



wwPDB EM Validation Summary Report ⓘ

Nov 4, 2024 – 05:45 AM EST

PDB ID : 7TZC
EMDB ID : EMD-26205
Title : A drug and ATP binding site in type 1 ryanodine receptor
Authors : Melville, Z.; Dridi, H.; Yuan, Q.; Reiken, S.; Anetta, W.; Liu, Y.; Clarke, O.B.; Marks, A.R.
Deposited on : 2022-02-15
Resolution : 2.45 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
Mogul : 2022.3.0, CSD as543be (2022)
MolProbity : 4.02b-467
buster-report : 1.1.7 (2018)
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

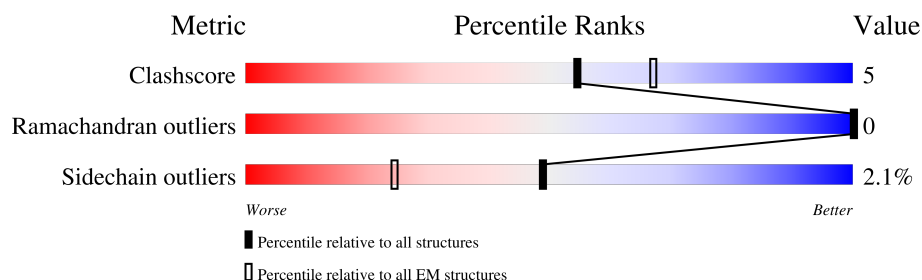
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

ELECTRON MICROSCOPY

The reported resolution of this entry is 2.45 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	C	150	<div> <div>33%</div> <div>77%</div> <div>22%</div> <div>..</div> </div>
1	D	150	<div> <div>30%</div> <div>79%</div> <div>20%</div> <div>..</div> </div>
1	E	150	<div> <div>33%</div> <div>79%</div> <div>20%</div> <div>..</div> </div>
1	K	150	<div> <div>33%</div> <div>78%</div> <div>20%</div> <div>..</div> </div>
2	F	107	<div> <div>86%</div> <div>14%</div> <div>.</div> </div>
2	H	107	<div> <div>85%</div> <div>15%</div> <div>.</div> </div>
2	J	107	<div> <div>88%</div> <div>11%</div> <div>.</div> </div>
2	O	107	<div> <div>86%</div> <div>13%</div> <div>.</div> </div>

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Mol	Chain	Length	Quality of chain
3	A	5037	<div><div></div><div>11%</div><div>75%</div><div>12%</div><div>13%</div></div>
3	B	5037	<div><div></div><div>11%</div><div>75%</div><div>12%</div><div>13%</div></div>
3	G	5037	<div><div></div><div>11%</div><div>75%</div><div>12%</div><div>13%</div></div>
3	I	5037	<div><div></div><div>11%</div><div>75%</div><div>12%</div><div>13%</div></div>

2 Entry composition

There are 9 unique types of molecules in this entry. The entry contains 149472 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 Calmodulin-1.

Mol	Chain	Residues	Atoms					AltConf	Trace
1	K	149	Total	C	N	O	S	0	0
			1174	719	190	255	10		
1	D	149	Total	C	N	O	S	0	0
			1174	719	190	255	10		
1	E	149	Total	C	N	O	S	0	0
			1174	719	190	255	10		
1	C	149	Total	C	N	O	S	0	0
			1174	719	190	255	10		

There are 4 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
K	-1	HIS	-	expression tag	UNP P0DP23
D	-1	HIS	-	expression tag	UNP P0DP23
E	-1	HIS	-	expression tag	UNP P0DP23
C	-1	HIS	-	expression tag	UNP P0DP23

- Molecule 2 is a protein called Peptidyl-prolyl cis-trans isomerase FKBP1A.

Mol	Chain	Residues	Atoms					AltConf	Trace
2	F	107	Total	C	N	O	S	0	0
			831	527	146	154	4		
2	H	107	Total	C	N	O	S	0	0
			831	527	146	154	4		
2	J	107	Total	C	N	O	S	0	0
			831	527	146	154	4		
2	O	107	Total	C	N	O	S	0	0
			831	527	146	154	4		

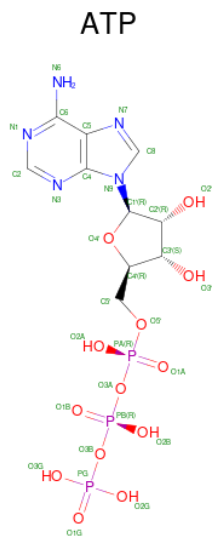
- Molecule 3 is a protein called Ryanodine receptor 1.

Mol	Chain	Residues	Atoms					AltConf	Trace
3	A	4404	Total	C	N	O	S	9	0
			35150	22365	6063	6485	237		
3	B	4404	Total	C	N	O	S	9	0
			35150	22365	6063	6485	237		
3	G	4404	Total	C	N	O	S	9	0
			35150	22365	6063	6485	237		
3	I	4404	Total	C	N	O	S	9	0
			35150	22365	6063	6485	237		

- Molecule 4 is CALCIUM ION (three-letter code: CA) (formula: Ca).

Mol	Chain	Residues	Atoms		AltConf
4	K	4	Total	Ca	0
			4	4	
4	D	4	Total	Ca	0
			4	4	
4	E	4	Total	Ca	0
			4	4	
4	C	4	Total	Ca	0
			4	4	
4	A	1	Total	Ca	0
			1	1	
4	B	1	Total	Ca	0
			1	1	
4	G	1	Total	Ca	0
			1	1	
4	I	1	Total	Ca	0
			1	1	

- Molecule 5 is ADENOSINE-5'-TRIPHOSPHATE (three-letter code: ATP) (formula: C₁₀H₁₆N₅O₁₃P₃) (labeled as "Ligand of Interest" by depositor).

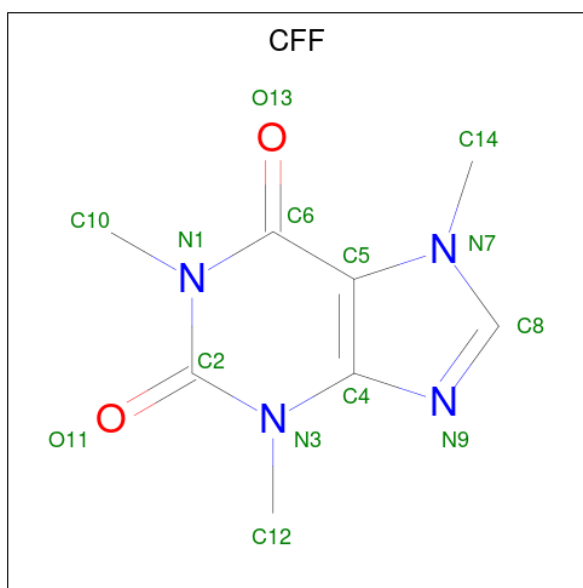


Mol	Chain	Residues	Atoms					AltConf
5	A	1	Total 31	C 10	N 5	O 13	P 3	0
5	A	1	Total 31	C 10	N 5	O 13	P 3	0
5	B	1	Total 31	C 10	N 5	O 13	P 3	0
5	B	1	Total 31	C 10	N 5	O 13	P 3	0
5	G	1	Total 31	C 10	N 5	O 13	P 3	0
5	G	1	Total 31	C 10	N 5	O 13	P 3	0
5	I	1	Total 31	C 10	N 5	O 13	P 3	0
5	I	1	Total 31	C 10	N 5	O 13	P 3	0

- Molecule 6 is ZINC ION (three-letter code: ZN) (formula: Zn).

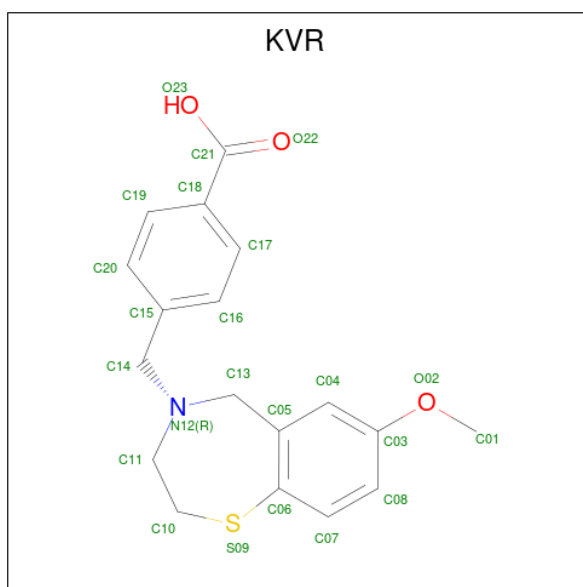
Mol	Chain	Residues	Atoms		AltConf
6	A	1	Total 1	Zn 1	0
6	B	1	Total 1	Zn 1	0
6	G	1	Total 1	Zn 1	0
6	I	1	Total 1	Zn 1	0

- Molecule 7 is CAFFEINE (three-letter code: CFF) (formula: $C_8H_{10}N_4O_2$).



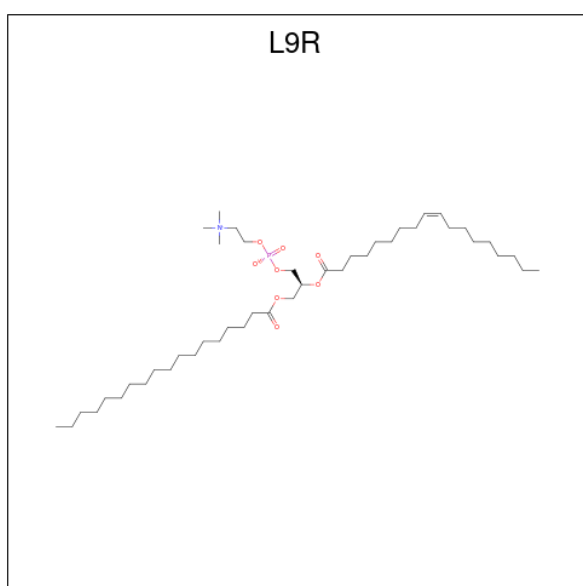
Mol	Chain	Residues	Atoms				AltConf
7	A	1	Total	C	N	O	0
			14	8	4	2	
7	B	1	Total	C	N	O	0
			14	8	4	2	
7	G	1	Total	C	N	O	0
			14	8	4	2	
7	I	1	Total	C	N	O	0
			14	8	4	2	

- Molecule 8 is 4-[(7-methoxy-2,3-dihydro-1,4-benzothiazepin-4(5H)-yl)methyl]benzoic acid (three-letter code: KVR) (formula: $C_{18}H_{19}NO_3S$).



Mol	Chain	Residues	Atoms					AltConf
8	A	1	Total	C	N	O	S	0
			23	18	1	3	1	
8	B	1	Total	C	N	O	S	0
			23	18	1	3	1	
8	G	1	Total	C	N	O	S	0
			23	18	1	3	1	
8	I	1	Total	C	N	O	S	0
			23	18	1	3	1	

- Molecule 9 is (2S)-3-(octadecanoyloxy)-2-[(9Z)-octadec-9-enoyloxy]propyl 2-(trimethylamm onio)ethyl phosphate (three-letter code: L9R) (formula: C₄₄H₈₆NO₈P).

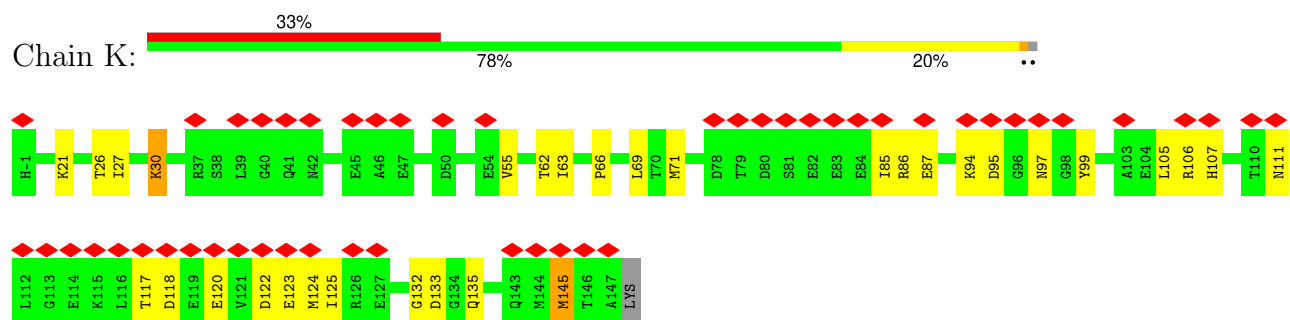


Mol	Chain	Residues	Atoms					AltConf
9	A	1	Total	C	N	O	P	0
			54	44	1	8	1	
9	A	1	Total	C	N	O	P	0
			54	44	1	8	1	
9	B	1	Total	C	N	O	P	0
			54	44	1	8	1	
9	B	1	Total	C	N	O	P	0
			54	44	1	8	1	
9	G	1	Total	C	N	O	P	0
			54	44	1	8	1	
9	G	1	Total	C	N	O	P	0
			54	44	1	8	1	
9	I	1	Total	C	N	O	P	0
			54	44	1	8	1	
9	I	1	Total	C	N	O	P	0
			54	44	1	8	1	

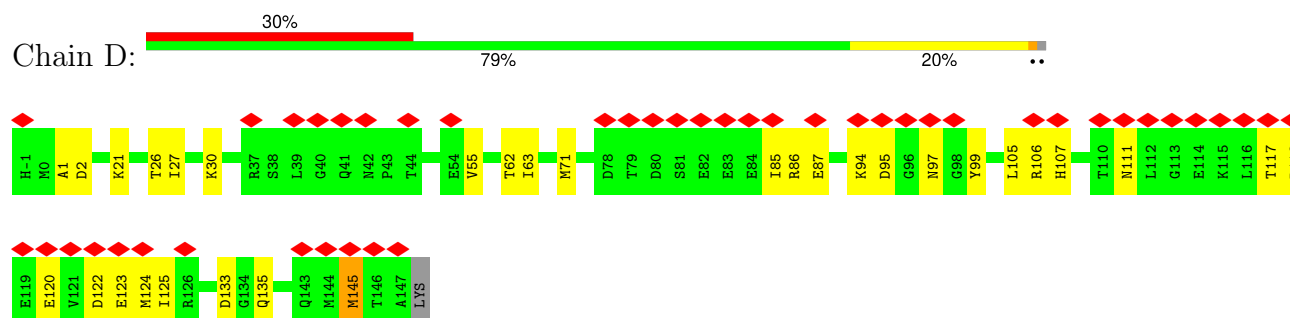
3 Residue-property plots

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

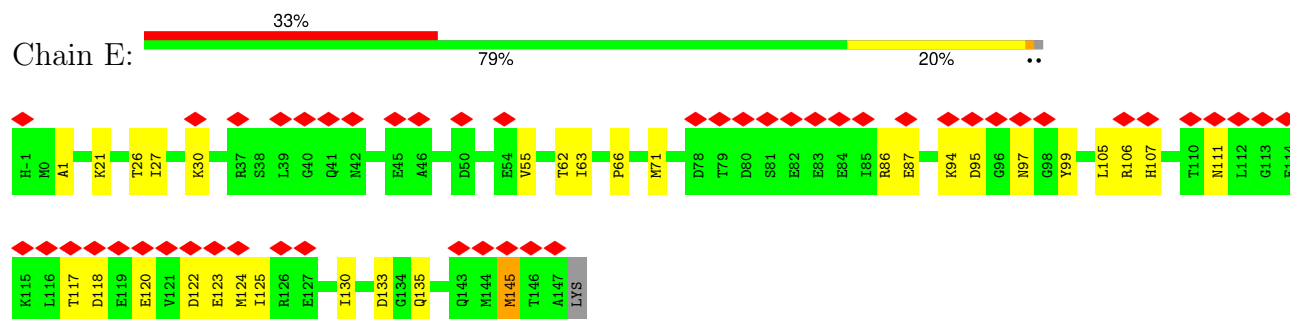
• Molecule 1: Calmodulin-1



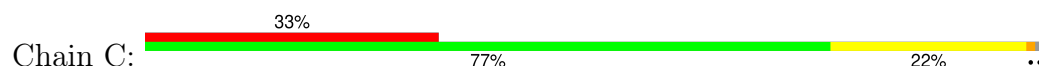
• Molecule 1: Calmodulin-1

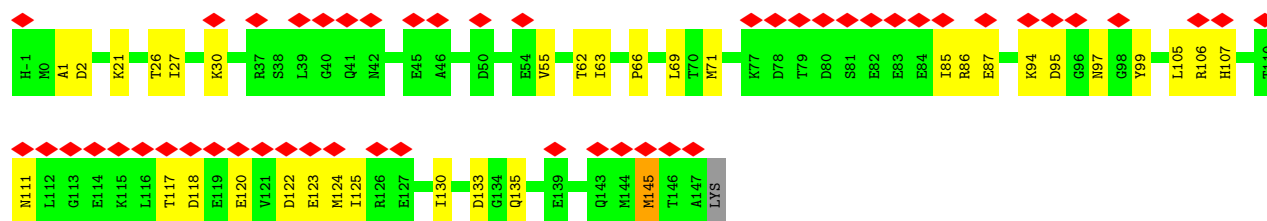


• Molecule 1: Calmodulin-1

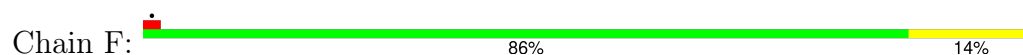


• Molecule 1: Calmodulin-1

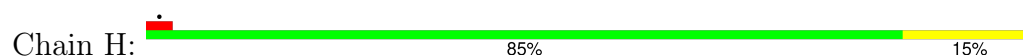




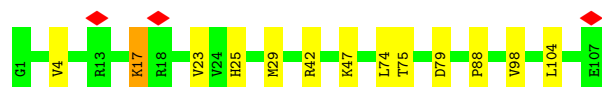
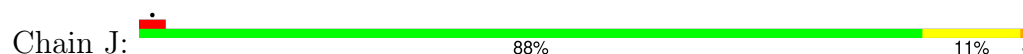
- Molecule 2: Peptidyl-prolyl cis-trans isomerase FKBP1A



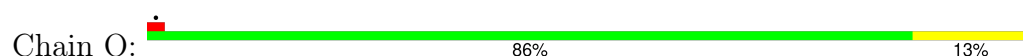
- Molecule 2: Peptidyl-prolyl cis-trans isomerase FKBP1A



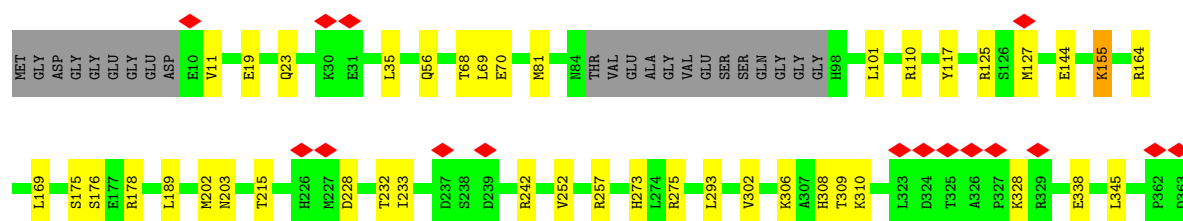
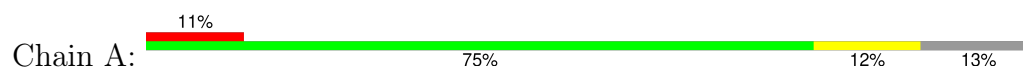
- Molecule 2: Peptidyl-prolyl cis-trans isomerase FKBP1A

