



## Full wwPDB EM Validation Report ⓘ

Nov 27, 2022 – 08:29 PM EST

PDB ID : 7KVE  
EMDB ID : EMD-23048  
Title : Cryo-EM structure of human Factor V at 3.3 Angstrom resolution  
Authors : Ruben, E.A.; Di Cera, E.  
Deposited on : 2020-11-27  
Resolution : 3.30 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](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.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) : 2.31.2

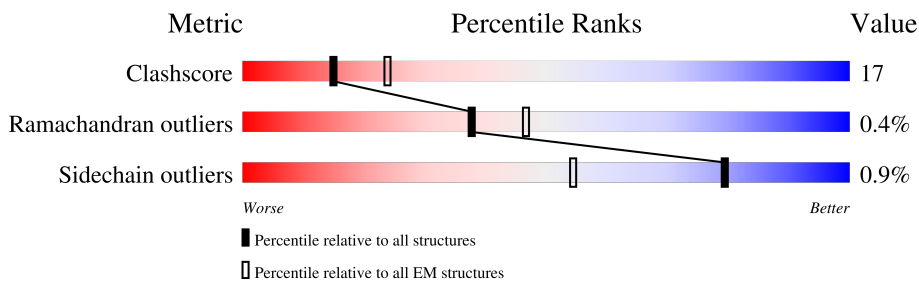
# 1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

*ELECTRON MICROSCOPY*

The reported resolution of this entry is 3.30 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	B	2196	

## 2 Entry composition

There is only 1 type of molecule in this entry. The entry contains 11142 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.

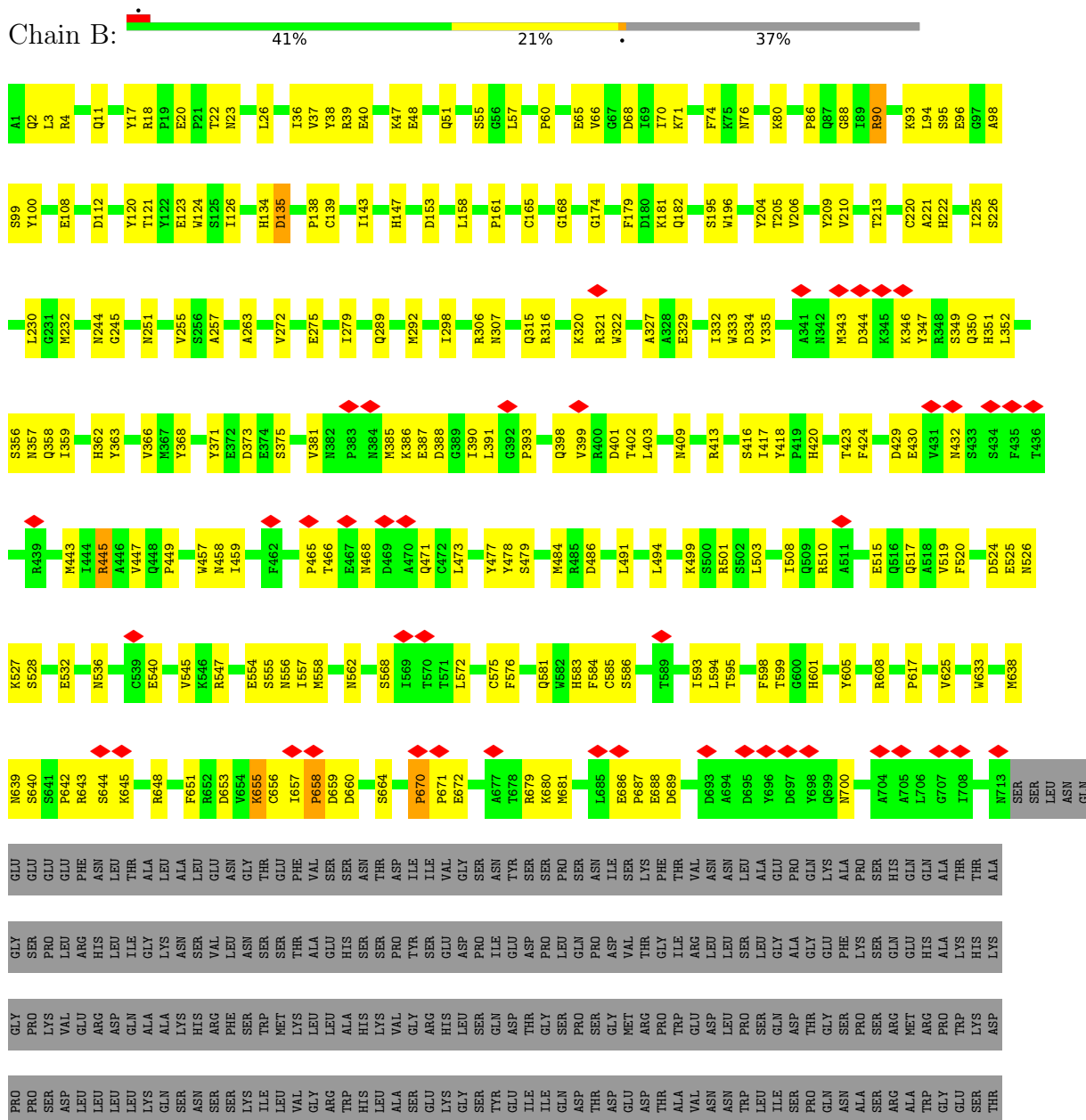
- Molecule 1 is a protein called Coagulation factor V.

Mol	Chain	Residues	Atoms					AltConf	Trace
			Total	C	N	O	S		
1	B	1374	11142	7074	1905	2108	55	0	0

