Research Article

Gas Chromatography-Mass Spectrometry Coupled with Multivariate Statistical Analysis to Identify the Alpha Glucosidase Inhibitors from Flesh of Salacca zalacca Fruits and Their Molecular Docking Studies

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Fruit of salak (Salacca zalacca) is traditionally used and commercialized as an antidiabetic agent. However, scientific evidence to prove this folk claim is quite lacking. Therefore, this research was aimed to evaluate the α-glucosidase inhibition activity of S. zalacca fruit and identify the bioactive compounds. The fruits were extracted by different ratios of ethanol and water (0, 20, 40, 60, 80, 100%, v/v) to get E0 (100% water), E20 (20% ethanol), E40 (40% ethanol), E60 (60% ethanol), E80 (80% ethanol), and E100 (100% ethanol) extracts. The extracts obtained were subjected to the α-glucosidase inhibitory assay. Gas chromatography-mass spectrometry-based metabolomics approach was used in profiling the bioactive metabolites present in the extracts. Orthogonal partial least square (OPLS) was used to correlate GC-MS data and α-glucosidase assay results to identify the possible chemical markers. All active compounds identified were subjected to molecular docking. The extracts from the S. zalacca fruit showed potent inhibition activity against α-glucosidase. The IC50 values from the α-glucosidase inhibitory assay ranged between 16 and 275 µg/ml. Overall, E60 displayed significantly higher α-glucosidase inhibition activity, while E0 showed the lowest α-glucosidase inhibition activity. Major compounds detected in S. zalacca fruits were sugars, fatty acids, and sterols, including myo-inositol, palmitic acid, stearic acid, and β-sitosterol. Moreover, the results obtained from molecular docking indicated that palmitic acid and β-sitosterol were close to the active side of the enzyme. Some of the residues that interacted include HID295, ASN259, LEU313, LYS125, PHE159, VAL216, PHE178, TYR72, TYR158, HIE315, ARG315, and PHE303. The bioassay result strongly suggests that E60 extract from S. zalacca fruits has potential α-glucosidase inhibitory activity. The hydrophobic compounds, including palmitic acid and β-sitosterol, were found to induce the α-glucosidase inhibition activity.
1. Introduction

Diabetes mellitus (DM) is a chronic metabolic disorder that may lead to other macro- and microvascular complications. The highest number of DM cases that have been reported to date was recorded in Asian countries, which cover about 60% of the world’s DM patients [1]. World populations majorly practice conventional medicines relying on plant-based medicines due to the availability [2]. Various plants have been utilized as a traditional remedy for multiple diseases and conditions for centuries [3]. The effect and mechanism of most plants for the treatment of diabetes can be related to their inhibitory activities against α-glucosidase enzyme. This intestinal enzyme is found in the brush border of the small intestine, responsible for catalysing the breakdown of polysaccharides into monosaccharides, thus reducing the postprandial glucose level, thereby decreasing the hyperglycaemia in DM patients [4]. In recent times, various plants and herbs have been extensively studied, providing scientific proof that the phytoconstituents present in plant matrix have various pharmacological effects on the human body system, such as fermented Chinese Ge-Gen-Qin-Lian decoction and Radix Salvia Miltiorrhiza [5–7]. *Salacca zalacca* is one example of a plant that is traditionally used as a hypoglycemic remedy, more commonly popular in Asian countries, including Malaysia, Myanmar, Indonesia, and Thailand [8, 9].

*Salacca zalacca* Gaertn. Voss. is one of the palm species belonging to the family Arecaceae and is ubiquitously found all around Southeast Asia. *S. zalacca* fruits are a good source of vitamins, fibre, minerals, and carbohydrates [10]. Moreover, *S. zalacca* possesses antioxidant, anticancer, and anti-diabetic effects [11–14], attributable to their high levels of vitamin C, Phenolics acids, and flavonoids [10]. In recent times, it has been claimed to possess anti-diabetic activity [12, 14]. Therefore, this study intends to initially analyse the α-glucosidase (EC 3.2.1.20, α-D-glucoside glucohydrolase) inhibitory activity of the *S. zalacca* fruit extracts. It is one of the pathways involved in the management of DM. One of the most appropriate methods to analyse these constituents is by using the metabolomics approach.

