The Potential of EDTA – Modified Rice Husk Ash as Solid Phase Extraction Resin in Seawater

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2018

Abstract

Solid Phase Extraction (SPE) is one of the methods used in extraction of elements and metals with the aid of preconcentration process. The development of sorbents and their application in preconcentration is a subject of great importance in the environment. This review summarizes the use of solid phase extraction in developing resin for the extraction of trace elements in seawater. On the right hand, Rice Husk Ash (RHA) is a cost-effective agricultural material that has the great properties and capability to be a sorbent for tracing elements. The important properties and components of the RHA will be further specify. This journal review will provide general information regarding solid phase extraction, rice husk ash (RHA), tracing elements and resin.

Keywords: solid phase extraction, trace elements, rice husk ash, resin
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Introduction

Trace metals play an important role in daily life. Thus, determination of trace elements or metals in water, soil, plant and other things is becoming increasingly important. Instrumental methods including spectrophotometry, inductively coupled plasma-atomic emission spectrometry (ICP-AES), atomic absorption spectrometry (AAS), have been used for the determination of traces of heavy metal ions. Separation and preconcentration of analyte are important aspects in the determination of trace metal ions. [1] Despite the selectivity and sensitivity of analytical techniques, there is a crucial need for the preconcentration of trace elements before their analysis due to their frequent low concentrations in numerous samples. Determination of trace metals in seawater is a hard task, not only because of the concentration of the water or detection limits of instruments that will be used but also the seawater matrix that may interfere with the determination of trace metals.

Preconcentration can overcome these problems. Preconcentration methods include ion exchange, precipitation, solvent extraction, and solid phase extraction. Above all, solid phase extraction is the most commonly used. [2]

For the extraction of trace elements, sorbent is needed. Sorbents are commercially available, but they come in an expensive price. In this study, rice husk ash was used as a cheaper and efficient sorbent for extraction of trace elements. Rice husk (RH) is available and widely produced throughout the world. These rice husks are waste product of every rice milling sites. The high silica content of RH makes it insoluble in water, having good chemical stability and structural strength. Thus, it serves an important role as a sorbent in detecting and tracing metals. Making use of them lessens the waste accumulated and contamination of the environment aside from being an efficient alternative sorbent from expensive ones. Solid Phase Extraction (SPE) is a widely used technique for fast and rapid sample preparations. Solid Phase Extraction (SPE) has many purposes, such as purification, fractionation, extraction, and adsorption. For the past few years, it has been a great interest and many journals regarding SPE have been published. [3]
Objective

The overall purpose of the journal is to provide information regarding the utilization of the rice husk (RH) into rice husk ash (RHA) and its usage in extracting trace elements. The rice husk ash is chemically modified with EDTA to improve the property of the sorbent.

1. Solid Phase Extraction

In most instances, environmental samples tend to be too complex with chromatographic systems. This fact is often overlooked in analytical development, thus causing sample preparation techniques to be a limiting factor in many analyses. In comparison with other developments, it has been suggested that sample preparation advances have come about due more to need rather than desire. Regardless of original sample concentration and matrix, the ultimate goal is a final solution with an enriched sub-fraction of analytes of interest at easily separated and detected concentrations that are compatible with the chromatographic system of choice. [4]

1.1 Advantages

1.1.1 Separation

The solid-phase extraction (SPE) approach as rapid acceptance separation procedure has advantage over classical liquid extraction technique in term of rapidity, simplicity, amenability to automation, low disposal costs and lower extraction time, and high preconcentration factor. SPE procedure characteristic performance significantly can be improved by efficient selection of a suitable sorbent. Silica gel immobilized with various organic compounds was widely used in this technique due its good mechanical and thermal stability, lower swelling susceptibility, shrinking, and microbial and radiation decay. [5]

Solid phase extraction (SPE) has several advantages over other method, such as simple and fast extractor system. Solid phase extraction procedures combine the advantages of high selectivity of solvent extraction and high efficiencies of chromatographic separation [6, 7] and they separate elements with high purity. [8]

1.1.2 Preconcentration

Solid phase extraction is an effective method of preconcentration for trace heavy metal ions. In solid phase extraction, many sorbents such as, Amberlite XAD resins [9]
and silica gel are used. [10] Diaion HP-20, fullerenes, naphthalene, activated carbon and other sorbents have also been used for separation and preconcentration of trace amounts of heavy metal ions from various matrices. [11] It is easily adaptable to the preconcentration and to the determination of trace metal ions by flow injection analysis technique. In solid phase extraction, analyte ions are adsorbed on an adsorbent, and then desorbed with a suitable eluent. Metal determinations were performed in this solution.

