APPLICATION OF A BACTERIAL EXTRACELLULAR POLYMERIC SUBSTANCE IN HEAVY METAL ADSORPTION IN A CO-CONTAMINATED AQUEOUS SYSTEM

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ABSTRACT

The application of a bacterial extracellular polymeric substance (EPS) in the bioremediation of heavy metals (Cd, Zn and Cu) by a microbial consortium in a hydrocarbon co-contaminated aqueous system was studied. At the low concentrations used in this work (1.00 ppm of each metal), it was not observed an inhibitory effect on the cellular growing. In the other hand, the application of the EPS lead to a lower concentration of the free heavy metals in solution, once a great part of them is adsorbed in the polymeric matrix (87.12% of Cd; 19.82% of Zn; and 37.64% of Cu), when compared to what is adsorbed or internalized by biomass (5.35% of Cd; 47.35% of Zn; and 24.93% of Cu). It was noted an increase of 24% in the consumption of ethylbenzene, among the gasoline components that were quantified, in the small interval of time evaluated (30 hours). Our results suggest that, if the experiments were conducted in a larger interval of time, it would possibly be noted a higher effect in the degradation of gasoline compounds. Still, considering the low concentrations that were evaluated, it is possible that a real system could be bioremediated by natural attenuation process, demonstrated by the low effect of those levels of contaminants and co-contaminants over the naturally present microbial consortium.

Key words: EPS, co-contamination, heavy metals, natural attenuation.

INTRODUCTION

Accidental and deliberate crude oil spills have been a significant source of environmental pollution, as well as metal based and metal finishing industries. Determination and characterization of the nature of combined pollution is a problem of current interest. Waterways, rivers and ultimately the seas are contaminated with organic waste containing high concentrations of heavy metals, which are of general concern as they are persistent and able to accumulate in the ecosystem posing serious environmental and health hazard (1,17,20). Still, the presence of heavy metals can inhibit the natural microbiota involved in the degradation of organic compounds and affect biodegradation rates. The level of inhibition will depend on the concentration and availability of the heavy metals and the nature of the metals, media and microbial species. Heavy metals inhibit microorganisms by blocking essential functional groups or interfering with essential metal ions incorporation of biological molecules. In some cases microorganisms are resistant to some heavy metals through different possible mechanisms (2).

Conventional processes used for removal of heavy metals from industrial wastewaters include chemical precipitation, oxireduction, filtration, electrochemical techniques and sophisticated separation processes using membranes. These processes are usually expensive, and this characteristic stimulates the use of alternative biotechnologies, due to their reduced cost and lower aggressiveness to the environment (4).

Effective strategies to enhance organic biodegradation in the presence of toxic metals include reducing the bioavailable concentration of the toxic metal and reducing interactions of the toxic metal with the cell. In their work, Sandrin et al. (17) carried out an experimental system to determine the effect of a rhamnolipid, which is an EPS produced by Pseudomonas aeruginosa 9027, on the capability of a metal-sensitive microorganism to degrade...
organic contaminant. Their results suggested that rhamnolipid reduces Cd induced inhibition of naphthalene degradation through a combination of Cd complexation and release of lipopolysaccharide (LPS) from the cell.

Extracellular polymeric substances (EPS) play a very important role in biosorption of heavy metals, and are produced by most bacteria. EPS comprise a mixture of polysaccharides, mucopolysaccharides and proteins, which depends on the strain and its culture conditions. They contain ionizable functional groups such as carboxyl, phosphoric, amine, and hydroxyl groups, which enable EPS to sequester heavy metals. Ion exchange, complexation with functional groups of negatively charged, adsorption and precipitation are the mechanisms involved in metal biosorption onto EPS (25).

The clean-up of gasoline-contaminated groundwater continues to be a challenge. The major gasoline constituents (benzene, toluene, ethylbenzene and xylene - BTEX) are relatively soluble in water and are considered human carcinogens. Thus, the level of BTEX in water must be highly controlled and limited to low values (6). According to Van Stemvooort et al. (24), one approach that has been recently recognized for groundwater applications is natural attenuation. Natural attenuation as a management strategy for contaminated soils and sediments hinges on the notion that some transformation processes involving the in situ microbial metabolism of the target compounds are self-sustaining, appropriate with regard to type and sufficient in magnitude to control the risk associated with resident pollutants (19). Biotechnological approaches to the abatement of toxic metal and organic pollution consist of selectively using and enhancing the natural processes to treat particular wastes (22).

