Optimization and characterization of dextranucrase production by local *Leuconostoc mesenteroides*

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**Abstract**

The dextranucrase production by local *Leuconostoc mesenteroides* was optimized and characterized to reach the optimum conditions of producing this enzyme which used in many food applications such as dextran production. The optimum conditions were using sugar as a carbon source with 20% concentration for giving the highest specific activity of dextranucrase, yeast extract as a nitrogen source, 25°C as incubation temperature, 6.5 optimum production pH and 10^6 inoculum size. The crude enzyme was characterized for pH stability and heat stability, they were 4-5 and 30°C for an hour, respectively. The enzyme was purified by using sephadex G-100 with a fold of 4.56 and overall yield of 20.5% of activity, then precipitate by ethanol with a fold of 17.88 and overall yield of 20.64% of activity.

**Key Words:** Dextranucrase, *Leuconostoc mesenteroides*, Optimization, Characterization.

**Introduction**

Dextranucrase (E. C. 2.4.1.5) from Leuconostoc species are used in industry for the production of oligosaccharides and dextran. Because of their desirable physiochemical properties in food and their prebiotics effect on intestinal bacteria, these oligosaccharides are very useful as food additives. Oligosaccharides produced by dextranucrase find several applications in food, feed, pharmaceutical, or cosmetic industries (5). Dextranucrase (glucansucrase or glucosyltranferase) is an extracellular enzyme that synthesizes dextran from sucrose (3). Many authors have described the effects of sucrose concentration, aeration rate, agitation speed, medium pH, incubation temperature, nature of the yeast extract and other nutritional requirements on the production of dextranucrase, dextran and fructose from sucrose using different strains of *Leuconostoc mesenteroides* (3,5). This enzyme is produced by many varieties of lactic acid bacteria, Leuconostoc and Streptococcus (6) Pediococcus, another lactic acid bacterium may also produce dextranucrase (9). Many authors purified this enzyme by different methods, The results confirmed the multimeric entity of this enzyme, which remains in single molecular form in the native state and separates into multiple bands when denatured by boiling, SDS or 2-mercaptoethanol (8). The aim of this study is to optimize the growth conditions for dextranucrase production by a local isolate of *Leuconostoc mesenteroides* from fermented cabbage.
Material and methods

Many effected factors in dextransucrase production were studied, which include carbon source and its percentages, nitrogen source, pH, temperature, to determine the optimal conditions for dextransucrase production.

Production media: The base media was as follows:

<table>
<thead>
<tr>
<th>The Substance</th>
<th>The weight: gm/100 ml. D. W.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sucrose</td>
<td>10</td>
</tr>
<tr>
<td>Yeast Extract</td>
<td>0.5</td>
</tr>
<tr>
<td>Peptone</td>
<td>0.5</td>
</tr>
<tr>
<td>K2HPO4</td>
<td>1.5</td>
</tr>
<tr>
<td>NaCl</td>
<td>0.001</td>
</tr>
<tr>
<td>MgSO4.7H2O</td>
<td>0.001</td>
</tr>
<tr>
<td>MnCl2. H2O</td>
<td>0.001</td>
</tr>
<tr>
<td>CaCl2</td>
<td>0.01</td>
</tr>
</tbody>
</table>

The pH was adjusted to 7.2 before sterilization by autoclave at 121⁰C for 15 minutes. The changes in this media were done as the optimal results respectively.

Optimal Carbon source:
Many carbon sources were used, whey, maltose, fructose, glucose, sugar, sucrose, malt and date syrup.

Optimal nitrogen source:
Ammonium sulphate, Sodium nitrate, peptone, tryptone, ammonium chloride, ammonium nitrate, yeast extract and thermal yeast extract were used as nitrogen sources.

Optimal temperature:
Temperature degrees (10, 25, 30, 35, 40)⁰c were chosen to determine the optimal temperature for dextransucrase production.

Optimal pH:
Five pH values (5, 6, 6.5, 7 and 8) of fermentation media were determined to stand on the optimal pH for dextranosecrase production.

Optimal inoculum's size:
Six inoculum's size of Leuconostoc mesenteroides (103, 104, 105, 106, 107,108) were used to emphasize the optimal size for dextranase production.

Dextranusecrase assay:
Dextranusecrase activity was measured according to Michelena et al. method (4) by using sucrose (1.25mg/ml) as a substrate at pH= 5 (0.1M Buffer) for 30 min. at 30⁰c., the reaction was stopped and detected by DNSA reagent.
The enzyme unit was defined as the amount of enzyme that could alter 1 mg of sucrose to dextran and fructose during one hour under test circumstances.