Solid phase extraction offers a major advantage over solvent extraction methods. [12] There are two broad types of solid phase extraction. In one, seawater is passed slowly over a resin containing immobilized chelating ligands (i.e., chelating ion exchange) while in the other hand, chelating ligands are added to the sample and the resultant metal–organic complexes sorbed to a resin. [13]

1.2 Basic Principles

Solid Phase Extraction has 4 steps:

First, the solid sorbent should be conditioned using an appropriate solvent, followed by the same solvent as the sample solvent. This step is difficult, as it enables the wetting of the packing material and the combination of the functional groups. In addition, it removes possible impurities initially contained in the sorbent or the packaging. This step removes the air in the column and fills the void volume with solvent. The nature of the conditioning solvent depends on the nature of the solid sorbent. Typically, for reversed phase sorbent (such as octadecyl-bonded silica), methanol is frequently used, followed with water or aqueous buffer whose pH and ionic strength are similar to that of the sample. Care must be taken not to allow the solid sorbent to dry between the conditioning and the sample treatment steps, otherwise the analytes will not be retained and poor recoveries will be obtained. If the sorbent dries for more than several minutes, it must be reconditioned.

The second step is the percolation of the sample through the solid sorbent. Depending on the system used, volumes can range from 1 ml to 1 l. The sample may be applied to the column by gravity, pumping, or by an automated system. The sample flow rate through the sorbent should be low enough to have efficient retention of the analytes, and high enough to avoid excessive duration. During this step, the analytes are
concentrated. Even though matrix components may also be retained by the solid sorbent, some of them pass through, thus enabling some purification of the sample.

The third step is the washing of the solid sorbent with an appropriate solvent, to have low elution strength and to eliminate matrix components that have been retained by the solid sorbent, without removing the analytes. A drying step may also be advisable, especially for aqueous matrices, to remove traces of water from the solid sorbent. This will remove the water in the final extract, which may hinder the succeeding concentration of the extract and the analysis.

The final step consists in the elution of the analytes by an appropriate solvent, without removing retained matrix components. The solvent volume should be adjusted so that required recovery of the analytes is achieved with subsequent low dilution. In addition, the flow rate should be correctly adjusted to ensure efficient elution. It is often recommended that the solvent volume be fractionated into two aliquots, and before the elution to let the solvent soak the solid sorbent. [14]

Fig. 1. Solid Phase Extraction standard operation

1.3 Adsorption/Extraction

Industrial activities are major source of water pollution due to industrial chemicals which contain heavy metals, hazardous waste which can affect not only our environment but also our health. Common methods for the removal of heavy metals comprise membrane separation, electrochemical precipitation, ion exchange, pre-concentration, fertilization and adsorption. [15] These methods differ depends on the cost, complexity and efficiency.

Among these methods, adsorption is a user-friendly technique for the removal of heavy metal. This process seems to be most versatile and effective method for removal of heavy metal. The adsorption process is being widely used by various researchers for the removal of heavy metals from waste streams.
and activated carbon has been frequently used as an adsorbent. [16]

Active carbon adsorption was considered to be common and effective, but the cost is high [17]. For this reason, many studies have been conducted in order to find out effective and low-cost adsorbents.

2. Sorbents

The development of sorbents is a topic of great importance for the determination of trace elements. Sorbent is used to effectively collect residues in various kinds of surfaces with the use of different methods. Sorbents such as coconut coir [18], coconut shell [19], sawdust, [20] corn cob [21] are effective sorbents for trace and heavy metals.

2.1 Rice Husk

Rice Husk (RH) is widely available throughout the world. Every year, approximately 500 million tons of rice is produced by world and 2-3 million tons are produced in the Philippines. Rice husk ash (RHA) is a by-product of RH. This is a versatile material with good adsorption properties.

