Isolation and identification of azo dye degrading microbes using 16s r RNA sequencing

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Abstract

The aim of the present study was to isolate and identify azo dye degrading microbes using 16s r RNA sequencing. For this, ten effluent and fifty soil samples were collected from textile industry from a discharge panel of textile industries near Peenya and Magadi road Industrial area, Bengaluru, and dying industries near Kerur, Bagalkot, Karnataka, India. The pH, temperature, BOD, COD, Odor, Total dissolved solid (TDS), Total suspended solids (TSS), Chemical oxygen demand (COD), Biological oxygen demand (BOD), Dissolved Oxygen (DO), and Total Hardness values were very high when compared with the values given by the Bureau of Indian Standards. Pure cultures were screened on the basis of colony morphology. Three different types of unique cultures were selected and named as isolates S1, S2 and S3. Outof 6 dyes viz. Corafast blue, Corafast red, Red 3BN, Solophenyl Brown, RemazoleRG , Rubin M2B used, isolate S1 showed degradation on the maximum number of dyes Red 3BN, Corafast blue, Corafast red, in comparison to other isolates (isolates S2 and S3). Thus, isolate S1 was used for the further studies. For this study Red 3BN dye was chosen. The isolated bacterium was gram positive Bacilli. In the biochemical characterization, the isolate was partially confirmed as Bacillus sp. Further, the selected isolate was identified by sequencing the 16S rRNA gene, partial sequence of the isolated strain and species related to Bacillus species in the gene bank database. Thus, this organism may be used significantly in effluent treatment such as textile, paper, ink and other industries.

Keywords: Azo dye, Bacillus species, Degradation, Effluent, Fermentation, Pure culture

INTRODUCTION

Azo dyes have been widely used as dyestuffs in cosmetics, foods, printing and textile materials. These dyes are the largest class of commercially available synthetic dyes and wide applications in textiles, food, cosmetics, plastic, laboratories, leather, paper printing, colour photography, pharmaceutical and toy industries(Mathur and Bhatnagar, 2007; Pant et al., 2008; Laowansiri et al.,2008). Several physico-chemical techniques have been proposed for treatment of colored textile effluents. These include adsorption on different materials, oxidation and precipitation by Fenton’s reagent, bleaching with chloride or ozone photo degradation or membrane filtration (Andre et al.,2007) coagulation, flocculation, flotation, electrochemical destruction and mineralization (Gogate and Pandit, 2004). All these physical or chemical methods are very expensive and result in the production of large amounts of sludge, which creates the secondary level of land pollution. Therefore, Bioremediation through microorganisms has been identified as a cost effective and environment friendly alternative for disposal of textile effluent (Robinson Tim et al., 2001). Microbial degradation of azo dyes is mediated by enzymes. The predominant enzymes are azoreductase, laccases, lignin peroxidase, manganese peroxidase, and hydroxylases. Laccase
and azoreductase have been shown to degenerate azo dyes. But the stable operation of continuous fungal bioreactors for the treatment of synthetic dye solutions have been achieved, application of fungi for the removal of dyes from textile wastewaters faces many problems such as production of large volumes, the nature of synthetic dyes, and control of biomass (Nigam et al., 2000; Robinson et al., 2001; Stolz, 2001). The bacterial reduction of the azo dye is usually nonspecific and bacterial decolorization is faster (Sudha et al., 2014). The objective of the present study was to analyze the physico-chemical characterization of textile dye effluents and isolate and characterize the dye degrading bacteria from dye effluents and soil.

**MATERIALS AND METHODS**

**Sample collection:** For the isolation of dye degrading microorganisms, 50 soil samples were collected in sterilized ziplock polythene covers and 10 effluent samples in sterilized screw capped bottles from a discharge panel of textile industries near Peenya and Magadi road industrial area, Bengaluru, and dying industries near Kerur, Bagalkot, Karnataka, India. Collected effluent samples were named as S1, S2, S3, S4, S5, S6, S7, S8, S9, S10 and brought to laboratory.

