Theoretical search of crystal polymorphs of temozolomide

David Stephen Arputharaj a,*, Meenashi Rajasekaran b, Christian Jelsch c, Saravanan Kandasamy d, Abdullah G. Al-Sehemi e

a Department of Physics, PSG College of Arts and Science, Coimbatore, 641014, Tamil Nadu, India
b Department of Physics, Kandaswami Kandar’s College, Velur, 638182, India
c Université de Lorraine, CNRS, CRM2, F-54000, Nancy, France
d Faculty of Chemistry, University of Warsaw, Lubelska Pasteura 1, Warsaw, 02093, Poland
e Research Center for Advanced Material Science (RCAMS), King Khalid University, Abha, 61413, Saudi Arabia

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ABSTRACT

Possible polymorphic forms of the chemotherapy drug, temozolomide were predicted from the \textit{ab initio} and DFT methods. The lattice minimization via distributed multipole analysis was carried out for the hypothetical generated structures. A crystal with unit cell parameters close to the real one and of same space group was retrieved, with partly similar packing and interactions. The analysis of inter molecular interaction (through Hirshfeld surface) and electrostatic potential reveals the complementary sites in the molecule. The 26 predicted structures were analyzed with respect to two computed lattice energies and hydrogen-bond propensity. The lattice energy of the real crystal [EXP] packing ranked number 6 compared on the basis of DMACRYS software and number 3 on the basis of the total lattice energy issued from the Crystalexplorer17 software at the B3LYP/6-31G** level of theory. The molecule has two strong hydrogen bond donors and five strong acceptors. The predicted packings are stabilized by one or two strong N–H…O/N–H…N as well as weak C–H…O/C–H…N and H…π hydrogen bonds. While the real structure with \(Z' = 1\), EXP, forms only one strong H-bond (N–H…O = C), several of the predicted packings form two strong H-bonds. Two predicted crystal packings have unit cell parameters close to the real structure, one of them shares several common intermolecular interactions.

1. Introduction

The phenomena of Polymorphism has acquired great academic and industrial interest [1] and the physical and chemical properties of polymorphic structures make a significant impact in both experimental and theoretical investigations. The polymorphic behavior in pharmaceutical compounds leads to different bio-availability. The theoretical search for unknown polymorphic structures has gained its importance as their experimental confirmation for their existence is the difficult procedure [2]. In this view, the current study aims to explore the different crystal phases of temozolomide drug which are thermodynamically stable, along with the experimentally known polymorphs. For the treatment of malignant glioma, the United States Food and Drug and Administration (US FDA) [3] approved the drug Temozolomide, an oral alkylating agent. The structure of temozolomide molecule [Figure 1] has tetrazine and imidazole rings, labelled as 1 and 2 with ketone and amide functional groups, which have the potential to form hydrogen bonds. The N–H…N and N–H…O hydrogen bonds are the lead interactions for the polymorphic formation. Ten polymorphs of TMZ were reported in the patent literature [4, 5]. Lowe et al. (2008) reported a first crystal structure of temozolomide (called ExpB) with \(Z' = 2\). Babu et al. [5] reported the same crystal form, one polymorph with \(Z' = 1\) (EXP) and one polymorph with \(Z' = 2\) (EXPb). And the protonated form of TMZ was also reported [6] in which the crystal structure was solved with one neutral temozolomide, one temozolomide-H^+–Cl^-, one H_3O^+–Cl^- and three water molecules. They described also three cocrystals of temozolomide. In addition, temozolomide polymorphs, characterized only by powder diffraction, were disclosed in two patents [7] and several of these polymorphs lack crystal

* Corresponding author.
E-mail address: stevepearlin@gmail.com (D.S. Arputharaj).
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structural information. To gain structural insights, various polymorphic forms of temozolomide were predicted theoretically via \textit{ab initio} calculations [8].