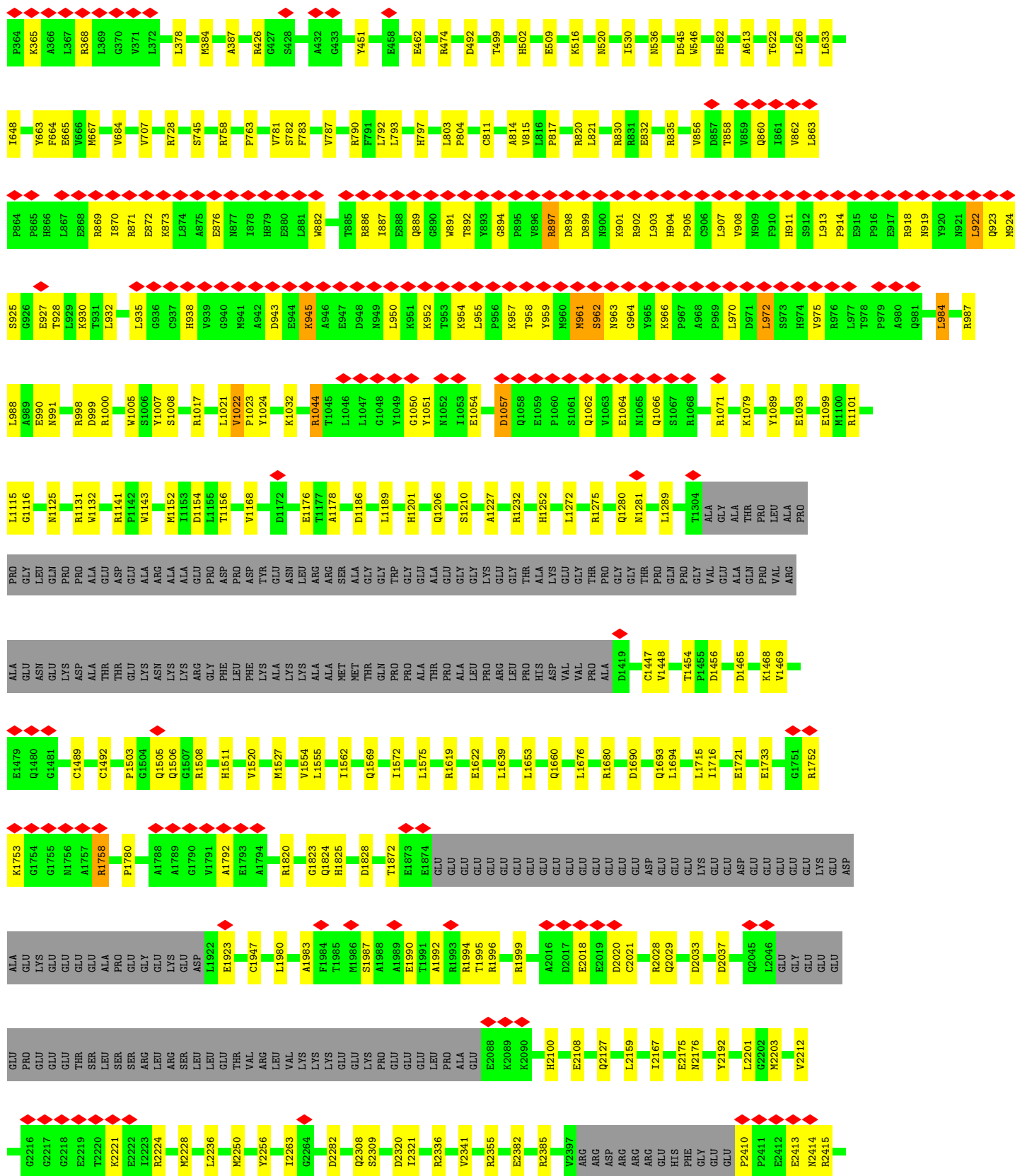


- Molecule 2: Peptidyl-prolyl cis-trans isomerase FKBP1A



- Molecule 3: Ryanodine receptor 1



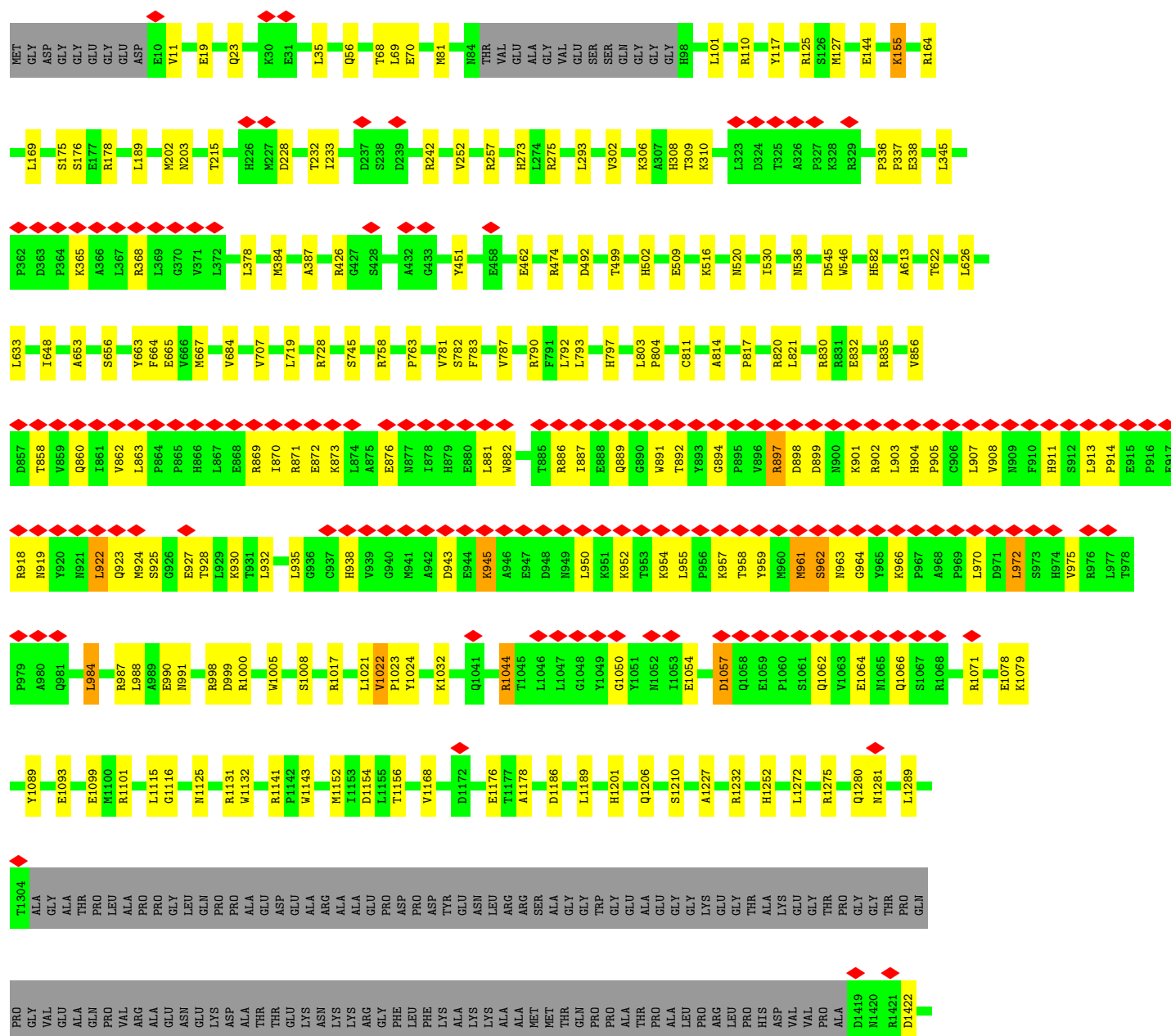
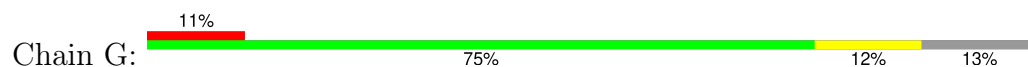


GLU	M4000	K3760	R3628	LYS	E3388	M3234	F5017	V2906	THR	L2785	K2786	L2785	K2725	E2449
ALA	M4001	Q3761	R3629	R3498	E3389	S3235	F5017	P2907	ALA	K2786	K2787	L2786	LYS	R2452
GLU	K4002	R3762	E3630	R3499	G3390	V3236	T3020	Y2908	THR	T2787	H2788	L2787	THR	
ASP	D4006	S3768	E3631	G3500	E3391	E3238	K3023	D2909	TVR	H2789	P2789	L2788	VAL	R2481
GLY	Q4020	H3771	V3633	D3501	L3393	L3249	K3036	T2912	PRO	M2790	P2790	L2789	ASP	D2482
MET	K4021	T3772	V3632	V3394	V3394	Y3263	R3053	A2913	ARG	L2791	L2791	L2790	ALA	G2483
GLY	D4022	R3773	R3637	R3395	R3395	I3270	V3107	K2914	GLU	R2792	R2792	L2791	GLY	V2495
ALA	D4063	L3805	M3638	L3514	V3400	V3269	R3053	E2915	GLY	N2794	F2735	N2794		H2498
ALA	M4064	Q3813	T3639	N3523	R3403	I3270	V3107	K2916		D2736	D2736	P2737		
GLU	S4074	L3842	A3680	L3535	L3412	T3273	L3112	A2917		P2793	P2793	P2737		
GLY	K4090	E3682	E3682	T3538	T3413	L3274	G3113	R2918		K2795	K2795	P2737		E2513
GLU	Q3850	Q3683	Q3683	T3538	R3414	L3277	G3113	D2919		T2796	T2796	R2738		N2514
GLY	Q3851	E3684	E3684	T3545	L3434	L3277	V3114	R2920		P2797	P2797	R2738		Q2515
GLY	K3852	E3685	E3685	E3548	L3434	L3277	S3116	R2921		S2798	S2798	P2739		D2516
ALA	Q3857	E3686	E3686	E3548	F3442	M3284	S3116	E2921		E2740	E2740	V2740		L2519
GLY	M3858	E3687	E3687	F3552	W3445	G3288	GLN	K2922		E2741	E2741	E2741		T2538
ALA	V3859	E3688	E3688	F3552	W3445	G3288	ALA	K2923		T2742	T2742	T2742		
GLY	N3860	E3689	E3689	N3556	S3446	P3294	THR	Q2924		L2743	L2743	L2743		F2541
GLY	E3861	V3690	F3451	L3559	F3451	P3294	GLN	E2925		N2744	N2744	N2744		
ALA	D3862	E3691	K3452	L3559	K3452	A3295	VAL	L2926		V2745	V2745	V2745		R2576
GLY	Q3863	E3692	E3692	E3564	E3455	L3296	K3123	L2927		I2746	I2746	I2746		
THR	T3864	K3693	E3693	L3579	Q3456	P3297	G3124	K2928		I2747	I2747	I2747		D2580
ALA	V3865	K3694	K3694	L3579	Q3456	A3298	V3125	F2929		P2748	P2748	P2748		Y2587
GLY	L3866	P3695	P3695	R3582	V3459	G3299	T3132	L2930		E2749	E2749	E2749		
ALA	N3867	D3696	D3696	R3582	V3460	G3299	T3133	L2931		K2750	K2750	K2750		S2590
THR	Q3870	L3710	L3710	E3583	I3464	A3300	K3136	M2932		L2751	L2751	L2751		H2621
ALA	E3872	D3717	D3717	E3583	A3472	P3303	L3136	N2933		S2752	S2752	S2752		L2626
ALA	K3873	K3731	K3731	E3583	D3473	G3304	L3136	N2933		F2754	F2754	F2754		L2633
ALA	V3874	H3734	H3734	E3583	S3474	P3304	L3136	N2933		I2755	I2755	I2755		
ALA	L3890	L3735	L3735	E3583	K3475	V3324	L3136	N2933		N2756	N2756	N2756		P2640
ALA	F3899	E3736	E3736	E3583	S3476	I3329	L3136	N2933		K2757	K2757	K2757		L2644
ALA	S3929	Q3608	Q3608	E3583	K3477	A3332	L3136	N2933		F2758	F2758	F2758		
ALA	F3933	Q3609	Q3609	E3583	H3478	M3335	L3136	N2933		E2759	E2759	E2759		R2650
ALA	K3940	Q3610	Q3610	E3583	L3479	L3332	L3136	N2933		A2761	A2761	A2761		C2651
ALA	E3944	Q3611	Q3611	E3583	ALA	F3341	L3136	N2933		T2762	T2762	T2762		W2661
ALA	Q3945	Q3612	Q3612	E3583	GLY	A3342	L3136	N2933		E2763	E2763	E2763		F2664
ALA	Q3946	Q3613	Q3613	E3583	ASP	Q3343	L3136	N2933		K2765	K2765	K2765		K2677
ALA	P4254	Q3614	Q3614	E3583	GLN	P3344	L3136	N2933		W2766	W2766	W2766		L2678
VAL	GLY	Q3615	Q3615	E3583	SER	V3346	L3136	N2933		A2767	A2767	A2767		K2689
GLY	GLY	Q3616	Q3616	E3583	GLY	V3346	L3136	N2933		D2769	D2769	D2769		D2716
ARG	ARG	Q3617	Q3617	E3583	GLY	V3346	L3136	N2933		K2770	K2770	K2770		S2720
ARG	ARG	Q3618	Q3618	E3583	SER	T3361	L3136	N2933		I2771	I2771	I2771		S2721
ARG	ARG	Q3619	Q3619	E3583	ASP	E3382	L3136	N2933		Q2772	Q2772	Q2772		K2722
VAL	VAL	Q3620	Q3620	E3583	GLN	A3383	L3136	N2933		N2773	N2773	N2773		A2723
ARG	ARG	Q3621	Q3621	E3583	ARG	K3384	L3136	N2933		N2774	N2774	N2774		E2724
ARG	ARG	Q3622	Q3622	E3583	THR	K3385	L3136	N2933		W2775	W2775	W2775		
ARG	ARG	Q3623	Q3623	E3583	LYS	E3386	L3136	N2933		S2776	S2776	S2776		
ARG	ARG	Q3624	Q3624	E3583	LYS	E3387	L3136	N2933		Y2777	Y2777	Y2777		
ARG	ARG	Q3625	Q3625	E3583	LYS	A3387	L3136	N2933		G2778	G2778	G2778		
ARG	ARG	Q3626	Q3626	E3583	LYS	A3387	L3136	N2933		E2779	E2779	E2779		
ARG	ARG	Q3627	Q3627	E3583	LYS	A3387	L3136	N2933		N2780	N2780	N2780		
ARG	ARG	Q3628	Q3628	E3583	LYS	A3387	L3136	N2933		V2781	V2781	V2781		
ARG	ARG	Q3629	Q3629	E3583	LYS	A3387	L3136	N2933		D2782	D2782	D2782		
ARG	ARG	Q3630	Q3630	E3583	LYS	A3387	L3136	N2933		E2784	E2784	E2784		
ARG	ARG	Q3631	Q3631	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3632	Q3632	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3633	Q3633	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3634	Q3634	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3635	Q3635	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3636	Q3636	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3637	Q3637	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3638	Q3638	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3639	Q3639	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3640	Q3640	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3641	Q3641	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3642	Q3642	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3643	Q3643	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3644	Q3644	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3645	Q3645	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3646	Q3646	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3647	Q3647	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3648	Q3648	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3649	Q3649	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3650	Q3650	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3651	Q3651	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3652	Q3652	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3653	Q3653	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3654	Q3654	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3655	Q3655	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3656	Q3656	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3657	Q3657	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3658	Q3658	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3659	Q3659	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3660	Q3660	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3661	Q3661	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3662	Q3662	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3663	Q3663	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3664	Q3664	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3665	Q3665	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3666	Q3666	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3667	Q3667	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3668	Q3668	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3669	Q3669	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3670	Q3670	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3671	Q3671	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3672	Q3672	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3673	Q3673	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3674	Q3674	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3675	Q3675	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3676	Q3676	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3677	Q3677	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3678	Q3678	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3679	Q3679	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3680	Q3680	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3681	Q3681	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3682	Q3682	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3683	Q3683	E3583	LYS	A3387	L3136	N2933						
ARG	ARG	Q3684	Q3684	E3583	LYS									



