



# wwPDB X-ray Structure Validation Summary Report ⓘ

Mar 10, 2024 – 12:58 AM EST

PDB ID : 3SYV  
Title : Crystal structure of mPACSIN 3 F-BAR domain mutant  
Authors : Bai, X.  
Deposited on : 2011-07-18  
Resolution : 3.10 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](mailto:validation@mail.wwpdb.org)

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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>.

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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  
Xtriage (Phenix) : 1.13  
EDS : 2.36  
Percentile statistics : 20191225.v01 (using entries in the PDB archive December 25th 2019)  
Refmac : 5.8.0158  
CCP4 : 7.0.044 (Gargrove)  
Ideal geometry (proteins) : Engh & Huber (2001)  
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)  
Validation Pipeline (wwPDB-VP) : 2.36

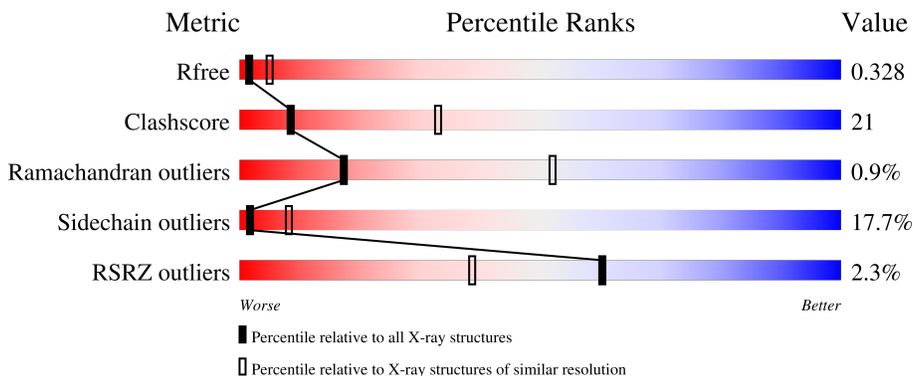
# 1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

*X-RAY DIFFRACTION*

The reported resolution of this entry is 3.10 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.



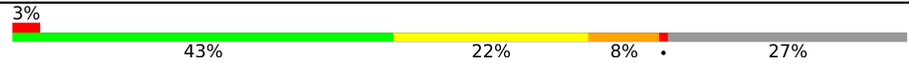
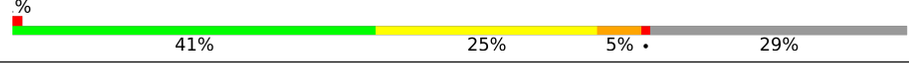
Metric	Whole archive (#Entries)	Similar resolution (#Entries, resolution range(Å))
$R_{free}$	130704	1094 (3.10-3.10)
Clashscore	141614	1184 (3.10-3.10)
Ramachandran outliers	138981	1141 (3.10-3.10)
Sidechain outliers	138945	1141 (3.10-3.10)
RSRZ outliers	127900	1067 (3.10-3.10)

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 of the lower bar indicate the fraction of residues that contain outliers for  $\geq 3$ , 2, 1 and 0 types of geometric quality criteria respectively. A grey segment represents the fraction of residues that are not modelled. The numeric value for each fraction is indicated below the corresponding segment, with a dot representing fractions  $\leq 5\%$ . The upper red bar (where present) indicates the fraction of residues that have poor fit to the electron density. The numeric value is given above the bar.

Mol	Chain	Length	Quality of chain
1	A	347	 43% 24% 5% 28%
1	B	347	 44% 25% 5% 26%
1	C	347	 41% 26% 7% 26%
1	D	347	 39% 29% 5% 27%
1	E	347	 39% 29% 6% 27%

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Mol	Chain	Length	Quality of chain
1	F	347	 <p>3% 43% 22% 8% 27%</p>
1	G	347	 <p>2% 41% 25% 7% 27%</p>
1	H	347	 <p>% 41% 25% 5% 29%</p>

## 2 Entry composition [i](#)

There are 2 unique types of molecules in this entry. The entry contains 16470 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 Protein kinase C and casein kinase II substrate protein 3.