The aim of this work was to investigate the effect caused by the application of a bacterial EPS in bioremediation of an aqueous system co-contaminated with gasoline and heavy metals (Cd, Zn and Cu) by a microbial consortium.

MATERIALS AND METHODS

Production, extraction and partial purification of EPS

In order to produce EPS, it was used the Gram-positive spore-forming bacterium Paenibacillus polymyxa, isolated from fermented sausages, as described by Piuri et al. (13). The microorganism was cultivated in a solid medium of following composition (adapted from Sharma et al. (20)): sucrose 5.00 g/L; yeast extract 0.15 g/L; KH₂PO₄ 0.50 g/L; (NH₄)₂SO₄ 1.0 g/L; MgSO₄.7H₂O 0.41 g/L; agar 30.00 g/L; and pH 7.0, over 2 days at 30°C. After this time, the culture was kept under refrigeration at 10°C. In order to activate and to obtain a pre-inoculum, subsequent cultivation took place in 400 mL of a medium of same composition as described above, in absence of agar, and incubated for at 30°C on a rotatory shaker at 150 rpm, for 24 hours. The initial cell concentration was 0.01 g/L.

The grown cells were heated until 100°C for 15 minutes in order to release the EPS adsorbed on cell surface. The medium was concentrated with a rotary vaporizer and further centrifuged at 3000 rpm for 30 minutes, for removing the cells. The volume of supernatant was measured and three volumes of cold ethanol were added for EPS precipitation, and the mixture was kept under refrigeration for 24 hours. After new centrifugation, at same conditions described above, precipitated EPS was dialyzed overnight in distilled water under refrigeration. Partial purified EPS was dried in a stove at 50°C until constant weight.

Obtainment of soil

The soil sample used to propagate the microbial consortium was obtained in the garden of Escola de Química - UFRJ. This soil has no history of contamination with hydrocarbons and heavy metals, and because of that it would be necessary the adaptation of the microbial consortium to those conditions.

Obtainment of microbial consortium

Microbial consortium used in the experiments was obtained as described: 10 g of a soil sample were added to an Erlenmeyer flask with 100 mL of mineral medium, whose composition was NaCl 5.00 g/L; KNO₃ 3.00 g/L; KH₂PO₄ 1.00 g/L; (NH₄)₂SO₄ 1.00 g/L; NH₄H₂PO₄ 1.00 g/L; MgSO₄.7H₂O 0.20 g/L; and pH 7.0. Gasoline 1.00% (v/v) was used as carbon source. The mixture was incubated at 30°C in a rotatory shaker at 150 rpm, for 4 days. After that, a sample of 10 mL was transferred to another Erlenmeyer flask, with 100 mL of same mineral medium and gasoline 1.00% (v/v). This new mixture was incubated at the same conditions described above. This procedure was repeated twice, and the last flask with the microbial consortium was kept under refrigeration.

A simple macroscopic characterization of the microbial consortium colonies was carried out with plates from the last propagation step, in TSA medium (5), of following composition: gasoline 0.10% (v/v), NaCl 5.00 g/L; K₂HPO₄ 2.50 g/L, tryptone 17.00 g/L, yeast extract 2.50 g/L, agar 20.00 g/L, and pH 7.0. Some dilutions of culture after 72 hours were done at 30°C, and some macroscopic characteristics of colonies were observed.

Obtainment of inoculum

In order to obtain the inoculum for the conduction of the experiments, a pre-inoculum was previously prepared in an Erlenmeyer flask containing 400 mL of mineral medium, amended with glucose 0.50 g/L and gasoline 0.10% (v/v). Twenty milliliters of the system containing the microbial consortium previously obtained was added to the medium, and it was cultivated for 4 days, under 30°C in a rotatory shaker at 150 rpm. From this culture it was obtained a volume of inoculum to start the experiments, with initial optical density of 0.05, measured at 440 nm, with a CamSpec spectrophotometer, model M302.
**Effect of the presence of heavy metals and/or EPS in cellular growing**