### 3 Residue-property plots [i](#)

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: Coagulation factor V



P2070	P2071	R2074	A2077	N2082	A2083	W2084	Q2085	A2086	K2087	A2088	N2089	Q2093	L2095	E2096	L2099	I2102	K2103	T2110	Q2111	G2112	C2113	K2114	E2119	M2120	Y2121	K2122	S2124	S2130	E2131	Q2132	C2133	W2134	E2135	R2140	L2141	K2142	S2143	S2144	M2145	K2148	M2153	T2156	K2157	P1851	G1867	T1870	G1981	M1982	S1983	T1984	R1985	M1986	G1987	R1988	L1885	S1886	T1887	G1888	I1889	Q1894	A1897	S1898	E1899	F1900	L1901	G1902	Y1903	W1904	R1910	A1919	W1920	S1921	V1922	E1923	K1924	L1925	A1926	A1927	E1928	F1929	P1933	W1934	I1935	Q1936	V1937	D1938	M1939	Q1949	K1958	S1959	C1960	Y1961	T1962	Y1756	E1757	R1761	F1777	M1781	I1784	Y1785	S1786	L1787	M1792	Y1793	E1794	Q1795	W1797	V1798	R1799	L1800	H1801	L1802	L1803	M1804	I1805	G1806	G1807	I1811	H1812	W1813	V1814	L1822	E1823	M1824	K1827	Q1828	H1829	Q1830	L1831	G1832	V1833	L1836	L1837	P1838	D1839	G1839	S1840	F1841	K1842	K1847	A1848	M1665	F1666	K1667	D1669	M1670	P1674	S1677	Y1678	H1682	T1685	E1686	S1692	P1693	G1694	C1697	R1698	A1699	Y1703	S1704	P1708	H1713	P1719	L1720	L1721	I1722	K1725	L1728	H1729	K1730	M1733	M1734	P1735	M1736	D1737	M1738	R1739	E1740	L1743	M1746	D1749	E1750	K1751	K1752	W1569	Q1570	R1571	E1572	T1573	D1574	I1575	D1576	A1577	S1578	D1579	P1582	E1583	D1584	T1585	T1586	V1590	D1597	F1600	T1601	Y1609	L1616	I1619	I1620	R1621	A1622	D1625	D1626	V1627	R1631	F1632	K1633	M1634	L1635	A1636	S1641	L1642	H1645	Y1649	S1652	S1653	E1654	G1655	K1656	P1663	E1664	P1536	D1537	M1538	I1539	A1540	Y1543	L1544	R1545	R1551	M1552	Y1553	Y1554	Y1555	A1557	E1560	S1652	S1653	E1654	G1655	K1656	P1663	E1664	P1568	P1569	P1570	P1571	P1572	P1573	P1574	P1575	P1576	P1577	P1578	P1579	P1580	P1581	P1582	P1583	P1584	P1585	P1586	P1587	P1588	P1589	P1590	P1591	P1592	P1593	P1594	P1595	P1596	P1597	P1598	P1599	P1600	P1601	P1602	P1603	P1604	P1605	P1606	P1607	P1608	P1609	P1610	P1611	P1612	P1613	P1614	P1615	P1616	P1617	P1618	P1619	P1620	P1621	P1622	P1623	P1624	P1625	P1626	P1627	P1628	P1629	P1630	P1631	P1632	P1633	P1634	P1635	P1636	P1637	P1638	P1639	P1640	P1641	P1642	P1643	P1644	P1645	P1646	P1647	P1648	P1649	P1650	P1651	P1652	P1653	P1654	P1655	P1656	P1657	P1658	P1659	P1660	P1661	P1662	P1663	P1664	P1665	P1666	P1667	P1668	P1669	P1670	P1671	P1672	P1673	P1674	P1675	P1676	P1677	P1678	P1679	P1680	P1681	P1682	P1683	P1684	P1685	P1686	P1687	P1688	P1689	P1690	P1691	P1692	P1693	P1694	P1695	P1696	P1697	P1698	P1699	P1700	P1701	P1702	P1703	P1704	P1705	P1706	P1707	P1708	P1709	P1710	P1711	P1712	P1713	P1714	P1715	P1716	P1717	P1718	P1719	P1720	P1721	P1722	P1723	P1724	P1725	P1726	P1727	P1728	P1729	P1730	P1731	P1732	P1733	P1734	P1735	P1736	P1737	P1738	P1739	P1740	P1741	P1742	P1743	P1744	P1745	P1746	P1747	P1748	P1749	P1750	P1751	P1752	P1753	P1754	P1755	P1756	P1757	P1758	P1759	P1760	P1761	P1762	P1763	P1764	P1765	P1766	P1767	P1768	P1769	P1770	P1771	P1772	P1773	P1774	P1775	P1776	P1777	P1778	P1779	P1780	P1781	P1782	P1783	P1784	P1785	P1786	P1787	P1788	P1789	P1790	P1791	P1792	P1793	P1794	P1795	P1796	P1797	P1798	P1799	P1800	P1801	P1802	P1803	P1804	P1805	P1806	P1807	P1808	P1809	P1810	P1811	P1812	P1813	P1814	P1815	P1816	P1817	P1818	P1819	P1820	P1821	P1822	P1823	P1824	P1825	P1826	P1827	P1828	P1829	P1830	P1831	P1832	P1833	P1834	P1835	P1836	P1837	P1838	P1839	P1840	P1841	P1842	P1843	P1844	P1845	P1846	P1847	P1848	P1849	P1850	P1851	P1852	P1853	P1854	P1855	P1856	P1857	P1858	P1859	P1860	P1861	P1862	P1863	P1864	P1865	P1866	P1867	P1868	P1869	P1870	P1871	P1872	P1873	P1874	P1875	P1876	P1877	P1878	P1879	P1880	P1881	P1882	P1883	P1884	P1885	P1886	P1887	P1888	P1889	P1890	P1891	P1892	P1893	P1894	P1895	P1896	P1897	P1898	P1899	P1900	P1901	P1902	P1903	P1904	P1905	P1906	P1907	P1908	P1909	P1910	P1911	P1912	P1913	P1914	P1915	P1916	P1917	P1918	P1919	P1920	P1921	P1922	P1923	P1924	P1925	P1926	P1927	P1928	P1929	P1930	P1931	P1932	P1933	P1934	P1935	P1936	P1937	P1938	P1939	P1940	P1941	P1942	P1943	P1944	P1945	P1946	P1947	P1948	P1949	P1950	P1951	P1952	P1953	P1954	P1955	P1956	P1957	P1958	P1959	P1960	P1961	P1962	P1963	P1964	P1965	P1966	P1967	P1968	P1969	P1970	P1971	P1972	P1973	P1974	P1975	P1976	P1977	P1978	P1979	P1980	P1981	P1982	P1983	P1984	P1985	P1986	P1987	P1988	P1989	P1990	P1991	P1992	P1993	P1994	P1995	P1996	P1997	P1998	P1999	P2000	P2001	P2002	P2003	P2004	P2005	P2006	P2007	P2008	P2009	P2010	P2011	P2012	P2013	P2014	P2015	P2016	P2017	P2018	P2019	P2020	P2021	P2022	P2023	P2024	P2025	P2026	P2027	P2028	P2029	P2030	C2033	E2034	W2035	N2036	S2039	M2044	K2048	N2051	T2055	A2056	S2057	S2058	F2059	K2060	K2061	S2062	W2063	W2064	G2065	D2066	Y2067	P1569	Q1570	R1571	E1572	T1573	D1574	I1575	D1576	A1577	S1578	D1579	P1582	E1583	D1584	T1585	T1586	V1590	D1597	F1600	T1601	Y1609	L1616	I1619	I1620	R1621	A1622	D1625	D1626	V1627	R1631	F1632	K1633	M1634	L1635	A1636	S1641	L1642	H1645	Y1649	S1652	S1653	E1654	G1655	K1656	P1663	E1664	P1536	D1537	M1538	I1539	A1540	Y1543	L1544	R1545	R1551	M1552	Y1553	Y1554	Y1555	A1557	E1560	S1652	S1653	E1654	G1655	K1656	P1663	E1664	P1568	P1569	P1570	P1571	P1572	P1573	P1574	P1575	P1576	P1577	P1578	P1579	P1580	P1581	P1582	P1583	P1584	P1585	P1586	P1587	P1588	P1589	P1590	P1591	P1592	P1593	P1594	P1595	P1596	P1597	P1598	P1599	P1600	P1601	P1602	P1603	P1604	P1605	P1606	P1607	P1608	P1609	P1610	P1611	P1612	P1613	P1614	P1615	P1616	P1617	P1618	P1619	P1620	P1621	P1622	P1623	P1624	P1625	P1626	P1627	P1628	P1629	P1630	P1631	P1632	P1633	P1634	P1635	P1636	P1637	P1638	P1639	P1640	P1641	P1642	P1643	P1644	P1645	P1646	P1647	P1648	P1649	P1650	P1651	P1652	P1653	P1654	P1655	P1656	P1657	P1658	P1659	P1660	P1661	P1662	P1663	P1664	P1665	P1666	P1667	P1668	P1669	P1670	P1671	P1672	P1673	P1674	P1675	P1676	P1677	P1678	P1679	P1680	P1681	P1682	P1683	P1684	P1685	P1686	P1687	P1688	P1689	P1690	P1691	P1692	P1693	P1694	P1695	P1696	P1697	P1698	P1699	P1700	P1701	P1702	P1703	P1704	P1705	P1706	P1707	P1708	P1709	P1710	P1711	P1712	P1713	P1714	P1715	P1716	P1717	P1718	P1719	P1720	P1721	P1722	P1723	P1724	P1725	P1726	P1727	P1728	P1729	P1730	P1731	P1732	P1733	P1734	P1735	P1736	P1737	P1738	P1739	P1740	P1741	P1742	P1743	P1744	P1745	P1746	P1747	P1748	P1749	P1750	P1751	P1752	P1753	P1754	P1755	P1756	P1757	P1758	P1759	P1760	P1761	P1762	P1763	P1764	P1765	P1766	P1767	P1768	P1769	P1770	P1771	P1772	P1773	P1774	P1775	P1776	P1777	P1778	P1779	P1780	P1781	P1782	P1783	P1784	P1785	P1786	P1787	P1788	P1789	P1790	P1791	P1792	P1793	P1794	P1795	P1796	P1797	P1798	P1799	P1800	P1801	P1802	P1803	P1804	P1805	P1806	P1807	P1808	P1809	P1810	P1811	P1812	P1813	P1814	P1815	P1816	P1817	P1818	P1819	P1820	P1821	P1822	P1823	P1824	P1825	P1826	P1827	P1828	P1829	P1830	P1831	P1832	P1833	P1834	P1835	P1836	P1837	P1838	P1839	P1840	P1841	P1842	P1843	P1844	P1845	P1846	P1847	P1848	P1849	P1850	P1851	P1852	P1853	P1854	P1855	P1856	P1857	P1858	P1859	P1860	P1861	P1862	P1863	P1864	P1865	P1866	P1867	P1868	P1869	P1870	P1871	P1872	P1873	P1874	P1875	P1876	P1877	P1878	P1879	P1880	P1881	P1882	P1883	P1884	P1885	P1886	P1887	P1888	P1889	P1890	P1891	P1892	P1893	P1894	P1895	P1896	P1897	P1898	P1899	P1900	P1901	P1902	P1903	P1904	P1905	P1906	P1907	P1908	P1909	P1910	P1911	P1912	P1913	P1914	P1915	P1916	P1917	P1918	P1919	P1920	P1921	P1922	P1923	P1924	P1925	P1926	P1927	P1928	P1929	P1930	P1931	P1932	P1933	P1934	P1935	P1936	P1937	P1938	P1939	P1940	P1941	P1942	P1943	P1944	P1945	P1946	P1947	P1948	P1949	P1950	P1951	P1952	P1953	P1954	P1955	P1956	P1957	P1958	P1959	P1960	P1961	P1962	P1963	P1964	P1965	P1966	P1967	P1968	P1969	P1970	P1971	P1972	P1973	P1974	P1975	P1976	P1977	P1978	P1979	P1980	P1981	P1982	P1983	P1984	P1985	P1986	P1987	P1988	P1989	P1990	P1991	P1992	P1993	P1994	P1995	P1996	P1997	P1998	P1999	P2000	P2001	P2002	P2003	P2004	P2005	P2006	P2007	P2008	P2009	P2010	P2011	P2012	P2013	P2014	P2015	P2016	P2017	P2018	P2019	P2020	P2021	P2022	P2023	P2024	P2025	P2026	P2027	P2028	P2029	P2030	C2033	E2034	W2035	N2036	S2039	M2044	K2048	N2051	T2055	A2056	S2057	S2058	F2059	K2060	K2061	S2062	W2063	W2064	G2065	D2066	Y2067	P1569	Q1570	R1571	E1572	T1573	D1574	I1575	D1576	A1577	S1578	D1579	P1582	E1583	D1584	T1585	T1586	V1590	D1597	F1600	T1601	Y1609	L1616	I1619	I1620	R1621	A1622	D1625	D1626	V1627	R1631	F1632	K1633	M1634	L1635	A1636	S1641	L1642	H1645	Y1649	S1652	S1653	E1654	G1655	K1656	P1663	E1664	P1568	P1569	P1570	P1571	P1572	P1573	P1574	P1575	P1576	P1577	P1578	P1579	P1580	P1581	P1582	P1583	P1
-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	-------	----

R2171	I2176	M2181	A2185
F2172	P2177	Q2182	L2186
	K2178	S2183	R2187
	T2179	I2184	
	N2180		

## 4 Experimental information

Property	Value	Source
EM reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, Not provided	
Number of particles used	299182	Depositor
Resolution determination method	FSC 0.143 CUT-OFF	Depositor
CTF correction method	NONE	Depositor
Microscope	FEI TITAN KRIOS	Depositor
Voltage (kV)	300	Depositor
Electron dose ( $e^-/\text{\AA}^2$ )	1.65	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	2.803	Depositor
Minimum map value	-1.629	Depositor
Average map value	0.000	Depositor
Map value standard deviation	0.067	Depositor
Recommended contour level	0.43	Depositor
Map size ( $\text{\AA}$ )	325.6, 325.6, 325.6	wwPDB
Map dimensions	296, 296, 296	wwPDB
Map angles ( $^\circ$ )	90.0, 90.0, 90.0	wwPDB
Pixel spacing ( $\text{\AA}$ )	1.1, 1.1, 1.1	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	B	0.31	0/11436	0.51	0/15488

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	B	11142	0	10800	369	0
All	All	11142	0	10800	369	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 17.

All (369) 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:656:CYS:H	1:B:660:ASP:HB2	1.16	1.10
1:B:2060:LYS:CG	1:B:2087:LYS:HA	1.84	1.08
1:B:2061:LYS:HD2	1:B:2067:TYR:HB3	1.10	1.08
1:B:2060:LYS:HG3	1:B:2087:LYS:HA	1.39	1.03
1:B:2061:LYS:CD	1:B:2067:TYR:HB3	1.91	0.99

*Continued on next page...*



*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:2061:LYS:HD2	1:B:2067:TYR:CB	1.94	0.95
1:B:1750:GLU:HG3	1:B:1756:TYR:CD1	2.02	0.94
1:B:1899:GLU:HB3	1:B:1933:PRO:HB3	1.53	0.89
1:B:656:CYS:N	1:B:660:ASP:HB2	1.87	0.88
1:B:139:CYS:HB2	1:B:165:CYS:HA	1.55	0.86
1:B:656:CYS:H	1:B:660:ASP:CB	1.90	0.83
1:B:1901:LEU:HA	1:B:1925:LEU:HD23	1.65	0.78
1:B:1900:PHE:HB3	1:B:1922:VAL:H	1.48	0.78
1:B:1609:TYR:H	1:B:1984:THR:HG21	1.49	0.78
1:B:1958:LYS:NZ	1:B:1960:CYS:SG	2.57	0.77
1:B:656:CYS:O	1:B:660:ASP:HA	1.85	0.76
1:B:2131:GLU:HG2	1:B:2133:GLY:H	1.51	0.76
1:B:349:SER:O	1:B:358:GLN:NE2	2.20	0.74
1:B:659:ASP:HB3	1:B:672:GLU:HG3	1.71	0.73
1:B:94:LEU:HB3	1:B:108:GLU:HG2	1.71	0.73
1:B:289:GLN:HE22	1:B:484:MET:HG3	1.54	0.73
1:B:1750:GLU:CG	1:B:1756:TYR:CD1	2.72	0.73
1:B:99:SER:HB2	1:B:112:ASP:HB3	1.71	0.72
1:B:659:ASP:CB	1:B:672:GLU:HG3	2.19	0.72
1:B:1958:LYS:HG2	1:B:1960:CYS:HB2	1.71	0.72
1:B:2070:PRO:HA	1:B:2084:TRP:HB2	1.72	0.71
1:B:1750:GLU:HG3	1:B:1756:TYR:HD1	1.56	0.71
1:B:37:VAL:HG11	1:B:57:LEU:HD22	1.73	0.70
1:B:655:LYS:HA	1:B:660:ASP:HB2	1.74	0.70
1:B:2114:LYS:HB3	1:B:2119:GLU:HG2	1.73	0.69
1:B:659:ASP:HB3	1:B:672:GLU:CD	2.12	0.69
1:B:1743:LEU:HD23	1:B:1781:ASN:HD22	1.58	0.69
1:B:659:ASP:HB3	1:B:672:GLU:CG	2.23	0.68
1:B:2130:SER:HB3	1:B:2172:PHE:HB2	1.75	0.68
1:B:88:GLY:HA3	1:B:90:ARG:NH2	2.09	0.68
1:B:2034:GLU:HB2	1:B:2039:SER:HB2	1.76	0.67
1:B:2112:GLY:O	1:B:2187:ARG:NH2	2.27	0.67
1:B:1877:ARG:O	1:B:1877:ARG:HG2	1.95	0.67
1:B:181:LYS:HB2	1:B:225:ILE:HG13	1.77	0.66
1:B:639:ASN:HB2	1:B:643:ARG:HD2	1.77	0.66
1:B:2102:ILE:HD13	1:B:2132:GLN:HE21	1.60	0.66
1:B:2060:LYS:HG2	1:B:2087:LYS:HA	1.75	0.66
1:B:1904:TRP:HE3	1:B:1919:ALA:HB3	1.60	0.65
1:B:40:GLU:HB2	1:B:60:PRO:HB3	1.79	0.65
1:B:1894:GLN:HB2	1:B:1938:ASP:HB3	1.78	0.65
1:B:388:ASP:HA	1:B:391:LEU:HD13	1.79	0.64