Mol	Chain	Residues	Atoms					ZeroOcc	AltConf	Trace
			Total	C	N	O	S			
1	A	250	Total 2047	C 1284	N 384	O 372	S 7	0	0	0
1	B	257	Total 2115	C 1317	N 403	O 387	S 8	0	0	0
1	C	257	Total 2061	C 1284	N 390	O 379	S 8	0	0	0
1	D	255	Total 2085	C 1304	N 396	O 378	S 7	0	0	0
1	E	255	Total 2044	C 1278	N 387	O 374	S 5	0	0	1
1	F	255	Total 2050	C 1283	N 392	O 369	S 6	0	0	0
1	G	254	Total 1978	C 1231	N 374	O 368	S 5	0	0	0
1	H	247	Total 1981	C 1236	N 378	O 364	S 3	0	0	0

There are 56 discrepancies between the modelled and reference sequences:

Chain	Residue	Modelled	Actual	Comment	Reference
A	-5	HIS	-	expression tag	UNP Q99JB8
A	-4	HIS	-	expression tag	UNP Q99JB8
A	-3	HIS	-	expression tag	UNP Q99JB8
A	-2	HIS	-	expression tag	UNP Q99JB8
A	-1	HIS	-	expression tag	UNP Q99JB8
A	0	HIS	-	expression tag	UNP Q99JB8
A	121	GLN	PRO	engineered mutation	UNP Q99JB8
B	-5	HIS	-	expression tag	UNP Q99JB8
B	-4	HIS	-	expression tag	UNP Q99JB8
B	-3	HIS	-	expression tag	UNP Q99JB8
B	-2	HIS	-	expression tag	UNP Q99JB8
B	-1	HIS	-	expression tag	UNP Q99JB8
B	0	HIS	-	expression tag	UNP Q99JB8

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Chain	Residue	Modelled	Actual	Comment	Reference
B	121	GLN	PRO	engineered mutation	UNP Q99JB8
C	-5	HIS	-	expression tag	UNP Q99JB8
C	-4	HIS	-	expression tag	UNP Q99JB8
C	-3	HIS	-	expression tag	UNP Q99JB8
C	-2	HIS	-	expression tag	UNP Q99JB8
C	-1	HIS	-	expression tag	UNP Q99JB8
C	0	HIS	-	expression tag	UNP Q99JB8
C	121	GLN	PRO	engineered mutation	UNP Q99JB8
D	-5	HIS	-	expression tag	UNP Q99JB8
D	-4	HIS	-	expression tag	UNP Q99JB8
D	-3	HIS	-	expression tag	UNP Q99JB8
D	-2	HIS	-	expression tag	UNP Q99JB8
D	-1	HIS	-	expression tag	UNP Q99JB8
D	0	HIS	-	expression tag	UNP Q99JB8
D	121	GLN	PRO	engineered mutation	UNP Q99JB8
E	-5	HIS	-	expression tag	UNP Q99JB8
E	-4	HIS	-	expression tag	UNP Q99JB8
E	-3	HIS	-	expression tag	UNP Q99JB8
E	-2	HIS	-	expression tag	UNP Q99JB8
E	-1	HIS	-	expression tag	UNP Q99JB8
E	0	HIS	-	expression tag	UNP Q99JB8
E	121	GLN	PRO	engineered mutation	UNP Q99JB8
F	-5	HIS	-	expression tag	UNP Q99JB8
F	-4	HIS	-	expression tag	UNP Q99JB8
F	-3	HIS	-	expression tag	UNP Q99JB8
F	-2	HIS	-	expression tag	UNP Q99JB8
F	-1	HIS	-	expression tag	UNP Q99JB8
F	0	HIS	-	expression tag	UNP Q99JB8
F	121	GLN	PRO	engineered mutation	UNP Q99JB8
G	-5	HIS	-	expression tag	UNP Q99JB8
G	-4	HIS	-	expression tag	UNP Q99JB8
G	-3	HIS	-	expression tag	UNP Q99JB8
G	-2	HIS	-	expression tag	UNP Q99JB8
G	-1	HIS	-	expression tag	UNP Q99JB8
G	0	HIS	-	expression tag	UNP Q99JB8
G	121	GLN	PRO	engineered mutation	UNP Q99JB8
H	-5	HIS	-	expression tag	UNP Q99JB8
H	-4	HIS	-	expression tag	UNP Q99JB8
H	-3	HIS	-	expression tag	UNP Q99JB8
H	-2	HIS	-	expression tag	UNP Q99JB8
H	-1	HIS	-	expression tag	UNP Q99JB8
H	0	HIS	-	expression tag	UNP Q99JB8

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Chain	Residue	Modelled	Actual	Comment	Reference
H	121	GLN	PRO	engineered mutation	UNP Q99JB8

