Pharmaceutical wastewater treatment using free and immobilized Cyanobacteria

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Abstract:

An attempt has been made to study the feasibility of pharmaceutical wastewaters (laboratories waste (WI) and production waste (WII)) treatment using free and immobilized cyanobacterium Phormidium fragile. After preliminary test with different concentrations, WI showed high toxicity, only 0.5% conc. was effective while WII was used at high conc. of 40% after 10 days incubation (from growth curve experiment) in optimum growth conditions. Role of Phormidium cells in bioremediation of pharmaceutical wastewaters was effective whereas maximum percentage removal in tested parameters were 51.56% and 58.89% for ammonia, 66.67% and 61.23% for phosphorus, 51.74% and 54.55% for COD and 37.05% and 59.85% for BOD in WI and WII respectively with respect to treatments without microalgae. N-starvation for 36 hours prior to cultivation caused increase in percentage removal of ammonia, phosphorus and COD with increase in algal dose than unstarved cells, 100%, 100% and 72.56% for WI and 87.28%, 100% and 71.98% for WII, respectively with 40 ml algal dose. It was found that 2500 lux and 25 ºC were the best for nutrients removal. Chlorophyll a, dry weight, protein and total carbohydrate contents were also estimated. During this study, pH levels increased and remained in the range 7.3 to 9.1. Significant removal of ammonia, phosphorus and COD were observed in algal alginate beads treatments than blank beads. Starvation before immobilization recorded the highest removal percentage when compared to unstarved beads or free cells in both wastes types and their mixture (1:1). When starved and unstarved Phormidium beads were incubated in WI, WII and a mixture of both in a semi-continuous system for five consecutive cycles (10 days each), there was increase in nutrients percentage removal up to the third cycle. Microalga in the beads survived long enough which gives a promise to upgrading immobilization technique for wastewater treatment with low cost.

Keywords: Cyanobacteria, Phormidium, Pharmaceutical wastewater treatment, starvation, immobilization, alginate beads.

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Introduction

The pharmaceutical industry is one of the most important for modern civilization. Increasing number of pharmaceutical industries leads to hazardous impact on water quality and thus affects the surrounding environment and human health. Thus, the pharmaceutical industry has become one of the major causes of concern. The day by day increased level of water pollution highlights the need for time to time assessment/characterization of pharmaceutical industrial wastewater (Rana et al., 2014). Many pharmaceuticals and personal care products (PPCPs) have been designed to stay active for long periods of time to fulfill their therapeutic function; thus, they can persist in the environment, remaining active and affecting aquatic life (Ricart et al., 2010; Kvarnryd et al., 2011).

In fact, the pharmaceutical industry is characterized by a diversity of products, processes, plant sizes, as well as wastewater quantity and quality. Hence, it is almost impossible to describe a “typical” pharmaceutical effluent because of such diversity, wastewater treatment and disposal problems have also increased as a result (Dixit and Parmar, 2013). Though the volume of untreated or incompletely treated pharmaceutical industry wastewater is small, it contains a high level of pollutants because of the presence of nonbiodegradable organic matter and other pollutants (Vuppala et al., 2012; Ramola and Singh, 2013). Untreated pharmaceutical wastewater discharge into the natural environment causes health hazards to existing flora and fauna and therefore, treatment of the effluents is necessary to bring down the concentration of toxic substances to desired limits, before they are finally discharged into the natural systems (UNESCO, 2005). In order to design an appropriate treatment system the characteristics of the generated wastewater need to be followed and analyzed. Nowadays, pharmaceutical companies employ a variety of treatment methods, which includes primary: chemical and physicochemical, secondary: biological process and tertiary: advanced oxidation processes which utilize costly chemicals and treatment units, which are difficult to manage at the industrial unit level (Amin et al., 2013; Vanerkar et al., 2013).
Phycoremediation is a novel, low cost and an effective technique that uses algae to clean up wastewater offering very simple and economical method as compared to the other conventional treatments (Elumalai et al., 2013). Also microalgae have an effective role in the uptake of the pollutants to use inorganic nitrogen and phosphorus for their growth as well as their ability to produce valuable biomass that can be used in different industrial applications (Tam and Wong, 1995). Successful treatment of wastewater with microalgae requires understanding of the factors that affect growth such as physical, chemical and biological factors (Larsdotter, 2006). The major problem in utilization of microalgae in any industrial or wastewater treatment is harvesting of the biomass. This is solved by the strategy of immobilization. This precludes the use of supporting material obtained either naturally including agar, alginate and carrageenan or synthetics such as polyacrylamide and polyurethane (Vijayakumar, 2012). The aim of the present study is treatment of the pharmaceutical wastewaters with free and immobilized cyanobacteria under different conditions in a trial for bioremediation and also testing the growth and synthesis of some metabolites of the used cyanobacterium were undertaken for further applications.

**Materials and Methods**

Organism and growth conditions:

The filamentous fresh water cyanobacterium *Phormidium fragile* was selected for the current study, it was kindly supplied from the culture collection of Botany and Microbiology Department, Faculty of Science, Cairo University, Egypt. It was routinely grown in BG 11 medium (Rippka et al., 1979) at continuous illumination with light intensity of 48.75 µEm⁻²s⁻¹ (2500 lux) at 25 ± 2 ºC. The start inoculum was adjusted to be approximately 1.297 µg ml⁻¹ chlorophyll a. Optimum growth period was found to be 10 days (cells in the exponential phase) as determined from growth curve experiment.
Pharmaceutical wastewater:

The pharmaceutical wastewaters used in this study were obtained from a pharmaceutical company in 6th of October city, 1st industrial zone on February 2017 in one liter capacity bottles. They are two types: Wastewater I (Laboratories waste WI), it was collected from chemical laboratories safety cans, it includes inorganic acids and bases, organic solvents, metals, unused chemicals and chemical reactions products and Wastewater II (Production Waste WII), it was collected by mixing equal volumes from the washing rinse of different production departments, it include parenteral, solids (Tablets and Capsules), semisolids, (cream, ointment, gel and suppository), syrups, antibiotics, instant and powders. Both waste types were separately collected and analyzed pre and post experiments within few hours from collection according to standard procedures of APHA (2012). Wastes were immediately preserved in refrigerator at 4 ºC. Thereafter the primary treated wastes were kept in a deep freezer at -20 ºC until used (Fedorova et al., 2014).