*Continued on next page...*

*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1616:LEU:HD23	1:B:1719:PRO:HD3	1.79	0.64
1:B:1750:GLU:CD	1:B:1756:TYR:HD1	2.01	0.64
1:B:2077:ALA:O	1:B:2082:ASN:ND2	2.30	0.64
1:B:679:ARG:HG3	1:B:680:LYS:H	1.63	0.64
1:B:36:ILE:HG22	1:B:158:LEU:HD21	1.81	0.63
1:B:429:ASP:HB2	1:B:443:MET:HB2	1.80	0.63
1:B:255:VAL:HG22	1:B:257:ALA:H	1.64	0.63
1:B:2061:LYS:HG3	1:B:2066:ASP:O	1.98	0.63
1:B:1983:SER:OG	1:B:1984:THR:N	2.29	0.63
1:B:2057:SER:HB3	1:B:2093:GLN:HE22	1.64	0.63
1:B:320:LYS:HG3	1:B:321:ARG:H	1.64	0.63
1:B:2140:ARG:NH1	1:B:2145:MET:O	2.32	0.63
1:B:2060:LYS:HG3	1:B:2087:LYS:CA	2.24	0.62
1:B:315:GLN:HG3	1:B:316:ARG:HG2	1.82	0.62
1:B:2061:LYS:HE2	1:B:2067:TYR:HD2	1.64	0.62
1:B:1750:GLU:CG	1:B:1756:TYR:HD1	2.10	0.62
1:B:1621:ARG:NH1	1:B:1740:GLU:OE2	2.32	0.61
1:B:2062:SER:OG	1:B:2063:TRP:N	2.32	0.61
1:B:2130:SER:HB2	1:B:2135:GLU:O	2.00	0.61
1:B:499:LYS:HB2	1:B:501:ARG:HE	1.65	0.61
1:B:670:PRO:HB2	1:B:672:GLU:H	1.65	0.61
1:B:65:GLU:OE2	1:B:168:GLY:N	2.24	0.61
1:B:503:LEU:HD13	1:B:508:ILE:HG13	1.83	0.60
1:B:585:CYS:SG	1:B:586:SER:N	2.74	0.60
1:B:1544:LEU:HD13	1:B:1627:VAL:HG11	1.83	0.60
1:B:1728:LEU:HD12	1:B:1733:ASN:HB3	1.82	0.60
1:B:1750:GLU:O	1:B:1756:TYR:HB2	2.00	0.60
1:B:2122:VAL:HG21	1:B:2186:LEU:HD23	1.82	0.60
1:B:1698:ARG:NH2	1:B:1699:ALA:O	2.35	0.59
1:B:659:ASP:CB	1:B:672:GLU:CD	2.71	0.59
1:B:471:GLN:OE1	1:B:581:GLN:NE2	2.36	0.59
1:B:465:PRO:HB3	1:B:473:LEU:HD11	1.85	0.59
1:B:1633:LYS:HE3	1:B:1635:LEU:HD21	1.84	0.59
1:B:1902:GLY:HA2	1:B:1924:LYS:HD2	1.85	0.58
1:B:2000:LYS:HE3	1:B:2002:ASN:HD21	1.67	0.58
1:B:1568:PHE:HA	1:B:1572:GLU:HB3	1.85	0.58
1:B:605:TYR:N	1:B:608:ARG:O	2.37	0.58
1:B:1738:MET:SD	1:B:1799:ARG:NH1	2.76	0.57
1:B:1751:LYS:HG2	1:B:1756:TYR:CZ	2.39	0.57
1:B:1656:LYS:O	1:B:1670:ASN:ND2	2.37	0.57
1:B:645:LYS:O	1:B:648:ARG:NH2	2.36	0.57

*Continued on next page...*

*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1664:GLU:HA	1:B:1667:LYS:HB2	1.87	0.57
1:B:38:TYR:O	1:B:39:ARG:NH1	2.38	0.57
1:B:575:CYS:SG	1:B:576:PHE:N	2.77	0.57
1:B:47:LYS:HG3	1:B:48:GLU:H	1.68	0.57
1:B:524:ASP:OD2	1:B:527:LYS:NZ	2.37	0.56
1:B:275:GLU:HA	1:B:298:ILE:HB	1.87	0.56
1:B:1828:GLN:O	1:B:1829:HIS:ND1	2.38	0.56
1:B:515:GLU:OE1	1:B:517:GLN:NE2	2.26	0.56
1:B:1597:ASP:N	1:B:1597:ASP:OD1	2.38	0.56
1:B:1698:ARG:HH22	1:B:1841:PHE:HB2	1.69	0.56
1:B:1713:HIS:HD2	1:B:1749:ASP:H	1.53	0.56
1:B:356:SER:HB2	1:B:358:GLN:NE2	2.22	0.55
1:B:251:ASN:OD1	1:B:1829:HIS:NE2	2.36	0.55
1:B:517:GLN:HE21	1:B:583:HIS:HD2	1.54	0.55
1:B:1831:LEU:O	1:B:1833:VAL:N	2.36	0.55
1:B:545:VAL:HG12	1:B:547:ARG:HG2	1.88	0.55
1:B:139:CYS:HB2	1:B:165:CYS:CA	2.33	0.55
1:B:1823:GLU:OE2	1:B:1842:LYS:NZ	2.36	0.55
1:B:93:LYS:NZ	1:B:100:TYR:OH	2.40	0.54
1:B:18:ARG:HG2	1:B:213:THR:HG21	1.89	0.54
1:B:1980:LYS:HA	1:B:1983:SER:HB2	1.88	0.54
1:B:680:LYS:HG3	1:B:681:MET:H	1.72	0.54
1:B:1554:TYR:OH	1:B:1621:ARG:O	2.21	0.54
1:B:1904:TRP:H	1:B:1921:SER:HB3	1.71	0.54
1:B:386:LYS:HG3	1:B:387:GLU:HG3	1.88	0.54
1:B:2130:SER:OG	1:B:2131:GLU:N	2.41	0.54
1:B:1811:ILE:HG12	1:B:1838:PRO:HD3	1.90	0.54
1:B:1569:VAL:HA	1:B:1574:ASP:HB2	1.89	0.54
1:B:1785:TYR:CE1	1:B:1867:GLY:HA2	2.42	0.54
1:B:2099:LEU:HD13	1:B:2103:LYS:HD2	1.90	0.54
1:B:655:LYS:CA	1:B:660:ASP:HB2	2.36	0.54
1:B:1544:LEU:HD23	1:B:1551:ARG:HE	1.73	0.54
1:B:1750:GLU:HG3	1:B:1750:GLU:O	2.06	0.54
1:B:1539:ILE:HD11	1:B:1678:TYR:HA	1.89	0.54
1:B:1904:TRP:O	1:B:1921:SER:N	2.41	0.54
1:B:1697:CYS:SG	1:B:1799:ARG:NH2	2.81	0.53
1:B:181:LYS:O	1:B:226:SER:OG	2.25	0.53
1:B:2085:GLN:HA	1:B:2184:ILE:O	2.08	0.53
1:B:1949:GLN:HG2	1:B:2001:GLU:HB3	1.89	0.53
1:B:205:THR:OG1	1:B:209:TYR:N	2.40	0.53
1:B:599:THR:HG22	1:B:633:TRP:HD1	1.74	0.53

*Continued on next page...*

*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1897:ALA:HB1	1:B:1920:TRP:HE1	1.73	0.53
1:B:2060:LYS:O	1:B:2060:LYS:HD3	2.09	0.53
1:B:593:ILE:HG22	1:B:617:PRO:HD3	1.91	0.53
1:B:1797:TRP:CE2	1:B:1847:LYS:HD3	2.44	0.53
1:B:22:THR:HG22	1:B:23:ASN:H	1.73	0.53
1:B:174:GLY:O	1:B:182:GLN:NE2	2.31	0.53
1:B:334:ASP:HA	1:B:352:LEU:HD11	1.91	0.53
1:B:356:SER:HB2	1:B:358:GLN:HE22	1.73	0.52
1:B:1938:ASP:OD1	1:B:1939:MET:N	2.41	0.52
1:B:1823:GLU:HG2	1:B:1824:ASN:H	1.75	0.52
1:B:335:TYR:HB2	1:B:347:TYR:CZ	2.45	0.52
1:B:1777:PHE:HB3	1:B:1784:ILE:HD11	1.91	0.52
1:B:358:GLN:N	1:B:358:GLN:OE1	2.43	0.52
1:B:2089:ASN:HD21	1:B:2183:SER:HA	1.74	0.52
1:B:195:SER:OG	1:B:196:TRP:N	2.42	0.52
1:B:329:GLU:OE2	1:B:371:TYR:OH	2.27	0.51
1:B:373:ASP:OD2	1:B:375:SER:OG	2.23	0.51
1:B:322:TRP:HB3	1:B:401:ASP:O	2.11	0.51
1:B:1622:ALA:HB3	1:B:1722:ILE:HD13	1.93	0.51
1:B:2060:LYS:HD3	1:B:2060:LYS:C	2.31	0.51
1:B:655:LYS:HA	1:B:660:ASP:CB	2.39	0.51
1:B:2089:ASN:ND2	1:B:2182:GLN:O	2.44	0.51
1:B:1904:TRP:CE3	1:B:1919:ALA:HB3	2.44	0.50
1:B:179:PHE:CD1	1:B:226:SER:HB3	2.46	0.50
1:B:2061:LYS:CE	1:B:2067:TYR:HB3	2.40	0.50
1:B:2063:TRP:O	1:B:2064:TRP:HD1	1.93	0.50
1:B:1560:GLU:HG3	1:B:1590:VAL:HG12	1.94	0.50
1:B:1746:MET:SD	1:B:1806:GLY:HA3	2.51	0.50
1:B:1803:LEU:HD22	1:B:1841:PHE:HD1	1.76	0.50
1:B:1991:ASN:OD1	1:B:1992:GLY:N	2.43	0.50
1:B:2061:LYS:HE2	1:B:2067:TYR:CD2	2.46	0.50
1:B:1685:THR:OG1	1:B:1686:GLU:N	2.43	0.49
1:B:3:LEU:HD12	1:B:71:LYS:HD3	1.94	0.49
1:B:1750:GLU:HG3	1:B:1756:TYR:CE1	2.45	0.49
1:B:499:LYS:HD2	1:B:501:ARG:HH11	1.78	0.49
1:B:385:MET:O	1:B:386:LYS:HG2	2.13	0.49
1:B:568:SER:O	1:B:568:SER:OG	2.30	0.49
1:B:638:MET:SD	1:B:1708:PRO:HB2	2.52	0.49
1:B:639:ASN:N	1:B:644:SER:OG	2.46	0.49
1:B:660:ASP:O	1:B:664:SER:HB3	2.13	0.49
1:B:232:MET:HA	1:B:263:ALA:H	1.78	0.48

