IN-SILICO ALANINE SCANNING ANALYSIS ON THE CATALYTIC RESIDUES OF A NOVEL β-GLUCOSIDASE FROM TRICHODERMA ASPERELLUM UC1

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Graphical abstract

Abstract

Currently, the catalytic residue of the highly prolific fungal β-glucosidase (BGL) of Trichoderma asperellum UC1 remains unvalidated. The study used the alanine scanning method to confirm the catalytic residues of the BGL as Glu165, Asp226, and Glu423. This method cancels out all intermolecular hydrogen bonds with substrates, lignin, hemicellulose, and cellulose. Results revealed an overall decline in the stability of the energy-minimized mutant enzymes’ compared to the wild-type BGL. The mutant enzyme registered lower PROCHECK (91.0%), ERRAT (93.09%), and Verify-3D (98.92%) values, in comparison to 90.2%, 92.09%, 98.06%, in the wild-type BGL, respectively. The mutant BGL UC1-substrate complexes were less stable than the wild-type enzyme, in which the mutant exhibited higher binding energies for docked lignin (−7.4 kcal mol⁻¹), cellulose (−7.2 kcal mol⁻¹), and hemicellulose (−7.2 kcal mol⁻¹). Binding energies of the wild-type BGL with the corresponding substrates were lower at −7.9 kcal mol⁻¹, −8.1 kcal mol⁻¹, and −7.8 kcal mol⁻¹. An interesting observation was that the alanine scanning changed the substrate preference order based on the calculated binding energies. The mutant BGL bound preferentially to lignin>cellulose=hemicellulose, while the wild-type BGL was selective to cellulose>lignin>hemicellulose. Hence, the findings convey the high likelihood of Glu165, Asp256, and Glu423 are the catalytic residues of the BGL of T. asperellum UC1.

Keywords: β-glucosidase from UC1, alanine scanning, catalytic triad, β-glucosidase, molecular docking, molecular dynamics
1.0 INTRODUCTION

Carbohydrate degrading enzymes secreted by the Trichoderma genus’ fungi are favored due to their ability to secrete large quantities of the enzymes extracellularly [1]. Recently, Ezeilo et al. [2] isolated a novel T. asperellum UC1 fungus (GenBank accession number MF774876) that secretes an array of synergistic carbohydrate hydrolyzing enzymes. The crude enzyme cocktail consisted of four endo-β-1,4-xylanases and endoglucanases, and one exo-glucanase and a β-glucosidase [1]. The enzymes are key bioremediation agents that recycle carbon from lignocellulosic biomass into the environment. Lignocellulose is the most abundant biopolymer on Earth, with cellulose, hemicellulose, and lignin being the major components [2–4]. However, this study focuses on the extracellular β-glucosidase (BGL) of T. asperellum UC1 since it is among the commonplace enzymes produced by organisms [1].

BGL hydrolyzes β-glycosidic bonds of terminal non-reducing residue in β-D-gluco- and oligosaccharides. This action liberates glucose or other sugars as products, which explains its valuable role in cellulose saccharification and degradation [5]. The BGL of T. asperellum UC1 has a higher enzymatic activity reportedly over other crude enzyme cocktails, in conjunction to better digests cellulose components than β-glucosidases of T. reesei [6, 7]. For this study to comprehend the catalytic workings of the BGL UC1, a resolved crystal structure of a comparable extracellular β-glucosidase called the ThBgl [8] was used as the enzyme template. The ThBgl falls under the β-retaining glycosyl GH1 family (E.C. 3.2.1.21), in which protein architecture is made up of a single (α/β) TIM barrel fold with two glutamate residues (Glu) as the catalytic residues that are placed at a certain distance to retain mechanism [8]. β-retaining glycosyl BGLs hydrolyze form glycosidic bonds in plants via a mechanism called deglycosylation. The mechanism is instigated by an attack on a water molecule by the general acid or base to discharge the glutamic acid (nucleophile) while the configuration of anomeric carbon is left unchanged [9]. Florindo et al. [10] deduced the catalytic residues of the BGL of ThBgl to be made up of nucleophilic Glu367 (E367) and Glu166 (E166). It is thought that the catalytic glutamate of BGL of T. asperellum UC1 is at positions, Glu165, and Glu423, alongside Asp226 [11]. However, this aspect is yet to be confirmed. Our earlier study only performed phylogenetic tree and sequence alignment studies to identify the species and characterized the biochemical- and catalytic behaviors of the BGL of T. asperellum UC1. We did not delve into the roles of the catalytic triad of the fungal enzyme. Although a crystal structure of another BGL is available, the catalytic residues of the BGL UC1 remain an educated guess. Hence, there is still a lacking of affirmative investigation to identify the BGL UC1 catalytic residues.

