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Original research article

Sucrase Activity and Exopolysaccharide Partial Characterization From Three *Weissella confusa* Strains

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ABSTRACT

Exopolysaccharides (EPSs) produced by lactic acid bacteria have been well known for their important economic value in food, pharmaceutical and health industries. Large extracellular enzyme sucraes are used by lactic acid bacteria to polymerize EPS, i.e. fructansucrase and glucansucrase. This study aimed to characterize sucrase activity of three *Weissella confusa* strains MBF8-1, MBF8-2 and MBFCNC-2(1), which were isolated previously from local beverages and their EPS products as well. All strains showed ability to form mucoid and slimy colonies by visual inspection on agar plate using raffinose as substrate suggesting that they possessed fructansucrase activity besides glucansucrase. Obtained EPS products were characterized by HPLC analysis after hydrolysis using 3% TCA at 100 °C for 1 hour, and by viscosity as well. All strains exhibited similar peak patterns, assuming that all of them possessed fructan EPS product. Supernatant and cell pellet were also analyzed by *in situ* activity assay performing periodic acid Schiff staining after polyacrylamide gel electrophoresis; only cell pellet showed sucrase activity. Viscosity observation showed that EPS products from all strains were able to increase the viscosity slightly.

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1. Introduction

Exopolysaccharides (EPSs) are versatile metabolic products of lactic acid bacteria (LAB). Due to the wide diversity in composition and variation in their high molecular weights (10–1000 kDa), EPSs have found a myriad of multifarious applications in many industrial and medical sectors, including food and pharmaceuticals (Nwodo *et al.* 2012; van Hijum *et al.* 2006; Velazquez-Hernandez *et al.* 2009).

EPSs produced by LAB are homopolysaccharide (HoPS) and heteropolysaccharide. HoPSs are composed of repeating units of a single monosaccharide, mainly glucose or fructose; the product is designated as glucan EPS and fructan EPS. Heteropolysaccharides are composed of repeating units consisting of two or more monosaccharides, mainly galactose, glucose and fructose (van Hijum *et al.* 2006). Glucan EPS and fructan EPS have been widely known, i.e.

dextran, inulin, and levan. Enzyme involved in this HoPS synthesis is a large extracellular enzyme known as sucrase.

The EPSs produced by LAB exert various functions including adherence of cells to surfaces during colonization, protection from attack by antimicrobial agents, and as the communal life of biofilms (Nwodo *et al.* 2012; van Hijum *et al.* 2006). They involve the microbial quorum sensing control through regulation of gene expression for proteins involved in EPS biosynthesis (Vu *et al.* 2009). Jellification of cane sugar syrups by microbial origin had been reported 150 years ago (Pasteur 1861). This effect was caused by an EPS named dextran due to its positive rotary power. As one of bacterial extracellular biopolymer, EPS exists as the most abundant biopolymers classified based on their location relative to the cell, i.e. loosely attached and unattached extracellular polymeric substances corresponding with the cell in structural and functional relationship. The loosely attached EPSs serve structural and protective purposes for the bacteria, and in addition, may take the form of a covalently bound cohesive layer. On the other hand, they may be unattached completely excreted into the environment as slime (Nwodo *et al.* 2012).

Gene coding for glucansucrase (GS; *gtf*) and fructansucrase (FS; *ftf*) has been identified from three *Weissella confusa* strains originally isolated from Indonesian beverages, i.e. MBF8-1, MBF8-2 and

Abbreviations: EPS, exopolysaccharide; *ftf*, fructosyltransferase; FS, fructansucrase; *gtf*, glucosyltransferase; GS, glucansucrase; LAB, lactic acid bacteria; PAS, periodic acid Schiff.

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MBFCNC-2(1) (Malik *et al.* 2009; Malik 2012). The strain MBF8-1 carried two *gtf*, whereas strain MBF8-2 exhibited one *gtf* and one *ftf*. Unique features of these genes were discussed in previous study, and thus generated a challenge for further characterization of the FTF/GTF proteins. The strain MBFCNC-2(1) interestingly possessed *ftf* which showed highest similarity to a putative inulosucrase of *Lactobacillus reuteri*. LAB strains carrying multiple GS genes (*gtf*) in their genome sequence have been reported (Van Hijum *et al.* 2006). Moreover, FS inulosucrases are exclusively present in LAB, unlike other FS enzymes, levansucrases, which are widely distributed in both gram-positive and gram-negative bacteria. Fructan EPSs are of high demand for industrial sectors, thus making the enzyme involved in high interest to be explored from various sources and to be bio-engineered as well.

In this study, we reported the characterization of those strains regarding the production of EPS by sucrase activity, as well as simple preliminary GS/FS identification. This study exhibits steps to narrow down the FS/GS identification by their enzyme activities performing *in situ* activity assay after visual inspection, as well as characterization of the EPS product by using HPLC. Cross-check of this report with previous studies reveals the possibility for more FS and/or GS genes to be identified and studied.

