Production of Sucrolytic Enzyme by Bacillus licheniformis by the Bioconversion of Pomelo Albedo as a Carbon Source

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Abstract: Recently, there has been increasing use of agro-byproducts in microbial fermentation to produce a variety of value-added products. In this study, among various kinds of agro-byproducts, pomelo albedo powder (PAP) was found to be the most effective carbon source for the production of sucrose hydrolyzing enzyme by Bacillus licheniformis TKU004. The optimal medium for sucrolytic enzyme production contained 2% PAP, 0.75% NH4NO3, 0.05% MgSO4, and 0.05% NaH2PO4 and the optimal culture conditions were pH 6.7, 35 °C, 150 rpm, and 24 h. Accordingly, the highest sucrolytic activity was 1.87 U/mL, 4.79-fold higher than that from standard conditions using sucrose as the carbon source. The purified sucrolytic enzyme (sleTKU004) is a 53 kDa monomeric protein and belongs to the glycoside hydrolase family 68. The optimum temperature and pH of sleTKU004 were 50 °C and pH = 6, respectively. SleTKU004 could hydrolyze sucrose, raffinose, and stachyose by attacking the glycoside linkage between glucose and fructose molecules of the sucrose unit. The Km and Vmax of sleTKU004 were 1.16 M and 5.99 μmol/min, respectively. Finally, sleTKU004 showed strong sucrose tolerance and presented the highest hydrolytic activity at the sucrose concentration of 1.2 M–1.5 M.

Keywords: agro-byproduct; Bacillus licheniformis; bioconversion; pomelo albedo; sucrolytic

1. Introduction

Sucrose (α-D-glucopyranosyl-1,2-β-D-fructofuranoside), the world’s most abundant disaccharide, comprises a glucose unit and a fructose unit linked together by an α1–β2 linkage [1]. Sucrose hydrolysis is catalyzed by sucrolytic enzymes to produce a mixture of glucose and fructose (also known as invert sugar). Sucrolytic enzymes are of various types such as invertases (sucrases) [2], levansucrases [3], dextranucrases [4], glucansucrases [5], and inulosucrases [6]. They use sucrose as the donor of fructosyl moiety to be transferred to various kinds of acceptors, such as water (sucrose hydrolysis), sucrose, and fructan [7]. Therefore, these enzymes are also responsible for the production of some compounds such as fructooligosaccharides (FOSs), inulins, and levans [8,9]. To date, sucrolytic enzymes have diverse applications in the food [10–12], beverage [13,14], medicine [1,14], cosmetic [5,15], bioenergy [16], paper [15], and biochemical sectors [15]. Thus, the use of sucrolytic enzymes is highly widespread, with a large number of industrial applications.

The worldwide market for enzymes is anticipated to reach $10.5 billion in 2024 [17]. Among these, enzymes from microbial sources are the most important group, comprising 90% of the global market [18]. Regardless of their high demand, the price of enzymes still remains relatively high, which thus increases the cost of the processes using enzymatic methods. Therefore, for the cost-effective production of enzymes, several strategies...
are being developed including using cheap and readily available carbon and nitrogen sources such as the wastes from food processing, fishery, and agriculture as the alternative substrates for the microbial fermentation process [19–22]. Besides, the utilization of by-products for enzyme production is also an environmentally friendly technique because it can reduce the amount of waste generated and released into the environment [23]. Accordingly, this green technique has received a great attention, and as a result, various kinds of enzymes have been effectively produced by using by-products as the major nutrient source such as proteases [22,24], xylanases [21,25], chitosanases [26,27], chitinases [28,29], pectinases [23,30], invertases [31,32], etc.

Bacillus licheniformis is a rod-shaped, facultatively anaerobic, Gram-positive, and spore-forming bacterium [33]. B. licheniformis is favored because of its outstanding fermentation properties, high protein yield, and completely toxin-free productivity [34]. To date, various applications of B. licheniformis have been explored for the production of probiotics [35], industrial enzymes [36,37], and biofuels [37]. B. licheniformis has been commonly used for the manufacture of amylase and protease [37–41], however, the production of sucrolytic enzymes by this genus has not been fully explored. In fact, there are only a few reports on the production of sucrose hydrolyzing enzymes by B. licheniformis, especially using low-cost materials for the fermentation process [42–44]. In comparison, several B. licheniformis strains are known to have a great capacity for processing agricultural byproducts to produce enzymes [39,45]. This leads to the idea of using agro-byproducts as economically effective nutritional supplements for the synthesis of sucrose hydrolytic enzymes by B. licheniformis.

The exploration of new enzymes, having a low-cost manufacturing method, and possessing desirable characteristics, has always been regarded as important. Accordingly, in this study, various kinds of agro-byproducts such as pomelo (Citrus maxima) albedo powder (PAP), pomelo flavedo powder (PFP), orange (Citrus sinensis) peel powder (OPP), banana (Musa paradisiaca) peel powder (BPP), rice bran powder (RBP), and wheat bran powder (WBP) were used as carbon sources for B. licheniformis TKU004 to produce sucrolytic enzyme. Besides, the medium composition and culture conditions were optimized for maximum sucrolytic enzyme production. In addition, the properties of the obtained enzyme, as well as its purification, were studied.

2. Materials and Methods

2.1. Materials

B. licheniformis TKU004 was isolated and described in an earlier report [46]. Sucrose, glucose, maltose, and fructose were bought from Katayama Chemical Industries Corporation (Osaka, Japan). 3,5-Dinitrosalicylic acid (DNS), and fructooligosaccharide (FOS) were purchased from Sigma Aldrich (Saint Louis, MO, USA). Rice bran and wheat bran were collected from Miaoli (Miaoli City, Taiwan). Orange, banana, and pomelo were bought from the local markets of Tamshui (New Taipei, Taiwan) and then peeled to prepare orange peel, banana peel, pomelo albedo, and pomelo flavedo. Finally, these materials were dried and ground to powder (particle size smaller than 0.297 mm). All other chemicals were of the highest possible quality.