Methods of experimentation:

After preliminary experiments serial dilutions of wastewaters were prepared by mixing primary treated wastewater and distilled water to obtain concentrations of 0.5%, 1%, 2%, 3% and 4% (V/V) for wastewater I and 10%, 20%, 40%, 60% and 80% (V/V) for wastewater II. Triplicate flasks from all treatments each containing 300 ml primary treated wastes in addition to control (BG11) were inoculated with 20 ml of Phormidium fragile culture. After 10 days incubation in optimum conditions cyanobacterial cells were harvested and growth parameters were measured in terms of chlorophyll a (Sartory and Grobbelaar, 1984), dry weight (APHA, 2012), total carbohydrates (Dubois et al., 1956) and protein (Lowry et al., 1951), in addition to BOD, COD (APHA, 2012), ammonia (Chaney and Marbach, 1962) and phosphorus using Fiske-Subbarow method (Clark and Switzer, 1977) of treated wastewaters were also analyzed. Next, the most efficient conc. selected from each waste but with no cyanobacterial inocula was prepared and used as negative control. The chosen wastes concentrations were used for the next experiment in which a series of cyanobacterial suspension inocula (5, 10, 20, 30, 40 ml) were inoculated in conical flasks containing 300 ml waste each. A parallel group of flasks in which Phormidium fragile was
previously nitrogen starved for 36 hours before inoculation. All treatments of starved and unstarved microalgal cells were incubated for 10 days after which growth parameters and wastewaters were analyzed. Two ecological factors, light intensity (2500, 8000 and 12000 lux) and temperature (15±2 °C, 25±2 °C and 40±2 °C) were tested to study their role in pharmaceutical wastewater bioremediation by *Phormidium fragile*.

For the formation of the immobilized cyanobacterial beads, the same inoculum of *Phormidium* was washed twice with double distilled water and initial chlorophyll *a* was determined. The required quantity was added as a suspension to the sodium alginate solution (5% W/V) in distilled water after sterilization and cooling under aseptic conditions, alginate beads were formed by using a peristaltic pump to obtain (bead average diameter ≈ 0.5 cm), by drop wise into calcium chloride CaCl$_2$ 0.2 M solution using magnetic stirrer during the preparation for two hours to maintain the shape stability (*Rai and Malick, 1992*). The same steps were repeated on the nitrogen starved alga and empty alginate beads (without microalgae). Fifty beads were added in each flask which containing 200 ml waste. The same steps were repeated on a mixture (1:1) from waste I (0.5%) and waste II (40%) freshly prepared for immobilized *Phormidium* cells either starved or unstarved, in addition to blank beads. At the end of incubation period microalgae beads were separated from the wastewater by filtration and the wastes were analyzed for ammonia, phosphorus and COD. *Phormidium* alginate beads used in the previous experiment were used in five consecutive cycles (10 days each) with fresh waste which was analyzed after each cycle in a semi-continuous mode in a trial to test the long term effect of microalgal beads in wastewater treatment.

**Statistical analysis:**

The means and standard deviation values of the triplicates for each treatment were calculated. The data were analyzed statistically on the basis of analysis of variance (ANOVA). All statistical analyses were carried out using SPSS program (IBM, version 23).
Results and Discussion

Some raw and treated pharmaceutical wastewaters (Laboratories WI and Production WII) characteristics were analyzed before the beginning of the experiments (Table 1). pH value of WI was 10.3 and became 7 after neutralization. Both wastes were filtered, total, suspended and dissolved solids were much lower in treated (filtered) wastes, all were higher in WII than in WI. Shabana (1994) mentioned that filtration helps in removal of solid particles which leads to decrease of some pollutants by adsorption on its surface. Decrease in ammonia, phosphorus, anions and cations, was observed in treated wastes. An observed decrease in COD and BOD by filtration with higher values in WII than WI, yet both values are still exceeding the official limits for wastewater disposal. At high pH (8.5 or above) especially in presence of calcium ion, some of the phosphorus will be precipitated in the form of orthophosphate which is the case for WII in this study (Jimenez-Perez et al., 2004).

The effect of different concentrations of pharmaceutical wastewaters WI and WII after preliminary experiment on the synthesis of pigment and metabolites in terms of chlorophyll a, dry weight, protein and total carbohydrates is shown in (Table 2). Chlorophyll a content for Phormidium fragile culture decreased significantly (P<0.05) with increase in waste I concentration to reach minimum value (1316.00 µgL⁻¹) at 4% conc. The highest chlorophyll a value was recorded at 0.5% conc. (2026.33 µgL⁻¹).

