



wwPDB X-ray Structure Validation Summary Report ⓘ

Jan 31, 2016 – 10:02 PM GMT

PDB ID : 1RUS
Title : CRYSTAL STRUCTURE OF THE BINARY COMPLEX OF RIBULOSE-1, 5-BISPHOSPHATE CARBOXYLASE AND ITS PRODUCT, 3-PHOSPHO-D-GLYCERATE
Authors : Lundqvist, T.; Schneider, G.
Deposited on : 1991-10-10
Resolution : 2.90 Å(reported)

This is a wwPDB X-ray Structure Validation Summary Report for a publicly released PDB entry.
We welcome your comments at 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.

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)
Xtrriage (Phenix) : **NOT EXECUTED**
EDS : **NOT EXECUTED**
Percentile statistics : 20151230.v01 (using entries in the PDB archive December 30th 2015)
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP) : trunk26865

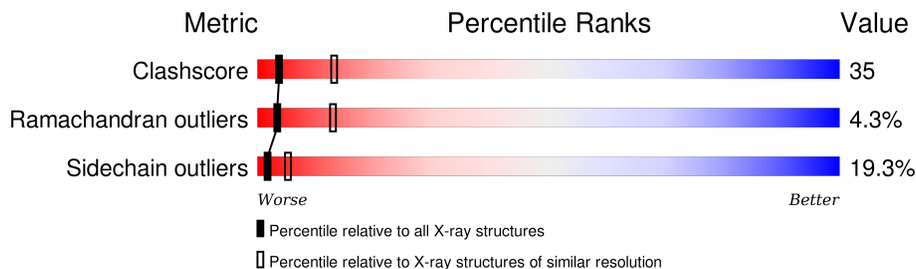
1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

X-RAY DIFFRACTION

The reported resolution of this entry is 2.90 Å.

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



Metric	Whole archive (#Entries)	Similar resolution (#Entries, resolution range(Å))
Clashscore	102246	1668 (2.90-2.90)
Ramachandran outliers	100387	1630 (2.90-2.90)
Sidechain outliers	100360	1632 (2.90-2.90)

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 ≥ 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.

Note EDS was not executed.

Mol	Chain	Length	Quality of chain
1	A	490	
1	B	490	

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
2	3PG	A	500	-	-	X	-
2	3PG	B	500	-	-	X	-

2 Entry composition [i](#)

There are 2 unique types of molecules in this entry. The entry contains 6677 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.

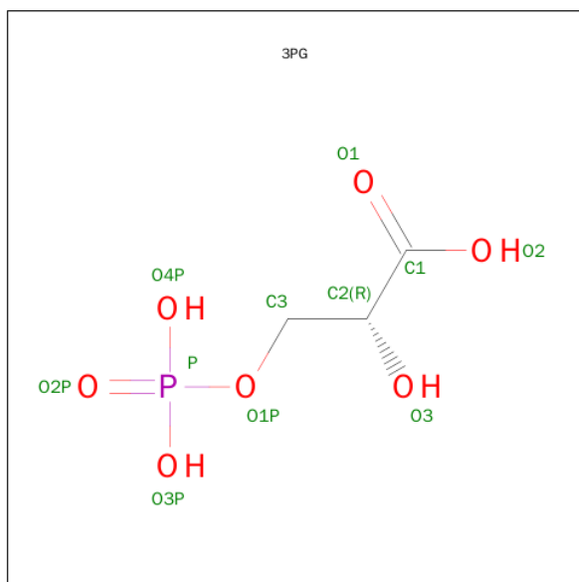
- Molecule 1 is a protein called RUBISCO (RIBULOSE-1,5-BISPHOSPHATE CARBOXYLASE(SLASH)OXYGENASE).

Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
			Total	C	N	O	S			
1	A	437	Total 3337	C 2115	N 589	O 617	S 16	7	0	0
1	B	435	Total 3318	C 2105	N 584	O 613	S 16	10	0	0

There are 2 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
A	91	ASP	HIS	CONFLICT	UNP P04718
B	91	ASP	HIS	CONFLICT	UNP P04718

- Molecule 2 is 3-PHOSPHOGLYCERIC ACID (three-letter code: 3PG) (formula: C₃H₇O₇P).



