



# Full wwPDB X-ray Structure Validation Report ⓘ

Jan 31, 2016 – 06:34 PM GMT

PDB ID : 1BG3  
Title : RAT BRAIN HEXOKINASE TYPE I COMPLEX WITH GLUCOSE AND  
INHIBITOR GLUCOSE-6-PHOSPHATE  
Authors : Mulichak, A.M.; Garavito, R.M.  
Deposited on : 1998-06-04  
Resolution : 2.80 Å(reported)

This is a Full wwPDB X-ray Structure 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  
<http://wwpdb.org/validation/2016/XrayValidationReportHelp>  
with specific help available everywhere you see the ⓘ symbol.

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

MolProbity : 4.02b-467  
Mogul : 1.7 (RC4), CSD as536be (2015)  
Xtriage (Phenix) : 1.9-1692  
EDS : rb-20026688  
Percentile statistics : 20151230.v01 (using entries in the PDB archive December 30th 2015)  
Refmac : 5.8.0135  
CCP4 : 6.5.0  
Ideal geometry (proteins) : Engh & Huber (2001)  
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)  
Validation Pipeline (wwPDB-VP) : trunk26865

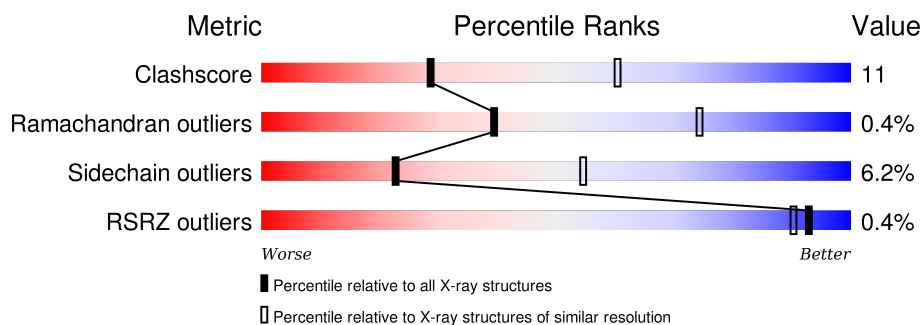
# 1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

*X-RAY DIFFRACTION*

The reported resolution of this entry is 2.80 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)	Similar resolution (#Entries, resolution range(Å))
Clashscore	102246	2827 (2.80-2.80)
Ramachandran outliers	100387	2782 (2.80-2.80)
Sidechain outliers	100360	2784 (2.80-2.80)
RSRZ outliers	91569	2404 (2.80-2.80)

The table below summarises the geometric issues observed across the polymeric chains and their fit to the electron density. The red, orange, yellow and green segments on the lower bar indicate the fraction of residues that contain outliers for  $\geq 3$ , 2, 1 and 0 types of geometric quality criteria. 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 electron density. The numeric value is given above the bar.

Mol	Chain	Length	Quality of chain
1	A	918	
1	B	918	

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

Mol	Type	Chain	Res	Chirality	Geometry	Clashes	Electron density
3	G6P	A	1002	X	-	-	-
3	G6P	A	1004	X	-	-	-

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Mol	Type	Chain	Res	Chirality	Geometry	Clashes	Electron density
3	G6P	B	1002	X	-	-	-
3	G6P	B	1004	X	-	-	-

## 2 Entry composition [i](#)

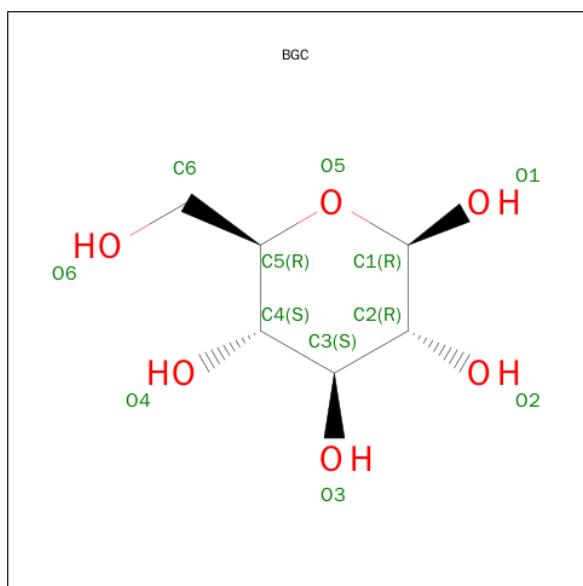
There are 5 unique types of molecules in this entry. The entry contains 13962 atoms, of which 0 are hydrogens and 0 are deuteriums.

In the tables below, the ZeroOcc column contains the number of atoms modelled with zero occupancy, 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 HEXOKINASE.

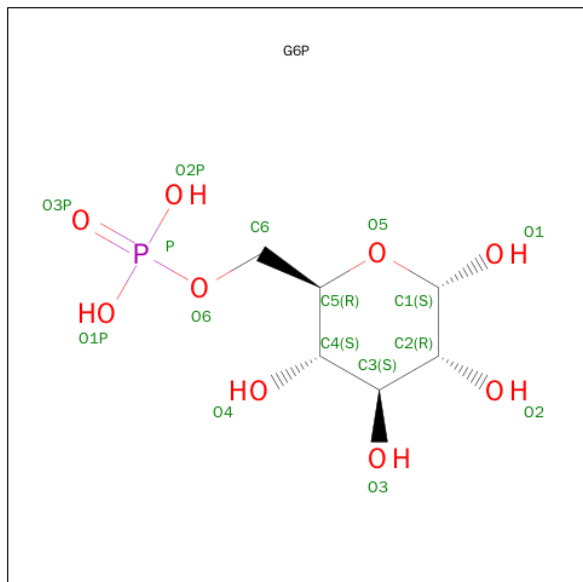
Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
1	A	902	Total	C	N	O	S	0	0	0
			6832	4304	1177	1296	55			
1	B	902	Total	C	N	O	S	0	0	0
			6783	4276	1157	1295	55			

- Molecule 2 is SUGAR (GLUCOSE) (three-letter code: BGC) (formula:  $C_6H_{12}O_6$ ).



Mol	Chain	Residues	Atoms			ZeroOcc	AltConf
2	A	1	Total	C	O	0	0
			12	6	6		
2	A	1	Total	C	O	0	0
			12	6	6		
2	B	1	Total	C	O	0	0
			12	6	6		
2	B	1	Total	C	O	0	0
			12	6	6		

- Molecule 3 is SUGAR (ALPHA-D-GLUCOSE-6-PHOSPHATE) (three-letter code: G6P) (formula:  $C_6H_{13}O_9P$ ).



Mol	Chain	Residues	Atoms				ZeroOcc	AltConf
3	A	1	Total	C	O	P	0	0
			16	6	9	1		
3	A	1	Total	C	O	P	0	0
			16	6	9	1		
3	B	1	Total	C	O	P	0	0
			16	6	9	1		
3	B	1	Total	C	O	P	0	0
			16	6	9	1		

