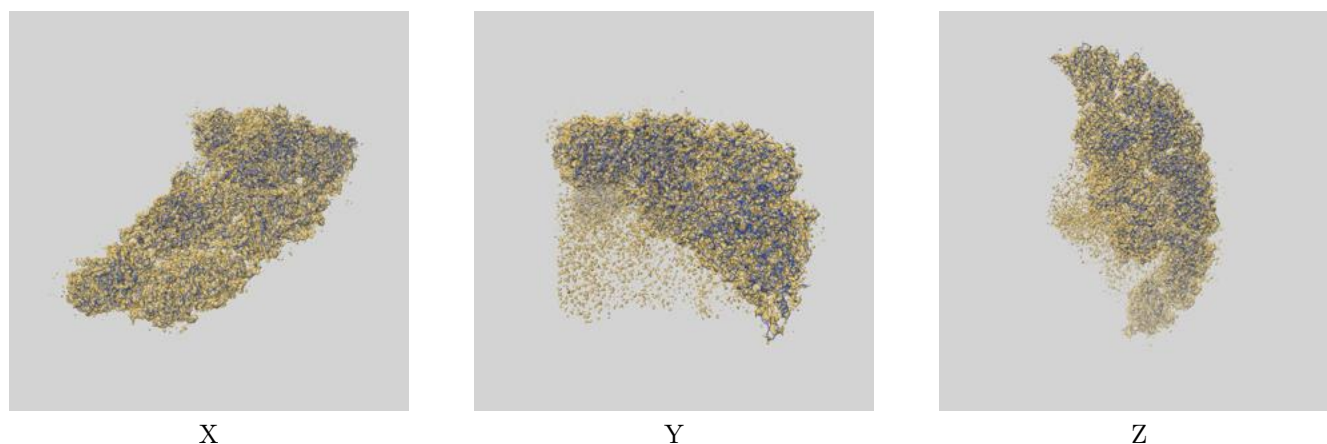
This section was not generated. No FSC curve or half-maps provided.

9 Map-model fit [i](#)

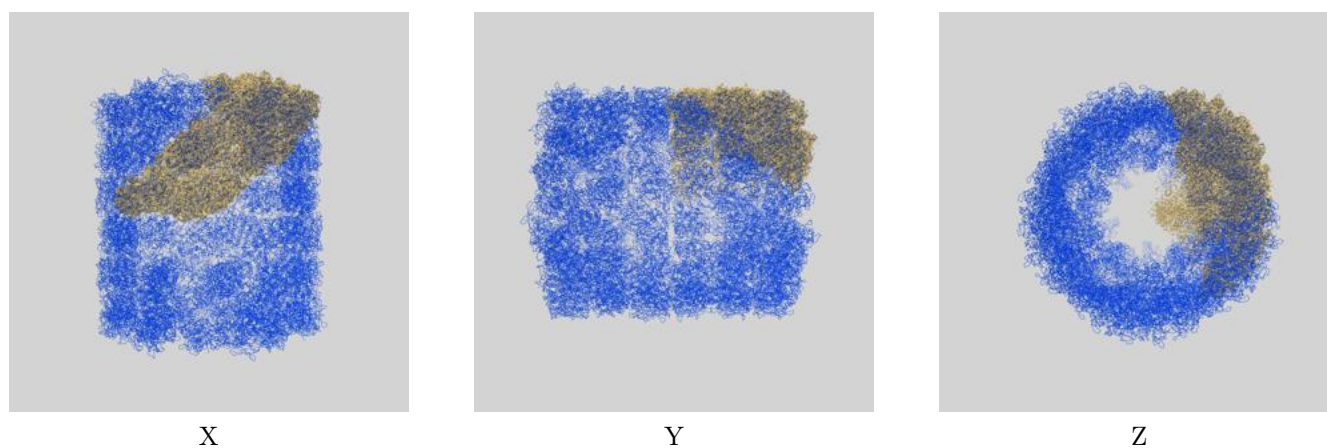
This section contains information regarding the fit between EMDB map EMD-5586 and PDB model 3J32. Per-residue inclusion information can be found in section 3 on page 4.

