

# wwPDB X-ray Structure Validation Summary Report (i)

#### Oct 5, 2023 – 03:49 AM EDT

PDB ID : 6VOT

Title : Crystal structure of Pseudomonas aerugonisa PBP3 complexed to gamma-

lactam YU253434

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Deposited on : 2020-01-31

Resolution : 2.40 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 (1)) were used in the production of this report:

MolProbity : FAILED

Mogul : 1.8.5 (274361), CSD as541be (2020)

Xtriage (Phenix) : 1.13

EDS : FAILED

buster-report : 1.1.7 (2018)

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) : 2.35.1

# 1 Overall quality at a glance (i)

The following experimental techniques were used to determine the structure: X-RAY DIFFRACTION

The reported resolution of this entry is 2.40 Å.

There are no overall percentile quality scores available for this entry.

MolProbity and EDS failed to run properly - the sequence quality summary graphics cannot be shown.



# 2 Entry composition (i)

There are 3 unique types of molecules in this entry. The entry contains 3951 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 Peptidoglycan D,D-transpeptidase FtsI.

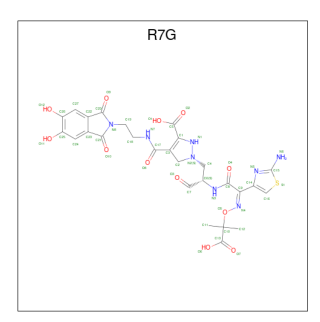
| Mo | l Chain | Residues | Atoms         |           |          |          | ZeroOcc | AltConf | Trace |   |
|----|---------|----------|---------------|-----------|----------|----------|---------|---------|-------|---|
| 1  | A       | 499      | Total<br>3818 | C<br>2413 | N<br>690 | O<br>702 | S<br>13 | 0       | 2     | 0 |

There are 8 discrepancies between the modelled and reference sequences:

| Chain | Residue | Modelled | Actual | Comment        | Reference  |
|-------|---------|----------|--------|----------------|------------|
| A     | 42      | MET      | -      | expression tag | UNP Q51504 |
| A     | 43      | GLY      | -      | expression tag | UNP Q51504 |
| A     | 44      | HIS      | -      | expression tag | UNP Q51504 |
| A     | 45      | HIS      | -      | expression tag | UNP Q51504 |
| A     | 46      | HIS      | -      | expression tag | UNP Q51504 |
| A     | 47      | HIS      | -      | expression tag | UNP Q51504 |
| A     | 48      | HIS      | -      | expression tag | UNP Q51504 |
| A     | 49      | HIS      | -      | expression tag | UNP Q51504 |

• Molecule 2 is 1-[(2S)-2-{[(2Z)-2-(2-amino-1,3-thiazol-4-yl)-2-{[(2-carboxypropan-2-yl)oxy]imino}acetyl]amino}-3-oxopropyl]-4-{[2-(5,6-dihydroxy-1,3-dioxo-1,3-dihydro-2H-isoindol-2-yl)ethyl]carbamoyl}-2,5-dihydro-1H-pyrazole-3-carboxylic acid (three-letter code: R7G) (formula:  $C_{27}H_{28}N_8O_{12}S$ ) (labeled as "Ligand of Interest" by depositor).





| Mol | Chain | Residues | Atoms       |         |  | ZeroOcc | AltConf |   |   |
|-----|-------|----------|-------------|---------|--|---------|---------|---|---|
| 2   | A     | 1        | Total<br>48 | C<br>27 |  | O<br>12 | S<br>1  | 0 | 0 |

• Molecule 3 is water.

| Mol | Chain | Residues | Atoms            | ZeroOcc | AltConf |
|-----|-------|----------|------------------|---------|---------|
| 3   | A     | 85       | Total O<br>85 85 | 0       | 0       |

 $\operatorname{MolProbity}$  and EDS failed to run properly - this section is therefore empty.