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

Mol	Chain	Residues	Atoms		ZeroOcc	AltConf
4	A	1	Total	Ca	0	0
			1	1		

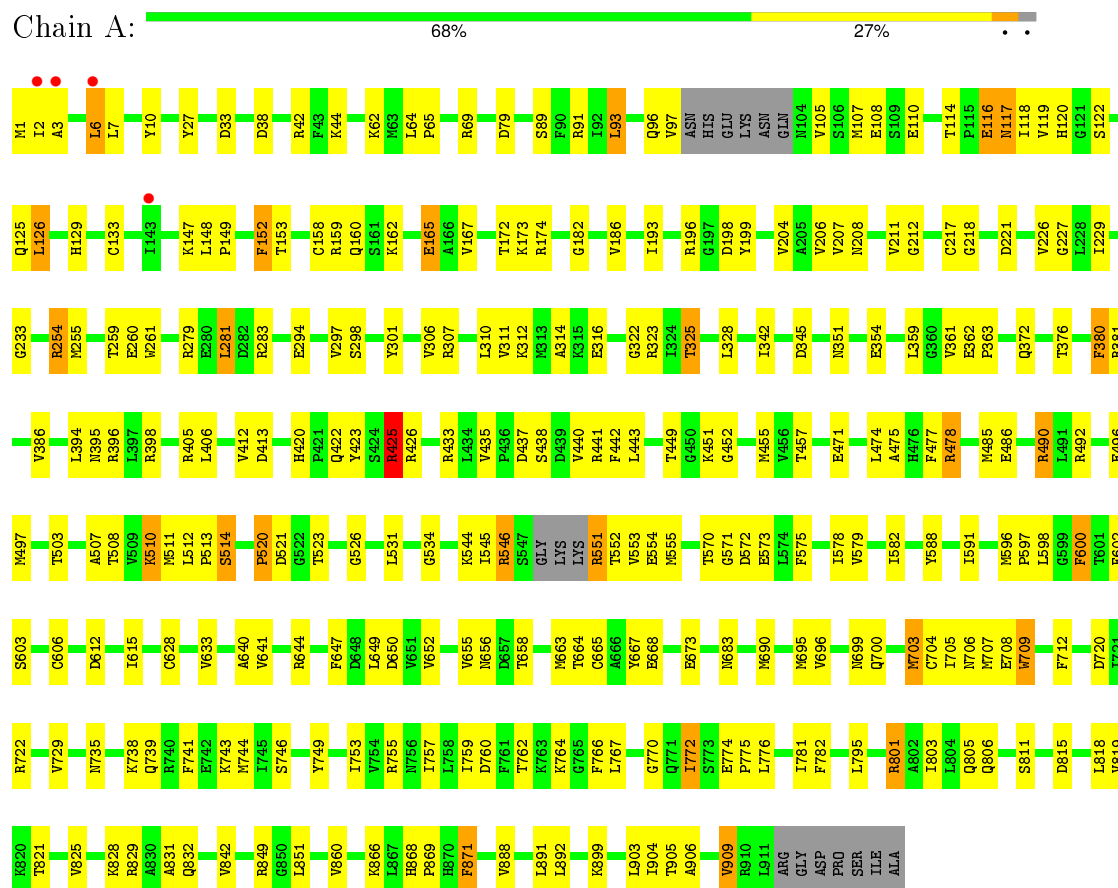
- Molecule 5 is water.

Mol	Chain	Residues	Atoms		ZeroOcc	AltConf
5	A	136	Total	O	0	0
			136	136		
5	B	98	Total	O	0	0
			98	98		

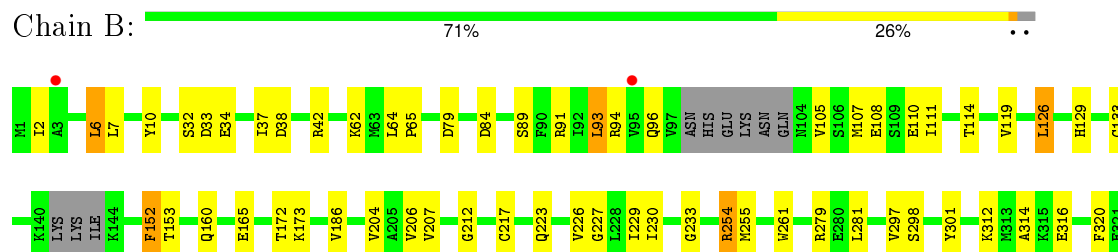
### 3 Residue-property plots

These plots are drawn for all protein, RNA and DNA chains in the entry. The first graphic for a chain summarises the proportions of errors displayed in the second graphic. The second graphic shows the sequence view annotated by issues in geometry and electron 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 dot above a residue indicates a poor fit to the electron density ( $\text{RSRZ} > 2$ ). 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: HEXOKINASE



