

wwPDB X-ray Structure Validation Summary Report (i)

Feb 4, 2024 – 07:22 AM EST

PDB ID	:	1NL3
Title	:	CRYSTAL STRUCTURE OF THE SECA PROTEIN TRANSLOCATION
		ATPASE FROM MYCOBACTERIUM TUBERCULOSIS in APO FORM
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		Consortium (TBSGC)
Deposited on	:	2003-01-06
Resolution	:	2.80 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 https://www.wwpdb.org/validation/2017/XrayValidationReportHelp with specific help available everywhere you see the (i) 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 (i)) were used in the production of this report:

MolProbity	:	4.02b-467
Xtriage (Phenix)	:	NOT EXECUTED
EDS	:	NOT EXECUTED
Percentile statistics	:	20191225.v01 (using entries in the PDB archive December 25th 2019)
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 (i)

The following experimental techniques were used to determine the structure: $X\text{-}RAY \, DIFFRACTION$

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	$egin{array}{c} { m Whole \ archive} \ (\#{ m Entries}) \end{array}$	${f Similar\ resolution}\ (\#{ m Entries,\ resolution\ range}({ m \AA}))$
Clashscore	141614	3569(2.80-2.80)
Ramachandran outliers	138981	3498 (2.80-2.80)
Sidechain outliers	138945	3500 (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 of the lower 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 <=5%

Note EDS was not executed.

Mol	Chain	Length	Quality of	chain	
1	А	922	60%	25%	6% • 9%
1	В	922	56%	27%	7% • 9%



2 Entry composition (i)

There are 2 unique types of molecules in this entry. The entry contains 14209 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.

Mol	Chain	Residues		Α	toms			ZeroOcc	AltConf	Trace
1	А	839	Total 6639	C 4157	N 1169	0 1288	S 25	0	0	0
1	В	838	Total 6640	C 4157	N 1171	0 1287	S 25	0	0	0

• Molecule 1 is a protein called PREPROTEIN TRANSLOCASE SECA 1 SUBUNIT.

Chain	Residue	Modelled	Actual	Comment	Reference					
А	-29	MET	-	cloning artifact	UNP P0A5Y8					
A	-28	LYS	-	cloning artifact	UNP P0A5Y8					
А	-27	GLU	-	cloning artifact	UNP P0A5Y8					
А	-26	THR	-	cloning artifact	UNP P0A5Y8					
А	-25	ALA	-	cloning artifact	UNP P0A5Y8					
A	-24	ALA	-	cloning artifact	UNP P0A5Y8					
А	-23	ALA	-	cloning artifact	UNP P0A5Y8					
А	-22	LYS	-	cloning artifact	UNP P0A5Y8					
А	-21	PHE	-	cloning artifact	UNP P0A5Y8					
А	-20	GLU	-	cloning artifact	UNP P0A5Y8					
А	-19	ARG	-	cloning artifact	UNP P0A5Y8					
А	-18	GLN	-	cloning artifact	UNP P0A5Y8					
А	-17	HIS	-	cloning artifact	UNP P0A5Y8					
А	-16	MET	-	cloning artifact	UNP P0A5Y8					
А	-15	ASP	-	cloning artifact	UNP P0A5Y8					
А	-14	SER	-	cloning artifact	UNP P0A5Y8					
А	-13	PRO	-	cloning artifact	UNP P0A5Y8					
A	-12	ASP	-	cloning artifact	UNP P0A5Y8					
А	-11	LEU	-	cloning artifact	UNP P0A5Y8					
А	-10	GLY	-	cloning artifact	UNP P0A5Y8					
А	-9	THR	-	cloning artifact	UNP P0A5Y8					
А	-8	LEU	-	cloning artifact	UNP P0A5Y8					
А	-7	VAL	-	cloning artifact	UNP P0A5Y8					
А	-6	PRO	-	cloning artifact	UNP P0A5Y8					
А	-5	ARG	-	cloning artifact	UNP P0A5Y8					

There are 62 discrepancies between the modelled and reference sequences:

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Chain	Residue	Modelled	Actual	Comment	Reference					
А	-4	GLY	-	cloning artifact	UNP P0A5Y8					
А	-3	SER	-	cloning artifact	UNP P0A5Y8					
А	-2	MET	-	cloning artifact	UNP P0A5Y8					
А	-1	ALA	-	cloning artifact	UNP P0A5Y8					
А	0	ASP	-	cloning artifact	UNP P0A5Y8					
А	1	ILE	-	cloning artifact	UNP P0A5Y8					
В	-29	MET	-	cloning artifact	UNP P0A5Y8					
В	-28	LYS	-	cloning artifact	UNP P0A5Y8					
В	-27	GLU	-	cloning artifact	UNP P0A5Y8					
В	-26	THR	-	cloning artifact	UNP P0A5Y8					
В	-25	ALA	-	cloning artifact	UNP P0A5Y8					
В	-24	ALA	-	cloning artifact	UNP P0A5Y8					
В	-23	ALA	-	cloning artifact	UNP P0A5Y8					
В	-22	LYS	-	cloning artifact	UNP P0A5Y8					
В	-21	PHE	-	cloning artifact	UNP P0A5Y8					
В	-20	GLU	-	cloning artifact	UNP P0A5Y8					
В	-19	ARG	-	cloning artifact	UNP P0A5Y8					
В	-18	GLN	-	cloning artifact	UNP P0A5Y8					
В	-17	HIS	-	cloning artifact	UNP P0A5Y8					
В	-16	MET	-	cloning artifact	UNP P0A5Y8					
В	-15	ASP	-	cloning artifact	UNP P0A5Y8					
В	-14	SER	-	cloning artifact	UNP P0A5Y8					
В	-13	PRO	-	cloning artifact	UNP P0A5Y8					
В	-12	ASP	-	cloning artifact	UNP P0A5Y8					
В	-11	LEU	-	cloning artifact	UNP P0A5Y8					
В	-10	GLY	-	cloning artifact	UNP P0A5Y8					
В	-9	THR	-	cloning artifact	UNP P0A5Y8					
В	-8	LEU	-	cloning artifact	UNP P0A5Y8					
В	-7	VAL	-	cloning artifact	UNP P0A5Y8					
В	-6	PRO	-	cloning artifact	UNP P0A5Y8					
В	-5	ARG	-	cloning artifact	UNP P0A5Y8					
В	-4	GLY	-	cloning artifact	UNP P0A5Y8					
В	-3	SER	-	cloning artifact	UNP P0A5Y8					
В	-2	MET	-	cloning artifact	UNP P0A5Y8					
В	-1	ALA	-	cloning artifact	UNP P0A5Y8					
В	0	ASP	-	cloning artifact	UNP P0A5Y8					
В	1	ILE	-	cloning artifact	UNP P0A5Y8					

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• Molecule 2 is water.

Mol	Chain	Residues	Ator	ns	ZeroOcc	AltConf
2	А	475	Total 475	O 475	0	0

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Mol	Chain	Residues	Atoms	ZeroOcc	AltConf
2	В	455	Total O 455 455	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. 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. 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: PREPROTEIN TRANSLOCASE SECA 1 SUBUNIT