In waste II treatments there was a significant increase in chlorophyll a content up to 40% conc. (2203.67 µgL⁻¹). Laboratories wastewater seems more toxic than production waste. In this connection, (Li et al., 1991) reported that the higher the content of algal chlorophyll a in the water, the more effective the removal of nitrogen and phosphorus from the water. Since the highest chlorophyll a content was observed in 0.5% for waste I and 40% conc. for waste II but all are still less than control value, these concentrations were selected to be used in all the following experiments in this investigation. Dry weight have similar trend of chlorophyll a for both types of wastes.
Table 1: Some characteristics of pharmaceutical wastes before and after filtration

| Parameter                        | Waste I (Laboratories) | Waste II (Production) |
|-----------------------------------|-------------------------|------------------------|
| mgL⁻¹ except for pH               | Raw waste | After neutralization and filtration | Raw waste | After neutralization and filtration |
| Temperature                       | 25 °C       | 25 °C                  | 25 °C       | 25 °C                  |
| Colour                            | Colorless | Colorless             | Orange      | Orange                 |
| pH                                | 10.3       | 7.0                   | 7.4         | 7.0                   |
| Total Solids (TS)                 | 12900      | 3382                  | 18300       | 4675                  |
| Total dissolved solids (TDS)      | 7900       | 3252                  | 13300       | 4325                  |
| Total suspended solids (TSS)      | 5000       | 130                   | 5000        | 350                   |
| Chemical Oxygen Demand (COD)      | 14624      | 7245                  | 24286       | 13705                 |
| Biological Oxygen Demand (BOD)    | 790        | 580                   | 1842        | 1223                  |
| Calcium (Ca⁺⁺)                    | 17         | 10                    | 395         | 101.22                |
| Chlorides (Cl⁻)                   | 570        | 143.5                 | 14          | 3.5                   |
| Sulphates (SO₄⁻²)                 | 2540       | 390                   | 8.5         | 2.6                   |
| Ammonia (NH₃)                     | 693        | 450                   | 43.6        | 39.67                 |
| Phosphorus (P)                    | 8.5        | 5.35                  | 11.8        | 6.9                   |

In waste I treatments the highest protein value (Table 2) was recorded at 0.5% conc. (25.97 mgL⁻¹). In waste II, the production of protein was favored by increase in waste concentration up to 40%, it recorded (22.74 mgL⁻¹) but all values are still more than control value. Total carbohydrates accumulation by Phormidium fragile culture was decreased by increase in WI concentrations till it reached (48.89 mgL⁻¹) at 4% conc. In WII treatments, there was an increase in...
total carbohydrates content up to 60% conc. (82.95 mgL\(^{-1}\)). In both wastes all recorded values are still less than control values grown on BG11 medium, still remain total carbohydrates in WII was higher than in WI cultures.

Table 2: Effect of different pharmaceutical wastewaters on chlorophyll a content, dry weight, protein and total carbohydrate contents of *Phormidium fragile* after 10 days incubation. Data are average of three replicates; each value represents the mean ± S.D.

| Waste I conc. (%) | Chlorophyll a content μgL\(^{-1}\) | Dry weight mgL\(^{-1}\) | Protein content mgL\(^{-1}\) | Total carbohydrate content mgL\(^{-1}\) |
|------------------|------------------------------------|------------------------|-----------------------------|----------------------------------------|
| Control (0.0)    | 2270.33±08.50\(^{a}\)             | 833.33±15.28\(^{f}\) | 14.02±01.58\(^{a}\)        | 98.90±07.05\(^{a}\)                    |
| 0.5              | 2026.33±07.09\(^{c}\)             | 420.00±10.00\(^{c}\)  | 25.97±00.91\(^{d}\)        | 61.74±03.71\(^{b}\)                    |
| 1                | 1793.00±10.82\(^{d}\)             | 366.67±05.77\(^{d}\)  | 21.78±01.75\(^{d}\)        | 59.16±02.15\(^{b}\)                    |
| 2                | 1547.33±07.23\(^{c}\)             | 340.00±10.00\(^{c}\)  | 18.43±00.72\(^{b}\)        | 57.43±01.60\(^{b}\)                    |
| 3                | 1455.67±09.45\(^{b}\)             | 306.67±05.77\(^{b}\)  | 14.18±02.00\(^{b}\)        | 55.63±02.01\(^{b}\)                    |
| 4                | 1316.00±08.19\(^{a}\)             | 283.33±05.77\(^{a}\)  | 13.45±01.72\(^{b}\)        | 48.89±03.11\(^{b}\)                    |
| Waste II conc. (%)
| Control (0.0)    | 2270.33±08.50\(^{f}\)             | 833.33±15.28\(^{f}\)  | 14.02±01.58\(^{a}\)        | 98.90±07.05\(^{d}\)                    |
| 10               | 1966.00±10.44\(^{a}\)             | 346.67±05.77\(^{a}\)  | 15.97±00.12\(^{ab}\)       | 53.61±02.21\(^{b}\)                    |
| 20               | 2157.33±10.60\(^{b}\)             | 413.33±11.55\(^{b}\)  | 18.73±00.77\(^{b}\)        | 59.01±02.82\(^{c}\)                    |
| 40               | 2203.67±14.57\(^{c}\)             | 493.33±15.28\(^{c}\)  | 22.74±01.63\(^{c}\)        | 69.45±02.32\(^{ab}\)                   |
| 60               | 2180.00±11.36\(^{d}\)             | 426.67±05.77\(^{b}\)  | 18.09±01.74\(^{b}\)        | 82.95±02.86\(^{c}\)                    |
| 80               | 2084.67±06.66\(^{b}\)             | 340.00±10.00\(^{a}\)  | 16.20±02.01\(^{ab}\)       | 65.56±01.97\(^{b}\)                    |
| Initial          | 1230.33±05.86                     | 71.83±02.21           | 12.62±00.65                | 27.02±01.21                            |

Means marked with the same superscript letters are not-significant (P>0.05), whereas others with different superscript letters are significant (P<0.05).
From the above mentioned experiment, the selected concentrations (0.5% for waste I and 40% for waste II) were incubated in optimum growth conditions with and without inoculation with *Phormidium fragile*. The microalgal efficiency for removing ammonia, phosphorus and reduction of COD and BOD from pharmaceutical wastewaters at certain concentrations was presented in (Table 3). The results demonstrated that removal percentage of ammonia, phosphorus, COD and BOD were higher after microalgal treatment than treatments without microalgae indicating effective role of *Phormidium fragile* in removal process. In the above parameters except phosphorus, percentage removal was higher in Pharmaceutical production waste (WII) than laboratories waste (WI). Whereas ammonia percentage removal by *Phormidium* in WI was 51.56% and 58.89% in WII while it was 22.67% for WI 0.5% and 25.02% for WII 40% in treatment without algae which is in agreement with *Lau et al.* (1995), they found that maximum ammonia removal percentage in the control (without algae) was less than 20%. Microalgae mainly use inorganic nitrogen as a nutrient source for their cell synthesis and ammonia is the first preferable form of inorganic nitrogen for microalgae. *Tam and Wong (2000)* found that the reductions in wastewater NH$_4^+$-N concentrations in all algal bioreactors were significantly better than those without algae, suggesting that uptake of ammonium and assimilation into algal biomass are essential processes. *Dubey et al.* (2011) investigated the potential degradation of industrial effluents by environmental species of cyanobacteria isolated from the pharmaceutical industries. Results indicated the potential of natural resources as efficient agents for pollution control.

In the results demonstrated in (Table 3), phosphorus was efficiently reduced in both wastes WI 0.5% conc. and WII 40% conc. by 66.67 % and 61.23% respectively. Also COD was reduced from 922 to 445 mgL$^{-1}$ and from 5460 to 2481.33 mgL$^{-1}$ for laboratories WI and production WII wastes respectively. Also BOD was reduced from 224 to 141 mgL$^{-1}$ and from 680 to 273 mgL$^{-1}$ for WI and WII with algae respectively, comparing with treatment without algae, this indicates the role of microalgal cells in the treatment. In previous study reported by *Kshirsagar (2010)*, COD of herbal and bulk drug pharmaceutical wastewaters was reduced from 13,090 to 3,927 mgL$^{-1}$ and from 34,452 to 13,333
mgL\(^{-1}\) respectively whereas BOD of herbal and bulk drug pharmaceutical wastewater was reduced from 7,420 to 2,003 mgL\(^{-1}\) and from 15,840 to 5,608 mgL\(^{-1}\) respectively.

Table 3: Analysis of filtered pharmaceutical WI and WII at certain concentrations before and after inoculation with *Phormidium fragile* for 10 days incubation. Data are average of three replicates; each value represents the mean ± S.D.

| Parameter mgL\(^{-1}\) | WI 0.5% conc. | | WII 40% conc. | |
|-----------------------------|-----------------|-----------------|-----------------|-----------------|
|                             | Initial | Without algae | After algal treatment | Initial | Without algae | After algal treatment |
| Ammonia                     | 2.25 ±0.03 | 1.74 ±0.05 | 1.09 ±0.07 | 11.87 ±0.31 | 8.90 ±0.34 | 4.88 ±0.10 |
| % Removal                   | ----    | 22.67         | 51.56         | ----    | 25.02         | 58.89         |
| Phosphorus                  | 0.03 ±0.01 | 0.02 ±0.01 | 0.01 ±0.01 | 2.76 ±0.05 | 2.04 ±0.07 | 1.07 ±0.08 |
| % Removal                   | ----    | 33.33         | 66.67         | ----    | 26.09         | 61.23         |
| COD                         | 922.00 ±03.61 | 690.00 ±09.54 | 445.00 ±07.00 | 5460.00 ±12.77 | 3983.00 ±14.53 | 2481.33 ±12.06 |
| % Reduction                 | ----    | 25.16         | 51.74         | ----    | 27.05         | 54.55         |
| BOD                         | 224.00 ±08.72 | 150.00 ±07.21 | 141.00 ±06.24 | 680.00 ±09.54 | 393.00 ±08.72 | 273.00 ±07.00 |
| % Reduction                 | ----    | 33.04         | 37.05         | ----    | 42.21         | 59.85         |

Nutrient starvation is a stressful condition for algae that changes biochemical compounds of biomass ([Malakootian et al., 2015](#)). The effect of N-starvation for *Phormidium* cells and a series of cyanobacterial suspension inocula on nutrient removal from WI and WII was investigated. The data generated in the present study (Fig.1) revealed that removal efficiency of ammonia from WI 0.5%
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conc. by *Phormidium fragile* ranged from 16 - 100% for unstarved cells, while it was 28.44-100% for starved cells and for WII 40% conc. ranged from 48.86-84.92% for unstarved cells, while it was 51.05-87.28% for starved cells. Increase in algal dose caused significant decrease in ammonia content from both pharmaceutical wastewaters in both unstarved and starved cell cultures and reached maximum removal percentage at 40 ml algal dose, it was 100% in WI and 87.28% in WII starved cells. *Lee et al. (2016)* reported that the increased algal biomass was accompanied by ammonia decrease during the first 5 days which indicates that algae take up ammonia for biomass production and as an energy source leading to increased nitrogen removal.

In the current study, Phosphorus conc. in the wastewaters exhibited high decrease with increase in algal inoculum of both unstarved, starved cultures (Fig.1). Phosphorus disappeared completely at higher algal doses (30 ml and 40 ml) recording 100% percentage removal in both unstarved and starved cells. These data are in agreement with *(Tam and Wong, 1989)*, who reported an increase in phosphorus removal with increasing inoculum dose of algal cells. *Amiri and Ahmadi (2019)* reported that phosphate removal during phycoremediation is due to the utilization of phosphorus for algal growth. The phosphorus, which is used in the algal cells mainly for the production of phosphorous-containing compounds such as adenosine triphosphate (ATP) and nicotinamide adenine dinucleotide phosphate-oxidase (NADPH). *Rao et al. (2011)* suggested that the chemical stripping of phosphorus may be regarded as an advantageous side-effect of the algal growth, with enhanced phosphorus removal as a result.