Typically, the analysis of metabolites can be achieved by herbal fingerprinting that requires the use of different chromatography and spectroscopy techniques, such as gas chromatography-mass spectrometry (GC-MS), and nuclear magnetic resonance (NMR) [15–18]. Therefore, the aim of the present study is to correlate the metabolic variation between *S. zalacca* extracts and α-glucosidase inhibitory activity using the GC-MS-based metabolomics approach. This study is planned to identify the related bioactive compounds in different ratios of ethanol-water and molecular docking were also applied to predict the inhibitory activity.

2. Materials and Methods

2.1. Authentication of the Sample. Fresh *S. zalacca* (Gaertn.) Voss) fruits were harvested from a local farm in Bukit Sagu plantation, Felda Sagu, Kuantan, Pahang DM, Malaysia, during the month of May 2017. A voucher specimen has been submitted to the Kulliyyah of Pharmacy, International Islamic University of Malaysia. It was identified by Dr. Shamsul Khamis (Botanist) and deposited in the herbarium of Kulliyyah of Pharmacy with the voucher specimen number, i.e., PIUM 0269–2 [VS-4].

2.2. Chemicals. Ethanol (analytical grade), dimethyl sulfoxide (DMSO) were obtained from Merck Germany (Darmstadt, Germany). α-Glucosidase enzyme was obtained from Megazyme (Wicklow, Ireland). 4-nitrophenyl β-D-glucopyranoside substrate, methoxyamine hydrochloride, pyridine, *N*-Methyl-N-(trimethylsilyl) trifluoroacetamide, and quercetin (≥95% HPLC) were obtained from Sigma-Aldrich (St. Louis, MO, USA).

2.3. Preparation of Extracts for Metabolomics Studies. The fruits of *S. zalacca* were cleaned with tap water and then peeled into peel and flesh. Immediately, the separated fruits flesh was cut into small cups to remove the seeds. Initially, the fruits flesh was weighed and dried using a freeze dryer. The freeze-dried samples were ground into a fine powder using a laboratory blender and then kept at −80°C prior to extraction. Five grams of dried powdered fruit was suspended in different ratios of ethanol and water (0, 20, 40, 60, 80, 100%, v/v). Approximately 150 mL of the desired solvent was mixed with powdered and shaken before being transferred to sonicator at 25°C for 1 h. A total of 36 extracts were obtained and filtered through a Whatman no. 1 filter paper. Finally, the filtered solutions were evaporated using a rotary evaporator at 40 ± 1°C. The stock concentrations of extracts were dried using a freeze dryer. The obtained amount of the extracts was weighed before keeping at −80°C prior to further analysis [4].

2.4. α-Glucosidase Inhibitory Activity. The α-glucosidase inhibitory activity was measured using a method described by Collins et al. [19] with slight modifications. Different dilutions of the *S. zalacca* extracts were prepared in triplicate (i.e., 320, 160, 80, 40, 20, 10, and 5 µg/mL). Then, 10 µL of each dilution was added to 100 µL of 30 mM phosphate buffer pH 6.5. After that, 15 µL of α-glucosidase enzyme solution (0.02 U/well) was added. The mixture was allowed to stand for 5 min before adding 75 µL of the substrate (PNPG) (6 mg in 20 mL of 50 mM phosphate buffer pH 6.5). A total of 50 µL of (2 M) glycine of pH 10 was added to the mixture to stop the reaction. The mixture was immediately measuring the absorbance at 405 nm. The α-glucosidase inhibition was decreased using the equation:

\[
\alpha - \text{glucosidase inhibitory activity (\%)} = \left[\frac{A_{\text{control}} - A_{\text{sample}}}{A_{\text{control}}}\right] \times 100\%.
\]  

IC$_{50}$ of the extract was determined from the graph of α-glucosidase inhibitory activity (%) against the amount of extract (µg).
2.5. GC-MS Analysis

2.5.1. Derivatization Procedure. The samples of extracts were derivatized using N-Methyl-N-(trimethylsilyl) tri-fluoroacetamide (MSTFA) and methoxyamine HCl for GC-MS analysis following the method described by Robinson et al. (2005) [20]. Briefly, about 12.5 mg S. zalacca fruits ethanolic extract was dissolved in 25 µL pyridine in a 1 mL Eppendorf microcentrifuge tube. The mixture was initially vortexed for 2 min then further sonicated for 20 min at 30 °C. About 50 µL (20 mg/mL in pyridine) was added to the sample solution and incubated for 2 h at 60 °C. About 150 µL of MSTFA was added and incubated again for 30 min at 60 °C. The derived sample was filtered using a microsyringe filter and equilibrated to room temperature overnight.