In this milieu, the method of alanine scanning comes in handy to validate active site residues. The technique has been regularly used to identify catalytic residues in various enzymes without disrupting their cores’ folding [12]. In this study, the technique systematically substitutes putative catalytic residues with alanine, thereby nullifying any possible interactions between the side-chains of amino and the ligand(s) [13]. It was expected that
the interaction of the mutant BGL-ligand complex (ΔGW) becomes destabilized and gives a higher Gibbs free energy when compared to the wild-type (ΔGW). The more negative the value, the stronger the interaction between the side chain of the enzyme and ligand [14]. The method was useful to demonstrate the binding of human growth hormone (hGH) to hGH-binding protein (hGHbp) [15], ligand-binding pocket for the human Vitamin D receptor [16], and complete alanine scanning of the Esherichia coli RbsB ribose binding protein involved in chemoreceptor signaling [17] amongst many others.

Hence, the molecular modeling approach using in silico alanine scanning performed by using the GROMACS software can further expedite the clarification of the catalytic residues in BGL UC1. It can overcome the time-consuming, expensive, and laborious shortcomings in the conventional empirical validation studies to identify the catalytic residues. GROMACS allows the rapid comprehension of the structure-activity relationships between the substrate and an enzyme's binding site. The software can rapidly gauge changes in the mutant enzyme [18][19]. This technique highlights the salient contribution of in silico alanine scanning to expediently identify and affirm the catalytic residues of the BGL of T. asperellum UC1 while complementing other empirical validation methods. It is hypothesized that the replacement of the putative catalytic residues in the BGL UC1 invalidates the essential intermolecular hydrogen bonds that anchor the substrates into the active site. This, in turn, renders the general increase in binding energies of the β-glucosidase-substrate complex. Pertinently, this is the first study detailing the in silico attempt to identify the catalytic residues of the BGL BGL of T. asperellum UC1.

Pertinently, the approach proposed here is feasible due to the availability of a similar three-dimensional (3D) molecular model of the BGL of T. asperellum in the National Center for Biotechnology Information (NCBI) database. In this work, the 3D structure of the β-glucosidase UC1 must first be constructed, followed by in silico substitution of Glu165, Asp226, and Glu423 into alanine, using the GROMACS package. Using AutoDock version 4.2.6, the enzyme's natural substrate was docked into the mutated catalytic site. Next, the study used molecular dynamics to estimate changes in binding energies before and post mutation.

2.0 METHODOLOGY

2.1 Construction of the 3D Structure of T. asperellum UC1 by Homology Modeling

The study first retrieved relevant data from the NCBI needed for protein Basic Local Alignment Search Tool (BLAST). This was necessary to identify homologous protein structures in the Protein Data Bank [20]. The amino acid sequences of β-glucosidases were retrieved from GenPept database (ID: XP_018660766.2) in NCBI. Based on the highest percentage of sequence similarity of alignment (90.06% similarity) for BGL (GenBank accession No. AW78142.1) from another T. asperellum, isolated by another study, was selected as the structural template. Next, the target enzyme's homology modeling used the SWISS-MODEL comparative protein modeling server to construct the three-dimensional (3D) structure of the BGL of Trichoderma asperellum UC1. The generated three-dimensional (3D) structure of the fungal β-glucosidase of T. asperellum UC1 was saved as a PDB file.

2.2 In Silico Site-directed Mutagenesis of BGL T. asperellum UC1

Based on a previous study, three structurally conserved active site residues of the BGL were reportedly located at Glu165, Asp226, and Glu423 [11]. In this study, mutations of catalytic residues of BGL to alanine (Ala) were introduced by Pymol 2.3 to yield mutant residues Glu165Ala, Asp226Ala and Glu423Ala, respectively. The mutant β-glucosidase was analyzed on an on-line Site Directed Mutator server (http://www-cryst.bioc.cam.ac.uk/~sdm/sdm.php) to assess for any changes in protein stability after the triple mutations. The analysis was based on a specific structural environment of a known 3D structure of another BGL [21]. The sequence of a highly similar BGL was obtained from a previously isolated T. asperellum (Genebank accession: ARW78142.1).