COD determined at the end of incubation period generally exhibited reduction with increase in algal inoculums (Fig.1). Percentage reduction of COD was 72.56% and 71.98% at higher starved algal dose for WI and WII respectively while it recorded percentage reduction 65.62% of initial conc. for unstarved algal cells for WI and 68.58% for WII. In the contrary, *Lau et al. (1995)* reported that *Chlorella vulgaris* with four initial inoculum sizes when applied to primary treated sewage, there were no significant differences between the four algal cultures in terms of COD removal.
The data presented in (Fig. 2) revealed that chlorophyll a content and dry weight values of *Phormidium* cells in both wastes (WI 0.5% conc. and WII 40% conc.) increased significantly by increasing the algal dose of both unstarved and starved *Phormidium* cells showing higher contents of chlorophyll a in the starved treatments than the unstarved. Chlorophyll a content was (3756.67 µgL⁻¹) and
(3690.00 µgL\(^{-1}\)) at 40 ml algal dose in starved Phormidium cells for WI 0.5% and WII 40% respectively. The same trend was demonstrated for dry weight data which was highly increased with increasing algal inoculum to reach highest values (546.67 mgL\(^{-1}\) and 480.00 mgL\(^{-1}\)) at 40 ml unstarved and starved Phormidium cells for WI 0.5% while recorded (626.67 mgL\(^{-1}\) and 533.33 mgL\(^{-1}\)) at 40 ml unstarved and starved algal cells for WII 40% at the end of incubation period. The data presented in (Fig.2) also revealed that protein production by Phormidium cultures was favored with increase in algal dose to reach the highest yields (45.88 mgL\(^{-1}\) and 49.81 mgL\(^{-1}\)) at 40 ml unstarved and starved cells for WI 0.5% while reached (29.79 mgL\(^{-1}\) and 34.80 mgL\(^{-1}\)) at 40 ml unstarved and starved for WII 40%. In addition, increase in the protein content in starved algal cells was more than unstarved for both waste types. The same trend was demonstrated for accumulation of total carbohydrates, since the highest values (76.32mgL\(^{-1}\) and 93.52 mgL\(^{-1}\)) were recorded at 40 ml unstarved and starved Phormidium cells for WI 0.5% while recorded (127.57 mgL\(^{-1}\) and 156.77 mgL\(^{-1}\)) at 40 ml unstarved and starved for WII 40%. Nitrogen starvation was also employed to trigger the accumulation of lipid and carbohydrate (Ho et al., 2012). It is clear that in contrary to protein production of total carbohydrates by Phormidium cultures in waste II at moderate and higher doses far exceeded that accumulated in waste I cultures.

Mandal and Mallick (2009) demonstrated that cultivation under nitrogen deficient conditions leads to a sharp increase in the lipid or carbohydrate content, because the condition of nitrogen-depletion probably transforms protein or peptides to lipids or carbohydrates. Therefore, the condition of nitrogen starvation causes an enhancement of energy-rich products, such as lipids and carbohydrates. Quantities of the available nitrogen in the culture directly alter cell growth. Nitrogen limitation in the microalgae culture can reduce growth and biomass productivity although they increase production of carbohydrates and lipids (Daliry et al., 2017). Worthy to be noted that, in the present study the protein and total carbohydrate contents of Phormidium cells in certain concentrations of both wastes tended to increase progressively by intensifying the algal dose with higher value in the starved than the unstarved algal dose. Salama and Shabana (1989) found that nitrogen starvation of Phormidium fragile for 36-44 hrs leads to elevation in the total carbohydrate and protein accumulation by the organism.
Fig. 2: Effect of pharmaceutical waste I (0.5 % conc.) and waste II (40 % conc.) on chlorophyll a content, dry weight, Protein and total carbohydrate contents of starved and unstarved *Phormidium fragile* cells with different doses after 10 days incubation.
An excessive cell density could limit the light penetration into the water column and thus reduce the growth, particularly in scarcely mixed cultures (Campos-Rudin and Silva-Benavides, 2018).

Apart from photosynthetic and chemosynthetic assimilation by microalgae, N and P removal from wastewaters are also affected by abiotic factors significantly. Therefore, the N and P removal in wastewater-based algae cultivation system were attributed to assimilation by microalgae as well as volatilization and precipitation caused by abiotic factors (Zhou, 2014). Culture growth depends on different factors such as light intensity, pH, nutrients, carbon dioxide, mixing, and temperature. In wastewater, growth depends in part by the active nutrient uptake of the photosynthetic cells and their transformation into biomass (Voltolina et al., 2005). Regarding light intensity (Fig.3), it is relevant to notice that 2500 lux was the optimum light intensity for ammonia and phosphorus removal from both pharmaceutical wastewaters by Phormidium cultures 52.89% and 66.67% in WI and 59.14% and 62.32% in WII. Microalgae need light as an energy source to convert the absorbed water and CO$_2$ into biomass (Ozkurt, 2009). Nutrient removal can also be further increased by NH$_3$ stripping or P precipitation due to the rise in the pH associated with photosynthesis (Larsdotter, 2006). Also the data in (Fig.3) revealed that the highest COD percentage reduction was in 2500 lux for WI 0.5% conc. and WII 40% conc. cultures (54.23% and 54.92%) respectively. Ho et al. (2012) observed in a study, the dependence of the specific growth rate of Scenedesmus obliquus on light intensity under continuous illumination. It was clearly seen that the specific growth rate increased dramatically with rising light intensity and then it gradually leveled off as the light intensity continued to rise. Suggesting that excessive illumination would inhibit the biomass production and CO$_2$ fixation ability, which is commonly recognized as the photo-inhibition effect.

Temperature is another important factor in the growth of microalgae and directly influences the biochemical processes, including photosynthesis, in the algal cell factory. Each species has its own optimal growth temperature. Increasing temperature to the optimum range exponentially increases algal growth, but an increase or decrease in the temperature beyond the optimal point retards or even stops algae growth and activity (Bechet et al., 2017).
Fig. 3: Effect of light intensity and temperature on pH, ammonia and phosphorus removal and COD reduction from pharmaceutical waste I (0.5 % conc.) and waste II (40% conc.) by Phormidium fragile after 10 days incubation.