2.5.2. GC-MS-Based Metabolomics. GC-MS analysis of the S. zalacca fruits ethanolic extracts was carried out following the method described by Saleh et al. [11]. GC-MS analysis was performed on an Agilent 6890 gas chromatograph equipped with a 5973-mass selective detector using an electron impact ionization. A DB-5MS, 5% phenyl methyl siloxane column with an inner diameter of 250 µm, and a film thickness of 250 µm attached with an autosampler (Agilent, Santa Clara, United States) was used for the separation. A volume of 1 µL sample was injected in splitless mode into GC-MS. The initial oven temperature was set at 180 °C and the hold time 10 min and then increased to 220 °C at a rate of 20 °C/min with the hold time of 5 min. The final oven temperature was 315 °C at a rate of 30 °C/min with a hold time of 10 min, and the total run time was 30.16 min. Helium gas was used as carrier gas at a flow rate of 1.5 mL/min. The injector and detector were set to 330 °C and 250 °C, respectively. Mass spectra were acquired using a full scan and a monitoring mode in a range of 50–550 m/ζ. Metabolites were identified mostly belonged to sugars, pattern and comparison with the GC-MS NIST14 database. Metabolites were converted into volatile compounds such as acetyl trimethyl silane (TMS) ethers or oxime-TMS ethers [21]. Flame ionization detectors or mass spectrometers are mostly used for detection. Confirmation of the identified metabolites was achieved by carefully examining the spectral pattern and comparison with the GC-MS NIST14 database library. The metabolites identified mostly belonged to sugars, sterols, as well as fatty acids that include D-(-)-fructofuranose, D-(-)-fructose, D-psicofuranose, D-glucose, sucrose, D-(-)-trehalose, myo-inositol, β-D-glucopyranoside, D-(+)-xyllose, D-(+)-talofuranose, α-D-glucopyranose, β-sitosterol, stearic acid, palmitic acid, and citric acid (Supplementary Figure S1).

All these metabolites have been labelled and indicated in each of the extracts’ chromatogram analyzed. Specifically, three metabolites were found to be abundant in E60, E80, and E100 extracts, namely stearic acid and palmitic acid, β-sitosterol when compared to E0, E20, and E40. It is pertinent to note that there was a decrease in the intensity of the metabolites in the extract as the polarity of the solvent increased by increasing the amount of water in ethanol. In the meantime, sucrose and D-glucose were found to be dominant in the highly polar E0, E20, and E40 extracts. It is observable that a consecutive decrease in water ratio resulted in a decrease in the polarity. This results in metabolites with less polarity abundantly found in E60, E80, and E100. These include palmitic acid, stearic acid, and β-sitosterol which might be attributed to the α-glucosidase inhibitory activity of S. zalacca fruits hydroethanolic extracts.

3. Results

3.1. Extraction Yield. Table 1 displays the trend of the extraction yield of the S. zalacca fruits extracts. The yield of extraction varied with the ratio of the water and ethanol of the solvents used, with the highest yield obtained from 80% hydroethanolic solvent (E80) as 75.5% and the least yield obtained from the ethanolic (100%, E100) solvent as 20.2% only. The trend of this result can be simplified in the order as 80% (E80) > 60% (E60) > 0% (E0) > 40% (E40) > 20% (E20) > 100% (E100).

3.2. Inhibitory Activity of S. zalacca Fruits Extracts against α-Glucosidase. The activity of S. zalacca extract on the α-glucosidase enzyme is described in Table 1. Results revealed that the highest α-glucosidase inhibition activity was found in E60 extract (IC50 16.48 µg/mL), followed by E100 extract (IC50 20.08 µg/mL), and followed by E80 (IC50 25.09 µg/mL). On the other hand, the E0 extract had the lowest α-glucosidase inhibition activity with IC50 280. 30 µg/mL.

