A HIGHLY SELECTIVE AND SENSITIVE ANALYTICAL TECHNIQUE FOR THE DETERMINATION OF ISOMALTULOSE IN PRESENCE OF ITS PROCESS RELATED IMPURITIES BY CAPILLARY ELECTROPHORESIS

Sri Rama Krishna Surapureddi1,2, Kunta Ravindranath1,*, Krishnna Darnasi2 and Suhashini Ramen2

1Department of Chemistry, Koneru Lakshmaiah Education Foundation, Green Fields, Vaddeswaram, Guntur District, Andhra Pradesh, India - 522 502
2Vimta Labs Ltd, Genome Valley, Hyderabad, Telangana, India-500085.

*E-mail: ravindranath.kunta@gmail.com

ABSTRACT
Isomaltulose is a rare disaccharide and has drawn the attention of the pharmaceutical and food industry due to its medical applications. In the present investigation, a simple first of its kind capillary electrophoresis (CE) method is developed and validated for the identification and quantification of isomaltulose and its process impurities (trehalulose, sucrose, d-glucose and d-fructose). The analysis is performed at pH: 12.6 using an electrolyte buffer containing 36 mM of Na₂HPO₄ and 130 mM NaOH. The calibration curves are plotted over a concentration range from 0.25 mM to 3.0 mM with the regression of 0.99 and with detection limits of 0.15 mM, 0.14 mM, 0.13 mM, 0.10 mM and 0.23 mM for isomaltulose, trehalulose, sucrose, d-glucose and d-fructose respectively. Concerning the internal standard, d-trehalose, the relative migration time is 1.32 min for isomaltulose. Better resolution is achieved under optimum conditions of 18 °C temperature, 16 kV capillary voltage and pH of 12.6. The method is found to be specific for the intended purpose and can be used as an orthogonal approach to the current existing United States Pharmacopoeia (USP) High-performance liquid chromatography (HPLC) monograph method.

Keywords: Isomaltulose /Palatinose, Capillary Electrophoresis Analysis, Disaccharide.

INTRODUCTION
Carbohydrates are the key macronutrients and constitute nearly half of the calories in a balanced diet and among which, sugars are the major constituents. Various food and health safety agencies identified the role of the added sugars and their impact on health issues like type 2 diabetes (T2D), obesity and developing dental caries. The World Health Organization (WHO) and the Dietary Guidelines Advisory Committee (DGAC) have set up an upper limit for free sugars as lesser than 10% of the total energy intake. However, in recent times, the Scientific Advisory Committee on Nutrition (SACN) has recommended to further reduce the free sugars to less than 5% of the total energy intake. Many studies have confirmed an alliance between the intake of sugars and dental caries. Rare disaccharides are widely used to avoid the ill effects of traditional sugars as they have low calories in comparison with regular sugars. These are non-carcinogenic and are found in small amounts in nature. Remarkably, many of them have great economic value in nutrition, cosmetics and pharmaceutical industries. They are also considered as potential raw materials in food processing and beverage production, and also in research areas. At present, the production of a rare disaccharide is through synthetics (chemical) routes and these methods are not economical due to expensive raw materials. In contrast, the productions of rare disaccharide through enzymatic methods are cost-effective. One of the rare disaccharides is isomaltulose, which is also called as palatinose. In 2008, the United States Food and Drug Administration (US FDA) approved the isomaltulose and registered under Generally Recognized as Safe (GRAS). When isomaltulose is consumed, it is hydrolyzed and absorbed in the form of glucose and fructose in the small
There are no side effects observed on human health when referred to the biological data, toxicological and metabolic studies of isomaltulose. This rare disaccharide is in use as a substitute for sugar in food since 1985 in Japan, 2005 in European Union (EU), 2006 in the United States (US), 2007 in Australia and New Zealand. Isomaltulose and trehalulose are two structural isomers of sucrose, present in small quantities in honey and sugar cane. The non-carcinogenic properties of isomaltulose and trehalulose have made them a frontline alternative to sucrose as a sugar substitute in food. As compared to sucrose, isomaltulose has a negligible impact on the glucose concentration in the blood, which is beneficial for both diabetic and non-diabetic people. The glycemic index (GI) of isomaltulose is 32 whereas that of sucrose is 72. As these rare disaccharides are present in very fewer amounts, there is a need to produce them. At present, the production of rare disaccharides is through synthetic (chemical) routes and these methods are not economical due to expensive raw materials. In contrast, the productions of rare disaccharides through enzymatic methods are cost-effective. Various microorganisms can produce rare disaccharides. Microorganisms like *Serratia plymuthica*, *Erwinia rhapontici*, *Pseudomonas mesoacidiphila* MX-45, *Agrobacterium radiobacter* MX-232 and *Klebsiella planticola* CCCB 19112. *A. radiobacter* MX-232 and *Pseudomonas mesoacidiphila* MX-45 produce more trehalulose than isomaltulose. Similarly, Isomaltulose is produced more by *S. plymuthica* NCIB 8285, *E. rhapontici* NCBB 1579, *K. planticola* CCCB 19112 and *Pseudomonas rubrum*. Current advances in molecular biology paved ways to use recombinant enzymes that can be used for the production of isomaltulose using sucrose as a substrate.