2.3 Model Refinement by Energy Minimization

Refinement of the generated model structure of the mutant BGL UC1 was then completed by allowing a subroutine within the energy minimization to replace missing atoms of altered residues before minimizing the structure using the appropriate parameters. Energy minimization is mandatory for protein structures with mutations. The process refines the generated protein model and improves any significant errors from its native structure [22]. Molecular dynamics (MD) simulation to refine the BGL of T. asperellum UC1 model was carried out on a GROMACS 2018.6 using the Gromos96 53a7 forcefield. Here, energy minimization by the steepest descent and conjugate gradient methods was done [23]. This step is important to establish the native state of the BGL UC1 protein structure and that it was truly at the global minimum [24]. The optimization process to obtain a better protein structure converges on achieving the model enzyme’s highest possible absolute performance [25].

2.4 Evaluation of Structural Validity

The default parameters of ERRAT checked the constructed mutant BGL UC1 model's reliability and
quality to evaluate non-bonded interactions [26]. Next, VERIFY3D checked the sequences’ compatibility to the structure [27, 28] using the SAVES server (http://services.mbi.ucla.edu/SAVES/). The PROCHECK examined the stereochemical quality and authenticated the constructed 3D model of the BGL protein in the Ramachandran plot. This plot typically evaluates the correctness of backbone conformation based on phi/psi distribution and the presence of non-GLY residues at the disallowed regions in the model [28].

2.5 Substrate Docking

The ligands' 3D structures were constructed in ChemSketch [29], and docking simulations were performed on the Autodock version 4.2.6 and AutoTools 1.5.6.[20]. The binding region was defined by the Autogrid tool in Autodock as ±1.000 Å from 39.214 Å, 39.185 Å, and 42.877 Å coordinates with the sizes 22, 16, and 14 (x, y, and z positions, respectively). The docking analysis was performed in triplicate using the Autodock Vina for comparison purposes. For each substrate in this work, the best result was taken as the largest conformation cluster showing the lowest binding energy. The “.pbdqt” file for each BGL-substrate complex was converted into the pdb format and visualized using Pymol version 2.3[30].

2.6 MD Simulation on Enzyme-ligand Complex

BGL-substrate complexes were converted into the pdb format and visualized using Pymol version 2. MD simulation for refining each mutant BGL 3D model was done on the parallel version of GROMACS 5.1.2 using the Gromo96 S3a6 force field. The mutant BGL protein was simulated in a cubic simulation box (10.0 x 10.0 x 10.0 nm³) and solvated with 18000 SPC/E water molecules through the addition of counterions to neutralize the net charge of the enzymes [11, 30]. A total of 7 Na⁺ were added for the BGL, and the system was energy-minimized using the steepest descent algorithm. A maximum of 10,000 steps was used to ensure the protein-ligand complex's solvated system was free from steric clashes or incorrect geometry. The structures of ligand-bound mutated β-glucosidase proteins were evaluated. NVT and NPT were used to equilibrate the complete system by compiling for 50,000 steps (100 ps) at 300K at 1 atm [30]. The equilibrated structures were subjected to MD simulations for 100 ns in triplicate, with an integration time steps of 2 fs. Each output was obtained as a finished Xmgrace graph to analyze the simulation trajectory. The established step-by-step configuration in this work was founded on published protocols [31]. Analysis of the dynamic behavior and structural changes of the BGL protein model was performed by calculating the root mean square deviation (RMSD) and root mean square fluctuation (RMSF) at the end of the 100 ns production simulation [30]. RMSD quantifies the deviation from the minimized crystal structure, where each part of the protein has altered from the initial conformation over the production simulation time. RMSF exemplifies the deviation from the mean structure over a dynamic production simulation and demonstrates the protein’s moving regions. In this work, the frame was stored at every 1.0 ps intervals during the simulation.

3.0 RESULTS AND DISCUSSION

3.1 In Silico Site-directed Mutagenesis

Previously, a BLAST search through the Protein Data Bank (PDB) using the target sequence as the query yielded the closest similarity to BGL from a T. asperellum species (ARW78142.1) [30]. BGL UC1 enzyme showed 90.04% % of sequence identities to two other T. reesei fungi with PDB identification number of 3AHY. The multiple sequence alignment of some BGL proteins was reported in our previous study, which showed their predicted conserved catalytic trial (Figure 1). The RMSD of the wild-type BGL model (0.51 °A) and the mutant enzyme (1.73 °A) were both in the acceptance range (RMSD < 2 °A). In this study, the putative catalytic residues, namely the Glu165, Asp226, and Glu423, were mutated into alanine by the Pymol 2.3 program to give mutants Glu165Ala, Asp226Ala, and Glu423Ala, respectively. The sequence was subjected to homology modeling using the SWISS-MODEL web server to generate the 3D structures of the mutant BGL. Figures 1a and 1b depict the SWISS-MODEL web server generated 3D structure of the BGL UC1 in cartoon. Literature has shown that a >30% degree of sequence similarity between the target-template is acceptable for homology modeling studies [27, 32].