2.2. Sucrolytic Assay

Sucrolytic activity was determined using the DNS method [21] with certain modifications. Briefly, a mixture of 50 µL enzyme and 150 mL sucrose (1%, w/v) was incubated at 37 °C for 60 min to allow the hydrolysis of sucrose. Later, 750 µL of DNS reagent was added to the reaction and the obtained mixture was heated at 100 °C for 10 min. Consequently, the absorbance of red color of the mixture was read at 515 nm using an ELISA reader (Bio-Rad, Hercules, CA, USA) and was used to calculate the amount of reducing sugar in the hydrolysis reaction. The amount of enzyme required to release 1 µM of reducing sugar in one minute was described as one unit of sucrolytic activity.
2.3. Agro-Byproducts as the Sole Carbon Source for Sucrolytic Enzymes Production

One percent of each carbon source (PAP, PFP, OPP, BPP, RBP, WBP, and sucrose) was added into a 250 mL flask containing 100 mL of basal medium (1% KNO₃, 0.1% KH₂PO₄, and 0.05% MgSO₄). The bacterial seed was prepared by adding a loopful of pure culture into a 250 mL flask containing 100 mL of nutrient broth and then culturing at 37 °C and 150 rpm for 24 h. The cultivation of *B. licheniformis* TKU004 on different carbon source-containing mediums was initiated by adding 1 mL of the bacterial seed solution (OD 660 nm = 1) to each medium and kept at 37 °C and 150 rpm. One milliliter of culture medium was withdrawn each 24 h and the sucrolytic activity was estimated. Different amounts of PAP (0.25 g, 0.5 g, 1.0 g, 1.5 g, 2.0 g, 2.5 g, and 3 g) were used to explore the optimal PAP concentration for sucrolytic enzyme production. Each experiment was performed in triplicate.

2.4. Effect of Other Conditions on Sucrolytic Enzymes Production

To explore the effect of other conditions on sucrolytic enzyme production by *B. licheniformis* TKU004, the one-factor-at-a-time method was used in the order of nitrogen source (KNO₃, NH₄NO₃, (NH₄)₂SO₄, casein, peptone, beef extract powder (BEP), squid pens powder (SPP), and shrimp heads powder (SHP)), NH₄NO₃ concentration (0%, 0.25%, 0.50%, 0.75%, 1%, 1.25%, 1.50%, and 2%), phosphate salts (NaH₂PO₄, Na₂HPO₄, KH₂PO₄, and K₂HPO₄), NaH₂PO₄ concentration (0%, 0.025%, 0.05%, 0.075%, and 0.1%), yeast extract powder concentration (0%, 0.05%, 0.1%, 0.15%, and 0.2%), pH (3.7, 4.7, 5.7, 6.7, 7.7, and 8.7), temperature (25 °C, 30 °C, 35 °C, 37 °C, and 40 °C), shaking speed (100 rpm, 125 rpm, 150 rpm, 175 rpm, and 200 rpm), and cultivation time (0 h, 12 h, 24 h, 36 h, 48 h, 60 h, and 72 h). The original conditions for the experiments were 2% PAP, 1% KNO₃, 0.1% KH₂PO₄, 0.05% MgSO₄, pH = 5.7, 37 °C, 100/250 mL, 150 rpm, and 24 h. At a time, only one factor was assessed while other factors were constant and the condition giving the highest sucrolytic activity was chosen for further experiments. Each experiment was performed in triplicate.

2.5. Enzyme Purification

To obtain cell-free supernatant, the culture medium of *B. licheniformis* TKU004 was centrifuged (9000 × g rpm for 30 min, 320 R, Hettich, Tuttingen, Germany) and then mixed with (NH₄)₂SO₄ (60% w/v, 4 °C, overnight). The precipitate was conveniently removed from the mixture by centrifugation (10.000 × g rpm, 30 min) and dissolved in sodium phosphate buffer (50 mM, pH = 7). The residual (NH₄)₂SO₄ in the crude enzyme solution was removed using a cellulose membrane (CelluSep T2, Intercim, Montluçon, France). The crude enzyme solution was then loaded onto a DEAE sepharose column pre-equilibrated with sodium phosphate buffer (50 mM, pH = 7). A linear gradient of NaCl (0 M–1.0 M) was applied to elute the target enzymes. Fractions exhibiting succrolytic activity were then pooled and concentrated by lyophilization. Finally, the obtained enzyme was purified by a high-performance liquid chromatography (HPLC, Hitachi Chromaster HPLC system, Hitachi, Tokyo, Japan) system consisting of a KW-802.5 column. The molecular weight (MW) of the obtained enzyme was determined using sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and matrix-assisted laser desorption ionization-time of flight mass spectrometry (MALDI-TOF MS, Bruker Daltonics, Bremen, Germany) methods, as described in early reports [22,47]. The identification of the obtained enzyme was determined by liquid chromatography-tandem mass spectrometry (LC-MS/MS, performed by Mission Biotech, Taipei, Taiwan) [48].