Experiments were carried out in duplicate, in 200 mL of mineral medium, amended with gasoline 0.10% (v/v), containing or not 1.00 ppm of each metal (Cd, Zn and Cu) and/or a solution of EPS at the final concentration of 0.05 g/L. “Experiment 1” was carried out only in presence of gasoline; “Experiment 2” was carried out in presence of gasoline and heavy metals; “Experiment 3” was carried out in presence of gasoline and EPS; and “Experiment 4” was carried out in presence of gasoline, heavy metals and EPS. A “Control 1” flask without those components was submitted to the same conditions of the experiments. Table 1 presents the compositions of the media used in different experiments. The volume of inoculum added to the media was such that the initial optical density was equal to 0.05. The flasks were incubated in a rotatory shaker at 30ºC and 150 rpm for 30 hours. Samples were taken from each system after 0, 6, 11 and 30 hours of cultivation.

**Biomass quantification**

Microbial growing was indirectly determined by intracellular protein quantification, according to Sandrin et al. (17), with the use of Lowry’s method, after biomass hydrolysis with 10% (v/v) of NaOH 1 M at 100ºC for 10 minutes. After reaction, absorbance was determined with an spectrophotometer CamSpec, model M302, at 660 nm.

In order to relate samples absorbance to protein concentration, a BSA (bovine serum albumine) standard curve was built. The results of protein concentration results were the average of two values obtained for each sample, with their respective standard errors.

**Effect of presence of EPS in heavy metals removal**

Experiments were carried out in 200 mL of mineral medium, amended with gasoline 0.10% (v/v), 1.00 ppm of each heavy metal (Cd, Zn, and Cu), and a solution of EPS at a final concentration of 0.05 g/L. The volume of inoculum added to the media was such that the initial optical density was equal to 0.05. The flasks were incubated in a rotatory shaker at 30ºC at 150 rpm for 30 hours. A “Control 2” flask without inoculum was used in order to evaluate gasoline abiotic losses, as well as to evaluate the heavy metals adsorption by EPS in absence of cells. A second flask without EPS (“Experiment 5”) was used in order to evaluate the heavy metals adsorption and absorption by cells in absence of EPS. “Experiment 6” involved the presence of inoculum, gasoline, heavy metals and EPS. Table 2 presents the compositions of media used in this step of the work.

**Table 1. Compositions of media related to experiments of evaluation of heavy metals removal by cell and/or EPS.**

| Inoculum     | Gasoline 0.10% (v/v) | Cd, Zn and Cu (1 ppm each) | EPS 0.05 g/L |
|--------------|----------------------|---------------------------|--------------|
| Control 1    | no                   | no                        | no           |
| Experiment 1 | yes                  | no                        | no           |
| Experiment 2 | yes                  | yes                       | no           |
| Experiment 3 | yes                  | no                        | yes          |
| Experiment 4 | yes                  | yes                       | yes          |

**Table 2. Compositions of media related to experiments of evaluation of heavy metals removal by cell and/or EPS.**

| Inoculum     | Gasoline 0.10% (v/v) | Cd, Zn and Cu (1 ppm each) | EPS 0.05 g/L |
|--------------|----------------------|---------------------------|--------------|
| Control 2    | no                   | yes                       | yes          |
| Experiment 5 | yes                  | yes                       | no           |
| Experiment 6 | yes                  | yes                       | yes          |

**Procedures to get samples for heavy metals and hydrocarbons of gasoline quantification**

In these experiments, heavy metals were quantified in five different fractions: C0, C1, C2, C3 and C4.

Before addition of inoculum, a sample from medium was obtained for initial heavy metals measurements (C0). After addition of the inoculum, experiments were conducted for 30 hours. After this time, a sample of 40 mL was taken and centrifuged at 3000 rpm and 30 minutes, in order to separate the cells. These cells were kept for further analysis. Supernatant was divided in three parts: the first one was used for total soluble metals measurements after 30 hours (C1), the second one was used to quantify EPS adsorbed metals (C2) after cold ethanol precipitation, and the third one to quantify the hydrocarbons. System was filtered with a 0.22 mm membrane, in order to separate EPS, and the filtered was used for soluble metals measurements (C2'). C2 was determined by C1 minus C2'.

Cells previously obtained were washed with 35 mL of a solution of EDTA 10 mM. System was centrifuged, at the same conditions described before, and supernatant was used for cells surface adsorbed metals (C3) measurements. Cells were resuspended in 35 mL of distilled water, and the suspension was hydrolyzed with 10% (v/v) of NaOH 1 M, at 100ºC for 10 minutes. Samples were then centrifuged as already described, and supernatant was used to quantify the metals inside cells (C4).

