Removal of Organic Pollutant from Waste Water

Abeer El-Saharty

National Institute of Oceanography & Fisheries (NIOF).

ABSTRACT

This study investigated the possibility of removing organic pollutants from water by using organo-Bentonite. The organo-bentonites were synthesized by ion exchange reaction using quaternary ammonium or phosphonium organic salts. The resultant organo-bentonite complexes were lyophilized, ground to 5μm and used as test materials. Bromoxynil, is a widely used herbicide was taken as a standard water pollutant in this study because it has been detected in many water samples around the globe. The experimental design of this study based on adsorption/desorption techniques. Bromoxynil concentrations in equilibrium solution were determined by HPLC. Results showed that the amounts of bromoxynil removed from water by using organo-Bentonite (Bentonite-NCP, and/or Bentonite-HDTBP) were several times higher than the amount removed by using hydrophilic bentonite. The removed amount of bromoxynil by using organo-Bentonite was further enhanced by lowering the pH of the solution, and optimizing the temperature for the adsorption reaction. Extraction of bromoxynil from NCP-, or HDTBP-bentonite was nearly similar to those removed from water. These results indicate the efficacy of organo-bentonite to remove organic pollutants from water, accordingly application of organo-bentonite may be advantageous materials for removing organic pollutants at a large scale.

INTRODUCTION

Bromoxynil is a nitrile herbicide, widely used in the region for annual weed control. Its application creates environmental contaminations. For instance, it was detected in air samples [1] water samples [2], and soil samples [3]. It application may create health problems [4]. Furthermore, bromoxynil has been reported as a toxic agent to important green algae in the ecosystem [5].

Application of commercial formulations of bromoxynil (emulsion concentrate, and/or suspension concentrate) in the ecosystem may induce species shifts within the communities and could affect the structure and the function of the aquatic communities [6].

Worldwide, very little information about removal of bromoxynil from water are available, beside the fact that no attempt has been made in Gaza to remove or attenuate bromoxynil concentration in the ecosystem. In this article we attempted to design suitable organo-clay complexes and optimize the aquatic pH to enhance removal of bromoxynil from water.

MATERIALS AND METHODS

2.1 Synthesis of organo-Bentonite complexes:

Ca$^{2+}$-bentonite clay (M48) with cation exchange capacity (CEC) equivalent to 960 mmol (+) kg$^{-1}$ (= 0.96 meq g$^{-1}$) [7] was used. N-cetylpyridinium bromide (NCP+Br$^{-}$) and N-hexadecyl tributylphosphonium bromide (HDTBP+Br$^{-}$) and Bromoxynil (HPLC grade, 3,5-dibromo-4-hydroxybenzonitrile) were purchased from Sigma Chemical Co., Germany.

The organo-Bentonite complexes were synthesized by simple ion-exchange reaction. In this procedure, 5 mmol of the solid organic salt was added to 1L of 1% (w/v) aqueous suspension of M48 under stirring conditions for 3 days [8]. Bromoxynil stock solution was prepared by dissolving 31 mg in 2-3 mL methanol and diluting to 1 L with deionized water.

© 2014 AENSI Publisher All rights reserved.
2.2 Removal of bromoxynil from water:

0.5 mg of bromoxynil was added to various glass centrifuge tubes containing 10 mg of different organo-Bentonite. The tubes were shaken for 48h and then the supernatants were collected by centrifugation at 20000g. The remaining concentration of bromoxynil in the supernatants were determined by HPLC as mentioned below.

2.3 Influence of bromoxynil concentrations on the removal process:

Thirty ml of bromoxynil concentrations ranged between 0.66-31 mg L⁻¹ were transferred to 30 mL glass centrifuge tubes containing 0.005 g organo -Bentonite. The tubes were kept under continuous rotary agitation for 48 hours to maintain aqueous suspension during the experimental time. This experiment was maintained at pH7. Control samples were made by performing the same procedures under the same conditions without adding the organo-Bentonite to each tube. The aqueous solutions were collected by centrifugation at 20,000 g for 0.5 h.