Analysis of carbohydrates is perplexing due to its complex structure, heterogeneity, lack of chromophore and highly polar nature. Existing analytical methods like HPLC, gas chromatography (GC), etc. have been used to quantify sugars after pre-column derivatization. Though these methods offer high resolution and are highly sensitive, they are time-consuming and the reagents are not so economical. Additionally, derivatization based analyses of sugars by HPLC or GC have certain limitations such as excess consumption of reagent. On the other hand, High-Performance Anion Exchange Chromatography coupled with Pulsed Amperometric Detection (HPAEC-PAD) is found to be a useful technique for sugar analysis. However, according to Oshima H et al 2006, HPAEC-PAD analysis is not sensitive enough due to the partial conversion of d-fructose to d-psicose. Despite the availability of different analytical methods for identification and quantification of rare disaccharides, an alternative approach, using capillary electrophoresis with many merits is currently being evaluated. It has many advantages like less buffer consumption and high throughput. Additionally, sample dilutions are simple and minimal sample quantity and reagents are consumed during the analysis. More importantly, good separation with a high degree of resolution is achieved in a wide range of pH, which is attributed to the repulsions of charged species or ions in an electric field. Recently capillary zone electrophoresis (CZE) methods are introduced to quantify the neutral sugars through indirect and direct UV detection methods with various background electrolytes (BGEs). Stella Rovio and co-workers validated the methods, where the saccharides can be analyzed using direct UV detection at 270 nm and high pH background. The indirect UV detection of CE has a drawback of sample recovery. Recently, some studies described that the analysis through direct UV detection is based on the typical alkaline degradation of sugars conjugated with enol carbonyl group that produces UV absorption at 265 ± 5nm. An attempt has been made earlier with capillary electrophoresis for the separation of process-related impurities of isomaltulose, in which trehalulose is co-migrated with d-fructose. The present investigation endeavors to develop a highly sensitive and selective analytical technique for identification and quantification of isomaltulose in presence of its process-related impurities, by capillary electrophoresis with a good and improved resolution between these rare disaccharides.

**EXPERIMENTAL**

**Chemicals**

D-glucose, d-fructose, sucrose, trehalulose, isomaltulose, d-trehalose, disodium hydrogen phosphate (Na₂HPO₄), and sodium hydroxide (NaOH) were procured from Sigma-Aldrich (India)/BOC Sciences (US). pH calibration buffers 1.68 and 12.00 were procured from Reagecon and other buffers of pH 4.00, 7.00 and 9.20 were procured from Merck. Acids (Hydrochloric acid and Acetic acid) were also procured.
from Merck. HPLC grade water and analytical grade chemicals were used in the present analysis. 25 mM concentration stock solutions of glucose, fructose, sucrose, isomaltulose and trehalulose were prepared. An effective length of 50.2 cm (60 cm actual length) fused-silica capillaries and capillary electrophoresis apparatus were purchased from Sciex, USA. Commercial samples were collected from the local market with three different suppliers.