*Continued on next page...*

*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:424:PHE:CD1	1:B:430:GLU:HB3	2.48	0.48
1:B:1750:GLU:CD	1:B:1756:TYR:CD1	2.83	0.48
1:B:95:SER:OG	1:B:124:TRP:NE1	2.46	0.48
1:B:532:GLU:HA	1:B:536:ASN:HB2	1.94	0.48
1:B:1925:LEU:HD12	1:B:1926:ALA:N	2.28	0.48
1:B:466:THR:HG22	1:B:468:ASN:H	1.77	0.48
1:B:333:TRP:CH2	1:B:390:ILE:HD12	2.48	0.48
1:B:368:TYR:CZ	1:B:417:ILE:HG22	2.49	0.48
1:B:1750:GLU:CG	1:B:1756:TYR:CE1	2.97	0.48
1:B:659:ASP:CB	1:B:672:GLU:CG	2.84	0.48
1:B:1963:THR:OG1	1:B:2019:ARG:O	2.26	0.48
1:B:134:HIS:CD2	1:B:135:ASP:H	2.32	0.48
1:B:343:MET:HE3	1:B:344:ASP:H	1.77	0.48
1:B:429:ASP:CB	1:B:443:MET:HB2	2.44	0.48
1:B:1641:SER:O	1:B:1704:SER:HA	2.14	0.47
1:B:1734:MET:O	1:B:1736:MET:HG2	2.14	0.47
1:B:2089:ASN:OD1	1:B:2183:SER:HA	2.14	0.47
1:B:402:THR:HG22	1:B:402:THR:O	2.14	0.47
1:B:1557:ALA:HB2	1:B:1600:PHE:HE2	1.78	0.47
1:B:1787:LEU:HB3	1:B:1870:THR:HG21	1.97	0.47
1:B:519:VAL:HG22	1:B:562:ASN:ND2	2.29	0.47
1:B:245:GLY:O	1:B:423:THR:OG1	2.24	0.47
1:B:351:HIS:HB3	1:B:363:TYR:OH	2.14	0.47
1:B:1981:GLY:O	1:B:1986:ASN:ND2	2.47	0.47
1:B:2084:TRP:O	1:B:2185:ALA:HA	2.14	0.47
1:B:327:ALA:HB3	1:B:371:TYR:HE1	1.79	0.47
1:B:368:TYR:OH	1:B:416:SER:O	2.32	0.47
1:B:520:PHE:HE1	1:B:584:PHE:HD1	1.60	0.47
1:B:595:THR:HG21	1:B:1838:PRO:O	2.14	0.47
1:B:1555:TYR:CE2	1:B:1631:ARG:HD2	2.49	0.47
1:B:1812:HIS:O	1:B:1836:LEU:N	2.38	0.47
1:B:2141:LEU:HD12	1:B:2142:LYS:H	1.79	0.47
1:B:222:HIS:H	1:B:272:VAL:HG23	1.79	0.47
1:B:679:ARG:HG3	1:B:680:LYS:N	2.29	0.46
1:B:657:ILE:HB	1:B:658:PRO:HD3	1.97	0.46
1:B:2023:ARG:O	1:B:2025:THR:N	2.49	0.46
1:B:2066:ASP:OD1	1:B:2066:ASP:N	2.38	0.46
1:B:432:ASN:CG	1:B:445:ARG:HG3	2.36	0.46
1:B:1692:SER:O	1:B:1694:GLY:N	2.48	0.46
1:B:1980:LYS:HA	1:B:1980:LYS:HD2	1.70	0.46
1:B:66:VAL:HG23	1:B:165:CYS:O	2.15	0.46

*Continued on next page...*

*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1842:LYS:HD2	1:B:1842:LYS:HA	1.80	0.46
1:B:126:ILE:HD12	1:B:126:ILE:H	1.80	0.46
1:B:153:ASP:O	1:B:158:LEU:HB2	2.16	0.46
1:B:1699:ALA:HB2	1:B:1801:HIS:ND1	2.30	0.46
1:B:2055:THR:HG23	1:B:2096:GLU:HB2	1.98	0.46
1:B:357:ASN:HB2	1:B:540:GLU:HB3	1.97	0.46
1:B:657:ILE:H	1:B:658:PRO:CD	2.29	0.46
1:B:1649:TYR:HB2	1:B:1653:SER:HB2	1.98	0.46
1:B:332:ILE:HG23	1:B:362:HIS:HA	1.98	0.46
1:B:413:ARG:N	1:B:449:PRO:HB3	2.31	0.46
1:B:457:TRP:NE1	1:B:459:ILE:HG12	2.31	0.46
1:B:1804:ASN:O	1:B:1839:GLY:N	2.49	0.46
1:B:572:LEU:HB2	1:B:651:PHE:HB2	1.97	0.45
1:B:1757:GLU:OE2	1:B:1761:ARG:NH2	2.44	0.45
1:B:1887:THR:OG1	1:B:1889:ILE:HG12	2.16	0.45
1:B:1936:GLN:HE22	1:B:1973:GLN:NE2	2.14	0.45
1:B:205:THR:HG1	1:B:209:TYR:C	2.19	0.45
1:B:1750:GLU:O	1:B:1756:TYR:CD1	2.69	0.45
1:B:1925:LEU:HD11	1:B:2023:ARG:NH1	2.31	0.45
1:B:391:LEU:HG	1:B:517:GLN:HB2	1.99	0.45
1:B:350:GLN:HA	1:B:358:GLN:HG2	1.99	0.45
1:B:2044:MET:O	1:B:2074:ARG:NE	2.50	0.45
1:B:2113:CYS:SG	1:B:2114:LYS:N	2.89	0.45
1:B:95:SER:OG	1:B:95:SER:O	2.34	0.45
1:B:138:PRO:HG2	1:B:179:PHE:CZ	2.52	0.45
1:B:1553:ASN:HB3	1:B:1631:ARG:HH22	1.82	0.45
1:B:2124:SER:HB3	1:B:2179:THR:OG1	2.17	0.45
1:B:96:GLU:HG3	1:B:98:ALA:H	1.82	0.45
1:B:244:ASN:ND2	1:B:279:ILE:O	2.32	0.45
1:B:346:LYS:HZ1	1:B:555:SER:HB2	1.82	0.45
1:B:598:PHE:CZ	1:B:625:VAL:HG21	2.52	0.45
1:B:1794:GLU:HG2	1:B:1795:GLN:HG2	1.98	0.45
1:B:343:MET:CE	1:B:344:ASP:H	2.30	0.45
1:B:2120:MET:HB2	1:B:2181:ASN:O	2.17	0.45
1:B:2:GLN:N	1:B:68:ASP:OD2	2.50	0.44
1:B:11:GLN:HE21	1:B:39:ARG:HG3	1.82	0.44
1:B:329:GLU:O	1:B:366:VAL:HA	2.17	0.44
1:B:1904:TRP:N	1:B:1904:TRP:CD1	2.84	0.44
1:B:134:HIS:CG	1:B:135:ASP:H	2.35	0.44
1:B:1652:SER:O	1:B:1652:SER:OG	2.32	0.44
1:B:2028:LEU:HD23	1:B:2028:LEU:H	1.82	0.44

*Continued on next page...*

*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:479:SER:OG	1:B:486:ASP:OD2	2.29	0.44
1:B:525:GLU:O	1:B:528:SER:OG	2.27	0.44
1:B:1568:PHE:O	1:B:1574:ASP:HB2	2.18	0.44
1:B:1575:ILE:HD12	1:B:1575:ILE:H	1.83	0.44
1:B:457:TRP:HE1	1:B:459:ILE:HG12	1.82	0.44
1:B:2051:ASN:HD21	1:B:2071:PHE:HB2	1.81	0.44
1:B:494:LEU:HD21	1:B:519:VAL:HG11	1.99	0.44
1:B:503:LEU:HD11	1:B:510:ARG:HD2	2.00	0.44
1:B:2051:ASN:ND2	1:B:2071:PHE:HB2	2.33	0.44
1:B:2110:THR:HG22	1:B:2111:GLN:N	2.33	0.44
1:B:143:ILE:HG22	1:B:230:LEU:HD13	2.00	0.44
1:B:147:HIS:O	1:B:147:HIS:ND1	2.49	0.44
1:B:1713:HIS:CD2	1:B:1749:ASP:H	2.35	0.43
1:B:381:VAL:HG11	1:B:385:MET:SD	2.59	0.43
1:B:1848:ALA:HB1	1:B:1874:ILE:HD12	2.01	0.43
1:B:491:LEU:HD23	1:B:491:LEU:HA	1.75	0.43
1:B:373:ASP:OD2	1:B:375:SER:N	2.50	0.43
1:B:390:ILE:HG21	1:B:562:ASN:O	2.18	0.43
1:B:1802:LEU:HD13	1:B:1814:VAL:HG21	2.00	0.43
1:B:1652:SER:HB2	1:B:1666:PHE:HA	2.01	0.43
1:B:2083:ALA:HB2	1:B:2187:ARG:HG3	2.01	0.43
1:B:656:CYS:O	1:B:660:ASP:OD1	2.36	0.43
1:B:657:ILE:N	1:B:658:PRO:CD	2.80	0.43
1:B:74:PHE:HE2	1:B:76:ASN:HB2	1.84	0.43
1:B:554:GLU:HA	1:B:557:ILE:HG12	1.99	0.43
1:B:594:LEU:HA	1:B:639:ASN:HD21	1.84	0.43
1:B:1579:ASP:N	1:B:1579:ASP:OD1	2.52	0.43
1:B:1663:PRO:O	1:B:1665:TRP:N	2.45	0.43
1:B:1792:MET:O	1:B:1875:MET:HG2	2.19	0.43
1:B:2048:LYS:HB2	1:B:2048:LYS:HE3	1.81	0.43
1:B:2070:PRO:HB3	1:B:2084:TRP:CD1	2.54	0.43
1:B:17:TYR:HB3	1:B:210:VAL:HG23	2.01	0.43
1:B:1663:PRO:C	1:B:1665:TRP:H	2.21	0.43
1:B:70:ILE:O	1:B:123:GLU:HA	2.19	0.43
1:B:418:TYR:HE1	1:B:424:PHE:HZ	1.67	0.43
1:B:1633:LYS:HG3	1:B:1677:SER:HB3	2.01	0.43
1:B:86:PRO:HB2	1:B:90:ARG:NH1	2.34	0.42
1:B:385:MET:HB2	1:B:388:ASP:OD2	2.18	0.42
1:B:2123:LYS:HE3	1:B:2181:ASN:HB2	1.99	0.42
1:B:51:GLN:OE1	1:B:55:SER:OG	2.37	0.42
1:B:1829:HIS:HB3	1:B:1831:LEU:HD23	2.01	0.42

*Continued on next page...*

*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:2060:LYS:CB	1:B:2087:LYS:HA	2.46	0.42
1:B:161:PRO:HG3	1:B:230:LEU:HD11	2.01	0.42
1:B:1634:ASN:OD1	1:B:1635:LEU:N	2.52	0.42
1:B:1887:THR:OG1	1:B:1888:GLY:N	2.53	0.42
1:B:80:LYS:NZ	1:B:153:ASP:OD1	2.41	0.42
1:B:335:TYR:CD2	1:B:558:MET:HG2	2.55	0.42
1:B:393:PRO:HG2	1:B:477:TYR:OH	2.19	0.42
1:B:599:THR:HG22	1:B:633:TRP:CD1	2.54	0.42
1:B:670:PRO:HD2	1:B:671:PRO:HA	2.00	0.42
1:B:1793:TYR:N	1:B:1796:GLU:OE2	2.35	0.42
1:B:1822:LEU:HD12	1:B:1829:HIS:O	2.19	0.42
1:B:1625:ASP:HB3	1:B:1725:LYS:HZ1	1.85	0.42
1:B:2130:SER:OG	1:B:2133:GLY:O	2.35	0.42
1:B:205:THR:OG1	1:B:206:VAL:N	2.53	0.42
1:B:351:HIS:HB3	1:B:363:TYR:HH	1.85	0.42
1:B:398:GLN:OE1	1:B:399:VAL:HG12	2.19	0.42
1:B:1885:LEU:O	1:B:1910:ARG:HD2	2.19	0.42
1:B:2089:ASN:ND2	1:B:2183:SER:HA	2.35	0.42
1:B:1654:GLU:N	1:B:1669:ASP:OD2	2.42	0.42
1:B:366:VAL:HG12	1:B:491:LEU:HD21	2.02	0.42
1:B:1564:ASP:HB3	1:B:1586:THR:HA	2.01	0.42
1:B:2102:ILE:HG21	1:B:2132:GLN:NE2	2.34	0.42
1:B:333:TRP:CZ2	1:B:390:ILE:HD12	2.55	0.42
1:B:1564:ASP:N	1:B:1564:ASP:OD1	2.53	0.41
1:B:1572:GLU:HG2	1:B:1786:SER:HB3	2.02	0.41
1:B:1619:ILE:HG23	1:B:1721:LEU:HD13	2.02	0.41
1:B:2111:GLN:HG2	1:B:2112:GLY:H	1.85	0.41
1:B:1694:GLY:O	1:B:1824:ASN:ND2	2.53	0.41
1:B:2132:GLN:OE1	1:B:2132:GLN:N	2.51	0.41
1:B:204:TYR:O	1:B:292:MET:HB3	2.20	0.41
1:B:4:ARG:HB2	1:B:70:ILE:HG12	2.03	0.41
1:B:2102:ILE:HA	1:B:2171:ARG:HB3	2.01	0.41
1:B:351:HIS:O	1:B:359:ILE:HD12	2.21	0.41
1:B:409:ASN:ND2	1:B:447:VAL:O	2.51	0.41
1:B:656:CYS:HA	1:B:657:ILE:HA	1.65	0.41
1:B:2013:ILE:HG21	1:B:2030:LEU:HD21	2.02	0.41
1:B:2058:SER:CB	1:B:2087:LYS:HB2	2.50	0.41
1:B:1752:LYS:HE3	1:B:1752:LYS:HB3	1.90	0.41
1:B:1805:ILE:HD13	1:B:1839:GLY:HA2	2.02	0.41
1:B:390:ILE:HG12	1:B:390:ILE:O	2.20	0.41
1:B:1584:ASP:OD1	1:B:1585:THR:N	2.53	0.41