3.3. Metabolite Profiling of S. zalacca Fruits Based on Metabolomics Approach

3.3.1. GC-MS-Based Metabolites Profiling. When chemical profiling using GC-MS, derivatization is required for the metabolites to be converted into volatile compounds such as acetyl trimethyl silane (TMS) ethers or oxime-TMS ethers [21]. Flame ionization detectors or mass spectrometers are mostly used for detection. Confirmation of the identified metabolites was achieved by carefully examining the spectral pattern and comparison with the GC-MS NIST14 database library. The metabolites identified mostly belonged to sugars, sterols, as well as fatty acids that include D-(-)-fructofuranose, D-(-)-fructose, D-psicofuranose, D-glucose, sucrose, D-(-)-trehalose, myo-inositol, β-D-glucopyranoside, D-(+)-xyllose, D-(+)-talofuranose, α-D-glucopyranose, β-sitosterol, stearic acid, palmitic acid, and citric acid (Supplementary Figure S1).

All these metabolites have been labelled and indicated in each of the extracts’ chromatogram analyzed. Specifically, three metabolites were found to be abundant in E60, E80, and E100 extracts, namely stearic acid and palmitic acid, β-sitosterol when compared to E0, E20, and E40. It is pertinent to note that there was a decrease in the intensity of the metabolites in the extract as the polarity of the solvent increased by increasing the amount of water in ethanol. In the meantime, sucrose and D-glucose were found to be dominant in the highly polar E0, E20, and E40 extracts. It is observable that a consecutive decrease in water ratio resulted in a decrease in the polarity. This results in metabolites with less polarity abundantly found in E60, E80, and E100. These include palmitic acid, stearic acid, and β-sitosterol which might be attributed to the α-glucosidase inhibitory activity of S. zalacca fruits hydroethanolic extracts.

2.6. Statistical Analysis. The results were calculated as the mean ± SD from six separate experiments (n = 6). The data were analyzed by one-way ANOVA test followed by Tukey’s test using Minitab version 17. 0.
RMSEE, on the other hand, measures the average deviation of the response values. Higher RMSEE values indicate lower predictive accuracy of the sample. Values represent the mean ± standard deviation (n = 6). The values that do not share a letter are significantly different at P < 0.05. nd = not determined.

### 3.3.2. Multivariate Data Analysis

One of the ways of identifying bioactive metabolites is through the GC-MS-based metabolomics approach using MVDA to develop an orthogonal partial least-squares (OPLS) model. Two results are usually automatically generated, namely OPLS components 1 and 2. The m/z (mass-to-charge ratio) of certain compounds identified was generated along with 1/IC⁵₀ (µg/mL) values of the analytes that were taken as variables x and y, respectively. The major aim of performing OPLS is to determine differences in the constituents among the analytes and to identify the m/z values responsible for the pharmacological activity.

The OPLS model can be assessed through different parameters, namely, the permutation test, the goodness of fit, and the capacity of this model in predicting the y-value via the predicted plot. Meanwhile, the cumulative values R²Y explains more on goodness of fit which indicates the variation percentage of response as explained by the model, whereas the Q²Y, the cumulative value, represents variation percentage of response based on model prediction in accordance with the cross validation. If the cumulative values of R²Y and Q²Y are observed greater than 0.5, then the fitness of the model and predictive ability are considered satisfactory [20]. The R²Y and Q²Y values obtained in this study were 0.856 and 0.925, respectively. In addition to the above-mentioned parameters, other ones that were also considered significant in determining the model accuracy and performance are the root-mean-square error of cross-validation (RMSECV) and root-mean-square errors of estimation (RMSEE). RMSECV simply measures the quality of the model in terms of new sample prediction. Low RMSECV values indicate higher predictive accuracy of the sample. RMSEE, on the other hand, measures the average deviation of the model from the data.

The RMSEE and RMSECV values obtained in this study were 0.356078 and 0.376476, respectively. The score scatter plot of the OPLS model is shown in Figure 1(a). It indicates good separation of the samples in terms of the chemical profile and pharmacological activity (IC⁵₀ value) of the samples. The positive OPLS component 1 consists of samples with higher activity (E00, E100, and E80) having IC⁵₀ value less than 30 µg/mL, whereas the less active samples (E40, E20, and E0) were found at the negative OPLS component 1.

The loading plot of different extracts of S. zalacca fruits hydroethanolic extracts is shown in Figure 1(b). The metabolites in each sample are represented by each peak. The compounds identification and details are shown in Table 2.