**Background Electrolyte**

An alkaline running buffer (electrolyte) solution containing 36 mM Na$_2$HPO$_4$.2H$_2$O and 130 mM NaOH was prepared by mixing 1.0 M sodium hydroxide (NaOH) solution with 450 mM stock solution of disodium hydrogen phosphate dihydrate. The pH of the electrolyte solution was measured with a Mettler Toledo (GMBH- model#8603) pH meter. Before measuring the electrolyte pH, the instrument was calibrated with standard buffers of pH 1.68 (±0.01), 4.00 (±0.01), 7.00 (±0.01), 9.20 (±0.01), and 12.00 (±0.01). The pH of the electrolyte solution was adjusted to 12.6 (±0.01) and ionic strength to 0.217 M$^2$. Standard stock solutions of 25 mM of each sugar, glucose, fructose, sucrose, isomaltulose and trehalulose and working standard solutions in the concentration range of 0.25, 0.5, 1.0, 2.0 and 3.0 mM were prepared with appropriate dilutions using water (HPLC grade). The internal standard: trehalose (1.0 mM) was employed to spike the working standards and commercial samples. The standard stock solutions were stored at 4 to 8 °C. As suggested by Stojakovic et al 2013, freshly prepared electrolyte solution was used to reduce the carbon dioxide influence on background electrolyte for better reproducibility$^{26}$.

**Instrumentation and Electrophoretic Procedures**

All experiments were carried out employing a computer loaded with a 32 karat software-controlled capillary electrophoresis system (CE PA800 plus, Beckman Coulter Inc.). An effective length of 50.2 cm (60 cm actual length) fused-silica capillary with a 50 µm inner diameter was employed for the investigation. A background electrolyte (BGE) consists of 36 mM Na$_2$HPO$_4$ and 130 mM NaOH was adjusted to a 12.6 pH with 1.0 M HCl. 10% (v/v) acetic acid was prepared in water. All sugars were effectively separated using a BGE and at a steady voltage of 16 kV and a cartridge temperature of 18 °C. New capillaries were pretreated by flushing with 1 M NaOH for 5 min before use, then treated with background electrolyte for 5 mins, again flushed with 1 M NaOH for 5 min and finally flushed with water for 5 min. Capillaries were preconditioned with background electrolyte for 2 min before injection. As suggested by Rovio et al 2007, with the pressure of 0.5 psi, the samples were injected for 5 seconds (approximately 5.9 nL) using hydrodynamic injection mode into a capillary at 18 °C temperature followed by background electrolyte inoculation with 0.5 psi for 5 seconds$^{24,27}$. The separation was carried out at 16 kV for 25 minutes with background electrolyte. The photodiode array (PDA) detector set at 270 nm was used for detection. After each sample run, the capillary was washed by flushing with 10% CH$_3$COOH for 4 minutes, water for 3 minutes and background electrolyte for 3 minutes and then the separation was carried out with background electrolyte for 1 minute. Rinsing was performed at a constant pressure of 20 psi.

**RESULTS AND DISCUSSION**

**Optimization of the Separation Technique**

The effect of different voltages and temperatures was investigated as part of optimization. It was observed that alkaline BGE buffer comprising of 36 mM Na$_2$HPO$_4$ and 130 mM NaOH at pH 12.6 was effective and a good separation was achieved when the voltage across the capillary was increased linearly from 0 to 16kV over two minutes and later maintained at a constant voltage of 16kV with a capillary temperature of 18 °C. Obtained results are presented in Figs.-1 and 2. Figure-1 is the 3D spectrum of all the sugars, while Fig.-2 is a two-dimensional electrophorogram of a mixture of mono and disaccharides.

