Role of microbubbles coupling fibrous-bed bioreactor in butyric acid production by Clostridium tyrobutyricum using Brewer’s spent grain as feedstock

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Research

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Abstract

Background: Microbubbles coupling fermentation is an efficient technology for improving oxygen transfer to aerobic microorganisms in a bioreactor containing an air sparger. However, the effect of this technology on anaerobic *Clostridium tyrobutyricum*, whose morphology develops viscous broth rheology and lumps impeding the nutrient mass transfer, has not been reported yet. Therefore, in this study, we evaluated the role of microbubbles coupling fibrous-bed bioreactor (MBFBB) on butyrate production by immobilized *Clostridium tyrobutyricum* cells obtained from renewable feedstock brewer’s spent grain (BSG).

Results: Compared with the conventional FBB fermentation, two-fold shorter fermentation time and two-fold higher butyrate productivity were achieved by MBFBB-immobilized *C. tyrobutyricum* cells. Furthermore, long-term stability and reliability for butyrate production were obtained in ten cycle process using BSG hydrolysate. Finally, fed-batch fermentation using BSG hydrolysate produced a high butyrate titer of 43.68 g/L, with a significantly higher selectivity of 11.67, reducing the production cost of downstream processing. The results indicated that butyrate productivity of 4.21 and 4.36 g/L·h were successfully obtained in BSG and glucose medium, thereby representing the highest productivity reported to date in *C. tyrobutyricum*. To our knowledge, this is also the first report related to microbial production of butyrate from brewer’s spent grain.

Conclusion: The MBFBB-based fermentation process with BSG is a robust and ecofriendly technique, which would provide insights into future development of commercial biobutyrate production for the animal feed supplement market.

Highlights

1. Production of butyrate from brewer's spent grain, an abundant industrial waste biomass
2. *tyrobutyricum* in MBFBB was used for butyrate fermentation
3. Cells in MBFBB yielded a higher ATPase activity, sugar utilization rate and productivity
4. The fermentation process from BSG hydrolysate has long-term stability and reliability
5. The MBFBB-based process was economic and attractive for waste biomass biorefinery

Background

Butyric acid, an oligomerize fatty acid, has been widely adopted in the fields of food, animal feed supplement, pharmaceutical, perfume, and industrial chemicals, with a global market of more than $8 \times 10^4$ metric tons per year [1]. However, the industrial production of butyrate is currently dominated by the chemical oxidation of petroleum-based butyraldehyde, which caused environmental pollutions and is also not favored for the food and feed industries [2,3]. In contrast, naturally derived butyrate through microbial fermentation has emerged as a promising alternative and extensively investigated for eco-friendly characteristics. *Clostridial* species are the most feasible strains for industrial butyrate bio-
production, especially, *Clostridium tyrobutyricum* due to its higher selectivity and tolerance to a high concentration of products [4-6]. Besides, the conventional fermentation technology for industrial production of butyric acid is not economically feasible due to cytotoxicity and byproducts generation, resulting in a low titer, yield, and productivity of butyrate [7]. By using an immobilized-cell bioreactor, the product inhibition could be partially relieved by *in situ* product removal and cell immobilization. However, the butyrate productivity in typical immobilized-cell fermentation process significantly decreases over extended operation periods, resulting from dead cell lumps accumulation and poor mass transfer under high cell density.

Microbubbles coupling fermentation is an efficient technology making breakthrough progress in recent years and has been widely adopted for the production of bioethanol [8], cellulase [9], and polyhydroxy butyrate [10]. Microbubbles exert outstanding characteristics than normal bubbles. The microbubbles dispersion generator can reduce the size of normal sparged gas bubbles from 3-5 mm to approximately 20-100 µm, leading to lower rising velocity than normal bubbles [11,12]. Additionally, the negatively charged surface prevents the microbubbles from coalescence, allowing them to persist in water for a longer period of two weeks [13]. Further, the microbubbles possess high internal pressure and a large specific surface area, which can significantly improve the mass transfer rate from bubbles into the aqueous phase resulting in higher dissolved gas concentration. Subsequently, the microbubbles can prolong the reactivity, increase the dissolved nutrient concentration, and the synthetic efficiency of target metabolites. It has been reported that microbubbles sparged fermenters could be energy efficient up to 0.01 kW per m$^3$ of fermentation capacity, lowering overall operating costs [14].