Mol	Chain	Residues	Atoms				ZeroOcc	AltConf
			Total	C	O	P		
2	A	1	Total 11	C 3	O 7	P 1	0	0

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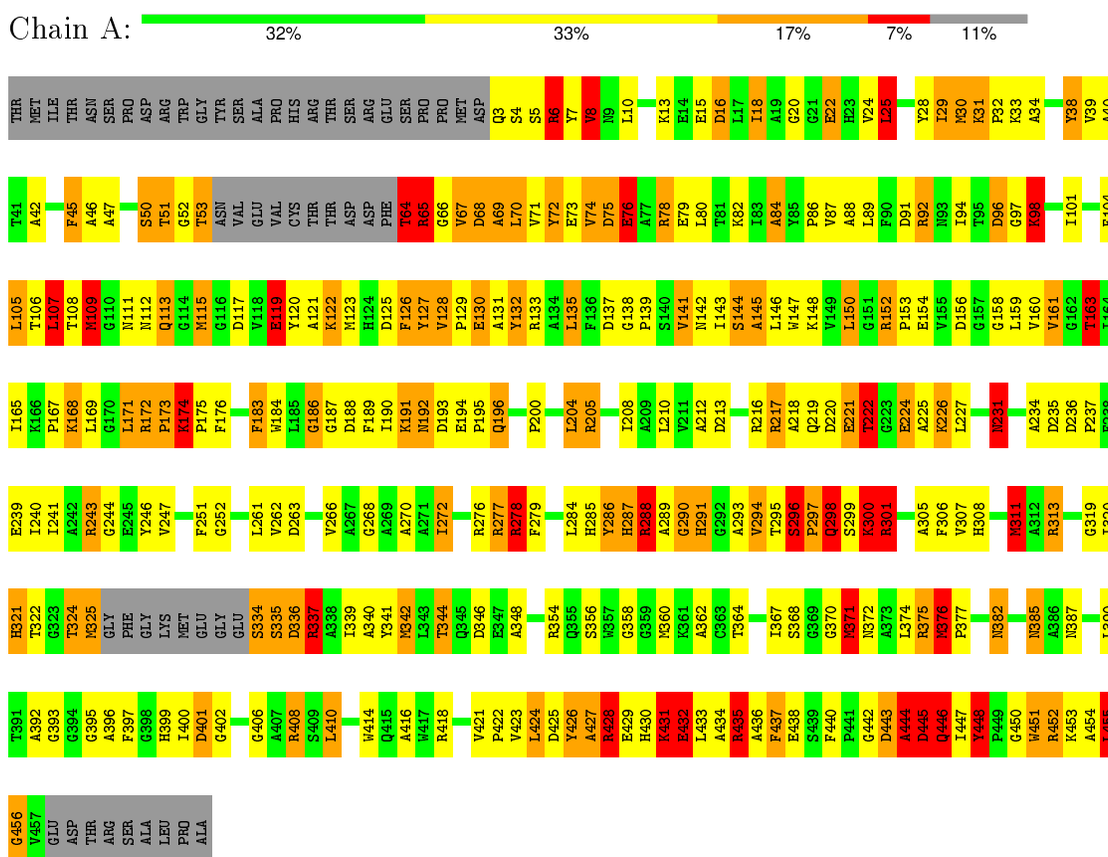
Mol	Chain	Residues	Atoms				ZeroOcc	AltConf
			Total	C	O	P		
2	B	1	11	3	7	1	0	0

3 Residue-property plots

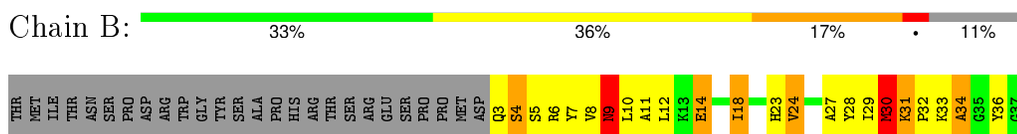
These plots are drawn for all protein, RNA and DNA chains in the entry. The first graphic for a chain summarises the proportions of errors displayed in the second graphic. The second graphic shows the sequence view annotated by issues in geometry and electron density. Residues are color-coded according 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 ($RSRZ > 2$). 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.

Note EDS was not executed.

