Production of isomaltulose obtained by Erwinia sp. cells submitted to different treatments and immobilized in calcium alginate

Produtio de isomaltulose obtida por células de Erwinia sp. submetidas a diferentes tratamentos e imobilizadas em alginato de cálcio

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1 Introduction

Sucrose is the most widely used sweetener in food production due to its physicochemical and sensorial characteristics. However, because of its high caloric value and cariogenic properties, alternative sweeteners are being studied. Foods that promote a state of well being associated with health improvement and low risks of disease have become popular, as consumers are becoming more aware and well informed. For this reason, substitutes of sucrose in are being researched.

In the past two decades, there has been an increasing interest in the production of isomaltulose, also known as Palatinose® (BUCHHOLZ; SEIBEL, 2008). Isomaltulose (6-O-α-D-glucopyranosyl-1-6-D-fructofuranose) is a disaccharide reducer obtained by the enzymatic conversion of sucrose - the α-glucosyltransferase enzyme. Different treatments were performed for the preparation of whole cells; lysed cells; and crude enzyme extract of Erwinia sp. D12 strain immobilized in calcium alginate. The packed bed column of granules, containing Erwinia sp. cells sonicated and immobilized in calcium alginate (CSI), reached a maximum conversion of 53-59% sucrose into isomaltulose and it presented activity for 480 hours. The converted syrup was purified and the isomaltulose crystallization was performed through the lowering of temperature. The isomaltulose crystals presented purity of 96.5%.

Keywords: calcium alginate; immobilized cells; Erwinia sp.; isomaltulose.
Many isomaltulose-derived products have potential industrial applications. Items such as intermediate disaccharides, polymers such as biodegradable detergents, and surfactants for industrial use, may be obtained from isomaltulose (Lichtenthaler; Peters, 2004). Isomaltulose can also be applied to produce isomaltulose oligomers, which act as prebiotics, stimulating proliferation of intestinal microbiota bifidobacteria (Kashimura; Kimura; Itokawa, 1996). The main isomaltulose derivative is isomalt, a sugar-alcohol obtained through hydrogenation, resulting an equimolar mixture of \([6-O-(\alpha-D-glucopyranosyl)-D-sorbitol]\) and \([1-O-(\alpha-D-glucopyranosyl)-D-mannitol]\), a non-cariogenic compound with low caloric value.

Microbial conversion of sucrose into isomaltulose has attracted great commercial interest due to its complexity of chemical synthesis. Isomaltulose is currently produced in large scale through the use of stable continuous columns of immobilized cells (Wu; Birch, 2005). Alginate is one of the most commonly used supports for immobilizing whole microbial cells (Vorlop; Klein, 1983), because its use is simple and cheap; it is also a reproducible technique and moderate during the immobilization process (Hulst; Trumper, 1989). The present paper depicts the conversion of sucrose into isomaltulose by immobilized Erwinia sp. D12 cells with calcium alginate. The cells were submitted to different treatments during and after immobilization with alginate support.

2 Materials and methods

2.1 Microorganism and culture maintenance

The Erwinia sp. D12 strain is a producer of the intracellular glucosyltransferase enzyme, which is capable of converting sucrose into isomaltulose, which was used in this study. The organism was grown in slants with medium composed of 6.0% sucrose (w/v), 4.0% peptone (w/v), 0.4% meat extract (w/v) and 2.0% agar (w/v) for 15 hours at 30 °C.

2.2 Production of glucosyltransferase from Erwinia sp. D12 in 6.6 L of fermentor

The wet cell mass of the Erwinia sp. D12 strain was obtained by the fermentation of the microorganism in culture medium composed of 160 g.L\(^{-1}\) of sugarcane molasses, 20 g.L\(^{-1}\) of bacteriological peptone and 15 g.L\(^{-1}\) yeast extract Prodex Lac SD\(^{+}\) (Kawaguti; MInRich; Sato, 2006) in 6.6 L of New Brunswick Bioflo IIc fermentor (New Brunswick Scientific, Edison, NJ, USA) (Kawaguti; MInRich; Sato, 2006). After fermentation, the cell mass was recovered by centrifugation at 9,600 × g for 15 minutes at 5 °C and then washed twice in aseptic conditions, with previously sterile distilled water.

2.3 Study of the conversion of sucrose into isomaltulose in a batch process

Different methods of immobilization using calcium alginate were investigated, as well as the influence of the conversion of sucrose into isomaltulose. The tests were initially performed in batches by using Erlenmeyer flasks under agitation and, subsequently, columned in a packed bed. The cell mass of Erwinia sp. D12 was obtained from the fermentation in the optimized culture medium as described in item 2.2.

For the immobilization, from cell suspension containing 40% (w/v) wet cells of Erwinia sp. D12, in sterile distilled water, the cells were mixed with a 2% sterile solution (w/v) of sodium alginate Synth PA\(^{+}\) (Labsynth products for laboratory Ltda. Diadema, SP, Brazil) containing 0.1% (w/v) Tween\(^{+}\) 80 (Riedel-de-Haën, Seeelze, Germany) in the proportion of 1:2 (v:v). Then, the suspension was dripped for the aid of a peristaltic pump MasterFlex\(^{+}\) L/S (Cole-Parmer Instruments Co., Vernon Hills, IL, USA) in 2% (w/v) solution of CaCl\(_2\), previously sterile, to form small granules, which were kept immersed in the solution at 5 °C for 12 hours. Subsequently, the granules were washed with distilled water to remove any CaCl\(_2\) excess. All steps were performed under aseptic conditions. For immobilization, samples of 25 mL of 40% (w/v) cell mass suspension were subjected to different treatments as described below:

- Immobilized Whole Cells (CII): The suspension of the cell mass was immobilized as described above; the granules were treated with a 0.06% (v/v) glutaraldehyde solution for 20 minutes, under agitation, and then washed with sterile distilled water;
- Dried Immobilized Whole Cells (CIIIS): The suspension of cell mass was immobilized as described for CII. After that, the granules containing immobilized cells were transferred to Petri dishes covered with a paper towel and kept for 24 hours refrigerated at 5 °C, 8 hours in a laminar flow, then at room temperature for a further 16 hours, and then back to 5 °C refrigeration for drying;
- Immobilized Sonicated Cells (CSI): The suspension of the cell mass was cooled to 5 °C and subjected to treatment in a Labline Ultra-Tip sonicator for 40 seconds, with 180-200 W. After the lysis of the cell wall, the cells and the extract containing glucosyltransferase were immobilized as described above, and the granules were treated with a glutaraldehyde solution as described for CII;
- Dried Immobilized Sonicated Cells (CSISIS): The suspension of cell mass was treated and immobilized as described for CSI, and the granules were dried as described in CSII;
- Immobilized Sonicated Cells Treated with Glutaraldehyde (CSGI): The mass of the cell suspension was sonicated as described for CSI. Then, 0.06% (v/v) of glutaraldehyde was added and the suspension was agitated for 20 minutes. The suspension containing lysed cells, glucosyltransferase and glutaraldehyde was immobilized as described above;
- Dried Immobilized Sonicated Cells Treated with Glutaraldehyde, (CSGIS): The wet cell mass was treated and immobilized as described for CSGI, and the granules were dried as described in CIIIS;
- Crudely Immobilized Enzyme Extract (EEI): The suspension of cell mass was cooled to 5 °C and subjected to treatment in a sonicator for 40 seconds, with
180-200 W. After the lysis of the cell wall, the suspension was centrifuged at 7,800 × g at 5 °C for 15 minutes. Subsequently, the crude enzyme extract was stopped and the granules were treated with a 0.06% glutaraldehyde solution (v/v), and then washed as described above;

- Dried Immobilized Crude Enzyme Extract (EEL): The suspension of cell mass was treated and the crude enzyme extract was immobilized as described for EEI, and the granules were dried as described in CIIS;

- Immobilized Crude Enzyme Extract Treated with Glutaraldehyde (EEGI): The suspension of cell mass was treated as described for EEI. Then 0.06% (v/v) glutaraldehyde was added to the crude enzyme extract and the solution was agitated for 20 minutes. The crude enzyme extract containing glutaraldehyde was then immobilized as previously described; and

- Dried Immobilized Crude Enzyme Extract Treated with Glutaraldehyde (EEGS): The crude enzyme extract was obtained, treated with glutaraldehyde, immobilized as described for EEGI, and dried as described in CIIS.