Rice husk was chosen as the local low-cost adsorbent since million tons of rice husks are produced annually as a result of paddy cultivation. Various modifications on rice husk have been published in order to enhance their sorption capacities for metal ions, dye and other pollutants. Most of the studies indicated that the modified adsorbents are efficient in binding either the cationic or anionic species but not both. In continued effort to use low-cost materials for the removal of organic pollutants, the performance of ethylenediaminetetraacetic acid (EDTA) modified rice husk as a sorbent have been investigated throughout the years. [22]

2.1.1 Composition of rice husk

RH contents are hemicelluloses 24.3%, cellulose 34.4%, lignin 19.2%, ash 18.85%, and the other trace elements 3.25%. Hemicelluloses used as recourse of activated carbon, xylose and silicon dioxide. RH contains elemental components as Carbon 37.05%, Hydrogen 8.80%, Nitrogen 11.06%, Silicon 9.01% and Oxygen 35.03 %. Husk contains 17-25% silica.[23] Rice husks have bulk density of 96-100 kg/m³, hardness (Mohr’s scale) 5-6, ash 22.29%, Oxygen 31-37%, Nitrogen 0.23-0.32%, Sulfur 0.04-
0.08%. Hydrogen 4-5%). The composition of RH depends on many factors such as rice variety, type of fertilizer used, soil chemistry, and even the geographic localization of the production. [24]

2.1.2 Composition of rice husk ash

RH is high in ash content as compared to other biomass fuels ranging 14-25%. [25] Silica content in RHA ranges from 83-98%. [26] Presence of high amount of silica makes it a valuable material for use in industrial applications. Chemical composition of RHA determined by x-ray fluorescence (XRF) as SiO$_2$ 89%, Al$_2$O$_3$ 1.20%, C 18.24%, CaO 1%, K$_2$O 1.22%, Fe$_2$O$_3$ 1.28%. The various factors which affects the ash properties are incinerating conditions (temperature and time), rate of heating, burning technique. [27]

2.1.3 Preparation of activated carbon

Various properties such as high surface area, large adsorption capacity and fast adsorption kinetics makes activated carbons as valuable material for different environmental applications. Production of activated carbon from rice husk is achieved through activation with chemical or physical means. The nanoporous activated carbon with large specific area of 2523.4 m$^2$ g$^{-1}$ was obtained from RH. [28] The effect of alkali carbonate in the activation process can be related to formation of porosity due to SiO$_2$ interaction with the carbonates, CO$_2$ reacts with the carbon phase with formation of CO, according following reaction:

$$M_2CO_3 + SiO_2 = M_2SiO_3 + CO_2$$

$$CO_2 + C = 2CO$$

In the study in which the disposal of dyes laden wastewater poses one of the industry’s major problems because such effluents contain a number of contaminants including acid, dissolved and suspended solids, toxic compound sand coloring pigments. Color is the first to be recognized because it is visibility. Safranin-T is one of the most commonly used azine dye, which are amongst the oldest known synthetic dyes. Safranin-T was considered as the model compound to represent the dyes that are released in effluents from the textile and food industries. Adsorption organic pollutants onto solid/water interfaces have been found to be an efficient in controlling the extent of water pollution. The efficiency of removal of
Safranin-T using activated rice husks and activated carbon is carried out. [29]

2.1.4 Water Purification

The high silica content of RH makes it insoluble in water, having good chemical stability and structural strength. Thus, it serves an important role in water purification as well as sorbent in tracing metals. Sorbent made by RH effectively used to removal of the six heavy metals such as Fe, Mn, Zn, Cu, Cd and Pb. [30] RH is also used as adsorbent to remove various pollutants, phenols, dyes, pesticides, inorganic anions, organic compounds and heavy metals. [31]

2.2 Rice Husk Ash

In particular, fly ash is generated by the air pollution control systems are hazardous waste materials for which landfilling is still considered the most appropriate management strategy. Indeed, this ash contains leachable toxic metals, as for example Pb and Zn. So, the use of silica extracted from rice husk ash (RHA) for heavy metals stabilization of fly ash. It is attributed to two different reactions, due to the amorphous silica and carbonation. Since the organic part of rice husk (RH) can easily be removed by thermal decomposition, thermal treatment of RH is an approach to obtain, not only energy, but also silica from rice husk. [32] Different optimization approaches have been used to improve the adsorption capacity of the mesoporous silica.