Peaks labelled as 1, 2, 3, 5, 6, 7, 9, 11, 13, 15, and 16 represent D-psicofuranose, D-(−)-fructofuranose, D-(−)-fructose, sucrose, D-glucose, D-(+)−trehalose, citric acid, D-(+)−talofuranose, β-D-glucopyranose, α-D-glucopyranose, D-(+)−xyllose, and D-psicopyranose, respectively. They are observed opposite to the 1/IC⁵₀ value. These compounds are identified as glycosylated compounds. The remaining peaks found in the same position aligned with the IC⁵₀ value were efficient in inhibiting α-glucosidase activity. These metabolites were identified as plant fatty acids, carboxylic sugar, and sterol, such as palmitic acid, stearic acid, myo-inositol, and β-sitosterol.

The permutation test plot shown in Figure 2 describes the validation of the developed OPLS model. The plot displayed the fraction intercepts of the sum of the squares Y-value and the predictive ability of the model intercept Y-value. In this study, the R²Y and Q²Y values obtained were 0.507 and −0.408, respectively. These values are acceptable because R² and Q² are less than or equal to −0.5 and 0.5, respectively. Meanwhile, the plot showing the observed versus predicted plot predicts the OPLS model accuracy along its validation using R² value obtained from the regression. This plot is valid if the R² value is greater than 0.9. Therefore, the result obtained is valid with the R² value of 0.9455.

### 3.4. Docking of Bioactive Compounds

All identified compounds responsible for the α-glucosidase inhibition activity were docked to the Saccharomyces cerevisiae isomaltase (SCI) crystal structure (PDB ID: 3A44A). The conformations showing the lowest binding energy for the compounds along with the interacting residues are summarised in Table 3. Based on the AutoDock 1.5.6 simulation result displayed in Table 3, the inhibitor complex of α-glucosidase-quercetin, which was used as positive control showed −8.25 kcal/mol binding energy having six hydrogen bonds with the interacting residues LYS156, ASN415, GLU411, THR310, LEU313, and PRO312 along with hydrophobic interactions involving ARG315 and PHE314. The stearic acid binding affinity in complex with α-glucosidase was observed to be −3.80 kcal/mol. One H-bond was displayed in the complex including CYS342 with hydrophobic interactions with LYS16 and AGR263. Meanwhile, for
α-glucosidase-myo-inositol complex, the binding affinity value of \(-5.68\) kcal/mol was observed. Three residues including PRO456, AGR467, and LYS406 were found to interact with myo-inositol through H-bonds, whereas TYR470 was found to interact via π-bond. It is interesting to know that β-sitosterol, despite interacting with just two residues by H-bonding with its O–H group, LEU313, and LYS125, it comparably displayed better binding energy of \(-9.55\) kcal/mol than the quercetin (positive control). The hydrophobic contacts appeared to dominate in α-glucosidase-β-sitosterol complex, perhaps as a result of its alky group and cyclic skeleton that is preferably bound to PHE159, VAL216, PHE178, TYR72, TYR158, HIE315, ARG315, and PHE303.

The crystallized ligand, α-D-glucose (ADG) demonstrated lower binding energy of 6.02 kcal/mol as compared to β-sitosterol and quercetin (positive control). However, the H-bond existing between ADG and the protein showed a stable interaction with nine amino acid residues, including ASH69, HIE112, ARG213, ASH215, GLH277, HIE351, ASP352, and ARG442 having hydrophobic interaction with TYR72. Unlike the docked ADG, results obtained from docking indicated that the binding of tested compounds was mostly supported through hydrophobic contacts with α-glucosidase. Our result corroborates with those reported by Seong et al. [21] in which the isoflavones, i.e., daidzein, genistein, and calycosin, exhibited more hydrophobic contacts with comparable residues, such as ILE262, ARG263,

Table 2: The metabolites identified in the S. zalacca fruits hydroethanolic extracts using GC-MS-based metabolomics.