In contrast, the replacement of pure carbon source sugar with low-cost waste biomass (e.g., corn fiber, waste paper, oilseed rape straw, and wheat straw) could notably reduce the butyrate bio-production cost [15-18]. Although various waste biomasses have been explored to produce butyrate, the complexity of lignocellulosic biomass, especially lignin and the crystalline structure of cellulose, requires an additional process, such as pretreatment and hydrolysis to remove biomass recalcitrance forces and polymeric carbohydrate breakdown. Therefore, suitable and sustainable biomasses with high monosaccharide contents are highly necessary to simplify pretreatment and increase the titer, yield, and butyrate productivity. Unlike biomasses from the seasonal agricultural crops, Brewer's spent grain (BSG) is a lignocellulosic waste biomass available throughout the year in beer factories, with low cost and large amounts. The worldwide annual production of BSG has been estimated to be approximately 38.6 × 10$^6$ MT per year [19]. BSG is mainly composed of fibers with its sugar content corresponding to half of the BSG composition on a dry weight basis. Lignin and proteins also contribute to the BSG composition in significant amounts. Hence it is mostly employed by local farms for cattle feed usage. Recently, BSG has been applied as a raw material to produce value-added compounds, such as xylitol, methane, ethanol, butanol, and biohydrogen. It should be noted that there is no report on the use of BSG as a substrate for biobutyrate production to reduce product cost.

The aim of this study was to evaluate the role of microbubbles coupling fibrous-bed bioreactor (MBFBB) on butyrate production by immobilized *Clostridium tyrobutyricum* cells obtained from renewable
feedstock brewer’s spent grain (BSG). This technology yielded a high cellular ATPase activity and sugar utilization rate of immobilized cells and enhanced butyrate productivity. Enhanced butyrate production from BSG hydrolysate was demonstrated in repeated-batch and fed-batch processes with *C. tyrobutyricum* cells immobilized in MBFBB. Finally, the techno-economic analysis proved the feasibility and advantages of our established fermentation process for butyrate production from industrial waste biomass BSG.

**Results**

**Fermentative production of butyric acid by immobilized *Clostridium tyrobutyricum* in FBB or MBFBB**

The fermentative productions of butyric acid by immobilized *C. tyrobutyricum* in FBB or MBFBB were evaluated and compared, taking glucose as the carbon source (Table 1). Although the titers and yields of butyrate and acetate, as well as the selectivity, were similar in both FBB and MBFBB, the fermentation time was significantly shortened in MBFBB (5 h) compared to FBB (12 h). This two-fold decrease led to an increment in butyrate productivity from 2.24 g/L·h to 4.36 g/L·h, and the glucose utilization rate was simultaneously improved from 1.25 to 2.35 g/L·h for about two folds. However, the selectivity (*g*\(_{\text{butyrate}}/g_{\text{acetate}}\)) in MBFBB did not vary from FBB, which might be due to the negligible butyrate inhibition and inefficient reutilization of acetate within a shorter fermentation time in MBFBB. Similarly, when BSG hydrolysate was applied as a substrate in MBFBB, the sugar utilization rate was 2.87 g/L·h, and the butyrate productivity was 4.21 g/L·h, which was the highest ever reported for butyrate fermentation [6]. Therefore, microbubbles exhibited positive effects on FBB fermentation and significantly improved the mass transfer rate between gas and liquid.

