Production of bacterial cellulose by agitation culture systems

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Abstract: An economical mass production system of bacterial cellulose (BC) on agitated culture was constructed. We first conducted screening of BC producers in agitated culture. A total of 2096 strains were isolated from natural sources and the best BC producing strain, BPR2001, was selected. According to the taxonomic examination, we concluded that the strain belongs to a new subspecies, Acetobacter xylinum subsp. sucrofermentans subsp. nov. Of the several organic nitrogen sources used to supplement the culture medium, corn steep liquor (CSL) was found to be the most suitable for BC production. A mutant strain derived from BPR2001, BPR3001E, resistant to an analogue of PABA, sulfaguanidine, showed increased cell growth and 40% higher BC productivity than BPR2001. A host-vector system for the BC producing Acetobacter strain was then developed using an indigenous plasmid, pAH4, detected in BPR2001. The complete nucleotide sequence of pAH4 was determined and the shuttle vector pSA19 was constructed by connecting pAH4 to pUC18. pSA19 was found to be very stable and the system was suitable for introducing genes into the BC producer in order to increase the yield of BC. Effective agitation culture systems essential for the BC fermentation with high yields were also developed. When the BC produced in this system was examined as a wet-end additive for papermaking processes, both the tensile strength and filler-retention of the resulting handsheets were found to be distinctly improved.

Key Words: Bacterial cellulose, Acetobacter xylinum, Agitation culture, Host-vector system, Papermaking process.

INTRODUCTION

BC producing by acetic acid bacteria is far superior to its counterpart from plants, because of its exceptional purity, ultra fine network structure, high biodegradability and unique mechanical strenght (ref.1). BC is expected to be used for many industrial applications as a high-strength construction material, food additive and a component of biodegradable products and paper. One of the problems that hindered the industrial application of BC is its low yield from static culture systems. Therefore, an economical mass production system based on agitated culture is necessary.

In previous works, several strains were grown by agitated cultures but all of them produced less BC than in static cultures (refs. 2,5). We thought that agitated culture was not suitable for BC production with regard to these strains tested but some strains suitable for BC production in agitated culture might exist in nature. In this study, we first established a screening system for BC producers that were isolated. We isolated a strain, designated BPR2001, from a cherry as one of the most potent BC producer (ref.4). The isolate BPR2001 was examined to determine its taxonomic characteristics (ref.5). We propose a new subspecies, Acetobacter xylinum subsp. sucrofermentans as a potent BC producer.

Few reports on chemical cultural conditions of BC production by agitated culture have been published. Dudman et al. used such a culture system and studied the effects of sugar and ammonia-nitrogen concentration on BC production (ref.6). In this study, corn steep liquor (CSL) was found to be the most suitable organic nitrogen source for BPR2001. Lactate in CSL promoted both BC production and cell growth (ref.7).
There have been some studies on breeding BC producing bacteria. Previously, BC producers with increased bacterial cellulose synthase activity have been bred by genetic engineering, with branches of their metabolic pathway blocked to decrease the amounts of by-products (refs. 8, 9). We bred mutant cells whose growth was increased to increase the enhancement of BC productivity (ref. 10).

Several host-vector systems for BC producing bacteria were already available and some of these use broad host range plasmids, such as pRK311 (ref. 11), pVK100 (ref. 12), pRK248 (ref. 13) and others have used plasmids isolated from related strains (ref. 8). As a cloning vector, the use of indigenous plasmids is often very useful, because they are stable. The constructed vector is also expected to be stably maintained. In this study, we describe the construction of a host-vector system in a BC producing strain, BPR2001, by using a cryptic plasmid pAH4 isolated from this strain (ref. 15).

Although the BC from static culture systems has superior properties such as an ultra fine network structure, high biodegradability and unique mechanical strength as compared with green plant cellulose, it has few commercial applications due to its high price. The BC produced in an agitated culture system was examined as a wet-end additive for papermaking processes (ref. 15).

**BODY OF RESEARCH WORK**

**Screening of BC producers from natural sources:** A total of 547 samples from many places in Japan were collected. Buffered Schramm & Hestrin's medium (ref. 16) with 0.2% (w/v) acetic acid, 0.5% (v/v) ethanol, and 0.01% (w/v) cycloheximide was used for the BC producing bacteria were detected in 126 of the samples (Table 1). The most efficient isolation was from fruit samples with an isolation ratio of producers to the total number of samples of over 30%. No producers were detected in soil samples. A final total of 2,096 BC producers were isolated.

The amounts of BC accumulated by many strains were higher than those of the BC producing ATCC strains under both static and shaken culture conditions. A strain isolated from a black cherry showed the highest BC accumulations, 4.4 g/l, with CSL-Fru medium and it was selected and named BPR2001.

**Taxonomic examination of strain BPR2001:** A scanning electron micrograph of the unwashed product of strain BPR2001 presented in Fig. 1. The isolate BPR2001 was characterized by the formation of dihydroxyacetone from

**TABLE 1 The Results of Screening for BC Producers from Natural Sources**

<table>
<thead>
<tr>
<th>Sources</th>
<th>Number of samples</th>
<th>Ratio of isolation of cellulose producer (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fruits</td>
<td>353</td>
<td>31.7</td>
</tr>
<tr>
<td>Nuts</td>
<td>28</td>
<td>10.7</td>
</tr>
<tr>
<td>Sugar Cane</td>
<td>124</td>
<td>7.3</td>
</tr>
<tr>
<td>Flowers</td>
<td>35</td>
<td>5.6</td>
</tr>
<tr>
<td>Soil</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>547</td>
<td>23.0</td>
</tr>
</tbody>
</table>
TABLE 2 DNA Homologies between the Isolate BPR2001 and the Type Strains of *Acetobacter* species

