

1        **BIOREMEDIATION OF INDUSTRIAL EFFLUENT USING CYANOBACTERIAL**

2                    **SPECIES: *Phormidium mucicola* AND *Anabaena aequalis*.**

3                    Sanjay Kumar Dubey\*, Preeti Vyas, Vaishali Gupta and Jaishree Dubey

4                                    Lab of Phycology, Department of Botany,

5                                    Dr. Hari Singh Gour University, Sagar, India - 470003

6                                    E-mail: dubeysanjay83@gmail.com

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8  
9        **ABSTRACT**

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Different Industries discharge effluent in different water bodies, which is the only reason of pollution. The main objective of the present study was to investigate the biodegradation and biosorption capacity of some potential cyanobacterial species; *Phormidium mucicola* and *Anabaena aequalis* in Textile and Pharmaceutical industries, Mandideep, Bhopal Madhya Pradesh, India. Industrial effluents are contaminated with heavy metal. The effluents were subjected to biological treatment using axenic cyanobacterial strains as batch system for 7 days. Removal efficiencies of the different contaminants were evaluated and compared. Results confirmed the high efficiencies of the investigated species for the removal of the target contaminants which were species and contaminant-dependent. BOD and COD recorded 91.18 and 82.54% as maximum removal efficiencies achieved by *Anabaena aequalis*. The highest removal efficiencies of the total suspended solids recorded 53.23% achieved by *Phormidium mucicola*, while 41.61% was recorded as the highest TDS. Concerning the contaminant metals, *Phormidium mucicola* showed the highest biosorption capacity where 86.12 and 94.63% removal efficiencies were achieved for Zn and Cu, respectively. In conclusion, results of the study confirmed the advantageous potential of using the tested cyanobacterial species for the bioremediation of industrial effluent and clearly showed the quality improvement of the discharged effluent which in turn will eliminate or at least minimize the expected deterioration of the receiving environment.

11        **Keywords:** Bioremediation, cyanobacterial species, heavy metals, industrial effluent

12        **1. INTRODUCTION**

13                    Water covers 71% of the earth and makes up 65% of the our bodies. All organisms  
14                    contain it, live in it, drink it. Plants and animals require water that is moderately pure and  
15                    they can not survive if their water is loaded with toxic chemical and harmful microorganisms  
16                    (Encarta Encyclopedia, 2009). Effluents discharge from different sources is directly

17 responsible for water pollution. This is mainly due to continuous discharge of huge quantities  
18 of the effluents lead to deterioration in the water quality of the rivers (El-Bestawy E., 1993;  
19 El-Bestawy E. et al., 2007; Mansy And El-Bestawy, 2002). Any change in pH of water bodies  
20 as a result of influx of effluent; can cause serious change in water chemistry, which can  
21 affect resources especially around the river area. These effects on water bodies can be very  
22 significant. Besides nutrients, the river water and sediments show terrible levels of organic  
23 matter, and heavy metals in worldwide. To overcome water pollution problem some physical  
24 and chemical methods are used, but they are expensive and generative concentrated waste  
25 that require subsequent treatment and disposal (Komori et al., 1990). Traditional methods for  
26 the cleanup pollutants usually involve the removal of unwanted materials through  
27 sedimentation and filtration, and subsequent chemical treatment such as flocculation,  
28 neutralization and electro-dialysis before disposal. Biological treatment may provide a  
29 suitable means for treatment from waste water (Lovely and Coates 1997 and Rittmann et al.,  
30 2004). Cyanobacterial species characterized by great ability to tolerate such high levels of  
31 pollution and proved high efficiency for degrading highly organic contaminants and  
32 accumulating heavy metals (El-Bestawy et al.,2007; Podda et al.,2000 and Palmer, 1980).  
33 Therefore, they could be efficiently used in advanced technologies for bioremediation of the  
34 industrial effluents.