ALA	GLY	ARG	THR	GLU	MET	D4022	L3805	R3637	S3508	L3392	L3230	L3003	P2903
GLY	GLU	PRO	SER	ALA	GLY	K4060	Q3813	M3638	M3638	V3394	P3233	P3004	L2904
VAL	GLY	ALA	VAL	ALA	ALA	D4063	S3840	T3639	L3514	R3395	M3234	L2905	
PRO	GLY	GLY	HIS	LEU	ALA	M4064	Y3841	E3682	N3523	V3400	V3235	F3017	V2906
ALA	GLY	LEU	GLY	ALA	GLY	D4070	L3842	Q3683	L3535	R3403	E3237	T3020	Y2907
ALA	PRO	GLY	GLY	LEU	GLU	S4074	Q3850	E3684	L3538	R3414	E3238	T3020	Y2908
PRO	PRO	ASP	PRO	LEU	GLU	E4075	N3851	E3685	T3538	L3412	V3263	K3023	I2909
GLY	GLY	MET	ALA	TRP	GLY	S4074	K3852	E3686	T3545	R3414	Y3263	K3036	
PRO	GLY	GLY	GLY	ALA	GLY	F4093	G3857	E3687	E3548	L3445	V3269	K3036	T2912
PRO	VAL	ASP	PRO	VAL	ALA	M4097	M3858	E3688	E3548	L3445	I3270	R3053	A2913
THR	VAL	THR	GLY	VAL	ALA	M4097	V3859	E3688	F3552	F3442	T3273	K2914	K2914
THR	VAL	THR	GLY	VAL	ALA	E4116	N3860	E3689	L3556	L3442	L3274	E2915	E2915
ALA	ALA	ALA	ALA	ALA	ALA	E4119	E3861	E3690	L3579	F3442	L3274	K2916	K2916
PRO	GLY	PRO	GLY	GLY	ALA	R4137	G3863	E3691	L3556	F3442	T3273	A2917	A2917
PRO	GLY	PRO	GLY	GLY	ALA	D4138	V3865	E3692	L3556	F3442	L3274	R2918	R2918
ALA	ALA	ALA	ALA	ALA	ALA	M4142	I3866	E3693	L3556	F3442	L3274	I2919	I2919
GLY	GLY	GLY	GLY	GLY	ALA	E4152	G3871	E3694	L3556	F3442	L3274	K3114	K3114
GLY	GLY	GLY	GLY	GLY	ALA	E4172	E3872	E3695	L3556	F3442	L3274	K3115	K3115
LEU	LEU	LEU	GLY	LEU	ALA	R4175	K3873	E3736	L3579	F3442	L3274	S3116	S3116
GLY	GLY	GLY	GLY	TRP	ALA	P4176	V3874	GLU	L3579	F3442	L3274	GLN	GLN
VAL	VAL	VAL	GLY	ASP	ALA	Y4177	GLY	GLY	L3579	F3442	L3274	ALA	ALA
ASP	ASP	ASP	GLY	SER	ALA	D3877	GLY	GLY	L3579	F3442	L3274	THR	THR
GLY	GLY	GLY	GLY	LEU	ALA	L3890	L3890	ASN	L3579	F3442	L3274	GLN	GLN
GLY	GLY	GLY	VAL	GLY	ALA	F4181	F3899	GLY	L3579	F3442	L3274	VAL	VAL
GLY	GLY	GLY	GLY	GLY	ALA	E4182	S3929	GLY	L3579	F3442	L3274	K3123	K3123
LEU	LEU	LEU	HIS	LEU	ALA	Y4194	S3929	ALA	L3579	F3442	L3274	G3124	G3124
VAL	VAL	VAL	ALA	VAL	ALA	E4199	F3933	GLY	L3579	F3442	L3274	V3125	V3125
PRO	PRO	PRO	GLY	GLY	TYR	E4199	F3933	GLY	L3579	F3442	L3274	T3132	T3132
GLY	GLY	GLY	GLY	GLY	GLY	E4224	K3940	V3749	L3579	F3442	L3274	T3133	T3133
GLY	GLY	GLY	GLY	GLY	GLY	G4225	D3941	E3750	L3579	F3442	L3274	V3134	V3134
PRO	PRO	PRO	GLY	VAL	ARG	G4226	D3941	E3750	L3579	F3442	L3274	A3135	A3135
PRO	PRO	PRO	ALA	VAL	ARG	E4227	E3944	V3751	L3579	F3442	L3274	L3136	L3136
GLY	GLY	GLY	THR	THR	ARG	E4227	E3944	S3752	L3579	F3442	L3274	L3137	L3137
GLY	GLY	GLY	GLY	THR	ARG	P4254	E3945	V3619	L3579	F3442	L3274	Y2935	Y2935
PRO	PRO	PRO	VAL	VAL	VAL	P4254	Q3946	ASP	L3579	F3442	L3274	A2936	A2936
GLY	GLY	GLY	VAL	VAL	ARG	GLY	Q3946	ASP	L3579	F3442	L3274	V2937	V2937
GLY	GLY	GLY	VAL	VAL	ARG	GLY	R3949	E3757	L3579	F3442	L3274	T2938	T2938
GLY	GLY	GLY	VAL	VAL	LEU	GLY	K3959	M3758	L3579	F3442	L3274	L3157	L3157
ALA	ALA	ALA	ALA	ALA	ARG	PRO	K3959	K3760	L3579	F3442	L3274	L3158	L3158
ASP	ASP	ASP	ASP	GLY	ARG	GLY	M4000	Q3761	L3579	F3442	L3274	D3159	D3159
ASP	ASP	ASP	GLY	MET	LEU	GLY	M4000	R3762	L3579	F3442	L3274	D3160	D3160
GLY	GLY	GLY	GLY	PRO	THR	ALA	M4001	Q3761	L3579	F3442	L3274	V3161	V3161
GLY	GLY	GLY	GLY	PRO	ALA	ASP	K4002	R3762	L3579	F3442	L3274	L3169	L3169
GLY	GLY	GLY	GLY	ASP	ALA	GLY	K4002	S3768	L3579	F3442	L3274	ASP	ASP
ASN	ASN	ASN	PHE	PRO	ARG	ASP	D4006	H3771	L3579	F3442	L3274	MET	MET
				GLY	GLY	GLY	D4006	T3772	L3579	F3442	L3274	GLU	GLU
				GLY	GLY	GLY	Q4020	R3773	L3579	F3442	L3274	L2946	L2946
							K4021		L3579	F3442	L3274	I2947	I2947
									L3579	F3442	L3274	L3194	L3194
									L3579	F3442	L3274	S2370	S2370
									L3579	F3442	L3274	F2973	F2973
									L3579	F3442	L3274	I2974	I2974
									L3579	F3442	L3274	A2975	A2975
									L3579	F3442	L3274	R3225	R3225
									L3579	F3442	L3274	E2978	E2978
									L3579	F3442	L3274	S2982	S2982
									L3579	F3442	L3274	S2983	S2983
									L3579	F3442	L3274	Q2984	Q2984
									L3579	F3442	L3274	R2985	R2985
									L3579	F3442	L3274	V2986	V2986
									L3579	F3442	L3274	E2987	E2987
									L3579	F3442	L3274	K2988	K2988

- Molecule 3: Ryanodine receptor 1











4 Experimental information

Property	Value	Source
EM reconstruction method	SINGLE PARTICLE	Depositor
Imposed symmetry	POINT, C4	Depositor
Number of particles used	153840	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 ($e^-/\text{\AA}^2$)	57.65	Depositor
Minimum defocus (nm)	1000	Depositor
Maximum defocus (nm)	2000	Depositor
Magnification	105000	Depositor
Image detector	GATAN K3 BIOQUANTUM (6k x 4k)	Depositor
Maximum map value	0.887	Depositor
Minimum map value	-0.013	Depositor
Average map value	0.013	Depositor
Map value standard deviation	0.034	Depositor
Recommended contour level	0.15	Depositor
Map size (Å)	426.496, 426.496, 426.496	wwPDB
Map dimensions	512, 512, 512	wwPDB
Map angles (°)	90.0, 90.0, 90.0	wwPDB
Pixel spacing (Å)	0.833, 0.833, 0.833	Depositor

5 Model quality [i](#)

5.1 Standard geometry [i](#)

Bond lengths and bond angles in the following residue types are not validated in this section: ZN, CFF, KVR, ATP, L9R, CA

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	C	0.24	0/1187	0.44	0/1594
1	D	0.24	0/1187	0.44	0/1594
1	E	0.24	0/1187	0.44	0/1594
1	K	0.24	0/1187	0.44	0/1594
2	F	0.32	0/850	0.52	0/1146
2	H	0.32	0/850	0.52	0/1146
2	J	0.32	0/850	0.52	0/1146
2	O	0.32	0/850	0.52	0/1146
3	A	0.25	0/35977	0.46	0/48726
3	B	0.25	0/35977	0.46	0/48726
3	G	0.25	0/35977	0.46	0/48726
3	I	0.25	0/35977	0.46	0/48726
All	All	0.25	0/152056	0.46	0/205864

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	C	1174	0	1099	20	0

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Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
1	D	1174	0	1099	17	0
1	E	1174	0	1099	17	0
1	K	1174	0	1099	19	0
2	F	831	0	831	7	0
2	H	831	0	831	8	0
2	J	831	0	831	7	0
2	O	831	0	831	8	0
3	A	35150	0	34797	347	0
3	B	35150	0	34797	347	0
3	G	35150	0	34797	347	0
3	I	35150	0	34797	353	0
4	A	1	0	0	0	0
4	B	1	0	0	0	0
4	C	4	0	0	0	0
4	D	4	0	0	0	0
4	E	4	0	0	0	0
4	G	1	0	0	0	0
4	I	1	0	0	0	0
4	K	4	0	0	0	0
5	A	62	0	24	2	0
5	B	62	0	24	2	0
5	G	62	0	24	2	0
5	I	62	0	24	2	0
6	A	1	0	0	0	0
6	B	1	0	0	0	0
6	G	1	0	0	0	0
6	I	1	0	0	0	0
7	A	14	0	10	0	0
7	B	14	0	10	0	0
7	G	14	0	10	0	0
7	I	14	0	10	0	0
8	A	23	0	0	0	0
8	B	23	0	0	0	0
8	G	23	0	0	0	0
8	I	23	0	0	0	0
9	A	108	0	172	9	0
9	B	108	0	172	6	0
9	G	108	0	172	9	0
9	I	108	0	172	8	0
All	All	149472	0	147732	1461	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 5.

The worst 5 of 1461 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)
3:I:4904:PRO:HB3	3:I:4913:ARG:HG2	1.57	0.86
3:G:4904:PRO:HB3	3:G:4913:ARG:HG2	1.57	0.86
3:A:4904:PRO:HB3	3:A:4913:ARG:HG2	1.57	0.84
3:B:4904:PRO:HB3	3:B:4913:ARG:HG2	1.57	0.83
3:A:2779:GLU:HG3	3:A:2792:ARG:HG2	1.66	0.78

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	C	147/150 (98%)	146 (99%)	1 (1%)	0	100	100
1	D	147/150 (98%)	146 (99%)	1 (1%)	0	100	100
1	E	147/150 (98%)	146 (99%)	1 (1%)	0	100	100
1	K	147/150 (98%)	146 (99%)	1 (1%)	0	100	100
2	F	105/107 (98%)	103 (98%)	2 (2%)	0	100	100
2	H	105/107 (98%)	103 (98%)	2 (2%)	0	100	100
2	J	105/107 (98%)	103 (98%)	2 (2%)	0	100	100
2	O	105/107 (98%)	103 (98%)	2 (2%)	0	100	100
3	A	4385/5037 (87%)	4273 (97%)	112 (3%)	0	100	100
3	B	4385/5037 (87%)	4273 (97%)	112 (3%)	0	100	100
3	G	4385/5037 (87%)	4274 (98%)	111 (2%)	0	100	100
3	I	4385/5037 (87%)	4273 (97%)	112 (3%)	0	100	100
All	All	18548/21176 (88%)	18089 (98%)	459 (2%)	0	100	100

There are no Ramachandran outliers to report.

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	C	127/128 (99%)	122 (96%)	5 (4%)	27	41
1	D	127/128 (99%)	122 (96%)	5 (4%)	27	41
1	E	127/128 (99%)	122 (96%)	5 (4%)	27	41
1	K	127/128 (99%)	122 (96%)	5 (4%)	27	41
2	F	89/89 (100%)	86 (97%)	3 (3%)	32	46
2	H	89/89 (100%)	86 (97%)	3 (3%)	32	46
2	J	89/89 (100%)	86 (97%)	3 (3%)	32	46
2	O	89/89 (100%)	86 (97%)	3 (3%)	32	46
3	A	3836/4276 (90%)	3760 (98%)	76 (2%)	50	65
3	B	3836/4276 (90%)	3760 (98%)	76 (2%)	50	65
3	G	3836/4276 (90%)	3760 (98%)	76 (2%)	50	65
3	I	3836/4276 (90%)	3760 (98%)	76 (2%)	50	65
All	All	16208/17972 (90%)	15872 (98%)	336 (2%)	49	64

5 of 336 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
3	G	3053	ARG
3	I	1758	ARG
3	G	3639	THR
3	I	897	ARG
3	I	2797	PHE

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 13 such sidechains are listed below:

Mol	Chain	Res	Type
3	B	3761	GLN
3	G	991	ASN
3	I	2881	ASN
3	I	991	ASN

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Mol	Chain	Res	Type
3	I	2180	GLN

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 oligosaccharides in this entry.

5.6 Ligand geometry [i](#)

Of 48 ligands modelled in this entry, 24 are monoatomic - leaving 24 for Mogul analysis.

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

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	# Z > 2	Counts	RMSZ	# Z > 2
7	CFF	G	5105	-	8,15,15	2.03	3 (37%)	8,23,23	1.45	1 (12%)
7	CFF	I	5105	-	8,15,15	2.03	3 (37%)	8,23,23	1.46	1 (12%)
8	KVR	B	5306	-	24,25,25	1.42	3 (12%)	31,34,34	1.55	4 (12%)
5	ATP	I	5102	-	28,33,33	0.63	0	34,52,52	0.85	2 (5%)
7	CFF	B	5304	-	8,15,15	2.02	3 (37%)	8,23,23	1.45	1 (12%)
7	CFF	A	5304	-	8,15,15	2.02	3 (37%)	8,23,23	1.45	1 (12%)
9	L9R	G	5108	-	53,53,53	1.24	5 (9%)	59,61,61	1.12	3 (5%)
9	L9R	B	5307	-	53,53,53	1.24	5 (9%)	59,61,61	1.12	3 (5%)
5	ATP	G	5102	-	28,33,33	0.61	0	34,52,52	0.85	2 (5%)
9	L9R	A	5307	-	53,53,53	1.24	5 (9%)	59,61,61	1.12	3 (5%)

Mol	Type	Chain	Res	Link	Bond lengths			Bond angles		
					Counts	RMSZ	# Z > 2	Counts	RMSZ	# Z > 2
5	ATP	B	5301	-	28,33,33	0.62	0	34,52,52	0.85	2 (5%)
8	KVR	G	5107	-	24,25,25	1.41	3 (12%)	31,34,34	1.56	4 (12%)
9	L9R	A	5308	-	53,53,53	1.21	4 (7%)	59,61,61	1.12	2 (3%)
8	KVR	A	5306	-	24,25,25	1.41	3 (12%)	31,34,34	1.55	4 (12%)
9	L9R	B	5308	-	53,53,53	1.21	4 (7%)	59,61,61	1.12	2 (3%)
5	ATP	G	5106	-	28,33,33	0.62	0	34,52,52	0.59	1 (2%)
5	ATP	I	5106	-	28,33,33	0.62	0	34,52,52	0.59	1 (2%)
5	ATP	B	5305	-	28,33,33	0.62	0	34,52,52	0.59	1 (2%)
9	L9R	I	5101	-	53,53,53	1.21	4 (7%)	59,61,61	1.12	2 (3%)
5	ATP	A	5305	-	28,33,33	0.62	0	34,52,52	0.59	1 (2%)
9	L9R	I	5108	-	53,53,53	1.24	5 (9%)	59,61,61	1.12	3 (5%)
5	ATP	A	5301	-	28,33,33	0.62	0	34,52,52	0.85	2 (5%)
8	KVR	I	5107	-	24,25,25	1.41	3 (12%)	31,34,34	1.55	4 (12%)
9	L9R	G	5101	-	53,53,53	1.21	4 (7%)	59,61,61	1.12	2 (3%)

In the following table, the Chirals column lists the number of chiral outliers, the number of chiral centers analysed, the number of these observed in the model and the number defined in the Chemical Component Dictionary. Similar counts are reported in the Torsion and Rings columns. '-' means no outliers of that kind were identified.

Mol	Type	Chain	Res	Link	Chirals	Torsions	Rings
7	CFF	G	5105	-	-	-	0/2/2/2
7	CFF	I	5105	-	-	-	0/2/2/2
8	KVR	B	5306	-	-	2/10/20/20	0/2/3/3
5	ATP	I	5102	-	-	8/18/38/38	0/3/3/3
7	CFF	B	5304	-	-	-	0/2/2/2
7	CFF	A	5304	-	-	-	0/2/2/2
9	L9R	G	5108	-	-	30/57/57/57	-
9	L9R	B	5307	-	-	30/57/57/57	-
5	ATP	G	5102	-	-	8/18/38/38	0/3/3/3
9	L9R	A	5307	-	-	30/57/57/57	-
5	ATP	B	5301	-	-	8/18/38/38	0/3/3/3
8	KVR	G	5107	-	-	2/10/20/20	0/2/3/3
9	L9R	A	5308	-	-	34/57/57/57	-
8	KVR	A	5306	-	-	2/10/20/20	0/2/3/3
9	L9R	B	5308	-	-	34/57/57/57	-
5	ATP	G	5106	-	-	6/18/38/38	0/3/3/3

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Mol	Type	Chain	Res	Link	Chirals	Torsions	Rings
5	ATP	I	5106	-	-	6/18/38/38	0/3/3/3
5	ATP	B	5305	-	-	6/18/38/38	0/3/3/3
9	L9R	I	5101	-	-	34/57/57/57	-
5	ATP	A	5305	-	-	6/18/38/38	0/3/3/3
9	L9R	I	5108	-	-	30/57/57/57	-
5	ATP	A	5301	-	-	8/18/38/38	0/3/3/3
8	KVR	I	5107	-	-	2/10/20/20	0/2/3/3
9	L9R	G	5101	-	-	34/57/57/57	-

The worst 5 of 60 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
8	B	5306	KVR	C06-S09	4.92	1.82	1.77
8	I	5107	KVR	C06-S09	4.91	1.82	1.77
8	A	5306	KVR	C06-S09	4.88	1.82	1.77
8	G	5107	KVR	C06-S09	4.86	1.82	1.77
9	A	5307	L9R	O2-C31	3.60	1.44	1.34

The worst 5 of 52 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
8	G	5107	KVR	C10-S09-C06	5.26	110.17	102.71
8	A	5306	KVR	C10-S09-C06	5.26	110.16	102.71
8	B	5306	KVR	C10-S09-C06	5.24	110.14	102.71
8	I	5107	KVR	C10-S09-C06	5.24	110.13	102.71
9	B	5308	L9R	O2-C31-C32	4.03	120.20	111.48

There are no chirality outliers.

5 of 320 torsion outliers are listed below:

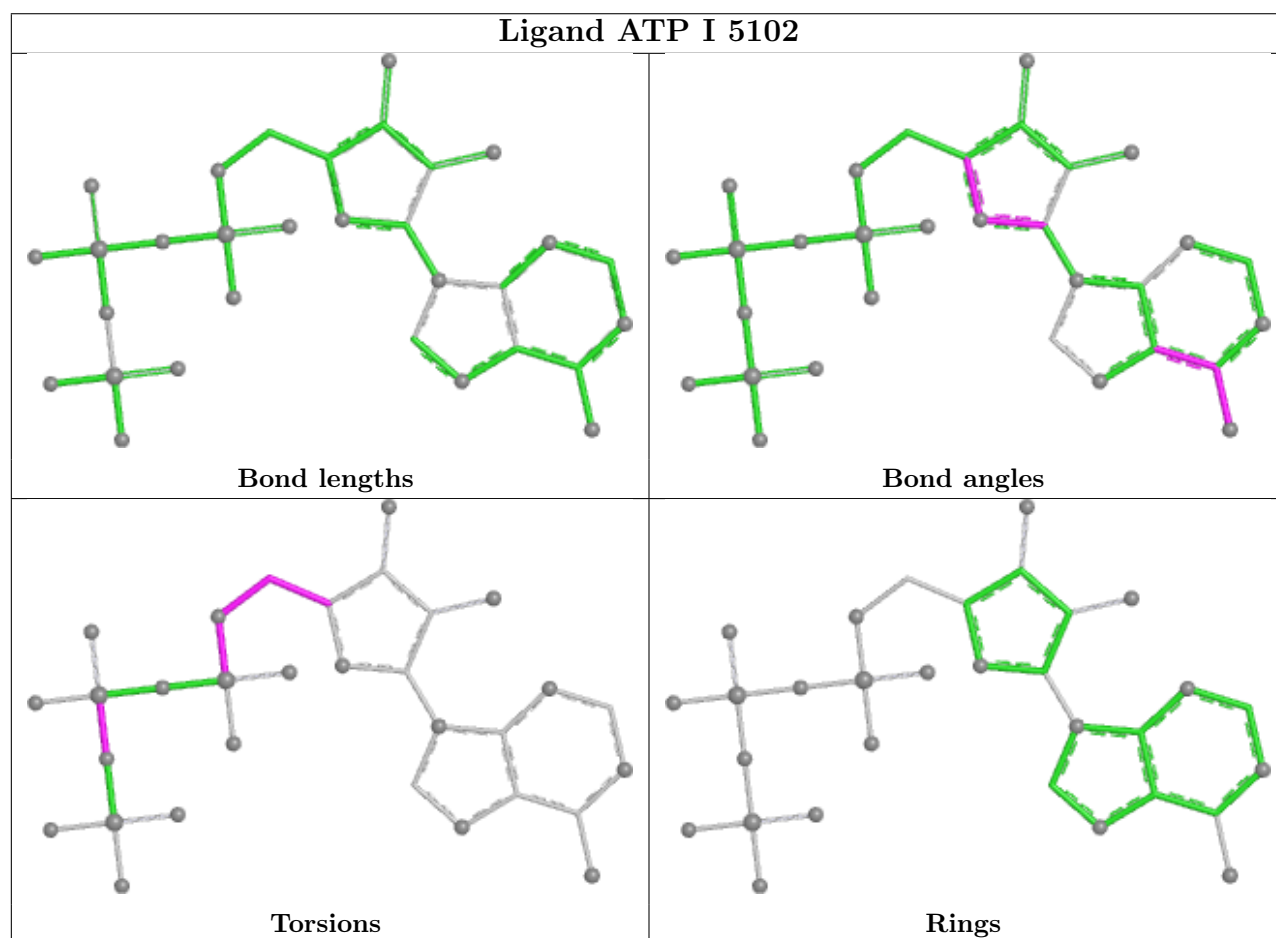
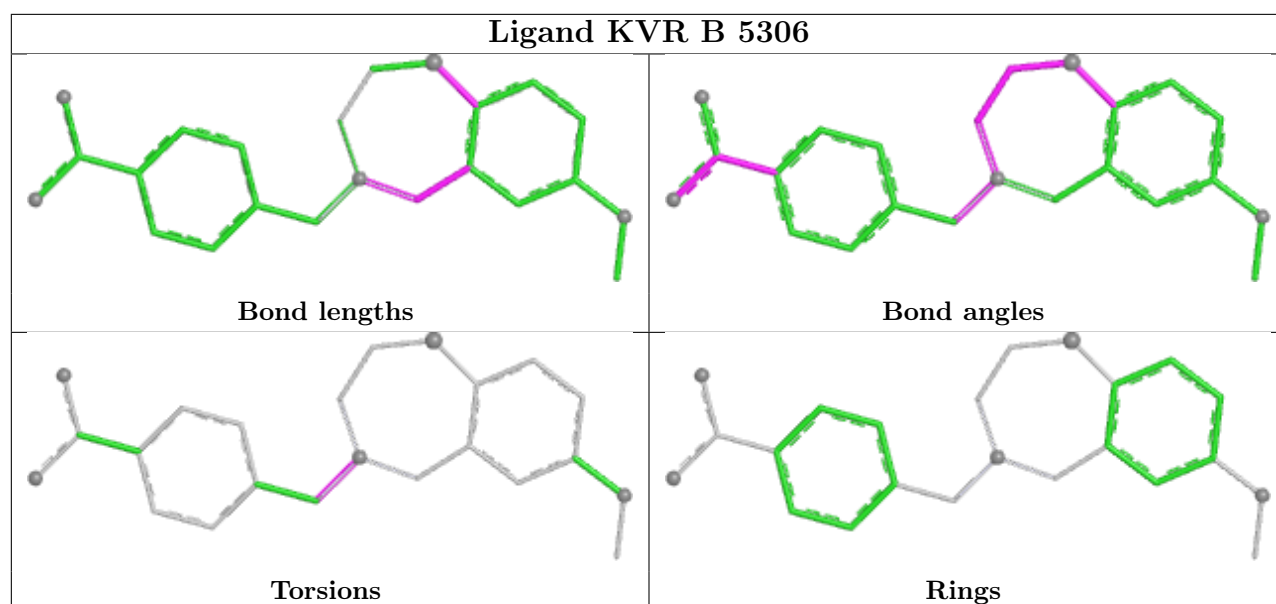
Mol	Chain	Res	Type	Atoms
5	A	5301	ATP	C5'-O5'-PA-O1A
5	A	5301	ATP	C5'-O5'-PA-O2A
5	A	5301	ATP	C5'-O5'-PA-O3A
5	A	5305	ATP	C5'-O5'-PA-O2A
5	A	5305	ATP	C5'-O5'-PA-O3A

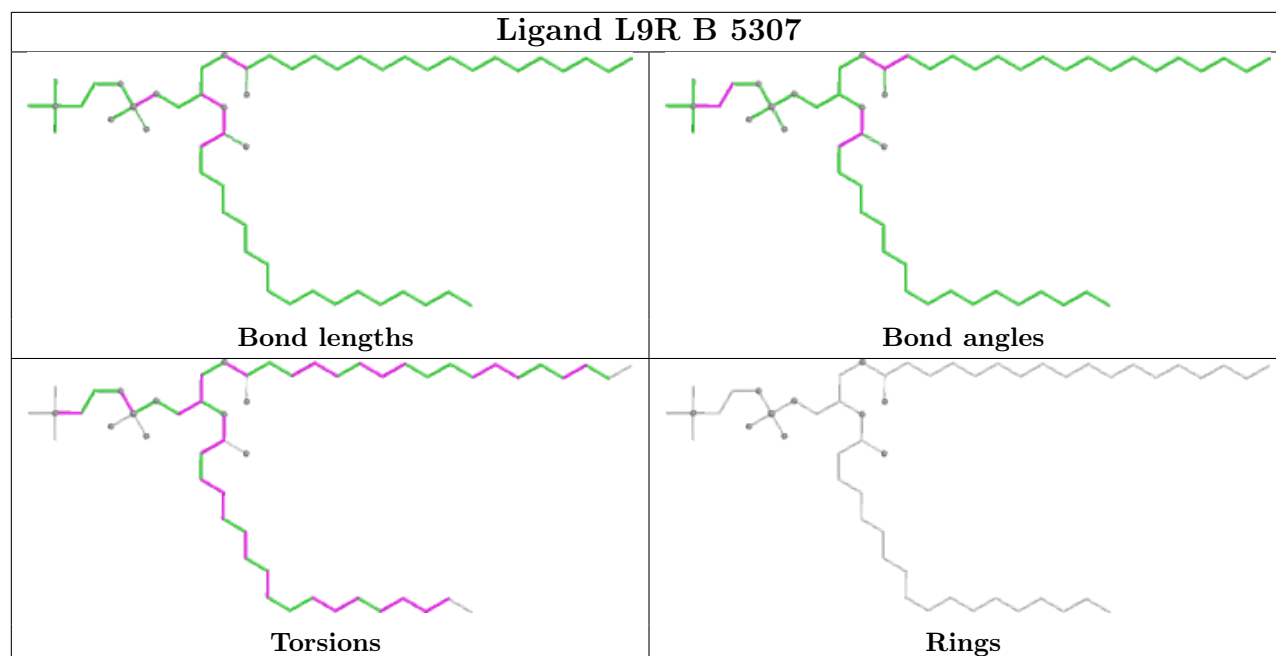
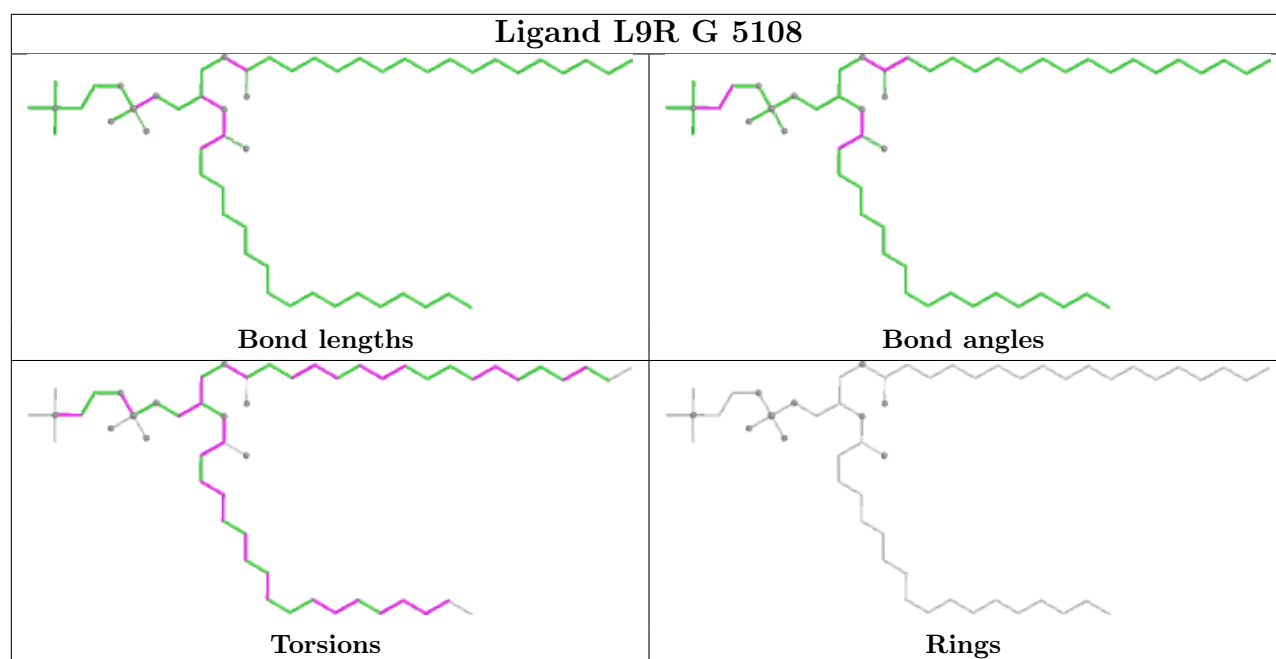
There are no ring outliers.

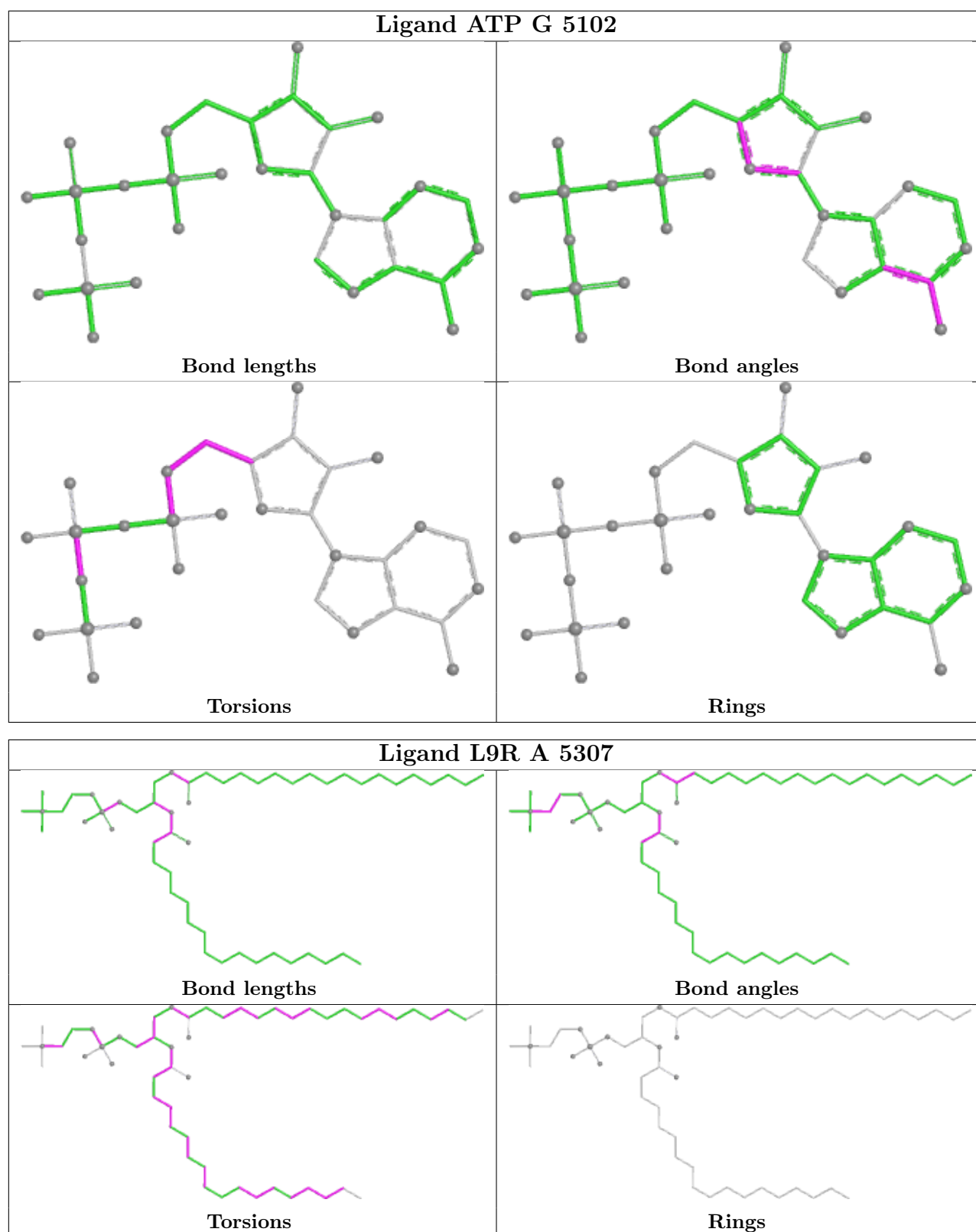
16 monomers are involved in 36 short contacts:

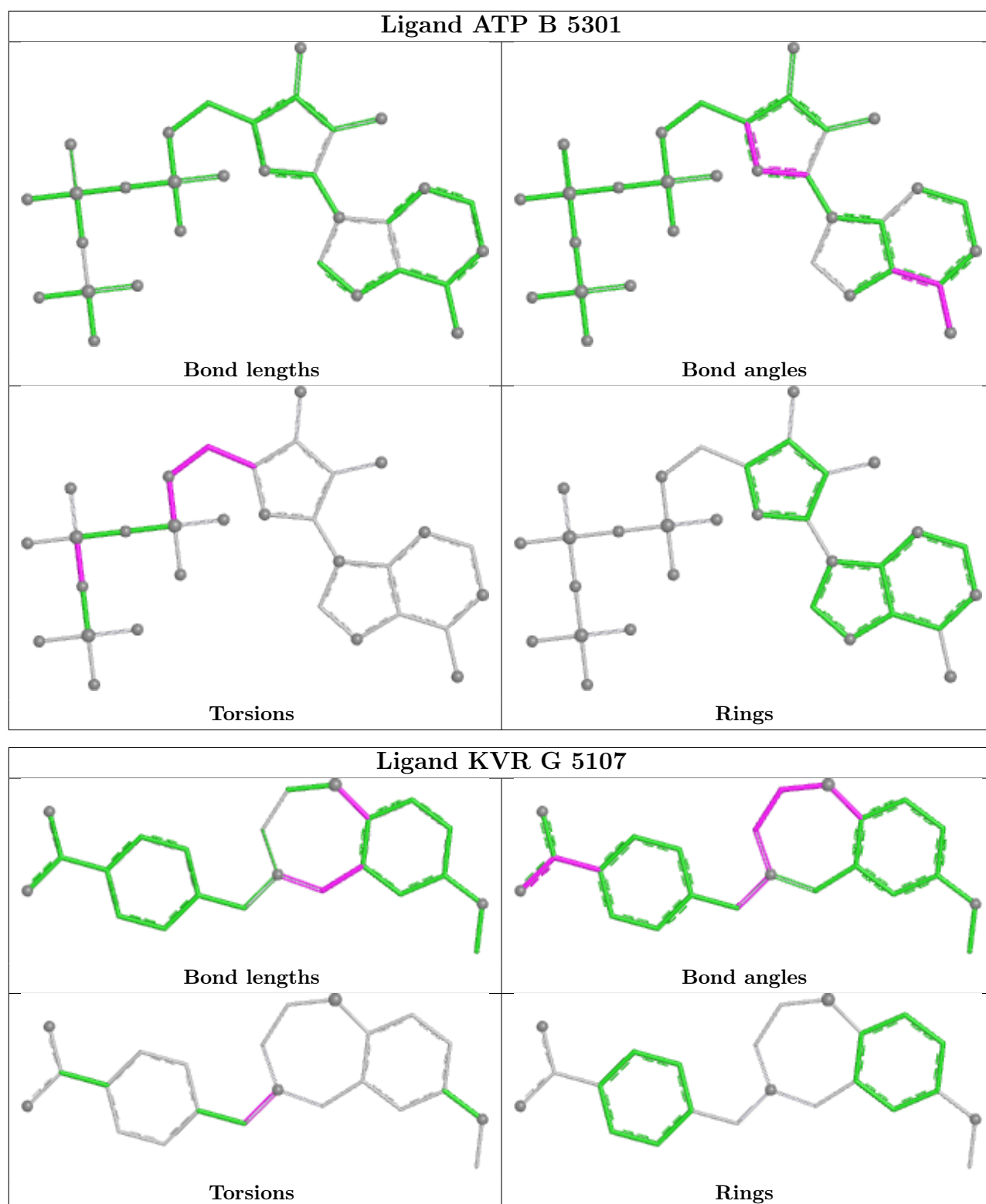
Mol	Chain	Res	Type	Clashes	Symm-Clashes
5	I	5102	ATP	1	0
9	G	5108	L9R	3	0
9	B	5307	L9R	2	0
5	G	5102	ATP	1	0
9	A	5307	L9R	3	0
5	B	5301	ATP	1	0
9	A	5308	L9R	6	0
9	B	5308	L9R	4	0
5	G	5106	ATP	1	0
5	I	5106	ATP	1	0
5	B	5305	ATP	1	0
9	I	5101	L9R	5	0
5	A	5305	ATP	1	0
9	I	5108	L9R	3	0
5	A	5301	ATP	1	0
9	G	5101	L9R	6	0