#### • Molecule 1: HEXOKINASE



L892	L893	L894	L895	L896	L897	L898	L899	L900	L901	L902	L903	L904	L905	L906	L907	L908	L909	L910	L911	ARG	GLY	ASP	PRO	SER	ILE	ALA												
I757	I758	I759	I760	I761	I762	I763	I764	I765	I766	I767	I768	I769	I770	I771	I772	I773	I774	I775	I776	I777	I778	I779	I780	I781	I782	I783	I784	I785	I786	I787	I788							
V651	V652	V653	V654	V655	V656	V657	V658	V659	V660	V661	V662	V663	V664	V665	V666	V667	V668	V669	V670	V671	V672	V673	V674	V675	V676	V677	V678	V679	V680	V681	V682	V683						
G322	G323	G324	G325	G326	G327	G328	G329	G330	G331	G332	G333	G334	G335	G336	G337	G338	G339	G340	G341	G342	G343	G344	G345	G346	G347	G348	G349	G350	G351	G352	G353	G354						
L443	L444	L445	L446	L447	L448	L449	L450	L451	L452	L453	L454	L455	L456	L457	L458	L459	L460	L461	L462	L463	L464	L465	L466	L467	L468	L469	L470	L471	L472	L473	L474	L475						
M553	M554	M555	M556	M557	M558	M559	M560	M561	M562	M563	M564	M565	M566	M567	M568	M569	M570	M571	M572	M573	M574	M575	M576	M577	M578	M579	M580	M581	M582	M583	M584	M585						
E596	E597	E598	E599	E600	E601	E602	E603	E604	E605	E606	E607	E608	E609	E610	E611	E612	E613	E614	E615	E616	E617	E618	E619	E620	E621	E622	E623	E624	E625	E626	E627	E628						
F503	F504	F505	F506	F507	F508	F509	F510	F511	F512	F513	F514	F515	F516	F517	F518	F519	F520	F521	F522	F523	F524	F525	F526	F527	F528	F529	F530	F531	F532	F533	F534	F535						
D612	D613	D614	D615	D616	D617	D618	D619	D620	D621	D622	D623	D624	D625	D626	D627	D628	D629	D630	D631	D632	D633	D634	D635	D636	D637	D638	D639	D640	D641	D642	D643	D644						
C813	C814	C815	C816	C817	C818	C819	C820	C821	C822	C823	C824	C825	C826	C827	C828	C829	C830	C831	C832	C833	C834	C835	C836	C837	C838	C839	C840	C841	C842	C843	C844	C845						
K328	K329	K330	K331	K332	K333	K334	K335	K336	K337	K338	K339	K340	K341	K342	K343	K344	K345	K346	K347	K348	K349	K350	K351	K352	K353	K354	K355	K356	K357	K358	K359	K360						
Q832	Q833	Q834	Q835	Q836	Q837	Q838	Q839	Q840	Q841	Q842	Q843	Q844	Q845	Q846	Q847	Q848	Q849	Q850	Q851	Q852	Q853	Q854	Q855	Q856	Q857	Q858	Q859	Q860	Q861	Q862	Q863	Q864	Q865					
V825	V826	V827	V828	V829	V830	V831	V832	V833	V834	V835	V836	V837	V838	V839	V840	V841	V842	V843	V844	V845	V846	V847	V848	V849	V850	V851	V852	V853	V854	V855	V856	V857	V858					
R358	R359	R360	R361	R362	R363	R364	R365	R366	R367	R368	R369	R370	R371	R372	R373	R374	R375	R376	R377	R378	R379	R380	R381	R382	R383	R384	R385	R386	R387	R388	R389	R390	R391					
M481	M482	M483	M484	M485	M486	M487	M488	M489	M490	M491	M492	M493	M494	M495	M496	M497	M498	M499	M500	M501	M502	M503	M504	M505	M506	M507	M508	M509	M510	M511	M512	M513	M514					
A507	A508	A509	A510	A511	A512	A513	A514	A515	A516	A517	A518	A519	A520	A521	A522	A523	A524	A525	A526	A527	A528	A529	A530	A531	A532	A533	A534	A535	A536	A537	A538	A539	A540					
T508	T509	T510	T511	T512	T513	T514	T515	T516	T517	T518	T519	T520	T521	T522	T523	T524	T525	T526	T527	T528	T529	T530	T531	T532	T533	T534	T535	T536	T537	T538	T539	T540	T541					
S514	S515	S516	S517	S518	S519	S520	S521	S522	S523	S524	S525	S526	S527	S528	S529	S530	S531	S532	S533	S534	S535	S536	S537	S538	S539	S540	S541	S542	S543	S544	S545	S546	S547					
P520	P521	P522	P523	P524	P525	P526	P527	P528	P529	P530	P531	P532	P533	P534	P535	P536	P537	P538	P539	P540	P541	P542	P543	P544	P545	P546	P547	P548	P549	P550	P551	P552	P553					
H420	H421	H422	H423	H424	H425	H426	H427	H428	H429	H430	H431	H432	H433	H434	H435	H436	H437	H438	H439	H440	H441	H442	H443	H444	H445	H446	H447	H448	H449	H450	H451	H452	H453	H454				
Y423	Y424	Y425	Y426	Y427	Y428	Y429	Y430	Y431	Y432	Y433	Y434	Y435	Y436	Y437	Y438	Y439	Y440	Y441	Y442	Y443	Y444	Y445	Y446	Y447	Y448	Y449	Y450	Y451	Y452	Y453	Y454	Y455	Y456	Y457				
R426	R427	R428	R429	R430	R431	R432	R433	R434	R435	R436	R437	R438	R439	R440	R441	R442	R443	R444	R445	R446	R447	R448	R449	R450	R451	R452	R453	R454	R455	R456	R457	R458	R459	R460	R461			
L541	L542	L543	L544	L545	L546	L547	L548	L549	L550	L551	L552	L553	L554	L555	L556	L557	L558	L559	L560	L561	L562	L563	L564	L565	L566	L567	L568	L569	L570	L571	L572	L573	L574	L575	L576			
F421	F422	F423	F424	F425	F426	F427	F428	F429	F430	F431	F432	F433	F434	F435	F436	F437	F438	F439	F440	F441	F442	F443	F444	F445	F446	F447	F448	F449	F450	F451	F452	F453	F454	F455	F456			
Q421	Q422	Q423	Q424	Q425	Q426	Q427	Q428	Q429	Q430	Q431	Q432	Q433	Q434	Q435	Q436	Q437	Q438	Q439	Q440	Q441	Q442	Q443	Q444	Q445	Q446	Q447	Q448	Q449	Q450	Q451	Q452	Q453	Q454	Q455	Q456			
V412	V413	V414	V415	V416	V417	V418	V419	V420	V421	V422	V423	V424	V425	V426	V427	V428	V429	V430	V431	V432	V433	V434	V435	V436	V437	V438	V439	V440	V441	V442	V443	V444	V445	V446	V447			
D413	D414	D415	D416	D417	D418	D419	D420	D421	D422	D423	D424	D425	D426	D427	D428	D429	D430	D431	D432	D433	D434	D435	D436	D437	D438	D439	D440	D441	D442	D443	D444	D445	D446	D447	D448			
L416	L417	L418	L419	L420	L421	L422	L423	L424	L425	L426	L427	L428	L429	L430	L431	L432	L433	L434	L435	L436	L437	L438	L439	L440	L441	L442	L443	L444	L445	L446	L447	L448	L449	L450	L451			
H420	H421	H422	H423	H424	H425	H426	H427	H428	H429	H430	H431	H432	H433	H434	H435	H436	H437	H438	H439	H440	H441	H442	H443	H444	H445	H446	H447	H448	H449	H450	H451	H452	H453	H454	H455	H456		
P421	P422	P423	P424	P425	P426	P427	P428	P429	P430	P431	P432	P433	P434	P435	P436	P437	P438	P439	P440	P441	P442	P443	P444	P445	P446	P447	P448	P449	P450	P451	P452	P453	P454	P455	P456	P457		
Q421	Q422	Q423	Q424	Q425	Q426	Q427	Q428	Q429	Q430	Q431	Q432	Q433	Q434	Q435	Q436	Q437	Q438	Q439	Q440	Q441	Q442	Q443	Q444	Q445	Q446	Q447	Q448	Q449	Q450	Q451	Q452	Q453	Q454	Q455	Q456	Q457	Q458	
Y423	Y424	Y425	Y426	Y427	Y428	Y429	Y430	Y431	Y432	Y433	Y434	Y435	Y436	Y437	Y438	Y439	Y440	Y441	Y442	Y443	Y444	Y445	Y446	Y447	Y448	Y449	Y450	Y451	Y452	Y453	Y454	Y455	Y456	Y457	Y458	Y459	Y460	
R426	R427	R428	R429	R430	R431	R432	R433	R434	R435	R436	R437	R438	R439	R440	R441	R442	R443	R444	R445	R446	R447	R448	R449	R450	R451	R452	R453	R454	R455	R456	R457	R458	R459	R460	R461	R462	R463	
L541	L542	L543	L544	L545	L546	L547	L548	L549	L550	L551	L552	L553	L554	L555	L556	L557	L558	L559	L560	L561	L562	L563	L564	L565	L566	L567	L568	L569	L570	L571	L572	L573	L574	L575	L576	L577	L578	
F421	F422	F423	F424	F425	F426	F427	F428	F429	F430	F431	F432	F433	F434	F435	F436	F437	F438	F439	F440	F441	F442	F443	F444	F445	F446	F447	F448	F449	F450	F451	F452	F453	F454	F455	F456	F457	F458	
Q421	Q422	Q423	Q424	Q425	Q426	Q427	Q428	Q429	Q430	Q431	Q432	Q433	Q434	Q435	Q436	Q437	Q438	Q439	Q440	Q441	Q442	Q443	Q444	Q445	Q446	Q447	Q448	Q449	Q450	Q451	Q452	Q453	Q454	Q455	Q456	Q457	Q458	
V412	V413	V414	V415	V416	V417	V418	V419	V420	V421	V422	V423	V424	V425	V426	V427	V428	V429	V430	V431	V432	V433	V434	V435	V436	V437	V438	V439	V440	V441	V442	V443	V444	V445	V446	V447	V448	V449	V450
D413	D414	D415	D416	D417	D418	D419	D420	D421	D422	D423	D424	D425	D426	D427	D428	D429	D430	D431	D432	D433	D434	D435	D436	D437	D438	D439	D440	D441	D442	D443	D444	D445	D446	D447	D448	D449	D450	
L416	L417	L418	L419	L420	L421	L422	L423	L424	L425	L426	L427	L428	L429	L430	L431	L432	L433	L434	L435	L436	L437	L438	L439	L440	L441	L442	L443	L444	L445	L446	L447	L448	L449	L450	L451	L452	L453	
H420	H421	H422	H423	H424	H425	H426	H427	H428	H429	H430	H431	H432	H433	H434	H435	H436	H437	H438	H439	H440	H441	H442	H443	H444	H445	H446	H447	H448	H449	H450	H451	H452	H453	H454	H455	H456	H457	
P421	P422	P423	P424	P425	P426	P427	P428	P429	P430	P431	P432	P433	P434	P435	P436	P437	P438	P439	P440	P441	P442	P443	P444	P445	P446	P447	P448	P449	P450	P451	P452	P453	P454	P455	P456	P457	P458	
Q421	Q422	Q423	Q424	Q425	Q426	Q427	Q428	Q429	Q430	Q431	Q432	Q433	Q434	Q435	Q436	Q437	Q438	Q439	Q440	Q441	Q442	Q443	Q444	Q445	Q446	Q447	Q448	Q449	Q450	Q451	Q452	Q453	Q454	Q455	Q456	Q457	Q458	
Y423	Y424	Y425	Y426	Y																																		

## 4 Data and refinement statistics

Property	Value	Source
Space group	P 1 21 1	Depositor
Cell constants a, b, c, $\alpha$ , $\beta$ , $\gamma$	132.10Å 77.10Å 137.10Å 90.00° 96.00° 90.00°	Depositor
Resolution (Å)	20.00 – 2.80 38.48 – 2.80	Depositor EDS
% Data completeness (in resolution range)	71.0 (20.00-2.80) 70.6 (38.48-2.80)	Depositor EDS
$R_{merge}$	0.08	Depositor
$R_{sym}$	(Not available)	Depositor
$\langle I/\sigma(I) \rangle$ <sup>1</sup>	2.83 (at 2.81Å)	Xtriage
Refinement program	X-PLOR 3.1	Depositor
R, $R_{free}$	0.196 , 0.251 0.195 , (Not available)	Depositor DCC
$R_{free}$ test set	No test flags present.	DCC
Wilson B-factor (Å <sup>2</sup> )	35.9	Xtriage
Anisotropy	0.426	Xtriage
Bulk solvent $k_{sol}$ (e/Å <sup>3</sup> ), $B_{sol}$ (Å <sup>2</sup> )	0.32 , 74.7	EDS
Estimated twinning fraction	0.045 for l,-k,h	Xtriage
L-test for twinning <sup>2</sup>	$\langle  L  \rangle = 0.43$ , $\langle L^2 \rangle = 0.25$	Xtriage
Outliers	0 of 48013 reflections	Xtriage
$F_o, F_c$ correlation	0.90	EDS
Total number of atoms	13962	wwPDB-VP
Average B, all atoms (Å <sup>2</sup> )	34.0	wwPDB-VP

Xtriage's analysis on translational NCS is as follows: *The largest off-origin peak in the Patterson function is 3.18% of the height of the origin peak. No significant pseudotranslation is detected.*

<sup>1</sup>Intensities estimated from amplitudes.