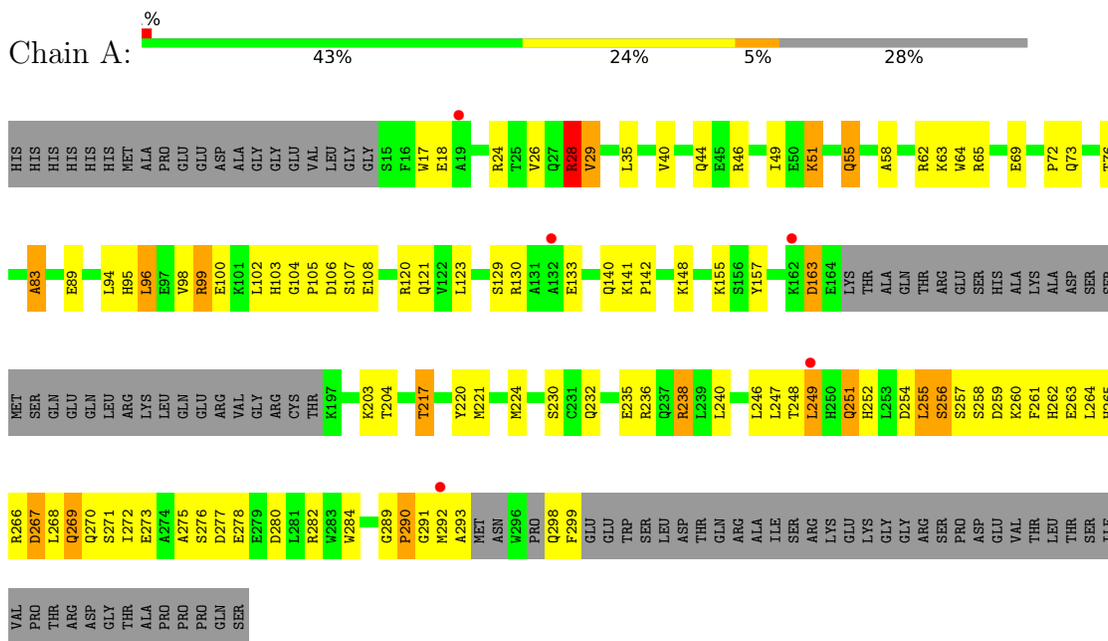
- Molecule 2 is water.

Mol	Chain	Residues	Atoms	ZeroOcc	AltConf
2	A	14	Total O 14 14	0	0
2	B	13	Total O 13 13	0	0
2	C	19	Total O 19 19	0	0
2	D	12	Total O 12 12	0	0
2	E	12	Total O 12 12	0	0
2	F	16	Total O 16 16	0	0
2	G	10	Total O 10 10	0	0
2	H	13	Total O 13 13	0	0

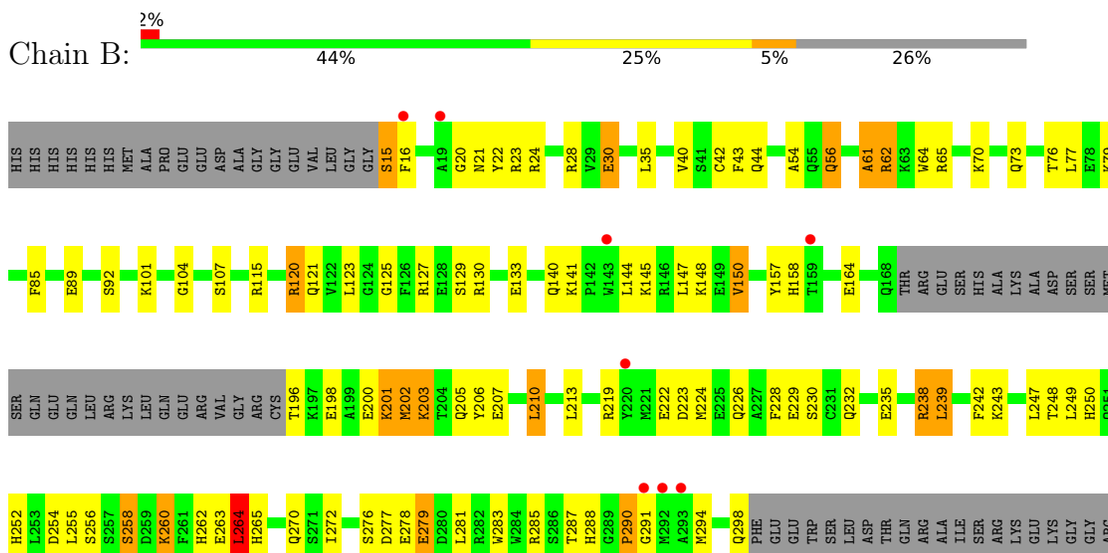
### 3 Residue-property plots i

These plots are drawn for all protein, RNA, DNA and oligosaccharide chains in the entry. The first graphic for a chain summarises the proportions of the various outlier classes displayed in the second graphic. The second graphic shows the sequence view annotated by issues in geometry and 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.