2. Materials and methods

2.1. Crystal structure prediction

The basic principle selected for the research to predict the stable nature of structures was to identify the structures with low lattice energy. The crystal phases of temozolomide at global energy minimum are determined from the first principle mentioned in Eq. (1).

\begin{equation}
E_{\text{total}} = E_{\text{lattice}} + \Delta E_{\text{intra}}
\end{equation}

The stable crystal structures of temozolomide were identified in the energy landscape plot. The stable hypothetical dense crystal phases of the molecule were generated using MOLPAK software [9]. The algorithm was designed to develop the possible packings by 3-dimensional rotations and repetitions of the parent molecule, incorporating the symmetry operators in all common coordination geometry. The global search was initiated through a gas phase optimization using the Becke 3-Parameter exchange method (B3LYP) [10]. An initial level optimization to minimize the packing density of the hypothetical structures with threshold interactions were selected for PMIN (Packaging minimization) optimizations [9] incorporating a repulsion alone potential field called UMD potential [9]. The current research specifically aimed to search the putative crystal forms of temozolomide within the commonly encountered space groups P1, P-1, P2, Pm, Pn, P21, P2/m, P21/c, Cc, C2, C2/c, Pn, Pna2, Pbc2, Pnma, Pna21, Pnma and Pbca; where 95% [11] of the crystal structures in the database were listed. The PMIN optimization carried out in the global search within the repulsion alone potential field neglected the electron correlation parameters. For the further lattice energy minimization using DMACRYS code [12], the densest hypothetical crystal structures generated from the MOLPAK search were considered, where the structures were subjected to rigid body optimization within a repulsion-dispersion potential field of the form of Eq. (2).

\begin{equation}
U_{\text{lattice}} = \sum_{i,j,k} \left( A_{ij} A_{kk} \right) \exp \left[ -\frac{1}{2} \left( B_{ij} + B_{kk} \right) R_{ik} \right] - \left( C_{ij} C_{kk} \right) / R_{ik}^6
\end{equation}

Figure 1. Chemical structure of Temozolomide.

Figure 2. Crystal energy landscape showing all possible stable crystal structures of temozolomide along with experimental structure [EXP] in black, Close to real [AM] in blue.
Table 1. List of predicted possible stable crystal structures of temozolomide molecule. The experimental and the predicted, close-to-real polymorphs are in bold.

| ID     | Crystal  | Space group | a (Å)  | b (Å)  | c (Å)  | alpha (°) | beta (°) | gamma (°) |
|--------|----------|-------------|--------|--------|--------|------------|----------|-----------|
| AM     | P21/c    | 6.721       | 7.552  | 16.101 | 90     | 96.1       | 90       |           |
| CC     | Pcca     | 7.498       | 7.752  | 27.439 | 90     | 90         | 90       |           |
| CB     | Pbc a    | 16.647      | 7.574  | 12.646 | 90     | 90         | 90       |           |
| CA     | P-1      | 6.233       | 6.119  | 12.506 | 119.4  | 97         | 75.9     |           |
| AB     | P-1      | 4.416       | 9.139  | 9.9    | 99.3   | 91.3       | 94.9     |           |
| EXP    | P21/n    | 6.672       | 7.875  | 16.02  | 90     | 95.3       | 90       |           |
| DC     | C2/c     | 19.079      | 4.563  | 18.783 | 90     | 102.7      | 90       |           |
| AH     | P21      | 8.679       | 6.265  | 7.661  | 90     | 106.9      | 90       |           |
| DB     | C2       | 7.715       | 6.552  | 16.963 | 90     | 105.9      | 90       |           |
| FA     | P21/c    | 5.854       | 25.756 | 7.855  | 90     | 78.4       | 90       |           |
| AQ     | P21/2121 | 14.121      | 6.554  | 8.787  | 90     | 90         | 90       |           |
| CE Z-2 | Pbcm     | 36.66       | 7.481  | 6.241  | 90     | 90         | 90       |           |
| FC     | P21/c    | 9.796       | 4.625  | 17.983 | 90     | 90         | 90       |           |
| Ai     | P21/c    | 10.636      | 4.13   | 19.359 | 90     | 68.7       | 90       |           |
| AL     | P21/c    | 7.92        | 7.011  | 15.387 | 90     | 90         | 90       |           |
| AA     | P1       | 12121       | 4.603  | 38.241 | 4.772  | 90         | 90       |           |
| AS     | Pna21    | 136.122     | 4.845  | 4.572  | 90     | 90         | 90       |           |
| BF Z-2 | Pmc      | 4.853       | 36.082 | 4.569  | 90     | 90         | 90       |           |
| AP     | P21/2121 | 4.603       | 38.241 | 4.772  | 90     | 90         | 90       |           |
| DA     | Cc       | 7.316       | 18.998 | 9.613  | 90     | 139.6      | 90       |           |
| BD     | Pna21    | 25.784      | 6.14   | 5.109  | 90     | 90         | 90       |           |
| AV     | Pna21    | 12.988      | 16.167 | 4.074  | 90     | 90         | 90       |           |
| BH     | Pca21    | 18.985      | 5.057  | 8.744  | 90     | 90         | 90       |           |
| EXPb Z-2 | P 21/c  | 17.332      | 7.351  | 13.247 | 90     | 109.56     | 90       |           |
| EXPc Z-2 | P -1   | 8.5        | 10.004 | 11.309 | 90     | 108.86     | 109.19   |           |