**Table 1** Fermentative production of butyric acid by immobilized *Clostridium tyrobutyricum* in FBB or MBFBB.
Effects of microbubbles on cellular ATPase activity and sugar utilization rate of *Clostridium tyrobutyricum* immobilized in FBB

The immobilized cells were harvested after 1, 3, 5, and 7 h in repeated batch fermentations of MBFBB and FBB to evaluate the potential effect of microbubbles on *Clostridium tyrobutyricum*. The cellular ATPase activities were measured, and the results are illustrated in Fig. 1a. It was observed that the immobilized cells in the microbubble coupling FBB had a significantly higher ATPase activity than the non-microbubbles coupling FBB (60.16 U/g vs. 18.2 U/g, 73.37 U/g vs. 35.54 U/g, 87.19 U/g vs. 45.76 U/g), and the cellular ATPase activities were increased by 230.5%, 106.4%, 90.5% after 1, 3, 5 h, respectively. When the fermentation ended at 7 h, the cellular ATPase activities of the immobilized cells in MBFBB significantly decreased. However, the cellular ATPase activities of the immobilized cells in FBB continuously increased with the prolonged fermentation time to maintain cell growth and proton transmembrane flux balance. Besides, the immobilized cells in MBFBB had a significantly higher cellular sugar utilization rate than FBB (0.050 g/g·h vs. 0.043 g/g·h, 0.067 g/g·h vs. 0.044 g/g·h, 0.076 g/g·h vs. 0.049 g/g·h), and the cellular sugar utilization rates increased by 16.2%, 52.3%, and 55.1%, respectively (Fig. 1b). With the progress of fermentation, both the cellular sugar utilization rates of the immobilized cells in microbubble coupling FBB and that in non-microbubbles coupling FBB significantly increased before the end of the fermentation.

*Fig. 1* Effects of microbubbles on cellular ATPase activity (A) and sugar utilization rate (B) of *Clostridium tyrobutyricum* immobilized in MBFBB. Results are expressed as the mean value and standard deviation.
Repeated-batch fermentation from BSG hydrolysate in MBFBB

The repeated-batch fermentation of BSG hydrolysate was first conducted in MBFBB to determine the long-term stability and reliability of the system. As depicted in Fig. 2a, no lag phase was found at the beginning of each cycle, which clearly indicated the high metabolic activities and adaptability of immobilized cells in the fresh medium. Most importantly, stable and reliable butyrate production was achieved for ten cycles and the titer, yield, and volumetric productivity were between 8.83 to 14.95 g/L, 0.30 to 0.48 g /g, and 2.85 to 6.00 g/L·h, with an average of 11.33 g/L, 0.40 g/g, and 4.21 g/L·h, respectively (Fig. 2b). Comparing to glucose, the fermentation using BSG hydrolysate similarly presented the reduced fermentation time and the improved productivity of butyrate. Notably, the selectivity significantly increased (6.77 g/g vs. 4.06 g/g) in the fermentation of BSG hydrolysate.

Fed-batch fermentation from BSG hydrolysate in MBFBB

When the fed-batch fermentation was applied, the titer of butyrate steadily increased with the addition of concentrated BSG hydrolysate throughout the whole process (Fig. 3). The butyrate productivity (3.45 g/L·h) and glucose consumption rates were high from beginning to the 16th hour, and then decreased gradually. At the end of the fermentation (~36 h), the highest titer of ~43.68 g/L was obtained. The overall yield (0.38 g/g) and productivity (2.33 g/L·h) of butyrate were only 95.0% and 55.3% of those from the repeated-batch fermentation, which might be resulted from the increased product inhibition of high concentrations butyrate. However, the fed-batch fermentation using concentrated BSG hydrolysate exhibited a significantly higher selectivity (11.67 g/g) for butyrate production than the batch fermentation (6.77 g/g), which would definitely benefit to the product recovery and significantly reduce the total cost for biobutyrate production.

Process design and techno-economic analysis for sodium butyrate production from BSG

The preparation of sodium butyrate from BSG includes the alkali pretreatment, followed by the enzymatic hydrolysis of BSG, the fermentation of Clostridium tyrobutyricum in MFBB, and the downstream separation and production of sodium butyrate. Since the strong offensive odor is not suitable for animal feed industries application, sodium butyrate should be additionally encapsulated by gelatin and maltodextrin and then spray-dried to granules [20]. The flowchart for manufacturing encapsulated sodium butyrate from BSG is illustrated in Fig. 4.
Fig. 4 A general process flowchart for manufacturing encapsulated sodium butyrate from BSG.