# 3 Data and refinement statistics (i)

EDS failed to run properly - this section is therefore incomplete.

| Property                               | Value                          | Source    |
|--|--------------------------------|-----------|
| Space group                            | P 21 21 21                     | Depositor |
| Cell constants                         | 68.11Å 83.81Å 89.74Å           | Depositor |
| a, b, c, $\alpha$ , $\beta$ , $\gamma$ | 90.00° 90.00° 90.00°           | Depositor |
| Resolution (Å)                         | 39.59 - 2.40                   | Depositor |
| % Data completeness                    | 99.4 (39.59-2.40)              | Depositor |
| (in resolution range)                  | 33.4 (33.03-2.40)              | Depositor |
| $R_{merge}$                            | 0.14                           | Depositor |
| $R_{sym}$                              | (Not available)                | Depositor |
| $< I/\sigma(I) > 1$                    | 2.87  (at  2.39Å)              | Xtriage   |
| Refinement program                     | REFMAC 5.8.0253                | Depositor |
| $R, R_{free}$                          | 0.182 , $0.236$                | Depositor |
| Wilson B-factor $(\mathring{A}^2)$     | 31.6                           | Xtriage   |
| Anisotropy                             | 0.133                          | Xtriage   |
| L-test for twinning <sup>2</sup>       | $ < L > = 0.49, < L^2> = 0.32$ | Xtriage   |
| Estimated twinning fraction            | No twinning to report.         | Xtriage   |
| Total number of atoms                  | 3951                           | wwPDB-VP  |
| Average B, all atoms (Å <sup>2</sup> ) | 36.0                           | wwPDB-VP  |

Xtriage's analysis on translational NCS is as follows: The largest off-origin peak in the Patterson function is 5.55% of the height of the origin peak. No significant pseudotranslation is detected.

<sup>&</sup>lt;sup>2</sup>Theoretical values of <|L|>,  $<L^2>$  for acentric reflections are 0.5, 0.333 respectively for untwinned datasets, and 0.375, 0.2 for perfectly twinned datasets.



<sup>&</sup>lt;sup>1</sup>Intensities estimated from amplitudes.

# 4 Model quality (i)

## 4.1 Standard geometry (i)

MolProbity failed to run properly - this section is therefore empty.

## 4.2 Too-close contacts (i)

MolProbity failed to run properly - this section is therefore empty.

## 4.3 Torsion angles (i)

#### 4.3.1 Protein backbone (i)

MolProbity failed to run properly - this section is therefore empty.

#### 4.3.2 Protein sidechains (i)

MolProbity failed to run properly - this section is therefore empty.

#### 4.3.3 RNA (i)

MolProbity failed to run properly - this section is therefore empty.

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

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

## 4.5 Carbohydrates (i)

There are no monosaccharides in this entry.

# 4.6 Ligand geometry (i)

1 ligand is modelled in this entry.

In the following table, the Counts columns list the number of bonds (or angles) for which Mogul statistics could be retrieved, the number of bonds (or angles) that are observed in the model and the number of bonds (or angles) that are defined in the Chemical Component Dictionary. The Link column lists molecule types, if any, to which the group is linked. 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| > 2 is considered an outlier worth inspection. RMSZ is the root-mean-square of all Z scores of the bond lengths (or angles).

| Mol Type |      | Chain |     | Chain | Chain    | Chain | Chain    | Chain    | Chain | Chain    | Chain | Chain | Chain | Chain | Chain | Chain | Chain | Chain | Chain | Res | Link | B | ond leng | ${ m ths}$ | Bond angles |  |  |
|----------|------|-------|-----|-------|----------|-------|----------|----------|-------|----------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-----|------|---|----------|------------|-------------|--|--|
| MIOI     | туре | Chain | nes | LIIIK | Counts   | RMSZ  | # Z  > 2 | Counts   | RMSZ  | # Z  > 2 |       |       |       |       |       |       |       |       |       |     |      |   |          |            |             |  |  |
| 2        | R7G  | A     | 601 | 1     | 44,51,51 | 4.50  | 18 (40%) | 57,75,75 | 2.93  | 13 (22%) |       |       |       |       |       |       |       |       |       |     |      |   |          |            |             |  |  |

In the following table, the Chirals column lists the number of chiral outliers, the number of chiral centers analysed, the number of these observed in the model and the number defined in the Chemical Component Dictionary. Similar counts are reported in the Torsion and Rings columns. '-' means no outliers of that kind were identified.