Table 2. Hydrogen bonding details in predicted close to real [AM] and experimental crystal structure [EXP]. Hydrogen bonds common in the two structures are in bold.

| Symmetry operation | H-Bond | H...A (Å) | D...A (Å) | D-H...A (°) |
|--------------------|--------|-----------|-----------|------------|
| Intra              | N6-H6..N3 | 2.340/2.257 | 2.756/2.730 | 103.5/107.1 |
| x/2+y/2,z/2-x,1/2,y,1/2-z | N6-H5..N2 | 1.845/1.834 | 2.836/2.824 | 166.4/166.0 |
| x+y/2,z/2-x,1/2,y,1/2-z | C4-H1..O1 | 2.442/2.370 | 3.432/3.390 | 151.3/156.2 |
| x+y/2, y+z/2, z/2-x,1/2,y,1/2-z | HS-H2..N3 | 2.559/2.700 | 3.332/3.357 | 127.6/118.7 |
| x+y/2, y+z/2, z/2-x,1/2,y,1/2-z | HS-H3..O1 | 2.935/2.935 | 3.409/3.277 | 106.7/98.6 |
| x+y/2, y+z/2, z/2-x,1/2,y,1/2-z | HS-H2..O1 | 2.941/2.713 | 3.409/3.277 | 106.7/112.0 |
| x+y/2, y+z/2, z/2-x,1/2,y,1/2-z | HS-H4..O2 | 2.564/2.562 | 3.528/3.571 | 147.9/154.6 |
| x+y/2, y+z/2, z/2-x,1/2,y,1/2-z | HS-H4..N4 | 2.656/2.704 | 3.464/3.498 | 130.9/120.8 |
| /x+1/2,y+z | HS-H3..O2 | -2.828 | -3.532 | /121.9 |
| /x+1/2,y+z | HS-H3..N4 | -2.760 | -3.571 | /154.6 |

where i and k represents the atomic species of 1 and 2 molecules. The FIT potential, calibrated by Williams and Cox [13] was implemented in the lattice energy minimization. The FIT potential was also included with potential, calibrated by Williams and Cox [13] was implemented in the lattice energy plot, where comparatively dense region found to be in between -112 kJ/mol to -142 kJ/mol. The most favorable crystal packings at 0K along with the experimentally known crystal structure by exhibiting the monoclinic packing of P21/c space group with cell density of 1.587 gm/cm3 at D_{min} = -141.8- kJ/mol. The unique hypothetical dense crystal packings of temozolomide molecule were ranked to construct the lattice energy landscape.

The lattice energy and density scatterplot (Figure 2) and the energy rank (Table 1) revealed that the predicted crystal structure AM resembles the experimental crystal structure by exhibiting the monoclinic packing of P21/c space group with cell density of 1.587 gm/cm3 at D_{min} = -141.8- kJ/mol. The most favorable crystal packings at 0K along with the experimentally known crystal structure of temozolomide were listed in the lattice energy plot, where comparatively dense region found to be in between -112 kJ/mol to -142 kJ/mol.
2.2. Total energy calculation

The total lattice energy $E_{\text{tot}}$ of the crystals was approximated with software Crystal Explorer19 [16] on a cluster of molecules surrounding the central one. The N–H and C–H bond distances were constrained to standard values obtained from neutron diffraction studies [17]. The $E_{\text{tot}}$ values, were calculated from the sum of scaled electrostatic, polarization, dispersion and repulsion components using the coefficients 1.057, 0.74 and 0.871 respectively using B3LYP/6-31G** level of theory. For the structures with two molecules per asymmetric unit ($Z' = 2$), the energy of the two molecules was averaged. Fingerprint plots of the intermolecular contacts were also generated with program Crystal Explorer19.

3. Results and discussion

3.1. Structure and intermolecular interactions

The temozolomide molecule comprises tetrazine, imidazole and acetamide groups which extends the possibilities of polymorphism. The predicted crystal AM has unit cell parameters close to the real crystal structure and the same space group. The bond lengths, bond angles and dihedral angles of the predicted structure [AM] agree well with that of X-ray geometry [EXP] [5]. There is no significant bond twist for all the bonds of the EXP structure. All the torsion angles in the predicted structures except AM which is close to real one were well agree with the experimentally [EXP] observed structures. The torsion angles of AM...
structure is varied with that of EXP structure with the minimum and maximum difference of 0.01° and 7.46°. The maximum torsion angle difference is noted for carbonyl (C=O) and methyl (-CH₃) attached C3–N2 bond of inner side of tetrazine ring. Figure S1 in supplementary material visualize the torsion angle difference distribution of AM with that of EXP structure. The tetrazine and imidazole rings are almost (with the maximum deviation of 0.02°) in same plane for EXP and this planarity was feebly distracted by 4.7° in AM predicted structure. This planarity was well established from the N6-H6…N3 intra-molecular interaction [Table 2] in both EXP and AM crystal structure. In the solid state confirmation, both the oxygen atoms [O1 and O2] and nitrogen atoms [N4 of tetrazine, N6 of acetamide and N3 of imidazole ring] participated in the intermolecular interactions. The two crystal forms share, in part, similar morphology [Figure 3a, b] and are both stabilized by the dimer interactions framed from N6–H5…O2 strong hydrogen bond [Table 2].

Some hydrogen bonds such as C–H…O, C–H…N and longer distance N–H…N and N–H…O are contributing to the AM and real crystal packings. The intermolecular interactions of the molecule with the neighbouring molecules in the crystal are shown in Figure 4. The Hirshfeld surface analysis gives the details of all the contribution of intermolecular contacts and especially major role played by H...O and H...N contacts. The fingerprint analysis (Figure 5) [18] shows that the N...H/H...N and O...H/H...O contacts give the largest contribution, which are about 28%[EXP]/28.1%[AM] and 25.5%[EXP]/25.9%[AM] respectively. Moreover the H...H contact also renders a significant contribution of 19.7%[EXP]/18.7%[AM] to the crystal packing of the molecule.

The fingerprint plot difference between the experimental and predicted molecule was displayed in Figure 5c. In general, the difference were represented by blue and red regions; Blue regions are more intense in the experimental structure and red region highlights for the predicted one. As both the crystal forms prevailed similar interactions with same intense, there is no red region in the difference plot. There is a blue region in the spike at de = 1.1 Å, di = 0.7 Å and de = 0.7 Å, di = 1.1 Å clearly illustrates that there is a greater prevalence of O...H/H...O interactions in experimental structure when compared with that of the predicted one.

From Figure 6, it appears that some of the predicted crystal structures have a two strong hydrogen bonds while the experimental
and AM crystal structures have only one. Up to ten predicted crystal structures have two strong hydrogens N–H…O/N involving the two H atoms of the NH₂ group. In the real crystal, one NH₂ hydrogen atom form one strong N–H…O hydrogen bond and the other one forms two longer distance weak H bonds. The crystal structure FA is characterized by the largest number of hydrogen bonds (2 strong and 7 weak). The BH packing with no strong H-bonds has the smallest energy magnitudes. The number of strong hydrogen bonds shows a correlation coefficient of 0.64 with both -U_lattice and -E_elec values. The crystal packings with two strong H-bonds appear to have generally the

| Table 3. Contacts types and their enrichment in the predicted [AM] and experimental [EXP] crystal structure of temozolomide. |
|---|---|---|---|---|---|---|---|---|---|---|
| structure | EXP | AM |
| atom | Hp | C | N | O | Hc | Hp | C | N | O | Hc |
| surface % | 13.5 | 23.2 | 22.9 | 15.9 | 24.6 | 12.9 | 22.8 | 23.9 | 15.9 | 24.6 |
| Nn | 0.3 | 0.3 | % contacts | 6.5 | 5.7 | % contacts |
| N | 8.1 | 11.1 | 4.3 | 8.6 | 10.9 | 4.7 |
| O | 7.4 | 8.3 | 2.1 | 0.0 | 7.5 | 7.9 | 2.2 | 0.1 |
| Hc | 4.9 | 7.3 | 14.9 | 15.0 | 4.0 | 3.4 | 7.4 | 15.5 | 14.9 | 4.5 |
| Hn | 0.16 | 0.16 | % contacts | 1.11 | 1.17 | % contacts |
| C | 1.04 | 1.17 | Enrichment | 1.11 | 1.17 | Enrichment |
| N | 1.32 | 1.11 | 0.86 | 1.39 | 1.06 | 0.87 |
| O | 1.64 | 1.13 | 0.28 | 0 | 1.73 | 1.1 | 0.28 | 0.05 |
| Hc | 0.72 | 0.65 | 1.33 | 1.83 | 0.64 | 0.51 | 0.67 | 1.33 | 1.81 | 0.72 |

