

Full wwPDB NMR Structure Validation Report (i)

Apr 22, 2024 – 01:01 PM JST

PDB ID : 6J37 BMRB ID : 36232

Title : DNA minidumbbell structure of two CTTG repeats

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Deposited on : 2019-01-04

This is a Full wwPDB NMR Structure Validation 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/NMRValidationReportHelp
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

Percentile statistics : 20191225.v01 (using entries in the PDB archive December 25th 2019)

wwPDB-RCI : v 1n 11 5 13 A (Berjanski et al., 2005)

PANAV : Wang et al. (2010)

wwPDB-ShiftChecker : v1.2

Ideal geometry (proteins) : Engh & Huber (2001) Ideal geometry (DNA, RNA) : Parkinson et al. (1996)

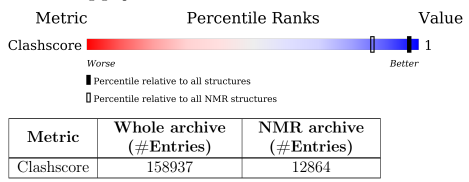
Validation Pipeline (wwPDB-VP) : 2.36.2

1 Overall quality at a glance (i)

The following experimental techniques were used to determine the structure: $SOLUTION\ NMR$

The overall completeness of chemical shifts assignment is 52%.

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.



The table below summarises the geometric issues observed across the polymeric chains and their fit to the experimental data. The red, orange, yellow and green segments indicate the fraction of residues that contain outliers for >=3, 2, 1 and 0 types of geometric quality criteria. A cyan segment indicates the fraction of residues that are not part of the well-defined cores, and 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%

Mol	Chain	Length	Quality of chain					
1	A	8	12%	38%	50%			



2 Ensemble composition and analysis (i)

This entry contains 20 models. This entry does not contain polypeptide chains, therefore identification of well-defined residues and clustering analysis are not possible. All residues are included in the validation scores.



3 Entry composition (i)

There are 2 unique types of molecules in this entry. The entry contains 260 atoms, of which 94 are hydrogens and 0 are deuteriums.

• Molecule 1 is a DNA chain called DNA (5'-D(*CP*TP*TP*GP*CP*TP*TP*G)-3').

Mol	Chain	Residues	Atoms						Trace
1	Λ	0	Total	С	Н	N	О	Р	0
1	A	0	253	78	94	24	50	7	U

• Molecule 2 is SODIUM ION (three-letter code: NA) (formula: Na).

Mol	Chain	Residues	Atoms
9	Λ	7	Total Na
	A	1	7 7



4 Residue-property plots (i)

4.1 Average score per residue in the NMR ensemble

These plots are provided for all protein, RNA, DNA and oligosaccharide chains in the entry. The first graphic is the same as shown in the summary in section 1 of this report. The second graphic shows the sequence where residues are colour-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 outliers are shown as green connectors. Residues which are classified as ill-defined in the NMR ensemble, are shown in cyan with an underline colour-coded according to the previous scheme. Residues which were present in the experimental sample, but not modelled in the final structure are shown in grey.

• Molecule 1: DNA (5'-D(*CP*TP*TP*GP*CP*TP*TP*G)-3')

Chain A: 12% 38% 50%

4.2 Scores per residue for each member of the ensemble

Colouring as in section 4.1 above.

4.2.1 Score per residue for model 1

 \bullet Molecule 1: DNA (5'-D(*CP*TP*TP*GP*CP*TP*TP*G)-3')