| No. | RT (min) | Similarity index (%) | Molecular weight | Molecular formula | Tentative metabolites |
|-----|---------|----------------------|------------------|------------------|----------------------|
| 1   | 20.032  | 90                   | 180.16           | C₆H₁₂O₆          | D-psicofuranose       |
| 2   | 20.638  | 87                   | 180.16           | C₆H₁₂O₆          | D-(-)-fructofuranose  |
| 9   | 21.175  | 91                   | 192.17           | C₆H₈O₇           | Citric acid          |
| 16  | 22.315  | 81                   | 180.16           | C₆H₁₂O₆          | D-psicopyranose       |
| 11  | 23.378  | 94                   | 180.16           | C₆H₁₂O₆          | D-(-)-talofuranose    |
| 3   | 25.719  | 91                   | 180.16           | C₆H₁₂O₆          | D-(-)-fructose        |
| 13  | 26.89   | 95                   | 180.16           | C₆H₁₂O₆          | α-D-glucopyranose     |
| 6   | 27.056  | 90                   | 180.16           | C₆H₁₂O₆          | D-glucose            |
| 14  | 28.685  | 93                   | 180.16           | C₆H₁₂O₆          | β-D-glucopyranose     |
| 10  | 29.331  | 99                   | 256.42           | C₁₈H₃₆O₂          | Palmitic acid         |
| 15  | 25.742  | 91                   | 150.13           | C₆H₁₀O₃           | D-(-)-xyllose         |
| 8   | 29.685  | 93                   | 180.16           | C₆H₁₂O₆          | Myo-inositol          |
| 4   | 30.657  | 99                   | 284.48           | C₁₈H₂₈O₃          | Stearic acid          |
| 7   | 32.817  | 95                   | 342.30           | C₂₉H₄₂O₁₁         | D-(-)-trehalose       |
| 12  | 37.195  | 99                   | 414.71           | C₂₉H₃₀O          | β-Sitosterol          |
| 5   | 37.601  | 91                   | 342.30           | C₁₈H₃₆O₁₁         | Sucrose               |

Figure 1: (a) OPLS score scatter plot of different extracts of S. zalacca fruits; (b) the PC1 loading column plot of different extracts of S. zalacca fruits.
ILE272, and VAL266. The 2D diagram of the interaction between the protein residues and ADG, positive control, stearic acid, palmitic acid, myo-inositol, and β-sitosterol is shown in Figure 3. Moreover, the 3D superimposed diagram of the compounds and ADG is shown in Figure 4(a). It was also observed that the binding site for palmitic acid and β-sitosterol was close to the active site pocket (Figures 4(b) and 4(c)).

4. Discussion

The α-glucosidase enzyme is one of the key enzymes involved in the digestion of carbohydrates in humans. Its main function is to hydrolyse the carbohydrate to release glucose, which may lead to an increase in postprandial blood glucose level. Inhibition of this enzyme is one of the effective ways of controlling the postprandial blood glucose level, which is considered beneficial for diabetic patients. There are several synthetic drugs available in the market for the management and treatment of diabetes, such as miglitol, acarbose, and voglibose. However, they are usually accompanied by serious side effects such as hepatic disorders and other negative gastrointestinal symptoms [22]. Therefore, plant-derived α-glucosidase inhibitors are envisaged as preferable and safer as a means of management and prevention of diabetes. Metabolites obtained from plants have provided a wide range of pharmacological benefits to human health [23]. In many bioactivity analyses of plants, in vitro techniques are applied to determine the pharmacological effectiveness of plant metabolites. In determining the in vitro antidiabetic activity, various procedures are employed. The most important of it is the α-glucosidase inhibitory assay. The pharmacological activity of plant metabolites is dependent on the type and numbers of functional groups present, the chemical structure, as well as the nature of substitution on the ring [24]. Currently, research on the use of medicinal plants has become a continuous effort due to the fact that numerous numbers of bioactive metabolites are distributed in plants with various pharmacological benefits. Water, a mixture of water and organic solvent and different organic solvents are employed to extract the potential metabolites from plants through different methods [4,11]. The polarity of the extracting solvent determines the type of metabolite and hence the bioactivities found in the extract. Due to this fact, different solvents were used in this study to obtain S. zalacca fruits extracts so as to determine different metabolites having α-glucosidase inhibition potential.