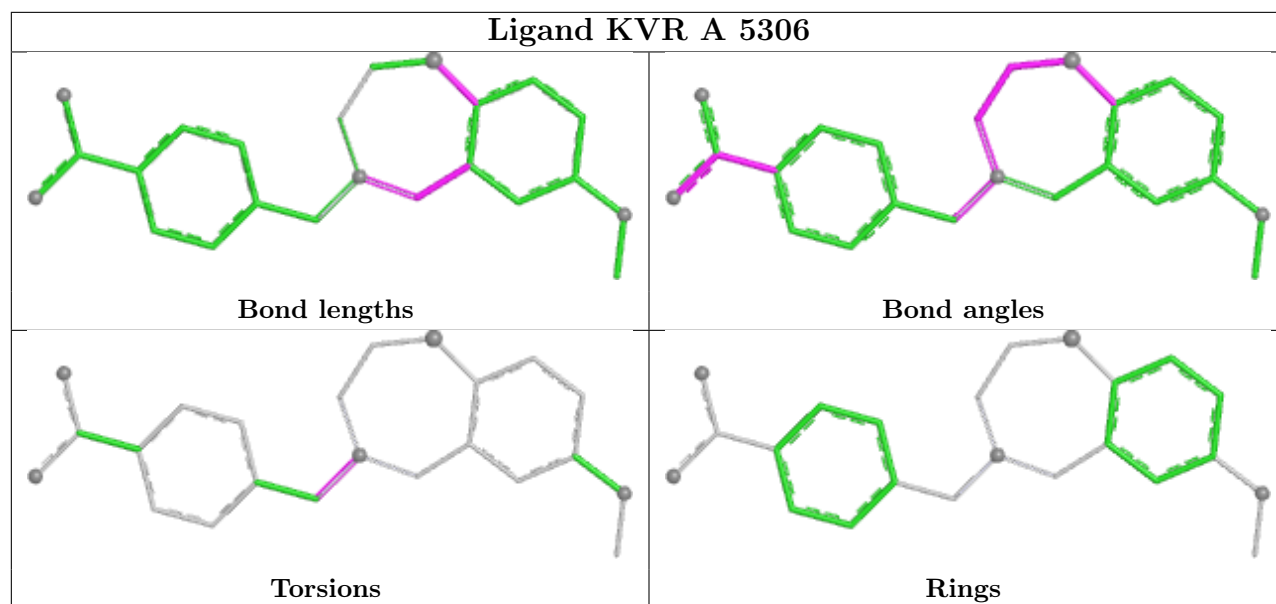
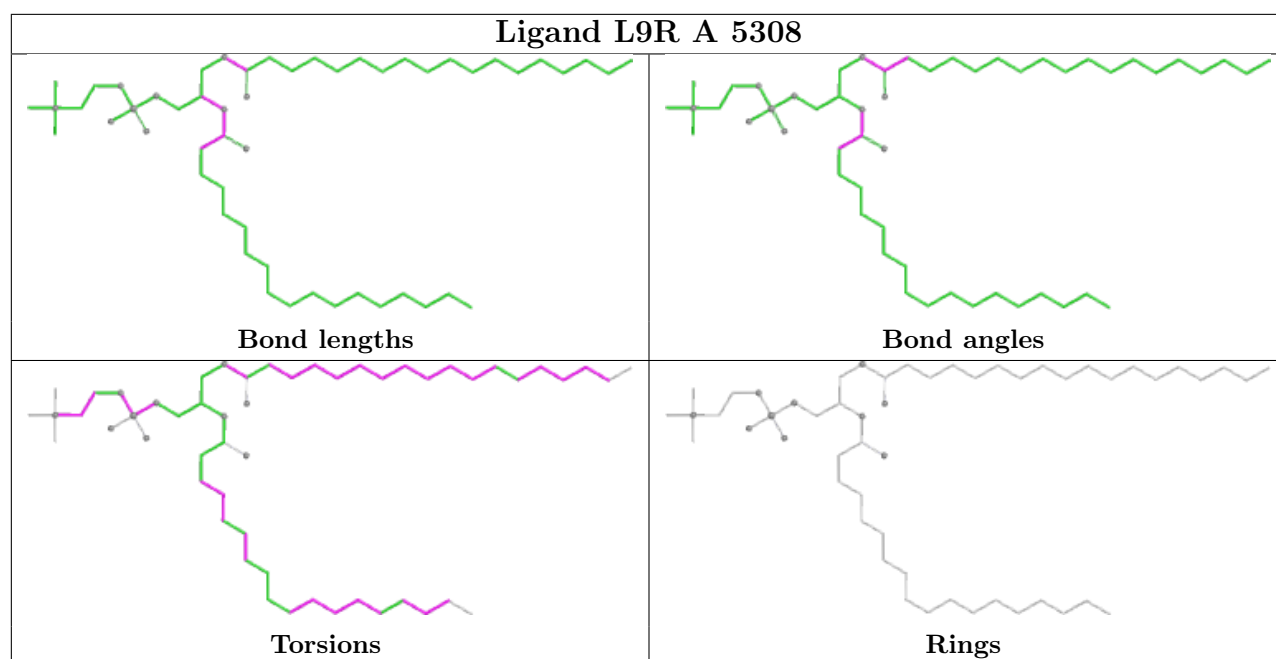
The following is a two-dimensional graphical depiction of Mogul quality analysis of bond lengths, bond angles, torsion angles, and ring geometry for all instances of the Ligand of Interest. In addition, ligands with molecular weight > 250 and outliers as shown on the validation Tables will also be included. For torsion angles, if less than 5% of the Mogul distribution of torsion angles is within 10 degrees of the torsion angle in question, then that torsion angle is considered an outlier. Any bond that is central to one or more torsion angles identified as an outlier by Mogul will be highlighted in the graph. For rings, the root-mean-square deviation (RMSD) between the ring in question and similar rings identified by Mogul is calculated over all ring torsion angles. If the average RMSD is greater than 60 degrees and the minimal RMSD between the ring in question and any Mogul-identified rings is also greater than 60 degrees, then that ring is considered an outlier. The outliers are highlighted in purple. The color gray indicates Mogul did not find sufficient equivalents in the CSD to analyse the geometry.

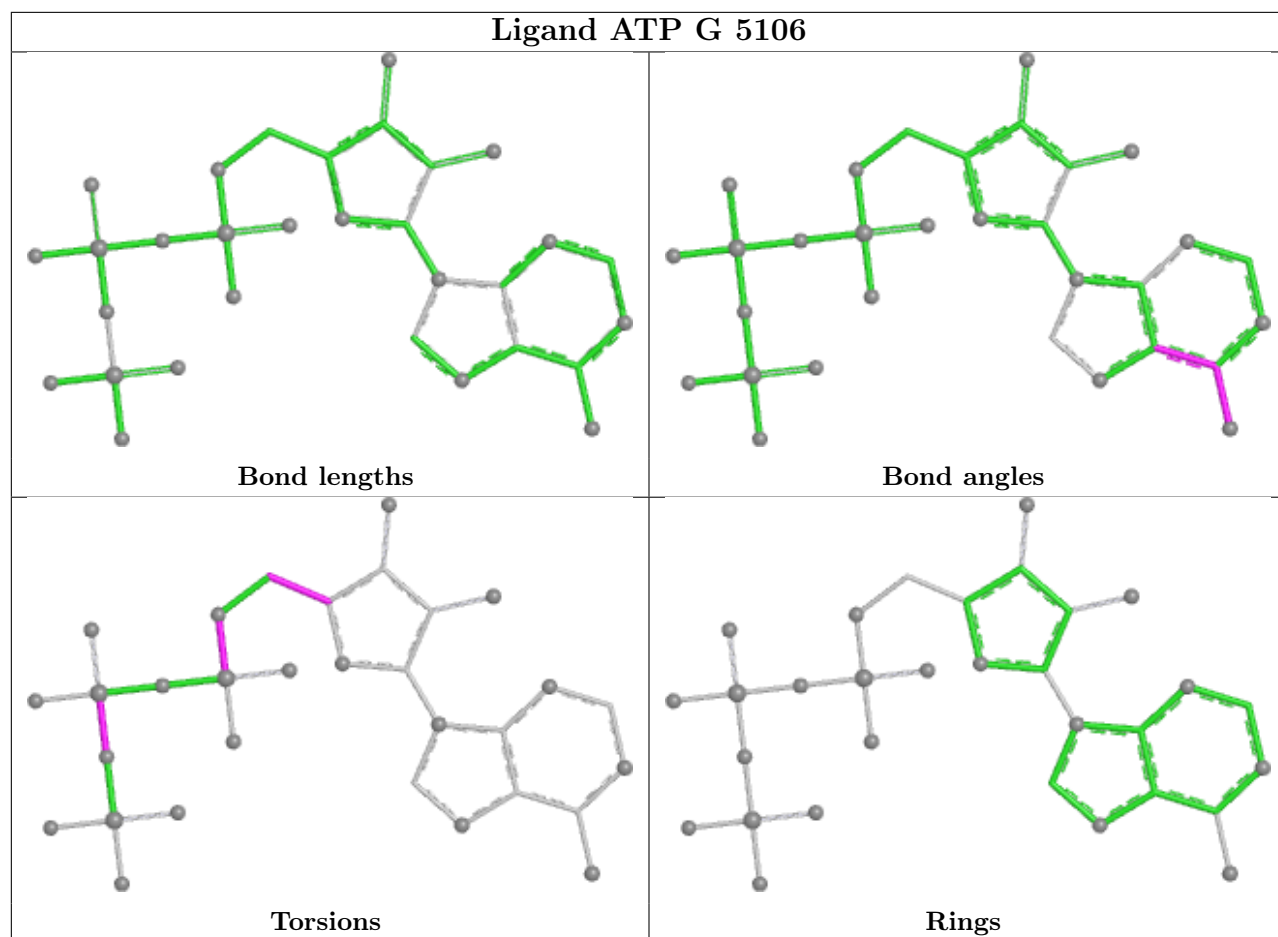
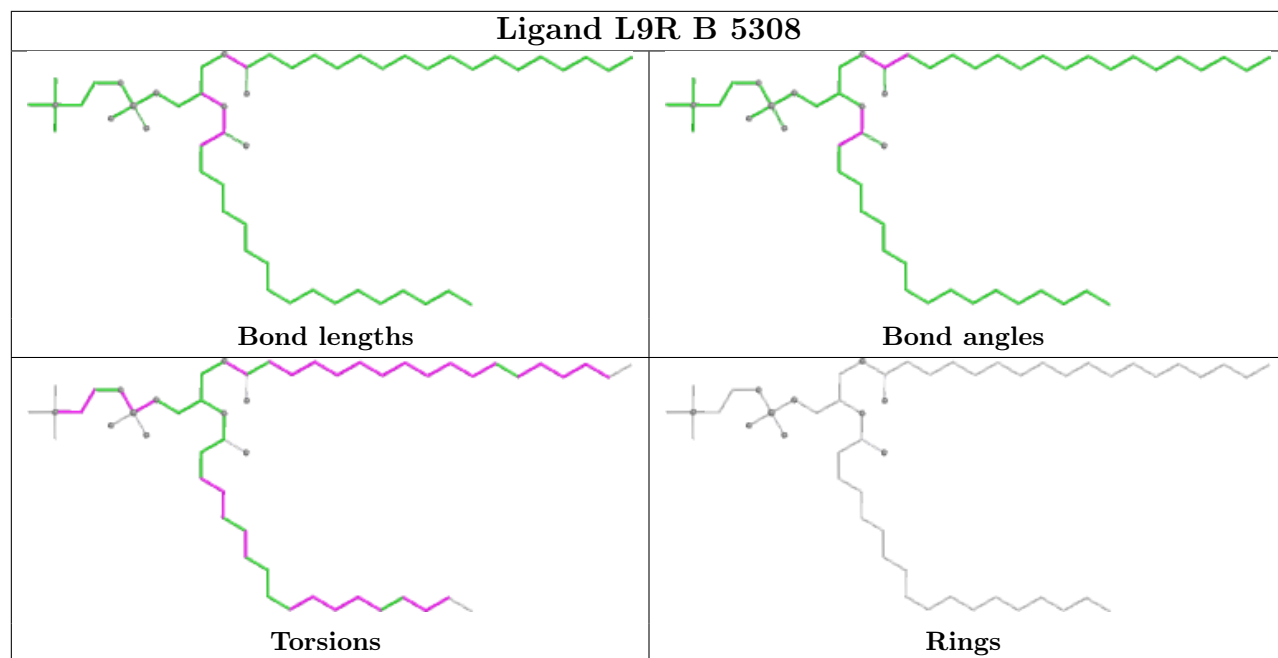