2.6. Effect of Temperature and pH

The optimum temperature of sleTKU004 activity was examined over a range of 30 °C to 80 °C. The thermal stability of sleTKU004 was accorded to its residual succrolytic activity after allowing the sleTKU004 solution to stand at 20 °C, 40 °C, 50 °C, and 60 °C for 0.5 h, 1 h, 2 h, and 4 h. The optimum pH and pH stability of sleTKU004 were explored at the
pH range of 3 to 10 using a system buffer (sodium carbonate buffer, pH 9–10; sodium phosphate buffer, pH 6–8; sodium acetate buffer, pH = 5; glycine HCl buffer, pH = 3–4). The pH stability was accorded to the residual sucrolytic activity after allowing the sleTKU004 solution to stand at different pH points for 1 h at 20 °C. Each experiment was performed in triplicate.

### 2.7. Effect of Various Chemicals

Various chemicals including KCl, BaCl$_2$, CaCl$_2$, CaSO$_4$, FeCl$_3$, FeCl$_2$, CuSO$_4$, ZnSO$_4$, MnSO$_4$, ethylenediaminetetraacetic acid (EDTA), Tween 20, Tween 40, Triton X-100, and sodium dodecyl sulfate (SDS) were used to examine their effect on the sucrolytic activity of sleTKU004. KCl, BaCl$_2$, CaCl$_2$, CaSO$_4$, FeCl$_3$, FeCl$_2$, CuSO$_4$, ZnSO$_4$, MnSO$_4$, and EDTA were tested at the final concentration of 1 mM, 5 mM, and 10 mM whereas Tween 20, Tween 40, Triton X-100, and SDS were assessed at 1%, 5%, and 10% concentrations. Each experiment was performed in triplicate.

### 2.8. Substrate Specificity

The specificity of the sleTKU004 enzyme was tested on substrates including sucrose, raffinose, stachyose, FOS, pectin, dialyzed pectin, dextran starch, gum arabic, and 1,3-β-glucan using the conditions of the sucrolytic activity assay (as described above). All the substrates were prepared at a concentration of 1% (w/v). Each experiment was performed in triplicate. The HPLC analysis of the sucrose, raffinose, stachyose hydrolysis was conducted under the following conditions: NH$_2$P-50 4E column; a mobile phase of 75% acetonitrile and 25% water; flowrate of 1.0 mL/min; infrared (IR) detector.

### 2.9. Effect of Sucrose Concentration

The effect of sucrose concentration on the sucrolytic activity of sleTKU004 was examined by mixing sleTKU004 solution with sucrose at different concentrations to gain the final sucrose concentration of 0.2 M to 1.5 M. Sucrolytic activity was then determined following the assay described above. Each experiment was performed in triplicate.

### 2.10. Statistical Analysis

The experiments pertaining to the culture parameters and effects of sucrose concentration on the activity of sleTKU004 were analyzed through one-way analysis of variance (ANOVA) and the post hoc Duncan’s Multiple Range Test. The experiment on the effects of various chemicals on the activity of sleTKU004 was performed using one-way ANOVA with the post hoc Tukey Honestly Significant Difference (HSD). Statistical significance was achieved when $p < 0.05$.

### 3. Results and Discussion

#### 3.1. Agro-Byproducts as the Sole Carbon Source for Sucrolytic Enzyme Production

Various carbon sources were examined for the production of sucrolytic enzymes by *B. licheniformis* TKU004. These included PAP, PFP, OPP, BPP, RBP, WBP, and sucrose. As shown in Figure 1a, the highest sucrolytic activity was observed in the culture supernatant containing PAP as the carbon source (0.81 U/mL on the first day of fermentation), followed by OPP (0.56 U/mL on the 1st day and 0.58 U/mL on the 2nd day of fermentation), RBP (0.59 U/mL on the 2nd day of fermentation), WBP (0.56 U/mL on the 2nd day of fermentation), BPP (0.46 U/mL on the 1st day and 0.46 U/mL on the 2nd day of fermentation), sucrose (0.38 U/mL on the 1st day of fermentation), and PFP (0.28 U/mL on the 1st day of fermentation). Pomelo albedo contains 72.62% carbohydrate, 16.13% moisture, 6.27% protein, 3.41% ash and 1.56% fat [49]. Huang et al., (2014) reported that the carbohydrates of pomelo peel include pectin (35.42%), cellulose (16.50%), hemicellulose (6.86%), lignin (3.16%), and soluble sugar (12.62%) [50]. The high carbohydrate ratio, especially the soluble sugar could make the pomelo albedo an attractive carbon source for microbial fermentation. Currently, bioethanol production is the primary commercial
operational process that utilizes pomelo peel waste as a raw material [51,52]. The potential of pomelo albedo in the manufacture of value-added products such as enzymes is not yet fully explored. In this study, among the tested byproducts, pomelo albedo exhibited the highest enzyme activity and was thus chosen as the most suitable carbon source for saccharolytic enzyme production by <i>B. licheniformis</i> TKU004.

![Figure 1](attachment:figure1.png)

**Figure 1.** Effect of different carbon sources (a) and amount of PAP (b) on the saccharolytic enzyme production by <i>B. licheniformis</i> TKU004. PAP, pomelo albedo powder; PFP, pomelo flavedo powder; OPP, orange peel powder; BPP, banana peel powder; RBP, rice bran powder; WBP, wheat bran powder. All data points are the mean ± standard deviation. Saccharolytic activity values with different letters (a–e) are significantly different based on Duncan’s Multiple Range Test at the level of 5%.