- Molecule 1: Protein kinase C and casein kinase II substrate protein 3



- Molecule 1: Protein kinase C and casein kinase II substrate protein 3



SER PRO ASP GLU VAL THR LEU THR SER SER ILE VAL LEU THR PRO ARG ASP THR ALA PRO PRO PRO GLN SER

● Molecule 1: Protein kinase C and casein kinase II substrate protein 3



HIS HIS HIS HIS HIS HIS HIS MET ALA PRO PRO GLU VAL LEU THR PRO ARG ASP G8 G14 S15 F16 W17 W18 E19 A19 G20 N21 Y22 R23 R24 T25

A80 W81 A82 A83 F84 F85 T86 E89 R90 G93 R99 E100 K101 K101 E108 Q114 R130 G20 N21 R137 Q140 K141 P142 W143 V29 V29 E30 D31 R34 K148 L35 C36 V40 S153 K154 Q44 S156 Y157 R46 H158 T159 D163 GLU LYS THR ALA GLN THR ARG GLU HIS ALA LYS ALA ASP

SER SER MET SER SER GLN MET LEU LYS LEU GLN ARG ARG VAL GLY ARG CYS THR LYS E198 A199 E200 M202 K203 T204 Q205 P142 W143 Y206 L144 K145 Q208 T209 L210 A211 E212 L213 N214 R215 Y216 T217 P218 R219 Y220 M221 M224 A227 F228 Q232 E235 R238 D244 V245 L247

Q251 R252 D254 L255 S256 S257 D259 K260 F261 H262 E263 H265 R266 D267 L268 Q269 Q270 S271 I272 L272 A273 A275 S276 D277 E278 E279 D280 L281 W284 R285 G289 F289 G291 M292 A293 M294 N295 Y296 P299 GLU GLY TRP SER LEU THR ASP GLN ARG ALA ILE SER ARG

LYS GLU LYS GLY GLY ARG MET PRO ASP GLU VAL THR LEU THR THR SER ILE VAL PRO THR ARG ASP GLY ALA PRO PRO GLN SER

● Molecule 1: Protein kinase C and casein kinase II substrate protein 3



HIS HIS HIS HIS HIS HIS MET ALA PRO PRO GLU VAL LEU THR ALA GLY GLY H14 S15 F16 W17 W18 E18 G20 N21 Y22 R23 R24 R28 R34 L35 D38 L39 V40 F43 Q44 A47 R48 E49 E50 K51 Q56 R62 R63 W64 R65 G66 A67 V68 E69 K70

G71 P72 T76 L77 W81 F84 E89 R90 E93 L94 H95 V98 R99 E100 K101 L102 D106 S107 E108 R109 V110 H119 R120 Q121 G124 G125 F126 R127 R130 E133 Q140 W143 L147 K148 E151 H158 T159 A160 R161 E164 K165 THR ALA

GLN THR ARG GLU SER HIS ALA LYS ALA ASP GLU SER MET SER SER GLN GLU GLN VAL LEU ARG LYS LEU LEU LYS L102 F196 T197 R201 M202 Q205 Y206 L210 A211 L213 E212 N214 R215 Y215 T217 P218 M221 E222 D223 M224 E225 Q226 A227 F228 C231 Q232 E235

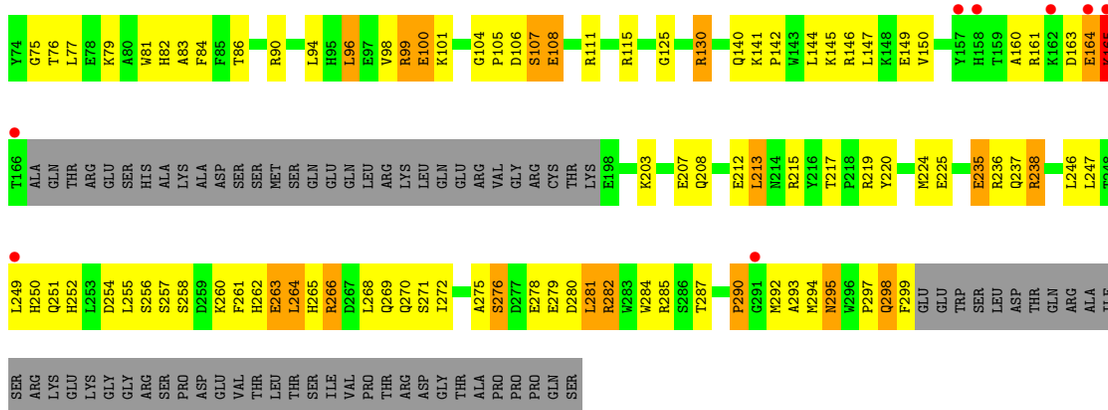
R236 Q237 L239 K243 D244 V245 L247 L248 L249 H250 L255 S256 S257 S258 D259 K260 F261 R266 Q269 A275 S276 D277 E278 E279 D280 L281 R282 S285 T287 H288 G289 P290 C291 M292 ALA M294 Q298 F299 GLU TRP M224 E225 Q226 A227 F228 C231 Q232 E235

ARG LYS GLU LYS GLY ARG SER PRO ASP GLU VAL THR LEU THR SER SER ILE VAL LEU THR PRO ARG ASP GLY ALA PRO PRO GLN SER

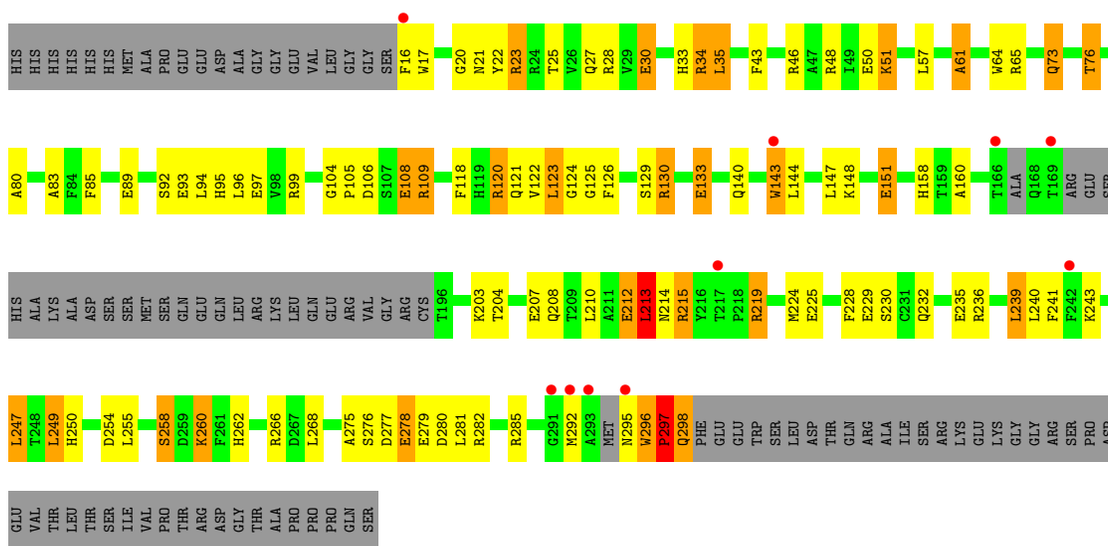
● Molecule 1: Protein kinase C and casein kinase II substrate protein 3