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In the current study, maximum ammonia and phosphorus removal were recorded at 25 ± 2°C for both wastes where it was 53.78% and 66.67% in WI and 59.65% and 64.13% in WII. Azeez (2010) reported that the highest removal of nitrogen and phosphorus is achieved at temperature 30°C and they were 92%, 89% respectively. The data recorded in (Fig.3) showed that maximum percentage reduction of COD was at 25 ± 2°C (53.15% and 55.01% of initial) was recorded at such degree for WI and WII respectively. On the other hand minimum percentage removal of COD was attained at 40 °C (47.47% and 49.48%) for WI and WII which is in agreement with (Azeez, 2010), who found that at temperature 30°C there is a decrease in COD with a removal efficiency 88%. While at temperature higher than 30°C the removal efficiency of COD decreases because the activity of microorganism becomes lower after this temperature which means that ability to remove pollutants decreases. However, in optimum temperature algal growth would not be affected by high light intensities (Borowitzka, 1998). Microalgae can tolerate the temperature range of 16–27 °C and the temperature lower than 16 °C slow down the growth whereas higher than 35 °C is harmful for the number of species (Butterwick et al. 2005).

The use of free cells is rather rare in comparison to immobilized cells, since immobilization of cells offers various advantages and the process is cost effective (Vijayakumar, 2012). The immobilized cells have many advantages in the bioremediation pollutants in wastewaters: nontoxic, transparent for light, stable in growth medium, a high population density can be entrapped in a small volume, it provides microorganisms protection against toxicity of the pollutants in wastewater, the minimization of inhibition and toxicity to microbial cells by diffusional constrains as well as avoiding cell harvesting problems. The immobilized cells can be reused for successive applications and this reduce the costs because it can remain viable for long time, it can be stored for long periods without loss of their activities for bioremediation (Mallick, 2002; Quek et al., 2006; Bikram et al., 2014). Protection against toxicity in immobilized cells is reported in different works. Immobilized cells are characterized by its stability because the cells are entrapped in the matrix and are protected from toxic compounds in wastewater; therefore, they have a higher efficiency for the removal of nutrients (Cassidy et al., 1996; EL-Sheekh et al., 2016), dye (Revathi et al., 2017), and heavy metals (EL-Sheekh and Mahmoud 2017) than free cells.
Despite these advantages, alginate beads have some disadvantages such as their low mechanical stability for long term use in bioreactor; more over calcium alginate gels are disrupted by phosphate ions (Boominathan, 2005; Vijayakumar and Manoharan, 2012).

In the present study, initial ammonia contents were (2.25, 11.87 and 7.33 mgL$^{-1}$) in waste I, waste II and their mixture. Ammonia content was significantly decreased in all treatments (Table 4) either in free or immobilized algal cells when incubated in waste I and waste II giving removal percentage (53.33% and 59.73%) in free cells. Removal percentage in all treatments increased (72.44%, 89.22% and 69.03%) for WI, WII and their mixture (1:1) which means that Phormidium in alginate beads was more efficient than free algal cells in the ammonia removal process. Higher nutrient removal efficiency has been recorded in the immobilized algal biomass than the freely suspended cells of the same algal species (De la Noüe and Proulx, 1988; Abdel Hameed, 2002). Moreover nitrogen starvation of algal cells before immobilization recorded the highest value of ammonia removal (80%, 91.15% and 79.54%) for WI, WII and their mixture, respectively. The success in employing immobilized microalgae for wastewater treatment depends upon many factors, including algal species, immobilization matrix, cell and bead concentration, bead morphology, etc. (Tam and Wong, 2000). This is in agreement with (Abdel Hameed, 2007), who reported that C. vulgaris immobilized in alginate beads were more effective in removing N and P from wastewater than blank alginate beads. (Table 4) showed that most efficient ammonia removal occurs in the treatments with starved immobilized Phormidium cultures for WI, WII and their mixture after 10 days incubation period.

Regarding to Phosphorus content, it was decreased in all treatments. Data in (Table 4) showed that immobilized Phormidium was more efficient than free algal cells in the phosphorus removal since percentage removal reached 100%, 73.91% and 63.33% for WI, WII and their mixture where as it was 66.67% and 63.41% for WI and WII free cultures. Starvation of algal cells before immobilization recorded the highest values of phosphorus removal (100.00%, 88.04% and 84.00%).
Tam and Wong (2000) reported 78% ammonium and 94% phosphate removal efficiencies with immobilized *C. vulgaris*, entrapped in calcium alginate beads, compared to the 40% ammonium and 59% phosphate removal with free cells. Lau et al. (1997) also observed significantly higher ammonium (95%) and phosphate (99%) removal efficiencies for *C. vulgaris* cells immobilized in alginate beads relative to their free counterparts.