*Continued on next page...*



*Continued from previous page...*

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:1600:PHE:O	1:B:1601:THR:OG1	2.27	0.41
1:B:686:GLU:HB2	1:B:687:PRO:HD3	2.02	0.41
1:B:1636:ALA:O	1:B:1674:PRO:HB3	2.21	0.41
1:B:1645:HIS:HE1	1:B:1703:TYR:OH	2.04	0.41
1:B:1797:TRP:CZ3	1:B:1847:LYS:HB2	2.56	0.41
1:B:220:CYS:SG	1:B:221:ALA:N	2.91	0.41
1:B:388:ASP:HB3	1:B:391:LEU:HB2	2.02	0.41
1:B:659:ASP:CG	1:B:672:GLU:HG3	2.41	0.41
1:B:679:ARG:CG	1:B:680:LYS:H	2.31	0.41
1:B:688:GLU:HG2	1:B:689:ASP:H	1.85	0.41
1:B:1827:LYS:HA	1:B:1827:LYS:HD3	1.90	0.41
1:B:2060:LYS:HG2	1:B:2087:LYS:HG3	2.03	0.41
1:B:2084:TRP:CZ3	1:B:2095:LEU:HB2	2.56	0.41
1:B:2111:GLN:HA	1:B:2153:ASN:HD21	1.85	0.41
1:B:2141:LEU:HD13	1:B:2148:LYS:HE2	2.03	0.41
1:B:20:GLU:HB3	1:B:26:LEU:HD11	2.03	0.41
1:B:420:HIS:NE2	1:B:478:TYR:OH	2.23	0.41
1:B:120:TYR:CG	1:B:121:THR:N	2.88	0.40
1:B:1556:ILE:HG22	1:B:1557:ALA:N	2.36	0.40
1:B:526:ASN:OD1	1:B:556:ASN:ND2	2.33	0.40
1:B:1544:LEU:C	1:B:1545:ARG:HD2	2.42	0.40
1:B:1642:LEU:HD23	1:B:1682:TRP:HH2	1.87	0.40
1:B:1962:THR:OG1	1:B:2025:THR:O	2.38	0.40
1:B:2156:THR:HG23	1:B:2157:LYS:H	1.86	0.40
1:B:1898:SER:HB3	1:B:1934:TRP:CE2	2.57	0.40
1:B:1997:SER:O	1:B:1997:SER:OG	2.33	0.40
1:B:2176:ILE:HG22	1:B:2178:LYS:HG2	2.02	0.40
1:B:213:THR:O	1:B:213:THR:HG22	2.22	0.40
1:B:601:HIS:NE2	1:B:653:ASP:OD2	2.54	0.40
1:B:640:SER:OG	1:B:642:PRO:HD2	2.20	0.40
1:B:1878:ASP:O	1:B:1880:ARG:N	2.53	0.40
1:B:306:ARG:O	1:B:307:ASN:ND2	2.55	0.40
1:B:403:LEU:HB3	1:B:458:ASN:HA	2.04	0.40
1:B:445:ARG:H	1:B:445:ARG:HD3	1.87	0.40
1:B:1851:PRO:HA	1:B:1874:ILE:HB	2.03	0.40
1:B:2034:GLU:HB3	1:B:2035:VAL:H	1.69	0.40

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	B	1370/2196 (62%)	1115 (81%)	250 (18%)	5 (0%)	34 66

All (5) Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	B	2060	LYS
1	B	135	ASP
1	B	700	ASN
1	B	658	PRO
1	B	670	PRO

### 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	B	1225/1989 (62%)	1214 (99%)	11 (1%)	78 87

All (11) residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	B	90	ARG
1	B	445	ARG
1	B	655	LYS
1	B	1698	ARG
1	B	1749	ASP
1	B	1750	GLU

*Continued on next page...*

*Continued from previous page...*

Mol	Chain	Res	Type
1	B	1920	TRP
1	B	2033	CYS
1	B	2034	GLU
1	B	2036	ASN
1	B	2061	LYS

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. All (22) such sidechains are listed below:

Mol	Chain	Res	Type
1	B	134	HIS
1	B	252	HIS
1	B	289	GLN
1	B	307	ASN
1	B	315	GLN
1	B	350	GLN
1	B	562	ASN
1	B	583	HIS
1	B	609	HIS
1	B	639	ASN
1	B	1570	GLN
1	B	1643	HIS
1	B	1645	HIS
1	B	1670	ASN
1	B	1713	HIS
1	B	1778	HIS
1	B	1781	ASN
1	B	1819	GLN
1	B	1936	GLN
1	B	2002	ASN
1	B	2093	GLN
1	B	2162	ASN

### 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.

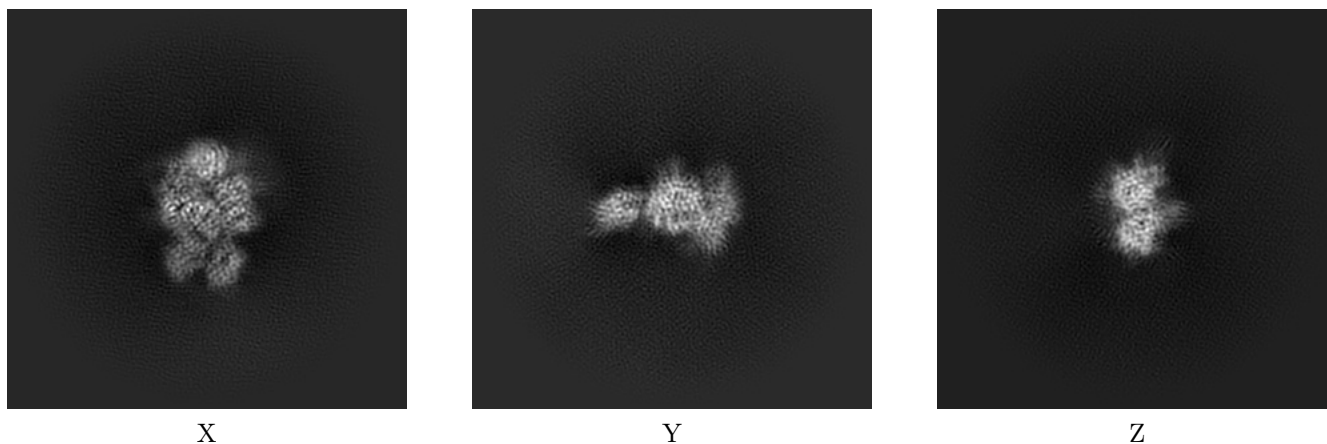
## 6 Map visualisation [i](#)

This section contains visualisations of the EMDB entry EMD-23048. These allow visual inspection of the internal detail of the map and identification of artifacts.

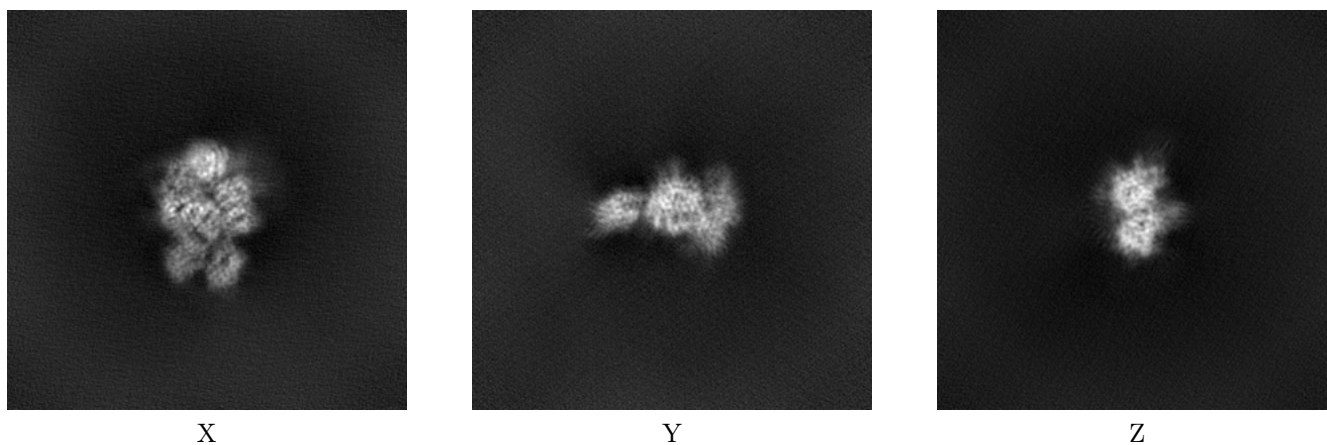
Images derived from a raw map, generated by summing the deposited half-maps, are presented below the corresponding image components of the primary map to allow further visual inspection and comparison with those of the primary map.

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

#### 6.1.1 Primary map



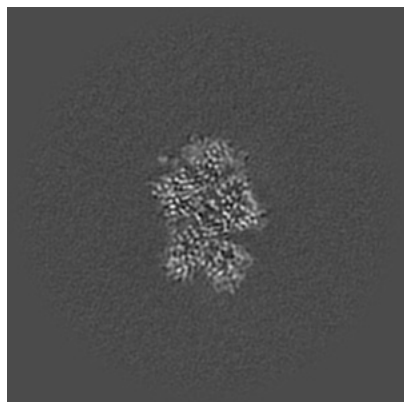
#### 6.1.2 Raw map



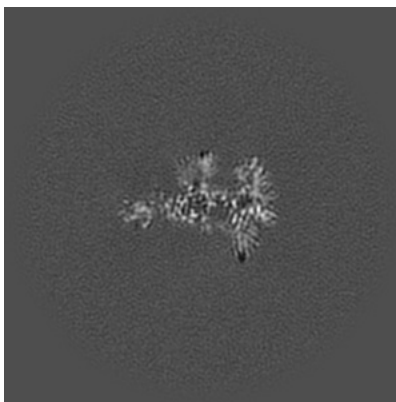
The images above show the map projected in three orthogonal directions.