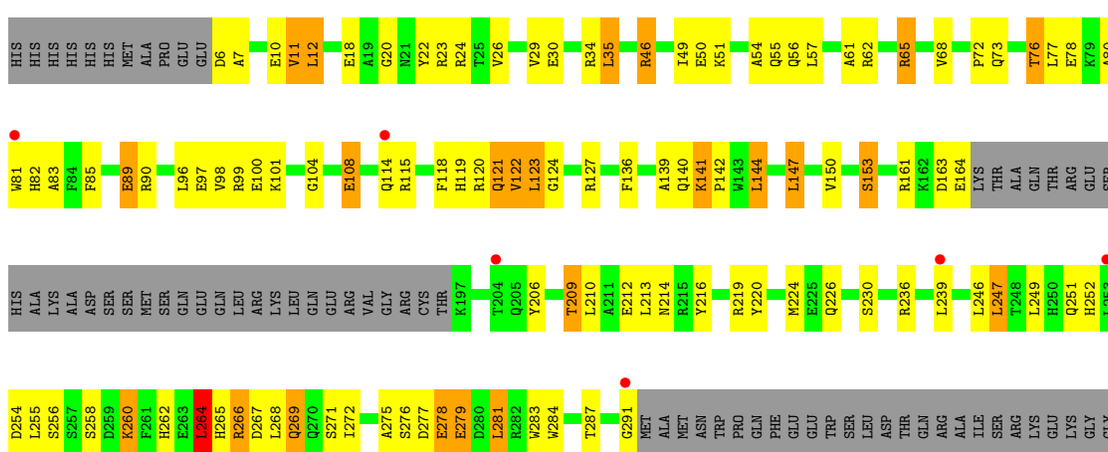
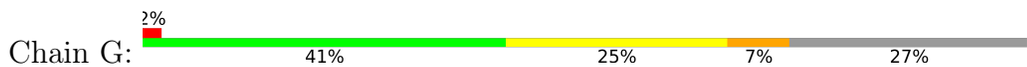
HIS HIS HIS HIS HIS HIS MET ALA PRO PRO GLU VAL LEU THR ALA GLY G14 S15 F16 W17 E18 A19 G20 N21 Y22 R23 R24 R28 V29 E30 R34 L35 V40 S41 C42 F43 Q44 A47 E50 Q55 R62 K63 W64 R65 E69 K70 G71 P72 Q73



● Molecule 1: Protein kinase C and casein kinase II substrate protein 3

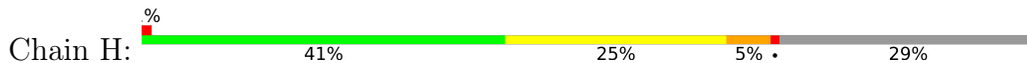


● Molecule 1: Protein kinase C and casein kinase II substrate protein 3



ARG  
SER  
PRO  
ASP  
GLU  
VAL  
THR  
LEU  
THR  
SER  
ILE  
VAL  
PRO  
THR  
ARG  
ASP  
GLY  
THR  
ALA  
PRO  
PRO  
PRO  
GLN  
GLN  
SER

● Molecule 1: Protein kinase C and casein kinase II substrate protein 3



HIS  
HIS  
HIS  
HIS  
HIS  
HIS  
HIS  
MET  
ALA  
PRO  
GLU  
GLU  
GLU  
ASP  
ALA  
ALA  
GLY  
GLY  
GLU  
VAL  
LEU  
GLY  
G14  
S15  
F16  
W17  
W17  
E18  
A19  
G20  
R23  
R24  
R28  
Y28  
E30  
D31  
L35  
C36  
L38  
V40  
S41  
A47  
R48  
I49  
E50  
K51  
A54  
Q55  
Q56  
L57  
R62  
K63  
W64  
A67  
V68  
E69

K70  
G71  
Q73  
T76  
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W81  
H82  
F85  
A88  
E89  
R90  
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S92  
E93  
L94  
H95  
L96  
E97  
V98  
R99  
E100  
G104  
P105  
D106  
S107  
E108  
R109  
V110  
R115  
R120  
Q121  
R127  
R130  
K141  
P142  
W143  
L144  
L147  
A160  
E164  
LYS  
THR  
ALA  
GLN  
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ARG  
G195  
K203  
T205  
Q205  
Y206  
E207  
Q208  
T209  
L210  
Y216  
THR  
P218  
R219  
E225  
F228  
E229  
S230  
Q231  
Q232  
E235  
R236  
Q237  
R238  
L239  
F242  
K243  
L247  
T248  
L249

H250  
Q251  
H252  
L253  
D254  
S258  
D259  
K260  
F261  
L264  
H265  
R266  
Q269  
A275  
S276  
D277  
E278  
E279  
D280  
L281  
S286  
T287  
H288  
G289  
P290  
G291  
MET  
ALA  
MET  
ASN  
TRP  
PRO  
GLN  
PHE  
GLU  
GLU  
TRP  
SER  
LEU  
ASP  
THR  
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ARG  
ASP  
GLY  
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## 4 Data and refinement statistics i