Samples of 12.5 g of dry or moist granules obtained from 12.5 g of wet granules were transferred to 250 mL Erlenmeyer flasks containing a solution of 100 mL with 35% (w/v) sucrose and then incubated on a rotary shaker at 150 rpm at 30 °C. Samples of 100 mL of sugar solution were removed five times after 24 hours and they were replaced by new samples of sucrose solution 35% to complete 120 hours of granule use. The conversion of sucrose into isomaltulose was analyzed as described below:

- Carbohydrate Analysis by Anion-Exchange Liquid Chromatography with a High-Efficiency for Pulse Amperometric Detection (HPAEC-PAD): The carbohydrate analysis was performed in a Dionex DX-600 (Dionex Corporation, 1228 Titan Way Sunnyvale, CA, USA) chromatograph equipped with IP25 isotropic pump and gold electrochemical detector ED50. The separation of sugars was performed using a column CarboPac™ PA 1 (4 × 250 mm), a guard column of CarboPac™ PA 1 (4 × 50 mm) and sodium hydroxide solution 250 mM as a mobile phase with a flow of 1 mL/minute at 20 °C. The carbohydrates were analyzed by retention time compared to standards of fructose, glucose, sucrose and isomaltulose (Sigma Ultra®, Sigma Chemical Co., St. Louis, MO, USA).

2.4 Production of isomaltulose using granules containing crude enzyme extract (EEI) and sonicated cells (CSI) from Erwinia sp. D12, immobilized in calcium alginate in packed bed columns

The continuous conversion of sucrose into isomaltulose using crude enzyme extract containing granules (EEI) and sonicated cells (CSI) from Erwinia sp. immobilized in calcium alginate in packed bed columns was tested.

The granules were prepared as described in Section 2.3, washed with sterile distilled water and packed into heat proof columns. A 35% (w/v) sucrose solution of P.A. Synth™ (Labsynth Laboratory Products Ltda. Diadema, Brazil) was circulated upwards continuously, with flow of 0.56 mL/minute. The column temperature was maintained at 30 °C and the conversion of sucrose into isomaltulose was analyzed, as described above, to determine the lifetime and stability of the column.

2.5 Isomaltulose obtainment by crystallization

The steps of clarification, purification and crystallization of isomaltulose were performed according to Moraes et al. (2005), with modifications.

- Syrup Clarification: the syrup containing high levels of isomaltulose was clarified with the aid of Romicon Ultrafiltration System® comprising a centrifugal pump of 20 gpm (76 L/minute) and an ultrafiltration column containing PM 50 (50,000 Da) membrane;

- Syrup Purification: the syrup clarified by ultrafiltration was purified through chromatography cation exchange, on Dowex 88 column (4 × 30 cm); and anionic exchange, on Dowex MSA1 column (4 × 30 cm), at room temperature. Subsequently, the deionized syrup was treated with activated charcoal (LF310) at a concentration of 0.1% (w/v). The syrup was in contact with coal for 30 minutes at 45-50 °C. Then, the syrup was vacuum pre-filtered using a Celite 508; finally, the resulting product was filtered in a 0.45 µm membrane; and

- Crystallization: The purified syrup was concentrated to about 69% (w/v) in a rotary evaporator (Rotavapor RE 120) and the crystallization was performed by slowly lowering the temperature. The syrup was incubated in a water bath, initially at 50 °C, and subjected to continuous agitation. The water bath was programmed to lower 5 °C/hour between 50 °C and 35 °C. The syrup was seeded with isomaltulose crystals to aid crystallization. The crystals formed were collected by continuous centrifugation in a bench centrifuge, equipped with a nylon basket. The drying of the isomaltulose crystals was performed in a vacuum oven.

3 Results and discussion

3.1 Study of the conversion of sucrose into isomaltulose in a batch process

The study of the conversion of sucrose into isomaltulose using the granules containing whole cells, lyed cells and crude enzyme extract of glucosyltransferase from Erwinia sp. D12 immobilized in calcium alginate, submitted to different treatments in the batch process, was conducted in accordance with item 2.3. Figure 1 shows the wet and dry cells granules and crude extract of glucosyltransferase from Erwinia sp. D12 immobilized in calcium alginate, which were tested for the conversion of sucrose into isomaltulose.

Figure 2 shows the conversion of sucrose into isomaltulose using wet or dry granules containing whole cells, and crude extracted sonicated cells of glucosyltransferase, immobilized in calcium alginate, with and without glutaraldehyde treatment, in
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conversion of sucrose into isomaltulose, were tested and then put in packed bed columns.