To examine the effect of PAP concentration on the production of saccharolytic enzymes, the amount of PAP was adjusted in a range of 0.25–3% (w/v). As shown in Figure 1B, the increase of PAP concentration from 0.25% to 2% could significantly increase the production of saccharolytic enzymes (from 0.08 U/mL to 1.09 U/mL, respectively). Besides, higher PAP concentration (2%, 2.5%, 3%) did not show any significant improvement in the production of saccharolytic enzymes (1.09 U/mL, 1.07 U/mL, and 0.99 U/mL). Due to the lower cost PAP material and the highest saccharolytic enzyme productivity, 2% PAP was chosen as the most suitable concentration for the production of saccharolytic enzymes by <i>B. licheniformis</i> TKU004. From the literature, various substances have been used as suitable carbon sources for producing saccharolytic enzymes, such as sucrose [43], glucose [53], fructose [54], orange peel [55], and pineapple crown [56].

3.2. Effect of Nitrogen Sources on Saccharolytic Enzymes Production

Pomelo albedo has a low nitrogen content and comprises only 6.27% protein [49]. Thus, supplementing the nitrogen source to a PAP-containing medium may be a necessity. Accordingly, different nitrogen sources such as KNO₃, NH₄NO₃, (NH₄)₂SO₄, casein, peptone, BEP, SPP, SSP, and SHP were supplemented to the PAP-containing medium to assess the most suitable nitrogen source for the production of saccharolytic enzymes by <i>B. licheniformis</i> TKU004. As shown in Figure 2a, a higher saccharolytic activity was observed using NH₄NO₃ (1.10 U/mL), followed by KNO₃ (0.94 U/mL), peptone (0.53 U/mL), casein (0.48 U/mL), and BEP (0.41 U/mL). The saccharolytic activity of SSP-, SPP-, and SHP-containing mediums were not significantly different from that of the control medium (0.33 U/mL, 0.31 U/mL, 0.31 U/mL, and 0.31 U/mL respectively), which only contained 2% PAP, 0.1% KH₂PO₄, and 0.05% MgSO₄. Besides, the use of 1% (NH₄)₂SO₄ could potentially inhibit the production of saccharolytic enzymes by <i>B. licheniformis</i> TKU004 (0.03 U/mL). These results indicate that NH₄NO₃ is the best nitrogen source as a supplement to PAP-containing medium for the production of saccharolytic enzymes by <i>B. licheniformis</i> TKU004. There are reports of various...
organic and inorganic nitrogen sources ideal for high sucrolytic enzyme production such as yeast extract, peptone, urea, NaNO₃, tryptone, NH₄Cl, and (NH₄)₂HPO₄ [15].

The effect of NH₄NO₃ concentration on sucrolytic enzyme production was studied by adding 0–2% NH₄NO₃ into the PAP-containing medium. As shown in Figure 2b, the sucrolytic activity of medium with NH₄NO₃ at 0%, 0.25%, 0.50%, 0.75%, 1%, 1.25%, 1.50%, 1.75%, and 2% was 0.31 U/mL, 0.79 U/mL, 0.93 U/mL, 1.08 U/mL, 1.07 U/mL, 0.76 U/mL, 0.53 U/mL, 0.03 U/mL, and 0.02 U/mL, respectively. The sucrolytic activity of the medium containing 0.75% NH₄NO₃ and 1% NH₄NO₃ were not significantly different, indicating that 0.75% NH₄NO₃ is the optimum concentration for the production of the sucrolytic enzyme by B. licheniformis TKU004.

3.3. Effect of Phosphate Salts and Yeast Extract Powder Concentration on Sucrolytic Enzyme Production

Phosphate salts also affect microbial bio-active productivity [57]. Thus, the effect of some phosphate salts such as NaH₂PO₄, Na₂HPO₄, KH₂PO₄, and K₂HPO₄ on the production of sucrolytic enzymes by B. licheniformis TKU004 was explored herein. Each of those phosphate salts was added to a medium containing 2% PAP, 0.75% NH₄NO₃, and 0.05% MgSO₄. As shown in Figure 3a, NaH₂PO₄ was screened as the most suitable form of phosphate for the optimum sucrolytic activity and was higher than that of the others (1.20 U/mL compared to 0.80 U/mL in Na₂HPO₄-containing medium, 1.08 U/mL in KH₂PO₄-containing medium, and 1.05 U/mL in K₂HPO₄-containing medium). As shown in Figure 3b, the sucrolytic activity of medium with NaH₂PO₄ at 0%, 0.025%, 0.05%, 0.075%, and 0.1% was, 0.16 U/mL, 0.95 U/mL, 1.30 U/mL, 1.27 U/mL, and 1.17 U/mL (respectively). The sucrolytic activity of the medium containing 0.05% NaH₂PO₄ and 0.075% NaH₂PO₄ were not significantly different, indicating that 0.05% NaH₂PO₄ is the optimum concentration for the production of the sucrolytic enzyme by B. licheniformis TKU004.
 Various microbial strains, for example, 

Pichia sp. [58], B. cereus TA-11 [59], B. macerans [60], Lactobacillus brevis Mm-6 [61], and Zymomonas mobilis CDBB-B 603 [53] need yeast extract to enhance the sucrolytic enzyme productivity. In this study, a range of yeast extract powder from 0% to 0.2% was added to a medium containing 2% PAP, 0.75% NH₄NO₃, 0.05% MgSO₄, and 0.05% NaH₂PO₄ to explore its effect on the production of sucrolytic enzymes from B. licheniformis TKU004. As shown in Figure 3c, yeast extract powder had a negative effect on the enzyme production and the highest sucrolytic activity was obtained from the medium without the addition of yeast extract powder (1.23 U/mL). In fact, yeast extract powder has a high ratio of proteins (48.52%), and carbohydrates (32.92%) [62]. According to the result, the supplement of yeast extract powder can provide nutrients to the bacteria without the necessity to use sucrose or sucrose-based substances, thereby drastically reducing the production of sucrolytic enzymes.