Property	Value	Source
Space group	P 1 21 1	Depositor
Cell constants a, b, c, $\alpha$ , $\beta$ , $\gamma$	120.57Å 108.90Å 222.32Å 90.00° 90.05° 90.00°	Depositor
Resolution (Å)	48.90 – 3.10 48.90 – 3.10	Depositor EDS
% Data completeness (in resolution range)	98.7 (48.90-3.10) 98.5 (48.90-3.10)	Depositor EDS
$R_{merge}$	(Not available)	Depositor
$R_{sym}$	(Not available)	Depositor
$\langle I/\sigma(I) \rangle$ <sup>1</sup>	3.60 (at 3.12Å)	Xtrriage
Refinement program	REFMAC 5.5.0110	Depositor
R, $R_{free}$	0.281 , 0.337 0.274 , 0.328	Depositor DCC
$R_{free}$ test set	5196 reflections (5.03%)	wwPDB-VP
Wilson B-factor (Å <sup>2</sup> )	64.1	Xtrriage
Anisotropy	0.371	Xtrriage
Bulk solvent $k_{sol}$ (e/Å <sup>3</sup> ), $B_{sol}$ (Å <sup>2</sup> )	0.34 , 31.0	EDS
L-test for twinning <sup>2</sup>	$\langle  L  \rangle = 0.55$ , $\langle L^2 \rangle = 0.40$	Xtrriage
Estimated twinning fraction	0.458 for h,-k,-l	Xtrriage
$F_o, F_c$ correlation	0.93	EDS
Total number of atoms	16470	wwPDB-VP
Average B, all atoms (Å <sup>2</sup> )	61.0	wwPDB-VP

Xtrriage's analysis on translational NCS is as follows: *The analyses of the Patterson function reveals a significant off-origin peak that is 50.83 % of the origin peak, indicating pseudo-translational symmetry. The chance of finding a peak of this or larger height randomly in a structure without pseudo-translational symmetry is equal to 6.1201e-05. The detected translational NCS is most likely also responsible for the elevated intensity ratio.*

<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, 0.333 respectively for untwinned datasets, and 0.375, 0.2 for perfectly twinned datasets.

## 5 Model quality i

### 5.1 Standard geometry i

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

Mol	Chain	Bond lengths		Bond angles	
		RMSZ	# Z  >5	RMSZ	# Z  >5
1	A	1.12	6/2094 (0.3%)	1.01	5/2811 (0.2%)
1	B	1.14	7/2160 (0.3%)	1.00	2/2895 (0.1%)
1	C	1.12	8/2104 (0.4%)	1.00	4/2821 (0.1%)
1	D	1.26	11/2133 (0.5%)	1.06	4/2862 (0.1%)
1	E	1.17	6/2092 (0.3%)	1.02	6/2813 (0.2%)
1	F	1.22	12/2097 (0.6%)	1.05	2/2821 (0.1%)
1	G	1.14	3/2021 (0.1%)	1.03	5/2724 (0.2%)
1	H	1.28	12/2025 (0.6%)	1.08	9/2725 (0.3%)
All	All	1.18	65/16726 (0.4%)	1.03	37/22472 (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	D	0	1

The worst 5 of 65 bond length outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(Å)	Ideal(Å)
1	F	108	GLU	CG-CD	10.46	1.67	1.51
1	H	108	GLU	CG-CD	10.14	1.67	1.51
1	D	160	ALA	C-O	9.96	1.42	1.23
1	H	97	GLU	CG-CD	9.93	1.66	1.51
1	A	108	GLU	CG-CD	8.23	1.64	1.51

The worst 5 of 37 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	A	238	ARG	NE-CZ-NH2	-9.19	115.71	120.30
1	D	161	ARG	NE-CZ-NH2	-7.60	116.50	120.30
1	A	28	ARG	NE-CZ-NH1	6.89	123.75	120.30

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Mol	Chain	Res	Type	Atoms	Z	Observed(°)	Ideal(°)
1	F	219	ARG	NE-CZ-NH1	6.88	123.74	120.30
1	E	238	ARG	NE-CZ-NH2	-6.84	116.88	120.30

There are no chirality outliers.

All (1) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	D	160	ALA	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	2047	0	1943	82	0
1	B	2115	0	2020	76	0
1	C	2061	0	1930	97	0
1	D	2085	0	1982	110	0
1	E	2044	0	1900	111	0
1	F	2050	0	1898	93	0
1	G	1978	0	1807	89	0
1	H	1981	0	1827	84	0
2	A	14	0	0	1	0
2	B	13	0	0	2	0
2	C	19	0	0	0	0
2	D	12	0	0	3	0
2	E	12	0	0	0	0
2	F	16	0	0	2	0
2	G	10	0	0	0	0
2	H	13	0	0	1	0
All	All	16470	0	15307	658	0

The all-atom clashscore is defined as the number of clashes found per 1000 atoms (including hydrogen atoms). The all-atom clashscore for this structure is 21.