A naked-eye inspection of the wild-type and mutant BGL 3D structures yielded non-discriminable structural differences. It appears that the generated 3-D structures of the mutant BGL protein preserved a similar fold as the wild-type enzyme (Figure 2). The study assumed that the backbone fold of the protein was retained after the substitution of the enzyme's catalytic residues with three alanine residues. There were subtle changes to the mutant BGL UC1 model structure, with slight rotations of side-chain torsions, revealing a minimum mixture of side-chain and main-chain shifts.
Figure 1 Multiple sequence alignment displaying the conserved catalytic triad of BGL UC1 protein (Glu165-Asp226-Glu423)

Figure 2 The SWISS-MODEL generated 3D structures of (a) wild-type BGL (b) mutant BGL UC1. Red, yellow, and green represent the α-helices, β-sheet, and loop of the enzyme

3.2 Energy Minimization and Evaluation on the 3D Structure of Mutant BGL UC1

All the models generated were energy minimized. Local strains within the generated 3D protein model for both wild-type and mutant BGL models were eliminated by energy minimization using Gromos97 to procure stable structures for the subsequent substrate docking. This is because minor errors from localized strain(s) can exist in the original protein structure, such as atomic overlap or bad Van der Waals interaction [30]. Energy minimization of enzymes helped remove any local strain after the addition of hydrogens. Also, a broken hydrogen bond network in water would lead to large forces and structure distortion. Therefore, PROCHECK, ERRAT, and Verify3D were used as indicators to verify and validate this data (Table 1).

In this study, the energy minimized mutant model of BGL UC1 was docked with substrates, cellulose, hemicellulose, and lignin before molecular dynamic (MD) simulations. This step is important to observe interactions between substrates with the new catalytic residues mutant BGL model, namely Glu165Ala, Asp226Ala, and Glu423Ala. This study anticipated an increase in the binding energy between the substrates and the mutant BGL catalytic residues. Crucially, a model structure showing higher binding energy signifies a negative or poorer interaction between the substrates and catalytic site. It also indicates the likelihood of catalytic failure. The energy minimized mutant BGL UC1 was subjected to several model evaluation tools to assess its quality (Table 1). In this study, evaluations were done on the local geometry (PROCHECK and ERRAT), while Verify3D gauged the mutant enzyme’s local environment [27, 30, 33].

Table 1 Summary of the validation result of wild-type and mutant BGL models before and after energy minimization

| Model Evaluation Tools | Normal Score Range (%) | Wild-type BGL (%) | Mutant BGL (%) |
|------------------------|------------------------|------------------|---------------|
| Procheck               | >90                    | 90.0             | 90.5          |
| Verify3D              | >80                    | 97.65            | 98.85         |
| ERRAT                  | >50                    | 93.41            | 92.09         |

3.2.1 PROCHECK

PROCHECK examines the geometrical properties of protein molecules by analyzing the residue-by-residue geometry of the structure. A protein model’s acceptable quality has over >90% residues within the most favored regions [33]. The quality of the mutant BGL UC1 was also evaluated based on the backbone conformation and overall stereochemical by observing the phi (Φ) and psi (ψ) torsion angles. The output of the analysis was demonstrated by Ramachandran plots (Figure 3).