**Physico-chemical property analysis:** The collected effluent samples were analyzed to determine their physico-chemical parameters. The various parameters such as Temperature, pH, Color, Odor, Total dissolved solid (TDS), Total suspended solids (TSS), Chemical oxygen demand (COD), Biological oxygen demand (BOD), Dissolved Oxygen (DO), and Total Hardness were analyzed in the laboratory by the standard protocol (Aneja, 2003).

**Isolation of bacteria from soil and effluent samples:** The organism was isolated from soil and effluent samples by pour plate method. In this process soil samples were dissolved in 8% saline and effluent samples were pipetted into sterile test tubes, 15ml of minimal agar media (Potassium dihydrogen phosphate 3.0g, Disodium hydrogen phosphate 6.0g, Sodium chloride 0.5g, Magnesium sulphate 0.1g, Agar 15g, Distilled water 1000ml) along with different dyes such as Corafast blue, Corafast red, Red 3BN, Solophenyl Brown, Remazole RG, Rubin M2B and incubated at 37°C for 24 hours. After incubation, maximum degradation showed colonies were inoculated into the minimal broth and analyzed by Spectrophotometric method. For this process, 100ml of minimal broth along with the dyes (10mg/100ml) was inoculated with loopful of isolated bacterial culture. The culture flasks were incubated on orbital shaker with 150rpm, at 37°C. Flasks without inoculum kept as control. Absorbance values were measured Spectrophotometrically at 580nm for blue dye and 530nm for Red dye to estimate the decolorization process. The rate of decolorization was calculated using the following formula:

\[ \text{% Decolorization} = \frac{\text{initial absorbance value - final absorbance value}}{\text{initial absorbance value}} \times 100 \]  

**Identification of bacteria:** Three different bacterial isolates showed degradation and were named as S1, S2 and S3. Among these S1 showed maximum degradation and was identified on the basis of morphological, biochemical and 16s rRNA sequence.

**Morphological characterization of isolated colonies from collected samples:** The morphological characterization gave an idea of the size, shape, arrangement of bacterial cells.

**Biochemical characterization of isolated bacteria:** Colonies were subjected to biochemical tests such as Indole production, Methyl red, Voges-Proskauer, Citrate utilization, Triple sugar iron utilization, Carbohydrate fermentation tests such as Glucose, lactose and enzymatic tests such as Starch hydrolysis, Gelatin hydrolysis, Casein hydrolysis, Urease utilization, Catalase production, Oxidase production, Nitrate reduction tests by following the methodology of Holding and Collee (1971).

**Molecular identification of the bacterial isolate:** Isolation of Genomic DNA: 25ml of LB broth was prepared, isolated organism was inoculated and incubated at 37°C for 24 hours. After the incubation 1ml of culture was taken in an appendorr tube, centrifuged at 6000rpm for 10 minutes. Pellets were taken and 500µl of lysis buffer (50mMTris HCL, 20mM EDTA, 1.25% SDS) added. Gently vortexed and incubated at 60–65°C for one hour. Allowed to room temperature and 200µl of 5 M Sodium acetate was added. Incubated in cold conditions for 10 minutes and centrifuged at 10000rpm for 10 minutes. Then, the double volume of ethanol was added to the collected supernatant and incubated for 10 minutes at room temperature. Centrifuged at 12000rpm for 10 minutes. Pellets were collected, dissolved in 50 µl of TE buffer (10mM TrisHCl, 1mM EDTA).

**PCR reaction:** The molecular signatures of the isolate were studied by partial sequencing of 16S rRNA gene and comparing it with known sequences. The set of universal primers; and
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F5’AGAGTTTGATCCTGGCTCAG3’R5’TACGGTTACCTTGTTACGACTT3’ was used to amplify the desired sequence.