- Molecule 1: RUBISCO (RIBULOSE-1,5-BISPHOSPHATE CARBOXYLASE(SLASH)OXYGENASE)



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H389	I400	D401	GLY	F251	E178	T106	A43
R408	S409	L410	PHE	M254	F183	L107	H44
R411	S411	R412	GLY	H257	M184	T108	F45
R418	Q412	R413	GLY	L261	L185	M109	A46
D419	Q413	R414	GLU	Y265	F189	G110	A47
P422	S414	R415	S334	V266	I190	M11	E48
V423	D335	R416	S335	G267	N192	M12	S49
L424	D336	R417	D336	A268	D193	Q113	S50
D425	W414	R418	R337	A269	E194	G114	T51
R428	R418	D419	A338	A270	F496	M15	G52
E429	D419	P422	I339	A271	Q196	G116	T53
H430	P422	V423	A340	I272	F200	D117	ASN
K431	L424	D425	T273	T274	P201	V118	VAL
E432	L433	D425	T274	A275	R208	E119	GLU
L433	D425	R428	T275	R276	D206	V120	VAL
A434	A434	E429	T276	R277	T207	A121	CYS
R435	A434	H430	T277	R278	T208	A125	THR
A436	F437	K431	T278	T279	L208	F126	THR
F437	E438	E432	A276	F279	L210	Y127	ASP
E438	S439	L433	R276	L284	L210	V128	PHE
S439	F440	A434	R277	R285	R211	E130	ARG
F440	D443	R435	R278	R286	L210	A131	G66
D443	A444	L433	F279	Y286	L211	V132	V67
D445	Q446	A434	R354	R287	A212	R133	D68
Q446	Y447	F437	R360	R288	A213	D137	A69
Y447	P449	R437	R361	A289	A214	G138	L70
P449	W451	L374	A362	G290	R215	P139	E73
W451	R452	R375	C363	R291	R216	S140	V74
R452	A454	R376	C364	G292	A218	M141	D75
A454	L455	P377	C365	A293	Q219	M142	E76
L455	G456	R377	C366	V294	D220	K148	A77
G456	V457	R378	C367	T295	E221	V149	R78
V457	GLU	R379	C368	S296	T222	L150	E79
GLU	ASP	R380	C369	P297	G223	G151	L80
ASP	THR	R381	C370	Q298	E224	R152	T81
THR	ARG	R382	C371	S299	A225	P153	R82
ARG	SER	L383	C372	K300	R226	D156	L83
SER	ALA	G384	C373	G302	L227	G157	A84
ALA	LEU	R385	C374	V307	M231	G158	P86
LEU	PRO	R386	C375	R308	L232	L159	V87
PRO	ALA	R387	C376	H308	T233	L160	A88
ALA		R388	C377	A234	A234	V161	L89
		R389	C378	M311	D235	G162	F90
		R391	C379	R313	E239	T163	D91
		T391	C380	L314	I240	I164	I93
		A392	C381	Q315	I241	I165	I94
		G393	C382	G316	I242	K166	T95
		G394	C383	A317	R243	P167	D96
		G395	C384	H321	G244	K168	G97
		F397	C385	T322	E245	G97	G97
		G396	C386	G323	V246	L169	K98
		G397	C387	T324	V247	R172	I101
		G398	C388	M325	T250	P173	A102
						K174	S103
						P175	

4 Data and refinement statistics

Xtrriage (Phenix) and EDS were not executed - this section will therefore be incomplete.

Property	Value	Source
Space group	P 1 21 1	Depositor
Cell constants a, b, c, α , β , γ	65.50Å 70.60Å 104.10Å 90.00° 92.10° 90.00°	Depositor
Resolution (Å)	10.00 – 2.90	Depositor
% Data completeness (in resolution range)	(Not available) (10.00-2.90)	Depositor
R_{merge}	(Not available)	Depositor
R_{sym}	(Not available)	Depositor
Refinement program	PROLSQ	Depositor
R, R_{free}	0.203 , (Not available)	Depositor
Estimated twinning fraction	No twinning to report.	Xtrriage
Total number of atoms	6677	wwPDB-VP
Average B, all atoms (Å ²)	21.0	wwPDB-VP

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: 3PG

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	1.04	2/3417 (0.1%)	2.34	189/4629 (4.1%)
1	B	1.13	6/3398 (0.2%)	2.37	182/4604 (4.0%)
All	All	1.08	8/6815 (0.1%)	2.35	371/9233 (4.0%)

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	B	0	1

The worst 5 of 8 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
1	B	14	GLU	CD-OE1	-6.38	1.18	1.25
1	B	195	PRO	N-CD	-5.90	1.39	1.47
1	B	194	GLU	CG-CD	5.88	1.60	1.51
1	B	288	ARG	CD-NE	5.87	1.56	1.46
1	B	288	ARG	NE-CZ	5.71	1.40	1.33

The worst 5 of 371 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	A	216	ARG	NE-CZ-NH1	31.64	136.12	120.30
1	B	418	ARG	NE-CZ-NH1	29.99	135.30	120.30
1	B	418	ARG	CD-NE-CZ	23.09	155.92	123.60
1	B	429	GLU	CA-CB-CG	22.29	162.44	113.40
1	A	428	ARG	NE-CZ-NH1	21.75	131.17	120.30

There are no chirality outliers.

All (1) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	B	111	ASN	Mainchain

5.2 Too-close contacts

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	3337	0	3242	258	2
1	B	3318	0	3220	237	0
2	A	11	0	4	7	0
2	B	11	0	4	5	0
All	All	6677	0	6470	464	2

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 35.

The worst 5 of 464 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:196:GLN:O	1:A:231:ASN:ND2	1.67	1.28
1:A:288:ARG:HD3	1:A:291:HIS:NE2	1.52	1.23
1:B:3:GLN:NE2	1:B:44:HIS:HA	1.50	1.22
1:B:321:HIS:HB3	2:B:500:3PG:O4P	1.41	1.17
1:B:335:SER:O	1:B:339:ILE:CG1	1.93	1.14

All (2) symmetry-related close contacts are listed below. The label for Atom-2 includes the symmetry operator and encoded unit-cell translations to be applied.

Atom-1	Atom-2	Interatomic distance (Å)	Clash overlap (Å)
1:A:346:ASP:OD2	1:A:408:ARG:NH2[2_646]	1.89	0.31
1:A:358:GLY:O	1:A:431:LYS:CD[2_646]	2.07	0.13

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	431/490 (88%)	374 (87%)	38 (9%)	19 (4%)	3	12
1	B	429/490 (88%)	365 (85%)	46 (11%)	18 (4%)	3	13
All	All	860/980 (88%)	739 (86%)	84 (10%)	37 (4%)	3	13

5 of 37 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	A	4	SER
1	A	298	GLN
1	A	372	ASN
1	B	4	SER
1	B	5	SER

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	330/376 (88%)	254 (77%)	76 (23%)	1	3
1	B	327/376 (87%)	276 (84%)	51 (16%)	3	10
All	All	657/752 (87%)	530 (81%)	127 (19%)	2	5

5 of 127 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	A	334	SER

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Mol	Chain	Res	Type
1	A	435	ARG
1	B	400	ILE
1	A	337	ARG
1	A	376	MET

Some sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 12 such sidechains are listed below:

Mol	Chain	Res	Type
1	A	415	GLN
1	B	9	ASN
1	B	315	GLN
1	A	385	ASN
1	B	231	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 carbohydrates in this entry.

5.6 Ligand geometry [i](#)

2 ligands are modelled in this entry.

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	3PG	A	500	-	7,10,10	4.40	6 (85%)	7,14,14	2.97	6 (85%)
2	3PG	B	500	-	7,10,10	1.84	1 (14%)	7,14,14	1.35	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	3PG	A	500	-	-	0/6/10/10	0/0/0/0
2	3PG	B	500	-	-	0/6/10/10	0/0/0/0

The worst 5 of 7 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
2	A	500	3PG	P-O3P	2.38	1.63	1.54
2	A	500	3PG	P-O4P	2.78	1.64	1.54
2	B	500	3PG	P-O2P	3.07	1.61	1.51
2	A	500	3PG	P-O2P	3.46	1.62	1.51
2	A	500	3PG	O1P-C3	3.54	1.59	1.44

The worst 5 of 6 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
2	A	500	3PG	O3-C2-C1	-3.77	99.88	111.44
2	A	500	3PG	O4P-P-O2P	-2.79	101.61	110.58
2	A	500	3PG	O3P-P-O2P	-2.01	104.11	110.58
2	A	500	3PG	O4P-P-O1P	2.43	113.55	106.56
2	A	500	3PG	O3P-P-O1P	3.38	116.29	106.56

There are no chirality outliers.

There are no torsion outliers.

There are no ring outliers.

2 monomers are involved in 12 short contacts:

Mol	Chain	Res	Type	Clashes	Symm-Clashes
2	A	500	3PG	7	0
2	B	500	3PG	5	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

6.1 Protein, DNA and RNA chains

EDS was not executed - this section will therefore be empty.

6.2 Non-standard residues in protein, DNA, RNA chains

EDS was not executed - this section will therefore be empty.

6.3 Carbohydrates

EDS was not executed - this section will therefore be empty.

6.4 Ligands

EDS was not executed - this section will therefore be empty.

6.5 Other polymers

EDS was not executed - this section will therefore be empty.