**Crude enzyme properties:**

1. **pH stability:**
   Crude enzyme was incubated with different pH (3, 4, 4.5, 5, 5.5, 6, 7, 8) For 1 hour, cooled with ice, then the activity was determined.

2. **Heat stability:**
   Crude enzyme was incubated with different temp. (20, 30, 40, 50, 60, 70, 80)°c at pH stability value for 1 hour, cooled with ice, the activity was estimated.

**Purification of enzyme:**

**Gel filtration**

The first step of purification was gel filtration with sephadex G-100 by using phosphate buffer 0.1M, the enzyme was gained by using phosphate buffer 0.1M, pH=5 containing 0.05M NaCl. The protein was determined by spectrophotometer at 280nm, enzyme activity was determined. Ethanol participation: the enzyme was participated by cooled ethanol, dextransucrase activity was determined by Michelena et. al. method (4) for the participate part, total activity was determined also. Protein was determined according to a modified method of lowery (2).

**Results and discussion:**

From fig.1 we can notice that the best carbon source was sugar for the highest activity and specific activity obtained with 20% concentrate (fig.2). Yeast extract recorded as a best nitrogen source as shown in figure 3. The optimal temperature for enzyme production was 25°c, and the optimal pH was 6.5 for the highest activity and specific activity (Fig.4 and Fig.5). The most suitable inoculum size for enzyme production was 106 as shown in figure 6. These results agreed with Bivolarski et al(1) when they studied the cell wall-bound of the bacteria by using different media, they found the best media for growth and enzyme production is by using glucose and fructose together.

The pH stability of the enzyme was determined at different pH value, it was stable at 4-5 (fig.8). The thermal stability of the enzyme was examined by assaying retained activities after incubation at various temperatures. The residual activity was stable after incubation for 1 hour at temperatures up to 30°C.(fig.9). These results was near to what Young-mo et. al.(11) gained of results. UIQader(10)said that the enzyme produced by *Leuconostoc mesenteroides* PCIR-4 was stable at 30°C for 240 hrs with 64.2 DSU/ml/hr. and the enzyme was most active in the pH range of 4.75 to 6.0. these results are so close to ours.
The purification step of gel filtration showed that there are 2 narrow peaks and a wide peak of protein, and there is only one sharp peak of dextranase enzyme (fig. 7), in another hand we can see the purification table in table 1 which showed that gel filtration step purified the enzyme with a fold of 4.56 and overall yield of 20.5% of activity and in the second step the enzyme retained his activity at the same yield. This is agreeing with the searcher Purama and Goyal (7).

![Diagram 1](image1.png)

**Fig.1** Dextransucrase activity and specific activity by using different carbon sources

![Diagram 2](image2.png)

**Fig.2** Dextransucrase activity and specific activity by using different Concentrations of sugar
Fig. 3 Dextranucrase activity and specific activity by using different Nitrogen sources.

Fig. 4 Dextranucrase activity and specific activity by using different degrees of temperature.
Fig. 5 Dextranucrase activity and specific activity by using different pH

Fig. 6 Dextranucrase activity and specific activity by using different Inoculum size
Fig. 7 Gel filtration of dextransucrase by Sephadex G-100 with phosphate buffer 0.1M, pH=5 containing 0.05M NaCl.

Table 1. Partial purification steps of dextransucrase

<table>
<thead>
<tr>
<th>Step of purification</th>
<th>Volume</th>
<th>Activity U/ml</th>
<th>Protein mg/ml</th>
<th>Qualitative activity U/mg</th>
<th>Total activity U</th>
<th>Folds of purification</th>
<th>Yields%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude enzyme</td>
<td>100</td>
<td>0.218</td>
<td>0.467</td>
<td>0.466</td>
<td>21.8</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>Gel filtration</td>
<td>30</td>
<td>0.149</td>
<td>0.070</td>
<td>2.128</td>
<td>4.47</td>
<td>4.56</td>
<td>20.5</td>
</tr>
<tr>
<td>Ethanol</td>
<td>3</td>
<td>1.5</td>
<td>0.18</td>
<td>8.333</td>
<td>4.5</td>
<td>17.88</td>
<td>20.64</td>
</tr>
</tbody>
</table>

Fig. 8 The pH stability of dextransucrase

Fig. 9 Heat stability of dextransucrase
References:


