



Central European Institute of Technology BRNO | CZECH REPUBLIC

Validation of biomacromolecular structures - motivation

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Validation: Why to validate?

Structural biology community found that some published structures contained serious errors

Nightmare before Christmas

Retraction

WE WISH TO RETRACT OUR RESEARCH ARTICLE "STRUCTURE OF MsbA from *E. coli*: A homolog of the multidrug resistance ATP binding cassette (ABC) transporters" and both of our Reports "Structure of the ABC transporter MsbA in complex with ADP•vanadate and lipopolysaccharide" and "X-ray structure of the EmrE multidrug transporter in complex with a substrate" (1-3).

The recently reported structure of Sav 1866 (4) indicated that our MsbA structures (1, 2, 5) were incorrect in both the hand of the structure and the topology. Thus, our biological interpretations based on these inverted models for MsbA are invalid.

An in-house data reduction program introduced a change in sign for anomalous differences. This program, which was not part of a conventional data processing package, converted the anomalous pairs (I+ and I-) to (F- and F+), thereby introducing a sign change. As the diffraction data collected for each set of MsbA crystals and for the EmrE crystals were processed with the same program, the structures reported in (*I*-3, 5, 6) had the wrong hand.

The error in the topology of the original MsbA structure was a consequence of the low resolution of the data as well as breaks in the elec-

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tron density for the connecting loop regions. Unfortunately, the use of the multicopy refinement procedure still allowed us to obtain reasonable refinement values for the wrong structures.

The Protein Data Bank (PDB) files 1JSQ, 1PF4, and 1Z2R for MsbA and 1S7B and 2F2M for EmrE have been moved to the archive of obsolete PDB entries. The MsbA and EmrE structures will be recalculated from the original data using the proper sign for the anomalous differences, and the new C α coordinates and structure factors will be deposited.

We very sincerely regret the confusion that these papers have caused and, in particular, subsequent research efforts that were unproductive as a result of our original findings.

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References

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- R. J. Dawson, K. P. Locher, Nature 443, 180 (2006).
- 5. G. Chang, J. Mol. Biol. 330, 419 (2003).
- 6. C. Ma, G. Chang, Proc. Natl. Acad. Sci. U.S.A. 101, 2852 (2004).

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Validation: Why to validate?

Garbage in, garbage out

Interesting articles:

- Matthews, B. W. (2007) Five retracted structure reports: inverted or incorrect? Protein science : a publication of the Protein Society, 16, 1013–6.
- Johnston, C. A., Kimple, A. J., Giguere, P. M., and Siderovski, D. P. (2008) RETRACTED: Structure of the Parathyroid Hormone Receptor C Terminus Bound to the G-Protein Dimer Gb1g2. Structure, 16, 1086–1094.



Can we still find wrong structures in PDB?



Can we still find wrong structures in PDB?

Example: Nipah G attachment glycoprotein (PDB ID 3D12, PNAS)

Contains 30 instances of 11 different carbohydrates, each with one ring and five chiral atoms.

Results:

- 13 of these ligands have incorrect chirality
- In a few cases, all chiral atoms exhibit incorrect chirality





Can we still find wrong structures in PDB?

Unfortunately yes



Validation approaches



PDB validation reports



Improved data quality

data

- Cleaned up mmCIF data
 - Standard vocabularies
 - Experimental details, binding sites, secondary structure, antibiotic/inhibitor information, nucleic-acid parameters
- Clean mmCIF files are used in production





Validation reports

Summary

- Quality vs. all PDB
- Quality vs. entries at similar resolution
- Overview of residuebased quality for every polymer
- Table of ligands that may need your attention

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 1.80 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 $(\#Entries)$	${f Similar\ resolution}\ (\#{ m Entries,\ resolution\ range}({ m \AA}))$
R_{free}	91344	4533 (1.80-1.80)
Clashscore	102246	5383 (1.80-1.80)
Ramachandran outliers	100387	5320 (1.80-1.80)
Sidechain outliers	100360	5319 (1.80-1.80)
RSRZ outliers	91569	4547 (1.80-1.80)