The optimum sucrolytic enzyme production by B. licheniformis TKU004 using pomelo albedo as the sole carbon source was in the medium containing 2% PAP, 0.75% NH₄NO₃, 0.05% MgSO₄, and 0.05% NaH₂PO₄. Then we explored the effect of the cultural conditions on the sucrolytic enzyme production by B. licheniformis TKU004. These included initial pH of the medium (3.7–8.7), cultivation temperature (25–40 °C), shaking speed (100 rpm, 125 rpm, 150 rpm, 175 rpm, and 200 rpm), and cultivation time (0–72 h). The sucrolytic activity produced in media of different pH (Figure 4a) were in the order of pH = 6.7 (1.29 U/mL), pH = 5.7 (1.09 U/mL), pH = 7.7 (0.89 U/mL), pH = 8.7 (0.89 U/mL), pH = 4.7 (0.23 U/mL), and pH = 3.7 (0.13 U/mL); at different temperatures (Figure 4b) were in the order of 35 °C (1.72 U/mL), 37 °C (1.12 U/mL), 40 °C (0.82 U/mL), 30 °C (0.18 U/mL), and 25 °C (0.02 U/mL); and at different shaking speeds (Figure 4c) were in the order of 150 rpm (1.60 U/mL), 125 rpm (1.09 U/mL), 175 rpm (0.97 U/mL), 200 rpm (0.16 U/mL), and 100 rpm (0.15 U/mL). The time course of cell growth and sucrolytic enzyme production of B. licheniformis TKU004 under the optimized conditions is shown in Figure 4d. The number of the cells of B. licheniformis TKU004 increased quickly at 0–24 h and then entered the stationary phase in the 24–60 h period. The sucrolytic activity of the culture medium also reached its maximum in the early stationary phase (1.87 U/mL at 24 h) and then decreased over time. Likewise, the time course for sucrolytic enzyme production by bacteria was around 12–72 h [15] for example Streptomyces sp. ALKC8 (24 h, 0.35 U/mL) [63], Brevibacterium divaricatum (12 h) [64], Lactobacillus brevis Mm-6 (72 h, 1399 U/mL) [61], and Arthrobacter sp.10137 (22.5 h, 26.69 U/mL) [65]. Another study found that the short fermentation period has more advantages than the long one [21]. This suggested that the time course for the production of sucrolytic enzyme by B. licheniformis

![Figure 3. Effect of phosphate salts (a), amount of NaH₂PO₄ (b), and amount of yeast extract powder (c) on the sucrolytic enzyme production by B. licheniformis TKU004. C, control. All data points are mean ± standard deviation. Sucrolytic activity values with the different letters (a–d) are significantly different based on Duncan’s Multiple Range Test at the level of 5%.](image-url)
TKU004 was acceptable. From the literature, the optimum medium for the production of sucrolytic enzymes may require the supplement of sugar (sucrose [43], fructose [54], and glucose [53]) and organic nitrogen (peptone [60], malt extract [61], and yeast extract [58]). However, the price of those materials may be a significant obstacle in the production of sucrolytic enzymes. Thus, to reduce the cost of the process of sucrolytic enzyme production, the use of agricultural wastes can be an effective alternative. In this study, we could successfully develop a low-cost and simple medium based on pomelo albedo and optimize the culture conditions for sucrolytic enzyme production by \textit{B. licheniformis} TKU004. The composition of the optimal medium was 2\% PAP, 0.75\% NH\textsubscript{4}NO\textsubscript{3}, 0.05\% MgSO\textsubscript{4}, and 0.05\% NaH\textsubscript{2}PO\textsubscript{4}, and the optimal culture conditions were observed to be initial medium pH of 6.7, cultivation temperature of 35 °C, shaking speed of 150 rpm, and cultivation time of 24 h. Accordingly, the highest sucrolytic activity was 1.87 U/mL, 4.79-fold higher than that in original conditions using sucrose as the carbon source.

![Figure 4](image)

**Figure 4.** Effect of initial pH (a), temperature (b), shaking speed (c), and incubation time (d) on the sucrolytic enzyme production by \textit{B. licheniformis} TKU004. All data points are the mean ± standard deviation. Sucrolytic activity values with different letters (a–f) are significantly different based on Duncan’s Multiple Range Test at the level of 5%.

### 3.5. Purification of the Produced Sucrolytic Enzyme

One-day culture supernatant (0.4 L) was used to purify the produced sucrolytic enzymes under the following condition: DEAE-Sepharose resin, 50 mM sodium phosphate (pH = 7.0), and NaCl gradient from 0 M to 1 M. The fractions showing sucrolytic activity from tube number 45 to 59 (Figure 5) were dialyzed against 50 mM sodium phosphate (pH = 7.0), and concentrated by a freeze-dryer. HPLC with size-exclusion chromatography mode was then used for the purification of the enzyme. After purification, approximately
1.34 mg of *B. licheniformis* TKU004 sucrolytic enzyme (sleTKU004) was obtained. The recovery yield of the obtained enzyme was 1.31% with 11.4-folds of the specific activity (Table 1). As shown in Table 2, the MW of 53 kDa (by SDS-PAGE method) of sleTKU004 is highly similar to sucrase from *B. stearothermophilus* NUB36 (51.519 Da) and levansucrases from *B. licheniformis* RN-01 (52 kDa), *B. licheniformis* 8-37-0-1 (51 kDa), *B. amyloliquefaciens* BH072 (55 kDa), and *B. amyloliquefaciens* KK9 (52974 Da). In general, the sucrolytic enzymes from the *Bacillus* genus have an MW range of 14–105 kDa (Table 2). The non-reducing SDS-PAGE analysis of sleTKU004 also revealed a single band at 53 kDa (Figure 6a), indicating that the enzyme is a monomer in its native form. Other reports show that the sucrose hydrolytic enzymes from the *Bacillus* genus could be monomers [66] or dimers [67]. The native form of sleTKU004 was also analyzed by MALDI-TOF MS and a peak was observed at 53,228 Da (Figure 6b).

