ABSTRACT

In this work, feasibility studies of using a natural and low cost adsorbent: saw dust for the removal of Cr(VI), Ni(II), Fe(II) and Cd(II) from aqueous solution was carried out. The efficiency of the adsorbent was judged from the variation of the % adsorption with (i) contact time, (ii) adsorbent dose, (iii) initial metal ion concentration, (iv) effect of pH variation of the adsorbate solution, (v) activation of the adsorbents and (vi) particle size. Changes in the concentrations of the metallic ions after the adsorption process were monitored using Shimadzu AA650 Double-Beam Digital Atomic Absorption/Flame Spectrophotometer. Results suggest that the adsorbent employed in this work have the ability to remove the tested ions from their aqueous solutions with adsorption order onto the adsorbent varying according to Cr > Fe > Ni > Cd. Also for each ion there is an optimum pH for maximum adsorption beyond which the efficiency declined.

Keywords: Saw dust, adsorption, adsorbent, metal ion, % removal

INTRODUCTION

Pollution can simply be defined as a harmful change in the natural environment caused by human activities. This may be as a result of the release of substances, which are toxic to either animals or plants, or it may be the release of energy (heat, light, radiation, or sound), which interferes with the development of animals or plants. In general pollution is usually harmful to one or more species of animal or plant. A pollutant on the other hand is any substance produced by human activities which, at an unacceptable level, causes damage to the environment or is harmful to living systems. As such, in order to maintain a healthy living environment, a careful study on the sources and nature of pollution and the ways of pollution control should be considered (Wong et al., 2002).

Water is one of the abundantly available substances in nature and it is distributed naturally in different forms, such as rainwater, river water, spring water and mineral water. Of these categories, rainwater though associated with some dissolved gases like CO₂, SO₂, NH₃ etc. is believed to be the purest. Water is an essential ingredient for the survival of any form of animal and plant life (Sharma, 2006). On average, a human being consumes about 2 litres of water everyday. Water accounts for about 70% of the weight of a human body, while about 80% of the earth surface is covered by water. Out of the estimated 1,011 million km² of the total water present on earth, only 33,400m³ of water is available for drinking, agriculture, domestic and industrial consumption. The rest of which is locked up in oceans as salt water, polar ice-caps and glaciers and underground. Owing to increasing industrialization on one hand and exploding population on the other, the demands of water supply have been increasing tremendously. Moreover, considerable part of this limited quantity of water is polluted by sewage, industrial waste and a wide array of synthetic chemicals (Dara, 2006). Water pollutants can be divided among some general classifications into organic pollutants, inorganic pollutants, suspended solids and sediments, radioactive materials and lastly heat. Among these, inorganic pollutants are of most concern for this research.

Chromium as an environmental pollutant enters into the ecosystem by electroplating, metal finishing, chromate preparation, leather tanning etc. The two forms of chromium ions predominantly present in the environment are Cr(III) and Cr(VI), in which Cr(VI) is soluble, toxic and carcinogenic (Ackerley et al., 2004). This fact warrants its regulation in the environment.

Nickel (II) containing wastewaters are common as it is used in a number of industries including electroplating, batteries manufacturing such as Ni/Fe storage batteries, mining, metal finishing and forging. Also it is used to produce ferrous steel cutlery (Greenwood and Earnshaw, 1993). Higher concentrations of nickel cause cancer of lungs, nose and bone. Dermatitis (Ni Itch) is the most frequent effect of exposure to Ni, such as coins and jewellery. High concentration of Ni(II) in ingested water may cause severe damage to lungs, kidneys, gastrointestinal distress, e.g., nausea, vomiting, diarrhoea, pulmonary fibrosis, renal edema and skin dermatitis (Meena et al., 2005 and Erdogan et al., 2005). Nickel is genotoxic and mutagenic as well. In industrial wastewaters, nickel concentration varies between 6 and 12mg/L, which is above safe limit (Bansal and Goyal, 2005).

Cadmium is mainly used in the industry for coating steel, glass and plastics (including polyvinyl chloride), and for nickel cadmium battery production, and automotive tires. It has special importance due to its long half life and it can exert toxic effects on almost all systems of the human body (Tsalev, 1993).
The main organ for long term cadmium accumulation is the kidney. Here, the half-life period for cadmium is approximately 10 years. A life-long intake can therefore lead to a cadmium accumulation in the kidney, consequently resulting in tubulobcell necrosis (Orlovski and Piotrowski, 2003). The primary targets of the toxic effects of cadmium on liver tissue are hepatocytes and the sinusoidal endothelium (Kayama et al., 1995). At the ultrastructural level, cadmium leads to nuclear condensation and expansion in liver cells. Cadmium could also be shown to be associated with occurrences of Itai-Itai, a disease under which patients show a wide range of symptoms such as low grade of bone mineralization, high rate of fractures, increased rate of osteoporosis, and intense bone associated pain (Fridberg et al., 1974; and Nogawa et al., 2004). Cadmium is of special concern because it is non-degradable and therefore persistent; it is highly toxic to humans, plants and animals (Tsezos, 2001).