Rice husk removal during rice refining, creates disposal problem due to less commercial interest. Also, handling and transportation of RH is problematic due to its low density. RHA is a great environment threat causing damage to land and surrounding area where it is dumped. Therefore, commercial use of rice husk and its ash is an alternative solution to the disposal problem. RHA has been found to be effective as an oil spill absorbent, and for use in waterproofing chemicals, flame retardants, and as a carrier for pesticides and insecticides. Presently, RH usually ends up being burnt in open space, thus causing environmental pollution. To conserve energy and resources, efforts have been made to burn the husk under controlled conditions and to utilize the resultant ash in building, semiconductor, composite, and abrasive materials. Also, ash is an active catalyst and a good material for catalyst support because of its high surface area. [33]

3. Trace Elements
Generally, toxic heavy metals have been reported to cause potential human health risk and ecological disturbances. Lots of studies have been performed for the determination of trace metal ions in various media including some body tissues and fluids, natural waters etc. The determination of heavy trace metals is important for human health. [34] Most heavy metal ions such as Ni^{2+}, Zn^{2+}, Pb^{2+}, Cu^{2+} and Co^{2+} are toxic and can exist for a long time in the environment. These metals cause serious environmental and public health problems even at very low concentrations.

Currently, research efforts have shifted towards the development of new, low cost adsorbents preferably from biogenic sources. An adsorbent may be regarded as low cost if it is abundant in nature, is a by-product, a waste material from industry or requires little processing. [35]

To curtail heavy metal pollution problems, some various techniques for wastewater treatment have been developed through decades. This includes coagulation, chemical precipitation, membrane separation, reverse osmosis, solvent extraction, ion exchange, filtration, and adsorption by activated carbon. Among these methods, adsorption by activated carbon is widely used as an effective adsorbent in many applications but metal ions removal by activated carbon is relatively expensive. Hence there is a need to search for the lowcost adsorbent. Rice husk an abundant agricultural waste material generated in rice producing countries is found to be capable of removing heavy metals. [36] It can be considered as an efficient and low-cost adsorbent for heavy metals and the potential of adsorbents to treat the wastewater in heavy metal ions. [37]

3.1 Seawater

Trace metals in seawater are widely studied to understand their geochemical and biological cycles in marine environment and for environmental monitoring.

Nowadays, seawater is continuous gaining of attention. The Philippine marine environment, located in the Asia-Pacific region, is no less than vulnerable to radiological threats. [38] It is because of accidents that are related to marine environment that keeps happening. Recently, studies for determination of elements in water samples have increased greatly. Due to low content of elements in water samples and complexity of matrix [39, 40] the most common sensitive analytical techniques used,
are inductively coupled plasma mass spectrometry (ICP-MS) and inductively coupled plasma optical emission spectrometry (ICP OES). [41]

Trace metal effects on marine primary production has driven a greater demand for accurate, precise metal determinations at the very low concentrations found in seawater. Bioactive metal cations like, Fe, Co, Ni, Cu, Zn and Cd, given the growing use of inductively coupled plasma mass spectrometry (ICP-MS) as a rapid multi-element instrumental detection system. The new extraction method must be applied easily to a range of sample volumes yet also be clean and precise enough to permit accurate trace metal determinations at picomolar levels in seawater. [42]

With the development of more sensitive analytical techniques, marine chemists are capable of detection and estimation of trace metals in seawater even at µg/l to ng/l level. Several methods are adopted to preconcentrate the trace metals and to separate the salt matrix from seawater prior to trace metal determination. With the growing use of ICP-MS as a rapid multi-element instrumentation detection system, a wide range of trace metals were determined in seawater by ICP-MS technique for preconcentration and matrix removal. [43]

3.1.1 Preconcentration of Seawater

![Figure 2: Sohrin's Diagram of Manual Preconcentration system](image)

The preconcentration and separation methods based on the sorption are considered to be superior to the liquid-liquid extraction as regards simplicity, rapidity and ability to attain a high preconcentration factor. Preconcentration methods based on adsorption of complexed metals on columns filled with activated carbon have found widespread application for this purpose. [44] Activated carbon was used for the purification of drinking water for the adsorbing of aromatic and other organic materials which cause bed smelling and taste in water for many years.