Chain A: 12% 38% 50%

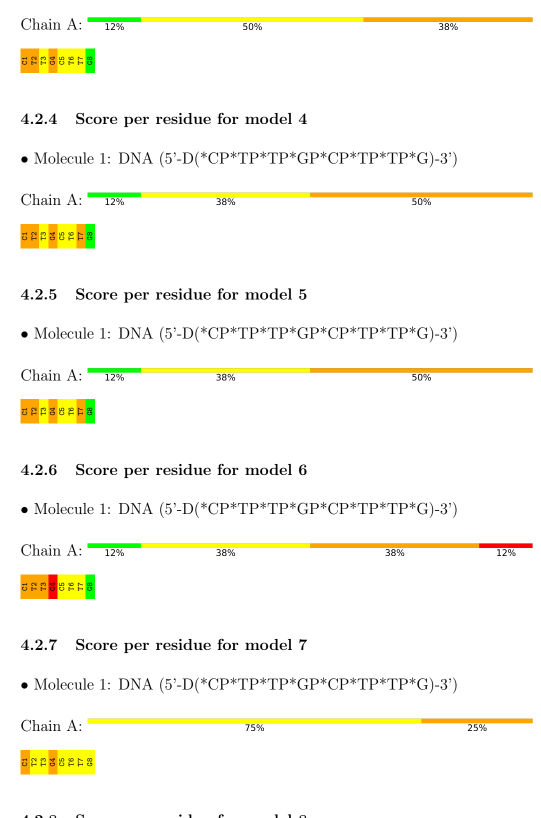
4.2.2 Score per residue for model 2

• Molecule 1: DNA (5'-D(*CP*TP*TP*GP*CP*TP*TP*G)-3')

Chain A: 12% 38% 50%

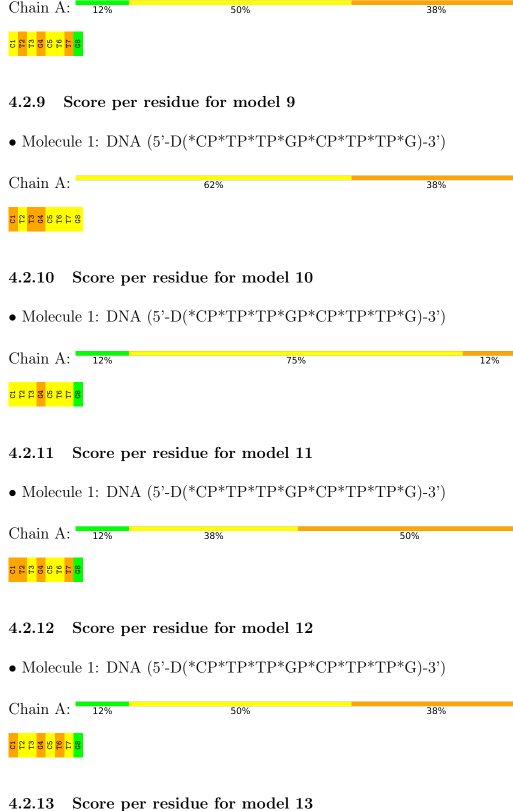
4.2.3 Score per residue for model 3



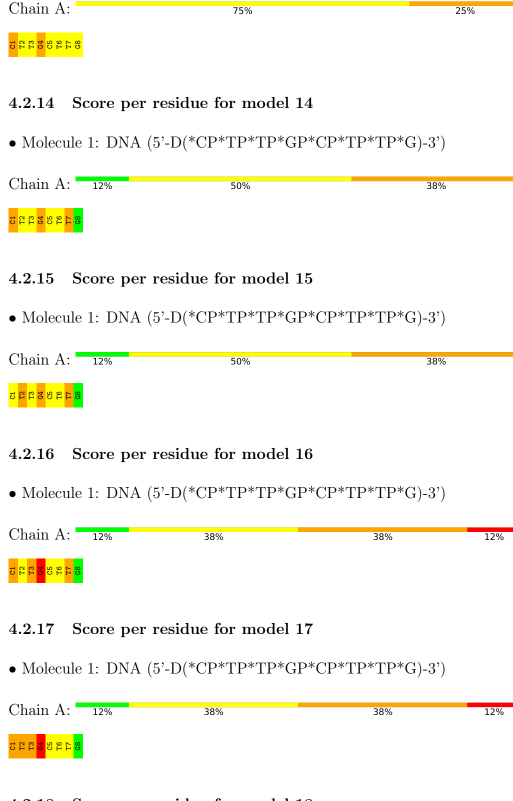


4.2.8 Score per residue for model 8









4.2.18 Score per residue for model 18



C1 T2 T3 C5 C5 C5 T6 T7 T7

Chain A: 12% 50% 38% C1 T2 T3 C5 C5 T6 T7 Score per residue for model 19 4.2.19• Molecule 1: DNA (5'-D(*CP*TP*TP*GP*CP*TP*TP*G)-3') Chain A: 12% 50% 38% C1 T2 T3 G4 C5 C5 T6 T7 Score per residue for model 204.2.20 \bullet Molecule 1: DNA (5'-D(*CP*TP*TP*GP*CP*TP*TP*G)-3') Chain A: 12% 38% 50%



5 Refinement protocol and experimental data overview (i)



The models were refined using the following method: not applicable.