Iron, although essential to biological systems and very widely abundant in the earth crust, its overload is known as hemochromatosis and is usually caused by a gene that enhances iron absorption. Common symptoms of iron overload include enlarged liver, skin pigmentation, joint diseases, loss of body hair, anemia, and impotence. Untreated hemochromatosis aggravates the risks of diabetes, liver cancer, heart disease and arthritis.

Vast number of work has been reported on the remediation of different toxic metals from the environment. The work of Chaiyasith et al., (2006) explores the feasibility of using treated fly ash for the removal of nickel and cadmium from aqueous solutions using batch adsorption techniques. Their results suggested that the adsorption equilibrium was usually attained within 30 min at a pH of 3 – 5. The presence of other cations such as lead was found to reduce the adsorption efficiency.

In the year 2009, Bansal et al., assessed the potentials of rice husk for the removal of nickel ions from solution. Their findings showed that the efficiency of the removal depends on the physico-chemical characteristics of the adsorbent, adsorbate concentration and other studied parameters. The efficiency of adsorbent for the nickel removal was found to be 51.8% for dilute solutions at 20g/L adsorbent dose.

Studies of iron(III) adsorption from aqueous solution by olive cake as an adsorbent was carried out by Al – Anber and Al - Anber, (2008), in which the influence of the initial pH, temperature, contact time and dosage of the adsorbent on adsorption performance were investigated. Their experimental results show that the kinetic model of pseudo-second order provided a good description of the adsorption process. The biosorption of iron from aqueous solution by dried biomass of activated sludge was also studied by Shokoohi et al., (2009). Their findings revealed that the equilibrium time for iron adsorption onto the biomass was 150 min and that Elowicz kinetic models fits well with experimental data.

Gomes et al., (2001) conducted a study to evaluate the selectivity sequence and estimate the competitive adsorption of several heavy metals in seven Brazilian soils with different chemical and mineralogical characteristics. The sorption affinity of metals for the solid phase, were obtained for each soil and heavy metal cation. On the basis of these, the selectivity sequence were evaluated and the most common were Cr > Pb > Cu > Cd > Zn > Ni and Pb > Cr > Cu > Cd > Ni > Zn. Chromium, lead and copper were the cations most strongly adsorbed by the soils, whereas cadmium, nickel and zinc were the least adsorbed in the competitive situation.

The work of Rajmond, 2005 utilised a mixed bed ion exchange column for the simultaneous determination of Cr(III) and Cr(VI) using UV detector at a wave length of 365nm from environmental samples such as rainfall and galloway sediments. The performance of his method was established from the linearity, limit of detection, limit of quantification and the influence of sample pH.

MATERIALS AND METHODS

All glasswares and plastic containers were washed with detergents, rinsed with distilled water and then soaked in a 10% HNO3 solution for 24hrs. They were then washed with deionised water and dried in an oven for 24hrs at 80°C (Todorov et al., 2001). Saw dust was collected from a local sawmill and it was air dried in sunlight until almost all the moisture evaporated, washed several times with distilled water in order to remove the water soluble tannins, after which it was dried in air and then in an oven at 80°C. The adsorbent was ground to the different working sizes and was kept in a plastic container for subsequent use.

Distilled, deionised water, AnalaR grade reagents were used without further purification for the preparation of all stock solutions, and were kept in a refrigerator. 6M HNO3 was prepared from conc. HNO3 (specific gravity 1.420; percentage purity 65% w/v) and 1000mg/L Cr(VI), Ni(II), Fe(II) and Cd(II) were respectively prepared by dissolving 2.829g of K2Cr2O7, 4.0503g of NiCl2.6H2O, 4.9780g of FeSO4.7H2O and 2.1930g of CdSO4.8H2O in small volumes of distilled deionised water in separate beakers and the solutions were transferred to a 1.0litre volumetric flasks followed by the addition of 100cm3 of 6M HNO3 and they were made to mark with more water (Svehla, 2006). Lower working concentrations were prepared daily from the stock solution by appropriate dilution.

Static adsorption experiments in which the same solution remains in contact with a given quantity of adsorbent till equilibrium between the adsorbate concentration in solution and the adsorbate adsorbed per unit weight of adsorbent is reached was employed in this work. This type of equilibrium established is static in character, as it does not change further with time.

Procedures described by Bhattacharya et al., (2008) and Isa et al., (2008) for testing the effects of variations of adsorbent dose, adsorbate initial concentration, adsorbate pH, surface area and activation of the adsorbent were followed. The extent of removal of the adsorbent and the amount of various metal ions adsorbed onto the adsorbent were calculated from the relations:

\[
\frac{C_i - C_e}{C_i} \times 100 = \frac{w}{m} \quad \text{and} \quad q_e = \frac{(C_i - C_e)V}{W}
\]

where CI and CE are the concentrations (mg/L) of metal ions initially and at equilibrium time, w is the weight of the adsorbent (g), while V the volume of the solution in litre (Bhattacharya et al., 2008; El-Nemr et al., 2008; Ibrahim et al., 2006; Wang and Lin, 2008).
RESULTS AND DISCUSSION

From Fig. 1, it is apparent that by increasing the adsorbent dose the removal efficiency of each metal ion increases. This is readily understood from the fact that the number of available adsorption sites increases by increasing the adsorbent dose and therefore, results in the increase in removal efficiency or the % adsorption of the respective ion. Also, at a low adsorbent dose, e.g. 2g, the available sorption sites were quite insufficient compared with the large amount of adsorbate ions in solution, resulting in low removal efficiency. However, at a higher adsorbent dose, e.g. 8g, the sorption sites were sufficient thereby resulting in a rapid removal of the different adsorbates. The adsorption order of the adsorbates onto the saw dust was observed to follow the pattern Cr > Fe > Ni > Cd.