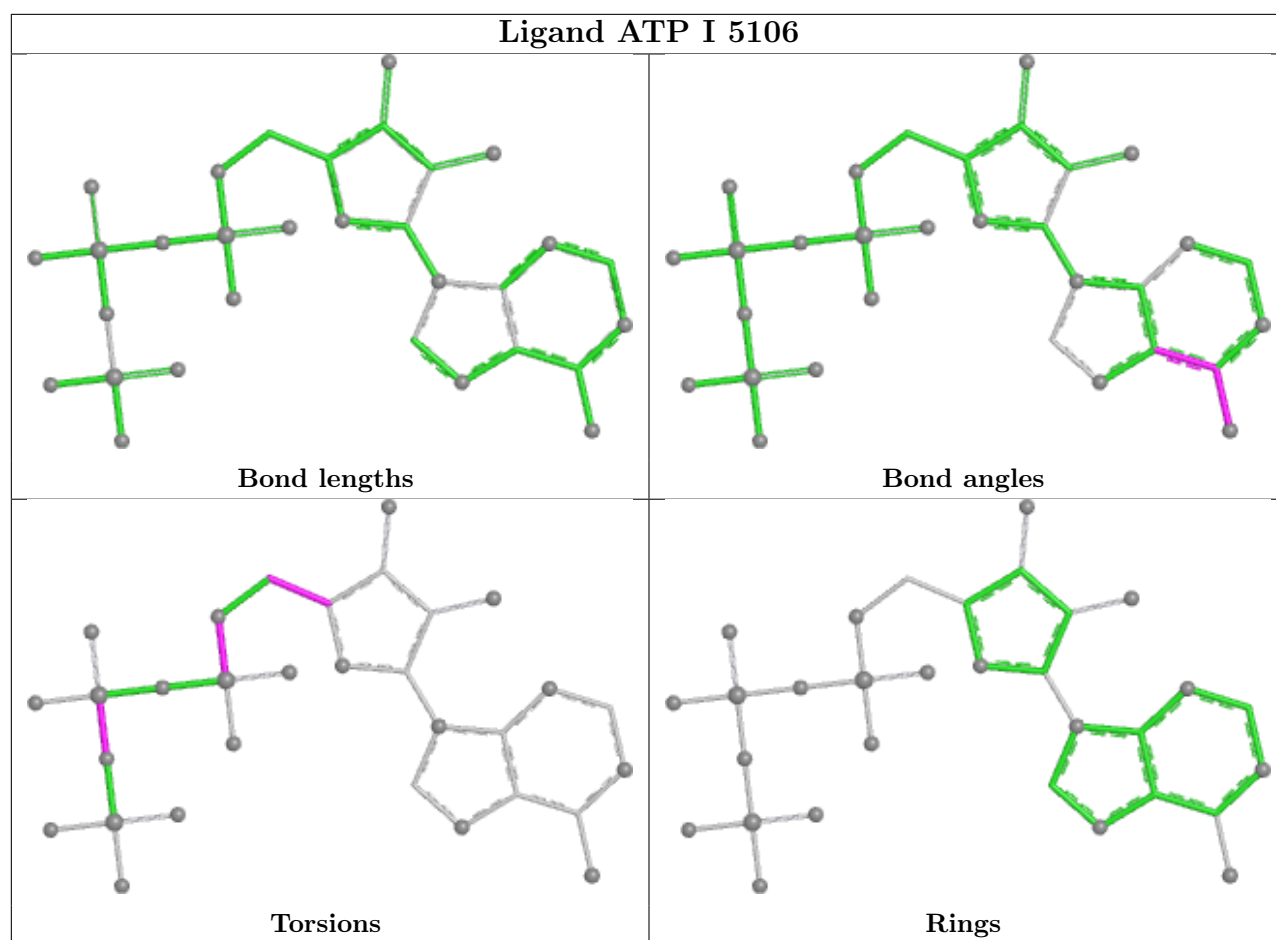


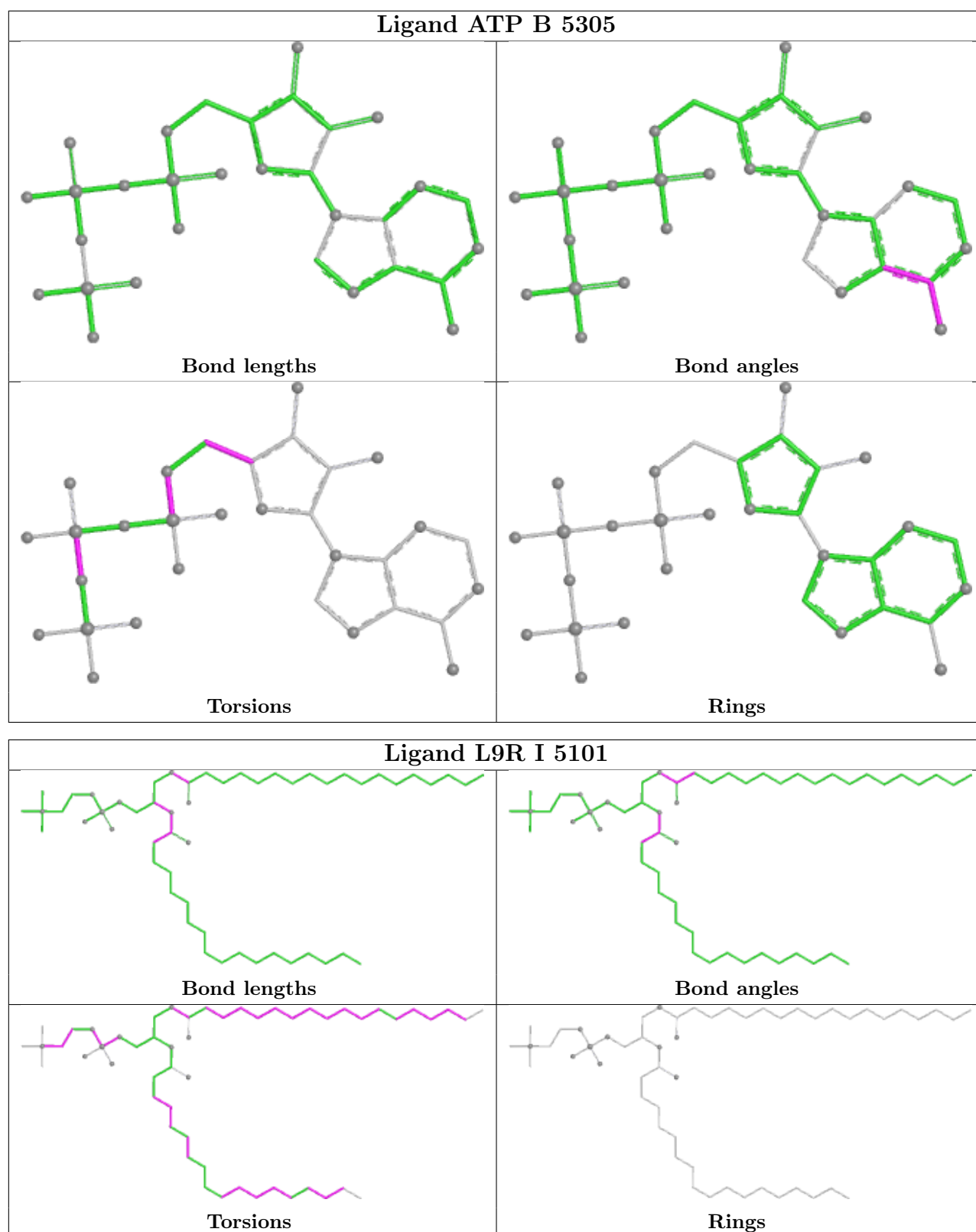


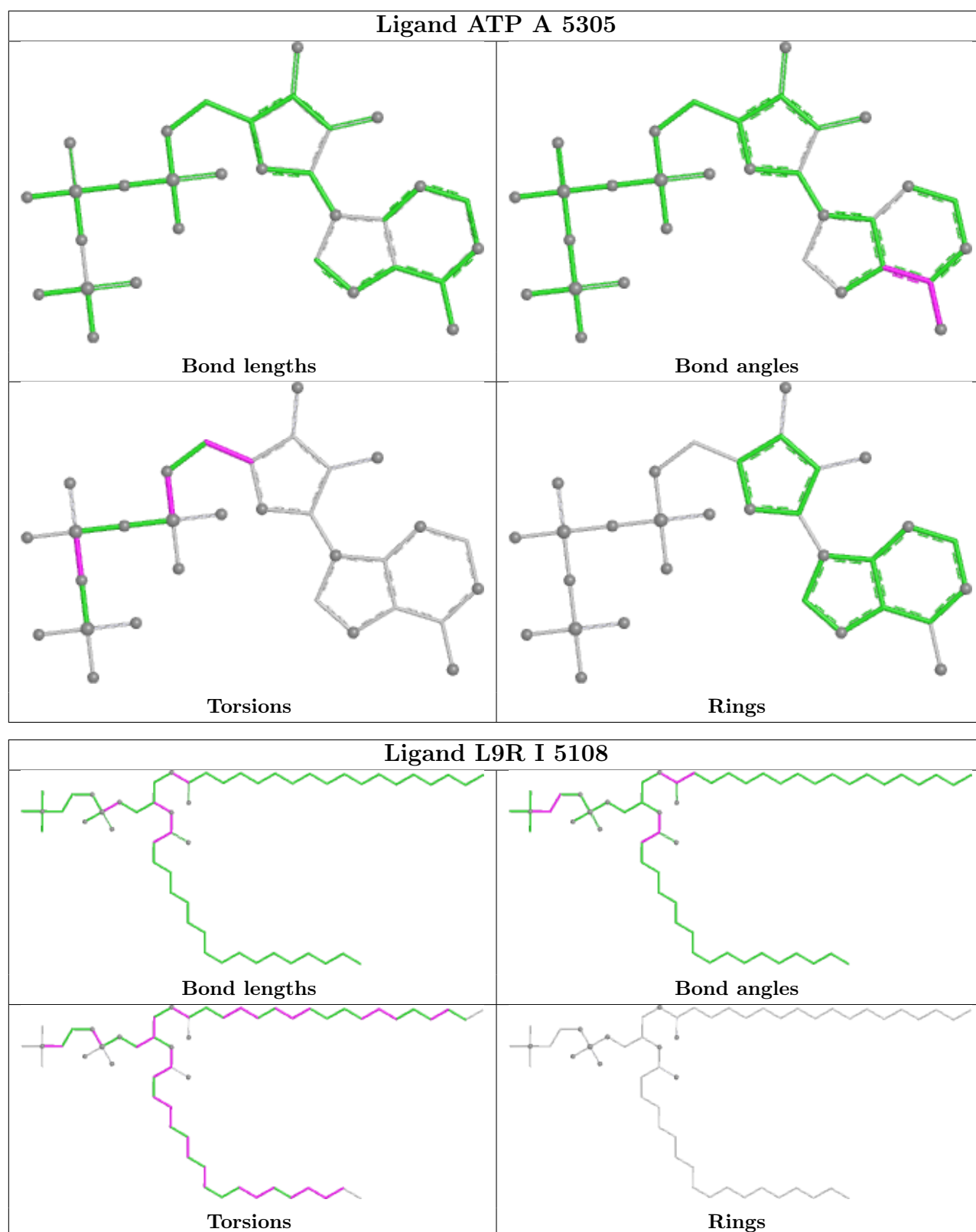


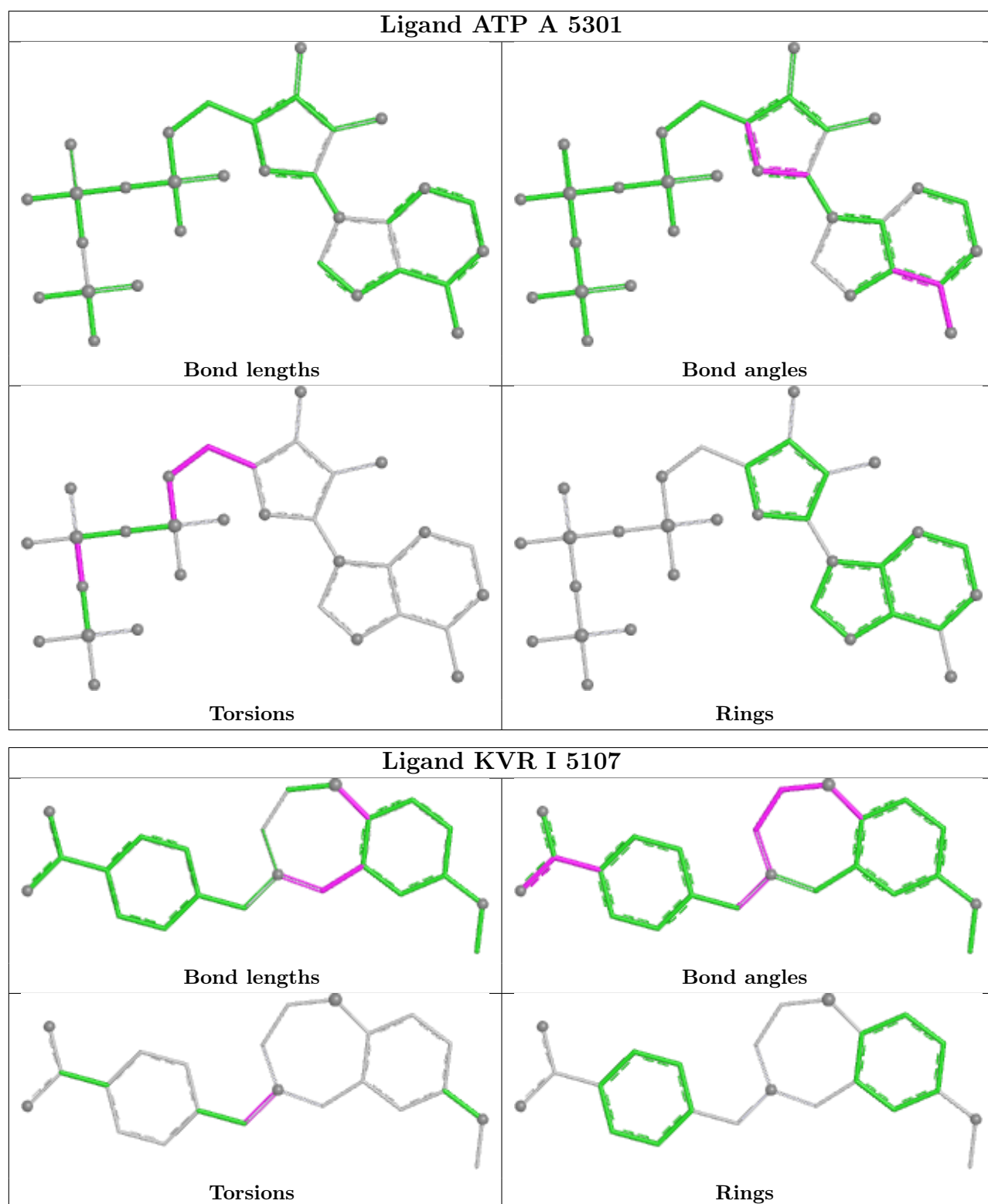


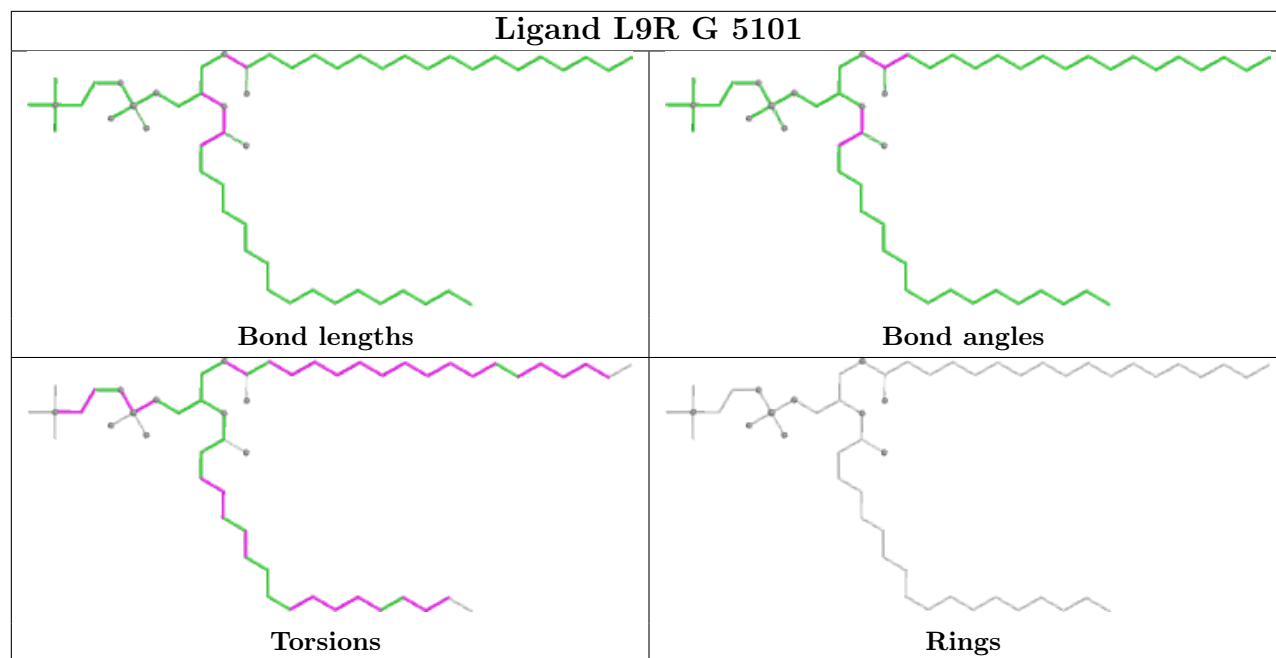












5.7 Other polymers [i](#)

There are no such residues in this entry.

5.8 Polymer linkage issues [i](#)

There are no chain breaks in this entry.

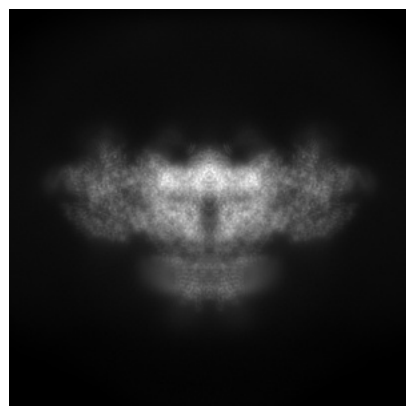
6 Map visualisation [i](#)

This section contains visualisations of the EMDB entry EMD-26205. These allow visual inspection of the internal detail of the map and identification of artifacts.

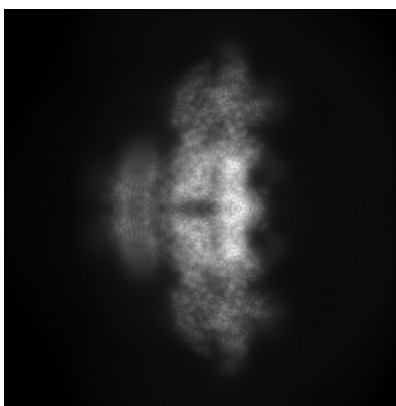
Images derived from a raw map, generated by summing the deposited half-maps, are presented below the corresponding image components of the primary map to allow further visual inspection and comparison with those of the primary map.

6.1 Orthogonal projections [i](#)

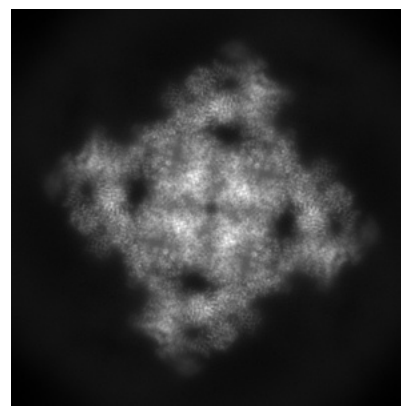
6.1.1 Primary map



X

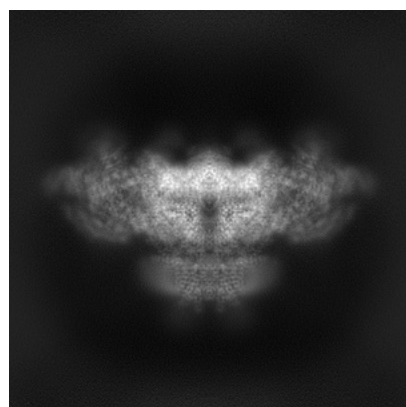


Y

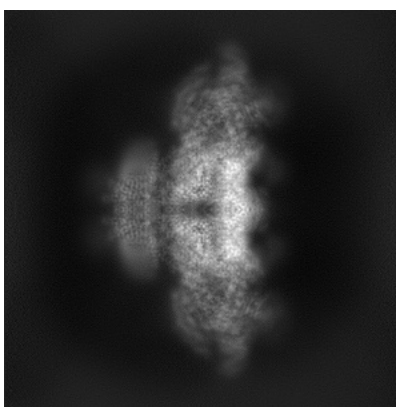


Z

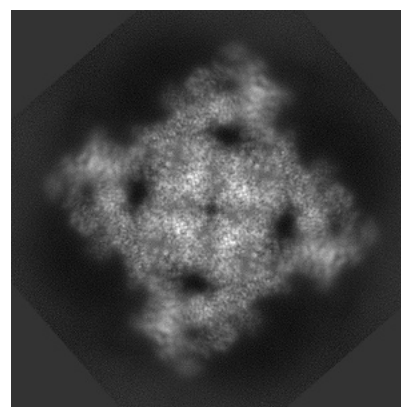
6.1.2 Raw map



X



Y



Z

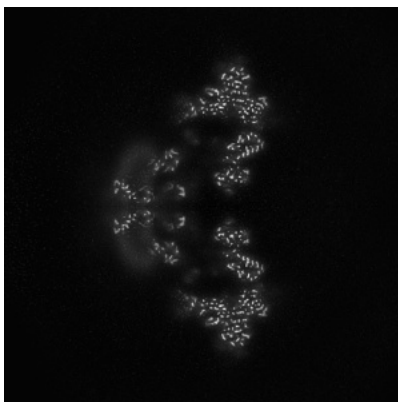
The images above show the map projected in three orthogonal directions.

6.2 Central slices [i](#)

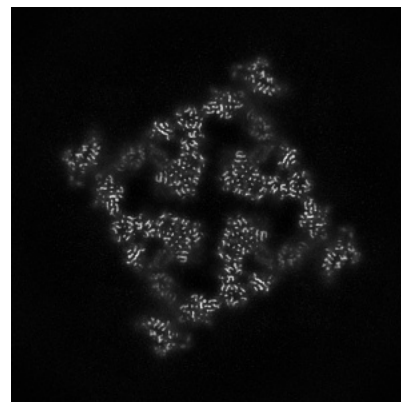
6.2.1 Primary map



X Index: 256

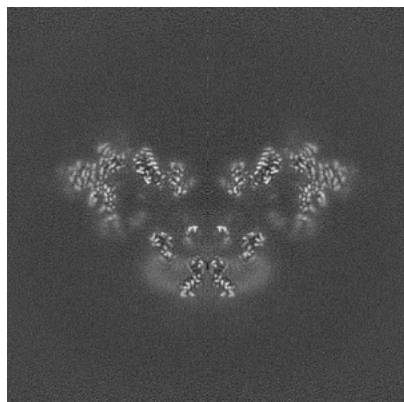


Y Index: 256

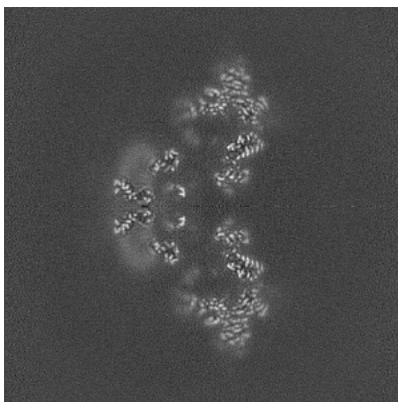


Z Index: 256

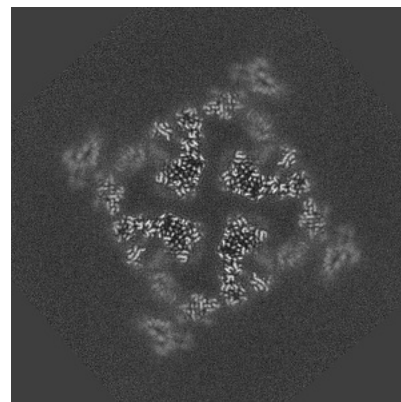
6.2.2 Raw map



X Index: 256



Y Index: 256

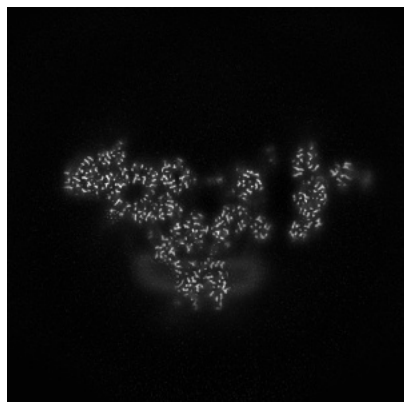


Z Index: 256

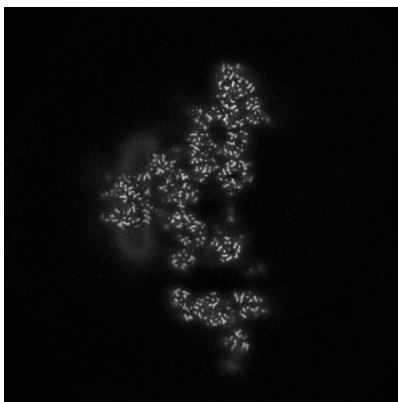
The images above show central slices of the map in three orthogonal directions.

