Original Research Article

Isolation, Identification and Screening of Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210 for Biosurfactant Production

Arjun Karmakar1*, Jyotsna Kiran Peter1, Ankit Singla2 and Anita Raisagar1

1Sam Higginbottom University of Agriculture Technology and Sciences (SHUATS), Prayagraj, Uttar Pradesh, India
2Regional Center of Organic Farming, Ministry of Agriculture & Frames Welfare, Bhubaneswar, Govt. of India

*Corresponding author

A B S T R A C T

In the present study on the basis of cultural, morphological, biochemical and 16s rRNA gene sequencing two rhizospheric bacteria were identified as Brevundimonas aurantiaca KY231210 and Enterobacter cloacae KY231211 and screened for their biosurfactant production potential by oil displacement assay, blood haemolysis assay and emulsification index with vegetable oils (soybean oil and sunflower oil) in different incubation period (0, 24, 48, 72 and 96h). In oil displacement assay, Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210 broth culture revealed positive oil displacement but Brevundimonas aurantiaca KY231210 showed highest displacement values than Enterobacter cloacae KY231211. Both cultures showed haemolysis on blood agar media. Soybean oil showed higher emulsification index with Enterobacter cloacae KY231211 culture while Brevundimonas aurantiaca KY231210 culture showed higher emulsification index with sunflower oil. Brevundimonas aurantiaca KY231210 revealed highest displacement values with Soybean oil. Brevundimonas aurantiaca KY231210 showed highest yield in biosurfactant production as compare to Enterobacter cloacae KY231211.

Keywords
Biosurfactant, Brevundimonas aurantiaca KY231210, Enterobacter cloacae KY231211, PPBS, 16s rRNA

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Introduction

Biosurfactant was first discovered as extracellular amphiphilic compounds of fermentation bacteria (Kitamoto et al., 2009). Biosurfactants are diverse groups of surface active molecules/chemical compounds synthesized by microorganisms (Desai and Banat 1997). These are amphiphilic biological compounds produced extracellularly or as part of the cell membrane by a variety of yeast, bacteria and filamentous fungi (Mata-Sandoval et al., 1999, 2000; Chen et al., 2007) from various substances including sugars, oils and wastes. All biosurfactant are amphiphiles, these amphiphilic compounds are produced on living surfaces, mostly on microbial cell surfaces, or excreted extracellularly. These are amphipathic molecules consist of two parts—a polar (hydrophilic) moiety and nonpolar (hydrophobic) group. A hydrophilic group
consists of mono-, oligo- or polysaccharides, peptides or proteins and a hydrophobic moiety usually contains saturated, unsaturated and hydroxylated fatty acids or fatty alcohols. A characteristic feature of biosurfactants is a hydrophilic-lipophilic balance (HLB) which specifies the portion of hydrophilic and hydrophobic constituents in surface-active substances. Both the hydrophilic and hydrophobic domains confer the ability to accumulate between fluid phases, thus reducing surface and interfacial tensions at the surface and interface respectively (Karanth et al., 1999). This property of biosurfactant makes them potential candidates for enhancing oil recovery (Sarkar et al., 1989).

*Brevundimonas aurantiaca* is a gram negative, rod shaped, aerobic bacteria. The *Brevundimonas* are a genus of proteobacteria. It has short wavelength flagella. It is motile and grows at 30°C. It is mostly found in the rhizosphere region of the soil (Euzeby, 1997).

*Enterobacter cloacae* is a significant Gram-negative, facultative-anaerobic, rod shaped bacterium, frequently grown at 30°C on nutrient agar or broth. It bears peritrichous flagella. *Enterobacter cloacae* have been used in a bioreactor based method for the biodegradation of explosives and in the biological control of plant diseases (Dalben et al., 2008).