Figure 3 Plot calculation on the 3D models of BGL UC1 computed by the PROCHECK program. The following is the comparison of Ramachandran’s plot for (a) wild-type BGL UC1, (b) energy minimized wild-type BGL UC1 model, (c) mutant BGL UC1 and (d) energy minimized mutant BGL UC1 model. The most favored regions [A, B, L], the additional allowed regions [a, b, l, p], the generously allowed regions [−a, −b, −l, −p] are colored in red, yellow, and pale yellow respectively. All non-glycine and proline residues are illustrated as a filled black square, and glycine (non-end) are indicated as filled black triangles; disallowed regions are colored in white.
Despite mutation of the catalytic residues, it appears that the 3D model of mutant BGL UC1 retained good stereochemical quality as that of the wild-type enzyme. This was apparent in the Ramachandran plots where the plotted (Φ, ψ) predicted model of the wild-type BGL UC1, refined BGL, mutant BGL, and refined Mut-BGL (Figure 3) shared distribution of 99.5% amino acid residues in the allowed regions. About 0.5% of residues resided in a disallowed region. This meant that little has changed in terms of the quality of the protein structure post-mutation. For brevity, the most favored regions are labelled as [A, B, L], the additional allowed regions as [a, b, l, p], and the generously allowed regions are [~a, ~b, ~l, ~p].

Summary of the Ramachandran plot statistics for BGL of T. asperellum UC1 by PROCHECK is presented in Table 2.

| Stereochemical parameter | Calculated Values (%) |
|--------------------------|------------------------|
|                         | Wild-type BGL (%)      | Refined Wild-type BGL | Mutant BGL | Refined Mutant BGL |
| Residue in most favoured regions [A, B, L] | 91.0%                  | 89.9%                  | 89.9%          | 90.2%          |
| Residue in the additionally allowed zones [a, b, l, p] | 8.3%                   | 9.3%                   | 9.3%         | 9.0%         |
| Residue in the generously allowed regions [~a, ~b, ~l, ~p] | 0.3%                | 0.3%                   | 0.3%       | 0.3%       |
| Residue in disallowed regions | 0.5%               | 0.5%                   | 0.5%       | 0.5%       |
| Non-glycine and non-proline residues | 100.0%             | 100.0%                 | 100.0%     | 100.0%     |

3.2.3 VERIFY-3D

Data calculated using Verify3D are useful in establishing an atomic model’s compatibility with its own amino acid sequence. This is done by assigning a structural class based on its location and environment. Kuriata et al. [34] deemed that a satisfactory Verify3D score should be >80%. In this study, the data of Verify3D revealed the wild-type BGL UC1 has a score of 98.92% while refined- and the refined mutant BGL UC1 showed a marginally reduced common score of 98.06% (Figure 5). The high scores for Verify3D seen here indicated the tested protein models achieved good side chain environments. In all, the integrated data of PROCHECK, ERRAT, and Verify3D for the mutant BGL UC1 exceeded the minimum cut-off score for a good 3-D structure. This conveyed that the mutant BGL UC1 maintained overall structural stability close to its wild-type counterpart.

![Figure 4 Overall quality of the models evaluated by the ERRAT for (a) non-refined BGL, (b) BGL after refinement, (c) mutated BGL, and (d) refined Mut-BGL. Black bars represent the poorly modeled regions, grey bars depict the error region, while white bars indicate the region with a lower error rate for protein folding.](image)
Figure 5 Results of Verify3D for (a) wild-type-, (b) refined-, (c) mutant-and (d) refined mutant BGL UC1 with the score corresponding to 98.92%, 98.06%, 98.06% and 98.06%, respectively.

3.4 The 3D Modeling of Refined Mutant BGL UC1 Structure

Comparison of 3D poses of refined BGL UC1 and mutant BGL UC1 is presented in Figure 6. There were subtle differences in the protein’s conformation in the proximity of the mutation sites, obtained from the superimposed proteins of the wild-type and mutant BGL UC1 (Figure 2 (a)). Protein side chains at positions 165, 226, and 423 of wild-type BGL UC1 appeared longer and more kinked. These changes were consistent with the presence of larger side chains of three- and two carbon carboxylic acids of glutamic acid (residue 165) and aspartic acid (residues 226 and 423), respectively [35]. Whereas residues E165A, D226A, and D423A of Mut-BGL UC1 adopted a more compact structure that corresponded well with the smaller methyl (−CH3) side chain of alanine.