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 >=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 <=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	А	137	91%	9%	

Mol	Type	Chain	Res	Chirality	Geometry	Clashes	Electron density
1	PAQ	А	471	Х	-	-	-
1	PAQ	В	471	Х	-	-	-
1	PAQ	С	471	Х	-	-	-
1	PAQ	D	471	Х	-	-	-
6	CU	А	1744	-	-	-	Х
6	CU	В	1748	-	-	-	Х



Topology validation (= validation of annotation)





Topology validation – basic terms

Residue

- Any component of a biomacromolecule
- Examples: amino acids, nucleotides, saccharides, ions, ...
- In PDB, residue is annotated via "Residue ID" a unique 3-letter code



PHE, phenylalanine



FUC, alpha-L-fucose

Topology validation – basic terms

Types of residues:

Standard residues:

- amino acids
- nucleotides

Non-standard residues:

modified amino acids and nucleotides

Ligands:

- Chemical compounds which form a complex with a biomacromolecule (e.g., sugar, drug, heme).
- Also ions are often referred as ligands



Topology validation – basic terms

Principles of topology validation:

- Subjects of topology validation are residues
- Validated residue is compared with a model residue, which has the same Residue ID
- The model residues are taken from a reference database
- Differences between the model residue and the validated residue are reported



Topology validation – approach

Complication with "shared atoms":

When some residues bind together, one of them can lose an atom:



 Solution: When we validate a residue, we must include also its close surrounding

Topology validation - approach

Input PDB entry



Mapping of **input motif** to the **model residue** (via subgraph matching)



Validated motif mapped to the Model Residue



Input motif = validated residue + surrounding

Topology validation – types of validation analyses

Completeness analyses

- Missing atoms
- Missing rings

Chirality analyses

 Chirality on C atom, metal atom, high bond order atom, planarity

Advanced analyses

- Substitution
- Different atom naming
- Foreign atoms



Topology validation – types of validation analyses Prerequisite: Can we map the validated and the model residue?



Degenerated structure

1IVG_17_7716 (MAN)





Topology validation – types of validation analyses ERRORS – INCOMPLETE STRUCTURE





Topology validation – types of validation analyses ERRORS – CHIRALITY



1E4M_16_4280 (MAN)





Topology validation – types of validation analyses ERRORS – CHIRALITY II

Wrong chirality on atom having high order bonds





Correct chirality (Tolerant)

Correct residues

residues having only these issues:

- Wrong chirality (planar)
- Wrong chirality on atom having high order bonds



Topology validation – types of validation analyses WARNINGS



 \square CEITEC

Different atom names - example

4ACQ_37_39697 (NAG)



Different Atom Names 7

Model	Motif	Туре
C1	C3 C 39699	N N
C3	C1 C 39697	N N'
C4	C6 C 39702	N N'
C6	C4 C 39700	N N'
03	O1 O 39706	N N'
04	O6 O 39709	N E
06	O4 O 39707	N E



0

Topology validation Software tools

PDB care (Lütteke et al., 2006):

- Tool focused on carbohydrates validation
- First application, which implements topology validation
- Performs basic validation analyses (missing atoms, missing rings, wrong chirality)

MotiveValidator (http://ncbr.muni.cz/MotiveValidator) :

- Tool, which allows validation of all residues
- Performs basic validation analyses + reports basic warnings (substitutions, foreign atoms, different naming)

ValidatorDB (http://ncbr.muni.cz/ValidatorDB):

- Database, containing validation results for all* ligands and non-standard residues in PDB (weekly updated)
- Performs basic validation analyses + advanced validation analyses (report degenerated residue, distinguish type of chirality error)
- Reports basic warnings + next warnings (Alternate locations, Zero RMSD with model)
- * Except amino acids, nucleotides, and small residues (<7 heavy atoms)



Thank you for your attention