9.1 Map-model overlays

9.1.1 Map-model overlay [i](#)

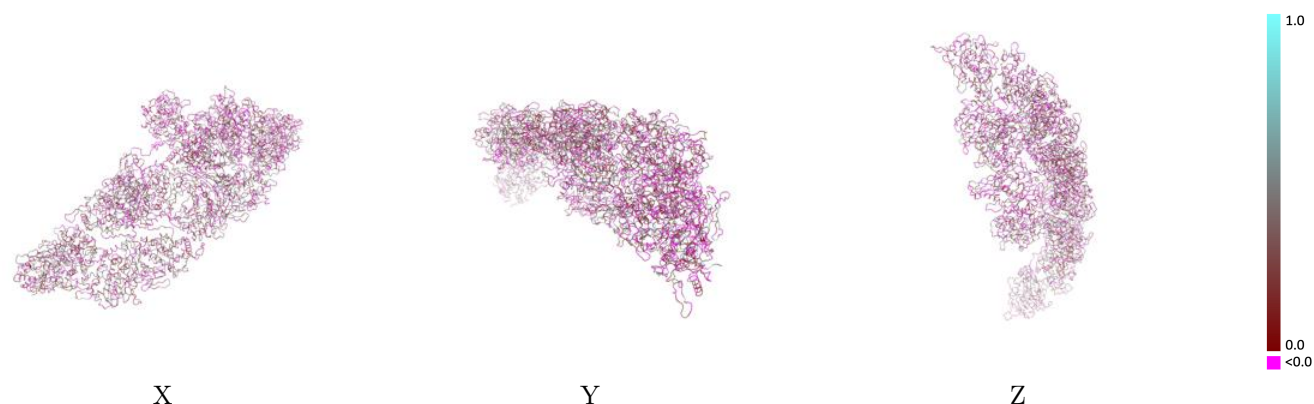


9.1.2 Map-model assembly overlay [i](#)



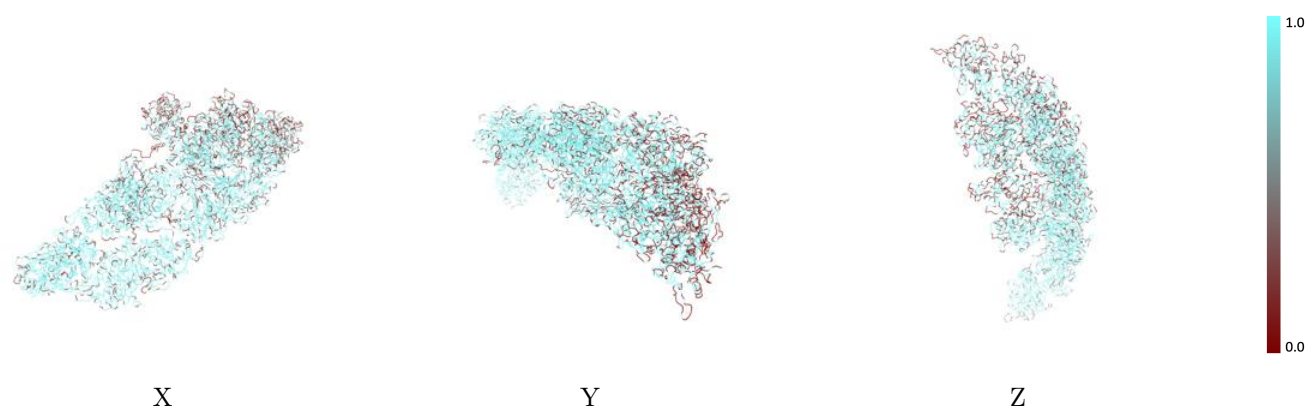
The images above show the 3D surface view of the map at the recommended contour level 1.5 at 50% transparency in yellow overlaid with a ribbon representation of the model coloured in blue. These images allow for the visual assessment of the quality of fit between the atomic model and the map.