The removal efficiency (η) was defined as:

$$\eta = \frac{C_0 - C_n}{C_0} \times 100$$

[1]

where n is 1, 2, or 3.
The metal uptake by cells or EPS \((q)\) was defined as:

\[ q = \frac{C_0 - C_m}{C_x} \]  

where \(m\) is 2 or 3, and \(C\) can be either EPS concentration or intracellular protein concentration, both in g/L.

**Gasoline fractions quantification**

The fractions of gasoline were quantified with a GC Trace Thermofinnigan, acceplled to a Headspace Tekmar Dohrmann 7000 self-sampling. The chromatograph was equipped with a PID detector and a 30 m x 0.53 mm (diameter) Megabore column. The make-up gas (N\(_2\)) flow rate was 6 mL/min, and the sheat gas (N\(_2\)) flow rate was 40 mL/min. The injection port temperature was 230ºC, and the PID detector temperature was 250ºC. The temperature program was: initial temperature 45ºC, held for 1 minute, programming rate 12ºC/min to 200ºC, held for 3 minutes. Peak areas and retention times were compared to reference standards. The injections were done in triplicate.

**Heavy metal measurements**

Soluble metals were quantified with an atomic absorption spectrometer Perkin Elmer, model 3100, with the following conditions: flame: air/acetylene, wave weight 228.8 nm (Cd), 213.9 nm (Zn), and 324.8 nm (Cu).

**RESULTS AND DISCUSSION**

**Effect of metals on microbial consortium growing**

Fig. 1 presents microbial consortium growing in mineral media, as described in Table 1.

As can be seen in Fig. 1, the maximum cellular growing is higher and it is reached faster in the medium containing only gasoline. The falling of the cellular concentration could be explained by the depletion of the energy source, and the posterior cellular growing is probably due to a possible consumption of the EPS or of some components released by cellular lysis in such conditions. In the medium amended with heavy metals at the concentrations used in these experiments, their presence caused a lengthening of the lag phase, probably due to an adaptation period of the microorganisms to the co-contaminated system, and the cellular growing seemed to be slower until it gets the stationary phase. This effect does not seem to greatly affect growing after ten hours of cultivation, and curves almost get the same final biomass values. Malik (11) reported that several bacterial isolated from effluents exhibited resistance to extreme conditions like pH peaks or high metals concentrations. According to Hassen et al. (7), at low concentrations, some metals (e.g. Co, Cu, Zn, Ni) are essential to microorganisms metabolism, since they provide vital cofactors for metallo-proteins and enzymes. In naturally polluted environments, the response of microbial communities to heavy metals depends on the concentration and availability of these metals and is dependent on the actions of complex process, controlled by multiple factors such as the type of metal, the nature of medium and microbial species.

In “Experiment 3”, containing gasoline and EPS, it is possible to observe that it does not occur an improvement of cellular growing, suggesting that the microorganisms do not use EPS as a carbon source. Experiments carried out with metals in presence of EPS (“Experiment 4”) resulted in a similar biomass final growing, indicating that the presence of EPS does not promote any growing improvement, as confirmed by Malik (11), although it seems that some little initial increase in the curve takes place when EPS is present. According to author, multispecies consortia are considered more efficient over the monospecies culture due to greater resistance against environmental fluctuations and the metabolic relations among the member strains. Sandrin et al. (17) observed a microbial growing improvement of *Bulkholderia* sp. in presence of Cd, caused by addition of a rhamnolipid produced by *Pseudomonas aeruginosa* strains. They confirmed that the presence of rhamnolipid reduced significantly the toxic effect caused by high concentrations of Cd (approximately 10000 ppm) and naphtalene (26% (v/v)) in cultivation medium.

The plattings for macroscopical characterization of microbial consortium presented two different species of bacteria (one of rosy color and other one of white color), and a specie of filamentous fungus of white color. These cultures seemed to be dominant along all experiments, once it was not observed any significant macroscopic variation of the colonies of the microbial consortium.

Results presented in Fig. 1 confirm that microbial consortium really seems to be adapted to heavy metals concentrations used in these experiments, once no negative effect was verified in
the microbial growing, and the same final biomass concentration was virtually attained.