35 Cyanobacteria group is one of the most diverse groups of prokaryotes. They are  
36 gram-negative oxygenic photosynthetic bacteria. They can be found in a wide range of water  
37 bodies especially where the excess nutrient input of anthropogenic activities. Many species  
38 of cyanobacteria have nitrogenase enzyme, they convert atmospheric nitrogen into  
39 ammonium. Their photoautotrophic nature and the ability of some species to fix atmospheric  
40 nitrogen enable them to be producers, as opposed to consumers, and make their growth and  
41 maintenance inexpensive [Castenholz et al., 1989; Somashekar, and Ramaswamy, 1983].  
42 Metabolic activities are not affected by the decrease in the levels of the biodegradable  
43 pollutants that they may break down. Cyanobacteria have been used efficiently as a low-cost  
44 method for remediating all industrial effluents as well as transformation and removal of  
45 heavy metals (Lefebvre et al., Budd 2007; Podda et al.,2000).

46 Recently, there have been increasing interest about using cyanobacteria as pollution control  
47 agents since they possess many advantages over other microorganisms isolated from soil.  
48 Remediation capabilities of cyanobacteria toward environmental pollutants can be improved  
49 and enhanced through genetic engineering technologies (Kuritz and Wolk 1995; Mansy and  
50 El-Bestawy, 2002 and Palmer, 1980). However, the beneficial application of cyanobacteria in  
51 remediation of contaminated waters and industrial effluents is still not optimally manipulated  
52 (Jeganathan, 2006 and Kannan, 2006).

53 **2. AIMS AND OBJECTIVES**

54 The main objective of the present study was to investigate the remediation capacity of some  
55 potential cyanobacterial species isolated from Textile and Pharmaceutical industrial effluent;

56 **3. MATERIAL AND METHODS**

57 Survey of different sites of industrial effluent for identification of different algal forms from  
58 taxonomic point of view will be undertaken. Samples were collected from two industries  
59 (Textile and pharmaceuticals industries, from Mandideep, Bhopal, M.P.), India. Effluents  
60 were collected in large sterilized containers and polythene bags respectively. Thus, it is  
61 expected that the effluents contain industrial pollutants such as heavy metals which are not  
62 likely to be removed by that primary treatment of the industries. Grab samples representing  
63 all effluents entering the plant during 24 h were collected from both plants to avoid the  
64 fluctuation in the flow and the strength of the effluent.

65 Physico-chemical characteristics of effluents were carried out by standard methods (APHA,  
66 1995). Such as biochemical oxygen demand (BOD); chemical oxygen demand (COD); total  
67 suspended solids (TSS); total dissolved solids (TDS); and Textile and pharmaceuticals  
68 effluents contain high Zn and Cu were characterized before and after treatment to determine  
69 the effectiveness of the remediation process (Bhoominathan 2005, Cairns, and Dickson,  
70 1971). All the investigated parameters were determined using the standard techniques  
71 described by (Celesseri et al., 1999) in the standard methods for the examination of water  
72 and effluent.

73 Standard microbiological methods were followed for the isolation of cyanobacteria. Algal  
74 samples were microscopically examined and the selected cyanobacterial species were  
75 grown in Chu No.10 (1942) media was used as culture media. and the pH was adjusted to  
76 7.2 with HCl. Media has sterilized by autoclaving separately at 121<sup>0</sup>C for 20 min.  
77 Micronutrient solution was sterilized by filtration through 0.22 mm polycarbonate membrane  
78 to avoid interaction and precipitation of heavy metals. Chu No.10 media was freshly  
79 prepared from A and B where 1.0 ml of each component of solution A and 1.0 ml of Solution  
80 B were combined and diluted to 1.0 l, sterilized as mentioned and used for selective culturing  
81 of the selected species. After inoculation, all the selected species were incubated at room  
82 temperature (28<sup>0</sup>C) and day light with manual shaking every 24 h to avoid adhesion of the  
83 algae on the walls of the glass vessels until heavy growth appeared within 3 weeks.  
84 Identification was confirmed based upon the keys given by (Geitler, 1932 and Desikachary,  
85 1959) for microscopic parameters. The isolated cyanobacteria were identified with the help  
86 of classical manuals. Two different cyanobacterial species; *Anabaena aequalis* and  
87 *Phormidium mucicola*; were selected for further study.

