

Integrative Structure Validation Report ?

March 27, 2025 - 10:11 AM PDT

The following software was used in the production of this report:

Integrative Modeling Validation Version 2.0

Python-IHM Version 1.8

MolProbity Version 4.5.2

| | |
|-------------------|---|
| PDB ID | 9A3K |
| PDB-Dev ID | PDBDEV_00000205 |
| Structure Title | Parathyroid hormone receptor type 1 in complex with a long-acting parathyroid hormone analog and arrestin 2 (5w0p-based template) |
| Structure Authors | Aydin, Y.; Bottke, T.; Lam, J.H.; Ernicke, S.; Fortmann, A.; Tretbar, M.; Zarzycka, B.; Gurevich, V.V.; Katritch, V.; Coin, I. |
| Deposited on | 2023-03-24 |

This is a PDB-IHM IM Structure Validation Report for a publicly released PDB-IHM entry.

We welcome your comments at helpdesk@pdb-ihm.org

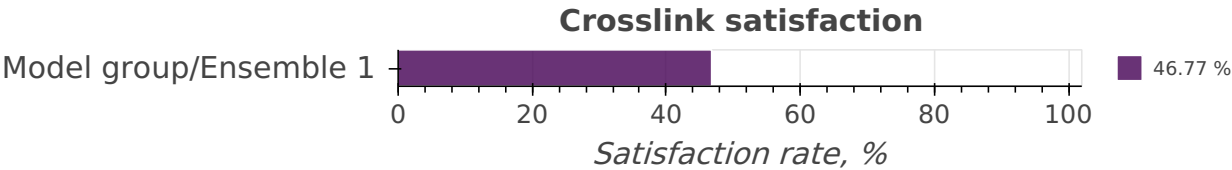
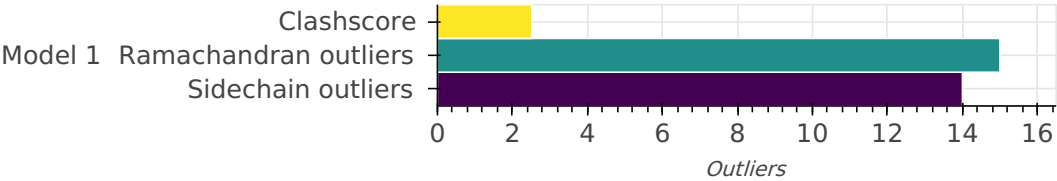
A user guide is available at https://pdb-ihm.org/validation_help.html with specific help available everywhere you see the ? symbol.

List of references used to build this report is available [here](#).

Overall quality ?

This validation report contains model quality assessments for all structures, data quality and fit to model assessments for SAS and crosslinking-MS datasets. Data quality and fit to model assessments for other datasets and model uncertainty are under development. Number of plots is limited to 256.

Model Quality: MolProbity Analysis



Ensemble information ?

This entry consists of 0 distinct ensemble(s).

Summary ?

This entry consists of 1 model(s). A total of 6 datasets were used to build this entry.

Representation ?

This entry has 1 representation(s).

| ID | Model(s) | Entity ID | Molecule name | Chain(s) [auth] | Total residues | Rigid segments | Flexible segments | Model coverage/ Starting model coverage (%) | Scale |
|----|----------|-----------|--|-----------------|----------------|----------------|-------------------|---|--------|
| 1 | 1 | 1 | Arrestin2 | A | 357 | - | 1-357 | 100.00 / 100.00 | Atomic |
| | | 2 | Long-acting parathyroid hormone analog | B | 32 | - | 1-32 | 100.00 / 100.00 | Atomic |
| | | 3 | PTH1R | C [P] | 504 | - | 1-504 | 100.00 / 100.00 | Atomic |

Datasets used for modeling ?