<table>
<thead>
<tr>
<th>Source of labeled DNA</th>
<th>% Relative binding of DNA</th>
<th>Isolate BPR2001</th>
<th><em>A. xylinum</em> IFO 15237™</th>
<th><em>A. hansenii</em> ATCC 35959™</th>
<th><em>A. pasteurianus</em> ATCC 33445™</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isolate BPR2001</td>
<td></td>
<td>100</td>
<td>58.2</td>
<td>15.6</td>
<td>0.3</td>
</tr>
<tr>
<td><em>A. xylinum</em> IFO 15237™</td>
<td></td>
<td>56.5</td>
<td>100</td>
<td>24.3</td>
<td>9.1</td>
</tr>
</tbody>
</table>

The isolate BPR2001 was distinguishable phenotypically from the type strain of *Acetobacter xylinum* by growing on ethanol, glycerol, dulcitol, and sucrose, by acid formation from D-galactose, glycerol, meso-erythritol, sucrose and trehalose, and by oxidation of glycerol, D-mannitol and lactose (Table 3). We proposed a new subspecies, *Acetobacter xylinum* subsp. *surafermentans* as a potent BC producer.

**Stimulation of BC productivity by addition of CSL:** We examined the effects of various organic nitrogen sources on BC production. When 0.24% w/v (of the total nitrogen) soytone, yeast extract (YE), CSL or peptone was added to the basal medium, cell growth and BC production were stimulated only by CSL. The amounts of BC produced with the other nitrogen sources were almost the same as that in basal medium without organic nitrogen. Therefore, CSL was found to be the most suitable organic nitrogen source for BC production by the strain BPR2001.

To investigate the effects of CSL in detail, the chemical composition of all the organic nitrogen sources used, soytone, YE, and peptone, was analyzed. The major difference was that CSL contained lactate. The addition of lactate stimulated cell growth and BC production. The amounts of BC produced in lactate-supplemented media containing soytone or YE were almost the same as that in basal medium containing only CSL. These results suggested that lactate in CSL stimulated cell growth and BC production.

**Increase in BC production by sulfaguanidine-resistant mutants:** The relationship between BC production and cell growth was investigated and the production was found to be growth associated. Thus, it was expected that enhanced cell growth would lead to enhanced BC production. On the other hand, factors which promote the growth of and BC production by strain BPR2001 were investigated. It was...
found that the addition of p-aminobenzoic acid (PABA) promoted these effects. Therefore, mutants resistant to sulfaguanidine (SG), which is an analogue of PABA, were bred from BPR2001. Forty-one SG resistant mutants were obtained from about 60,000 BPR2001 cells through NTG-treatment. The distribution of BC production among these SG-resistant mutants was tested. The best BC producer, BPR3001E, was selected.

Cell growth and BC production by the SG-resistant mutant BPR3001E were examined. The course of BC production by BPR3001E compared with BPR2001 in a jar fermentor was tested (Fig.2). Cell growth and BC production of BPR3001E were found to be higher than those of BPR2001. BPR3001E produced 9.7 g/l of BC from 44 g/l fructose. The BC production by the mutant was 40% greater than by BPR2001.

*Construction of a host-vector system:* An indigenous plasmid, named pAH4, was detected in a BC-producing strain BPR2001. This plasmid, consisting of 4002 base pairs contained an AT-rich region and encoded several open reading frames, as deduced by the complete nucleotide sequence. One of the putative open reading frames showed homology with replication proteins of other plasmids. A shuttle vector was constructed from *Escherichia coli* and this strain BPR2001 was constructed by connecting pAH4 to pVC18 (Fig.3). Electroporation of the shuttle vector into the strain BPR2001 yielded 1.7 x 10^5 ampicillin resistant transformants per μgDNA (Table 4). The shuttle vector was very stably maintained in the strain BPR2001.

**TABLE 4 Transformation Efficiency of BPR2001 with the Shuttle Vector**

<table>
<thead>
<tr>
<th>Plasmid (pSA 19)</th>
<th>From E. coli</th>
<th>From BPR2001</th>
</tr>
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<tbody>
<tr>
<td>Transformation Efficiency (cfu/μg DNA)</td>
<td>10</td>
<td>1.7x10^5</td>
</tr>
</tbody>
</table>

*BC as a new material for papermaking:* The application of BC to papermaking was investigated. The morphology of BC observed through a scanning electron micrograph comprises ultrafine fibrils which are approximately 1/100 width of pulp fibre. The BC can be obtained with two typical cultural methods, namely static and agitated culture. We have found some structural differences between these two types of BC. We, therefore, investigated the properties of the BCs for papermaking.

When these two types of BCs were added at the wet-end of paper production, both tensile strength and filler-retention of the handsheets were improved (Figs.4 & 5). In particular, the BC from agitated cultures had enhanced effects on filler-retention than that from static cultures.
Production of bacterial cellulose

CONCLUSIONS

The bacterial cellulose producer that yielded the highest polymer accumulation in agitated culture and obtained from natural sources was named BPR2001. According to the taxonomic examination, this strain BPR2001 is designated as a new subspecies, *Acetobacter xylinum* subsp. *sucrofermentas* subsp. nov. CSL was found to be the most suitable organic nitrogen source for BC production. Lactate in CSL stimulated cell growth and BC production. As sulfaguanidine resistant mutant BPR3001E derived from BPR2001 showed increased cell growth and 40% higher BC productivity. A host-vector system for a BC-producing strain was developed by using an indigenous plasmid, pAH4 found in BPR2001. The BC produced in the agitated culture when incorporated at the wet-end of the papermaking process enhanced both the tensile strength and filler-retention of the product.

REFERENCES


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