The techno-economic feasibility of sodium butyrate production from BSG was analyzed by SuperPro Designer. Based on 42.4% (w/w) sugar yield from BSG (~20% cellulose, ~20% hemicellulose) at $29 per dry ton and 20 g/L butyrate with a yield of 0.4 g/g sugar and productivity of 4.21 g/L·h in fermentation, butyrate can be produced at the cost of $1.5 per kilogram for a 5,000-metric ton (MT) per year plant. The production cost would increase to ~$2.49 per kilogram if the plant size decreased to 1,000 MT. In contrast, the butyrate production at 5,000-MT scale would cost ~$1.75 per kilogram from barley and ~$1.95 per kilogram from glucose due to their much higher feedstock cost than BSG. The cost breakdowns for sodium butyrate production, including raw material costs (carbon source, nitrogen sources, enzyme, sulfuric acid, NaOH, and solvent used in extraction), utility costs (nitrogen gas, electricity, steam, and water), equipment maintenance and depreciation costs, and labor costs are also shown in detail (Fig. 5). The raw materials accounted for 43% of the gross costs when glucose (95% glucose, $450 per MT) was used as a carbon source. Moreover, this ratio decreased to 37% for barley (65% glucose, $180 per MT) and 25% for BSG (43% glucose, $29 per MT), respectively.

Fig. 5 Sodium butyrate production costs for 500, 1,000, 2,000, 3,000 and 5,000 MT plants using glucose, barley, and BSG, respectively, as feedstock in MBFBB fermentation processes. Cost data on capital equipment, energy, tax, interest, and labor were collected from existing butyrate fermentation plant and literature data.

The economics of sodium butyrate production could be influenced by the yield, productivity, and production scale of butyrate (Fig. 6). The estimated manufacturing costs for 30% (w/v) sodium butyrate production from BSG was ~ $1500 per MT with 0.4 g/g yield, 0.421 g/L·h productivity and 5,000 MT/yr scale. This value is much lower than the current cost of ~$1,800/MT for the petroleum-derived butyrate. When the butyrate productivity is improved, the capital investment and product cost decrease, leading to an increased return of investment (ROI) [21]. At a sale price of $1.8 per kilogram butyrate, ROI is 33.6% with a payback period of ~3 years for the 5,000-MT plant, whose estimated total capital investment is ~$14.6 MM. Compared with the butyrate cost from corn husk hydrolysate [20], butyrate production from BSG would be economically beneficial because of its higher ROI.

Fig. 6 Effects of annual production, butyrate yield and productivity on manufacturing cost of sodium butyrate from glucose, barley and BSG. (a) Effect of annual production on manufacturing cost. (b) Effect of butyrate yield on manufacturing cost. (c) Effect of butyrate productivity on manufacturing cost and return of investment.

Discussion

Mass transfer processes have a significant effect on microbial growth in industrial fermentations. Nutrients must be continuously replenished in the liquid layers closest to the microorganisms. Nutrients, such as glucose or ammonia of 1 mol/L concentration in the fermentation might cause mass transfer issues [22]. In this study, the fermentation process with microbubbles coupling fibrous-bed bioreactor was
superior to the traditional fibrous-bed bioreactor and had a shorter fermentation time between 12 h to 5 h. It also had a two-fold higher butyric acid productivity from 2.24 g/L·h to 4.36 g/L·h. The fermentable sugar utilization rate significantly increased by two-fold from 1.25 to 2.35 g/L·h. It can be concluded that microbubbles in MBFBB exhibit a positive effect on the fermentation and significantly improve the mass transfer efficiency between gas and liquid due to their small size, long existence time, and higher interface zeta potential.

The cellular ATPase activities and sugar utilization rate variations were investigated during the fermentation of butyric acid with MBFBB to evaluate the effects of microbubbles in MBFBB on the immobilized cells. It was observed that the immobilized cells in MBFBB had a significantly increased ATPase activity and sugar utilization rate, indicating that bacteria require a higher amount of ATP and glucose to maintain a higher growth rate. The metabolic pathway in *C. tyrobutyricum* is as follows: each unit of glucose can produce 3 units of ATP when producing butyric acid through the butyrate metabolic pathway, and each unit of glucose can produce 4 units of ATP when producing acetic acid through the acetic acid metabolic pathway. Therefore, in the rapid bacterial growth stage, cell metabolism requires more ATP, and the metabolic pathway is more inclined to the acetate metabolic pathway producing more ATP [2]. Our experimental results were consistent with these results, where the acetic acid concentration of the exponential phase of the fermentation increased faster than the end of the exponential phase. Furthermore, the decoupling effect of main products (butyrate and acetate) oxidative phosphorylation will interfere with establishing and maintaining the cell transmembrane pH gradient [23]. In order to maintain a proper intracellular and extracellular proton gradient, ATP must be consumed, and these protons should be pumped out of the cell by transmembrane ATPase. An active transmembrane ATPase is essential to prevent main products from acidifying the cytoplasm and maintaining the relative stability of intracellular pH [24]. In the fermentative production of butyric acid with MBFBB, the production rates of butyrate and acetate were much higher than the conventional FBB. A larger amount of ATP was required to maintain the relative stability of intracellular pH.

Additionally, pure monosaccharides derived from crops are not sustainable and suitable substrates for the production of bulk biochemicals due to the high feedstock cost and impact on food supply. To overcome these limitations, the feedstocks for bio-butyrate production are gradually enlarging from food crops to inexpensive and renewable biomass, such as waste biomass [25]. Waste biomass as low-valued byproducts obtained from agriculture (wheat straw, oilseed rape straw, corn fiber, sugarcane bagasse, sorghum stalk, and corn husk) and forestry (waste paper) has been a focus of recent studies for butyric acid fermentation [5, 15-18, 20, 26]. Unlike seasonal crop wastes, BSG can be obtained in large quantities at a low cost throughout the year. BSG mainly composed of the barley malt grain husks in mixture with part of the pericarp and seed coat layers of these grains, rich in sugar, protein, and minerals, is the main by-product of the beer brewing mill, and the annual production of BSG in China has been estimated at approximately 10 million MT per year [27]. As a biomass feedstock, BSG can not be directly utilized by *Clostridium tyrobutyricum* cells and must be pretreated and hydrolyzed to generate monosaccharides. Sulfuric acid combined with high temperature is one of the most commonly employed methods to hydrolyze biomass. However, the biomass acid pretreatment could generate various toxic by-products,
including acids (coumaric, glucuronic, and formic acids), salts (neutralization products), 5-HMF, and furfural, which could exert inhibitory effects on the C. tyrobutyricum cells [17, 28]. In order to avoid these inhibitors, alkali and enzymatic hydrolysis pretreatments of BSG were performed, and the highest sugar yield of 42.4% was achieved under the optimum hydrolysis conditions. The hydrolysates presented low toxicity (0.001 mg/mL HMF and 0.006 mg/mL furfural), and thus, it could eliminate the detoxification stage before fermentation. Moreover, a stable and reliable production of butyrate was obtained in repeated-batch and fed-batch fermentation processes using BSG hydrolyzate. The average butyrate titer, yield, productivity, and selectivity were 11.33 g/L, 0.40 g/g, 4.21 g/L·h, and 6.77 g/g, for repeated-batch fermentation and 43.68 g/L, 0.38 g/g, 2.33 g/L·h, and 11.67 (gbutyrate/gacetate) for fed-batch fermentation. The butyrate titer and yield obtained from the fermentation of BSG hydrolyzate were comparable to those of traditional monosaccharide substrates, such as glucose, xylose, and fructose. Moreover, the productivity and selectivity exceeded higher levels than other low-cost feedstocks.

Overall, the production cost of encapsulated sodium butyrate from petroleum-derived butyrate ($1800 per MT) for animal feed application is currently sold at US $2380-$3174 per MT. A similar product can be produced from BSG at a cost of US $1517-$2100 per MT, assuming a similar encapsulation process cost for the fermentation-derived sodium butyrate. With a large gross margin of more than $1000 per MT, the valorization of BSG process is highly profitable and economically attractive.