2.4 Influence of pH on removal of bromoxynil:

Deionized water was used to perform the removal experiments at pH 7, whereas few drops of acetic acid were added to the stock solution of bromoxynil to adopt the pH to 3. The removal experiments were performed as described above using HDTBP-Bentonite complex.

2.5 Influence of Temperature on removal of bromoxynil:

Deionized water was used to perform the removal experiments at 5, 17 and 40 oC using NCP-Bentonite complex as a removing materials

2.6 Measurements of bromoxynil:

The concentrations of bromoxynil in the supernatants were determined by Waters 717 HPLC with UV detector (detection wavelength 283 nm). Column: Nova-Pak C18 (inner diameter 3.9 mm, length 150 mm), flow rate: 1 mL min⁻¹. The mobile phase was methanol/water 50/50 (v/v). The amount of bromoxynil adsorbed was calculated from the depletion of the bromoxynil concentration in the aqueous solutions.

2.7 Release of bromoxynil from organo-bentonite complexes:

To insure removal of bromoxynil from water, the used organo-bentonites in the above mentioned experiments were collected, air dried and used for bromoxynil extraction. In this experiments the air dried organo-bentonites were transferred to test tube containing 5 ml of water methanol mixture as described in mobile phase. The test tubes were transferred to an ultrasonic machine functioning at high speed for 3 min. the solution was collected and additional 5 ml of water methanol mixture was added again and the same procedure was repeated to insure maximum extraction. Concentration of bromoxynil in the solution was determined as mentioned above.

2.8 Data analysis:

The removal data were collected as an average of 3 replicate and the standard deviation was calculated and used as error bars to discriminate differences among treatments. Presenting the standard deviation as error bar is the best method to determine significant differences among adsorption isotherms. It is well known that overlapping of error bars indicating no difference whereas small or extremely small error bars indicate significant difference.

RESULTS AND DISCUSSION

The cationic quaternary ammonium/phosphonium salts used in this study are solid materials at room temperature, and surface-active agents (surfactants), The molecular structures include an aliphatic part and/or an aromatic ring. In a diluted solution (< 0.1 mmol/g) the adsorbed amounts of surfactant on bentonite surfaces were nearly similar (Data not shown), whereas as high added concentration (>0.5 mmol/g) the extent of adsorption became a function with the size and shape of the surfactant. Large organic cations can effectively displace inorganic cation such as Ca²⁺ and/or Na⁺ from mineral surfaces of clay by ion exchange [8,9].

Removal of bromoxynil from water by different organo-bentonite complexes are shown in Figure 1. It can be seen that the removed amounts of bromoxynil on raw Bentonite (clay) or modified Bentonite with NCP and/or HDTBP, are: (14.6); (84.65) and (46.48) mg/g respectively.

The largest removed amount of bromoxynil was observed on bentonite modified with NCP followed by HDTBP. The explanation of these results is that low removed amount using raw Bentonite is due to the hydrophilic surfaces of Bentonite and hydrophobic nature of bromoxynil. Modification of Bentonite surfaces with NCP or HDTBP may have created a microscopic organic phase on Bentonite surfaces as in hexadecyltrimethyl ammonium-smectite [10]. This situation may acts as a solubilizing (partitioning) medium for removing nonionic organic compounds from water [11].
Effects of various concentrations of bromoxynil on the removal process are shown in Figure 2. It is obvious that at low concentration of bromoxynil, the removed amount by using organo-bentonite complexes exceed 90% of bromoxynil from water, whereas at high bromoxynil concentrations (25 mg/l), the removed amount of bromoxynil reduced to 48.02% and 76.57% using Clay-HDTBP or Clay-NCP respectively. Using hydrophilic clay the removed amount of bromoxynil did not exceed 8% and was not affected by the concentration of bromoxynil.