| $\mathbf{Mol}$ | Type | Chain | Res | Link | Chirals | Torsions    | Rings   |
|----------------|------|-------|-----|------|---------|-------------|---------|
| 2              | R7G  | A     | 601 | 1    | -       | 18/37/72/72 | 0/4/4/4 |

The worst 5 of 18 bond length outliers are listed below:

| Mol | Chain | Res |     |         | Z      | Observed(A) | $\operatorname{Ideal}(	ext{\AA})$ |
|-----|-------|-----|-----|---------|--------|-------------|-----------------------------------|
| 2   | A     | 601 | R7G | C17-C3  | -12.49 | 1.32        | 1.49                              |
| 2   | A     | 601 | R7G | O5-N4   | 11.75  | 1.68        | 1.42                              |
| 2   | A     | 601 | R7G | C2-C3   | -11.44 | 1.37        | 1.50                              |
| 2   | A     | 601 | R7G | C22-C20 | -8.80  | 1.34        | 1.48                              |
| 2   | A     | 601 | R7G | C1-C5   | -8.58  | 1.34        | 1.48                              |

The worst 5 of 13 bond angle outliers are listed below:

| Mol | Chain | Res | Type | Atoms      | $\mathbf{Z}$ | $Observed(^o)$ | $Ideal(^{o})$ |
|-----|-------|-----|------|------------|--------------|----------------|---------------|
| 2   | A     | 601 | R7G  | C10-O5-N4  | 9.16         | 118.47         | 110.33        |
| 2   | A     | 601 | R7G  | C21-N8-C20 | -8.50        | 105.10         | 112.03        |
| 2   | A     | 601 | R7G  | C19-N8-C21 | 8.17         | 132.10         | 123.88        |
| 2   | A     | 601 | R7G  | C22-C20-N8 | 7.37         | 111.11         | 105.88        |
| 2   | A     | 601 | R7G  | C23-C21-N8 | 6.72         | 110.65         | 105.88        |

There are no chirality outliers.

5 of 18 torsion outliers are listed below:

| Mol | Chain | Res | Type | Atoms        |
|-----|-------|-----|------|--------------|
| 2   | A     | 601 | R7G  | O8-C17-C3-C1 |
| 2   | A     | 601 | R7G  | C3-C1-C5-O2  |
| 2   | A     | 601 | R7G  | C3-C1-C5-O1  |
| 2   | A     | 601 | R7G  | N1-C1-C5-O2  |
| 2   | A     | 601 | R7G  | N1-C1-C5-O1  |



There are no ring outliers.

No monomer is involved in short contacts.

The following is a two-dimensional graphical depiction of Mogul quality analysis of bond lengths, bond angles, torsion angles, and ring geometry for all instances of the Ligand of Interest. In addition, ligands with molecular weight > 250 and outliers as shown on the validation Tables will also be included. For torsion angles, if less then 5% of the Mogul distribution of torsion angles is within 10 degrees of the torsion angle in question, then that torsion angle is considered an outlier. Any bond that is central to one or more torsion angles identified as an outlier by Mogul will be highlighted in the graph. For rings, the root-mean-square deviation (RMSD) between the ring in question and similar rings identified by Mogul is calculated over all ring torsion angles. If the average RMSD is greater than 60 degrees and the minimal RMSD between the ring in question and any Mogul-identified rings is also greater than 60 degrees, then that ring is considered an outlier. The outliers are highlighted in purple. The color gray indicates Mogul did not find sufficient equivalents in the CSD to analyse the geometry.

# 4.7 Other polymers (i)

There are no such residues in this entry.

## 4.8 Polymer linkage issues (i)

There are no chain breaks in this entry.



# 5 Fit of model and data (i)

#### 5.1 Protein, DNA and RNA chains (i)

EDS failed to run properly - this section is therefore empty.

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

EDS failed to run properly - this section is therefore empty.

# 5.3 Carbohydrates (i)

EDS failed to run properly - this section is therefore empty.

## 5.4 Ligands (i)

EDS failed to run properly - this section is therefore empty.

## 5.5 Other polymers (i)

EDS failed to run properly - this section is therefore empty.