Initial COD content in WI (922.00 mgL⁻¹), in WII (5460.00 mgL⁻¹) and their mixture (3191.67 mgL⁻¹) decreased recording higher percentage reduction in *Phormidium* alginate beads treatments. Starvation before immobilization increased reduction percentage of COD in all wastes tested recording (60.48%, 61.31% and 64.63%) for WI, WII and their mixture, respectively. The lowest reduction percentages for ammonia, phosphorus and COD were observed in blank beads for all waste treatments indicating the role of algal cells in the process of bioremediation. The low efficiency of removal of COD in the blank beads flat-photobioreactor was due to the presence of a carboxyl functional group which binds to the organic material only (Singh et al., 2012). Lau et al. (1997) supported the idea that simple inorganic ions such as nitrate, ammonium and phosphate would be as freely available to immobilized algae as to their free counterparts, because nutrients must diffuse through the alginate pores to reach the algal cells. The higher phosphorus removal in immobilized cells may be directly related to the initial concentration of cells per ml of culture medium. The lower cells density may have caused a greater light diffusion and, thus, lower self-shading effect within cells. This could have allowed for greater cellular activity and consequently reach larger nutrients removal (Ruiz-Marin et al., 2010). It is worthy to mention that free algal cells were disintegrated in the mixture treatment giving no growth which means inhibitory effect of it when in a direct contact with the microalgal cells. The immobilization on alginates protects microorganisms from toxic substances, pH, and temperature extremes (Lessel, 1994). It increases the rate of biodegradation of pollutants through increasing cell loading and this also improve the catalytic stability as well as the tolerance against toxic pollutants (Wang et al., 2007). From the present data it is clear that immobilized *Phormidium* was more efficient than free algal cells in the removal process where as nitrogen starvation before immobilization increased removal percentage of ammonia, phosphorus and COD.
Table 4: Effect of immobilized starved and unstarved *Phormidium fragile* cells on ammonia and phosphorus removal and COD reduction from pharmaceutical wastewaters I, II and their mixture after 10 days incubation. Data are average of three replicates; each value represents the mean ± S.D.

| Parameter | Treatment | Ammonia Removal mgL⁻¹ | Phosphorus Removal mgL⁻¹ | COD Reduction mgL⁻¹ | pH          |
|-----------|-----------|------------------------|--------------------------|---------------------|-------------|
| WI 0.5%   | Blank Beads | 1.57±0.14             | 30.22 (%)                | 33.33 (%)           | 634.67±0.93  | 31.16  | 8.40±0.13  |
|           | Unstarved  | 0.62±0.10             | 72.44 (%)                | 100 (%)             | 410.33±0.28  | 55.50  | 8.69±0.14  |
|           | Starved    | 0.45±0.06             | 80.00 (%)                | 100 (%)             | 364.33±0.06  | 60.48  | 8.50±0.11  |
|           | Free cells | 1.05±0.10             | 53.33 (%)                | 66.67 (%)           | 438.00±0.31  | 52.49  | 8.58±0.17  |
|           | Initial    | 0.25±0.03             | 0.03±0.01 (%)            | 922.00±0.61 (%)     | 07.00±0.00  |        |           |
| WII 40%   | Blank Beads | 07.43±0.16            | 37.41 (%)                | 73.91 (%)           | 2323.67±0.28 | 57.44  | 8.53±0.25  |
|           | Unstarved  | 01.28±0.07            | 89.22 (%)                | 73.91 (%)           | 2323.67±0.28 | 57.44  | 8.53±0.25  |
|           | Starved    | 01.05±0.08            | 91.15 (%)                | 88.04 (%)           | 2112.33±0.86 | 61.31  | 8.41±0.24  |
|           | Free cells | 04.78±0.16            | 59.73 (%)                | 63.41 (%)           | 2546.00±14.00 | 53.37  | 8.21±0.09  |
|           | Initial    | 11.87±0.31            | 02.76±0.05 (%)           | 5460.00±12.77 (%)   | 07.00±0.00  |        |           |
| Mixture   | Blank Beads | 05.54±0.14            | 24.42 (%)                | 25.33 (%)           | 2293.00±0.54  | 28.16  | 7.40±0.11  |
|           | Unstarved  | 02.27±0.09            | 69.03 (%)                | 63.33 (%)           | 1213.33±0.63  | 61.98  | 7.53±0.22  |
|           | Starved    | 01.50±0.04            | 79.54 (%)                | 84.00 (%)           | 1129.00±0.58  | 64.63  | 7.43±0.23  |
|           | Free cells | ----                  | ---- (%)                 | ---- (%)            | ----         |        |           |
|           | Initial    | 07.33±0.10            | 01.50±0.09 (%)           | 3191.67±0.74 (%)    | 07.00±0.00  |        |           |

Means marked with the same superscript letters are not significant (P>0.05), whereas others with different superscript letters are significant (P<0.05).

Data in (Fig.4) showed that repeated incubation of starved and unstarved *Phormidium* alginate beads in WI, WII and their mixture in a semi-continuous system for 5 consecutive cycles (10 days each) caused higher ammonia and
phosphorus removal especially in the first three cycles with higher percentage removal in starved than unstarved beads treatments. After the third cycle percentage removal of ammonia and phosphorus began to decrease. Ammonia removal percentage reached 100% in waste I in both starved and unstarved beads where as it recorded 99.49% and 97.56% for waste II and for the mixture 97.82% and 94.54% in the second cycle. Phosphorus removal percentage reached 100% in starved WI in all cycles where as it reached 100% removal in cycle three for WII and mixture. Regarding to COD, higher percentage reduction was recorded especially in the first two cycles with higher percentage removal in starved than unstarved beads. The highest COD reduction percentage was obtained in the second cycle recording higher values in starved than unstarved Phormidium beads (86.73%, 69.26% and 66.52%) for WI, WII and their mixture. These data are in agreement with (Cruz et al., 2013) who reported that degradation of beads is relatively slow; hence, there is sufficient time to efficiently remove nutrients by microbial agents immobilized inside the beads. Still the beads survived long enough to allow completion of the removal of phosphorus, ammonia from wastewater; the contaminants in the wastewater significantly reduced the mechanical strength of the beads. Faafeng et al. (1994) observed the degradation of sodium alginate beads, used for the immobilization of Selenastrum capricornutum, after keeping them in polluted wastewater with high phosphorous (P) and nitrogen (N) content for longer than two weeks. This instability of alginate happens when the gels are in contact with cations that serve as chelating agents and anti-gelling cations, such as dissolved phosphorus, EDTA citrates, sodium bicarbonate, and several more (Moreira et al., 2006). Ruiz-Marin et al. (2010) found that Scenedesmus obliquus when tested in semi-continuous mode was more effective in removing N and P for longer periods than batch cultures. Immobilized S. obliquus removed 97% and 90% of ammonium in AWI (Artificial wastewater immobilization) and UWI (Urban wastewater immobilization), respectively, during the first 48 h. After 250 h the removal decreased to 30 and 10%. A similar pattern was observed for phosphorous removal where 85% removal in AWI was maintained for four cycles, after which the system had a removal capacity of only 30%. They concluded that practically the semi-continuous culture of S. obliquus does not appear to allow a sustainable culture to treat wastewater which is in contrary to our results.
Increasing algal cells entrapped within the beads didn't cause any significant improvement in nutrient removal (Lau et al., 1997). On the contrary, superconcentrated cell stockings in the beads posed serious leakage problem (Robinson et al., 1986). The study of (Abdel Hameed, 2007), indicated that the beads concentrations in wastewater did not affect the efficiency of phosphate removal. Also phosphate might be precipitated as calcium phosphate due to the presence of calcium ions in the alginate matrix (Jimenez- Perez et al., 2004) and wastewater together with elevated pH values. Finally one can say that the immobilized cells either starved or unstarved showed greater efficiency for nutrient removal even after five repeated cycles. The present study recommended the upgrading of traditional operation systems to algal cell immobilization technique with low cost.