**Table 1.** A summary of the purification of sleTKU004 from *B. licheniformis* TKU004.

| Step                           | Total Protein (mg) | Total Activity (U) | Specific Activity (U/mg) | Recovery (%) | Purification (Fold) |
|-------------------------------|--------------------|--------------------|--------------------------|--------------|---------------------|
| Cultural supernatant          | 1170.91            | 582.31             | 0.497                    | 100.00       | 1.00                |
| (NH₄)₂SO₄ precipitation        | 359.71             | 297.74             | 0.827                    | 51.13        | 5.02                |
| Ion-exchange chromatography    | 79.98              | 199.47             | 2.494                    | 34.26        | 5.02                |
| Size-exclusion chromatography | 1.34               | 7.63               | 5.670                    | 1.31         | 11.40               |

**Table 2.** A comparison of sleTKU004 with sucrolytic enzyme produced by other *Bacillus* strains.

| Enzyme/Strain | Opt. Temp. | Opt. pH | Carbon Source            | MW               | Ref    |
|---------------|------------|---------|--------------------------|------------------|--------|
| Levansucrase  | 50 °C      | 6       | PAP                      | 53 kDa           | This study |
| *B. licheniformis* TKU004 |           |         |                          |                  |        |
| β-d-fructofuranosidase | 30–60 °C   | 4–8     | wheat bran and molasses  | 66 kDa and 64,512.31 Da¹ | [68] |
| *B. subtilis* LYN12    |            |         |                          |                  |        |
| Levansucrase | 50 °C      | 6       | peptone                  | 52 kDa           | [42]   |
| *B. licheniformis* RN-01 |         |         |                          |                  |        |
| Levansucrase | 45–50 °C   | 6       | sucrose                  | 51 kDa           | [43]   |
| *B. licheniformis* 8-37-0-1 |       |         |                          |                  |        |
### Table 2. Cont.

| Enzyme/Strain                      | Opt. Temp. | Opt. pH | Carbon Source         | MW            | Ref  |
|------------------------------------|------------|---------|-----------------------|---------------|------|
| Fructan sucrase B. subtilis ZW019 | 50         | 5.6     | tryptone              | 58 kDa        | [69] |
| Sucrase and levansucrase B. subtilis Marburg |            |         |                       | 40 kDa        | [70] |
| Invertase B. subtilis HJ14         | 30–32.5 °C | 8       |                       | 58 kDa        | [71] |
| Invertase B. subtilis              |            |         | sucrose               | 73 kDa        | [72] |
| Levansucrase B. amyloliquefaciens BH072 | 40 °C     | 6       | sucrose and corn      | 55 kDa        | [73] |
| Invertase B. cereus TA-11          | 50 °C      | 7       | sucrose               | 23 kDa and 26 kDa | [59] |
| Sucrase B. stearothermophilus NUB36 | 55 °C      |         |                       | 51,519 Da 1, 105,000 Da 2 | [66] |
| α-glucosidase B. subtilis         |            |         |                       | 66 kDa        | [74] |
| levansucrase B. amyloliquefaciens KK9 |           |         |                       | 52,974 Da 3 | [75] |
| levansucrase B. subtilis NRC16     | 45 °C      | 8.2     | starch                | 14 kDa        | [76] |
| Levansucrase B. methylotrophicus SK 21.002 | 40 °C      | 6.5     | sucrose               | 60 kDa        | [77] |
| Levansucrase B. licheniformis ANT 179 | 60 °C      | 6       | sugarcane juice       | 25 kDa        | [44] |
| Levansucrase Bacillus sp.          | 60 °C      | 6       | starch                | 56 kDa        | [78] |

1, LC/MS; 2, gel filtration; 3, deduced polypeptide.

![Figure 6. SDS-PAGE (a) and MALDI-TOF MS profile (b) of the sleTKU004. M, protein markers; 1, with 2-mercaptoethanol (2-ME); 2, without 2-ME.](image-url)
3.6. Identification of sleTKU004 by LC-MS/MS Analysis

To identify sleTKU004 that appeared as a prominent 53-kDa band via SDS-PAGE, the band was excised and analyzed after tryptic digestion. The excised band from the gel was subjected to electrospray tandem mass spectrometry analysis. The fragment spectra were subjected to an NCBI non-redundant protein database search. As shown in Table 3, the spectra of sleTKU004 matched nine tryptic peptides that were identical to the levansucrase (sacB) from *B. subtilis* subsp. *subtilis* str. 168 with 30% sequence coverage. The peptide sequences indicate that sleTKU004 belongs to the family 68 glycosyl hydrolase based on the amino acid sequence similarity of the cited GH-68 enzymes from *B. subtilis*. Besides, the MW of sleTKU004 (53 kDa) was also similar to that of levansucrase from *B. subtilis* subsp. *subtilis* str. 16 (52.938 Da).

|| Matched Peptide Sequence | Identified Protein and Coverage Rate | Mass and pI | Strain |
|---|---|---|---|---|
| | 39ETYGISHITR48 | Levansucrase | 52,938 Da | B. subtilis subsp. subtilis strain 168 |
| | 63VQVFEDSSTR74 | | | |
| | 115NADDTIYMFQKVGETSIDSWK137 | | | |
| | 145DSKFDANDSLIK157 | | | |
| | 177LFYTDFSGK185 | | | |
| | 216SIFGDGKTYQNVQFIDEGNYSSGDNHTLR246 | | | |
| | 326KVMKPLASNVTDEIER343 | | | |
| | 421GNNVVITSYMTRN433 | | | |
| | 461DSILEQQILTVN473 | | | |

Superscripts (39 and 48, 63 and 74, 115 and 137, 145 and 157, 177 and 185, 216 and 246, 326 and 343, 421 and 433, 461 and 473) indicate regions in the protein sequence of the levansucrase from *B. subtilis* subsp. *subtilis* strain 168.