Of the 1000 calculated structures, 20 were deposited, based on the following criterion: structures with the least restraint violations.

The following table shows the software used for structure solution, optimisation and refinement.

Software name	Classification	Version
Amber	structure calculation	

The following table shows chemical shift validation statistics as aggregates over all chemical shift files. Detailed validation can be found in section 7 of this report.

Chemical shift file(s)	working_cs.cif
Number of chemical shift lists	1
Total number of shifts	92
Number of shifts mapped to atoms	92
Number of unparsed shifts	0
Number of shifts with mapping errors	0
Number of shifts with mapping warnings	0
Assignment completeness (well-defined parts)	52%



6 Model quality (i)

6.1 Standard geometry (i)

Bond lengths and bond angles in the following residue types are not validated in this section: NA

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 (average) root-mean-square of all Z scores of the bond lengths (or angles).

Mol	Chain	В	Sond lengths	Bond angles		
MIOI	Chain	RMSZ	#Z>5	RMSZ	#Z>5	
1	A	1.57 ± 0.01	$1\pm0/176$ ($0.6\pm$ 0.1%)	2.23 ± 0.04	$10\pm2/270$ ($3.8\pm$ 0.6%)	
All	All	1.57	21/3520 ($0.6%$)	2.23	206/5400 (3.8%)	

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	Planarity
1	A	0.0 ± 0.0	3.4 ± 0.6
All	All	0	68

All unique bond outliers are listed below. They are sorted according to the Z-score of the worst occurrence in the ensemble.

Mal	Chain	Pog	Type	Atoms	7	$Observed(\mathring{A})$	Ideal(Å)	Mod	dels
MIOI	Cham	nes	Type	Atoms		Observed(A)	Ideal(A)	Worst	Total
1	A	5	DC	C4'-O4'	-5.42	1.39	1.45	5	20
1	A	4	DG	C4'-O4'	-5.03	1.40	1.45	14	1

All unique angle outliers are listed below. They are sorted according to the Z-score of the worst occurrence in the ensemble.

Mol	Iol Chain Re		Type	Atoms	$f Z = f Observed(^o)$		$Ideal(^{o})$	Models	
IVIOI	Chain	1605	Type	Atoms		Obscived()	ideai()	Worst	Total
1	A	4	DG	O4'-C1'-N9	9.96	114.97	108.00	12	20
1	A	7	DT	C6-C5-C7	-9.23	117.36	122.90	9	20
1	A	6	DT	O4'-C1'-N1	9.10	114.37	108.00	1	20
1	A	6	DT	P-O3'-C3'	7.62	128.85	119.70	12	1
1	A	3	DT	C6-C5-C7	-7.22	118.57	122.90	10	20
1	A	6	DT	C6-C5-C7	-6.95	118.73	122.90	2	20

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Mol	ol Chain Re		Tuno	Atoms	\mathbf{z}	$Observed(^{o})$	$Ideal(^{o})$	Models	
IVIOI	Chain	Res	Type	Atoms		Observed()	ideai()	Worst	Total
1	A	2	DT	C6-C5-C7	-6.90	118.76	122.90	6	20
1	A	2	DT	P-O3'-C3'	6.72	127.77	119.70	1	20
1	A	1	DC	P-O3'-C3'	6.33	127.30	119.70	3	8
1	A	2	DT	O4'-C1'-N1	6.31	112.42	108.00	3	11
1	A	4	DG	O4'-C4'-C3'	6.16	109.69	106.00	7	15
1	A	5	DC	P-O3'-C3'	5.39	126.17	119.70	20	15
1	A	1	DC	N1-C2-O2	5.39	122.13	118.90	13	11
1	A	1	DC	O4'-C1'-N1	5.27	111.69	108.00	13	2
1	A	7	DT	C4-C5-C7	5.19	122.11	119.00	8	2
1	A	3	DT	O4'-C1'-N1	5.01	111.51	108.00	16	1

There are no chirality outliers.

All unique planar outliers are listed below. They are sorted by the frequency of occurrence in the ensemble.

Mol	Chain	Res	Type	Group	Models (Total)
1	A	1	DC	Sidechain	20
1	A	4	DG	Sidechain	20
1	A	2	DT	Sidechain	11
1	A	7	DT	Sidechain	10
1	A	8	DG	Sidechain	3
1	A	6	DT	Sidechain	2
1	A	3	DT	Sidechain	2

6.2 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 each 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 averaged over the ensemble.

Mol	Chain	Non-H	H(model)	H(added)	Clashes
1	A	159	94	94	0±0
All	All	3320	1880	1880	3

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 1.

All unique clashes are listed below, sorted by their clash magnitude.



Atom-1	Atom-2	Clash(Å)	Distance(Å)	Models	
Atom-1				Worst	Total
1:A:3:DT:H1'	1:A:4:DG:C5	0.41	2.51	6	3

6.3 Torsion angles (i)

6.3.1 Protein backbone (i)

There are no protein molecules in this entry.

6.3.2 Protein sidechains (i)

There are no protein molecules in this entry.

6.3.3 RNA (i)

There are no RNA molecules in this entry.

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

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

6.5 Carbohydrates (i)

There are no monosaccharides in this entry.

6.6 Ligand geometry (i)

Of 7 ligands modelled in this entry, 7 are monoatomic - leaving 0 for Mogul analysis.

6.7 Other polymers (i)

There are no such molecules in this entry.

6.8 Polymer linkage issues (i)

There are no chain breaks in this entry.



7 Chemical shift validation (i)

The completeness of assignment taking into account all chemical shift lists is 52% for the well-defined parts and 52% for the entire structure.

7.1 Chemical shift list 1

File name: working cs.cif

Chemical shift list name: cttg2_cs_deposit.txt

7.1.1 Bookkeeping (i)

The following table shows the results of parsing the chemical shift list and reports the number of nuclei with statistically unusual chemical shifts.

Total number of shifts	92
Number of shifts mapped to atoms	92
Number of unparsed shifts	0
Number of shifts with mapping errors	0
Number of shifts with mapping warnings	0
Number of shift outliers (ShiftChecker)	0

7.1.2 Chemical shift referencing (i)

No chemical shift referencing corrections were calculated (not enough data).

7.1.3 Completeness of resonance assignments (i)

The following table shows the completeness of the chemical shift assignments for the well-defined regions of the structure. The overall completeness is 52%, i.e. 80 atoms were assigned a chemical shift out of a possible 154. 0 out of 0 assigned methyl groups (LEU and VAL) were assigned stereospecifically.

	Total	$^{1}{ m H}$	$^{13}\mathbf{C}$	$^{15}{ m N}$
Sugar	56/96~(58%)	56/56 (100%)	0/40 (0%)	0/0 (%)
Base	24/58 (41%)	24/34 (71%)	0/14 (0%)	0/10 (0%)
Overall	80/154 (52%)	80/90 (89%)	0/54 (0%)	0/10 (0%)

The following table shows the completeness of the chemical shift assignments for the full structure. The overall completeness is 52%, i.e. 80 atoms were assigned a chemical shift out of a possible 154. 0 out of 0 assigned methyl groups (LEU and VAL) were assigned stereospecifically.



	Total	$^{1}{ m H}$	$^{13}\mathbf{C}$	$^{15}\mathbf{N}$
Sugar	56/96 (58%)	56/56 (100%)	0/40 (0%)	0/0 (%)
Base	24/58 (41%)	24/34 (71%)	0/14 (0%)	0/10 (0%)
Overall	80/154 (52%)	80/90 (89%)	0/54~(0%)	0/10 (0%)

7.1.4 Statistically unusual chemical shifts (i)

There are no statistically unusual chemical shifts.

7.1.5 Random Coil Index (RCI) plots (i)

No $random\ coil\ index(RCI)$ plot could be generated from the current chemical shift list. RCI is only applicable to proteins

