Conversion of Sucrose into Laminaribiose Using Sucrose Phosphorylase, Xylose Isomerase and Laminaribiose Phosphorylase

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Laminaribiose was synthesized from sucrose by coupling reactions of sucrose phosphorylase, xylose isomerase and laminaribiose phosphorylase in the presence of a catalytic amount of Pi. For the synthesis, a reaction mixture (2 ml) containing 200 mM sucrose, 20 mM P$_i$, 0.47 U/ml sucrose phosphorylase, 0.072 U/ml xylose isomerase, and 0.37 U/ml laminaribiose phosphorylase in 50 mM imidazole-HCl buffer (pH 7.0) was used. The concentration of laminaribiose reached 110 mM after 48 hr incubation at 37°C. Laminaritriose was also formed in the same mixture after 15 hr, and its concentration reached 22 mM.

Laminaribiose is $\beta$-1,3-linked glucobiose. This compound has been expected as one of new food or industrial materials, such as a soluble dietary fiber or a growth-promoting factor of bifidobacteria (unpublished results). It can be produced by partial hydrolysis or acetylation of $\beta$-1,3-glucan with enzymes or acids. However, it is very difficult to produce industrially laminaribiose by these methods because $\beta$-1,3-glucan is not inexpensive. Laminaribiose also can be synthesized from glucose-1-phosphate (G-1-P) and glucose using a crude extract of Euglena gracilis Z cells containing laminaribiose phosphorylase [EC 2.4.1.31] and $\beta$-1,3-oligoglucan phosphorylase [EC 2.4.1.30], but G-1-P is not also inexpensive. Furthermore, the yield of laminaribiose is not high, because of forming higher laminarioligosaccharides catalyzed by the latter enzyme. We developed a novel method to synthesize cellobiose from sucrose using sucrose phosphorylase, xylose isomerase and cellobiose phosphorylase. The method can be applied for synthesis of laminaribiose. The present paper describes the new method for synthesis of laminaribiose.

Sucrose phosphorylase from Leuconostoc mesenteroides [EC 2.4.1.8] was purchased from Sigma Chemical Co. (St. Louis, USA). Xylose isomerase (glucose isomerase, [EC 5.3.1.5] from Streptomyces was purchased from Kanto Chemicals (Tokyo, Japan) as dried cells and extracted by sonication. Laminaribiose phosphorylase was purified from a crude extract of E. gracilis Z (IAM E-6) cells by using hydrophobic chromatography to be separated from $\beta$-1,3-oligoglucan phosphorylase.

One unit of sucrose phosphorylase was defined as the amount of the enzyme which produced 1 $\mu$mol G-1-P per minute from 10 mM sucrose and 10 mM P$_i$ in 50 mM imidazole-HCl buffer (pH 7.0) at 37°C. That of laminaribiose phosphorylase was defined as the amount of the enzyme which produced 1 $\mu$mol P$_i$ from 10 mM glucose and 10 mM G-1-P in the same
condition. That of xylose isomerase was defined as the amount of the enzyme which produced 1 μmol glucose from 10 mM fructose in the same condition. The amount of G-1-P was measured by the phosphoglucomutase-glucose-6-phosphate dehydrogenase system. That of glucose was done by the glucose oxidase-peroxidase system with mutarotase using Glucose C Test Wako (Wako Pure Chemicals, Osaka, Japan). Those of oligosaccharides were measured by HPLC equipped with an RI detector using LiChrosorb NH₂ column (Cica-Merck, Tokyo, Japan) with 80% acetonitrile as a solvent. The products which formed in the reaction were identified by comparing their retention times on HPLC using both LiChrosorb NH₂ column (Cica-Merck, solvent: distilled water) with those of the standard laminarioligosaccharides (Yaizu Suisankagaku, Yaizu, Japan).

The new method to synthesize laminaribiose consists of three enzymatic reactions as follows:

\[
\text{Sucrose} + P_i \rightarrow \text{G-1-P} + \text{Fructose} \quad (1)
\]
\[
\text{Fructose} \rightarrow \text{Glucose} \quad (2)
\]
\[
\text{G-1-P} + \text{Glucose} \rightarrow \text{Laminaribiose} + P_i \quad (3)
\]

Sucrose \rightarrow Laminaribiose \quad (4)

At first, sucrose is phosphorolyzed into fructose and G-1-P by sucrose phosphorylase as in Formula (1). The resultant fructose is converted to glucose by xylose isomerase as in Formula (2). Then glucose and G-1-P are condensed into laminaribiose with release of \( P_i \) by laminaribiose phosphorylase as in Formula (3). When the reaction formulae are summed up, Formula (4) is obtained indicating that sucrose can be converted into laminaribiose in a single reaction mixture containing the three enzymes and a catalytic amount of \( P_i \). In the case of the enzymatic synthesis of cellobiose, laminaribiose phosphorylase using in Formula (3) is substituted with cellobiose phosphorylase.