6.3 Largest variance slices [i](#)

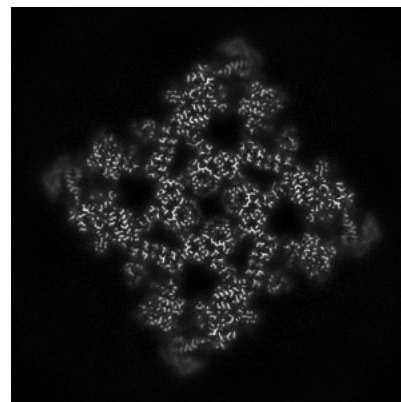
6.3.1 Primary map



X Index: 272

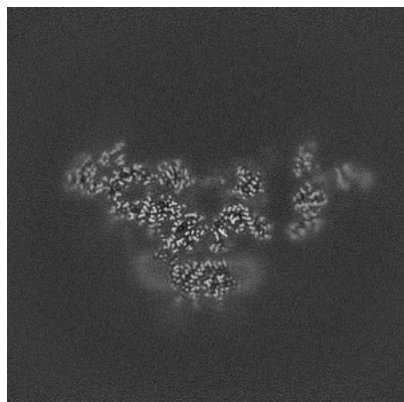


Y Index: 272

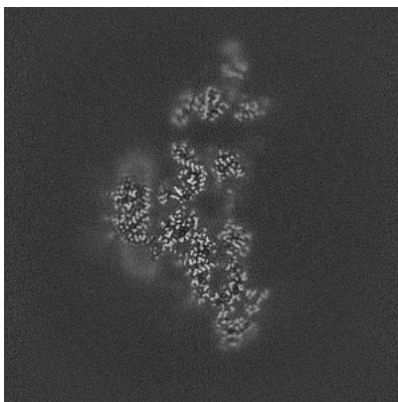


Z Index: 286

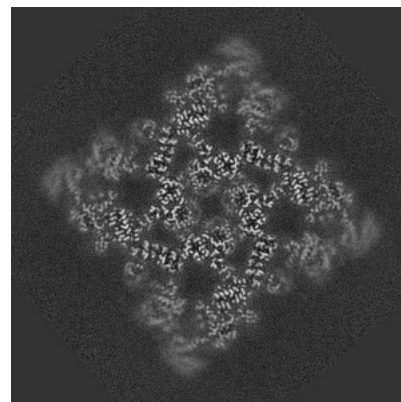
6.3.2 Raw map



X Index: 275



Y Index: 237

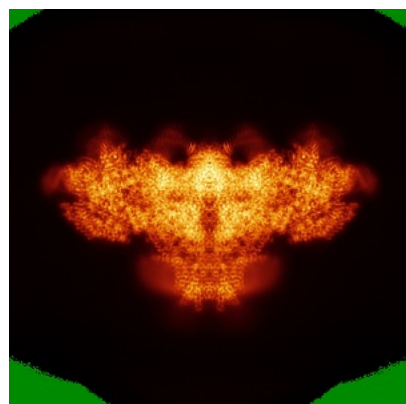


Z Index: 286

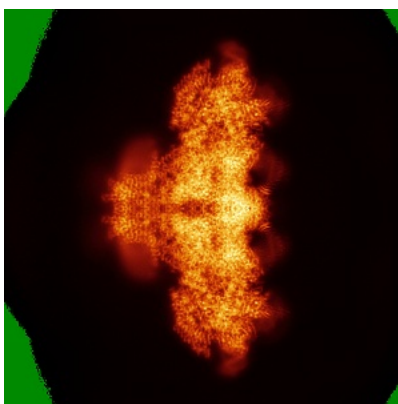
The images above show the largest variance slices of the map in three orthogonal directions.

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

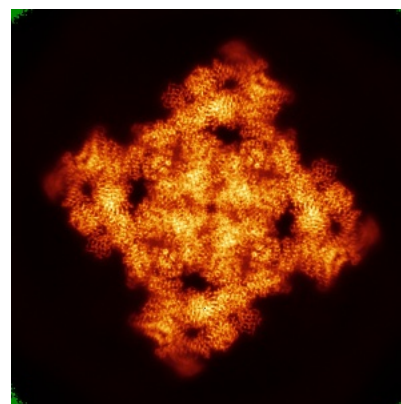
6.4.1 Primary map



X



Y

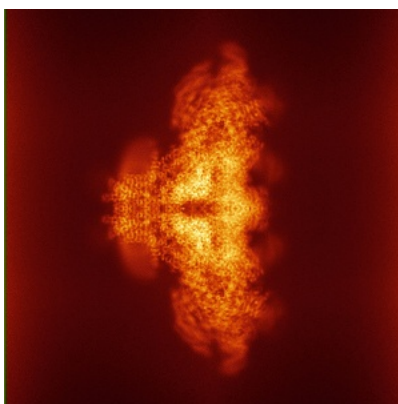


Z

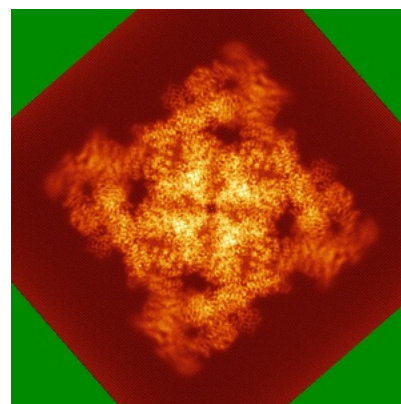
6.4.2 Raw map



X



Y

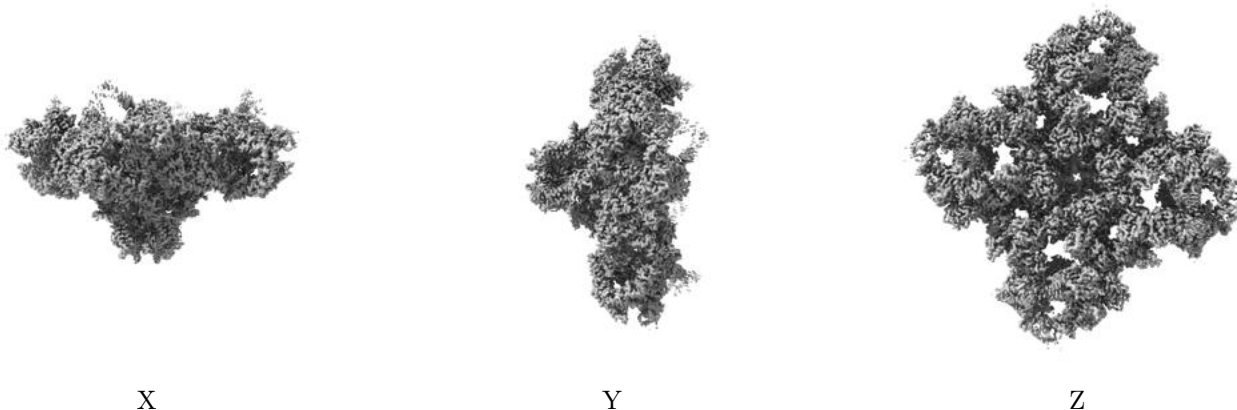


Z

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 0.15. 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.

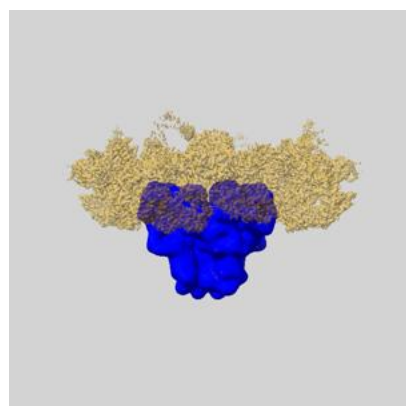
6.6 Mask visualisation [i](#)

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

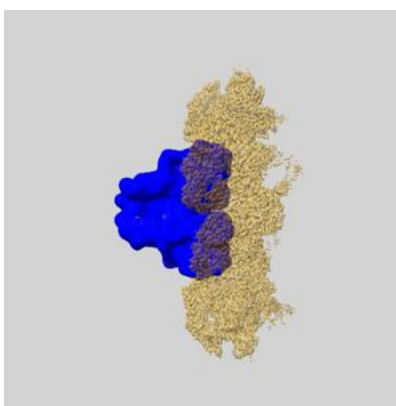
A mask typically either:

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

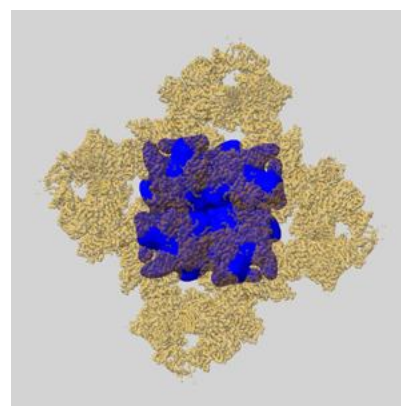
6.6.1 emd_26205_msk_1.map [i](#)



X

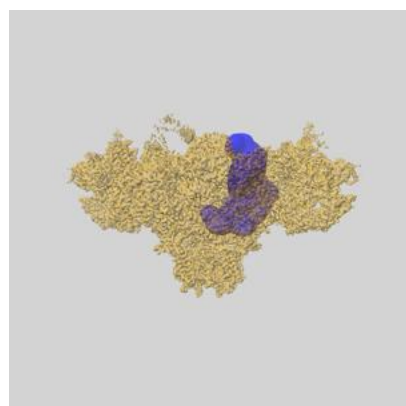


Y

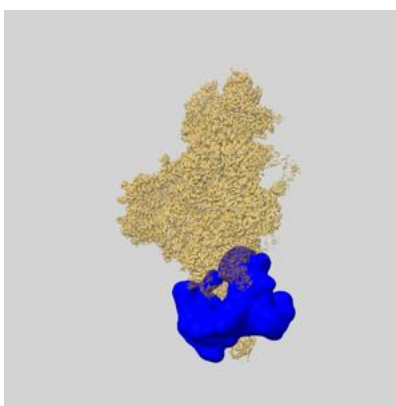


Z

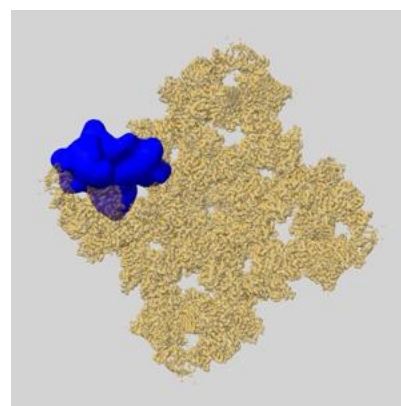
6.6.2 emd_26205_msk_2.map [i](#)



X

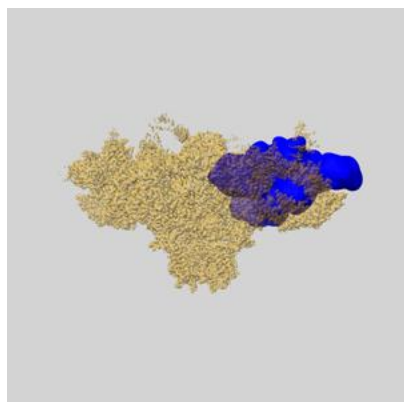


Y

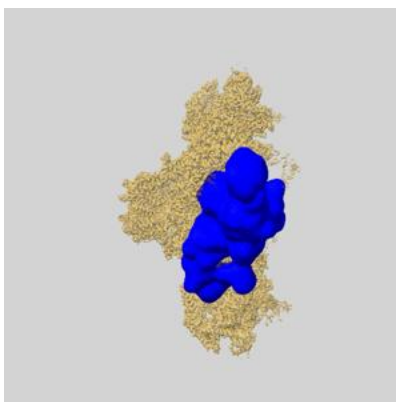


Z

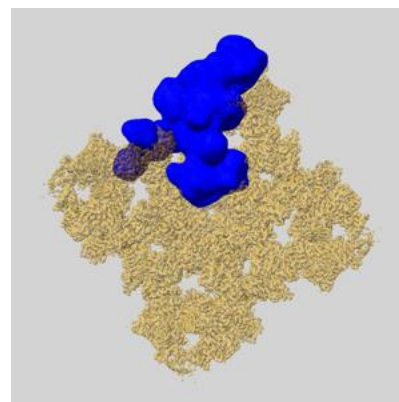
6.6.3 emd_26205_msk_3.map [i](#)



X



Y

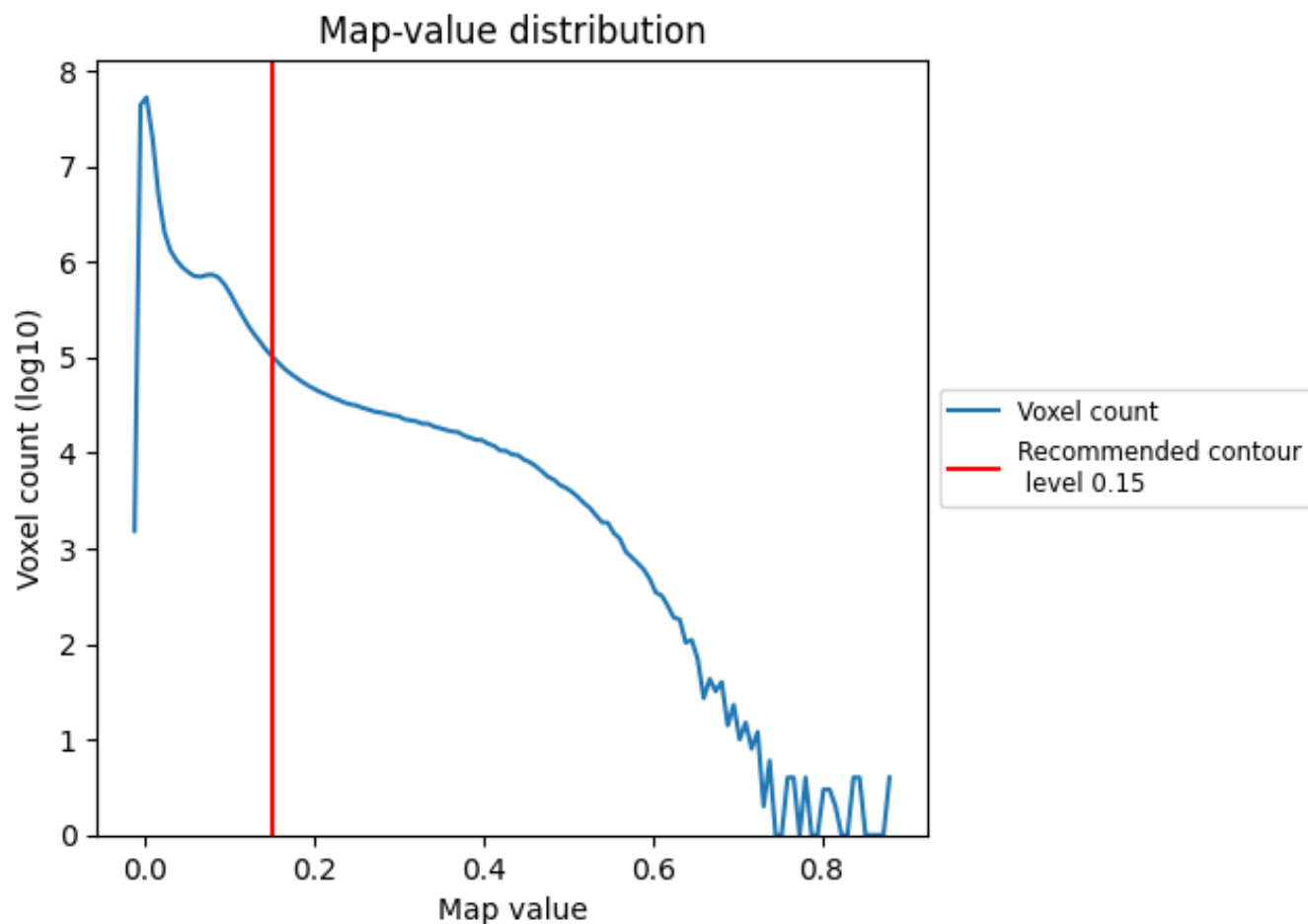


Z

7 Map analysis [i](#)

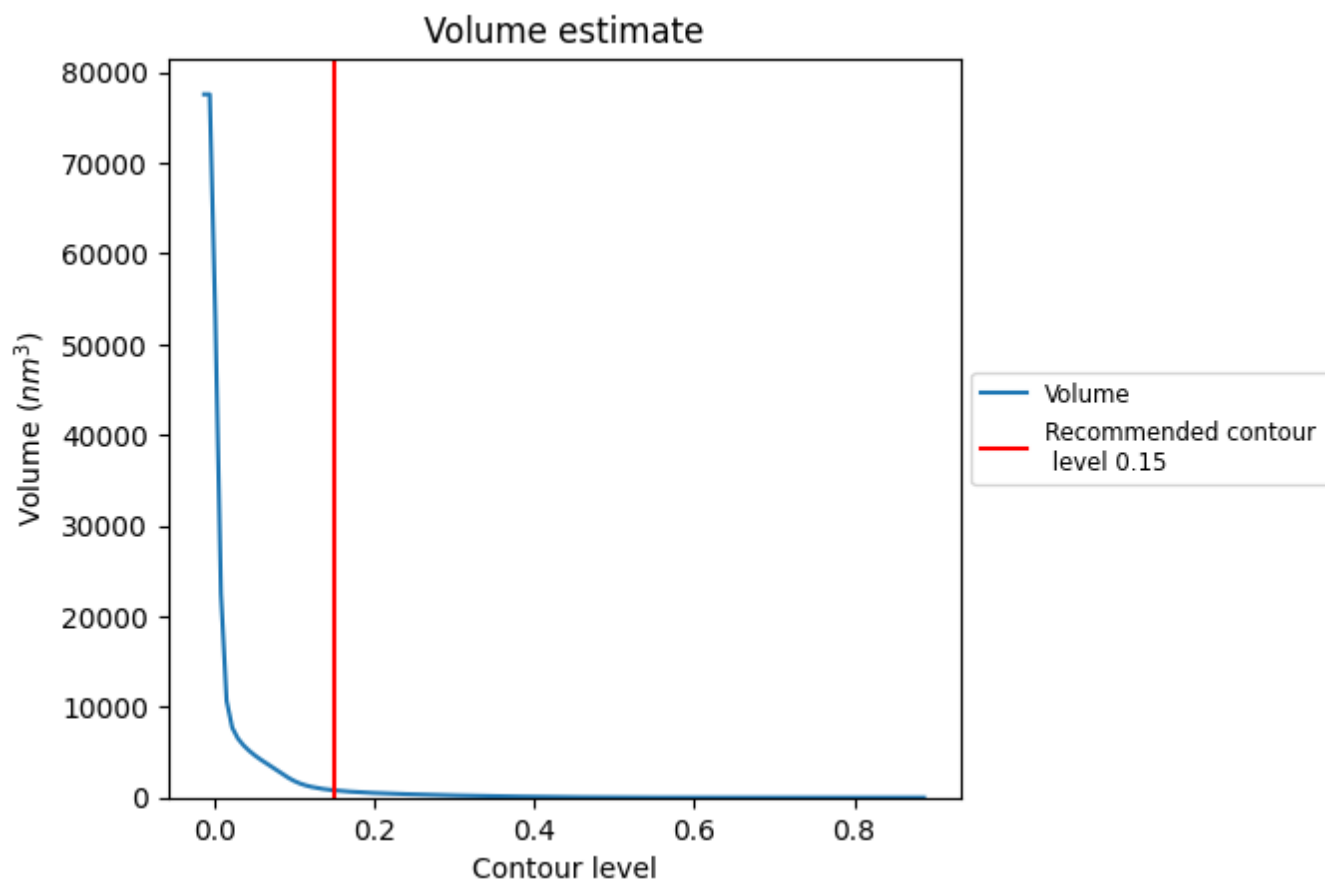
This section contains the results of statistical analysis of the map.