**Effect of the presence of EPS in metals removal**

Table 3 presents data concerning metal removal efficiency \((h)\) by adsorption by cells and EPS, and incorporated within the cells only.

Table 3. Heavy metals removal efficiency in experiments carried out in the absence and/or presence of EPS, with or without microorganisms.

|                          | Removal efficiency (%) |
|--------------------------|------------------------|
| Experiment in absence of EPS | Cd  | Zn  | Cu  |
| Metal adsorbed on cell surface | 4.33% | 39.05% | 12.36% |
| Metal internalized by cells   | 1.77% | 15.78% | 9.36%  |
| Experiment in presence of EPS | Cd  | Zn  | Cu  |
| Metal adsorbed on cell surface | 3.51% | 30.33% | 14.57% |
| Metal internalized by cells   | 1.84% | 17.02% | 10.36% |
| Metal adsorbed on EPS         | 87.12% | 19.82% | 37.64% |
| Experiment in presence of EPS and without microorganisms | Cd  | Zn  | Cu  |
| Metal adsorbed on EPS         | 98.00% | 53.25% | 51.69% |

It is possible to note in Table 3, in relation to experiments containing EPS or not, that at the experimental conditions of cellular growing and during time cultivation, the greatest part of Cd and Zn is adsorbed on cell surface than inside cells. The amount of Cd that is internalized by cells is lower than Zn and Cu, and can be explained by Cd higher toxicity to microorganisms (9,16). Once Zn is an essential metal to cells, so as Cu, the amount of both metals that was absorbed by cells was relatively high, and the quantity of Zn that is internalized is bigger probably because it is involved in many cell enzymatic activities (9).

Hassen et al. (7) presented results involving a decreasing order of toxicity in *Pseudomonas aeruginosa, Pseudomonas paucimobilis* and *Proteus mirabilis* growing: Hg>Cd>Cu=Cr>Zn. Castro-Silva (3) related that bacteria and fungi isolated from coal mining are resistant to high levels of Cd, Zn and Cu, being this resistance associated to the capacity to accumulate these elements.

The amount of cadmium adsorbed on cell surface was lower than Zn and Cu, also probably due its higher toxic effect. Once Zn is part of several biocatalytic functions, which include about 200 enzymes, acid nucleic metabolism and cell division (9), it probably explains why this metal was adsorbed and internalized in higher quantities when compared to Cu.

In regard to metals remotion by EPS, it is possible to verify that removal efficiency of Cd was higher than Zn and Cu, in media containing inoculum or not. This fact could be justified by a better metal affinity to EPS than cell surface. Figs. 2a and 2b show metal uptake by cell surface and EPS.

Metal uptake by soluble EPS was affected by presence of cells (Fig. 2a), probably due to a protector biofilm formation around the cells, reducing the amount of free EPS available to adsorption of soluble metals. According to Priester *et al.* (14) and Ramey *et al.* (15), bacterial biofilms are tri-dimension structures that consist on extracellular polymeric substances. In this way, it is also possible to explain why metals adsorption by cells was a little higher in medium containing EPS (Fig. 2b). Once cell surface was surrounded by EPS, the quantity of heavy metals adsorbed would be higher.
Gasoline hydrocarbons consumption

In order to evaluate gasoline hydrocarbons consumption by the microbial consortium in experiments carried out in absence or presence of EPS, the fractions of benzene, toluene, ethylbenzene, meta- and para-xylene, and ortho-xylene were quantified. Results are presented in Table 4. These results were calculated deducting losses by evaporation in relation to “Control 2”, carried out in absence of biomass. In regard to benzene and xlenes, it was not observed any decrease in chromatogram peaks, suggesting that these components were not consumed, the reason why these components are not presented in Table 4. The presence of EPS did not affect toluene consumption, once the results related to this hydrocarbon are statistically the same. In the other hand, it is possible to observe that, although cell growing was almost the same in media containing or not EPS, at the end of 30 hours, the presence of EPS lead to a higher consumption of ethylbenzene, which could be justified by the probable protector effect that EPS causes to cell. Merroun et al. (12) report that toxic heavy metals accumulate mainly within the EPS in experiments carried out with Myxococcus xanthus, and this fact would suggest that extracellular polymers could protect this myxobacterium in its habitats because it fixes heavy metals around the cell, avoiding their absorption. According to Tsuneda et al. (23), an EPS covering on a cell surface alters the physicochemical characteristics of the surface such as polymeric property, hydrophobicity and charge, what could favoring ionic exchange.