## 88 **2.1 Axenity and bioassay**

89 Unialgal cultures usually remained contaminated with bacteria and therefore to free them  
90 from bacteria is a pre requisite for further studies (Ash and Jenkins, 2006; Anagnostidis, El-  
91 Nahhal et al, 2013, Komárek, 1985 and Safi et al, 2014). The cultures were made bacteria  
92 free by ultraviolet irradiation for varying periods and inoculated in the medium. Axenic  
93 cultures were prepared of these isolate species. However, before using these strains in the  
94 bioremediation of the contaminated industries effluents, their axenity was checked using  
95 agar phototactic response method. Semi solid standard agar medium was prepared and  
96 liquated into test tubes and sterilized. Each tube was inoculated with 100 µl of  
97 cyanobacterial culture (two replicates per culture). Light was prevented to reach the top 10  
98 cm of the tube using aluminum foil. All the tubes were incubated in optimal conditions (28<sup>0</sup>C)  
99 in an illuminated incubator. Based on the phototactic response phenomena the  
100 cyanobacterial filaments were grown toward light direction through the semi solid agar, but  
101 bacteria did not grown. After 7days incubation of the agar column was dragged out the tube  
102 on sterilized Petri dish. The agar column was sliced into ten slices 1 cm per each. Each slice  
103 was stranded longitudinally and transversally cut under common sterilized conditions to  
104 separate the algal filaments surrounded with a small piece of agar. Each agar piece  
105 involving cyanobacterial growth was inoculated into standard selective liquid medium. After  
106 incubation, each inoculated culture was tested for contamination using general bacterial  
107 medium (nutrient agar). In bioremediation bioassay, the tested species were checked for  
108 their axenity, and the liquid cultures were tested by plating on bacterial nutrient medium and  
109 incubating at 28<sup>0</sup>C for 7days. Only axenic cultures were involved in the assays. The selected  
110 species were inoculated individually in 100 ml culturing medium (three replicates) and  
111 incubated for 2 weeks till heavy growth was obtained. Effluents from both industries (textile  
112 and pharmaceutical) was dispensed (900 ml each) in 18 sterilized conical flasks, nine flasks  
113 for each effluent. Each culture (100 ml) was separately seeded at a final volume of 1l each  
114 (three replicates/strain/effluent) and incubated under the previously mentioned conditions for  
115 7 days. Another six flasks (three flasks for each industry) were supplied by 1ml each of the  
116 effluent of both industries without seeding with cyanobacteria to serve as control for the  
117 bioassay (each experiment has been performed in triplicate). They were incubated under the  
118 same conditions. For the determination of heavy metals and other parameters residues,  
119 samples were collected at 24 h interval. At each sampling time, 130 ml from each flask were  
120 aseptically drawn, where all the investigated parameters were determined and their removal  
121 efficiencies using the selected species were determined by PERKIN ELMER OPTIMA 5300  
122 DV ICP-OES (Inductively Coupled Plasma Optical Emission Spectrophotometer) method

123 and result were obtained from SAIF (Sophisticated Analytical Instrument Facility) Indian  
 124 Institute of Technology Madras, Chennai.

125 **3. RESULTS AND DISCUSSION**

126 Following species have been collected from Textile and pharmaceutical industries.

127 **Table 1: Occurrence of cyanobacteria in effluents of different sites of**  
 128 **Pharmaceutical, Textile industries**

129

Cyanobacterial species	Industries					
	Pharmaceutical			Textile		
	Site I	Site II	Site III	Site I	Site II	Site III
<i>Chroococcus disperses</i>	+	+	+	+	+	+
<i>Rhabdoderma irregular</i>	-	-	-	-	-	+
<i>Aphanocapsa elachista</i>	-	-	-	+	-	-
<i>Holopedium irregular</i>	-	-	-	+	-	-
<i>Gloeotheca linearis</i>	-	-	-	-	-	-
<i>Gomphosphaeria aponina var.cordiformis</i>	-	-	-	+	-	-
<i>Microcystis aeruginosa</i>	+	+	+	+	+	+
<i>Spirulina major</i>	+	+	+	-	-	-
<i>Oscillatoria amoena</i>	-	-	-	+	+	+
<i>Phormidium inundatum</i>	+	+	+	+	+	+
<i>Phormidium mucicola</i>	+	+	+	+	+	+
<i>Schizothrix lacustris</i>	+	+	+	-	-	-
<i>Schizothrix tinctoria</i>	-	-	-	-	-	-
<i>Anabaena aequalis</i>	+	+	+	+	+	+
<i>Anabaena affinis</i>	+		+			
<i>Nostoc verrucosum</i>	+	+	+	+	+	+
<i>Nodularia spumigena</i>	-	-	-	-	-	-
<i>Aphanizomenon flos-aquae</i>	+	+	+	+	+	+
<i>Cylindrospermum catenatum</i>	-	-	+	-	-	-
<i>Scytonema crispum</i>	-	+	-	-	-	-
<i>Plectonema wollei</i>	-	-	-	+	+	+
<i>Stigonema mesentericum</i>	-	-	+	+	-	-
<i>Stigonema ocellatum</i>	+	+	+	+	+	+