There are 6 unique datasets used to build the models in this entry.

| ID | Dataset type | Database name | Data access code |
|----|----------------------|---------------|----------------------|
| 1 | Crosslinking-MS data | Not available | Not available |
| 2 | Experimental model | PDB | 6NBF |
| 3 | Experimental model | PDB | 5W0P |
| 4 | Comparative model | Not available | Not available |
| 5 | Comparative model | Not available | Not available |
| 6 | De Novo model | Not available | Not available |

Methodology and software ?

This entry is a result of 1 distinct protocol(s).

| Step number | Protocol ID | Method name | Method type | Method description | Number of computed models | Multi state modeling | Multi scale modeling |
|-------------|-------------|-------------|-------------|--------------------|---------------------------|----------------------|----------------------|
| 1 | 1 | None | None | None | None | False | False |

There is 1 software package reported in this entry.

| ID | Software name | Software version | Software classification | Software location |
|----|-------------------------|------------------|-------------------------|---|
| 1 | ICM-Pro | v.3.9.2c | Model building | https://www.molsoft.com/icm_pro.html |

Data quality ?

Crosslinking-MS

At the moment, data validation is only available for crosslinking-MS data deposited as a fully [compliant](#) dataset in the [PRIDE Crosslinking](#) database. Correspondence between crosslinking-MS and entry entities is established using [pyHMMER](#). Only residue pairs that passed the reported threshold are used for the analysis. The values in the report have to be interpreted in the context of the experiment (i.e. only a minor fraction of in-situ or in-vivo dataset can be used for modeling).

Crosslinking-MS dataset is not available in the [PRIDE Crosslinking](#) database.

Model quality ?

For models with atomic structures, MolProbity analysis is performed. For models with coarse-grained or multi-scale structures, excluded volume analysis is performed.

Standard geometry: bond outliers

There are 63 bond length outliers in this entry (0.86% of 7323 assessed bonds). A summary is provided below.

| Chain | Res | Type | Atoms | Z | Observed (Å) | Ideal (Å) | Model ID (Worst) | Models (Total) |
|-------|-----|------|---------|------|--------------|-----------|------------------|----------------|
| C | 477 | TPO | OG1-P | 9.77 | 1.52 | 1.72 | 1 | 1 |
| C | 463 | SEP | O3P-P | 4.82 | 1.51 | 1.61 | 1 | 1 |
| C | 467 | SEP | O3P-P | 4.82 | 1.51 | 1.61 | 1 | 1 |
| C | 493 | SEP | O1P-P | 4.82 | 1.51 | 1.61 | 1 | 1 |
| C | 478 | SEP | O2P-P | 4.81 | 1.51 | 1.61 | 1 | 1 |
| C | 478 | SEP | O1P-P | 4.81 | 1.51 | 1.61 | 1 | 1 |
| C | 493 | SEP | O2P-P | 4.80 | 1.51 | 1.61 | 1 | 1 |
| C | 467 | SEP | O2P-P | 4.80 | 1.51 | 1.61 | 1 | 1 |
| C | 463 | SEP | O2P-P | 4.80 | 1.51 | 1.61 | 1 | 1 |
| C | 493 | SEP | O3P-P | 4.79 | 1.51 | 1.61 | 1 | 1 |
| C | 467 | SEP | O1P-P | 4.78 | 1.51 | 1.61 | 1 | 1 |
| C | 478 | SEP | O3P-P | 4.78 | 1.51 | 1.61 | 1 | 1 |
| C | 463 | SEP | O1P-P | 4.75 | 1.51 | 1.61 | 1 | 1 |
| C | 493 | SEP | OG-P | 4.40 | 1.52 | 1.61 | 1 | 1 |
| C | 463 | SEP | OG-P | 4.39 | 1.52 | 1.61 | 1 | 1 |
| C | 478 | SEP | OG-P | 4.39 | 1.52 | 1.61 | 1 | 1 |
| C | 467 | SEP | OG-P | 4.36 | 1.52 | 1.61 | 1 | 1 |
| C | 117 | HIS | CE1-NE2 | 4.36 | 1.36 | 1.32 | 1 | 1 |
| C | 190 | HIS | CE1-NE2 | 4.36 | 1.36 | 1.32 | 1 | 1 |
| A | 111 | HIS | CE1-NE2 | 4.36 | 1.36 | 1.32 | 1 | 1 |
| B | 25 | HIS | CE1-NE2 | 4.36 | 1.36 | 1.32 | 1 | 1 |
| C | 88 | HIS | CE1-NE2 | 4.36 | 1.36 | 1.32 | 1 | 1 |
| A | 30 | HIS | CE1-NE2 | 4.35 | 1.36 | 1.32 | 1 | 1 |
| C | 197 | HIS | CE1-NE2 | 4.35 | 1.36 | 1.32 | 1 | 1 |
| A | 159 | HIS | CE1-NE2 | 4.34 | 1.36 | 1.32 | 1 | 1 |
| B | 9 | HIS | CE1-NE2 | 4.33 | 1.36 | 1.32 | 1 | 1 |
| C | 476 | HIS | CD2-NE2 | 4.33 | 1.33 | 1.37 | 1 | 1 |
| A | 111 | HIS | CD2-NE2 | 4.33 | 1.33 | 1.37 | 1 | 1 |
| B | 25 | HIS | CD2-NE2 | 4.32 | 1.33 | 1.37 | 1 | 1 |
| A | 198 | HIS | CE1-NE2 | 4.32 | 1.36 | 1.32 | 1 | 1 |