Biosurfactants, lead to an increasing interest as alternatives to chemical surfactants. Chemical surfactants are widely used in industries for cleanup of oil spills and heavy metal pollutants, enhancement of oil recovery, removal of oil sludge from storage tank, but these chemical surfactants are non-biodegradable and toxic for environment; while if, biosurfactant will be used instead of chemical surfactant these are biodegradable, low toxic, eco-friendly, compatible with human skin, environmentally acceptable and have ability to produced from cheap substrates like vegetable oils such as sunflower and soybean oils, oil wastes from vegetable oil refineries and the food industry industrial oil wastes such as tallow, soapstock, marine oils, lard and free fatty acids, plant-derived oils such as jatropha oil, mesua oil, castor oils, ramtil oil and jojoba oil etc. The present study was conducted to demonstrate the isolation, identification and screening of biosurfactant producing bacteria and production of biosurfactant.

**Materials and Methods**

**Place of work**

The present study was conducted at PG Laboratory, Department of Microbiology Industrial Microbiology, Jacob Institute of Biotechnology and Bioengineering, Sam Higginbottom University of Agriculture, Technology and Sciences, Allahabad, UP during July 2015 to June 2016.

**Isolation of biosurfactant producing bacteria**

Biosurfactant producing bacteria *Enterobacter cloacae* KY231211 and *Brevundimonas aurantiaca* KY231210 were isolated from the rhizospheric soil collected from agricultural field of SHUATS. Serial ten-fold dilutions were prepared from the soil sample and 1 ml of each dilution was added onto nutrient agar plates.

The plates were incubated for 24 hours at 30°C. Colonies with different morphological appearances was selected from the countable plates and re-streaked on a new plate containing the same media to obtain pure colonies. The isolates obtained in this manner were maintained on nutrient agar slants.
Identification of bacterial isolates

Isolates Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210 were identified by cultural, morphological, biochemical and molecular characterization. For colony morphology of isolates shape, edge, elevation, surface and pigmentation was recorded and for cellular morphology was cell shape and Gram staining was done. Biochemical characterization was done by using oxidase test, catalase test, indole test, citrate utilization assay, nitrate reduction test, urease test, esculin hydrolysis and sugar fermentation tests and molecular characterization was done by Sanger sequencing.

Screening of isolates for biosurfactant production

A loopful culture of Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210 was introduced to five ml Bushnell Haas broth in a flask and allowed to grow at 35°C for 10-15 days at 160 rpm. Cells in the flask were harvested through centrifugation at 6000 rpm for 15 minutes and supernatant was used as biosurfactant solution. The following tests were done to check the potency of biosurfactant production of isolates:

Oil displacement assay

30ml of distilled water was taken in a Petri-plate. 1ml of Sunflower oil was added to the centre of the plates containing distilled water. Then 20µl of the supernatant of the culture of isolates was poked into the oil drop.

An uninoculated oil drop in another petri-plate served as a negative control. The biosurfactant producing organisms displace the oil (increases in diameter) and spread in water (Anandraj and Thivakaran, 2010).

Blood haemolysis assay

Hemolytic activity was tested using Blood agar plate. Blood agar medium was prepared using sheep blood (5%) and blood agar base. Blood agar base was sterilized by autoclaving at 121°C at 15 lbs pressure for 15 min. The medium was poured into the plates and after solidify the medium; in center of each plate one well was bored with borer (6mm). Then culture of Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210 were placed in wells in separate plate. The plates were kept for incubation at 30°C for 48-72 hours and observed for zone of clearance around the colonies. The clear zone indicated the presence of biosurfactant producing organisms (Carrillo et al., 1996).

Determination of emulsification index for selective vegetable oils by biosurfactant producing isolates

Emulsification is a process in which an emulsion is formed, an emulsion being liquid containing fine droplets of another liquid without forming a solution. Emulsification activity was performed by adding 2ml of Sunflower oil and Soybean oil to 2 ml of aqueous solution of the isolates. The emulsion stability was determined at 0, 24, 48, 72 and 96h of incubation. The emulsification index was generated as E0, E24, E48, E72, and E96 (EI) at 0, 24, 48, 72 and 96 h to check the stability of the emulsion formed (Anandraj and Thivakaran, 2010) and calculated by following formula:

\[
\text{Emulsification Activity} = \frac{\text{Height of emulsion}}{\text{Total height}}
\]

\[E0=\text{emulsification index at 0h}\]
\[E24=\text{emulsification index after 24h}\]
\[E48=\text{emulsification index after 48h}\]
\[E72=\text{emulsification index after 72h}\]
\[E96=\text{emulsification index after 96h}\]
Production, extraction and partial purification of biosurfactant

Biosurfactant production was conducted in submerged batch mode. Bushnell Haas broth was used as the production medium for the biosurfactant.