The results obtained showed a decrease in the ratio of water, with a moderate decrease in polarity, suggesting that the metabolites responsible for the inhibitory activity are the less polar ones such as sterols and fatty acids. This outcome signifies that the extracting solvent has a greater influence on

| Compound control | Binding energy (kcal/mol) | H-bond interacting residues | Other interacting residues |
|------------------|---------------------------|----------------------------|---------------------------|
| α-D-glucose (control ligand) | −6.02 | ARG442, ASH69, HIE112, ASH215, GLH 277, ARG213, ASP352, HIE351 | |
| Quercetin (positive control) | −8.24 | LYS156, ASN415, GLU411, THR310, LEU313, PRO312 | ARG315, PHE314 |
| Stearic acid | −3.80 | CYS342 | |
| Myo-inositol | −5.68 | PRO456, ARG467, LYS406 | |
| Palmitic acid | −3.59 | HID295, ASN259 | |
| β-Sitosterol | −9.55 | LEU313, LYS125 | PHE159, VAL216, PHE178, TYR72, TYR158, HIE315, ARG315, PHE303 |

Figure 2: Permutation test of different extracts of S. zalacca fruits.
the bioactivity of the extract. Some of the compounds identified have previously been reported in the literature to exhibit in vitro α-glucosidase inhibitory effect. The investigation conducted on α-glucosidase inhibitory activity of palmitic acid and other fatty acids by Liu et al. [25] showed that palmitic acid exhibited inhibitory activity with the IC\textsubscript{50} value > 400 μg/mL which was in contradiction with the value earlier reported in other studies for the same compound, i.e., IC\textsubscript{50} 11.53 μg/mL and 24.09 μg/mL [26, 27]. Aside palmitic acid, other compound that was found active against α-glucosidase was stearic acid 5.3% inhibition at 10 μg/mL [28].

In another study, myo-inositol and β-sitosterol with other compounds were purified from the methanol extract of Tournefortia hartwegiana (Boraginaceae) (METH). An oral glucose tolerance test (OGTT) was conducted using a different substrate such as glucose, maltose, and sucrose through intragastric administration. Results obtained...
showed that the increased plasma glucose level was significantly ($P < 0.05$) suppressed by the extract containing myo-inositol and $\beta$-sitosterol after the administration of the substrates. Moreover, METH also exhibited an \textit{in vitro} $\alpha$-glucosidase inhibitory activity in a concentration-dependent manner (IC$_{50}$: 3.16 mg/mL). Another investigation was conducted by Kim et al. [29] on myo-inositol to determine its effect on the differentiation, lipolysis of adipocytes, insulin-stimulated glucose uptake and to test whether the antidiabetic activity of myo-inositol via is mediated through adipocytes. In the study, 3T3-L1 cells were treated with varying concentrations of myo-inositol. Lipid accumulation was increased after exposure to myo-inositol dose-dependently ($P < 0.01$). An increase in adipogenesis was observed along with insulin-stimulated glucose uptake in adipocytes through activation of insulin signalling. Furthermore, lipolysis in matured adipocytes was also inhibited. Therefore, myo-inositol possesses a promising therapeutic property for improving insulin sensitivities in adipocytes through the increase in lipid storage capacity, glucose uptake, and inhibition of free fatty acid formation.

Chemical and structural derivatives of plant sterols are phytochemicals that are found ubiquitously in many plant parts. Many of them have been previously identified from plant extracts; these include $\beta$-sitosterol. Research conducted on the antidiabetic and antioxidant potentials of $\beta$-sitosterol in STZ-induced rats showed that there was a significant decrease in glycated haemoglobin, nitric oxide, and serum glucose and with a simultaneous increase in serum insulin levels. In addition to that, pancreatic antioxidants levels were also increased with a drastic reduction in thiobarbituric acid-reactive substances [30]. In another study, $\beta$-sitosterol, which is an active phytosterol in the \textit{S. surattense} stem bark, has also been reported to be useful in the treatment of coronary artery disease, breast cancer, prostate cancer, and hypercholesterolemia [31]. Apoptosis induced by caspase 8 activity, FAS receptor levels, and phosphorylation of extracellular signal-regulated kinase and mitogen-activated protein kinase has been suggested to be responsible for the chemopreventive property of $\beta$-sitosterol with no cytotoxicity to normal cells [32, 33].