**Optimization of Voltage**

The voltage applied across the CE also has a profound effect on the peaks resolutions and also on the migration times. Generally, voltage is inversely proportional to the migration time. Hence, the optimum voltage needed for good resolution with less migration time is a critical parameter that has to be assessed.
In this investigation, the influence of voltage on the separation of sugar mixtures was assessed by performing experiments with varying voltages from 15.0 to 18.0 kV across the capillaries using the same electrolyte solution (130mM NaOH and 36mM Na$_2$HPO$_4$) and keeping the pH constant at 12.6 with cartridge temperature 24°C. The findings are presented in Fig.-3.

From the obtained data it is evident that with an increase in voltage, the migration time of sugars is reducing, which is as expected. At a voltage of 15 kV, except trehalulose and d-glucose, all the other sugars were well resolved. But with the increase in voltage to 16 kV, the resolution between trehalulose and d-glucose is progressive and a valley between the peaks of trehalulose and d-glucose is getting deepened. With a further increase in the voltage to 17 kV and 18 kV, the peaks of trehalulose and d-glucose were merged and co-migrated. So from the experiments that were performed with different voltages, it can be inferred that 16 kV is optimum, where better resolution is observed. This voltage is adopted to further investigate the effect of temperature on the resolution.

**Effect of Cartridge Temperature**

To improve the resolution between trehalulose and d-glucose peaks, experiments were performed at 16, 18, 20 and 22 °C (±1 °C) by maintaining pH of the background electrolyte at 12.6 and the optimized voltage of 16 kV. The results are presented in Fig.-4. From the obtained data it can be inferred that with a
decrease in temperature from 22°C to 18°C, the resolution between trehalulose and d-glucose is progressively increased and at 18°C, the resolution of all sugars is good. But when the temperature is further decreased to 16°C, isomaltulose and trehalulose peaks were getting merged. Hence, 18°C is considered as optimum temperature, where a considerable resolution was achieved, which is greater than 1.2 (Table-1).

Thus, the optimum separation conditions for the satisfactory separation of the sugars taken for the current study were found to be: applied voltage: 16kV, injection pressure 0.5 psi for 5 seconds, capillary: 50/66 cms, separation temperature 18°C, electrolyte solution: 130mM NaOH 36 mM and Na₂HPO₄·2H₂O and pH: 12.6.
Validation
To further strengthen the suitability of the method developed for the intended purpose, the method was validated with all the critical parameters that include linearity, reproducibility (intraday and interday), the limit of detection and specificity. The BGE solution consists of 36 mM of Na₂HPO₄ and 130 mM NaOH, with optimized conditions, viz. pH 12.6, cartridge temperature 18 °C and voltage of 16 kV were used for all the validation experiments. By analyzing all the sugars, specificity was determined as individual runs and independently post spiking of all sugars with the internal standard (a known quantity). Isomaltulose has shown a migration time of 15.99 minutes with a resolution (USP) of 16.19 concerning substrate (sucrose). Further, the resolution between all other sugars and Isomaltulose is found to be satisfactory. The results are presented in Table-1.

<table>
<thead>
<tr>
<th>Table-1: Migration Time, Relative Migration Time, Resolution and Asymmetry of Sugars</th>
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<tbody>
<tr>
<td>Name of the Sugars</td>
</tr>
<tr>
<td>---------------------</td>
</tr>
<tr>
<td>d-Trehalose (IS)</td>
</tr>
<tr>
<td>Sucrose</td>
</tr>
<tr>
<td>Isomaltulose</td>
</tr>
<tr>
<td>Trehalulose</td>
</tr>
<tr>
<td>d-Glucose</td>
</tr>
<tr>
<td>d-Fructose</td>
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</tbody>
</table>

Linearity
To find out the range of the method, a mixture of five sugars at known concentrations were prepared. The concentration of each sugar in a mixture was adjusted to 0.25, 0.5, 1.0, 2.0 and 3.0 mM and analyzed in the optimized experimental conditions. The coefficient of correlation was derived by plotting a graph between concentration versus response. The results of regression analyses are presented in Table-2 and Fig.-5. The correlation coefficient for Isomaltulose was found to be 0.998.