Figure 6. Total number of hydrogen bonds in the different crystal packings. For the Z' = 2 crystals, the number of H-bonds has been divided by two.
largest $E_{\text{elec}}$ lattice electrostatic energy values in magnitude (Figure S4 in supplementary material).

### 3.2. Contact enrichment ratio

The MoPro Viewer [19] package was used to calculate the enrichment ratio which compares the actual contact with equi-probable theoretical contacts. The interactions; N...C, N...Hc and O...Hc were considered as major interactions as they represent ~41% of the global Hirshfeld surface and their corresponding enrichment values are listed in Table 3. The strong interaction H5...O2 was represented with $E$ values 1.64 and 1.73 for EXP and AM respectively. The most enriched contact was noted for non-polar hydrogen and oxygen atom and their $E$ values were 1.83 [EXP] and 1.81 [AM]. The calculated values from the contact enrichment ratio reflects similar trend in the interaction energies. It is noted that there is >99% correlation similarity between the intermolecular interaction for predicted crystal structure [AM] and experimental structure [EXP] of temozolomide, which reveals the structures, were equivalent.

### 3.3. Closed and open shell interactions

The electron density and Laplacian of the electron density mappings of the molecules show the covalent bonding regions and lone pairs of the oxygen and nitrogen atoms that are picturized in Figure 7 (a,b). The topological properties of the electron density of the molecule in gas phase lifted from AM is calculated and the same was compared with that from EXP crystal structure in accordance with quantum theory of atoms in molecules (QTAIM) [20]. It is observed that electron density $\rho_{\text{cp}}$ and Laplacian $\nabla^2\rho_{\text{cp}}$ values show respectively a 93% and 95% correlation coefficient and the values are listed in Table S1 in supplementary material. Among the homogeneous nitrogen-nitrogen bonds, the bond N5–N4 has the highest electron density value $\rho_{\text{cp}} ~ 3 \text{ e/Å}^3$ at the critical point and it is stronger than the N2–N5 bond which has the value of $\sim 2.2e/\text{Å}^3$. The C5–N2 bond has low $\rho_{\text{cp}}$ electron density (AM $= 1.73e/\text{Å}^3$ and real $= 1.66e/\text{Å}^3$). The charges in their bonding region were highly depleted which was shown from less negative Laplacian values (AM $= -16.2e/\text{Å}^5$ and real $= -12.7e/\text{Å}^5$) in Table S1 (in supplementary material) and found to be weakest among all the C–N bonds with its low ellipticity value (0.04) suggesting the cylindrical nature of the bond.

### 3.4. Energy of intermolecular interactions

The stability of the molecules in the experimental and predicted crystal structure was established from C–H...O, N–H...O and C–H...N interactions.

#### Table 4. PIXEL lattice energies (kJ/mol) for predicted, AM and real crystal structure, EXP.

| Term    | AM    | EXP   |
|---------|-------|-------|
| $E_{\text{coul}}$ | -102.3 | -97.0 |
| $E_{\text{pol}}$  | -30.7  | -29.3 |
| $E_{\text{Disp}}$ | -111.2 | -117.2 |
| $E_{\text{Rep}}$  | 108.3  | 102.4 |
| $E_{\text{total}}$| -135.9 | -141.1 |

#### Table 5. Interaction energies (kJ/mol) of the molecular pairs, computed with program PIXEL for the real (first lines) and AM unit cell close to real (second lines) polymorphs.