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



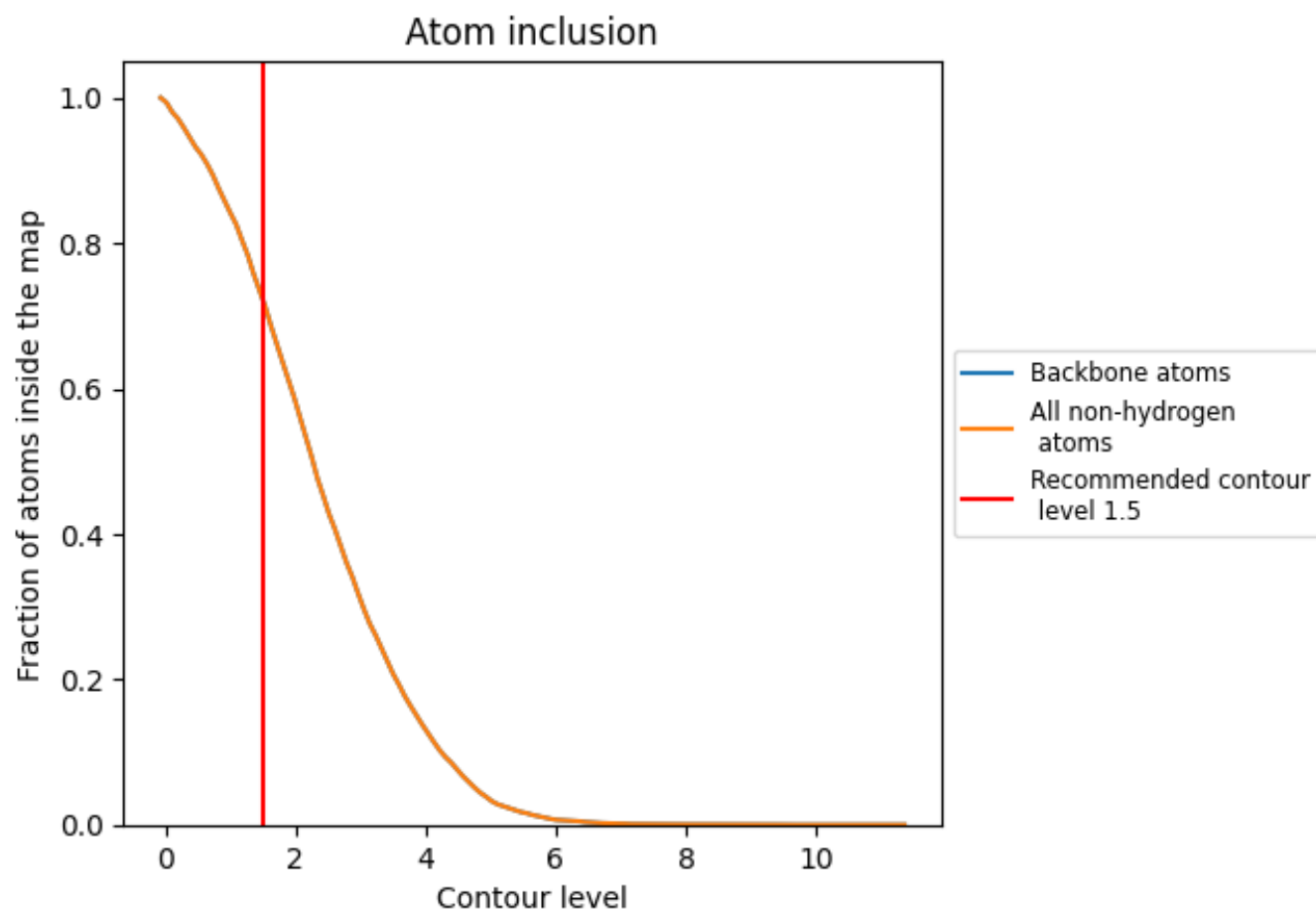
The images above show the model with each residue coloured according its Q-score. This shows their resolvability in the map with higher Q-score values reflecting better resolvability. Please note: Q-score is calculating the resolvability of atoms, and thus high values are only expected at resolutions 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 (1.5).

9.4 Atom inclusion [i](#)



At the recommended contour level, 72% of all backbone atoms, 72% 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 (1.5) and Q-score for the entire model and for each chain.

Chain	Atom inclusion	Q-score
All	<div></div> 0.7227	<div></div> 0.2030
A	<div></div> 0.7693	<div></div> 0.2160
B	<div></div> 0.6761	<div></div> 0.1890