In summary, the role of MBFBB in the fermentative production of butyrate by immobilized Clostridium tyrobutyricum cells from a renewable feedstock of BSG was evaluated in this study. Compared with the conventional FBB fermentation, two-fold shorter fermentation time and two-fold higher butyrate productivity were achieved. Furthermore, long-term stability and reliability for butyrate production were achieved and fed-batch fermentation using BSG hydrolysate produced a high butyrate titer with a significantly higher selectivity. The study results indicated that the MBFBB-based fermentation process with BSG is a robust and ecofriendly technique, which would provide insights into future development of commercial biobutyrate production derived from lignocellulosic waste biomass in beer factories.

Methods

Cultures and medium

Butyrate-producing strain Clostridium tyrobutyricum ZJUT1 was collected from strain Clostridium tyrobutyricum ZJU8235 provided by Professor Zhinan Xu from Zhejiang University. This strain was adapted in a fibrous bed bioreactor by the previously reported methods [29]. The fermentation culture medium contained (g/L): glucose 30; yeast extract 5; peptone 5; (NH₄)₂SO₄ 3; K₂HPO₄ 1.5; MgSO₄·7H₂O 0.6; FeSO₄·7H₂O 0.03. The medium pH was adjusted to ~6.5-7.0 and sterilized at 121 °C for 20min.

Pretreatment and hydrolysis of brewer’s spent grain

Brewer’s spent grain was provided by Hangzhou Cheerday Brewery Co. Ltd, Hangzhou, China. The BSG was dried in an oven at 45 °C prior to the experimental runs. After alkaline pretreatment (1M NaOH
solution for 40 min at 121 °C) of BSG, the pH of the solid fraction was controlled at 4.8 by adding 4 M of hydrochloric acid. Glucose was extracted by enzymatic hydrolysis using cellulase (12000 U/mL, Hunan Hongying Biotech Co. Ltd) and β-glucosidase (110 U/g, Jiangsu Ruiyang Biotech Co. Ltd) for 24 h at 50 °C and 200 rpm. Three-level orthogonal tests were designed to evaluate the effects of BSG biomass (5, 10, 15%, w/v) and NaOH concentration (1, 2, 3%, w/v) in the alkaline pretreatment step, as well as the BSG biomass (5, 10, 15%, w/v), enzyme loading (45, 67.5, 90, U_{cellulase}/g_{substrate} and 45, 67.5, 90, U_{β-glucosidase}/g_{substrate}) in the enzymatic hydrolysis step for better glucose yield. A maximum glucose yield of 42.4% (g_{glucose}/g_{BSG}) was achieved (Table S1) under the optimized hydrolysis conditions. Then, the hydrolysate sugar compositions were determined and the generated monosaccharides contained 50.2 g/L glucose and 3.0 g/L xylose. A small amount of xylose in the hydrolysate might have been produced by the acid neutralization process. The BSG hydrolysate was collected by vacuum filter and stored in the refrigerator for future use.

**Fermentation in microbubbles coupling fibrous-bed bioreactor**

The MBFBB was composed of a 5-L stirred-tank fermenter containing an N_{2} sparger as microbubbles generator (The bubble generator diameter was 6 cm and the thickness was 1.2 cm, the pore size was 40 µm, and the plate had 680 pores per cm²) and a recirculation loop connected a 0.5-L fibrous-bed bioreactor (FBB). The FBB was made of a glass column packed with a spirally wound cotton towel (300 × 300 mm; 5 mm in thickness; with >95% porosity) overlaid with a stainless-steel mesh. The bioreactor system was operated at 37 °C, and the pH was controlled at 6.0 with 2 M NaOH through an auto-sensing and dosing system. Before use, the bioreactor containing the medium was sterilized by autoclaving at 121 °C for 30 min, and then flushed with N_{2} for 30 min. The fermenter was inoculated with 100 mL cells cultured in serum bottles and allowed to reach a cell density of 6.0 (OD_{600}) to start the fermentation. Later, the fermentation broth was circulated to immobilize cells through the fibrous bed at a pumping rate of 30 mL/min. After 48 h of continuous circulation, most of the cells were immobilized, and the cell density in the broth became constant. The fermentation broth in the fermenter was then removed, replaced with fresh medium, and increased the medium circulation rate to 100 mL/min to start the fermentation. The batch fermentations were carried out with glucose to evaluate the role of microbubbles in the fermentative production of butyric acid with MBFBB. When sugars were almost depleted, the fermentation broth in the fermenter was removed and replaced with a fresh medium. Repeated batch fermentation and fed-batch fermentation with BSG hydrolysate were then carried out to evaluate the kinetics and possible effects of hydrolysate inhibitors on long-term process performance. N_{2} was supplied to MBFBB at 1 VVM (L/L·min) throughout the fermentation process. Samples were taken at regular intervals for the analysis of biomass, substrate, and product concentrations.