The following reaction mixture was used to amplify the 16S rRNA of isolates. The total reaction volume was 50 μl for each reaction. The PCR reaction mixture was as follows: DNS sample 3μl, 5X Phusion HF buffer 10μl, and Forward primer2μl (0.05 μM) Reverse primer 2μl(0.05 μM), dNTPs (200μM) 5 μl, Phusion Polymerase 0.5 l Distilled water29.5μl.

PCR conditions: PCR amplification was performed in 95˚C. Different annealing temperatures were tested (from 46 to 56 ˚C). The best annealing temperature was 30 second at 55˚C where it produced only one high intense DNA ampilicon. The optimum program was one cycle at95˚C for five minutes, and then 30cycles were performed as follows: thirty seconds at 98˚C for denaturation, thirty seconds at 55˚C for annealing, one minute at 72˚C for elongation, and then 10 minutes at 72˚C for final extension. The reaction mixtures were held at 4˚C until used. After completion of PCR run 5μl PCR product on 1.2% agarose gel to confirm amplification and the amplified sample was sent to Eurofins Genomics India Pvt Ltd Bangalore for Sequencing.

RESULTS AND DISCUSSION

Total dissolved solids were present in S8 (3200 mg/l) and S9 (1279mg/l) samples respectively which were above the BIS standards (2100mg/l). High concentration of dissolved solids affects the density of water and influences solubility of gases like oxygen in water and osmoregulation of freshwater organisms (Sriram et al 2015, Thoker Farook et al 2012). The minimum total suspended solids (TDS) were recorded at S5 (120 mg/l) while the maximum at S10 (210 mg/l), the standard limit prescribed is 100mg/l. The maximum chemical oxygen demand (COD) were observed from S7 (929 mg/l) and minimum in S5(667mg/l) which was much higher than maximum recommended limit of Federal Environmental Quality and Bureau Indian Standards, it’s impacted the receiving water body to some extent and its effects on the quality of freshwater and subsequently cause harm to aquatic life (Thoker Farook et al., 2012, Manikandan et al., 2012,Rajeswariet al., 2013). The maximum biological oxygen demand (BOD) was observed from S5 (310 mg/l) and minimum in S8 (123mg/l) and the standard recommendation (BIS) is 350mg/l. Dissolved oxygen is a fundamental requirement for aquatic life (Sofia Nosheen 2000). The maximum dissolved oxygen (DO) was recorded in S9 (190 mg/l) and minimum in S3 (106mg/l) samples comparatively with other effluent samples (Table 1).

Isolation of bacteria from soil samples: About 50 different bacterial isolates were isolated from
dye contaminated soil and effluent samples.

**Identification of bacteria:** In the present study, bacterial strains were isolated from effluents and soil samples near textile industries. On the basis of morphological, biochemical and 16s rRNA sequencing, the organism is identified as *Bacillus* species.

**Morphological characterization of isolated colonies:** The morphological characteristics of the isolated colonies were large gram-positive rods, arranged in chains.

**Biochemical characterization of isolated *Bacillus* strains:** Isolate subjected to biochemical tests and liquified gelatin, hydrolyzed starch, casein and lipid. It also fermented dextrose and sucrose (gas production), reduced nitrate, gives positive result to oxidase and catalase activity. It gives negative results for hydrogen sulphide production, Indole production, Methyle red reaction, Vogesproskauer test. Lactose fermentation and Citrate utilization. So by observing the results of biochemical tests the isolate is partially confirmed that belongs to *Bacillus sp.* (Table 2).

Based on morphological and biochemical reactions, the isolated organism belongs to *Bacillus* species. Similarly, HauwaTahir et al., (2016), isolated *Staphylococcus aureus, Bacillus cereus, Escherichia coli* and *Klebsiella pneumonia* and identified based on their colonial, morphological and biochemical characterization. AmminiParavthi et al., (2009) isolated and identified *Bacillus pumilus* by detailed conventional biochemical methods, fatty acid methyl ester (FAME) analysis. Ali Ramadan Ali (2014) identified the bacteria based on molecular and biochemical characterization of bacterial isolates from food stall vegetables. Azharet et al. (2017) identified cellulose degrading bacteria *Anoxybacillus flavithermus*, Van Veen et al. (2009) identified *Staphylococci* species and Sujatha et al. (2012) identified *Bacillus thuringiensis*, *Streptococcus pneumoniae* and Beuving et al. (2011) identified b-hemolytic streptococci that belong to the viridans group. Isabel Sanchez, et al. (2000) identified *Lactobacillus plantarum, L. brevis, L. fermentum, L. brevis, L. spersus*. Molecular identification of the bacterial isolate: The selected isolate was identified by 16S rRNA sequencing. The results cleared that the bacterial isolate is identified as *Bacillus* species exhibiting 99% similarity. Fig.1 shows the phylogenetic relationship of 16S ribosomal RNA gene, partial sequence of the isolated strain and species related to *Bacillus* species in the gene bank database. Similarly, Azharet et al., (2017) identified cellulose degrading 4 bacterial isolates such as isolate 1 as 99% similarity with *Anoxybacillus flavithermus*, isolate 2 has 99% similarity with *Bacillus megaterium*, isolate 3 has 99% similarity with *B. amyloliquefaciens* and isolate 4 was 99% similar with *Bacillus subtilis*. Trupti (2009) isolated *Tributylphosphate degrading organism* and sequenced and compared using BLAST, Clusta IW and PHYLIP, and the organism was identified as *Pseudomonas alcaligenes* strain DSM50018T while Sujatha et al. (2012) identified *B. thuringiensis* strain IAM 12077 using 16s rRNA sequencing.

**Conclusion**

The industrial effluents and soils contained enriched nutrients to grow and spread microbial population. Bacterial species have at least one copy of the 16S rRNA gene containing highly con-

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**Table 2. Biochemical characterization of Isolated *Bacillus* species.**

<table>
<thead>
<tr>
<th>S.N.</th>
<th>Biochemical tests</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Gelatin liquification</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>Starch hydrolysis</td>
<td>+</td>
</tr>
<tr>
<td>3</td>
<td>Lipid hydrolysis</td>
<td>+</td>
</tr>
<tr>
<td>4</td>
<td>Lactose fermentation</td>
<td>-</td>
</tr>
<tr>
<td>5</td>
<td>Dextrose fermentation</td>
<td>Acid production</td>
</tr>
<tr>
<td>6</td>
<td>Sucrose fermentation</td>
<td>Acid production</td>
</tr>
<tr>
<td>7</td>
<td>TiS</td>
<td>Acid production</td>
</tr>
<tr>
<td>8</td>
<td>H2S production</td>
<td>-</td>
</tr>
<tr>
<td>9</td>
<td>NO3 reduction</td>
<td>+</td>
</tr>
<tr>
<td>10</td>
<td>Indole production</td>
<td>-</td>
</tr>
<tr>
<td>11</td>
<td>Methyle red reaction</td>
<td>-</td>
</tr>
<tr>
<td>12</td>
<td>Vogesproskauer test</td>
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</tr>
<tr>
<td>13</td>
<td>Citrate utilization</td>
<td>-</td>
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<tr>
<td>14</td>
<td>Urease activity</td>
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<tr>
<td>15</td>
<td>Catalase activity</td>
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<tr>
<td>16</td>
<td>Oxidase activity</td>
<td>+</td>
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<tr>
<td>17</td>
<td>Casein hydrolysis</td>
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</tr>
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</table>

Fig. 1. Phylogenic tree of *Bacillus* species.
served regions together with hyper variable regions. The use of 16S rRNA gene sequences to identify new strains bacteria is gaining momentum in recent years. We showed the use of 16S rRNA gene sequence to characterize the bacterial isolate from the textile effluent and soil were found to be Bacillus sp.. Thus, the genotyping method using 16S rRNA gene sequence is both simple and effective in strain identification and the study concluded that the isolate had adequate potential to decolorize the azo dyes- Red and Blue dye. Thus, the isolate could be exploited for its bioremediation ability.

REFERENCES