<sup>2</sup>Theoretical values of  $\langle |L| \rangle$ ,  $\langle L^2 \rangle$  for acentric reflections are 0.5, 0.375 respectively for untwinned datasets, and 0.333, 0.2 for perfectly twinned datasets.



## 5 Model quality

### 5.1 Standard geometry

Bond lengths and bond angles in the following residue types are not validated in this section: CA, BGC, G6P

The Z score for a bond length (or angle) is the number of standard deviations the observed value is removed from the expected value. A bond length (or angle) with  $|Z| > 5$  is considered an outlier worth inspection. RMSZ is the root-mean-square of all Z scores of the bond lengths (or angles).

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	# $ Z  > 5$	RMSZ	# $ Z  > 5$
1	A	0.62	1/6935 (0.0%)	0.89	15/9367 (0.2%)
1	B	0.62	0/6885	0.90	15/9310 (0.2%)
All	All	0.62	1/13820 (0.0%)	0.90	30/18677 (0.2%)

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

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

All (1) bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
1	A	158	CYS	CB-SG	-5.23	1.73	1.81

All (30) bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	A	425	ARG	NE-CZ-NH2	-14.98	112.81	120.30
1	B	279	ARG	NE-CZ-NH2	-14.97	112.81	120.30
1	A	279	ARG	NE-CZ-NH2	-14.74	112.93	120.30
1	B	425	ARG	NE-CZ-NH2	-14.09	113.26	120.30
1	B	279	ARG	NE-CZ-NH1	14.07	127.34	120.30
1	B	381	ARG	NE-CZ-NH2	-14.07	113.27	120.30
1	A	381	ARG	NE-CZ-NH1	13.93	127.26	120.30
1	A	381	ARG	NE-CZ-NH2	-13.71	113.44	120.30
1	B	381	ARG	NE-CZ-NH1	13.63	127.12	120.30
1	A	254	ARG	NE-CZ-NH1	13.56	127.08	120.30

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	A	279	ARG	NE-CZ-NH1	12.88	126.74	120.30
1	A	254	ARG	NE-CZ-NH2	-12.60	114.00	120.30
1	B	254	ARG	NE-CZ-NH2	-12.57	114.02	120.30
1	B	425	ARG	NE-CZ-NH1	11.87	126.23	120.30
1	B	254	ARG	NE-CZ-NH1	11.44	126.02	120.30
1	A	425	ARG	NE-CZ-NH1	10.96	125.78	120.30
1	A	490	ARG	NE-CZ-NH1	7.39	123.99	120.30
1	A	254	ARG	CD-NE-CZ	7.34	133.88	123.60
1	B	279	ARG	CD-NE-CZ	6.90	133.26	123.60
1	A	381	ARG	CD-NE-CZ	6.55	132.77	123.60
1	A	279	ARG	CD-NE-CZ	6.48	132.67	123.60
1	B	514	SER	N-CA-C	-6.26	94.11	111.00
1	A	514	SER	N-CA-C	-6.24	94.14	111.00
1	A	425	ARG	CD-NE-CZ	6.11	132.15	123.60
1	B	254	ARG	CG-CD-NE	5.98	124.35	111.80
1	B	425	ARG	CD-NE-CZ	5.98	131.97	123.60
1	B	381	ARG	CD-NE-CZ	5.72	131.61	123.60
1	B	254	ARG	CB-CG-CD	-5.58	97.09	111.60
1	A	281	LEU	CA-CB-CG	5.15	127.14	115.30
1	B	254	ARG	CD-NE-CZ	5.08	130.72	123.60

There are no chirality outliers.

All (1) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	A	490	ARG	Sidechain

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

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

Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
1	A	6832	0	6711	177	0
1	B	6783	0	6623	136	0
2	A	24	0	23	2	0
2	B	24	0	24	1	0
3	A	32	0	22	0	0
3	B	32	0	22	0	0

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Mol	Chain	Non-H	H(model)	H(added)	Clashes	Symm-Clashes
4	A	1	0	0	0	0
5	A	136	0	0	6	0
5	B	98	0	0	0	0
All	All	13962	0	13425	311	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 11.

All (311) close contacts within the same asymmetric unit are listed below, sorted by their clash magnitude.

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:A:361:VAL:O	1:A:363:PRO:HD3	1.62	1.00
1:B:361:VAL:O	1:B:363:PRO:HD3	1.67	0.93
1:A:312:LYS:O	1:A:316:GLU:HG3	1.80	0.81
1:A:162:LYS:HG3	1:A:165:GLU:HB2	1.62	0.81
1:A:849:ARG:HB2	1:A:851:LEU:HD11	1.62	0.81
1:A:69:ARG:HD3	5:A:2044:HOH:O	1.81	0.78
1:A:477:PHE:HB2	1:A:825:VAL:HG12	1.65	0.76
1:B:397:LEU:O	1:B:401:LYS:HG3	1.85	0.76
1:B:906:ALA:O	1:B:909:VAL:HG12	1.85	0.76
1:B:204:VAL:HG22	1:B:457:THR:HG23	1.68	0.75
1:B:477:PHE:HB2	1:B:825:VAL:HG12	1.68	0.75
1:A:520:PRO:HD3	1:A:663:MET:SD	2.27	0.74
1:B:658:THR:HG21	1:B:683:ASN:HD22	1.53	0.73
1:A:212:GLY:HA3	1:A:449:THR:HG23	1.70	0.73
1:A:97:VAL:HG13	1:A:105:VAL:HG22	1.71	0.72
1:B:520:PRO:HD3	1:B:663:MET:SD	2.29	0.72
1:B:212:GLY:HA3	1:B:449:THR:HG23	1.71	0.72
1:A:578:ILE:O	1:A:582:ILE:HG13	1.89	0.72
1:A:172:THR:HG22	1:A:173:LYS:HG3	1.70	0.72
1:A:571:GLY:HA2	1:A:628:CYS:SG	2.30	0.72
1:A:652:VAL:HB	1:A:905:THR:HG23	1.71	0.72
1:A:762:THR:O	1:A:770:GLY:HA2	1.91	0.71
1:A:2:ILE:O	1:A:6:LEU:HB2	1.91	0.70
1:A:658:THR:HG21	1:A:683:ASN:HD22	1.56	0.70
1:B:172:THR:HG22	1:B:173:LYS:HG3	1.71	0.70
1:B:545:ILE:O	1:B:546:ARG:HD3	1.92	0.70
1:A:665:CYS:SG	1:A:891:LEU:HD23	2.34	0.68
1:B:380:PHE:CE1	1:B:426:ARG:HB3	2.28	0.68
1:B:578:ILE:O	1:B:582:ILE:HG13	1.95	0.67
1:A:159:ARG:HB2	1:A:167:VAL:HB	1.77	0.67