130 Present (+) Absent (-)

131

132 *Phormidium mucicola* and *Anabaena aequalis* investigated as free-living cells for their ability  
 133 for organic matter biodegradation and heavy metal removal from the effluent. They were  
 134 selected based on their dominance and survival in the highly polluted water of  
 135 Pharmaceutical industries and textile industries where they acquired high resistance and  
 136 acclimatized to deal with high loads of different contaminants. They were also proven high  
 137 ability for degradation of the heavy metals. Therefore, the selected species were considered  
 138 promising candidates for biological treatment of the industrial effluents.

139 Table 2: Residue concentrations (RC) of the quality parameters from the contaminated  
 140 industrial effluents

141 Textile industry effluent (Before treatment value in mg/l)

Time(day)	BOD	COD	TSS	TDS	ZN	CU
Raw water	140	360	167	1,150	0.11	0.04
2	89	289	132	887	0.09	0.01
3	80	243	100	766	0.05	0.02
4	81	200	101	756	0.12	0.04
5	70	187	98	611	0.09	0.05
6	120	387	154	900	0.14	0.06
7	143	331	143	998	0.12	0.07

142

143

144 Pharmaceutical industry effluent (Before treatment value in mg/l)

Time(day)	BOD	COD	TSS	TDS	ZN	CU
Raw water	198	445	387	1430	0.01	0.03
2	187	465	411	1457	0.04	0.01
3	199	412	376	1432	0.03	0.04
4	176	398	311	1200	0.01	0.04
5	166	345	298	1100	0.03	0.06
6	178	411	321	1289	0.05	0.02
7	200	378	365	1008	0.02	0.01

145 Table 3: Residue concentrations (RC) of the quality parameters from the contaminated  
 146 industrial effluents using the selected cyanobacteria at different exposure time icpoes.

147 Textile industries effluent (After treatment value in mg/l)

Days	<i>Anabaena aequalis</i>						<i>Phormidium mucicola</i>					
	BOD	COD	TSS	TDS	ZN	CU	BOD	COD	TSS	TDS	ZN	CU
Raw	149	312	200	1012	0.11	0.05	148	324	234	1134	0.12	0.07

water													
2	141	301	199	1011	0.09	0.05	143	321	223	1124	0.10	0.07	
3	132	298	189	1001	0.06	0.03	132	309	212	1120	0.5	0.05	
4	121	288	170	998	0.05	0.02	122	289	206	1103	0.5	0.03	
5	100	198	150	988	0.03	0.02	101	281	189	1087	0.3	0.02	
6	76	122	132	986	0.03	0.02	98	256	167	1079	0.1	0.1	
7	62	98	112	983	0.02	0.01	78	249	156	1012	0.1	0.1	

148 Pharmaceutical industries effluent (After treatment value in mg/l)

Days	<i>Anabaena aequalis</i>						<i>Phormidium mucicola</i>					
	BOD	COD	TSS	TDS	ZN	CU	BOD	COD	TSS	TDS	ZN	CU
Raw water	254	432	321	1349	0.08	0.07	231	421	401	1401	0.06	0.07
2	233	428	311	1323	0.06	0.05	211	421	387	1400	0.04	0.06
3	212	412	306	1321	0.06	0.04	201	401	381	1387	0.04	0.04
4	198	398	285	1287	0.04	0.04	192	356	372	1345	0.02	0.03
5	167	378	265	1287	0.03	0.02	187	241	324	1302	0.01	0.03
6	121	321	209	1270	0.02	0.01	123	209	283	1265	0.01	0.01
7	98	243	187	1201	0.02	0.00	98	198	223	1230	0.00	0.01