## 6.2 Central slices [i](#)

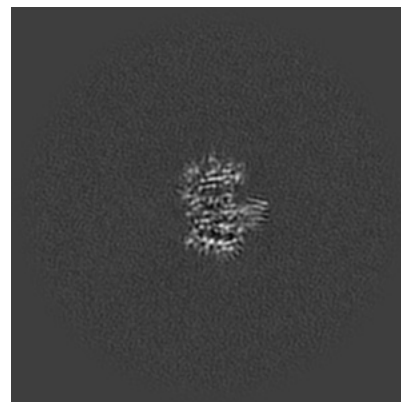
### 6.2.1 Primary map



X Index: 148

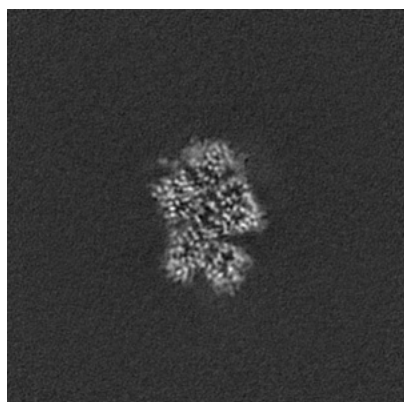


Y Index: 148

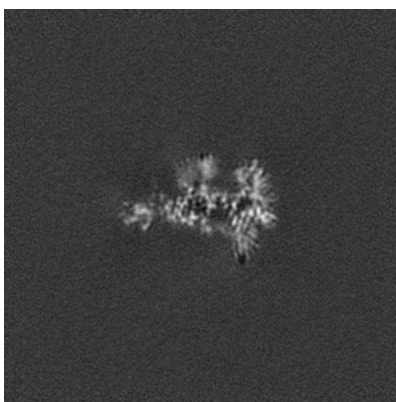


Z Index: 148

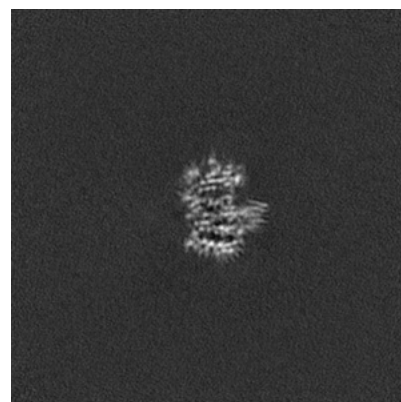
### 6.2.2 Raw map



X Index: 148



Y Index: 148

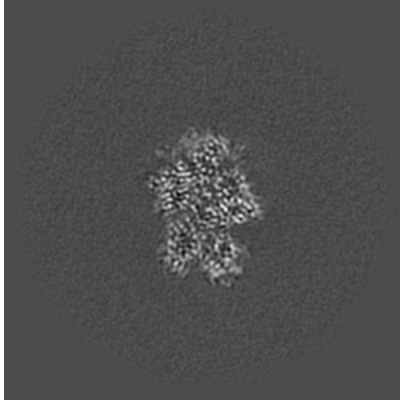


Z Index: 148

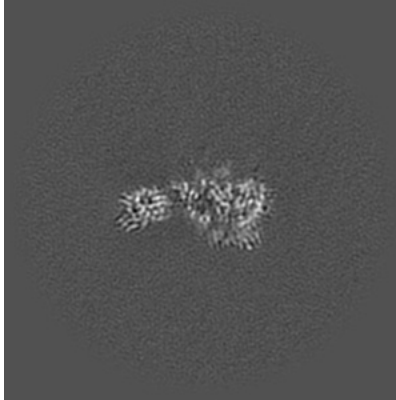
The images above show central slices of the map in three orthogonal directions.

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

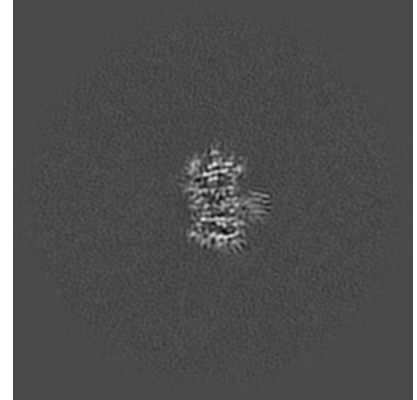
### 6.3.1 Primary map



X Index: 149

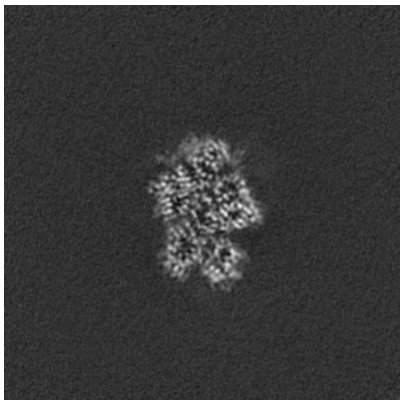


Y Index: 157

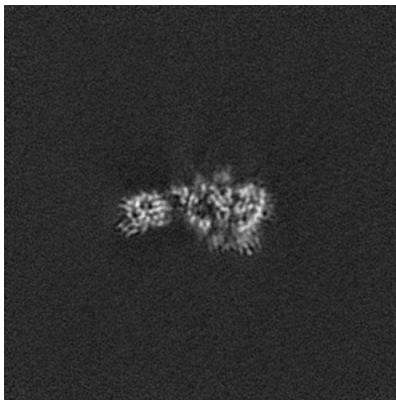


Z Index: 147

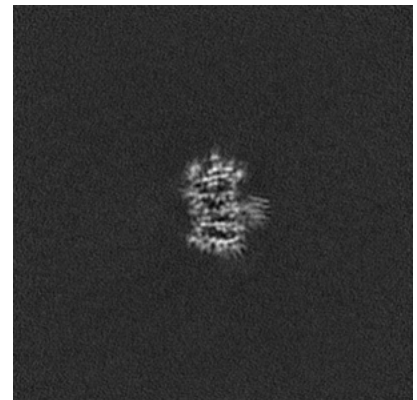
### 6.3.2 Raw map



X Index: 149



Y Index: 157

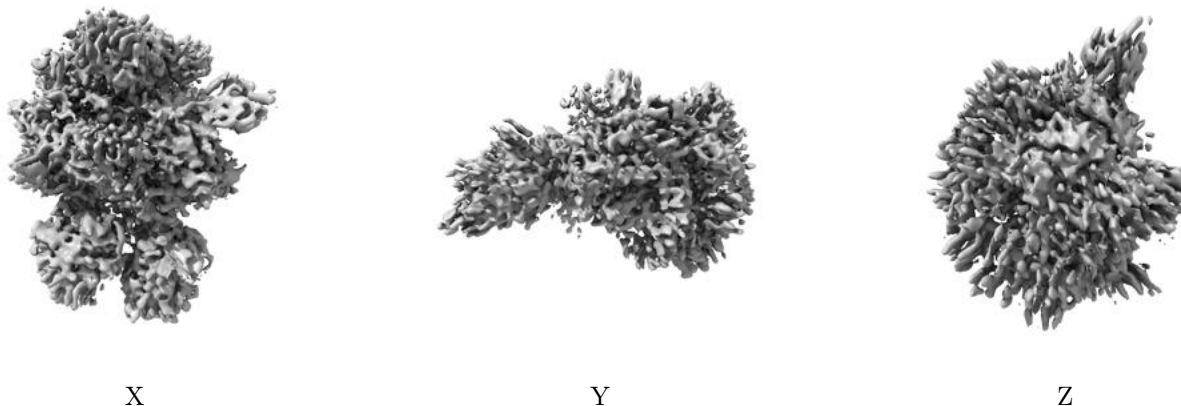


Z Index: 147

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

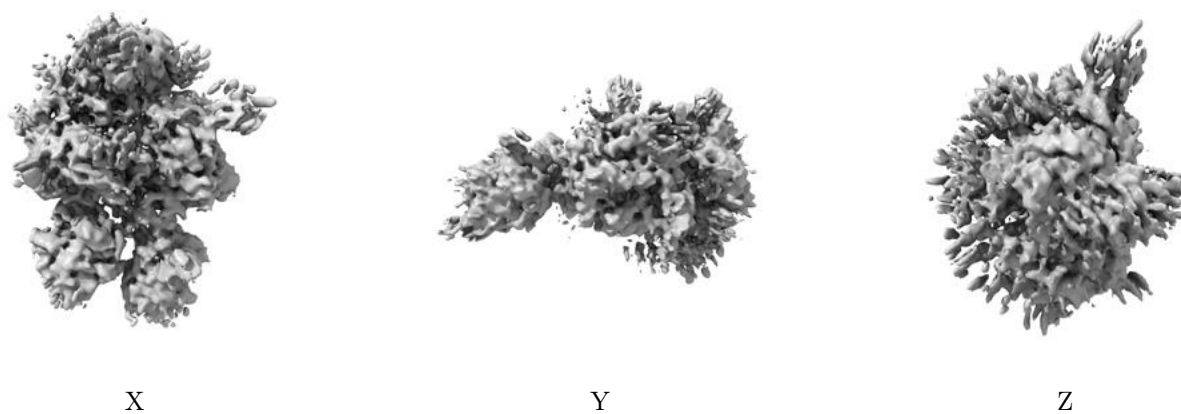
## 6.4 Orthogonal surface views [i](#)

### 6.4.1 Primary map



The images above show the 3D surface view of the map at the recommended contour level 0.43. 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.



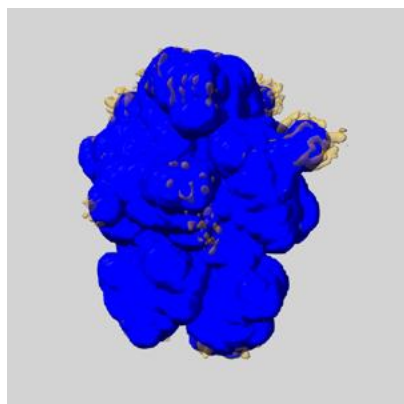
## 6.5 Mask visualisation [i](#)

This section shows the 3D surface view of the primary map at 50% transparency overlaid with the specified mask at 0% transparency

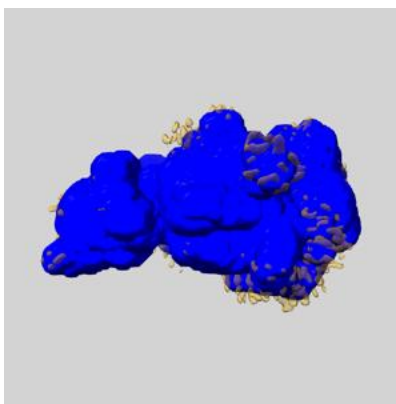
A mask typically either:

- Encompasses the whole structure
- Separates out a domain, a functional unit, a monomer or an area of interest from a larger structure

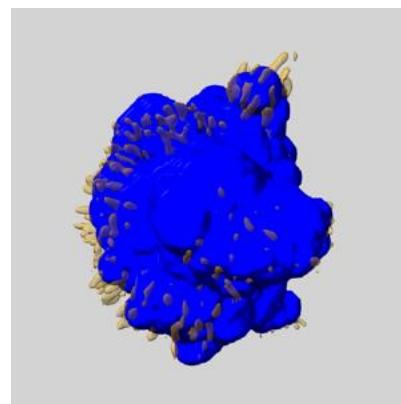
### 6.5.1 emd\_23048\_msk\_1.map [i](#)