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:89:SER:HB2	1:B:91:ARG:NH1	2.10	0.67
1:A:89:SER:HB2	1:A:91:ARG:NH1	2.11	0.66
1:B:107:MET:SD	1:B:451:LYS:HB2	2.36	0.65
1:B:204:VAL:CG2	1:B:457:THR:HG23	2.25	0.65
1:A:545:ILE:O	1:A:546:ARG:HD3	1.97	0.65
1:A:118:ILE:HG23	1:A:126:LEU:HA	1.77	0.65
1:B:481:LYS:O	1:B:485:MET:HB2	1.96	0.64
1:A:849:ARG:HB2	1:A:851:LEU:CD1	2.25	0.64
1:A:204:VAL:HG22	1:A:457:THR:HG23	1.79	0.64
1:A:435:VAL:HG12	1:A:438:SER:OG	1.97	0.64
1:A:325:THR:HG21	1:A:359:LEU:O	1.97	0.63
1:A:906:ALA:O	1:A:909:VAL:HG12	1.98	0.63
1:B:571:GLY:HA2	1:B:628:CYS:SG	2.38	0.63
1:A:695:MET:HG2	5:A:2029:HOH:O	1.97	0.62
1:A:570:THR:HG22	1:A:572:ASP:H	1.65	0.62
1:B:217:CYS:HB3	1:B:443:LEU:HD23	1.81	0.62
1:A:755:ARG:HD3	1:A:781:ILE:HG22	1.81	0.62
1:B:501:LYS:HG2	1:B:695:MET:SD	2.40	0.61
1:B:828:LYS:O	1:B:832:GLN:HG3	2.01	0.61
1:B:762:THR:O	1:B:770:GLY:HA2	2.00	0.61
1:B:2:ILE:O	1:B:6:LEU:HB2	2.00	0.60
1:B:325:THR:HG21	1:B:359:LEU:O	2.01	0.60
1:B:866:LYS:HE3	1:B:892:LEU:HD11	1.82	0.59
1:A:204:VAL:CG2	1:A:457:THR:HG23	2.31	0.59
1:A:866:LYS:HE3	1:A:892:LEU:HD11	1.84	0.59
1:A:351:ASN:HA	1:A:354:GLU:HG2	1.84	0.59
1:A:298:SER:HB3	1:A:301:TYR:HD2	1.68	0.58
1:A:477:PHE:HB2	1:A:825:VAL:CG1	2.34	0.58
1:A:545:ILE:HG23	1:A:553:VAL:HG22	1.85	0.58
1:A:767:LEU:HD13	1:A:818:LEU:HD23	1.85	0.58
1:B:569:GLY:O	1:B:625:ALA:HA	2.04	0.58
1:B:605:PRO:HB2	1:B:617:ILE:HB	1.87	0.57
1:A:591:ILE:HG21	1:A:596:MET:SD	2.44	0.57
1:A:507:ALA:O	1:A:510:LYS:HE3	2.03	0.57
1:B:520:PRO:HG2	1:B:903:LEU:HD13	1.86	0.56
1:A:570:THR:HB	1:A:573:GLU:H	1.69	0.56
1:B:542:LEU:HD13	1:B:591:ILE:CD1	2.35	0.56
1:B:298:SER:HB3	1:B:301:TYR:HD2	1.68	0.56
1:A:196:ARG:HG2	1:A:198:ASP:HB2	1.86	0.56
1:A:107:MET:SD	1:A:451:LYS:HB2	2.45	0.56
1:B:65:PRO:HB3	1:B:255:MET:HE3	1.87	0.56

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:570:THR:OG1	1:B:573:GLU:HG3	2.06	0.56
1:B:767:LEU:HD13	1:B:818:LEU:HD23	1.88	0.56
1:A:755:ARG:O	1:A:759:ILE:HG13	2.06	0.55
1:B:755:ARG:HD3	1:B:781:ILE:HG22	1.87	0.55
1:A:849:ARG:CB	1:A:851:LEU:HD11	2.34	0.55
1:A:652:VAL:CB	1:A:905:THR:HG23	2.37	0.55
1:B:507:ALA:O	1:B:510:LYS:HE3	2.06	0.55
1:B:114:THR:HG22	1:B:119:VAL:HG23	1.88	0.55
1:A:2:ILE:O	1:A:6:LEU:HD23	2.07	0.54
1:B:223:GLN:OE1	1:B:441:ARG:NH2	2.40	0.54
1:A:712:PHE:O	1:A:741:PHE:HB2	2.07	0.54
1:B:91:ARG:HA	1:B:110:GLU:O	2.08	0.54
1:B:477:PHE:HB2	1:B:825:VAL:CG1	2.36	0.54
1:B:755:ARG:O	1:B:759:ILE:HG13	2.07	0.54
1:A:570:THR:HG22	1:A:571:GLY:N	2.23	0.54
1:A:114:THR:HG22	1:A:119:VAL:HG23	1.90	0.54
1:B:544:LYS:HB2	1:B:554:GLU:HB3	1.90	0.53
1:A:544:LYS:HB2	1:A:554:GLU:HB3	1.91	0.53
1:B:531:LEU:HD12	1:B:600:PHE:CD1	2.44	0.53
1:B:579:VAL:HG13	1:B:647:PHE:HZ	1.73	0.53
1:B:62:LYS:HB3	1:B:64:LEU:HG	1.90	0.53
1:A:602:PHE:CE2	1:A:633:VAL:HG11	2.44	0.53
1:B:545:ILE:HD13	1:B:903:LEU:HD23	1.90	0.53
1:A:91:ARG:HA	1:A:110:GLU:O	2.09	0.53
1:A:811:SER:OG	1:A:815:ASP:HB2	2.08	0.53
1:A:152:PHE:HB3	1:A:206:VAL:HG22	1.91	0.53
1:A:591:ILE:CG2	1:A:596:MET:SD	2.97	0.52
1:B:422:GLN:O	1:B:426:ARG:HG3	2.09	0.52
1:B:354:GLU:O	1:B:358:ARG:HG3	2.09	0.52
1:A:147:LYS:O	1:A:148:LEU:HD23	2.10	0.52
1:A:665:CYS:O	1:A:668:GLU:HG2	2.10	0.52
1:B:323:ARG:NH1	1:B:362:GLU:H	2.08	0.52
1:B:815:ASP:O	1:B:819:VAL:HG23	2.09	0.51
1:A:376:THR:HG23	1:A:426:ARG:HH12	1.75	0.51
1:B:579:VAL:HG13	1:B:647:PHE:CZ	2.45	0.51
1:A:551:ARG:NH2	1:A:667:TYR:O	2.42	0.51
1:B:226:VAL:HG12	1:B:227:GLY:N	2.25	0.51
1:A:531:LEU:HD12	1:A:600:PHE:CD1	2.45	0.51
1:B:38:ASP:O	1:B:42:ARG:HG3	2.10	0.51
1:B:152:PHE:HB3	1:B:206:VAL:HG22	1.92	0.51
1:A:193:ILE:HG23	1:A:199:TYR:CE1	2.45	0.51