### 149 3.1 Industrial effluent characteristics

150 Effluent produced by the two industries were characterized (Table2, control).BOD, COD,  
 151 TSS, TDS, Zn and Cu recorded averages of 140, 360, 167, 1150, 0.11and 0.04 mg/l,  
 152 respectively, in the effluent of the textile industry. Significantly higher levels for almost all the  
 153 tested parameters were detected in the pharmaceutical effluent where 198, 445, 387, 1430,  
 154 0.01, and 0.03 mg/l were recorded as average. However, Zn recorded much lower average  
 155 in the pharmaceutical effluent (0.01mg/l) compared to that of the textile effluent (0.11 mg/l)  
 156 while no significant differences were recorded in the Cu levels among the two industry (0.04  
 157 and 0.03 mg/l in the textile and pharmaceutical effluents). Nitrogen and phosphorus content  
 158 in both effluents (El-Bestawy et al., 2005 and Ellis 1977) along with the toxic industrial  
 159 contaminants suppressed the growth of cyanobacteria or any other algae.

### 160 3.2 Treatability and removal efficiency of effluent

#### 161 3.2.1 Contaminants

162 Residue levels of the selected quality parameters were determined (Table 2) and the  
 163 removal efficiencies (RE %) as results of the biological treatment using the selected species  
 164 were calculated. As a general trend, the two tested species exhibited positive correlation

165 between their RE% of all the tested parameters and the exposure time up to the last  
166 exposure day for both types of effluents.

### 167 **3.2.2 Organic matter removal**

168 **Biochemical oxygen demand Removal of BOD from industrial effluents of both**  
169 **industries using the selected algae revealed the following points:**

170 1. High REs% were obtained for BOD removal from industrial effluent by the selected  
171 species with *Anabaena aequalis* (90.65%) and finally *Phormidium mucicola* (81.9%).

172 2. Despite the RE variations of BOD achieved by the tested species, RC(s) of the BOD in  
173 the industrial effluent reached acceptable limits (19, 20 and 32 mg/l by *Anabaena aequalis*  
174 and *Phormidium mucicola*, respectively) after 7 days exposure which is much lower than the  
175 maximum permissible limit (APHA 1998) of 60 mg/l stated by the Environmental Laws for  
176 safe discharge into surface water courses. When these figures compared with those  
177 obtained by the control it was showed that the natural microbial population of the effluent  
178 achieved a maximum removal of 50% after 5 days equivalent to 60.7 mg/l (MPL of the BOD)  
179 after which there was a sharp decline in the efficiency associated by increasing the RC  
180 reaching 140 mg/l and 3.20% RE after 7 exposure of days.

181 3. Comparing BOD removal by the selected cyanobacteria revealed very high efficiency for  
182 all of them in the degradation of biodegradable organic matter which is stimulated by  
183 increasing the levels of the pollutant in the wastewater.

184

185 **Chemical oxygen demand Removal of COD from the industrial effluents using the**  
186 **selected species revealed the following points:**

187 1. *Anabaena aequalis* considered the most effective for removing COD from the industrial  
188 effluent achieving the maximum RE of 83.68% compared to the RE achieved by *Phormidium*  
189 *mucicola* (45.00%) after 7 exposure days. However, *Phormidium mucicola* exhibited higher  
190 COD RE% within the first 24 h compared to *Anabaena aequalis*.

191 2. The lowest residue concentration of 100 mg/l was achieved by *Anabaena aequalis* which  
192 is the maximum acceptable limit stated by the law (MPL for COD = 100 mg/l) while  
193 *Phormidium mucicola* could not bring the COD levels of the effluent to better quality. They  
194 recorded 180 and 246 mg/l, respectively, and required longer exposures. The highest RE%  
195 achieved by the control culture recorded 50.11% (170.59 mg/l) after 5 exposure days.

196 3. Similar to BOD removal, the natural microorganisms in the control culture were inhibited  
197 by the high Strength of the industrial effluent leading to reduction in the COD removal.

