## Phosphorus and ammonia release experiments in the sewage-impacted coastal marine sediments of Alexandria, Egypt.

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## Introduction

Introduction Sorption and desorption of nutrient salts by the sediment particles probably account for about 20-30% of the nutrients turnover in the marine environment (MARTIN, 1970; SEITZINGER, 1987) which play important role in nutrients contribution to the overlying water column. In this paper, our target is to study, experimentally, the amount of phosphorus adsorbed on sediments as well as to measure the sediment bound ammonium-nitrogen characteristic to the marine environment of Alexandria. Study area The Eastern Harbour (E.H) of Alexandria is a relatively shallow semi-closed basin,

The Eastern Harbour (E.H) of Alexandria is a relatively shallow semi-closed basin, sheltered from the sea by an artificial break-water, leaving two openings through which the exchange of the harbour water and the neritic Mediterranean water take place. About 35.2x10<sup>6</sup> m<sup>3</sup> of domestic sewage are discharged into the Eastern Harbour of Alexandria through 11 outfalls, distributed along the coast. This quantity is about 2.3 times the water volume of this basin. Material and Method

Material and Method Representative bottom sediment samples were collected from 3 stations in the Eastern Harbor. The sediment bound ammonium-nitrogen from each station was mechanically released by centrifugation (3600 RPM) a 1 cc wet sediment sample for ten minutes in 150 ml of ammonia free distilled water (KAMIYAMA *et al.*, 1977). The most suitable sediment to water ratio was found to be 1 cc sediment to 150 cc distilled water (MAHMOUD, 1985). The concentration of ammonia in the supernatant distilled water (MAHMOUD, 1985). The concentration of ammonia in the supernatant distilled water was measured. The liberation of ammonia-nitrogen from the sediment sample may result from the disruption of the equilibrium between the dissolved form in the interstitial water and adsorbed form on the particles by addition of the distilled water, and a new equilibrium state reached. The results were expressed as ug/lcc/l. Phosphorus adsorbed on sediments was measured by shaking a 1 cc sediment for one hour with 150 ml of distilled water in an electric shaker. The phosphate released was measured. The results were expressed in ug at/l cc/l. As the desorption of phosphate is a function of pH, the same experiment was carried out using phosphate distilled water of different pH. The range used was 6-9, representing that of the harbor water, to yield maximum adsorption of phosphate for the harbor sediments. **Results and discussion** In the present study, the amount of phosphorus released from 1 cc sediment sample

In the present study, the amount of phosphorus released from 1 cc sediment sample after shaking for one hour at the *in situ* pH, was obtained for different stations (Table 1). It was found that the amount of phosphorus released by shaking were higher than the corresponding *in situ* concentrations in the bottom water. This is obviously due to 1) It was robuint that the automic prosphere in the bettom water. This is obviously due to the strong tendancy of phosphate to be adsorbed in sediments and suspended particles. The *in situ* concentration in near bottom sea water was proportional to the concentration of phosphorus in surficial sediments. The ratio is almost constant (Table 1). This indicates that the distribution of the water column caused by winds and/or vertical mixing processes would increase the concentration of phosphorus in the overlying water column by appreciable amounts even at the same pH. The equilibrium state between phosphorus adsorbed on the sediment particles and that dissolved in the overlying water seems to be grately governed by the oxidation (Table 1). This indicates that the distribution of the water column caused by winds and/or vertical mixing processes would increase the concentration of phosphorus in the overlying water column by appreciable amounts even at the same pH. The equilibrium state between phosphorus absorbed on the sediment particles and that dissolved in the overlying water seems to be grately governed by the oxidation state of the bottom sediments. At pH range between 6-8, a complete absorption takes place. According to El-Sayed (1977), an increase in salinity values from 27-39% had no significant effect on the desorption processes.

Table 1: Amount of phosphate released by shaking (ug aVcc. sediment), in situ concentrations in near bottom sea water, and the ratio between in situ concentration and that released from schments, as well as pH and salinity (%o) for selected stations in the harbot basin

Station	Phosphate released per liter (1)	in situ conc. in near shore bottom sca water (2)	Ratio {(2)/(1)]	рН	5%0
п	2.07	1.65	0.80	7.43	37.98
ш	3.33	2.35	0.71	7,12	36.17

Experimentally, it was found that ammonia-nitrogen concentration in bottom waters (in situ) was less than that release of ammonia-nitrogen to the overlying water. Also, the in situ concentration of ammonia near the bottom was inversely proportional to the ambient salinity (Table 2).

Table 2: Amount of ammonia released from sediments by centrifugation versus in situ ammonia concentrations (ug at/l) and salinity (%o) for selected stations in the harbor basin

Station	Ammonia released (1)	In situ conc. In pear shore bottom sea water (2)	Ratio [(2)/(1)]	рН	S%o
n	10.00	5.85	0.59	7.43	37.98
ш	15.33	9_30	0.61	7.12	36.17

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