The sterilized medium was seeded with 24-48h old bacterial culture and kept in a shaking incubator maintained at 35\(^\circ\)C for 10 days interval at 160 rpm. After 10 days the broth contained the biosurfactant that was further extracted at two sub stages viz. crude extraction of biosurfactant and solvent extraction of biosurfactant (partially purified biosurfactant).

**Crude extraction of biosurfactant**

The broth culture was centrifuged at 10,000 rpm for 30 minutes at 4\(^\circ\)C and supernatant was collected. The pellet was discarded that contained the bacterial cell fractions.

The supernatant was collected as crude biosurfactant and was further purified through solvent extraction method.

**Solvent extraction (chloroform: methanol)**

The supernatant was subjected to acid precipitation by adding concentrated HCl (drop wise) to achieve a final pH of 2.0 and kept at 4\(^\circ\)C overnight. The biosurfactant was extracted with mixed solvent system *i.e.* chloroform: methanol in 2:1 ratio.

Results and Discussion

**Isolation of biosurfactant producing bacteria**

Different bacterial isolates were isolated from rhizospheric soil and further identification was done.

Identification of isolates

Isolates were identified by as *Brevundimonas aurantiaca* KY23121 and *Enterobacter cloacae* KY23121 by cultural, morphological, biochemical and molecular characterization (16S rRNA sequencing).

**Cultural, morphological and biochemical characterization of Enterobacter cloacae KY231211**

The colony of *Enterobacter cloacae* KY231211 was white circular, with irregular margin, flat elevation and yellow to brown pigmentation. *Enterobacter cloacae* KY231211 is Gram negative rods and Catalase positive, oxidase negative, citrate positive, Esculin hydrolysis negative and showed negative for salicin and ribose sugar fermentation (Table 1; Fig. 1).

**Cultural, morphological and biochemical identification of Brevundimonas aurantiaca KY231210**

The colony of *Brevundimonas aurantiaca* KY231210 was orange circular, with round margin, convex elevation and no pigmentation. It is a Gram negative rod and Catalase negative, citrate positive, Esculin hydrolysis negative and showed negative for salicine and ribose sugar fermentation (Table 1; Fig. 2).

**Screening of Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210 for biosurfactant activity**

**Oil displacement assay**

Oil displacement assay is a screening technique to identify biosurfactant activity by microorganism or by biosurfactants. Displacement of oil was measured for two different concentrations *viz.* 10 µl/drop and
20µl/drop oil. In each case displacement of oil was more by 20 µl broth/drop oil. *Brevundimonas aurantiaca* KY231210 showed highest zone of oil displacement in comparison with *Enterobacter cloacae* KY231211 in all cases (Table 2).

**Blood haemolysis assay**

Blood haemolysis pattern of broth culture of *Enterobacter cloacae* KY231211 and *Brevundimonas aurantiaca* KY231210 were examined on Sheep Blood agar medium. Both broth culture (24 h old) of *Enterobacter cloacae* KY231211 and *Brevundimonas aurantiaca* KY231210 revealed β-haemolysis pattern on Blood agar plates. Zone of blood haemolysis was recorded higher with *Enterobacter cloacae* KY231211 in compared with *Brevundimonas aurantiaca* KY231210 (Table 3).

**Emulsification index**

Soybean oil gave highest emulsion at 0h incubation in comparison to Sunflower oil at 0h in case of *Enterobacter cloacae* KY231211, but in case of *Brevundimonas aurantiaca* KY231210 culture emulsification Sunflower oil gave highest emulsion at 0h incubation in comparison to Soybean oil at 0h respectively. Emulsification index was decreased with incubation time in all cases (Table 4; Fig 3).