| H-Bond     | Centroid distance | $E_{\text{coul}}$ | $E_{\text{pol}}$ | $E_{\text{Disp}}$ | $E_{\text{Rep}}$ | $E_{\text{total}}$ |
|------------|------------------|-------------------|-----------------|------------------|-----------------|------------------|
| N6–H5...O2 | 9.7              | -72.4             | -24.8           | -19.6            | 57.7            | -59              |
| N6–H5...O2 | 9.6              | -87.5             | -30.2           | -20.3            | 85              | -53              |
| C4–H1...O1 | 7.4              | -20.3             | -5.6            | -13.2            | 16.4            | -22.7            |
| C4–H1...O1 | 7.5              | -15.5             | -4.6            | -11.8            | 12.1            | -19.7            |
| C5–H2...N3 | 8.2              | -4.6              | -2.7            | -12.7            | 8.4             | -11.5            |
| C5–H2...N3 | 8.4              | -5.2              | -2.2            | -11.5            | 7.4             | -11.6            |
| C5–H2...N1 | 9.6              | -3.7              | -1.3            | -6.0             | 2.8             | -8.3             |
| C5–H2...N1 | 9.5              | -4.0              | -1.3            | -6.2             | 3.6             | -8.0             |
| C5–H4...O2 | 5.9              | -20.2             | -5.6            | -19.3            | 10.8            | -34.3            |
| C5–H4...N4 | 6.0              | -19.1             | -5.5            | -18.8            | 11.4            | -32.1            |
| C5–H3...N4 | 6.7              | -20.9             | -6.3            | -21.5            | 16.9            | -31.9            |
Figure 8. Energy frameworks along (a) a-axis, (b) b-axis, (c) c-axis for predicted [AM] and experimental [EXP] structure, showing separate electrostatic (left, red), and dispersion (middle, green) components, and total energy interactions (right, blue). The tube size (scale factor) used in all energy frameworks was 50 and the cut-off was 3.00 kJ/mol.
It appears that most other interactions are predominantly stabilized by dispersion energies in predicted crystal structure [AM] and experimental structure (Table 4). Also, the net energies were also calculated from Crystal Explorer 17 using B3LYP/6-31G(d,p) and found the same trend with net electrostatic energies; -98.5 KJ/mol [AM]/-90.1 KJ/mol[EXP] and net dispersion energies; -106.8 KJ/mol[AM]/-101.7 KJ/mol[EXP]. From the PIXEL method [21], the intermolecular energy of predicted [AM] and real crystal [EXP] were calculated and are listed in Table 5. Among the interactions, the N6–H5...O2 hydrogen bond is found to be the strongest in the real and predicted AM crystal structures and the corresponding total energy values are -59 kJ/mol and -53 kJ/mol respectively. The similar trend was observed for antipyrine-like derivatives [22]. Figure 8 clearly indicates that the electrostatic, dispersion and total energy frameworks for AM and EXP packed along the a-axis, b-axis and c-axis respectively. This was well reflected from the large and red regions labelled 1 in dnorm maps [Figure 9]. Next to this, the C5–H4...O2 and C5–H4...N4 hydrogen bonds exhibit total energy values of around -33 kJ/mol, and this combined

Table 6. The $U_{lattice}$ and scaled $E_{tot}$ energy values from DMACRYS and Crystal Explorer software.

| Crystal ID | $U_{lattice}$ KJ/mol | $E_{tot}$KJ/mol |
|------------|----------------------|-----------------|
| AM         | -141.8               | -207.9          |
| CC         | -140.9               | -192.2          |
| CB         | -140.7               | -168.7          |
| CA         | -134.9               | -190.3          |
| AB         | -132.3               | -189.4          |
| EXP        | -132.2               | -189.4          |
| DC         | -130.9               | -159.7          |
| AH         | -130.9               | -135.1          |
| DB         | -130.2               | -179.2          |
| EXPc Z' = 2| -129.6               | -142.9          |
| FA         | -128.0               | -179.5          |
| EXPc Z' = 2| -128.0               | -145.5          |
| AQ         | -128.0               | -127.4          |
| EC         | -127.4               | -161.8          |
| CE [Z = 2] | -127.4               | -122.9          |
| AI         | -127.3               | -133.2          |
| AL         | -126.7               | -137.7          |
| AY         | -126.5               | -115.2          |
| AA         | -125.9               | -116.9          |
| CD[Z = 2]  | -125.6               | -150.8          |
| AZ         | -123.7               | -114.0          |
| AP         | -123.6               | -113.5          |
| AS         | -123.1               | -111.8          |
| BF[Z = 2]  | -121.0               | -91.6           |

| Crystal ID | $U_{lattice}$ KJ/mol | $E_{tot}$KJ/mol |
|------------|----------------------|-----------------|
| AP         | -121.1               | -109.0          |
| DA         | -119.1               | -116.7          |
| BD         | -118.4               | -109.5          |
| AV         | -117.4               | -106.1          |
| BH         | -112.6               | -103.8          |

Figure 9. Hirshfeld surfaces mapped with dnorm for both (a) EXP and (b) AM crystal structures.
participation was evident from the white regions apart from each other labelled 2 in dnorm maps. The small red regions labelled 3 represents the O1…H1 interactions generating R2(10) motif predicted with total interaction energies of -22.7 kJ/mol and -19.7 kJ/mol for real [EXP] and predicted [AM] crystal structures respectively.

The total energy, $E_{\text{tot}}$ quantified as scaled sum of electrostatic, dispersion, polarization and exchange-repulsion terms from CrystalExplorer program is listed in Table 6 along with $U_{\text{lattice}}$. The $E_{\text{tot}}$ energy for the real crystal is ordered in third position (-189.4 kJ/mol). The structure AM, close to real, comes, in first position for both $E_{\text{tot}} = -207.9$ kJ/mol and $U_{\text{lattice}} = -141.8$ kJ/mol.

3.5. Lattice energy of crystal packings

Concerning the $U_{\text{lattice}}$ energy descriptor, the structure AM, close to real, has the largest value (-141.8 kJ/mol) and the real structure is in fourth position.

The CE crystal packing, having two molecules in asymmetric unit, holds the strongest $E_{\text{tot}}$ lattice value from CrystalExplorer17 at -224.8 kJ/mol. Compared to the real crystal, there are a few additional H-bond interactions prevailing in CE, but the corresponding $U_{\text{lattice}}$ energy value -127.4 kJ/mol is around ~10 kJ/mol smaller in magnitude than the real structure.
There is a group of predicted structures which have both weak $U_\text{lattice}$ energy (less than -130.9 kJ/mol in magnitude) and weak $E_{\text{tot}}$ values (less than -135.1 kJ/mol). They present no strong H-bond (BV, AV) or one (BD, DA, AP, AS, AF, AZ, AY, AA, AQ, AH), like the real structure. On the whole, the $E_{\text{tot}}$ values of the structures correlated with the $U_\text{lattice}$ energy values by 47%. The computed $E_{\text{tot}}$ energies show variations which are double compared to $U_\text{lattice}$ but their average magnitudes are similar ($-142$ vs. -$125$ kJ/mol). Globally the $U_\text{lattice}$ energies show a good correlation of 0.58 with the lattice $E_{\text{elec}}$ values (Figure S3 in supplementary material) obtained from a transferred multipolar atom model, using UBDB database [23]. The 9 most stable polymorphs according to the Crystalexplorer scaled sum $E_{\text{tot}}$ energy are shown in Figure 10. The crystal CE has the strongest $E_{\text{tot}}$ value ($-238$ kJ/mol) and the real crystal appears next within a group formed by (CA, AB, real) crystals for which $E_{\text{tot}} = 189$ kJ/mol. Despite the crystal CE has a strong repulsive energy, this component is strongly attenuated (coefficient 0.618) when the $E_{\text{tot}}$ descriptor is considered. Only the EXP polymorph corresponds to one of the crystal structures of TMZ reported in the patent literature [4, 7], which may include some cocrystals. The list of predicted crystal structures in SHELX format were listed in Table S2 in supplementary material.

3.6. Molecular and packing electrostatic potential

The three-dimensional molecular electrostatic potential (MESP) plot for both experimental and predicted AM structure of temozolomide is shown in Figure 11. The polar nature of the molecule was well established from the positive and negative extrema of the potential on the molecular surface. As expected, in both experimental and predicted AM structure, the strongest negative potential is found for the oxygen atoms O1/O2 and their corresponding potential values were -22.3/-54.4 kcal/mol [EXP] and -22.5/-53.1 kcal/mol [AM] respectively. The maximum positive potential was found near the hydrogen atoms of amine group and their average values were +32.7 kcal/mol [EXP] and +35.0 kcal/mol [AM] respectively. The hydrogen bond donor and acceptor capability was well characterized from potential of oxygen atoms and –NH2 groups.

Figure 12 represents a scatterplot of the inner and outer electrostatic potential $V$ on the Hirshfeld surface for the experimental crystal structure. The contacts related to weak C–H...O and C–H...N hydrogen bonds are in color while those related to a strong hydrogen bond N–H...O or N–H...N are in black.
Figure 13. (a) Average percentage $C_{xy}$ of actual contacts on the Hirshfeld surface in the predicted crystal packings (in blue). The values for the real packing are shown in black. The errors bars represent the standard deviation in the sample of predicted structures, (b) Percentage of actual contacts in the real and predicted crystal packings, grouped by types.
3.7. Contact types in the crystal packings

The difference in contact on the Hirsheld surface between the real crystal and predicted crystals with respect to the proportion of contact types is picturized in Figure 13. In terms of contact surface, the real crystal is more involved in weak hydrogen bonds of N...Hc, O...Hc and C...Hn types, compared to the average of predicted structures. Strong H-bonds O...Hn and N...Hn are also, on average, more represented in the real crystal. Only the AQ crystal structure has significantly more hydrogen bond surface than the real crystal structure, while the AM close to real structure has slightly less H-bonding. On the other hand, the amount of O...C contacts and hydrophobic interaction Hc...Hc and C...Hc is diminished in the real crystal compared to the average of predictions. Percentage of actual contacts in the experimental and predicted crystal packing are shown in Figure S2 in supplementary material.

3.8. Hydrogen propensity analysis

The H-bond propensity [HBP] was calculated for the real and predicted crystal structures to get the possible inter and intramolecular interactions. The propensity calculations were performed from the statistical model established for donor and acceptor pairs, and the training dataset utilized for this model was extracted from Cambridge Structural Database (CSD). The HBP values of possible donor-acceptor pairs of real [EXP] and predicted [AM] structures are listed in Table 7. In general the HBP can take values between 0. and 1.; 0 represents the no likelihood of H-bond formation and 1 indicates the maximal likelihood of H-bond interaction.

From Table 6, it appears that the atom N6 of acetamide groups was calculated as active donor. On the other hand, the nitrogen atoms [N4 and N5 of tetrazine] and the oxygen atoms [O1 and O2] were calculated as active acceptor atoms in the theoretical calculated H-bond interactions of temozolomide. Among them, the H-bond interactions having highest propensity values [mean value: 0.85] were observed in real [EXP] and predicted is close to real structure [AM], whose mean H-bond propensity value is 0.81.

Figure 14 represents the putative structure landscape for real and predicted temozolomide crystal structures. The crystal structure having the maximum number of H-bond interactions thus having the maximum HBP value were represented by the points situated at the right bottom corner of the landscape plot and the points in the top left represents the unstable polymorphs (Figure S3 in supplementary material).

4. Conclusion

Ab initio prediction of the possible stable crystal polymorphs of temozolomide molecule, an oral alkylating pharmaceutical drug, was carried out. By analyzing the lattice energy landscape, unresolved polymorphs with theoretical greater stability were found to exist in the lowest energy regions. The lattice energy landscape was generated via global search through the repulsion alone potential field and incorporated lattice minimization using distributed multipole analysis associated with the hypothetical structures. The polymorph AM shows unit cell

| Donor | Acceptor | EXP | AM |
|-------|----------|-----|----|
| N6    | N3       | 0.89| 0.89|
| N6    | O2       | 0.81*| 0.81*|
| N6    | N5       | 0.80| 0.80|
| N6    | N4       | 0.78| 0.78|
| N6    | O1       | 0.69| 0.69|

Figure 14. The putative structure landscape for temozolomide real [EXP] and predicted polymorphs.
parameters close to the experimental crystal form of temozolomide (EXP) and shares some common contacts. The stability of the predicted structure AM was studied by PIXEL energy approach and by evaluating the hydrogen bonds related with the structure. Further studies on the intermolecular interactions were carried out through the Hirshfeld surface analysis. The thermodynamically stable polymorph is not in first position but ranks among the structures with strongest $U_{\text{lattice}}$ and $E_{\text{tot}}$ lattice energy descriptors.

Declarations

Author contribution statement

David Stephen Arputharaj: Concepted and designed the experiments; Performed the experiments; Wrote the paper.

Meenashi Rajasekaran, Christian Jelsch: Analyzed and interpreted the data.

Saravanan Kandasamy, Abdullah G Al-Sehemi: Contributed reagents, materials, analysis tools or data.

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Data will be made available on request.

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The authors declare no conflict of interest.

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