**Characterization of microbubbles generated by N_{2} sparger**

Microbubbles generated by N_{2} sparger were characterized with the images captured by a digital camera (EOS M5, Canon) in an illuminating field created by a halogen lamp (500 W). The captured images were
analyzed by image processing software Digimizer [30]. Microbubbles image and the microbubble size
distribution are illustrated in Fig. S1. The microbubbles size distribution was relatively concentrated and
uniform and the bubble diameter was mainly distributed in the range of 0.2-0.25 mm with an average
diameter of 0.22 mm.

**Cellular ATPase activity Assay**

*Clostridium tyrobutyricum* cells were collected after 1, 3, 5, and 7 h in repeated batch fermentation with
glucose as the carbon source using microbubbles coupling or non-microbubbles coupling fibrous-bed bioreactors. *Clostridium tyrobutyricum* cell extract was prepared, and the cellular ATPase activity assay
was strictly performed following the instructions of the ATPase test kit (Ca$^{2+}$/Mg$^{2+}$ATPase) [29]. The
ATPase activity unit was defined as the amount of Ca$^{2+}$/Mg$^{2+}$-ATPase decomposing ATP to produce 1
μmol of inorganic phosphorus per gram of cell per hour is one enzyme activity unit.

**Cellular sugar utilization rate assay**

Cell pellets were harvested by centrifugation at 8000 rpm and 4 °C for 5 min, then the pellets were washed
twice with equal volume of distilled water and resuspended in 10 g/L glucose solution. After incubation
at 37 °C for 6 h, the suspension aliquots were discarded to determine the glucose concentration [16,17].
Cellular sugar utilization rate was expressed as follows: cellular sugar utilization rate (g/g·h) = ($W_0$ -
$W_G$)/$W$/6, where $W_0$ is the initial weight of glucose, $W_G$ is the final weight of glucose, and $W$ is the weight
of wet bacteria.

**Analytical methods**

During fermentation, the cell concentration was determined at 600 nm (OD$_{600}$) by a spectrophotometer,
and the glucose concentration was determined by SBA Biosensor Analyzer (Biology Institute of Shandong
Academy of Science. Shandong, China). The concentrations of other sugars in the hydrolysate were
determined by the previously reported methods [16,17], where HPLC-ELSD was used with Bio-Rad Aminex
HPX-87H column. The butyrate and acetate concentrations were determined by gas chromatography with
flame ionization detector and Stabilwax column (Restek 10624, USA) (30 m × 0.32 mm × 0.25 μm, $d_f$ =
0.25).

**Techno-economic analysis**

The economic feasibility for producing encapsulated sodium butyrate from BSG was analyzed using
SuperPro Designer.

**Declarations**

**Authors’ Contributions**
Linqi Zhao: Conceptualization, Validation, Formal analysis, Investigation, Writing original draft. Gaoya Sun: Investigation. Jing Jiang: Conceptualization, Validation, Writing - review & editing. Li Chen: Resources, Writing - review & editing. Jin Huang: Conceptualization, Validation, Writing - review & editing, Supervision, Project administration.

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Competing Interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Availability of data and materials

All data generated or analyzed during this study are included in this published article and its supplementary information files.

Consent for publication

Not applicable.

Ethics approval and consent to participate

Not applicable.

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