To further examine the crucial key catalytic residues of mutant BGL UC1 for protein-chemical interactions, the in silico site-directed mutagenesis showing mutations on the catalytic amino acid residues was used to create substitutions, followed by docking simulations [36]. Herein, the conserved residues 165, 265, and 423 of the mutant BGL UC1 were mutated from Glu and Asp to Ala. The introduction of non-polar residues will hypothetically nullify hydrogen bonds forming between the docked substrates and the catalytic residues of BGL UC1. In this investigation, the results of the on-line Site Directed Mutator data showed that residues E165 (ΔΔG =), D226 (ΔΔG=) and E423 (ΔΔG =) yield values of ΔΔG corresponding to − 0.51 kcal mol−1, − 0.14 kcal mol−1 and − 1.13 kcal mol−1. The predicted data was that the resultant alanine scanning mutation on BGL UC1 had destabilized the 3D structure, proven by the decrease in protein stability post mutation. A positive and negative sign corresponds to destabilizing and stabilizing mutations predicted to decrease and increase the binding affinity of an enzyme-substrate complex. Literature has shown that destabilizing mutations at catalytic residues typically result in gains in binding energies of proteins or enzyme-protein complexes [37].
3.5 Molecular Docking of Mutant BGL UC1 with Cellulose, Hemicellulose, and Lignin

This study's subsequent investigation involved molecular docking of the natural substrates, cellulose, hemicellulose, and lignin into the refined structures of the wild-type- and mutant BGL UC1. This investigation compared the interaction between the enzyme and the three ligands using the AutoDock version 4.2.6 and AutoTools 1.5.6, followed by AutoGrid tools. Considering that mutant BGL UC1 has three alanine residues replacing the putative catalytic residues in BGL UC1 (Glu165, Asp226, and Glu423), hydrogen bonds between the carbonyl oxygen atom of the substrates were not expected to occur. Another scenario that may transpire was forming another type of hydrophobic interaction (in lieu of alanine's neutral nature) with a ‘neighboring’ amino acid. This scenario is possible for the mutant BGL UC1 as residues in its catalytic pocket are tightly-packed as any other enzymes. Close-ranging residues are bound to be attracted to one another, as ‘likes attract likes.’ Furthermore, the complexity of a protein structure is determined by the sequence of amino acids. Meanwhile, the available group side chains' chemical nature and the polar and nonpolar side chains, alongside the twisting and conformations of proteins, are caused by hydrophobic and hydrophilic interactions. Hence, alanine’s hydrophobicity enables the residue to be ‘wedged’ into tight loops or chains.

A previous study has shown that the wild-type BGL UC1 docked with cellulose (−8.1 kcal mol⁻¹) recorded the most favorable binding energy followed by lignin (−7.9 kcal mol⁻¹) and hemicellulose (−7.5 kcal mol⁻¹). As anticipated, mutation with alanine yields the BGL UC1-substrate complex showing increased binding energies to the substrate lignin (−7.4 kcal mol⁻¹), cellulose (−7.2 kcal mol⁻¹), and hemicellulose (−7.2 kcal mol⁻¹) (Table 2). Next, the docking study's binding energy was used to calculate the equilibrium dissociation constant (Kd) as described by Manas et al. [38]. Low binding energy is related to a low Kd value that signifies high binding affinity, and this can measure substrate-binding strength with the enzyme. From the docking analysis of the mutant GBL model, lignin exhibited the lowest Kd value (2.8239 X 10⁻⁴), followed by cellulose and hemicellulose (3.9887 X 10⁻⁴). Conversely, the Kd was constant for cellulose and hemicellulose (Table 3). The data proved that lignin was bound tightly to BGL UC1. On the other hand, the wild-type BGL showed that cellulose displayed the lowest Kd value (8.4322 X 10⁻⁷), followed by hemicellulose (1.415 X 10⁻⁷) and cellulose (1.191 X 10⁻⁷) (Table 3). The results seen here indicated the poor interaction between the mutant BGL UC1-substrate complex after the triple substitutions of Glu165, Asp265, and Glu423 with alanine. The data was also affirmative of the residues' important role in the hydrolysis of the three aforesaid natural substrates of BGL. Interestingly, the mutation led to a change in the substrate-binding preference order of the mutant BGL UC1. The mutant enzyme preferred lignin over hemicellulose and cellulose. Conversely, the wild-type BGL favorably hydrolyzed cellulose, followed by lignin and hemicellulose [11]. Thus, our in silico study's outcome supported the catalytic role of residues Glu165, Asp265, and Glu423 in BGL UC1 to hydrolyze cellulose, lignin, and hemicellulose [1]. Thus, the increased binding energies strongly indicated that replacing the catalytic residues with alanine has marked adverse repercussions on hydrolytic activity and substrate specificity of the mutant of BGL UC1.