Although from the structure-activity relationship studies, the presence of multiple hydroxyl groups in a compound has been reported to be responsible for the $\alpha$-glucosidase inhibitory effect, yet, compounds identified using GC-MS in this study were mostly nonpolar compounds which include sterol, fatty acids, excluding myo-inositol. These compounds containing a few numbers of hydroxyl groups except myo-inositol. Despite the same fact, these compounds were found to exhibit a certain level of $\alpha$-glucosidase inhibitory activity based on the literature. This could be as a result of interactions existing between protein and the inhibitors, thereby forming probable conformational complexes that may result in positive enzyme inhibitory effect of the identified inhibitors; myo-inositol, palmitic acid and stearic acid, and $\beta$-sitosterol.

With these results, it is deduced that the hydrophobic and $\pi$-interactions existing between the cyclic part of the compounds structure with hydrophobic side chains (TYR, VAL, and PHE) of the enzyme could be responsible for the much higher activities of these nonpolar compounds, which is agreeable to the finding reported by Nussinov [34] and Du et al. [35]. In addition, the strong hydrogen bond also occurs between the compounds’ hydroxyl group with the residues having both the polar (GLN and SER) and electrically charged side chains (ARG, ASP, and HIE (protonated HIS). This has been observed in quercetin and other similar compounds. Interaction between the residues and that of ADG was observed as shown through its binding energy. As illustrated in Table 3, there was an interaction between ADG and ASH215 residue that acted as a catalytic nucleophile. Furthermore, a strong H-bond was also formed with GLU277 protonated GLU residue, which served as the general acid-base catalyst. Apart from this, ADG also showed interaction with ASP352, which stabilises the substrate during the catalytic reaction so as to strengthen the acid-base hydrolysis reaction [34, 36]. The 3D diagram in Figure 4(a) also indicated that the protein might have more than one site for ligand binding. This could induce allosteric control for catalytic reaction, which may cause changes in the active site, thereby resulting in the enzyme inhibition. Allosteric inhibitors are capable of modifying protein activity [35, 37]. In general view, the conjugated behaviour of a compound in which the conformational perturbations produced at one region of a macromolecule affects the active site and consequently, the enzyme activity confers the allosteric binding ability. In some occasions where the allosteric interaction is unstable, the binding residues stabilises it. This will eventually result in an altered functional site due to the equilibrium shifting toward this conformer or the allosteric ligand, thus causing the inhibition activity [38, 39]. Additionally, Figure 4(a) (the superimposed 3D structure) also suggested that the identified inhibitors using GC-MS displayed noncompetitive inhibition mode when compared to the control ligand, ADG, the known competitor. For instance, $\beta$-sitosterol, previously reported as a potential inhibitor binds away from the active site indicating the noncompetitive and allosteric binding mode. The binding of $\beta$-sitosterol was assisted through hydrophobic interactions that contributed to its binding energy, hence presumably responsible for its inhibitory activity. In this phenomenon, the conformational changes will be observed in the active site of the enzyme and therefore slows down the enzyme activity [39–41].

The predicted binding area in myo-inositol could be capable of inhibiting protein activity via a noncompetitive mode and those compounds with similar nature. Hydrophobic and $\pi$-interactions play a crucial role in the enzyme inhibitory activity of compounds with a lack of hydroxyl groups. The control ligand’s binding site on $\alpha$-glucosidase within the active site, whereas the compounds such as quercetin and palmitic acid were observed to occupy the entrance of the pocket and not directly interacting with the residues within the active site. This suggests the existence of an allosteric binding site, which could probably cause alterations in the enzyme activity. Clef closure may be induced in this arrangement to avoid the entrance of substrate,
which then acts as noncompetitive inhibitors. In conclusion, molecular docking conducted in this study has enabled the prediction and provided supportive data for α-glucosidase inhibitory activity of the bioactive compounds.

5. Conclusion

The study showed that E60 S. zalacca fruit revealed potent α-glucosidase inhibitory activity. By GC-MS analysis, many compounds were detected in S. zalacca fruits, and the molecular docking results showed that palmitic acid and β-sitosterol were close to the active side of the enzyme. This study has provided the appropriate information on the potential antidiabetic action of S. zalacca fruits via α-glucosidase inhibitory effect and the phytochemical constituents that may be responsible for this activity. This finding can be applied for further investigations of the S. zalacca fruits’ beneficial uses and the development of the medicinal preparations for the management of diabetes.

Data Availability

The data used to support the findings of this study are included within the article and the supplementary information files. Any other data can be made available upon request.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

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Supplementary Materials

Figure S1: TIC of S. zalacca extracts. (Supplementary Materials)

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