| Chain | Res | Type | Atoms | Z | Observed (Å) | Ideal (Å) | Model ID (Worst) | Models (Total) |
|-------|-----|------|---------|------|--------------|-----------|------------------|----------------|
| C | 134 | HIS | CE1-NE2 | 4.32 | 1.36 | 1.32 | 1 | 1 |
| C | 114 | HIS | CE1-NE2 | 4.32 | 1.36 | 1.32 | 1 | 1 |
| A | 219 | HIS | CE1-NE2 | 4.31 | 1.36 | 1.32 | 1 | 1 |
| A | 159 | HIS | CD2-NE2 | 4.31 | 1.33 | 1.37 | 1 | 1 |
| C | 114 | HIS | CD2-NE2 | 4.31 | 1.33 | 1.37 | 1 | 1 |
| A | 353 | HIS | CD2-NE2 | 4.31 | 1.33 | 1.37 | 1 | 1 |
| C | 16 | HIS | CE1-NE2 | 4.30 | 1.36 | 1.32 | 1 | 1 |
| C | 88 | HIS | CD2-NE2 | 4.30 | 1.33 | 1.37 | 1 | 1 |
| C | 197 | HIS | CD2-NE2 | 4.30 | 1.33 | 1.37 | 1 | 1 |
| C | 394 | HIS | CE1-NE2 | 4.30 | 1.36 | 1.32 | 1 | 1 |
| C | 134 | HIS | CD2-NE2 | 4.29 | 1.33 | 1.37 | 1 | 1 |
| C | 476 | HIS | CE1-NE2 | 4.29 | 1.36 | 1.32 | 1 | 1 |
| C | 190 | HIS | CD2-NE2 | 4.29 | 1.33 | 1.37 | 1 | 1 |
| C | 16 | HIS | CD2-NE2 | 4.28 | 1.33 | 1.37 | 1 | 1 |
| A | 353 | HIS | CE1-NE2 | 4.28 | 1.36 | 1.32 | 1 | 1 |
| A | 210 | HIS | CE1-NE2 | 4.26 | 1.36 | 1.32 | 1 | 1 |
| A | 295 | HIS | CE1-NE2 | 4.26 | 1.36 | 1.32 | 1 | 1 |
| A | 198 | HIS | CD2-NE2 | 4.26 | 1.33 | 1.37 | 1 | 1 |
| A | 219 | HIS | CD2-NE2 | 4.26 | 1.33 | 1.37 | 1 | 1 |
| C | 281 | HIS | CE1-NE2 | 4.26 | 1.36 | 1.32 | 1 | 1 |
| C | 416 | HIS | CE1-NE2 | 4.26 | 1.36 | 1.32 | 1 | 1 |
| A | 210 | HIS | CD2-NE2 | 4.26 | 1.33 | 1.37 | 1 | 1 |
| C | 199 | HIS | CD2-NE2 | 4.25 | 1.33 | 1.37 | 1 | 1 |
| C | 117 | HIS | CD2-NE2 | 4.25 | 1.33 | 1.37 | 1 | 1 |
| B | 32 | HIS | CD2-NE2 | 4.25 | 1.33 | 1.37 | 1 | 1 |
| A | 295 | HIS | CD2-NE2 | 4.24 | 1.33 | 1.37 | 1 | 1 |
| B | 32 | HIS | CE1-NE2 | 4.24 | 1.36 | 1.32 | 1 | 1 |
| A | 30 | HIS | CD2-NE2 | 4.24 | 1.33 | 1.37 | 1 | 1 |
| C | 199 | HIS | CE1-NE2 | 4.23 | 1.36 | 1.32 | 1 | 1 |
| C | 281 | HIS | CD2-NE2 | 4.22 | 1.33 | 1.37 | 1 | 1 |
| C | 416 | HIS | CD2-NE2 | 4.22 | 1.33 | 1.37 | 1 | 1 |
| C | 394 | HIS | CD2-NE2 | 4.22 | 1.33 | 1.37 | 1 | 1 |
| B | 9 | HIS | CD2-NE2 | 4.21 | 1.33 | 1.37 | 1 | 1 |