The strong affinity of the SD adsorbent for Cr agrees closely with that reported by Baral et al., (2006) for the removal of hexavalent chromium using alumina, goethite, alum sludge and treated saw dust derived from Shorea robusta (Sal). Also the general trend of increase of % removal with increase in adsorbent dose has been reported by various workers such as Bansal et al., (2009) for the removal of Ni using rice husk.

Initial metal ion concentration is one of the effective factors in removal efficiency. Fig. 2 depicts the effect of the increase in the initial metal ion concentration on the adsorption of the various metal ions onto SD adsorbent. The strong affinities of Cr, Fe and Ni the saw dust, give way to a marginal variation in % adsorption with increase in initial metal ion concentration, leaving Cd with a linear variation with increase in concentration.

The pH of aqueous solution and hence wastewater effluents affects a number of reactions that takes place in these solutions. For this reason, in this work the pH of the test solution was varied from 2 to 10 in order to determine the optimum pH value for the adsorption of the various metal ions onto the adsorbent. The % adsorption of the various metal ions analysed with particular reference to Ni adsorption onto saw dust adsorbent (Figs. 3) shows a gradual increase from pH 2 to 6 and then followed by a decrease. This indicates that there is an optimum pH value required for the adsorption of the metal ion beyond which a decrease will be experienced.

The observed lower % removal efficiency at lower pH may be related to the competition existing between metal ions and H+ ions for the available adsorption sites on the adsorbents, while the decrease in the % removal efficiency after the optimum values has been reached according to Ibrahim et al., (2006) may be attributed to the precipitation of the metals which occur at higher pH values.

Comparing Figs. 1 and 4, the effect of increase in surface area by using powdered adsorbent on the % adsorption can be seen. As in the case where powdered adsorbent was used, the % adsorption for the respective metals onto the adsorbent increased. The same increase (though with some initial decrease) in % removal efficiency can be observed in Figs. 2 and 5 for a situation where powdered adsorbent was used with increase in initial metal ion concentration. These observed increases are obvious as with reduction in particle size of the adsorbent, the active site available for complexation with metal ions increases thereby leading to improved % removal efficiency.

Treatment of an adsorbent with acid solution helps in the removal of any debris or soluble biomolecules that might interact with the metal ions during sorption process. This treatment, according to Abia and Asuquo (2006), is called the chemical activation of the adsorbent. Comparing the % adsorptions of the metal ions from Figs. 1 and 6, a general improvement in the % removal efficiency is observed. Of particular interest is the % removal of Cd ion where almost over 30% improvement was recorded. This generally may be explained as the activation has made large percentage of the adsorption sites available for the adsorption of the metal ions.

**Amounts Adsorbed**

Changes in the amount of metal ion adsorbed on the adsorbent with variation in different parameters studied can generally be summarised as follows;

- The metal ion uptake capacity (mg/g), except for the adsorption of cadmium on saw dust, decreases with increase in adsorbent weight (regardless of whether it is powdered or granular, activated or unactivated) for a fixed adsorbate concentration. This trend was similarly observed by Donmez et al., (1999); and Baral et al., (2006).
- For all cases where the uptake capacity was judged with variation in adsorbate concentration, it has been found that the amount adsorbed (mg/g) increases with increase in the initial metal ion concentration until an equilibrium value is reached. This is in agreement with what has been observed earlier by Ibrahim et al., (2006).

The pattern of the adsorption behavior with changes in pH show no difference between those of % adsorption for the different adsorbates.

**CONCLUSION**

The work has demonstrated the possibility of using Saw dust in the treatment of heavy metal containing effluents. Also surmised in the work is the benefit that would open-up to food agriculture and states in Nigeria with abundant wood processing industries in converting waste to wealth. Similarly, the work has attempted to make available for industries, a low cost and efficient method of effluent treatment.
Fig. 1: Variation of the % Removal with Weight of Adsorbent (Saw Dust)

Fig. 2: Variation of % Removal with Increase of Initial Metal Ion Conc. using Saw Dust Adsorbent

Fig. 3: Variation of % Removal with pH of the Metal Ion for One Hour Agitation with Saw Dust Adsorbent
Fig. 4: Variation of % Removal with Increase in Weight of Powdered Adsorbent (SD)

Fig. 5: Variation of % Removal with Increase in Initial Metal Ion Conc. Using Powdered Adsorbent (SD)

Fig. 6: Variation of % Removal with Increase in Weight of the Activated SD Adsorbent
REFERENCES