The heavy metals and gasoline concentrations used in this work were low; however, they were higher than the concentrations allowed by Brazilian laws to several classes of sweet and salted waters. It was possible to observe that such concentrations did not affect significantly microbial growing, suggesting that natural attenuation of the system is a very appropriate alternative in cases where contamination by heavy metals and hydrocarbons does not reach high levels. Although metal residual concentrations still are higher than the allowed for discharges by legislation, the short time in which these experiments were carried out, and the possibility of continuous cellular growing verified in Fig. 1, allow us to suppose that it would be possible to occur a higher consumption of organic compounds, and the system could reach the limits of discharge for both kinds of pollutants.

According to Malik (11), in many times biosorption alone may not be sufficient for effective metal remediation. Under such situation, application of active and growing cells might be a better option due to their ability of self-replenishment, promoting a continuous metabolic uptake of metals after physical adsorption, and the potential for optimization through development of resistant species and cell surface modification.

In the other hand, there is still a possibility of production of EPS by the microbial consortium used in these experiments, what could also justify its better resistance to heavy metals. Roane et al. (16) report that strains of Pseudomonas H1 and Bacillus H9, isolated from soil, showed evidences of EPS production, and were resistant to high Cd concentrations.

Our results also showed that the metal uptake and removal efficiency by EPS were relatively high, and it could represent a potential application in co-contaminated systems with higher concentrations of hydrocarbons and metals, avoiding the negative effect caused by heavy metals in the biodegradation of the organic pollutant, as related by Hoffman et al. (8), Lin et al. (10) and Sandrin et al. (17).

**CONCLUSIONS**

In this study, the effect of application of a bacterial EPS in bioremediation of a co-contaminated system was evaluated. Our results showed that the small concentrations of heavy metals and hydrocarbons used in these experiments did not affect the microbial growing, as the EPS application did not seem to influence such growing. However, its presence in medium increased about 24% ethylbenzene consumption. EPS presented good sorption of soluble metals, demonstrating its potential application in metal detoxification, permitting an improvement of microbial activity over hydrocarbons. In the other hand, our results also indicated that a system with low concentrations of these organic and inorganic pollutants, which could be found in natural systems after a contamination with hydrocarbons, could possibly be susceptible to natural attenuation, if process is conducted in a larger period of time, without the necessity of intervention, once the microbial consortium presented good resistance and adaptation to the conditions used in these experiments.

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RESUMO
Avaliação de diferentes tecnologias de biorremediação de metas pesados em um sistema aquoso co-contaminado

A aplicação de uma substância polimérica extracelular (EPS) bacteriana na bioremedição de metas pesados (Cd, Zn e Cu) por um consórcio microbiano em um sistema aquoso co-contaminado com hidrocarbonetos foi estudada. Nas baixas concentrações usadas neste trabalho (1,00 ppm de cada metal), não foi observado um efeito inibitório no crescimento celular. Por outro lado, a aplicação da EPS bacteriana levou a uma menor concentração de metas livres em solução, uma vez que grande parte destes fica adsorvido na matriz polimérica (87,12% de Cd; 19,82% de Zn; e 37,64% de Cu) quando comparado ao que é adsorvido ou interiorizado pela biomassa (5,35% de Cd; 47,35% de Zn; e 24,93% de Cu). No pequeno intervalo de tempo avaliado (30 horas) e na baixa concentração de gasolina utilizada (0,1% (v/v)), foi percebido um aumento de 24% no consumo de etilbenzeno, entre os componentes da gasolina que foram quantificados. Nossos resultados sugerem que, se os experimentos fossem conduzidos em um intervalo de tempo maior, possivelmente poderia ter sido observado um maior efeito na degradação dos componentes da gasolina. Ainda, considerando as baixas concentrações avaliadas, é provável que um sistema real pudesse ser biorremediado pelo processo de atenuação natural, tendo em vista o baixo efeito desses níveis de contaminantes e co-contaminantes sobre o consórcio microbiano naturalmente presente.

Palavras-chave: EPS, co-contaminação, metas pesados, atenuação natural.

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