Standard geometry: angle outliers ?

There are no bond angle outliers.

Too-close contacts ?

The following all-atom clashscore is based on a MolProbity analysis. All-atom clashscore is defined as the number of clashes found per 1000 atoms (including hydrogen atoms). The table below contains clashscores for all atomic models in this entry.

| Model ID | Clash score | Number of clashes |
|----------|-------------|-------------------|
| 1 | 2.52 | 36 |

There are 36 clashes. The table below contains the detailed list of all clashes based on a MolProbity analysis. Bad clashes are ≥ 0.4 Angstrom.

| Atom 1 | Atom 2 | Clash(Å) | Model ID (Worst) | Models (Total) |
|---------------|----------------|----------|------------------|----------------|
| A:150:CYS:O | A:151:ALA:CB | 0.73 | 1 | 1 |
| A:149:PHE:CD1 | A:149:PHE:N | 0.61 | 1 | 1 |
| A:89:PRO:O | A:90:ALA:HB3 | 0.60 | 1 | 1 |
| C:62:TYR:N | C:63:PRO:HD2 | 0.59 | 1 | 1 |
| C:165:TYR:C | C:165:TYR:CD1 | 0.59 | 1 | 1 |
| A:197:LEU:C | A:197:LEU:HD23 | 0.58 | 1 | 1 |
| C:71:ALA:N | C:72:PRO:CD | 0.57 | 1 | 1 |
| A:150:CYS:O | A:151:ALA:HB2 | 0.56 | 1 | 1 |
| C:165:TYR:CD1 | C:166:THR:N | 0.54 | 1 | 1 |
| C:339:VAL:N | C:340:PRO:CD | 0.53 | 1 | 1 |
| C:305:LEU:N | C:306:PRO:CD | 0.53 | 1 | 1 |
| C:459:ARG:O | C:460:LYS:C | 0.52 | 1 | 1 |
| A:90:ALA:HB1 | A:91:PRO:HD2 | 0.51 | 1 | 1 |
| C:369:GLY:O | C:370:ARG:CB | 0.49 | 1 | 1 |
| A:150:CYS:O | A:151:ALA:HB3 | 0.49 | 1 | 1 |
| C:366:THR:O | C:367:ASN:C | 0.49 | 1 | 1 |
| A:95:LYS:N | A:96:PRO:HD2 | 0.48 | 1 | 1 |
| C:62:TYR:N | C:63:PRO:CD | 0.47 | 1 | 1 |
| A:298:THR:OG1 | A:299:ASN:N | 0.47 | 1 | 1 |
| A:314:ILE:N | A:314:ILE:HD12 | 0.46 | 1 | 1 |
| A:328:VAL:O | A:329:VAL:HG23 | 0.46 | 1 | 1 |
| A:89:PRO:O | A:90:ALA:CB | 0.46 | 1 | 1 |
| C:62:TYR:HB3 | C:63:PRO:HD3 | 0.45 | 1 | 1 |
| C:478:SEP:OG | C:479:VAL:N | 0.45 | 1 | 1 |
| C:467:SEP:O2P | C:469:SER:N | 0.44 | 1 | 1 |
| C:321:ALA:O | C:322:ASN:C | 0.44 | 1 | 1 |
| A:148:ALA:C | A:149:PHE:CG | 0.43 | 1 | 1 |