X



Y

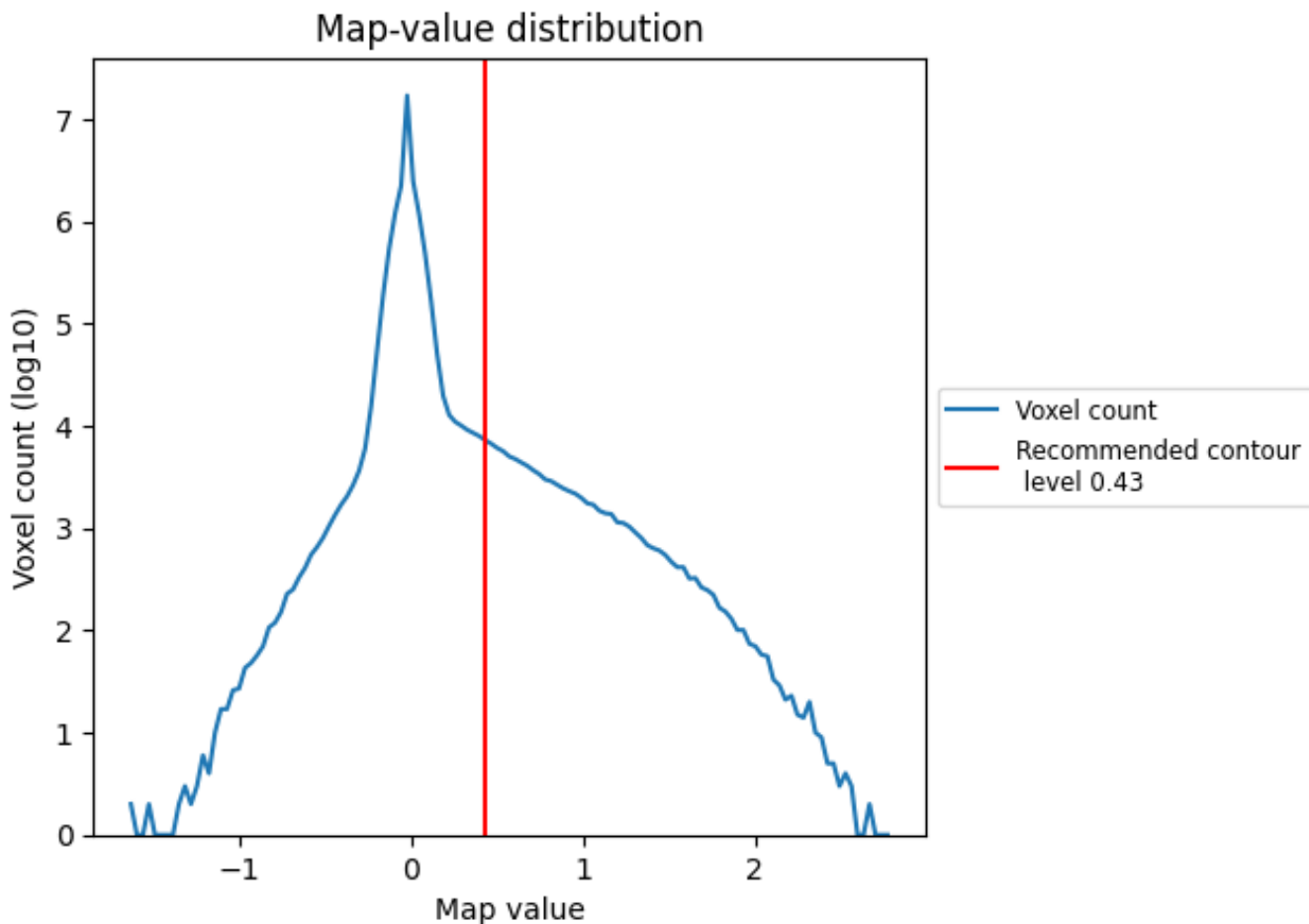


Z

## 7 Map analysis [i](#)

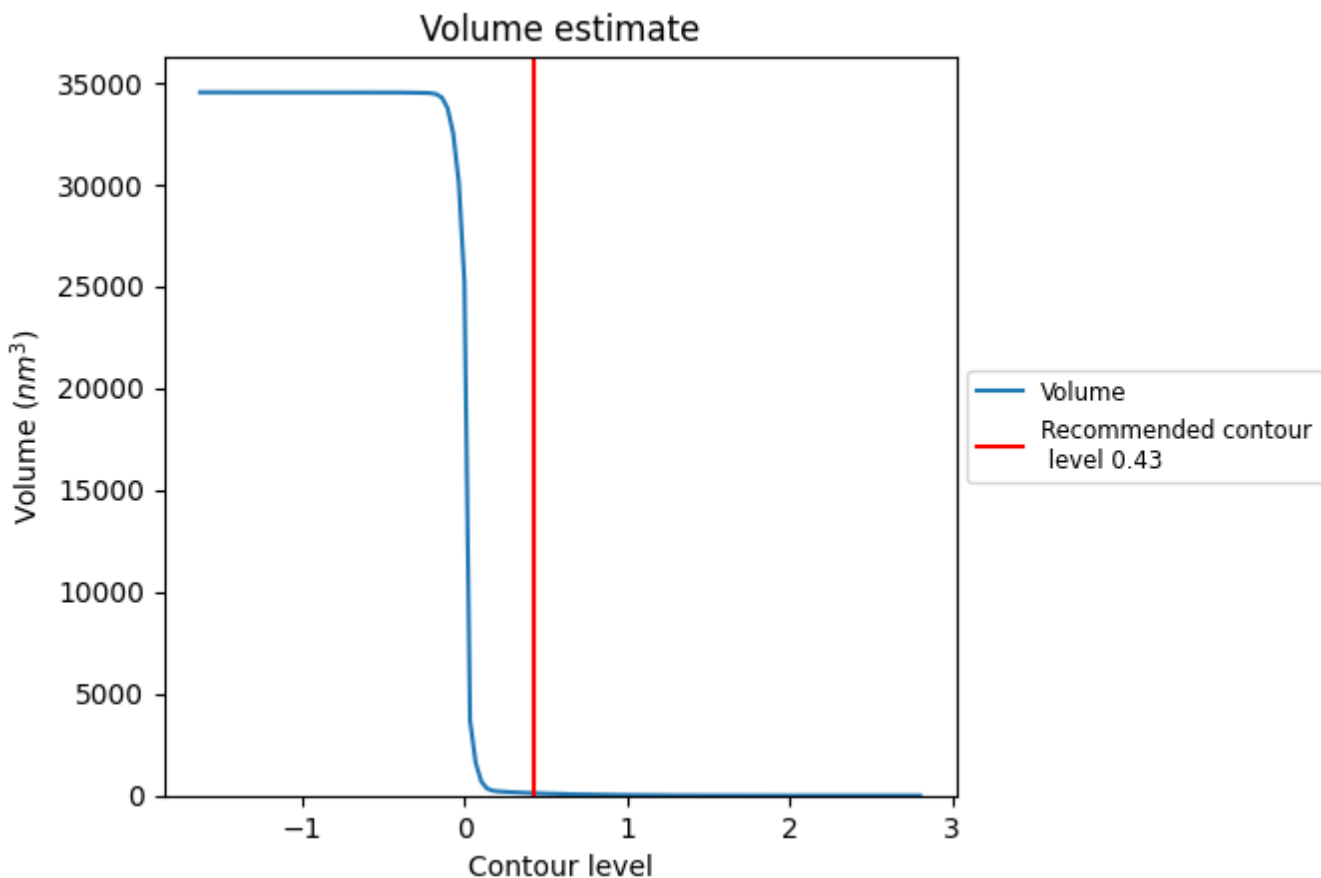
This section contains the results of statistical analysis of the map.

### 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.

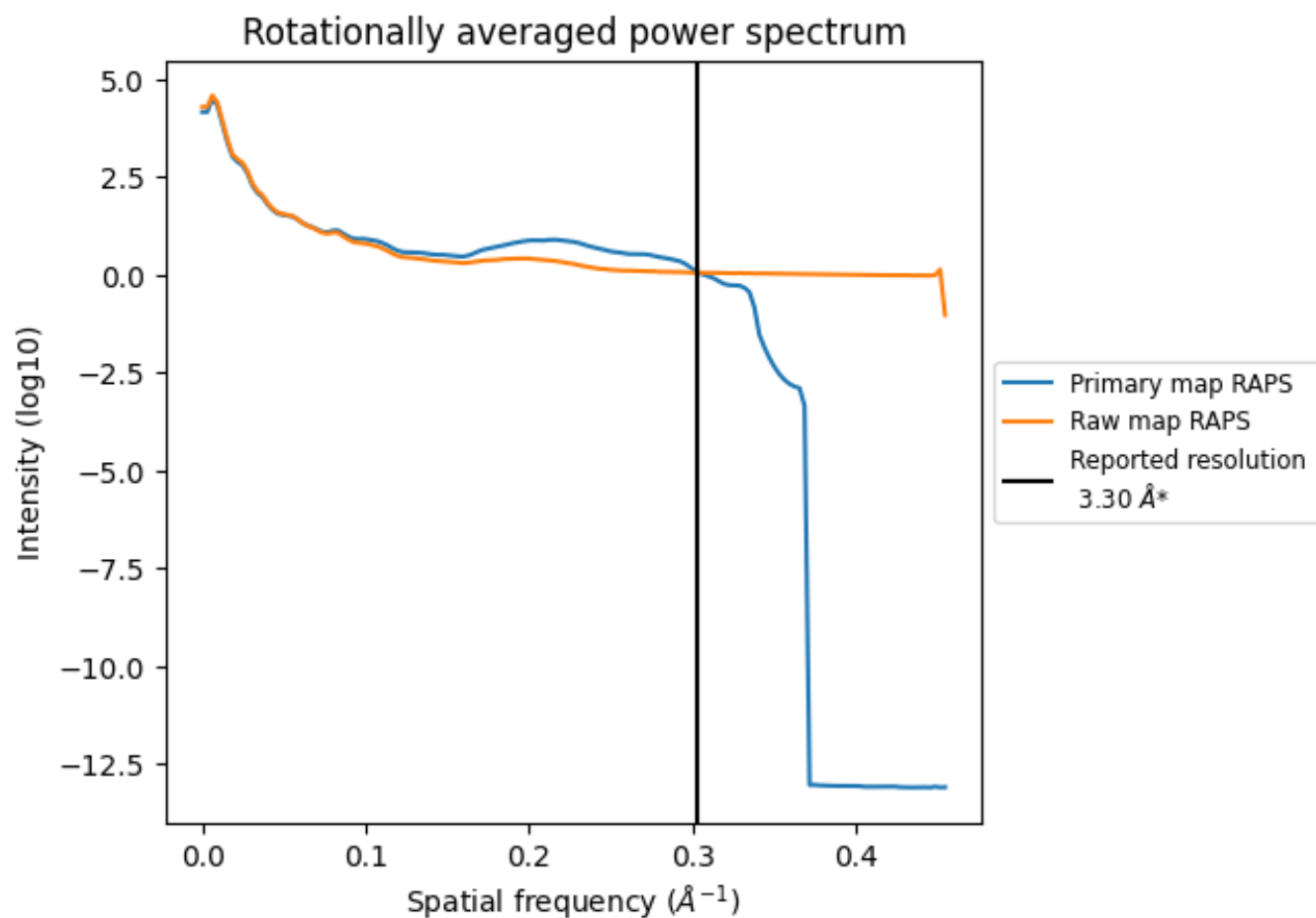
## 7.2 Volume estimate [i](#)



The volume at the recommended contour level is 117 nm<sup>3</sup>; this corresponds to an approximate mass of 106 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](#)