3.2 Production of isomaltulose using granules containing crude enzyme extract (EEI) and sonicated cells (CSI) from Erwinia sp. D12, immobilized in calcium alginate, in packed bed columns

Figure 3 shows that, in the first three days, there was a higher conversion using EEI, which was obtained at 59.6, 62.9 and 62.1% of isomaltulose. However, after the third day, there was a gradual decrease of the sucrose conversion into isomaltulose. Moreover, after nineteen days of continuous use, the granules of immobilized crude enzyme extract (EEI) showed no further activity.

In the first three days the CSI column showed a conversion of 58.4, 57.8 and 56.9% of sucrose into isomaltulose. However, after the third day, there was a gradual decrease of the sucrose conversion into isomaltulose. Moreover, after nineteen days of continuous use, the granules of immobilized crude enzyme extract (EEI) showed no further activity.
Isomaltulose-CII: % of Isomaltulose obtained from immobilized whole cells; Isomaltulose-CSI: % of Isomaltulose obtained from sonicated immobilized whole cells; Isomaltulose-CSGIS: % of Isomaltulose obtained from sonicated cells treated with glutaraldehyde and immobilized; Isomaltulose-CSGIS: % of Isomaltulose obtained from sonicated cells treated with glutaraldehyde and immobilized; Isomaltulose-EEI: % of Isomaltulose obtained from crude enzyme extract; Isomaltulose-EEI: % of Isomaltulose obtained from immobilized enzyme extract; Isomaltulose-CSGI: % of Isomaltulose obtained from sonicated, immobilized and dry cells; Isomaltulose-CSGI: % of isomaltulose obtained from the Immobilized and Dry Cells; Isomaltulose-CSGIS: % of isomaltulose obtained from sonicated, immobilized and dry cells; Isomaltulose-CSGIS: % of isomaltulose obtained from sonicated dry cells; Isomaltulose-CSGIS: % of isomaltulose obtained from the sonicated immobilized cells in calcium alginate; Isomaltulose-EEI: % of isomaltulose obtained from the immobilized crude enzyme extract in calcium alginate).

Figure 2. Conversion of sucrose into isomaltulose using wet or dry granules, containing whole cells, and cells treated with glucosyltransferase crude extract, immobilized in calcium alginate, with and without treatment with glutaraldehyde, in the batch process. [□] Isomaltulose-CII: % of isomaltulose obtained from immobilized whole cells; [■] Isomaltulose-CSI: % of isomaltulose obtained from the Immobilized and Dry Cells; [▲] Isomaltulose-CSGI: % of isomaltulose obtained from sonicated Immobilized Cells; [■] Isomaltulose-CSGIS: % of isomaltulose obtained from sonicated immobilized cells; [■] Isomaltulose-EEI: % of isomaltulose obtained from the Immobilized enzyme extract; [■] Isomaltulose-CSGIS: % of isomaltulose obtained from sonicated immobilized cells in calcium alginate, such as cells of Erwinia rhapontici (CHEETHAM; GARRETT; CLARK, 1985) and Erwinia rhapontici BN68689 (AHN et al., 2003). However, to compare the results with reported literary data, the methodology for analysis of carbohydrates and the conditions for conversion of sucrose into isomaltulose should be considered.

Cheetham, Garret and Clark (1985) reported that cells of Erwinia rhapontici NC PPB 1578, immobilized in calcium alginate granules, showed conversion of approximately 90% of sucrose into isomaltulose. The carbohydrates were analyzed by the liquid chromatograph Waters Prep LC/system 500A with a refractive index detector and a column PrepPak 500/C18. The high amount of isomaltulose could be due not to the chromatographic separation of isomaltulose and trehalulose, because the old equipment did not have the ability to separate as the new ion exchange liquid chromatography does. In this case, the two isomers may have been regarded as isomaltulose. In this study, the detention cell was made up by mixing the cell suspension to a 20% solution of 5% sodium alginate. The half-life of immobilized cells in the 200 × 15 cm column was approximately 8,600 hours in continuous process at 30 °C, using 1.6 M sucrose solution adjusted to a pH of 7.0.