| Atom 1 | Atom 2 | Clash(Å) | Model ID (Worst) | Models (Total) |
|--------------|----------------|----------|------------------|----------------|
| C:123:ASP:C | C:123:ASP:OD1 | 0.43 | 1 | 1 |
| A:87:PHE:HA | A:88:PRO:C | 0.42 | 1 | 1 |
| C:1:ASP:C | C:1:ASP:OD1 | 0.42 | 1 | 1 |
| C:388:MET:N | C:389:PRO:CD | 0.42 | 1 | 1 |
| A:197:LEU:O | A:197:LEU:HD23 | 0.41 | 1 | 1 |
| C:370:ARG:O | C:371:CYS:C | 0.41 | 1 | 1 |
| C:47:SER:O | C:48:THR:C | 0.41 | 1 | 1 |
| C:490:LEU:HA | C:491:PRO:C | 0.40 | 1 | 1 |
| C:85:GLU:OE2 | C:120:ARG:NH1 | 0.40 | 1 | 1 |

Torsion angles: Protein backbone ?

In the following table, Ramachandran outliers are listed. The Analysed column shows the number of residues for which the backbone conformation was analysed.

| Model ID | Analysed | Favored | Allowed | Outliers |
|----------|----------|---------|---------|----------|
| 1 | 874 | 811 | 48 | 15 |

There are 15 unique backbone outliers. Detailed list of outliers are tabulated below.

| Chain | Res | Type | Models (Total) |
|-------|-----|------|----------------|
| A | 89 | PRO | 1 |
| A | 90 | ALA | 1 |
| A | 149 | PHE | 1 |
| A | 151 | ALA | 1 |
| A | 152 | GLU | 1 |
| A | 329 | VAL | 1 |
| A | 335 | LEU | 1 |
| C | 149 | THR | 1 |
| C | 322 | ASN | 1 |
| C | 331 | GLY | 1 |
| C | 370 | ARG | 1 |
| C | 392 | GLY | 1 |
| C | 459 | ARG | 1 |
| C | 470 | TYR | 1 |
| C | 485 | ARG | 1 |

Torsion angles : Protein sidechains ?

In the following table, sidechain rotameric outliers are listed. The Analysed column shows the number of residues for which the sidechain conformation was analysed.

| Model ID | Analysed | Favored | Allowed | Outliers |
|----------|----------|---------|---------|----------|
| 1 | 766 | 725 | 27 | 14 |

There are 14 unique sidechain outliers. Detailed list of outliers are tabulated below.

| Chain | Res | Type | Models (Total) |
|-------|-----|------|----------------|
| A | 18 | LEU | 1 |
| A | 56 | THR | 1 |
| A | 75 | PHE | 1 |
| A | 100 | LEU | 1 |
| A | 102 | GLU | 1 |
| A | 138 | LYS | 1 |
| A | 149 | PHE | 1 |
| A | 159 | HIS | 1 |
| A | 215 | SER | 1 |
| A | 329 | VAL | 1 |
| C | 371 | CYS | 1 |
| C | 462 | ARG | 1 |
| C | 480 | THR | 1 |
| C | 503 | ASN | 1 |

Fit of model to data used for modeling ?