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.

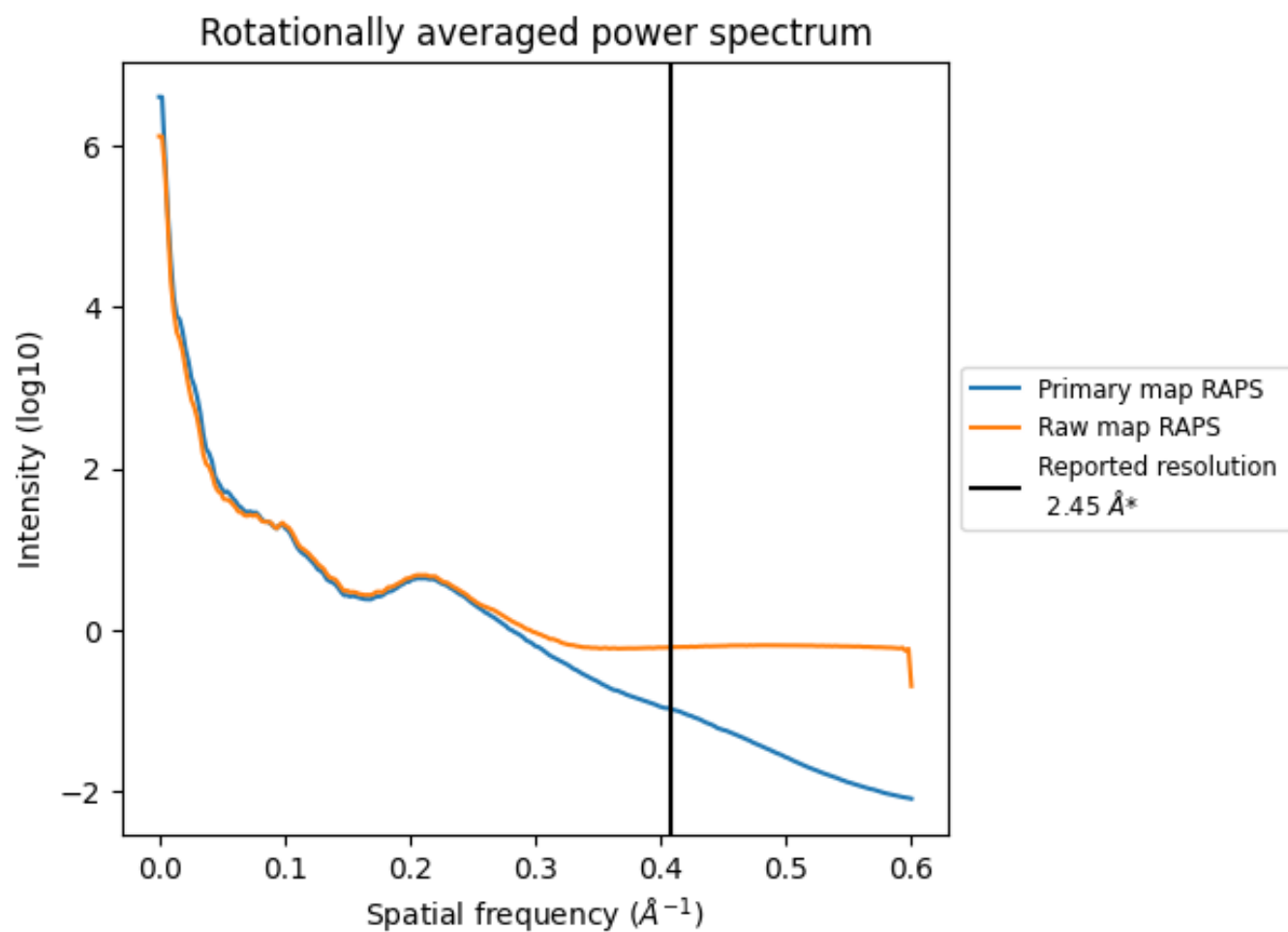
7.2 Volume estimate [i](#)



The volume at the recommended contour level is 800 nm³; this corresponds to an approximate mass of 723 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 ⓘ

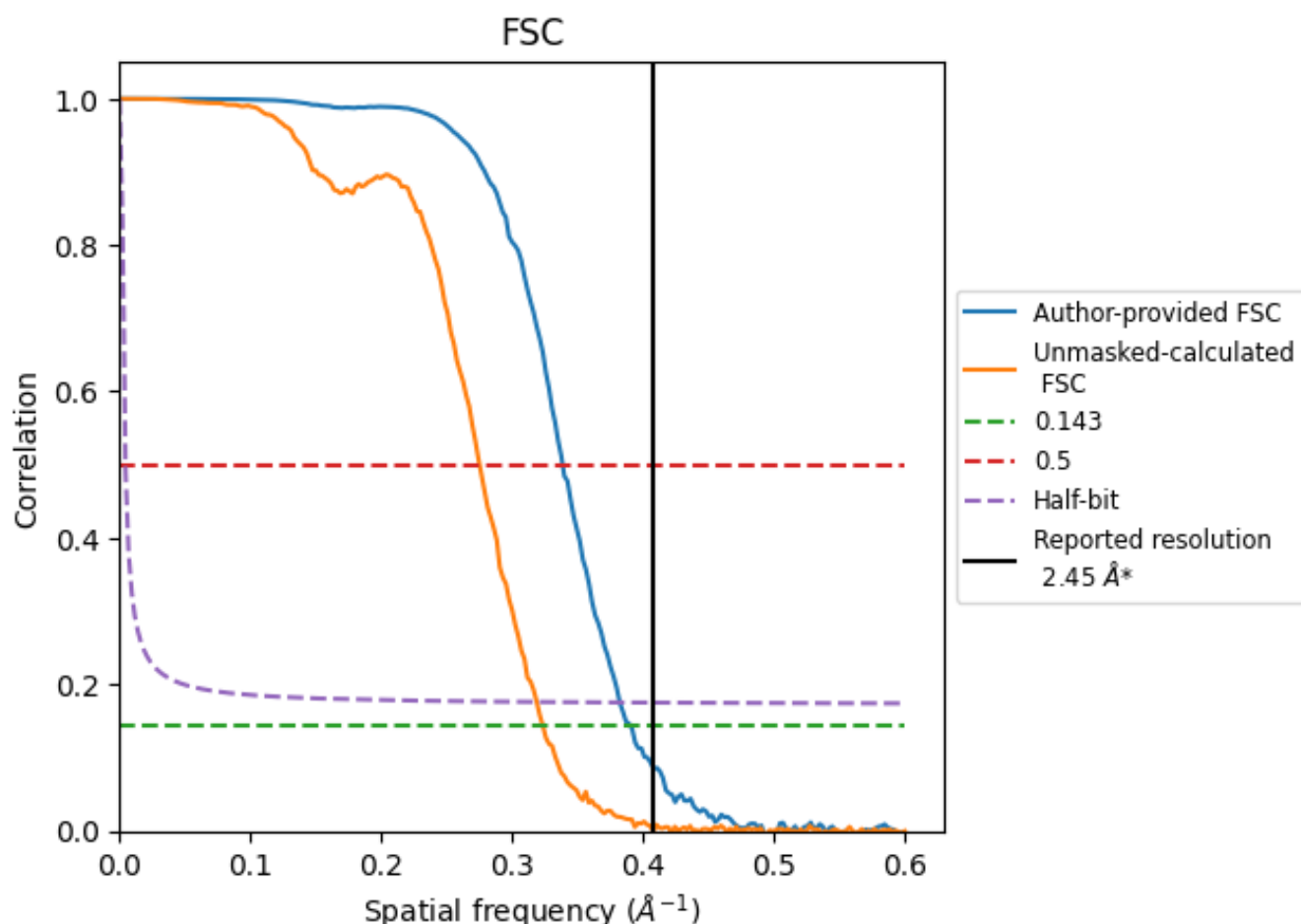


*Reported resolution corresponds to spatial frequency of 0.408 Å⁻¹

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 0.408 Å⁻¹

8.2 Resolution estimates

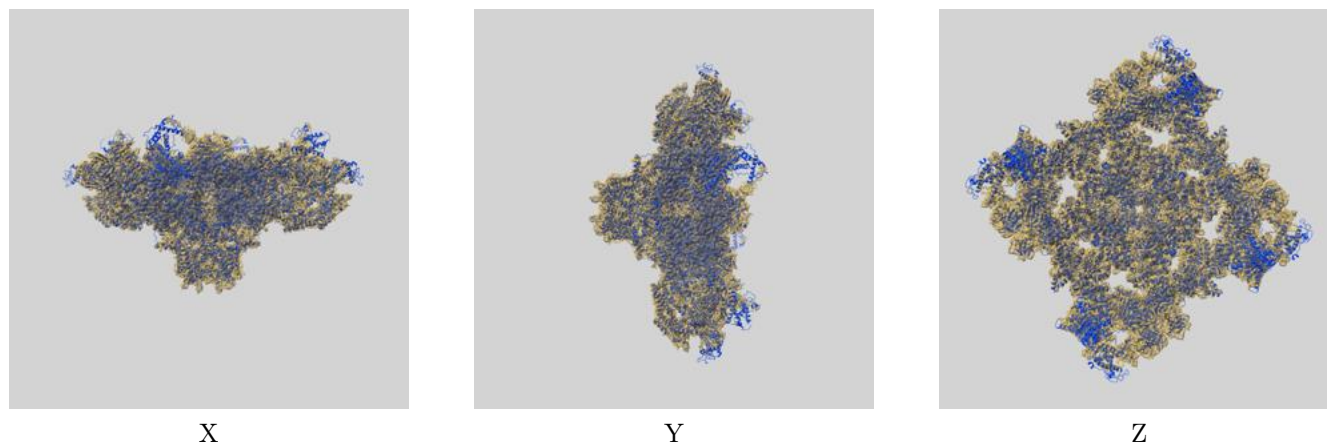
Resolution estimate (Å)	Estimation criterion (FSC cut-off)		
	0.143	0.5	Half-bit
Reported by author	2.45	-	-
Author-provided FSC curve	2.56	2.95	2.62
Unmasked-calculated*	3.09	3.63	3.14

*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 3.09 differs from the reported value 2.45 by more than 10 %

9 Map-model fit [i](#)

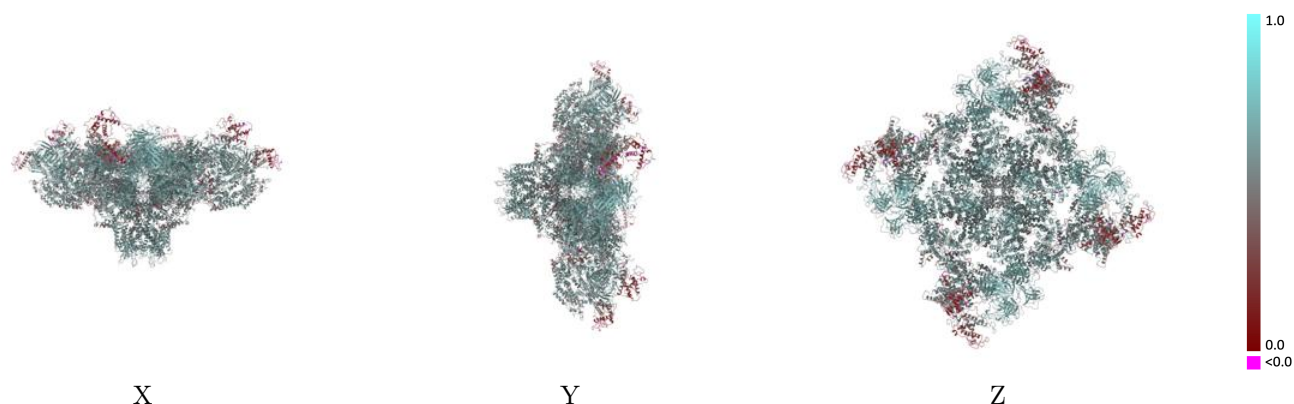
This section contains information regarding the fit between EMDB map EMD-26205 and PDB model 7TZC. Per-residue inclusion information can be found in section [3](#) on page [10](#).

9.1 Map-model overlay [i](#)



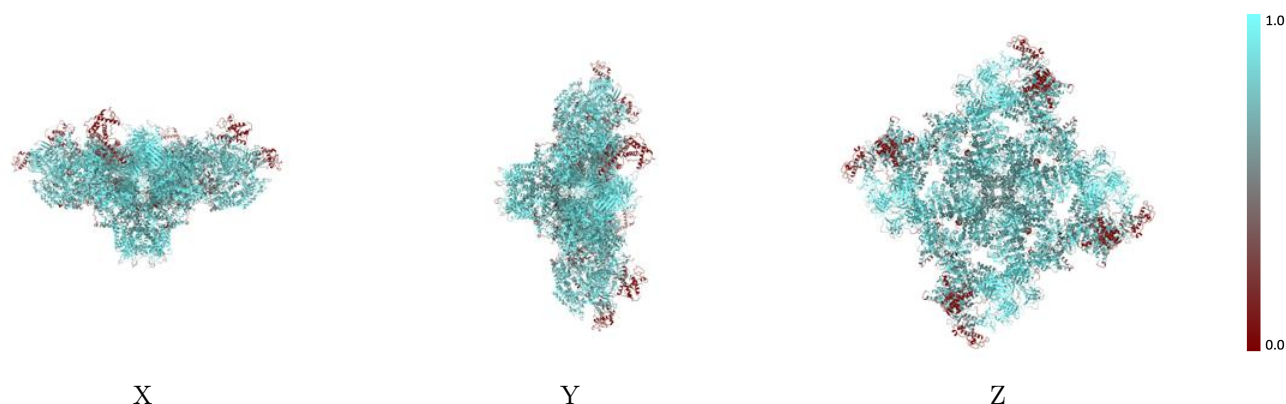
The images above show the 3D surface view of the map at the recommended contour level 0.15 at 50% transparency in yellow overlaid with a ribbon representation of the model coloured in blue. These images allow for the visual assessment of the quality of fit between the atomic model and the map.

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



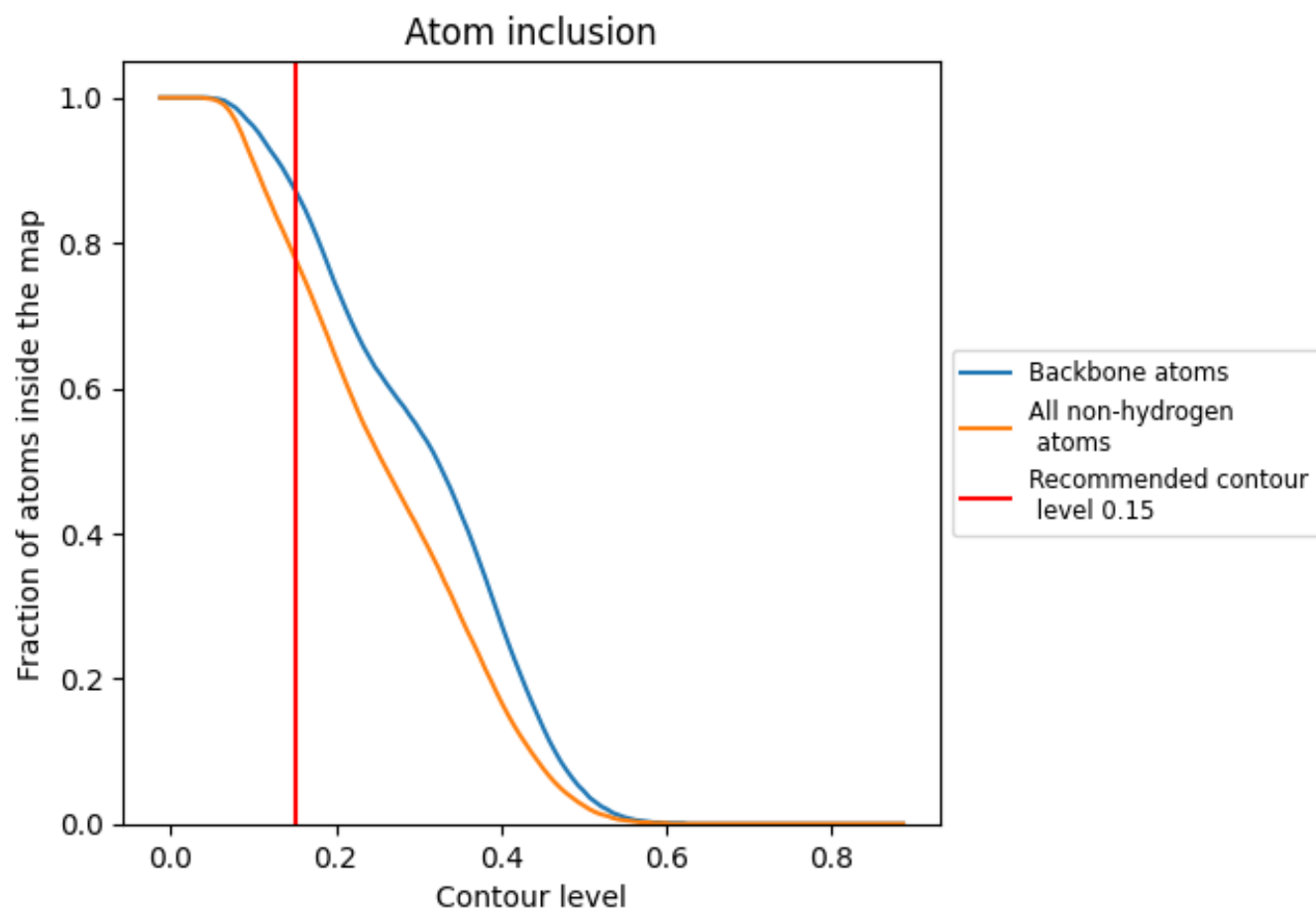
The images above show the model with each residue coloured according to its Q-score. This shows their resolvability in the map with higher Q-score values reflecting better resolvability. Please note: Q-score is calculating the resolvability of atoms, and thus high values are only expected at resolutions 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 (0.15).

9.4 Atom inclusion [i](#)



At the recommended contour level, 87% of all backbone atoms, 78% 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 (0.15) and Q-score for the entire model and for each chain.

Chain	Atom inclusion	Q-score
All	<div></div> 0.7810	<div></div> 0.5400
A	<div></div> 0.7900	<div></div> 0.5430
B	<div></div> 0.7890	<div></div> 0.5400
C	<div></div> 0.5240	<div></div> 0.4370
D	<div></div> 0.5310	<div></div> 0.4320
E	<div></div> 0.5310	<div></div> 0.4390
F	<div></div> 0.8040	<div></div> 0.6020
G	<div></div> 0.7890	<div></div> 0.5410
H	<div></div> 0.7930	<div></div> 0.6020
I	<div></div> 0.7890	<div></div> 0.5420
J	<div></div> 0.7890	<div></div> 0.6020
K	<div></div> 0.5280	<div></div> 0.4400
O	<div></div> 0.8020	<div></div> 0.6030