3.7. Effects of Temperature and pH on the Activity and Stability of sleTKU004

The optimum temperature for sucrolytic activity of sleTKU004 was examined in a temperature range of 30–80 °C. The sucrolytic activity of the enzyme increased with a rise in temperature from 30 °C to 50 °C, and then sharply decreased at higher temperatures (Figure 7a). The optimum temperature for sucrolytic activity of sleTKU004 was 50 °C. Likewise, sucrolytic enzymes from *Bacillus* genus have also been found to work best in the temperature range of 30 °C to 60 °C. To explore the thermal stability of sleTKU004, the enzyme was treated at 20 °C, 40 °C, 50 °C, and 60 °C for 0.5–4 h. The result showed that the sucrolytic activity of sleTKU004 could be retained to 100% at 20 °C in 4 h, 92% at 40 °C in 3 h, and 73% at 40 °C in 4 h (Figure 7b). At higher temperatures, the half-life of sleTKU004 was estimated to be around 2 h (at 50 °C), and 0.3 h (at 60 °C). The pH activity profile sleTKU004 is shown in Figure 7c,d. The optimum enzyme activity was observed at pH = 6 and it stable at the pH range of pH = 5–9. Likewise, many sucrolytic enzymes from other *Bacilli* expressed the highest activity at pH = 6 (Table 2).

3.8. Effect of Various Chemicals on the Activity of sleTKU004

Generally, metal ions have a great influence on protein folding and catalytic processes [58]. As shown in Figure 8, some metal salts such as KCl, BaCl₂, CaCl₂, and CaSO₄ had no significant effect or only slightly inhibited the sucrolytic activity of sleTKU004. At 1 mM, FeCl₃, FeCl₂, FeSO₄, CuSO₄, ZnSO₄, and EDTA did not affect sleTKU004, however, at higher concentrations (5 mM, and 10 mM), they could inhibit the sucrolytic activity of the enzyme. On the contrary, MgSO₄ and MnSO₄ in the concentration range of 1 mM–10 mM slightly promoted the sucrolytic activity of sleTKU004. Tween 20, Tween 40, Triton X-100, and SDS were also tested for their effect on sleTKU004 sucrolytic activity. A slight inhibiting effect was found on tween 20 (10%), tween 40 (10%), and triton X-100 (5% and 10%). At 1%, SDS did not affect sleTKU004, however, at higher concentrations (5% and 10%) it completely inhibited the enzyme activity.
3.8. Effect of Various Chemicals on the Activity of sleTKU004

Generally, metal ions have a great influence on protein folding and catalytic processes [58]. As shown in Figure 8, some metal salts such as KCl, BaCl$_2$, CaCl$_2$, and CaSO$_4$ had no significant effect or only slightly inhibited the sucrolytic activity of sleTKU004. At 1 mM, FeCl$_3$, FeCl$_2$, FeSO$_4$, CuSO$_4$, ZnSO$_4$, and EDTA did not affect sleTKU004, however, at higher concentrations (5 mM, and 10 mM), they could inhibit the sucrolytic activity of the enzyme. On the contrary, MgSO$_4$, and MnSO$_4$ in the concentration range of 1 mM–10 mM slightly promoted the sucrolytic activity of sleTKU004. Tween 20, Tween 40, Triton X-100, and SDS were also tested for their effect on sleTKU004 sucrolytic activity. A slight inhibiting effect was found on tween 20 (10%), tween 40 (10%), and triton X-100 (5% and 10%). At 1%, SDS did not affect sleTKU004, however, at higher concentrations (5% and 10%) it completely inhibited the enzyme activity.

3.9. Substrate Specificity of sleTKU004

Substrate specificity sleTKU004 is summarized in Table 4. In the case of disaccharide and oligosaccharide substrates, sleTKU004 expressed the greatest activity in the order of sucrose (100%) > raffinose (85.74%) > stachyose (73.60%) > FOS (19.15%). No activity was observed against maltose and sucralose. In the case of polysaccharide substrates, sleTKU004 seemingly showed high pectinolytic activity and expressed a relative activity of 81.38% on pectin. Surprisingly, a significant amount of residual sucrose was found in the pectin substrate and was the primary contributor to the production of reducing sugar which was then used to calculate the enzyme’s relative activity (data not shown). Thus, 2-days dialyzed pectin was used to test the hydrolysis activity of sleTKU004, and the enzyme only expressed 15.26% relative activity on this type of pectin. Likewise, the relative activity on dextran was 16.94%. These results indicate that sleTKU004 may have a slight pectinolytic, and dextranolytic activity. Besides, the activity of sleTKU004 was insignificant on starch (8.82%), gum arabic (9.96%), and 1,3-β-glucan (3.59%).

Figure 7. Effects of Temperature and pH on the activity and stability of sleTKU004. (a), optimum temperature; (b), thermal stability; (c), optimum pH; (d), pH stability. All data points are the mean ± standard deviation.

Figure 8. Effect of various chemicals on the activity of sleTKU004. All data points are the mean ± standard deviation. All data points are the mean ± standard deviation. Asterisks were used to highlight the statistical differences between control and treatment groups using the Tukey HSD Test. *, and ** indicate statistical significance at $p < 0.05$ and $p < 0.01$, respectively.
3.9. Substrate Specificity of sleTKU004

Substrate specificity sleTKU004 is summarized in Table 4. In the case of disaccharide and oligosaccharide substrates, sleTKU004 expressed the greatest activity in the order of sucrose (100%) > raffinose (85.74%) > stachyose (73.60%) > FOS (19.15%). No activity was observed against maltose and sucralose. In the case of polysaccharide substrates, sleTKU004 seemingly showed high pectinolytic activity and expressed a relative activity of 81.38% on pectin. Surprisingly, a significant amount of residual sucrose was found in the pectin substrate and was the primary contributor to the production of reducing sugar which was then used to calculate the enzyme’s relative activity (data not shown). Thus, 2-days dialyzed pectin was used to test the hydrolysis activity of sleTKU004, and the enzyme only expressed 15.26% relative activity on this type of pectin. Likewise, the relative activity on dextran was 16.94%. These results indicate that sleTKU004 may have a slight pectinolytic, and dextranolytic activity. Besides, the activity of sleTKU004 was insignificant on starch (8.82%), gum arabic (9.96%), and 1,3-β-glucan (3.59%).

Table 4. Substrate specificity of sleTKU004.

| Substrate       | Relative Activity (%) |
|-----------------|-----------------------|
| Sucrose         | 100.00 ± 4.44         |
| Raffinose       | 85.74 ± 3.45          |
| Stachyose       | 73.60 ± 5.39          |
| FOS             | 19.15 ± 2.47          |
| Starch          | 8.82 ± 1.93           |
| Pectin          | 81.38 ± 2.60          |
| Dialyzed pectin | 15.26 ± 3.28          |
| Dextran         | 16.94 ± 5.04          |
| Gum arabic      | 9.96 ± 1.65           |
| 1,3-β-glucan    | 3.59 ± 2.24           |
| Maltose         | N.A.                  |
| Sucralose       | N.A.                  |

N.A.: no activity. All data points are the mean ± standard deviation.

The mechanism of high hydrolytic activity of sleTKU004 on sucrose, raffinose, and stachyose was further analyzed by HPLC. The results showed that fructose was one of the major products of the hydrolysis, indicating that sleTKU004 strictly attacked the glycoside linkage between glucose and fructose molecules of the sucrose unit (Figure 9). The other products of sucrose, raffinose, and stachyose hydrolysis were found to be glucose, melibiose, and manninotriose.

3.10. Effect of Sucrose Concentration and Kinetic Characterization

The effect of sucrose concentration on the activity of sleTKU004 was shown in Figure 10a. With an increase in sucrose concentration from 0.15 M to 1.2 M, there was a gradual increase in the sucrolytic activity of sleTKU004 (1.98–8.39 U/mL, respectively). Higher sucrose concentrations (1.2–1.5 M) maintained the high sucrose hydrolytic activity (8.39–8.79 U/mL). It was shown that the enzymatic method is preferable in the production of high-quality invert syrups compared with the acidic method. However, high sucrose concentration, which is regarded as one of the inhibitors that block the preparation of inverted sugar [2], could be a significant obstacle for the application of the enzymatic method. For this reason, candidates producing sucrolytic enzymes, which are tolerant to high-concentration sucrose, are being explored for the production of high-quality invert sugar syrups [2]. Together, the result indicates that sleTKU004 could be a potential candidate for the inverted sugar preparation.
The effect of sucrose concentration on the activity of sleTKU004 (Figure 9). The other products of sucrose, raffinose, and stachyose hydrolysis were found to be fructose and manninitol. The mechanism of high hydrolytic activity of sleTKU004 on sucrose, raffinose, and stachyose is shown in Figure 10a. With an increase in sucrose concentration from 0.15 M to 1.2 M, there was a gradual increase in the sucrolytic activity of sleTKU004 (1.98–8.39 U/mL, respectively).

$K_m$ and $V_{max}$ of sleTKU004 were estimated to be 1.16 M and 5.99 µmol/min based on a Lineweaver-Burk plot (Figure 10b). $K_m$ value of sleTKU004 was higher than that of sucrolytic enzymes from other Bacilli such as $B.\ methylotrophicus$ SK 21.002 (117.2 mM) [77], $B.\ amyloliquefacien$ BH072 (0.71 mM) [73], $B.\ cereus$ TA-11 (370 mM) [59], and $B.\ subtilis$ (33.96 g/L) [69]. A high $K_m$ value indicates that high concentration of sucrose must be present to saturate sleTKU004. This suggests that the affinity of sleTKU004 towards sucrose is low [59].
4. Conclusions

The current study aimed to establish bioprocessing of agro-byproducts for sucrolytic enzyme production by *B. licheniformis* TKU004. A medium containing PAP as the sole carbon source was found to be the most suitable for sucrolytic enzyme production. Then, the culture condition for PAP-containing mediums was successfully optimized with a sucrolytic activity 4.79-fold higher than that from original conditions. A 53 kDa sucrolytic enzyme (sleTKU004) was isolated from the PAP culture supernatant and purified and was then determined to belong to the GH 68 family. Furthermore, the characteristics of sleTKU004 were also determined. Finally, sleTKU004 showed strong sucrose tolerance and presented the highest hydrolytic activity at sucrose concentration of 1.2–1.5 M, indicating that sleTKU004 may be a good candidate for industrial applications requiring sucrolytic enzymes.

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