However, separation and preconcentration procedures have said to be
applied for ultratrace amounts determination in waters. Methods based on solid phase extraction (SPE) are widely used for this purpose. The efficiency of SPE procedure and such SPE characteristics as analysis time), preconcentration factor, selectivity, regeneration ability, chemical and mechanical stability depend mainly on the nature and properties of the sorbent material. Sorbents used for this purpose are chemically modified silica gel, activated carbon, inorganic–organic hybrid materials as C 18–silica (8, 9) and synthetic chelating polymers resins. [45]Preconcentration procedures, especially solid phase extraction (SPE), have been indispensable tools to analyte enrichment to bring the elemental concentrations in solution to the detection range of the ICP-OES instrument. [46]

4. **Resin**

The most common packing used today in Solid Phase Extraction (SPE) and HighPerformance Liquid Chromatography (HPLC) is silica gel. The gel is a network of silica atoms bound to one another by siloxane bonds. This gel is then modified with any number of chemical entities such as C 8 or C 18 reacted at terminal hydroxyl groups to facilitate reverse-phase separations. [47]

5. **EDTA**

EDTA is a remarkable reagent not only because it forms chelates with all cations but also because most of these chelates are sufficiently stable for titrations. EDTA can form a highly soluble complex with a wide variety of metal ions in an aqueous solution and can greatly reduce the efficiency of metal removal by conventional chemical precipitation, such as hydroxide and sulfide precipitations. [48]

One of the common structures for EDTA complexes is shown in Fig 2.

![Figure 3: EDTA Structure](image-url)
EDTA forms very stable complexes with metal ions and prevents them from catalyzing air oxidation reactions that can lead to decomposition of proteins and other compounds. [49]

5.1 EDTA – Modified

Various materials from biomass to inorganic oxides have been functionalized with EDTA. The usage of EDTA arises from its strong metal chelating ability, local availability, and economically-friendly price. Silica gel is an amorphous form of silicon dioxide where silicon atoms are linked together via oxygen atoms with siloxane bonds. Silica gel surface contains hydroxyl groups that can be further functionalized using a familiar silanization procedure. [50]

Different modifications on rice husk have been reported in order to enhance their sorption capacities for metal ions, dye and other pollutants. [51] Most of the studies indicated that the modified adsorbents are efficient in binding either the cationic or anionic species but not both. Therefore, there is a need to have adsorbents capable of removing different types of dyes either singly or simultaneously. Many studies have investigated the performance of ethylenediaminetetraacetic acid (EDTA) modified rice husk as a sorbent for basic and reactive dyes and other metals.

Some trace metals have been determined by ICP-MS just after dilution of seawater. Although this technique is simple, the sensitivity is not adequate to determine the concentrations in the open ocean and the signals usually suffer from drift and suppression due to the seawater matrix. Thus, preconcentration from major constituents is crucial to accurate and precise determination of trace metals. EDTA can form a maximum of five coordinate bonds to a metal ion. This resin has excellent affinity to trace metal ions. This resin is usually used in a closed column to preconcentrate trace metals, which reduces the risk of contamination. The trace metals in the eluate were determined by ICP-MS. This method was successfully applied to determine trace metals in seawater. [52]

Methodology

1. Seawater Samples

Polyethylene bottles that will contain surface seawater samples will pre-wash with seawater itself from insert seawater sample location. The bottles will be pre-cleaned with
detergent, followed by deionized (Milli Q) water, dilute nitric acid, and lastly, with double distill deionized water.

2. Procedure for the Preconcentration of Seawater

Seawater samples will be acidified first with nitric acids at <2.0 pH in order to prevent metal adsorption onto the inner bottles' walls. These seawater samples will then transferred into a pre-cleaned polyethylene bottles by first filtering it through 0.45 um polycarbonate membrane Nucleopore filter (Millipore) [53]. The samples will be analyzed within 2 weeks after the collection, following the preconcentration procedure that is given above. For the analysis, the sample containing 600ml of the acidified sea water will be neutralized first and then buffer to the desired pH. [54]

3. Rice Husk Samples

The rice husks from C4 Navotas will be modified. These rice husk samples will be authenticated at Bureau of Plant Industry, Pandacan, Manila.

4. EDTA Modification of Rice Husk Ash

The rice husk ash will wash thoroughly with water then dry it in an oven at 105 °C for 12 hours. [55] 8g of grinded rice husk with 0.5g of (Ethylenediaminetetraacetic acid) EDTA and soaking it in 300 mL of 1.0 M Sodium hydroxide (NaOH) for 3 hours at 70 °C, the optimum conditions for the modification of rice husk according to the study. The effect of functional groups on the surface of rice husk will be studied and investigated. In order to do that, the rice husk will treat first with NaOH, as stated on the reported method. [56] This sample of treated rice husk will filter, wash with excess water, dry in an oven at 60 °C, and label as ERH.

5. Characterization of RHA
   a. X-Ray Diffraction (XRD)

   X – Ray Diffraction analysis is use to determine the particle size and shape of the rice husk ash.

   b. Scanning Electron Microscopy

   Visual confirmation of surface morphology of the rice husk ash is analyze
and determine by the use of scanning electron microscopy.

6. **Preconcentration System**

The EDTA-modified rice husk ash will be contained and stored in a propylene syringe. It will initially clean using methyl alcohol, acetone, 3 M HNO₃, and 1 M HCl-1x10⁻² M ascorbic acid by allowing the reagents to drip with gravity through the syringe. After that, the resin will take out of syringe and suspend in 1 M HCl in a Teflon perfluoroalkoxy (PFA) beaker for 24 hours in order to remove small resin particles. After decantating, the resin will packed in the column of the PFA tube.

The inside diameter of the preconcentration system's columns are 6 mm and has a bed height 30 mm, containing 500 mg of the resin sandwich between ethylene-tetrafluoroethylene meshes. Another set up of this will prepared again and use in order to purify the buffer solution.

The column will mount on either manual or automated preconcentration system. The system will compose of chelating resin columns, low-density polyethylene (LDPE) bottles, PFA tubes, poly(tetrafluoroethylene) (PTFE) joints, PTFE three-way valves, peristaltic pump, and a nitrogen gas line. The system will place in a clean box, which will construct with plastic props and polyethylene sheets. All the plastic materials are clean in the same way as the bottles prior to construction. The PTFE three-way valves switch the flow line. The peristaltic pump sends sample and buffer solutions to the column. The eluent will send by nitrogen gas pressure to avoid contamination from the tubings. The nitrogen gas will introduce into the eluent bottle after passing through deionized water (MQW) trap and a cartridge filter. The system use three parallel lines and which can process three samples simultaneously. After construction, the chelating resin column will clean by passing 1 M HNO₃ as an eluent. The whole solution line will fill with 1M HNO₃ and leave to stand for 24 hours. Lastly, it will clean with ultrapure MQW water.

The automated system has a similar flow line design from the manual system. PTFE solenoid three way valves are use to switch the flow line and two peristaltic pumps are use to send all solutions. The valves and pumps are control using a programmable controller. The solution line is pre-clean in a similar manner to that of the manual system. In this system, the tube is require to be clean with eluent before
proceeding to the elution step of every sample.

A stock solution of concentrated HAcO-NH4AcO buffer with 3.6 M and pH 6.0 will prepare by mixing 100 g of HAcO and 44.3 g of NH3. A working solution 0.05 M HAcO-NH4AcO will prepare by diluting the concentrated buffer. It will purify by passing it twice through a EDTA-modified solid phase extraction resin column for purification. As stated, the preconcentration column will clean using 15 mL of the eluent. After that, the column will condition by passing 30 mL MQW and 40 mL of working solution of GAcO-NH4AcO through it, prior to sample being loaded. After loading the seawater samples in the column for ~40 minutes, 40 mL of 0.05 M HAcO-NH4AcO are use to wash away the remaining sea salts that accumulate in the column.