The best interaction poses for the protein-ligand complex of mutant BGL UC1 model are illustrated in Figure 7. The lower binding energy recorded for lignin indicated a stronger binding in the protein-ligand complex interaction, as the carboxylate group of BGL UC1 formed hydrogen bonds to Asn224 and Trp424 at distances of 2.3 Å and 3.0 Å (< 3.0 Å) (Figure 6c), respectively. Mutation to alanine led to weaker hydrogen bond interactions in mutant BGL UC1 model with cellulose through residues Asn164 (2.3 Å), Asn224 (2.0 Å) (Figure 7a), and Trp424 (1.8 Å). The same was also observed for hemicellulose through Asn164 (2.1 Å) and Asn224 (1.6 Å) (Figure 7b). Intermolecular distances of less than 3 Å are universally accepted, as which hydrogen bonds are formed [39, 40]. Pertinently, the appreciable increase in binding energies for all mutant BGL UC1-substrate complexes, alongside the reduced number of hydrogen bonds in the enzyme-hemicellulose and enzyme-lignin complexes, were clear indications of their reduced stability of interactions. The findings thus hypothetically validated the reduced ability of the mutant BGL UC1 to hydrolyze all three substrates while arguably, affirming the catalytic role of Glu165, Asp226, and Glu423.

Table 3 Comparison of the minimum binding energies in kcal/mol, as estimated for the wild-type- and mutant BGL UC1 as calculated by AutoDock.

| Ligand      | Binding Energy (kcal/mol) | Residues                        |
|-------------|---------------------------|---------------------------------|
|             | Wild-type                 | Mutant                          |
| Cellulose   | −8.1                      | −7.2 Asn224 - Asn164 - Trp424   |
| Hemicellulose| −7.8                      | −7.2 Asn164 - Asn224            |
| Lignin      | −7.9                      | −7.4 Asn224 - Trp424            |
Figure 7 The best interaction poses for the protein-ligand complex by molecular docking: (a) mutant BGL-cellulose, (b) mutant BGL-hemicellulose, and (c) mutant BGL UC1-lignin complexes. (a) mutant BGL-cellulose formed hydrogen bond at Asn224, Asn164, and Trp424, (b) mutant BGL interact with hemicellulose by hydrogen bonding at Asn164 and Asn224 residue, and (c) mutant BGL UC1-lignin complexes formed the hydrogen bond at Asn224 and Trp424. The hydrogen bond distances are illustrated as yellow dashed lines.

3.6 MD Simulations of the BGL UC1-ligand Complex

Protein-ligand interactions are a prerequisite to visualizing the structural and dynamic characteristics of the mutant BGL UC1 model. In this study, the initial protein-ligand molecular interactions were docked by AutoDock 4.2 using the classical Lamarckian genetic algorithm (LGA) as energy optimization algorithms. The best conformations of each docking algorithm were then subjected to molecular dynamic (MD) simulations to analyze the molecular mechanisms of protein-ligand interactions further. Here, we analyzed the binding energy between the protein receptors and ligands, hydrogen bonds’ interactions in the docking region, and the structural changes when the wild-type- and mutant BGL UC1 model interact with all three substrates. In this study, MD simulation was performed by looking at the total root mean square deviation (RMSD) and total root mean square fluctuation (RMSF) of the Ca backbone-backbone. We assessed the RMSD to study the residue behavior of the protein during the simulations. In general, a residue’s RMSD value represents the local flexibility of a protein. It reflects the mobility of an atom during the MD simulation trajectory. Therefore, a higher residue RMSD value indicates a higher mobility. Conversely, a lower residue RMSD value indicates lower mobility. A relatively lower RMSD value indicates the complex structures’ high stability and vice versa [41, 42]. Therefore, a complex’s stability is affected by the binding affinity between a ligand and its target molecule.