198 4. Although high REs% of the COD was achieved by the selected species, none of them  
199 could bring the COD levels in the effluent below the MPL during the investigated exposure  
200 time (1 week). This may be attributed to the need for longer time for achieving the proper

201 quality. It could also result from the inhibition in cyanobacterial growth due to the higher COD  
202 levels in the pharmaceutical effluent compared to that of the textile effluent.

### 203 **3.2.3 Solids removal**

204 **Total suspended solids (TSS) Removal of TSS from the industrial effluents using the**  
205 **selected species revealed the following points:**

206 1. The highest recorded TSS REs% in the effluent recorded 42.0 and 29.12% achieved by  
207 *Anabaena aequalis*, and *Phormidium mucicola* (125 and 133 mg/l RC), respectively, after 7  
208 days.

209 2. According to the law, 60 mg/l is stated as the MPL of the TSS; therefore none of the  
210 tested species reached the required efficiency to bring the TSS in the effluents below the  
211 MPL during the tested exposure time. This indicates that they required longer time, heavier  
212 biomass or different application using the same species to achieve that quality.

213 3. In contrast to cyanobacteria control culture achieved higher TSS removal form the textile  
214 effluent compared to that of the pharmaceutical effluent.

215

216 **Total dissolved solids (TDS) Removal of TDS from the industrial effluent using the**  
217 **selected species revealed the following points:**

218 1. The maximum TDS REs% obtained for the effluent by the tested species ranged between  
219 a maximum of 16.66% (1,078 mg/l) achieved by *Phormidium mucicola* and a minimum of  
220 12.00% (1,119 mg/l) obtained by *Anabaena aequalis* after 7 exposure days. *Phormidium*  
221 *mucicola* exhibited higher TDS RE% at the shorter exposures (up to 2nd day) compared to  
222 *Anabaena aequalis*.

223 2. Similar behavior for TSS removal was shown by bacteria of the control culture where  
224 higher TDS removal was achieved form the textile effluent compared to that of the  
225 pharmaceutical effluent.

226 3. Since the TDS content in the effluents were lower than the MPL of the TDS (2,000 mg/l),  
227 the residual concentrations of the TDS produced in the final effluents by all the tested  
228 species as well as the two controls improved and still within the safe range for discharging.

### 229 **3.2.4 Heavy metals removal**

230 Results revealed the following points:

231 1. *Phormidium mucicola* recorded the highest REs% for Zn from EWTP (86.12) and  
232 *Anabaena aequalis* (70.88%) recording RCs of 0.0247, and 0.0370 mg/l by three species,  
233 respectively, after 7 days.

234 2. Although low Zn levels were detected in the pharmaceutical effluent, lower Zn REs were  
235 achieved compared to those obtained for the textile effluent. Zn removal recorded 78.2, and

236 65.00% achieved as the highest Zn REs% by *Phormidium mucicola*, and *Anabaena*  
237 *aequalis*, respectively (0.0123, and 0.0182 mg/l, respectively) after 7 days.

238 3. Although all the average levels of Zn for both effluents were below the MPL of 5 mg/l  
239 before the treatment, Zn levels were reduced producing much better effluent quality. Zinc  
240 removal was stimulated by increasing its level in the wastewater.

241 4. Concerning Cu , much higher REs% were recorded for effluent compared to those  
242 obtained for Zn removal regardless it's high toxicity. This may be attributed to the high  
243 resistance of the selected members which was stimulated by increasing Cu levels in the  
244 wastewater. 94.63, 90.99 and 90.64% RE of Cu were achieved by *Phormidium mucicola*,  
245 and *Anabaena aequalis* (0.0031, and 0.0054 mg/l RC), respectively, after 7 days.

#### 246 **4. CONCLUSION**

247 In conclusion, results confirmed that *Phormidium mucicola* and *Anabaena aequalis* are the  
248 effective species for BOD, COD, TSS, TDS, Zn and Cu removal from the effluents of the  
249 textile and pharmaceutical industries. Zn and Cu both are toxic heavy metals. Effluents  
250 discharge in water bodies can affect the living organism. These cyanobacterial species are  
251 able to remove these metals from water body. Absorption of metals increase in the higher  
252 concentration. Presence of these cyanobacteria can low BOD, COD, TSS, TDS.

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