The reaction was carried out in 2 ml of 50 mM imidazole-HCl buffer (pH 7.0) containing the following compositions: 200 mM sucrose, 20 mM \( P_i \), 0.47 U/ml sucrose phosphorylase, 0.072 U/ml xylose isomerase, 0.37 U/ml laminaribiose phosphorylase and 5 mM MgCl₂. The result is shown in Fig. 1. Sucrose decreased during the reaction and disappeared after 48 hr. On the other hand, laminaribiose increased with decrease in sucrose, but not stoichiometrically. Its concentration reached 110 mM after 48 hr. Laminaritriose appeared after 18 hr and finally reached 22 mM. Concentrations of fructose and glucose increased during the reaction to become 46 mM and 31 mM, respectively, after 48 hr. At the final state of the reaction, a trace amount of laminaritetraose was detected on the chromatogram.

The retention times of the laminaribiose synthesized in the reaction on the two different columns corresponded to those of the standard laminarioligosaccharides. To confirm its structure, the laminaribiose (25 mg) was isolated from 1 ml of the reaction mixture using a column of Toyopearl HW40S (Tohsoh, Tokyo, Japan, 2.5 cm × 40 cm) after treating the mixture with invertase, and its \(^{13}\text{C}-\text{NMR} \) spectrum was taken in \( \text{D}_2\text{O} \) using a JNM-GX400 spectrometer (JEOL, Tokyo, Japan) as shown Fig. 2. The spectrum obtained supports that the structure is \( \beta-1,3 \) linked glucobiose, and it corresponds to that of a standard laminaribiose.

In the case of the cellobiose synthesis, cellobiose increased stoichiometrically with decrease in sucrose, and no cellotriose or higher oligosaccharides appeared in the reaction mixture. The cellobiose phosphorylase cannot use cellobiose as a glucosyl acceptor.
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Fig. 2. A $^{13}$C-NMR spectrum on the laminaribiose synthesized.

Values were indicated in $\delta$ ppm. Dioxane was used as an internal standard (67.4 ppm, indicated by an arrow).

However, the laminaribiose phosphorylase can use laminaribiose as a glucosyl acceptor to form laminaritriose with a lower rate compared with glucose.$^{7,11}$ Formation of laminaritriose (and laminaritetraose) is probably due to the following by-reaction.

$$\text{laminaribiose} + G-1-P \rightleftharpoons \text{laminaritriose} + P_i \quad (5)$$

$$\text{(laminaritriose} + G-1-P \rightleftharpoons \text{laminaritetraose} + P_i) \quad (6)$$

Laminaribiose was synthesized at 55% yield from sucrose. This result suggests that laminaribiose can be produced effectively from inexpensive materials. Xylose isomerase is produced industrially and is commercially available. Recently, cloning and overexpression of sucrose phosphorylase were reported.$^{12}$ However, laminaribiose phosphorylase is not commercially available so far. If commercial production of this enzyme is realized, the present method can be used for industrial production of laminaribiose.

In addition, 1 mole of laminaritriose was formed from 2 moles of sucrose in the reaction. The yield of laminaritriose, 22 mM, means that 44 mM (22%) of sucrose were consumed to form laminaritriose. If a laminaribiose phosphorylase that does not use laminaribiose as a glucosyl acceptor is obtained, the yield of laminaribiose will be significantly improved.

REFERENCES


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スクロースホスホリラーゼ、キシロースイソメラーゼおよびラミナリビオースホスホリラーゼを用いたスクロースのラミナリビオースへの変換

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ラミナリビオースを、スクロースを原料として触媒量のリン酸存在下でスクロースホスホリラーゼ、キシロースイソメラーゼおよびラミナリビオースホスホリラーゼを同時に作用させることにより合成した。50 mM イミダソール-塩酸緩衝液（pH 7.0）中、20 mM リン酸存在下において、200 mM スクロースに 0.47 U/ml スクロースホスホリラーゼ、0.072 U/ml キシロースイソメラーゼ、0.37 U/ml ラミナリビオースホスホリラーゼの混合酵素系を37℃で作用させたところ、反応48時間後に反応済液中のラミナリビオース濃度は110 mM（収率55％）に達した。反応15時間後からラミナリントリオースの生成がみられ、反応48時間後には22 mM に達した。また微量のラミナリントリオースの生成も確認された。セロビオースホスホリラーゼを用いた同様なセロビオースの合成実験では、三糖以上のオリゴ糖は生成しなかった。この違いはセロビオースホスホリラーゼが三糖以上にはまったく作用しないのに対して、ラミナリビオースホスホリラーゼは弱いながらも三糖以上に作用するため生成されるものと考えられた。