The MD simulations of BGL-cellulose complexes were run in a 10 Å cubic box containing water molecules at 300K, and the spc216 template was applied to solvate the protein [11]. All MD simulations were carried out by the GROMACS 5.1.2 package using the Gromas96 53a6 force field. The BGL UC1-substrate complexes’ trajectories were appreciably stable for the most part of the 100 ns production simulation run. The trajectory stability was checked and corroborated by the RMSD analysis (Figure 8) as the function of time for the mutant BGL UC1-substrate complexes. The RMSD values for all mutant BGL complexes increased in the first 10 ns and stabilized for most of the simulation (Figure 8a). RMSD value of the mutant BGL UC1-cellulose revealed that the system first reached equilibrium at 30 ns (2.4 Å) and fluctuated closely between 1.2 Å to 2.5 Å. Notably, MD simulation for mutant BGL UC1-lignin equilibrated soon after 42 ns (2.5 Å), and its corresponding RMSD value fluctuated between 1.2 Å to 2.8 Å. The complex incrementally deviated (RMSD ~ 2.5 Å) and fluctuated again to 2.8 at 98 ns (Figure 8a). Conversely, the mutant β-glucosidase-hemicellulose complex RMSD value fluctuated between 1.1 Å to 2.5 Å while equilibrated at 72 ns (2.4 Å).
Confirms an early study by Cheng et al. [43]. A RMSD value that fluctuates > 3.0 Å is acceptable for stable protein structure [44]. However, the MD simulations results for mutant BGL UC1 negated the wild-type BGL UC1 substrate docking data which showed lignin as the preferred substrate. Again, our results proved that alanine scanning on the BGL UC1 catalytic triad did alter the enzyme’s specificity and catalytic properties, as stated by [44]. Nonetheless, such a change requires further empirical study since this study focused on the consequence of the in silico mutation on BGL UC1.

To identify the flexible residues in the mutant BGL UC1-substrate complexes, root mean square fluctuations (RMSF) were calculated from the MD trajectories. The residue number is shown in the abscissa axis, and the RMSF for the Ca of each residue is the inordinate axis. It is worth mentioning that the threshold value of RMSF at > 0.5 Å represents a significant change in structural movements that correspond with enzyme stability [45]. As can be seen, the RMSF plots appeared stable for most of the simulation trajectory, fluctuating between 0.5 Å to 3.1 Å for all mutant BGL UC1-substrate complexes (Figure 9).

The highest RMSF peak for the mutant BGL UC1-cellulose complex was sited at residue 315 at 3.1 Å. Several other fluctuations were also identified at residues 23, 50, 218, and 440, with values ranging between 2.4 Å to 3.0 Å. Comparingly, the wild-type BGL UC1-cellulose highest peak was recorded differently compared to its mutant-cellulose complex for residue 48 at 3.0 Å [11]. This was an apparent distinction in the mutant BGL UC1 protein's flexibility compared to the wild-type, despite their close structural similarity seen in earlier PROCHECK, ERRAT, and Verify3D assessments.

As shown in Figure 8, a small fluctuation is incidental of the a-helix region's low flexibility compared to its average position, which implied a rigid secondary structure conformation. RMSF value of wild-type BGL UC1 appeared stable for most of the simulation trajectory, fluctuating only between 0.5-3.1 Å. Comparingly, mutant BGL UC1 complexes recorded the same fluctuation average. However, the wild-type BGL UC1-cellulose complex's highest peak was recorded differently compared to its mutant-cellulose complex for residue 48 at 3.0 Å [11]. This was an apparent distinction in the flexibility of the mutant BGL UC1 protein compared to the wild-type. The protein fluctuation was then related to the catalytic residue of mutant BGL UC1 to recognize and interact with all three substrates as mutation took place. This can further influence the complexes' behavior and catalytic activity, resulting in the enzyme's poor ability to bind all three substrates.

Hence, it can be construed that the observed disparity in protein fluctuation was related to the mutant BGL UC1 catalytic residues' ability to recognize and interact with the substrates (cellulose, hemicellulose, and lignin). This was consistent with the markedly poorer ability of the enzyme to bind the substrates. Finally, this study's findings collectively affirmed the crucial role of Glu165, Asp26, and Glu423 as the catalytic residues of BGL secreted by the fungus T. asperellum UC1.
4.0 CONCLUSION

Based on this research, this study successfully constructed a 3D structure of mutant BGL UC1 using the SWISS-MODEL data extracted from NCBI based on the BGL (ARW78142.1) amino acid sequence. This allowed the subsequent in silico site-directed mutation on the catalytic triad (Glu165, Asp226, and Glu423) with the neutral and chargeless amino acid, alanine. The in-silico docking of substrates, cellulose, lignin, and hemicellulose (ligand) into the wild-type and mutant BGL UC1 active sites corroborated the crucial catalytic role of Glu165, Asp226, and Glu423 in the enzyme. This was proven after comparing the binding energies of the wild-type BGL UC1 with its mutant counterpart. A notable increase in the enzyme-substrates complexes’ binding energies proved the diminished catalytic role of the active site of mutant BGL UC1 to bond the substrates tightly for further hydrolysis. Thus, we demonstrated that the alanine scanning could assert that the Glu165, Asp226, and Glu423 were pertinent in hydrolyzing the substrates.

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