2. Materials and methods

2.1. Bacterial strains and growth condition

All *W. confusa* strains, i.e. MBF8-1, MBF8-2 and MBFCNC-2(1) (Malik *et al.* 2009; Malik 2012) were cultivated at 32 °C in MRS (De Man *et al.* 1960) medium (Difco, Franklin Lakes, NJ), or in MRS-s medium (i.e. MRS medium with 100 g/L sucrose instead of 20 g/L glucose) for EPS production. For visual inspection, MRS medium modified with 5% raffinose (MRS-r), i.e. MRS medium with 50 g/L raffinose (Merck), and with 5% sucrose (MRS-s) was used specifically for FS enzyme activity detection (Malik *et al.* 2009; Malik 2012). Agar plates were made by adding 1.5% agar to the MRS medium.

2.2. Visual inspection assay for GS and FS activities

Visual inspection for FS enzyme activity was done as described previously (Malik *et al.* 2009).

2.3. EPS production and purification

The EPS production of three *W. confusa* strains were done by using MRS medium containing 10% sucrose as growth medium at 32 °C without agitation for 24 hours. Sucrose was separately autoclaved and then added to the sterilized MRS medium. Bacterial cell cultures were harvested by centrifugation at 4 °C and 5 000g for 30 minutes, the supernatants were collected and were kept at 4 °C for 2–3 hours until chilled. Two volumes of 96% ethanol were added to the supernatants and were incubated back at 4 °C for 8 hours. Cloudy appearance of coagulated EPS was collected by centrifugation at high speed (minimum 10,000 g) for 1 hour; the EPS precipitant was decanted from the clear liquid part, and redissolved in pure water by warming it up at 37 °C on a water bath. After cooling, this suspension was repeated for one more precipitation process in an attempt to purify the EPS. EPS precipitation process was repeated until no precipitate formed. All EPS precipitant suspensions obtained were pooled, and were then lyophilized and kept as dried purified EPS.

2.4. Identification of sugar composition

Composition of EPS sugar was analyzed by TLC and HPLC. For TLC analysis, silica gel GF350 plate was used as stationary phase, and a mixture of organic solvents were employed, i.e. n-butanol, ethanol, water (5:3:2). For HPLC analysis, a Prominence-20AB

(Shimadzu, Japan) equipped with RI detector and an ion exchange CA-bonded SCR 101-C column have been employed with water only as moving phase, with resin polystyrene divinylbenzene as support. Hydrolysis of EPS was done by 3% TCA at 100 °C for 1 hour, and then was treated by deionization using cationic and anionic resins to remove all ion contaminants. HPLC was performed at 80°C and 1 mL/min with water as liquid phase. Reference samples used were glucose and fructose suspensions.

2.5. Measurement of viscosity

Observation of physical property (i.e. rheology/viscosity) was conducted using the LAB extracts harvested from the same fermentation process mentioned above, employing Ostwald viscometer. All *W. confusa* cultures in MRS-sucrose 10%, after incubation at 32 °C without agitation for 24 hours, were subjected to OD₆₀₀ measurement in an attempt to adjust all cell cultures at the same concentration. Cultures were centrifuged at 4 °C and 5 000g for 30 minutes, and the supernatant was decanted. Medium MRS-sucrose 10% was used as reference, and the density of the reference as well as all sample supernatants were measured by a Pycnometer before use in viscosity measurement.

2.6. *In situ* sucrase activity assay by SDS PAGE and periodic acid Schiff (PAS) staining

Supernatant and cell pellet obtained from liquid cultures were analyzed by performing polyacrylamide gel electrophoresis. The molecular mass was determined using the Laemmli system (Laemmli 1970) on 10% acrylamide (BioRad, USA) gel using electrophoresis AE-6530 mPAGE (Atto, Japan). Proteins were stained with Coomassie blue R-250 (BioRad), and molecular weight was estimated using protein marker PageRuler^R Prestained Protein Ladder^R (Fermentas, USA). Gel was run in duplicate, one for CBB staining and the other one for PAS staining. Raffinose buffer was used as substrate when the gel was incubated overnight, followed by staining with PAS reagent (Kralj *et al.* 2003). PAS staining consisted of incubation steps, first with 12.5% TCA for 30 minutes with slow and gentle agitation, followed by incubation with 1% periodic acid in 3% acetic acid for 50 minutes, and then with Schiff's Fuchsin-sulfite for 50 minutes, and last with 0.5% sodium metabisulfite for 30 minutes. Positive result can be observed as intensive pink to red band on the gel.

3. Results

3.1. GS and FS activities

The simple screening method for FS activity was done by visual inspection on agar plate using raffinose as substrate. Three *W. confusa* strains MBF8-1, MBF8-2 and MBFCNC-2(1), which were reported previously to possess mainly as FS gene carrier LAB as well as fructan EPS producers (Malik *et al.* 2009; Malik 2012), were able to produce a mucoid colony on plate as an indication of EPS product (Figure 1). The sucrase activity on MRS-sucrose 10% agar showed abundance of EPS product as seen on plate as mucoid and ropy colonies, whereas on MRS-raffinose 5% or MRS-raffinose 5% + glucose 5% showed lesser amount.

3.2. Isolation of EPS product

For the EPS production, the MRS-sucrose 10% liquid medium was chosen. Consecutive repeated simple purification steps were performed in this study to collect a rather pure EPS precipitate; no advanced technique needed for EPS product isolation. The dried EPS product obtained from isolation, purification and lyophilization appears as yellowish-white powdery precipitate as presented in Figure 2.

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