<table>
<thead>
<tr>
<th>Table-2: Calibration Ranges, Correlation Coefficient, Precision (Intra and Inter Day) and Limit of Detection.</th>
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<tbody>
<tr>
<td>Name of the Sugars</td>
</tr>
<tr>
<td>---------------------</td>
</tr>
<tr>
<td>Sucrose</td>
</tr>
<tr>
<td>Isomaltulose</td>
</tr>
<tr>
<td>Trehalulose</td>
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<tr>
<td>Glucose</td>
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<td>Fructose</td>
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Preparation and Assay of Standard and Commercial Samples

Standard Samples
To evaluate the optimized method, experiments were performed using Isomaltulose standard that was purchased from Sigma Aldrich. Five different preparations with a concentration of 2.0 mM were prepared using HPLC grade water. Each standard solution was injected (spiked with internal standard) and analyzed. The average peak response ratios were calculated.

Commercial Sample
To further evaluate the applicability of this optimized method for samples from multiple sources purchased three samples of different makes through the amazon market with the trade name of palatinose.
from 3 different vendors. The sample solutions were prepared with a 2 mM concentration of sugar (n=3) and these samples were spiked with internal standard individually for the analysis. Each sample solution was injected in triplicates and calculated the average peak response ratio using the equations:

Corrected Peak Area = \frac{Area\ of\ Standard/\ Sample}{Migration\ Time\ of\ Standard/\ Sample}

Peak response ratio = \frac{Corrected\ Peak\ Area\ of\ Standard/\ Sample}{Corrected\ Peak\ Area\ of\ Internal\ Standard}

% Assay = \frac{Peak\ Response\ ratio\ of\ Sample}{Peak\ Response\ ratio\ of\ Standard} \times 100

The results obtained from commercial samples are presented in Table-3. It can be inferred from the Table that the observed content of isomaltulose in commercial samples concerning the standard is in agreement with the label assert and GRAS requirements of USFDA. Thus we claim that the present method is highly successful in determining the content of isomaltulose in sugar mixtures.

Table-3: % Assay for Isomaltulose in Commercial Samples

<table>
<thead>
<tr>
<th>Name</th>
<th>Average Peak Response Ratio of Isomaltulose</th>
<th>% Assay</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average Peak Response Ratio of Isomaltulose Standard = 0.989</td>
<td>99.80</td>
<td></td>
</tr>
<tr>
<td>Isomaltulose Vendor – 1</td>
<td>0.987</td>
<td>99.80</td>
</tr>
<tr>
<td>Isomaltulose Vendor – 2</td>
<td>0.982</td>
<td>99.29</td>
</tr>
<tr>
<td>Isomaltulose Vendor – 3</td>
<td>0.986</td>
<td>99.70</td>
</tr>
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</table>

**CONCLUSION**

The current study has shown the development of an efficient method for identification and quantification of isomaltulose along with other related impurities and successfully analyzed the commercially available isomaltulose as well. The method is further validated at 0.25 mM to 3.0 mM conc. series, for all the related sugars. Better resolution is achieved under optimum conditions of a temperature of 18 °C, the capillary voltage of 16 kV and a pH of 12.6. This method is applicable for the identification and quantification of enzymatically converted isomaltulose from sucrose. In addition to this, better resolution (R=16) can be obtained between the substrate (sucrose) and product (isomaltulose), even in the presence of other sugars like glucose, fructose and trehalulose. Commercially available isomaltulose samples were
analyzed using the current method and observed a purity of >99%. These obtained findings are in agreement with the label claim and GRAS requirements of USFDA.

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