• Molecule 1: PREPROTEIN TRANSLOCASE SECA 1 SUBUNIT

Chain B: 56% 27% 7% 9%



MET	GLU	THR	ALA	ALA ALA	LYS	GLU	ARG	GLN	MET	D-15	P-13	D-12		К-5 G-4		L2	K4	L5	L6	R7 1.8	2	R12	K15	-	K18	A21		N.24	D30	V31	T35	D36	L39	R40	K42	T43	D44	R48	R49	A51	D52 D53	K54
N55	P64		L78	D79 080	R81	M88		TOTM	K107	T108	C111	V112		Y116 L117	N118		V130		Y133	K136	R137		M141 M142	G143	R144 V145	H146	R147	F148	L151	Q152	V155	D163		R166	N178	N179	D184	Y185	L186	A191	DIGE	D196
-	R200	A205		V210 D211	S212	D216		K219	L222	1223	122 4 S225	G226	P227	A228 D229		S232	N233 W234	Y235		R240 1241	A242	P243	E246	K247	D248 V249	H250	Y251	D254	L255	R256 K257	R258	T259 V260		K265	V270		1276 D277	N278	1004	1821	N284	S289
Y290	L291	L295	0 0 1 1 1	E299	R303	K305		V309 R310	D311	0104	D318 E319	F320	T321	G322 R323	V324	L325	1326 6327	R328	R329	0336		K341	E342	E345	I346 K347	A348	E349	0351	T352	L353 A354	T355	I356 T357	L358	Q 359	L364	Y365	D366 K367		M371 T270	13/2	A375	E381
	E384	V391	2000	M397	K410 T41	E412	E413	D421	D422	V423	A4.24 E4.25	R426		K429 G430	-	V433	L434 T435	G436	T437	T438 S439	V440	E441	R442 S443	E444	S447	R448	0449	F450	R453	R454 T455	P456	H457	L460	V163	Y464		E468	1472	04 70	R4 / 8	N486	G489
R490	D493		D501	D505	0506 0506		E510	R511	D514	4	E51/ T518	P519	E520	E521 Y522	E523		H527 S528	E529	L530	P531 T532	V533	K534	E536	A537	8538 K539	E540		E943	E546	V551	V552	TSSS	E556	R557 UEEO	E559		D564 N565	0566 0566		8571	G572 B573	
E579	L586		R592	R593	L599 E600	E000 T601	L602	L603 T604	R605	000	P609	D610	1	E615 A616	K617	M618		R621		K624	0629	V630	E636	V637	R639 K639		M648	0650 0650	Q651	R652 K653	V654	I 655	R659	R660	1662	L663	KEEO	D670	0671	A072 L673	D674	R677
D678	Y683	V684	D685	G686 A687	T688	A693	E694	D695 W696	D697	L698	A700	L701	W7 02	L705	-	60LX	1713	T714	A715	D716 S717	L718	T7 19	LYS	ASP	HIS GLII	PHE	GLU	ARG	ASP	LEU THR	ARG	E733 F734	L735	L736 E737	, i	L740	K741 D742		R750	L754	E755 E756	1757
A758	G759 E760		M763	R764	E767	N769	0227		V774	1775	B/77	K778	W779	R/80	Y784	E785	V788		E791	G792 1793	G794	L795	A797	M798	A799 D800	R801	D802	P803	E806	7807 0808	R809	E810	M814		G821	M822	K823 F824	E825	S826 11007	1281	F831 N832	
V835	GLU ALA	VAL	PR0	ALA PRO	PRO	ALA	PRO	ALA ALA	GLU	PRO	GLU	TEU	ALA	GLU	ALA	ALA	ALA AI.A	ALA	ALA	ALA	GLN	GLN	SER	ALA	VAL	GLY	GLY	ALA ARG	GLU	ARG ALA	PRO	SER ALA	LEU	ARG	TYS	GLY	VAL AT A	SER	GLU	PRO	ALA	



4 Data and refinement statistics (i)

Xtriage (Phenix) and EDS were not executed - this section is therefore incomplete.

Property	Value	Source			
Space group	P 62 2 2	Depositor			
Cell constants	206.20Å 206.20Å 295.41Å	Depositor			
a, b, c, α , β , γ	90.00° 90.00° 120.00°	Depositor			
Resolution (Å)	8.00 - 2.80	Depositor			
% Data completeness	99.4 (8.00-2.80)	Depositor			
(in resolution range)	55.4 (0.00 2.00)	Depositor			
R_{merge}	(Not available)	Depositor			
R _{sym}	0.10	Depositor			
Refinement program	REFMAC $5.1.25$	Depositor			
R, R_{free}	0.193 , 0.259	Depositor			
Estimated twinning fraction	No twinning to report.	Xtriage			
Total number of atoms	14209	wwPDB-VP			
Average B, all atoms $(Å^2)$	48.0	wwPDB-VP			



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).

Mal	Chain	Bond	lengths	Bond angles							
	Unam	RMSZ	# Z > 5	RMSZ	# Z > 5						
1	А	0.59	0/6743	0.87	26/9116~(0.3%)						
1	В	0.59	0/6744	0.88	25/9116~(0.3%)						
All	All	0.59	0/13487	0.87	51/18232~(0.3%)						

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	А	0	1

There are no bond length outliers.

The worst 5 of 51 bond angle outliers are listed below:

Mol	Chain	Res	Type	Atoms		$Observed(^{o})$	$Ideal(^{o})$
1	В	820	ASP	CB-CG-OD2	7.49	125.04	118.30
1	В	79	ASP	CB-CG-OD2	7.42	124.98	118.30
1	А	248	ASP	CB-CG-OD2	7.18	124.76	118.30
1	А	820	ASP	CB-CG-OD2	6.83	124.45	118.30
1	А	318	ASP	CB-CG-OD2	6.68	124.31	118.30

There are no chirality outliers.

All (1) planarity outliers are listed below:

Mol	Chain	Res	Type	Group
1	А	320	PHE	Peptide



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	А	6639	0	6576	197	0
1	В	6640	0	6585	220	0
2	А	475	0	0	57	0
2	В	455	0	0	67	0
All	All	14209	0	13161	409	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 15.

The worst 5 of 409 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:615:GLU:HG3	1:B:323:ARG:HG3	1.37	1.06
1:A:702:TRP:NE1	1:A:714:THR:HA	1.73	1.03
1:A:660:ARG:HG2	2:A:1748:HOH:O	1.60	1.01
1:B:359:GLN:H	1:B:359:GLN:HE21	1.08	1.00
1:A:734:GLU:O	1:A:735:LEU:HB3	1.64	0.94

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	Perce	entiles
1	А	835/922~(91%)	765 (92%)	57 (7%)	13 (2%)	9	31
1	В	834/922~(90%)	758 (91%)	60 (7%)	16 (2%)	8	26

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Mol	Chain	Analysed	Favoured	Allowed	Outliers	Percentiles
All	All	1669/1844~(90%)	1523 (91%)	117 (7%)	29(2%)	9 29

5 of 29 Ramachandran outliers are listed below:

Mol	Chain	Res	Type
1	А	264	GLU
1	А	714	THR
1	А	715	ALA
1	А	716	ASP
1	В	695	ASP

5.3.2 Protein sidechains (i)

In the following table, the Percentiles column shows the percent side chain 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	А	696/755~(92%)	577~(83%)	119 (17%)	2 6
1	В	697/755~(92%)	572 (82%)	125~(18%)	2 5
All	All	1393/1510~(92%)	1149 (82%)	244 (18%)	2 6

5 of 244 residues with a non-rotameric sidechain are listed below:

Mol	Chain	Res	Type
1	А	818	MET
1	В	702	TRP
1	В	219	ARG
1	В	694	GLU
1	В	798	MET

Sometimes side chains can be flipped to improve hydrogen bonding and reduce clashes. 5 of 26 such side chains are listed below:

Mol	Chain	Res	Type
1	В	118	ASN
1	В	284	ASN
1	В	671	GLN

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Mol	Chain	Res	Type
1	В	250	HIS
1	В	293	ASN

5.3.3 RNA (i)

There are no RNA molecules in this entry.

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

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

5.5 Carbohydrates (i)

There are no monosaccharides in this entry.

5.6 Ligand geometry (i)

There are no ligands in this entry.

5.7 Other polymers (i)

There are no such residues in this entry.

5.8 Polymer linkage issues (i)

There are no chain breaks in this entry.



6 Fit of model and data (i)

6.1 Protein, DNA and RNA chains (i)

EDS was not executed - this section is therefore empty.

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

EDS was not executed - this section is therefore empty.

6.3 Carbohydrates (i)

EDS was not executed - this section is therefore empty.

6.4 Ligands (i)

EDS was not executed - this section is therefore empty.

6.5 Other polymers (i)

EDS was not executed - this section is therefore empty.