The explanation of these results is that at low concentration of bromoxynil the molar concentration of the adsorption sites equal to 0.0096 meql\(^{1}\). Under these condition, the molar ratio between bromoxynil molecules and the adsorption sites is nearly similar. At a high concentration of bromoxynil the molar ratio between bromoxynil molecules and the adsorption sites tend to increase. Accordingly, the number of free adsorption sites tends to decease, under this condition the removed amount of bromoxynil tend to decease. Furthermore, one can realized from the data in Figure 2 that Clay-NCP is more potent than Clay-HDTBP for removal of bromoxynil from water. The explanation of these results is that Clay-NCP contains an aromatic ring in its chemical structure which may enhance the interaction between bromoxynil molecules and Clay-NCP molecules. The adsorption of organic molecules from water could be enhanced by the efficacy of the chemical structure [12]. Under this condition, hydrophobic interaction between the adsorbed molecules of bromoxynil and those in water may be enhanced. Similar phenomena for HDTMA [13].

Using different fractions of organo-bentonite for removing bromoxynil showed optimal removal at low fraction of organo-bentonite complexes [14]. The explanation of these results is that at low fraction of organo-bentonite complex the adsorption sites are diluted in the solution and available for interaction with bromoxynil from water whereas at high concentration of organo-bentonite complexes the percent removal tends to decrease due to possible aggregation of organo-bentonite complexes in the water clay suspension. Accordingly, the availability of free adsorption sites is limited for removal of bromoxynil. Consequently, the concentration of the adsorption site in water should be optimized. It was shown that the best results were obtained at a very diluted organo-Bentonite complexes (Data not shown).

Influence of pH values in the removed amount of bromoxynil from water is shown in Figure 3.
It is obvious that the removed amount of bromoxynil at pH 3 is several times higher than the removed amount at pH 7. The explanation of these results is that bromoxynil has an acid dissociation constant pKa value of 4.06 [15], accordingly the degree of ionization in the solution is depends on both the pKa value and the pH of the solution in which it is dissolved, a relation described by the Henderson-Hasselbalch equation for weak acids:

\[
pK_a - pH = \log\frac{[\text{nonionized}]}{[\text{ionized}]} \tag{1}
\]

Employing Eq 1 at pH 3, one can realized that the nonionized fraction equals to 11.48 times higher than the ionized fraction whereas at pH 7 the nonionized fraction equals to 0.001 of the ionized faction. As obvious at pH 3 bromoxynil molecules remain in the nonionized from (Hydrophobic form) and adsorbed directly to the organo-bentonite (Figure 3) whereas at pH 7 bromoxynil molecules tend to be ionized (Hydrophilic form) in this case it remains in the solution due to formation on anionic form (Figure 4). This explanation is in accord with the higher removed amount of bromoxynil at pH 3 (Figure3).

The presented results agree with previous reports [3,7,16], who found high adsorbed amount of bromoxynil in soil and wheat char at low pH.

Removal of bromoxynil at various temperatures are shown in Figure 5. It can be seen that removed amount of bromoxynil is more pronounced at 17 °C than at 40 or 5 °C (Figure 5).

The explanation of these results is that at a low temperature (e.g. 5 °C) the chemical potential of bromoxynil molecules is reduced and the molecules tend to from crystals due to low solubility in water (0.13 g.l⁻¹, 20 °C). This agrees with the general concept of solubility in chemistry. In contrast, at high temperature the system absorbs heat energy and an increase in the chemical potential may have occurred. This step makes more bromoxynil molecules available for removal from water due to dynamic mobility in the aqueous solution. The optimal removed amount appears to be at 17 °C used in this study.

These results indicate the efficacy of the used matrixes in removing of bromoxynil. These results are in agreement with recent report [17], who showed that bromoxynil was strongly encapsulated in organo-clay complex and used for slow-release formulations under field conditions.
Fig. 5: Removal of bromoxynil at different temperatures. Error Bars represent standard deviation.

**Conclusion:**

The study reveals that exchanging Bentonite surfaces with organic cations increases their capacity to remove bromoxynil (organic pollutant) from water. The rationale of this work is that removal of bromoxynil from water can be enhanced by modifying the Bentonite surfaces with NCP and HDTBP. The results showed that highest removed amount of bromoxynil was obtained at pH 3 and at 17 °C (Figures 3, 5). Extraction of bromoxynil from the used matrixes was nearly similar to the amount removed in Figure 1. The environmental relevance of this work emerges from the fact that organo-bentonite complexes can be used to remove organic pollutants from water and develop environmentally acceptable herbicide encapsulation for safe application.

**REFERENCES**