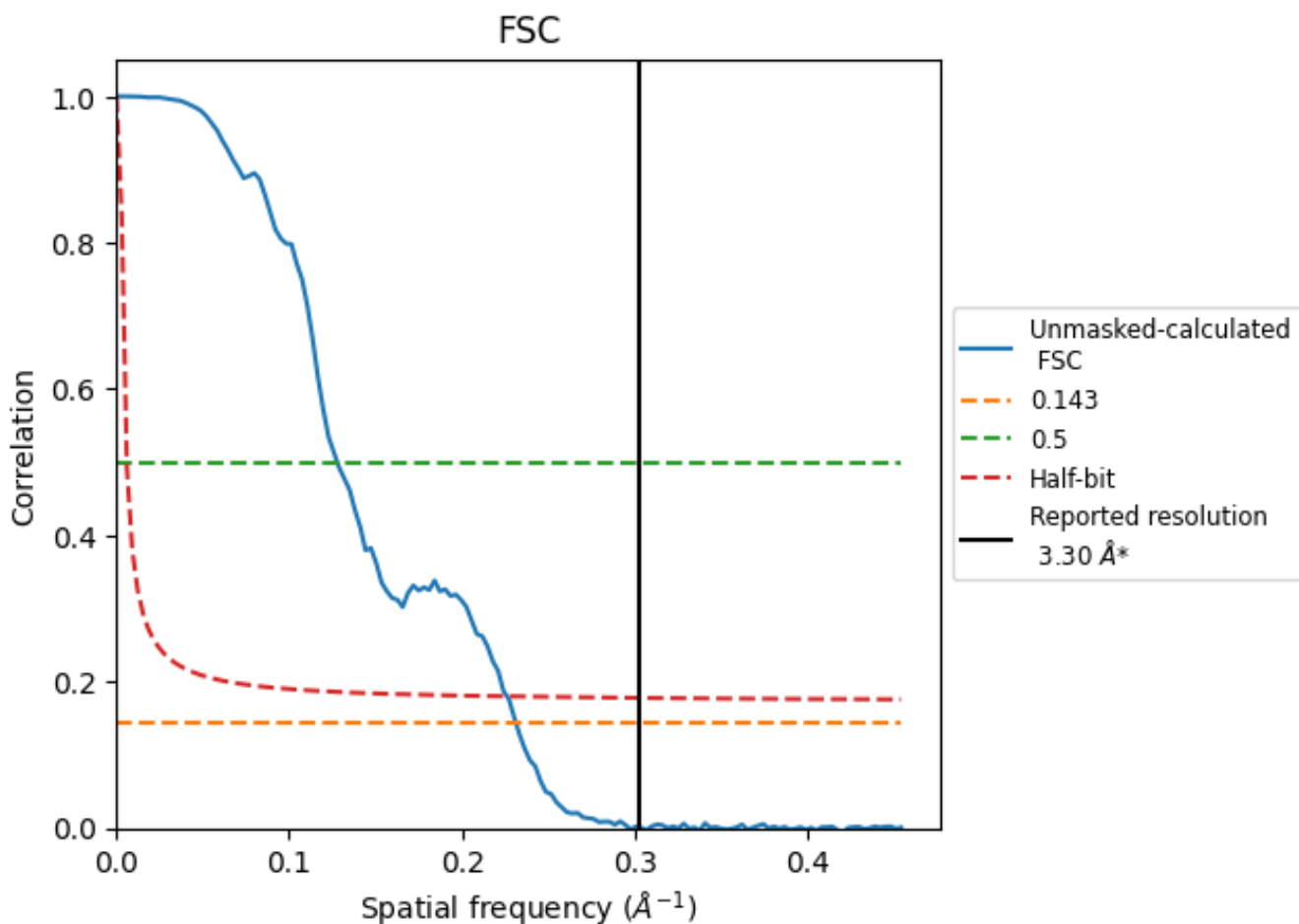


\*Reported resolution corresponds to spatial frequency of 0.303 Å<sup>-1</sup>

## 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 0.303 Å<sup>-1</sup>

## 8.2 Resolution estimates [i](#)

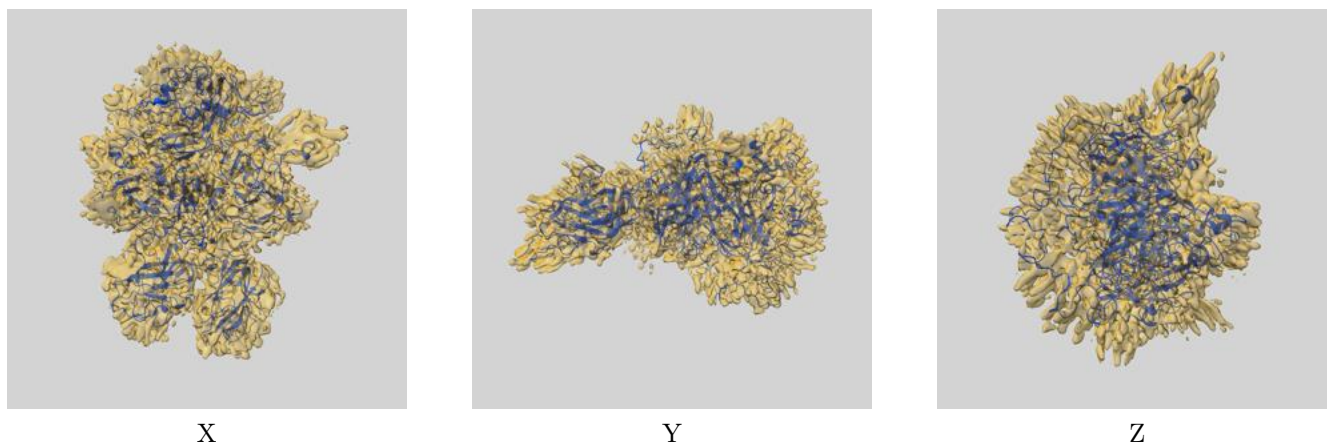
Resolution estimate (Å)	Estimation criterion (FSC cut-off)		
	0.143	0.5	Half-bit
Reported by author	3.30	-	-
Author-provided FSC curve	-	-	-
Unmasked-calculated*	4.32	7.82	4.42

\*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 4.32 differs from the reported value 3.3 by more than 10 %

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

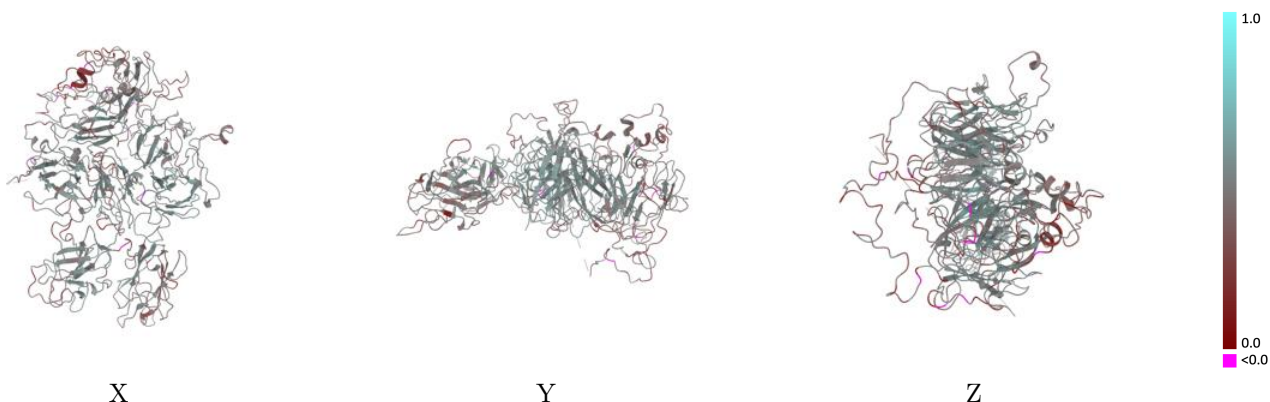
This section contains information regarding the fit between EMDB map EMD-23048 and PDB model 7KVE. Per-residue inclusion information can be found in section 3 on page 4.

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



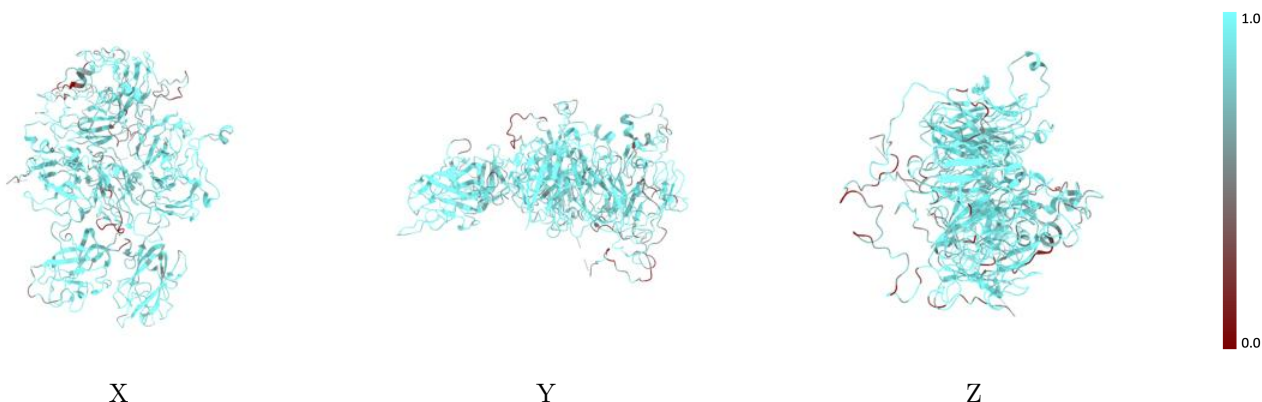
The images above show the 3D surface view of the map at the recommended contour level 0.43 at 50% transparency in yellow overlaid with a ribbon representation of the model coloured in blue. These images allow for the visual assessment of the quality of fit between the atomic model and the map.

## 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.

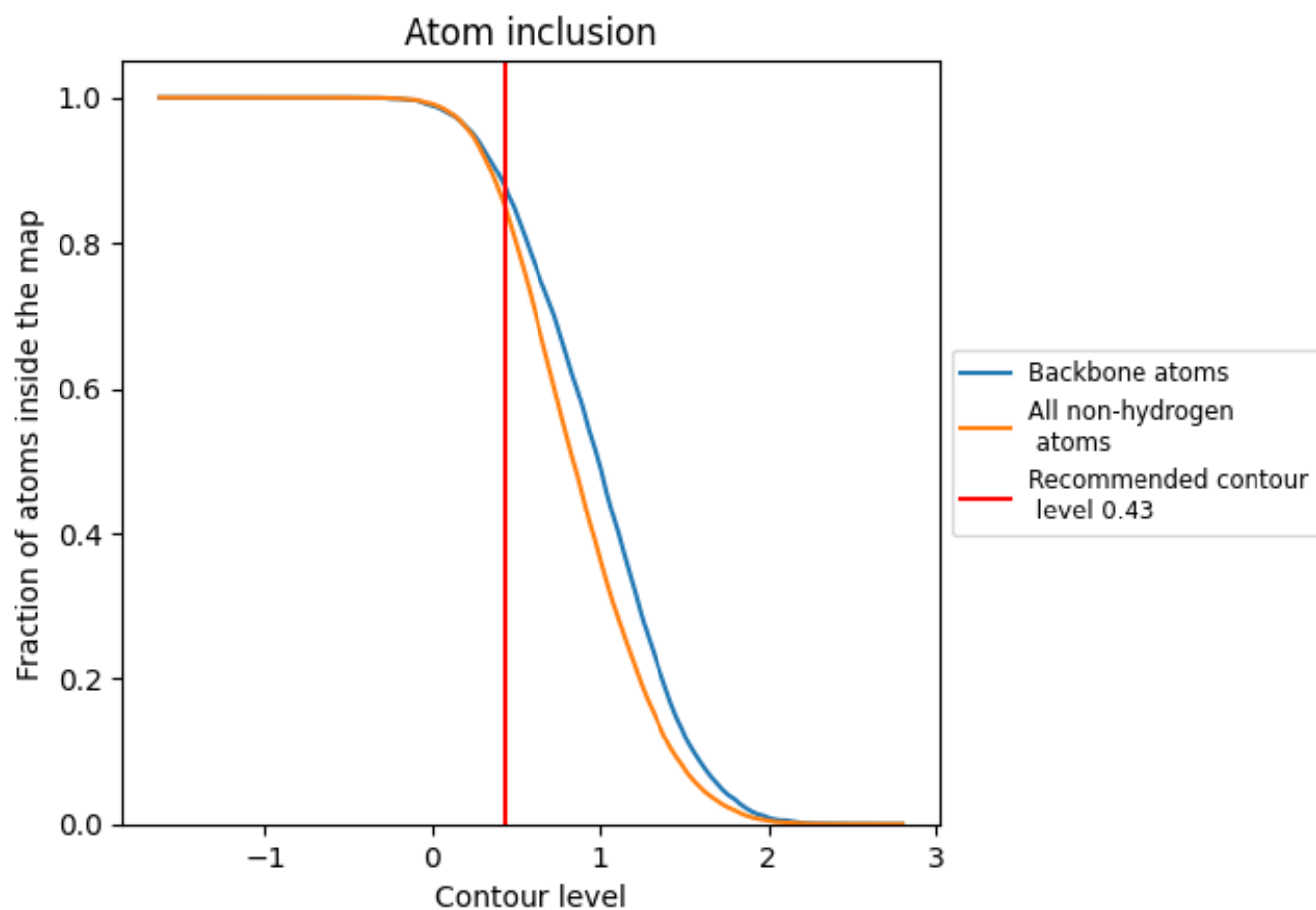
## 9.3 Atom inclusion mapped to coordinate model [i](#)



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 (0.43).




## 9.4 Atom inclusion [i](#)



At the recommended contour level, 88% of all backbone atoms, 85% 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 (0.43) and Q-score for the entire model and for each chain.

Chain	Atom inclusion	Q-score
All	 0.8512	 0.4510
B	 0.8512	 0.4510