During the present study, pH levels increased initially and thereafter remained in the range 7.3 to 9.1 (Fig.3). Increase in pH during phycoremedation is the usual behavior of blue green algae which favors their growth (Promya et al., 2008). Increasing pH represent the indirect role of microalgae to improve the treatment process. High levels of pH alter the physicochemical environment of the wastewater and cause phosphate and metals precipitation, ammonia stripping and disinfection (Abdulsada, 2014). pH is considered as indicator of several biochemical activities, whose the photosynthesis and the biodegradation of the organic matter, the high pH values are attributed to higher photosynthetic rates of algae (Mahapatra et al., 2013). Microalgae species have different pH requirements. Chlorella vulgaris can grow in broad range of pH however the maximum growth rate and biomass productivities are reported at pH 9–10 (Daliry et al., 2017).
Fig. 4: Effect of repeated treatments of immobilized starved and unstarved *Phormidium fragile* cells on ammonia and phosphorus removal and COD reduction from pharmaceutical wastes I, II and their mixture (5 consecutive cycles, each cycle 10 days) incubation in a semi-continuous system.
**Conclusion**

- If cyanobacterium *Phormidium fragile* is applied to natural discharge of pharmaceutical wastewater, it can utilize the pharmaceutical wastewater as nutrient source and will help in reducing the toxicity and facilitate recycling and reutilization of polluted water before discharging into surface aquatic systems providing a low cost, with no secondary pollution and naturally renewable technology.
- Starved *Phormidium* alginate beads are more effective in removing N and P from wastewater than unstarved and free cells or blank alginate beads.
- Repeated incubation of immobilized starved and unstarved *Phormidium* beads in W1, WII and their mixture in a semi-continuous system (5 consecutive cycles) caused increase in nutrients percentage removal which gives a promise to upgrading immobilization technique for wastewater treatment with low cost.

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معالجة مياه الصرف الصيدلانية باستخدام طحالب السيانوبكتيريا الحرة والمقيدة

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تعتبر الطحالب من أهم عوامل المعالجة البيولوجية و التي يتم استخدامها في معالجة مياه الصرف.

يعتبر هذا البحث محاولة لدراسة إمكانية معالجة مياه الصرف الصيدلانية الناتجة عن المعامل WI والعمليات الإنتاجية WII في مصنع للأدوية باستخدام السيانوبكتيريا فورميديا فراجيل الحر والمقيدة. بعد التجربة الأولية أظهرت مياه الصرف الصيدلانية تراوح نسبة عالية، فقط تركز 5% كمية من الرياح، بينما تم استخدام WI و WII الجغرافية أظهر الرياح عن تركز 50% في ظروف الرياح الواحة. وكان للطحالب دوراً فعالاً في المعالجة البيولوجية حيث كانت أعلى نسبة إزالة للعوامل المختبرة 51.56%، 58.89%، 66.67%، 61.23% للأساتذة، 54.55% للأساتذة، 37.05% للأساتذة، 25% للأساتذة والمختبرات بالمختبرات المختلفة.

أظهرت المعاملات التي دون طحالب تسبب التيزواج من النباتات لمدة 36 ساعة قبل الزراعة في زيادة نسبة إزالة الأدوية للفوسفور والأساتذة الكيميائي المستهلك مع زيادة جرعة الطحالب. مع زيادة الأدوية المستهلك بالخلايا غير المعالجة حيث سجلت 100% و 72.56% للمختبر الأول و WI و 100% و 72.8% للمختبر الثاني و 71.98% للمختبر الثاني و 71.98% للمختبر الثاني على الترتيب مع زيادة جرعة الطحالب إلى 40 مل. كما وجد أن شدة الإضاءة 2500 lux و درجة حرارة 25 °C كانت الأفضل في إزالة المغذيات.

خلال هذه الدراسة ارتفعت مستويات الأكسجين والكربونات الكلية للطحالب في المعالات المختلفة. خلال هذه الدراسة أظهرت مياه الصرف الصيدلانية تراوح بين 7.3 إلى 9.1. أظهرت المعالات باستخدام كريات الأنجيبات للمياه زادت معالجة في إزالة كل من الأمونيا، الفوسفور، الأكسجين، السكر، الكربونات الكلية و تراجعت بين 7.3 إلى 9.1. أظهرت المعالات ينعتي مع كريات الأنجيبات الحرة في كريات الأنجيبات المخلوطة، والخلايا الحرية في كلا المختبرين، وكذلك المخلوطة منها. عند تحضر كل من كريات المخلوطة و غير المخلوطة في مختبر WI و مختبر WII، بالإضافة إلى إمكانية معالجة مصر للصرف لتكنولوجيا أقل.