Fit of model(s) to crosslinking-MS data

Restraint types

Restraint types are summarized in the table below. Restraints assigned "*by-residue*" are interpreted as between CA atoms. Restraints between coarse-grained beads are indicated as "*coarse-grained*".

Restraint group represents a set of crosslinking restraints applied collectively in the modeling.

There are 136 crosslinking restraints combined in 136 restraint groups.

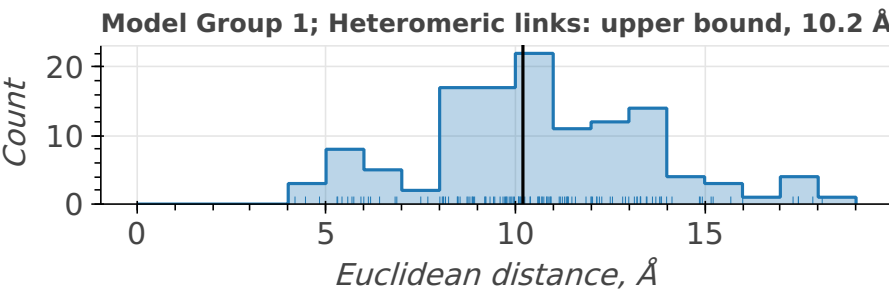
| Linker | Residue 1 | Atom 1 | Residue 2 | Atom 2 | Restraint type | Distance, Å | Count |
|--------|-----------|--------|-----------|--------|----------------|-------------|-------|
| BrEtY | ALA | CB | TYR | CB | upper bound | 10.2 | 1 |
| BrEtY | ALA | CB | ASP | CB | upper bound | 10.2 | 1 |
| BrEtY | ALA | CB | LYS | CB | upper bound | 10.2 | 3 |
| BrEtY | ALA | CB | LEU | CB | upper bound | 10.2 | 1 |
| BrEtY | ALA | CB | VAL | CB | upper bound | 10.2 | 1 |
| BrEtY | ALA | CB | HIS | CB | upper bound | 10.2 | 1 |

| Linker | Residue 1 | Atom 1 | Residue 2 | Atom 2 | Restraint type | Distance, Å | Count |
|--------|-----------|--------|-----------|--------|----------------|-------------|-------|
| BrEtY | ALA | CB | ASN | CB | upper bound | 10.2 | 1 |
| BrEtY | CYS | CB | THR | CB | upper bound | 10.2 | 1 |
| BrEtY | ASP | CB | CYS | CB | upper bound | 10.2 | 2 |
| BrEtY | CYS | CB | PHE | CB | upper bound | 10.2 | 1 |
| BrEtY | CYS | CB | LYS | CB | upper bound | 10.2 | 1 |
| BrEtY | ASP | CB | ASP | CB | upper bound | 10.2 | 1 |
| BrEtY | ASP | CB | PHE | CB | upper bound | 10.2 | 1 |
| BrEtY | ASP | CB | LYS | CB | upper bound | 10.2 | 1 |
| BrEtY | ASP | CB | VAL | CB | upper bound | 10.2 | 1 |
| BrEtY | ASP | CB | GLU | CB | upper bound | 10.2 | 1 |
| BrEtY | GLU | CB | GLU | CB | upper bound | 10.2 | 1 |
| BrEtY | GLU | CB | PHE | CB | upper bound | 10.2 | 1 |
| BrEtY | GLU | CB | LYS | CB | upper bound | 10.2 | 1 |
| BrEtY | GLU | CB | LEU | CB | upper bound | 10.2 | 1 |
| BrEtY | GLU | CB | VAL | CB | upper bound | 10.2 | 1 |
| BrEtY | PHE | CB | TYR | CB | upper bound | 10.2 | 1 |
| BrEtY | ASP | CB | GLY | CB | upper bound | 10.2 | 1 |
| BrEtY | GLY | CB | PHE | CB | upper bound | 10.2 | 1 |
| BrEtY | GLY | CB | VAL | CB | upper bound | 10.2 | 1 |
| BrEtY | GLY | CB | LYS | CB | upper bound | 10.2 | 5 |
| BrEtY | ASN | CB | GLY | CB | upper bound | 10.2 | 1 |
| BrEtY | GLY | CB | PRO | CB | upper bound | 10.2 | 1 |
| BrEtY | ARG | CB | GLY | CB | upper bound | 10.2 | 2 |
| BrEtY | LEU | CB | LYS | CB | upper bound | 10.2 | 3 |
| BrEtY | ARG | CB | LEU | CB | upper bound | 10.2 | 8 |
| BrEtY | LEU | CB | PRO | CB | upper bound | 10.2 | 1 |
| BrEtY | LYS | CB | MET | CB | upper bound | 10.2 | 1 |
| BrEtY | ASN | CB | ASP | CB | upper bound | 10.2 | 2 |
| BrEtY | ASN | CB | PHE | CB | upper bound | 10.2 | 1 |
| BrEtY | ASN | CB | LYS | CB | upper bound | 10.2 | 2 |
| BrEtY | ASN | CB | VAL | CB | upper bound | 10.2 | 2 |
| BrEtY | ASN | CB | LEU | CB | upper bound | 10.2 | 1 |
| BrEtY | ARG | CB | ASN | CB | upper bound | 10.2 | 2 |
| BrEtY | LYS | CB | PRO | CB | upper bound | 10.2 | 4 |
| BrEtY | ARG | CB | PRO | CB | upper bound | 10.2 | 8 |

| Linker | Residue 1 | Atom 1 | Residue 2 | Atom 2 | Restraint type | Distance, Å | Count |
|--------|-----------|--------|-----------|--------|----------------|-------------|-------|
| BrEtY | GLN | CB | PHE | CB | upper bound | 10.2 | 1 |
| BrEtY | GLN | CB | LYS | CB | upper bound | 10.2 | 1 |
| BrEtY | ARG | CB | ASP | CB | upper bound | 10.2 | 1 |
| BrEtY | ARG | CB | PHE | CB | upper bound | 10.2 | 2 |
| BrEtY | ARG | CB | LYS | CB | upper bound | 10.2 | 5 |
| BrEtY | ARG | CB | VAL | CB | upper bound | 10.2 | 4 |
| BrEtY | ARG | CB | HIS | CB | upper bound | 10.2 | 1 |
| BrEtY | ARG | CB | ARG | CB | upper bound | 10.2 | 4 |
| BrEtY | HIS | CB | SEP | CB | upper bound | 10.2 | 2 |
| BrEtY | LYS | CB | SEP | CB | upper bound | 10.2 | 5 |
| BrEtY | PRO | CB | SEP | CB | upper bound | 10.2 | 3 |
| BrEtY | LYS | CB | SER | CB | upper bound | 10.2 | 7 |
| BrEtY | PRO | CB | SER | CB | upper bound | 10.2 | 3 |
| BrEtY | HIS | CB | SER | CB | upper bound | 10.2 | 1 |
| BrEtY | SER | CB | SER | CB | upper bound | 10.2 | 1 |
| BrEtY | ASN | CB | SEP | CB | upper bound | 10.2 | 1 |
| BrEtY | ARG | CB | SEP | CB | upper bound | 10.2 | 2 |
| BrEtY | ASP | CB | THR | CB | upper bound | 10.2 | 1 |
| BrEtY | PHE | CB | THR | CB | upper bound | 10.2 | 2 |
| BrEtY | LYS | CB | THR | CB | upper bound | 10.2 | 2 |
| BrEtY | LEU | CB | THR | CB | upper bound | 10.2 | 1 |
| BrEtY | THR | CB | VAL | CB | upper bound | 10.2 | 1 |
| BrEtY | LYS | CB | TPO | CB | upper bound | 10.2 | 2 |
| BrEtY | ARG | CB | THR | CB | upper bound | 10.2 | 1 |
| BrEtY | LYS | CB | VAL | CB | upper bound | 10.2 | 2 |
| BrEtY | LYS | CB | TYR | CB | upper bound | 10.2 | 4 |
| BrEtY | ASN | CB | TYR | CB | upper bound | 10.2 | 2 |
| BrEtY | PRO | CB | TYR | CB | upper bound | 10.2 | 2 |

Distograms of individual restraints

Restraints with identical thresholds are grouped into one plot. Only the best distance per restraint per model group/ensemble is plotted. Inter- and intramolecular (including self-links) restraints are also grouped into one plot. Distance for a restraint between coarse-grained beads is calculated as a minimal distance between shells; if beads intersect, the distance will be reported as 0.0. A bead with the highest available resolution for a given residue is used for the assessment.



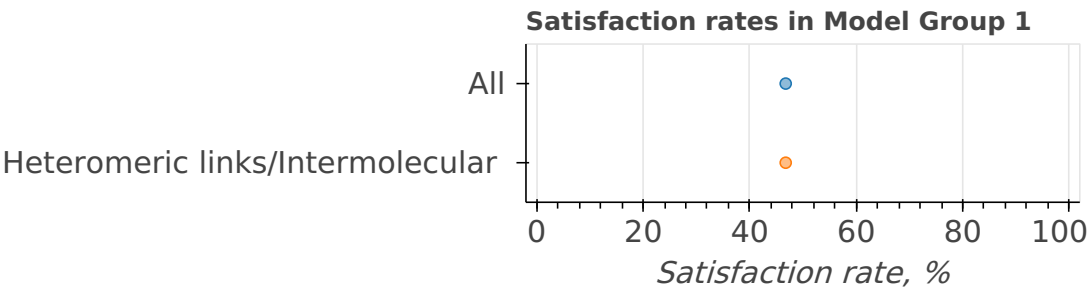
Satisfaction of restraints

Satisfaction of restraints is calculated on a [restraint group](#) (a set of crosslinking restraints applied collectively in the modeling) level. Satisfaction of a restraint group depends on satisfaction of individual restraints in the group and the conditionality (all/any). A restraint group is considered satisfied, if the condition was met in at least one model of the model group/ensemble. The number of measured restraints can be smaller than the total number of restraint groups if crosslinks involve non-modeled residues. Only deposited models are used for validation right now.

| State group | State | Model group | # of Deposited models/Total | Restraint group type | Satisfied (%) | Violated (%) | Count (Total=136) |
|-------------|-------|-------------|-----------------------------|-----------------------------------|---------------|--------------|-------------------|
| 1 | 1 | 1 | 1/1 | All | 46.77 | 53.23 | 124 |
| | | | | Heteromeric links/ Intermolecular | 46.77 | 53.23 | 124 |

Per-model satisfaction rates in ensembles

Every point represents one model in a model group/ensemble. Where possible, boxplots with quartile marks are also plotted.



Fit of model to data used for validation ?

Validation for this section is under development.

The development of integrative model validation metrics, implementation of a model validation pipeline, and creation of a validation report for integrative structures are funded by NSF awards to the [PDB-IHM team](#) (DBI-1756248, DBI-2112966, DBI-2112967, DBI-2112968, and DBI-1756250) and awards from NSF, NIH, and DOE to the [RCSB PDB](#) (DBI-2321666, R01GM157729, and DE-SC0019749). The PDB-IHM team and members of the [Sali lab](#) contributed model validation metrics and software packages.

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