Tsuyuki et al. (1992) reported the detention of a suspension of 40% of plant cells of Klebsiella 10 MX 5% solution of alginate to convert sucrose into isomaltulose. The granules were treated with 2% solution of polyethyleneimine for 5 minutes and, later, with 0.5% glutaraldehyde solution for 20 minutes. The immobilized cells, packed into columns (50 × 180 mm), fully converted the 25% sucrose solution and obtained 65.4% of isomaltulose and 29.7% of trehalulose. Ahn et al. (2003) used the strain Erwinia rhapontici BN68689 to convert sucrose into isomaltulose. The detention was made from 40% cell suspension (w/v), mixed with equal volume of 5% sodium alginate solution. The granules were packed into columns (180 × 50 mm and capacity of 300 mL) at 30 °C and fed with sucrose solution of 55% (w/v) pH 7.0. Temperatures were tested at 27 °C, 30 °C and 33 °C, and the initial profits of isomaltulose were respectively 79, 89 and 85%, and then decreased to 67, 79 and 57% after 60 days, respectively. The carbohydrates were analyzed by a Waters 410 liquid chromatograph with a detection refractive index.

Protaminobacter rubrum cells immobilized in calcium alginate have been used to obtain isomaltulose (HASHIMOTO; YAMADA; OSHIMURA, 1987). The maximum immobilized cell activity, at a pH of 5.5, was observed after 3 hours and it was proportional to the amount of cells, yielding conversions of 18, 30 and 44% for 10, 20 and 40 mg of cells mL⁻¹ of solution, respectively.

Moraes et al. (2005) obtained the maximum conversion of sucrose into 56.4% isomaltulose when a 19.4% sucrose solution was circulated through packed bed columns (25 × 200 mm)
containing immobilized cells of Erwinia sp. D12 (20% cell suspension in alginate solution of 1%) at a substrate flow of 21.88 mL/hour and temperature of 35 °C. Mudra, Desai and Lele (2007) studied the effect of various immobilization parameters on converting sucrose into isomaltulose by immobilized cells of Erwinia rhapontici NCPPB 1578. Using a 30% sucrose solution, the maximum production of 140 mg.mL⁻¹ was obtained in a batch process when cells were suspended at 5 g.L⁻¹ and immobilized in 5% sodium alginate.

News matrix and others methodologies of immobilization have been used to convert sucrose into isomaltulose in continuous process. Serratia plymuthica cells were immobilized in chitosan (KRASTANOV et al., 2006). The effect of the substrate sucrose concentration, the temperature and the residence time of the substrate solution on the specific volumetric productivity of the biocatalyst in a tubular fixed bed reactor was studied. A residence time of 1.7-3.0 hours was necessary for the achievement of 98-100% conversion of sucrose at substrate concentration of 30-50%. A longer residence time (4-5 hours) was needed for reaching the same conversion rate of concentrated sucrose solutions (60-70%). Krastanov, Blazheva and Stanche (2007) used immobilized cells of Serratia plymuthica in hollow fiber bioreactor to produce isomaltulose from sucrose solution. The specific productivity of the membrane reactor was 16.8 g m⁻²/hours using a flow rate of 1.3 cm³/minutes and 40% substrate concentration in continuous mode of action. The activity of the biocatalyst decreased with the increase of the operation time.

In this study, the continuous conversion of 35% solution of sucrose into isomaltulose in a packed bed column, using cells of Erwinia sp. lysed by ultrasonication and immobilized in calcium alginate (CSI), obtained 50-60% of isomaltulose for nine days.

3.3 Syrup crystallization

The steps of clarification, purification and crystallization of the isomaltulose syrup were performed as described in item 2.5. The sugar crystals were analyzed and showed 96.5% of isomaltulose. The yield of isomaltulose was obtained only through a process of crystallization and it can be obtained from a pure product if recrystallized. Cheetham (1987) obtained crystals containing 99% of isomaltulose after two crystallization processes.

4 Conclusions

The conversion of sucrose into isomaltulose was performed in a batch process using granules containing immobilized crude enzyme extract (EEI), which presented a conversion rate between 59.7-63.3%; and granules containing immobilized cells lysed by ultrasonication (CSI), which showed a conversion rate between 47.6-62.6%. The packed bed column containing pellets of cells of Erwinia sp. sonicated and immobilized in calcium alginate (CSI), showed higher stability obtained of 53-59% isomaltulose after seven days. More studies should be performed to obtain greater stability of the immobilized glucosyltransferase in the form of whole cells, lysed cells or enzyme extract. The crystals of isomaltulose after syrup clarification and purification converted to 96.5% purity.

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References