The worst 5 of 658 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:D:196:THR:CB	1:D:196:THR:CG2	1.76	1.59
1:D:121:GLN:HE22	1:D:130:ARG:CD	1.42	1.33
1:D:121:GLN:NE2	1:D:130:ARG:HD3	1.46	1.30
1:E:76:THR:CG2	1:E:275:ALA:HA	1.69	1.23
1:A:76:THR:CG2	1:A:275:ALA:HA	1.74	1.18

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	243/347 (70%)	229 (94%)	10 (4%)	4 (2%)	9	37
1	B	253/347 (73%)	236 (93%)	14 (6%)	3 (1%)	13	44
1	C	251/347 (72%)	231 (92%)	20 (8%)	0	100	100
1	D	249/347 (72%)	230 (92%)	17 (7%)	2 (1%)	19	54
1	E	249/347 (72%)	232 (93%)	16 (6%)	1 (0%)	34	69
1	F	247/347 (71%)	224 (91%)	21 (8%)	2 (1%)	19	54
1	G	250/347 (72%)	222 (89%)	25 (10%)	3 (1%)	13	44
1	H	241/347 (70%)	224 (93%)	14 (6%)	3 (1%)	13	44
All	All	1983/2776 (71%)	1828 (92%)	137 (7%)	18 (1%)	17	52

5 of 18 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	F	213	LEU
1	F	297	PRO
1	B	290	PRO
1	D	71	GLY
1	H	71	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	201/294 (68%)	173 (86%)	28 (14%)	3 15
1	B	210/294 (71%)	177 (84%)	33 (16%)	2 11
1	C	197/294 (67%)	154 (78%)	43 (22%)	1 4
1	D	205/294 (70%)	172 (84%)	33 (16%)	2 10
1	E	196/294 (67%)	157 (80%)	39 (20%)	1 5
1	F	193/294 (66%)	157 (81%)	36 (19%)	1 7
1	G	182/294 (62%)	143 (79%)	39 (21%)	1 4
1	H	186/294 (63%)	159 (86%)	27 (14%)	3 13
All	All	1570/2352 (67%)	1292 (82%)	278 (18%)	2 8

5 of 278 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	G	147	LEU
1	G	230	SER
1	H	89	GLU
1	C	259	ASP
1	C	246	LEU

Sometimes sidechains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 69 such sidechains are listed below:

Mol	Chain	Res	Type
1	G	269	GLN
1	H	82	HIS
1	H	250	HIS
1	C	288	HIS
1	C	270	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 monosaccharides in this entry.

#### 5.6 Ligand geometry [i](#)

There are no ligands in this entry.

#### 5.7 Other polymers [i](#)

There are no such residues in this entry.

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

There are no chain breaks in this entry.

## 6 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	250/347 (72%)	0.36	5 (2%) 65 44	37, 61, 103, 129	0
1	B	257/347 (74%)	0.41	8 (3%) 49 26	25, 55, 120, 137	0
1	C	257/347 (74%)	0.38	3 (1%) 79 61	42, 59, 97, 112	0
1	D	255/347 (73%)	0.38	5 (1%) 65 44	34, 54, 120, 135	0
1	E	255/347 (73%)	0.40	8 (3%) 49 26	39, 59, 108, 127	0
1	F	255/347 (73%)	0.46	10 (3%) 39 20	32, 55, 101, 114	0
1	G	254/347 (73%)	0.34	6 (2%) 59 37	41, 59, 83, 98	0
1	H	247/347 (71%)	0.39	2 (0%) 86 72	35, 53, 89, 102	0
All	All	2030/2776 (73%)	0.39	47 (2%) 60 39	25, 57, 103, 137	0

The worst 5 of 47 RSRZ outliers are listed below:

Mol	Chain	Res	Type	RSRZ
1	B	292	MET	7.4
1	C	293	ALA	6.4
1	F	293	ALA	5.8
1	E	291	GLY	5.0
1	E	166	THR	4.9

### 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 monosaccharides in this entry.

## 6.4 Ligands

There are no ligands in this entry.

## 6.5 Other polymers

There are no such residues in this entry.