**Production, extraction and partial purification of biosurfactant**

Biosurfactant was produced through submerged batch mode fermentation using 2% Mustard oil and Sunflower oil as a SSCU supplemented to Bushnell Hass broth seeded with Cfu/ml bacterial inoculums of *Enterobacter cloacae* KY231211 and *Brevundimonas aurantiaca* KY231210 followed by incubation 30±5°C for 15-20 days at 160 rpm. Yield of PPBS was measured after solvent extraction and drying process (Fig. 4).

**Yield of biosurfactant in g/l of production medium**

Yield was obtained after 15-20 days of incubation. Among the two isolates *Brevundimonas aurantiaca* KY231210 gave highest yield (4.31g/l) followed by yield of *Enterobacter cloacae* KY231211 (2.94g/l) (Fig. 5).

The single screening method is unsuitable for identifying all types of biosurfactant, and recommended that more than one screening method should be included during primary screening to identify potential biosurfactant producers (Kiran *et al.*, 2010 and Satpute *et al.*, 2008). Therefore, hemolytic activity assay, oil displacement assay, and emulsification activity measurement were used to screen the biosurfactant producer. If biosurfactant is present in supernatant, oil is displaced and a clearing zone is formed. The diameter of this zone on the oil surface correlates to surfactant activity. For pure biosurfactant a linear correlation between quantity of surfactant and clearing zone is obtained. The oil displacement assay was used as indicator for biosurfactant production for the screening of biosurfactant microorganisms (Anandaraj and Thivakaran, 2010; Priya and Usharani 2009; Urum *et al.*, 2004).

Positive oil displacement was recorded in present study. This result was correlated with study of Bhat *et al.*, (2015) and Rodrigues *et al.*, (2006); whereas in blood haemolysis assay, β haemolysis was recorded with both broth cultures. There is an association between hemolytic activity and surfactant production and because of this blood agar lysis is used for a primary method to screen
biosurfactant production (Carrillo et al., 1996). Emulsification activities (E24) are one of the criteria to determine the potential of biosurfactant and determine the productivity of bio-emulsifier (Bonilla et al., 2005).

In the present study soybean oil and sunflower oil were used for emulsification and highest emulsification index was recorded in 0h of incubation. Emulsification index was recorded in decreasing order when incubation periods were increased.

The production of biosurfactant from various carbon sources such as glycerol, glucose and hydrocarbons has been reported earlier by Jayanti and Joshi (1992) (Fiebig et al., 1997; Haba et al., 2000; Raza et al., 2007). In present study mustard oil and sunflower oil were used for biosurfactant production.

These substrates are low priced, high in purity and as hydrophobic substrates they can possibly enhance the production of biosurfactant (Van Hamme et al., 2006). For extraction of biosurfactant centrifugation, sedimentation and organic solvent, chloroform and methanol was used (Peter et al., 2014). also reported the solvent extraction method for partially purification of biosurfactant and obtained highest yield of biosurfactant in Serratia spp.

**Table.1 Taxonomic identification of Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210**

<table>
<thead>
<tr>
<th>S.no.</th>
<th>Characteristics</th>
<th>Enterobacter cloacae</th>
<th>Brevundimonas aurantiaca</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>KY231211</td>
<td>KY231210</td>
</tr>
<tr>
<td></td>
<td>Cultural characteristics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.</td>
<td>Colour of colony on Nutrient agar</td>
<td>White</td>
<td>Orange</td>
</tr>
<tr>
<td>2.</td>
<td>Shape of colony</td>
<td>Circular</td>
<td>Circular</td>
</tr>
<tr>
<td>3.</td>
<td>Elevation</td>
<td>Flat</td>
<td>Convex</td>
</tr>
<tr>
<td>4.</td>
<td>Margin</td>
<td>Irregular</td>
<td>Round</td>
</tr>
<tr>
<td>5.</td>
<td>Pigmentation</td>
<td>Yellow to brown</td>
<td>No pigmentation</td>
</tr>
<tr>
<td></td>
<td>Morphological Characteristics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.</td>
<td>Gram’s reaction</td>
<td>Negative</td>
<td>Negative</td>
</tr>
<tr>
<td>2.</td>
<td>Shape</td>
<td>Rod</td>
<td>Rod</td>
</tr>
<tr>
<td>3.</td>
<td>Catalase test</td>
<td>Positive</td>
<td>Positive</td>
</tr>
<tr>
<td>4.</td>
<td>Oxidase test</td>
<td>Negative</td>
<td>Negative</td>
</tr>
<tr>
<td>5.</td>
<td>Citrate test</td>
<td>Positive</td>
<td>Positive</td>
</tr>
<tr>
<td>6.</td>
<td>Escculin hydrolysis</td>
<td>Negative</td>
<td>-</td>
</tr>
<tr>
<td>7.</td>
<td>Indole test</td>
<td>-</td>
<td>Negative</td>
</tr>
<tr>
<td>8.</td>
<td>Nitrate reduction test</td>
<td>-</td>
<td>Negative</td>
</tr>
<tr>
<td>9.</td>
<td>Urease test</td>
<td>-</td>
<td>Negative</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate fermentation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.</td>
<td>Salicin fermentation</td>
<td>Negative</td>
<td>-</td>
</tr>
<tr>
<td>2.</td>
<td>Ribose fermentation</td>
<td>Negative</td>
<td>-</td>
</tr>
<tr>
<td>3.</td>
<td>Arabinose fermentation</td>
<td>-</td>
<td>Positive</td>
</tr>
<tr>
<td>4.</td>
<td>Xylose fermentation</td>
<td>-</td>
<td>Negative</td>
</tr>
</tbody>
</table>
Table.2 Oil displacement assay by broth culture

<table>
<thead>
<tr>
<th>Name of bacteria</th>
<th>Sunflower oil</th>
<th>Soybean oil</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10 μl</td>
<td>20 μl</td>
</tr>
<tr>
<td><strong>Enterobacter cloacae KY231211</strong></td>
<td>6.15</td>
<td>8.8</td>
</tr>
<tr>
<td><strong>Brevundimonas aurantiaca KY231210</strong></td>
<td><strong>8.8</strong></td>
<td><strong>12</strong></td>
</tr>
</tbody>
</table>

Table.3 Blood haemolysis assay by broth culture

<table>
<thead>
<tr>
<th>Name of bacteria</th>
<th>Zone of haemolysis (mm)</th>
<th>Type of haemolysis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Enterobacter cloacae KY231211</strong></td>
<td>5</td>
<td></td>
</tr>
<tr>
<td><strong>Brevundimonas aurantiaca KY231210</strong></td>
<td>4.55</td>
<td></td>
</tr>
</tbody>
</table>

Table.4 Emulsification index for biosurfactant producing culture with Sunflower oil and Soybean oil

<table>
<thead>
<tr>
<th>Oil</th>
<th><strong>Enterobacter cloacae KY231211</strong></th>
<th><strong>Brevundimonas aurantiaca KY231210</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>E₀</td>
<td>E₄₀</td>
</tr>
<tr>
<td>Sunflower</td>
<td>22.5</td>
<td>18.75</td>
</tr>
<tr>
<td>Soybean</td>
<td><strong>55</strong></td>
<td><strong>47.5</strong></td>
</tr>
</tbody>
</table>

Fig.1 Phylogenetic tree of *Enterobacter cloacae KY231211*
**Fig.2** Phylogenetic tree of *Brevundimonas aurantiaca* KY231210

![Phylogenetic tree](image)

**Fig.3** (a) and (b) Emulsification index for biosurfactant producing culture with Sunflower oil and Soybean oil

(a) *Enterobacter cloacae* KY231211  
(b) *Brevundimonas aurantiaca* KY231210
In conclusion, in this present study, biosurfactant is produced by using rhizospheric bacterial isolates i.e. Enterobacter cloacae KY231211 and Brevundimonas aurantiaca KY231210. Selected microorganisms showed positive result for all the method use for screening. Brevundimonas aurantiaca KY231210 is found to be higher yield of biosurfactant as compared to Enterobacter cloacae KY231211.

References


Satpute, S.K., Bawsar, B.D., Dhakephalkar,


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