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:A:690:MET:HE3	1:A:703:MET:HB2	1.93	0.51
1:A:441:ARG:HG2	1:A:442:PHE:N	2.25	0.51
1:A:323:ARG:NH1	1:A:362:GLU:H	2.08	0.51
1:A:62:LYS:HB3	1:A:64:LEU:HG	1.93	0.51
1:A:116:GLU:HG2	1:A:120:HIS:ND1	2.25	0.51
1:B:606:CYS:HA	1:B:615:ILE:O	2.10	0.51
1:B:325:THR:HG23	1:B:328:LEU:HB2	1.93	0.51
1:A:93:LEU:N	1:A:93:LEU:HD12	2.26	0.51
1:A:673:GLU:CD	1:A:849:ARG:HH22	2.14	0.51
1:B:7:LEU:O	1:B:10:TYR:HB3	2.11	0.51
1:A:217:CYS:HB3	1:A:443:LEU:HD23	1.92	0.51
1:B:93:LEU:HD12	1:B:93:LEU:N	2.26	0.50
1:A:186:VAL:HG22	1:A:206:VAL:HG21	1.93	0.50
1:B:553:VAL:HG11	1:B:899:LYS:HG3	1.93	0.50
1:A:606:CYS:HA	1:A:615:ILE:O	2.12	0.50
1:A:65:PRO:HB3	1:A:255:MET:HE3	1.93	0.50
1:B:93:LEU:HA	1:B:108:GLU:O	2.12	0.50
1:B:172:THR:HG21	2:B:1001:BGC:O1	2.12	0.50
1:B:105:VAL:HG11	1:B:451:LYS:HG3	1.94	0.50
1:A:33:ASP:OD2	1:A:433:ARG:NH1	2.45	0.50
1:B:129:HIS:CE1	1:B:133:CYS:SG	3.05	0.49
1:B:534:GLY:HA3	1:B:603:SER:HB2	1.94	0.49
1:B:541:LEU:HG	1:B:557:ASN:HB3	1.95	0.49
1:B:521:ASP:O	1:B:523:THR:HG23	2.13	0.49
1:A:441:ARG:HG2	1:A:442:PHE:H	1.76	0.49
1:B:712:PHE:O	1:B:741:PHE:HB2	2.12	0.49
1:B:186:VAL:HG22	1:B:206:VAL:HG21	1.94	0.49
1:A:1:MET:C	1:A:3:ALA:N	2.64	0.49
1:B:664:THR:HG23	1:B:899:LYS:HD2	1.93	0.49
1:A:521:ASP:O	1:A:523:THR:HG23	2.13	0.49
1:A:422:GLN:O	1:A:426:ARG:HG3	2.13	0.49
1:A:597:PRO:HA	1:A:650:ASP:HB3	1.95	0.49
1:A:520:PRO:HG2	1:A:903:LEU:HD13	1.95	0.49
1:A:579:VAL:HG13	1:A:647:PHE:CZ	2.48	0.49
1:B:614:GLY:O	1:B:633:VAL:HG22	2.13	0.49
1:A:1:MET:C	1:A:3:ALA:H	2.15	0.49
1:B:545:ILE:HG22	1:B:546:ARG:N	2.28	0.48
1:A:534:GLY:HA3	1:A:603:SER:HB2	1.95	0.48
1:B:312:LYS:O	1:B:316:GLU:HG3	2.13	0.48
1:B:690:MET:CE	1:B:703:MET:HB2	2.44	0.48
1:A:380:PHE:CE1	1:A:426:ARG:HB3	2.48	0.48

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:A:105:VAL:HG11	1:A:451:LYS:HG3	1.96	0.48
1:A:325:THR:HG23	1:A:328:LEU:HB2	1.96	0.48
1:A:511:MET:HB3	1:A:705:ILE:HG21	1.96	0.48
1:A:160:GLN:HG2	1:A:165:GLU:O	2.13	0.48
1:B:602:PHE:CE2	1:B:633:VAL:HG11	2.48	0.48
1:A:815:ASP:O	1:A:819:VAL:HG23	2.14	0.48
1:B:722:ARG:HG2	1:B:744:MET:CE	2.44	0.48
1:A:118:ILE:HG21	1:A:129:HIS:HD2	1.79	0.47
1:A:307:ARG:O	1:A:311:VAL:HG23	2.13	0.47
1:B:870:HIS:ND1	1:B:873:ARG:NH2	2.62	0.47
1:A:739:GLN:HG3	5:A:2017:HOH:O	2.14	0.47
1:A:172:THR:HG21	2:A:1001:BGC:O1	2.13	0.47
1:B:380:PHE:CD1	1:B:426:ARG:HD3	2.49	0.47
1:A:656:ASN:HB2	5:A:2111:HOH:O	2.13	0.47
1:A:153:THR:HA	1:A:207:VAL:O	2.15	0.47
1:B:683:ASN:OD1	1:B:708:GLU:HA	2.15	0.47
1:B:65:PRO:HB3	1:B:255:MET:CE	2.44	0.47
1:B:33:ASP:OD2	1:B:433:ARG:NH1	2.48	0.47
1:B:297:VAL:HG11	1:B:386:VAL:CG2	2.45	0.47
1:A:118:ILE:HG21	1:A:129:HIS:CD2	2.50	0.47
1:B:690:MET:HE3	1:B:703:MET:HB2	1.95	0.47
1:A:7:LEU:O	1:A:10:TYR:HB3	2.14	0.47
1:B:652:VAL:HB	1:B:905:THR:HG23	1.95	0.47
1:A:394:LEU:HD13	1:A:440:VAL:HG21	1.97	0.47
1:B:828:LYS:HG3	1:B:874:ILE:HD13	1.96	0.47
1:B:511:MET:HB3	1:B:705:ILE:HG21	1.97	0.47
1:A:664:THR:HG23	1:A:899:LYS:HD2	1.97	0.47
1:A:452:GLY:O	1:A:455:MET:HB2	2.15	0.47
1:A:722:ARG:HG2	1:A:744:MET:CE	2.45	0.47
1:B:314:ALA:O	1:B:322:GLY:HA2	2.15	0.46
1:A:229:ILE:HD13	1:A:413:ASP:HB3	1.97	0.46
1:A:314:ALA:O	1:A:322:GLY:HA2	2.16	0.46
1:A:803:ILE:O	1:A:806:GLN:HB3	2.16	0.46
1:A:755:ARG:HD2	1:A:776:LEU:O	2.16	0.46
1:A:147:LYS:C	1:A:148:LEU:HD23	2.36	0.46
1:B:452:GLY:O	1:B:455:MET:HB2	2.15	0.46
1:B:229:ILE:HD13	1:B:413:ASP:HB3	1.98	0.46
1:A:79:ASP:HA	1:A:96:GLN:HA	1.98	0.46
1:A:38:ASP:O	1:A:42:ARG:HG3	2.16	0.46
1:A:571:GLY:CA	1:A:628:CYS:SG	3.01	0.45
1:B:451:LYS:O	1:B:455:MET:HG2	2.16	0.45

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:755:ARG:HD2	1:B:776:LEU:O	2.16	0.45
1:B:160:GLN:HG2	1:B:165:GLU:O	2.16	0.45
1:A:828:LYS:HD3	5:A:2125:HOH:O	2.16	0.45
1:B:773:SER:O	1:B:777:LYS:HG3	2.16	0.45
1:A:683:ASN:OD1	1:A:708:GLU:HA	2.16	0.45
1:B:153:THR:HA	1:B:207:VAL:O	2.16	0.45
1:A:735:ASN:HB2	1:A:738:LYS:HB2	1.97	0.45
1:A:471:GLU:O	1:A:475:ALA:HB2	2.16	0.45
1:B:545:ILE:CD1	1:B:903:LEU:HD23	2.46	0.45
1:A:699:ASN:OD1	1:A:700:GLN:HG2	2.17	0.45
1:A:474:LEU:O	1:A:478:ARG:HG3	2.16	0.45
1:B:529:LEU:HD11	1:B:586:LEU:HD21	1.99	0.45
1:A:129:HIS:CE1	1:A:133:CYS:SG	3.10	0.45
1:B:526:GLY:O	1:B:545:ILE:N	2.50	0.44
1:B:542:LEU:HD13	1:B:591:ILE:HD13	1.99	0.44
1:A:757:ILE:O	1:A:760:ASP:HB3	2.17	0.44
1:A:93:LEU:HA	1:A:108:GLU:O	2.16	0.44
1:A:764:LYS:HE2	1:A:766:PHE:CZ	2.52	0.44
1:B:477:PHE:N	1:B:477:PHE:CD1	2.85	0.44
1:B:866:LYS:CE	1:B:892:LEU:HD11	2.48	0.44
1:A:602:PHE:HD2	1:A:606:CYS:HG	1.66	0.44
1:A:323:ARG:HH11	1:A:362:GLU:H	1.66	0.44
1:A:652:VAL:CG1	1:A:905:THR:HG23	2.48	0.44
1:B:688:GLU:OE1	1:B:848:ASN:ND2	2.51	0.44
1:A:406:LEU:HB3	1:A:438:SER:HB3	2.00	0.44
1:B:665:CYS:O	1:B:668:GLU:HG3	2.18	0.44
1:A:422:GLN:OE1	1:A:425:ARG:NH2	2.51	0.44
1:A:772:ILE:HG23	1:A:776:LEU:HD23	1.99	0.44
1:B:501:LYS:HE3	1:B:694:GLU:O	2.18	0.44
1:B:323:ARG:HH11	1:B:362:GLU:H	1.65	0.44
1:A:706:ASN:ND2	1:A:708:GLU:OE1	2.51	0.43
1:A:283:ARG:HH21	1:B:558:LYS:HE3	1.82	0.43
1:B:420:HIS:HB3	1:B:423:TYR:HB2	2.00	0.43
1:A:801:ARG:O	1:A:805:GLN:HG3	2.18	0.43
1:A:709:TRP:C	1:A:709:TRP:CD1	2.92	0.43
1:A:553:VAL:HG11	1:A:899:LYS:HG3	2.00	0.43
1:B:605:PRO:HD2	1:B:618:SER:O	2.19	0.43
1:A:690:MET:CE	1:A:703:MET:HB2	2.48	0.43
1:A:579:VAL:HG21	1:A:640:ALA:CB	2.48	0.43
1:A:696:VAL:HG21	1:A:703:MET:CE	2.48	0.43
1:A:497:MET:HB3	1:A:507:ALA:HB2	2.01	0.43

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:B:706:ASN:ND2	1:B:708:GLU:OE1	2.51	0.43
1:A:226:VAL:HG12	1:A:227:GLY:N	2.33	0.43
1:B:230:ILE:HG22	1:B:416:LEU:HB3	2.00	0.43
1:A:376:THR:HG23	1:A:426:ARG:NH1	2.34	0.43
1:A:64:LEU:HA	1:A:65:PRO:HD3	1.78	0.43
1:B:757:ILE:O	1:B:760:ASP:HB3	2.18	0.43
1:A:821:THR:O	1:A:825:VAL:HG23	2.19	0.43
1:A:571:GLY:O	1:A:575:PHE:HD1	2.02	0.43
1:A:297:VAL:HG11	1:A:386:VAL:CG2	2.49	0.43
1:A:398:ARG:NH1	1:A:437:ASP:OD2	2.51	0.43
1:B:525:HIS:NE2	1:B:546:ARG:HD2	2.34	0.43
1:A:526:GLY:O	1:A:545:ILE:N	2.52	0.43
1:B:233:GLY:HA2	1:B:298:SER:OG	2.19	0.43
1:B:401:LYS:C	1:B:403:THR:H	2.21	0.42
1:B:350:GLN:HG3	1:B:354:GLU:OE2	2.19	0.42
1:B:84:ASP:HA	1:B:153:THR:HB	2.01	0.42
1:A:44:LYS:HE3	1:A:395:ASN:HD22	1.83	0.42
1:B:846:ARG:CD	1:B:885:LYS:HB3	2.49	0.42
1:A:729:VAL:HG21	1:A:753:ILE:HG12	2.01	0.42
1:A:97:VAL:HG11	1:A:455:MET:HE1	2.00	0.42
1:A:828:LYS:O	1:A:832:GLN:HG3	2.20	0.42
1:A:735:ASN:O	1:A:743:LYS:NZ	2.52	0.42
1:B:37:ILE:HD13	1:B:434:LEU:HD22	2.01	0.42
1:B:689:GLU:O	1:B:693:VAL:HG23	2.19	0.42
1:A:117:ASN:O	1:A:125:GLN:NE2	2.52	0.42
1:B:571:GLY:O	1:B:575:PHE:HD1	2.03	0.42
1:B:579:VAL:HG21	1:B:640:ALA:CB	2.50	0.42
1:A:764:LYS:HD3	1:A:766:PHE:CE2	2.54	0.42
1:A:259:THR:O	1:A:260:GLU:HB2	2.20	0.42
1:B:471:GLU:O	1:B:475:ALA:HB2	2.19	0.42
1:B:32:SER:OG	1:B:34:GLU:HG2	2.20	0.42
1:A:842:VAL:HG11	1:A:888:VAL:HG22	2.02	0.42
1:A:514:SER:OG	1:A:704:CYS:HB3	2.20	0.42
1:A:294:GLU:OE2	2:A:1001:BGC:O1	2.38	0.41
1:B:729:VAL:HG21	1:B:753:ILE:HG12	2.01	0.41
1:A:746:SER:OG	1:A:749:TYR:HD2	2.03	0.41
1:A:148:LEU:HA	1:A:149:PRO:HD3	1.74	0.41
1:B:842:VAL:HG11	1:B:888:VAL:HG22	2.02	0.41
1:B:91:ARG:CG	1:B:111:ILE:HG12	2.50	0.41
1:A:233:GLY:HA2	1:A:298:SER:OG	2.20	0.41
1:B:596:MET:HA	1:B:597:PRO:HD3	1.75	0.41

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Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:A:588:TYR:CE1	1:B:331:ARG:HB2	2.55	0.41
1:A:602:PHE:CD2	1:A:606:CYS:SG	3.13	0.41
1:A:380:PHE:HZ	5:A:2205:HOH:O	2.01	0.41
1:A:655:VAL:HG11	1:A:904:ILE:HG21	2.01	0.41
1:A:65:PRO:HB3	1:A:255:MET:CE	2.51	0.41
1:B:690:MET:HG3	1:B:700:GLN:O	2.21	0.41
1:B:342:ILE:HG12	1:B:352:ALA:HA	2.03	0.41
1:A:3:ALA:O	1:A:6:LEU:N	2.54	0.41
1:A:420:HIS:HB3	1:A:423:TYR:HB2	2.02	0.41
1:B:79:ASP:HA	1:B:96:GLN:HA	2.01	0.41
1:B:853:HIS:HE1	1:B:887:THR:OG1	2.02	0.41
1:A:868:HIS:ND1	1:A:869:PRO:HD2	2.36	0.41
1:B:710:GLY:O	1:B:739:GLN:HA	2.21	0.41
1:A:118:ILE:HD12	1:A:125:GLN:HE21	1.86	0.41
1:A:477:PHE:CD1	1:A:477:PHE:N	2.88	0.41
1:A:167:VAL:HG13	1:A:182:GLY:O	2.21	0.41
1:A:781:ILE:HG23	1:A:782:PHE:CD2	2.56	0.41
1:B:737:GLY:HA2	1:B:740:ARG:HH21	1.86	0.41
1:A:306:VAL:O	1:A:310:LEU:HG	2.21	0.41
1:B:669:GLU:HA	1:B:670:PRO:HD2	1.73	0.41
1:A:512:LEU:HA	1:A:513:PRO:HD3	1.80	0.41
1:B:497:MET:HB3	1:B:507:ALA:HB2	2.03	0.41
1:A:376:THR:O	1:A:380:PHE:HB2	2.21	0.41
1:A:774:GLU:N	1:A:775:PRO:HD2	2.36	0.41
1:B:655:VAL:HG11	1:B:904:ILE:HG21	2.02	0.41
1:A:641:VAL:O	1:A:644:ARG:HB3	2.22	0.40
1:B:735:ASN:O	1:B:743:LYS:NZ	2.54	0.40
1:A:218:GLY:HA2	1:A:221:ASP:O	2.22	0.40
1:B:126:LEU:O	1:B:129:HIS:HB3	2.22	0.40
1:A:868:HIS:HA	1:A:869:PRO:HD3	1.92	0.40
1:A:342:ILE:O	1:A:372:GLN:HG3	2.20	0.40
1:A:208:ASN:HB3	1:A:211:VAL:HG23	2.02	0.40
1:A:831:ALA:HB2	1:A:871:PHE:HA	2.03	0.40
1:A:602:PHE:HD2	1:A:606:CYS:SG	2.44	0.40
1:B:882:LEU:HA	1:B:882:LEU:HD23	1.89	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 X-ray entries followed by that with respect to entries of similar resolution.

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

Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles	
1	A	896/918 (98%)	829 (92%)	65 (7%)	2 (0%)	52	84
1	B	896/918 (98%)	835 (93%)	55 (6%)	6 (1%)	26	62
All	All	1792/1836 (98%)	1664 (93%)	120 (7%)	8 (0%)	39	74

All (8) Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	B	345	ASP
1	A	345	ASP
1	A	871	PHE
1	B	871	PHE
1	B	404	PRO
1	B	651	VAL
1	B	348	GLY
1	B	593	GLY

### 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 X-ray entries followed by that with respect to entries of similar resolution.

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

Mol	Chain	Analysed	Rotameric	Outliers	Percentiles	
1	A	722/791 (91%)	676 (94%)	46 (6%)	22	52
1	B	714/791 (90%)	671 (94%)	43 (6%)	24	56
All	All	1436/1582 (91%)	1347 (94%)	89 (6%)	23	54

All (89) residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	A	6	LEU
1	A	27	TYR
1	A	93	LEU
1	A	116	GLU
1	A	117	ASN
1	A	122	SER
1	A	126	LEU
1	A	152	PHE
1	A	165	GLU
1	A	174	ARG
1	A	254	ARG
1	A	261	TRP
1	A	281	LEU
1	A	325	THR
1	A	380	PHE
1	A	396	ARG
1	A	405	ARG
1	A	412	VAL
1	A	425	ARG
1	A	478	ARG
1	A	485	MET
1	A	486	GLU
1	A	492	ARG
1	A	496	GLU
1	A	503	THR
1	A	508	THR
1	A	510	LYS
1	A	520	PRO
1	A	546	ARG
1	A	551	ARG
1	A	552	THR
1	A	555	MET
1	A	598	LEU
1	A	600	PHE
1	A	612	ASP
1	A	649	LEU
1	A	703	MET
1	A	707	MET
1	A	709	TRP
1	A	720	ASP
1	A	772	ILE
1	A	795	LEU
1	A	801	ARG

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Mol	Chain	Res	Type
1	A	829	ARG
1	A	860	VAL
1	A	909	VAL
1	B	6	LEU
1	B	93	LEU
1	B	94	ARG
1	B	126	LEU
1	B	152	PHE
1	B	254	ARG
1	B	261	TRP
1	B	281	LEU
1	B	320	PHE
1	B	325	THR
1	B	349	ILE
1	B	380	PHE
1	B	412	VAL
1	B	425	ARG
1	B	428	HIS
1	B	468	ARG
1	B	478	ARG
1	B	485	MET
1	B	486	GLU
1	B	492	ARG
1	B	496	GLU
1	B	508	THR
1	B	510	LYS
1	B	520	PRO
1	B	546	ARG
1	B	555	MET
1	B	592	LYS
1	B	598	LEU
1	B	612	ASP
1	B	626	THR
1	B	649	LEU
1	B	671	THR
1	B	703	MET
1	B	707	MET
1	B	709	TRP
1	B	720	ASP
1	B	736	SER
1	B	801	ARG
1	B	813	CYS

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Mol	Chain	Res	Type
1	B	829	ARG
1	B	844	LYS
1	B	860	VAL
1	B	911	LEU

Some sidechains can be flipped to improve hydrogen bonding and reduce clashes. All (7) such sidechains are listed below:

Mol	Chain	Res	Type
1	A	125	GLN
1	A	384	ASN
1	A	692	ASN
1	B	125	GLN
1	B	384	ASN
1	B	692	ASN
1	B	853	HIS

### 5.3.3 RNA [i](#)

There are no RNA molecules in this entry.

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

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

## 5.5 Carbohydrates [i](#)

There are no carbohydrates in this entry.

## 5.6 Ligand geometry [i](#)

Of 9 ligands modelled in this entry, 1 is monoatomic - leaving 8 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
2	BGC	A	1001	-	12,12,12	0.61	0	17,17,17	1.23	1 (5%)
3	G6P	A	1002	-	16,16,16	0.79	0	23,24,24	0.70	0
2	BGC	A	1003	-	12,12,12	0.39	0	17,17,17	0.67	1 (5%)
3	G6P	A	1004	-	16,16,16	0.71	0	23,24,24	0.68	0
2	BGC	B	1001	-	12,12,12	0.48	0	17,17,17	1.14	1 (5%)
3	G6P	B	1002	-	16,16,16	0.74	0	23,24,24	0.83	0
2	BGC	B	1003	-	12,12,12	0.37	0	17,17,17	0.73	1 (5%)
3	G6P	B	1004	-	16,16,16	0.76	0	23,24,24	0.72	0

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
2	BGC	A	1001	-	-	0/2/22/22	0/1/1/1
3	G6P	A	1002	-	1/1/6/6	0/6/26/26	0/1/1/1
2	BGC	A	1003	-	-	0/2/22/22	0/1/1/1
3	G6P	A	1004	-	1/1/6/6	0/6/26/26	0/1/1/1
2	BGC	B	1001	-	-	0/2/22/22	0/1/1/1
3	G6P	B	1002	-	1/1/6/6	0/6/26/26	0/1/1/1
2	BGC	B	1003	-	-	0/2/22/22	0/1/1/1
3	G6P	B	1004	-	1/1/6/6	0/6/26/26	0/1/1/1

There are no bond length outliers.

All (4) bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
2	A	1001	BGC	C1-C2-C3	-4.26	104.09	110.43
2	B	1001	BGC	C1-C2-C3	-3.75	104.85	110.43
2	A	1003	BGC	C1-C2-C3	-2.37	106.91	110.43
2	B	1003	BGC	C1-C2-C3	-2.31	106.99	110.43

All (4) chirality outliers are listed below:

Mol	Chain	Res	Type	Atom
3	B	1002	G6P	C1
3	A	1002	G6P	C1
3	A	1004	G6P	C1
3	B	1004	G6P	C1

There are no torsion outliers.

There are no ring outliers.

2 monomers are involved in 3 short contacts:

Mol	Chain	Res	Type	Clashes	Symm-Clashes
2	A	1001	BGC	2	0
2	B	1001	BGC	1	0

## 5.7 Other polymers [i](#)

There are no such residues in this entry.

## 5.8 Polymer linkage issues [i](#)

There are no chain breaks in this entry.



## 6 Fit of model and data [i](#)

### 6.1 Protein, DNA and RNA chains [i](#)

In the following table, the column labelled ‘#RSRZ> 2’ contains the number (and percentage) of RSRZ outliers, followed by percent RSRZ outliers for the chain as percentile scores relative to all X-ray entries and entries of similar resolution. The OWAB column contains the minimum, median, 95<sup>th</sup> percentile and maximum values of the occupancy-weighted average B-factor per residue. The column labelled ‘Q< 0.9’ lists the number of (and percentage) of residues with an average occupancy less than 0.9.

Mol	Chain	Analysed	<RSRZ>	#RSRZ>2	OWAB(Å <sup>2</sup> )	Q<0.9
1	A	902/918 (98%)	-0.43	4 (0%) 93 90	14, 33, 47, 59	0
1	B	902/918 (98%)	-0.46	3 (0%) 94 92	15, 33, 47, 59	0
All	All	1804/1836 (98%)	-0.44	7 (0%) 93 90	14, 33, 47, 59	0

All (7) RSRZ outliers are listed below:

Mol	Chain	Res	Type	RSRZ
1	B	3	ALA	3.3
1	B	95	VAL	3.2
1	A	3	ALA	3.0
1	A	2	ILE	2.9
1	A	6	LEU	2.8
1	A	143	ILE	2.2
1	B	593	GLY	2.2

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

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

### 6.3 Carbohydrates [i](#)

There are no carbohydrates in this entry.

### 6.4 Ligands [i](#)

In the following table, the Atoms column lists the number of modelled atoms in the group and the number defined in the chemical component dictionary. LLDF column lists the quality of electron density of the group with respect to its neighbouring residues in protein, DNA or RNA chains. The B-factors column lists the minimum, median, 95<sup>th</sup> percentile and maximum values of B factors

of atoms in the group. The column labelled 'Q< 0.9' lists the number of atoms with occupancy less than 0.9.

Mol	Type	Chain	Res	Atoms	RSCC	RSR	LLDF	B-factors( $\text{\AA}^2$ )	Q<0.9
2	BGC	B	1003	12/12	0.96	0.23	1.22	21,26,29,31	0
2	BGC	B	1001	12/12	0.96	0.18	1.16	23,29,31,31	0
2	BGC	A	1003	12/12	0.96	0.25	0.84	21,26,31,33	0
2	BGC	A	1001	12/12	0.97	0.18	0.56	25,27,29,29	0
3	G6P	B	1004	16/16	0.98	0.19	0.40	35,37,42,43	0
3	G6P	A	1002	16/16	0.97	0.14	0.17	28,31,36,37	0
3	G6P	A	1004	16/16	0.97	0.19	-0.42	36,38,41,45	0
3	G6P	B	1002	16/16	0.97	0.11	-0.71	30,34,42,44	0
4	CA	A	1005	1/1	0.96	0.18	-	10,10,10,10	0

## 6.5 Other polymers [i](#)

There are no such residues in this entry.