The analytes are then elute with 15 mL of 1 M HNO3. LPDE bottle is use to store the target metals until determination by ICP-MS. The flow rate is 3 mL/min for both sample and buffer solution, while eluent's is 1 mL/min in the opposite direction to the sample loading. The exact volume of the loaded sample and eluate is determined by weighing for the flow rates vary with regards to the condition of the column. It find out that one cycle of preconcentration usually took 2 hours to be accomplished.

7. Instrument for Tracing Elements

Inductively coupled plasma-mass spectrometry (ICP-MS) offers exceptional sensitivity and multi-element capability for trace metal analysis. [42] An ICP MS instrument is use for analysis of the eluate.

Results

I. Parameters in Solid Phase Extraction

1. pH [12, 52]

For Lutfi’s experiment, they will adjust the sample solution and column with HNO3 or CH3COOH-NH3 buffer solution. A 25 ml of seawater sample is spike with the analyte and pass through the TSK-8HQ column. Adsorption of metals onto the TSK-8HQ resin is dependent on both in metals and in 8-hydroxyquinoline group which changes depending on the pH.

Sohrin’s experiment on the other hand, the pH of each sample solution will adjust with HNO3 to pH 1-3, 0.05 M HAcO-NH4AcO to pH4-5, 0.05M2-(N-morpholino)ethanesulfonic acid (MES, Dojindo) and NaOH to pH 6, 0.05 M N-2-hydroxyethylpiperazine-N′-2-
ethanesulfonic acid and NaOH to pH 7, or 0.05M-N-tris(hydroxymethyl)methyl-3-aminopropanesulfonic acid and NaOH to pH 8.5.

In Lutfi’s experiment, all of the metals are quantitatively collected with the TSK-8HQ column between pH 5 and pH 6. Therefore, further experiments are carried out at pH 5.2 ± 0.2. While on Sohrin experiment, all metals in the graph have increased their % collection by pH 2 and fully at pH 4. In their experiment, alkali is involved, namely, Al, Mn, Fe, Co, Ni, Cu, Zn, Cd, and Pb, are quantitatively collected at pH 6 and 7. On the other hand, the collection percentage of alkaline earth metals increase above pH 7. Thus, the optimum sample pH is at 6.0 and it goes down at pH 9.

These experiments show that the higher the pH that will be used, the higher the
recovery or percent collection of the trace metals, using the same method.

2. Flow Rate [57,58]

In Hossein’s experiment, 50 mL portions of sample solution are pass through the column at different flow rates.

Huseyin discuss that the retention time of each metal is examine under optimum conditions by passing 100 ml of metal solution though a column with a peristaltic pump.

Therefore, a flow rate of 2mL min-1 is choose to compromise between the analytical time and the quantitative adsorption in Hossein’s experiment while in Huseyin’s experiment, the flow rates of the solutions are adjust from time to time. In this experiment, the optimum flow rate is 3 mL min-1 for copper and nickel.

As we can see, the percent retention or collection decrease with increase in flow rate, which is due to the shorter contact time between the adsorbent and the metal ions and subsequently the decrease in complex formation.
3. Eluent

The choice of a suitable eluent is another important task in the recovery of analytes. As can be seen from Figure I.3, the extraction of these metal ions are negligible at pH < 3, in the presence of dilute HCl and HNO₃. The elution of the adsorbed metal-chelates on the solid phases are important step for solid phase extraction studies.

Due to this point, various eluent solutions, given in Table 1, are used for desorption of ions complex within diane-1,2,3-trione 1,2-dioxime chelates from SDS coated alumina. Quantitative recoveries for copper, nickel, and cobalt ions are obtained only with 1 mol L⁻¹ HNO₃. Recoveries are not quantitative for other eluents used in the present study. For this reason, HNO₃ is the best eluent and choose as eluent and various concentrations and volumes of HNO₃ are attempted to elute the analytes retained in the modified column.

Table I.3.1

<table>
<thead>
<tr>
<th>Method</th>
<th>Chelating agent</th>
<th>Analytes of interest</th>
<th>Precision (%) Recovery</th>
<th>Detection Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid phase extraction</td>
<td>Chel</td>
<td>ex-100</td>
<td>Zr, Hf</td>
<td>2.5-7.2, 8.9-22</td>
</tr>
<tr>
<td>isotope dilution-ICP-MS</td>
<td>TSK-8HQ</td>
<td>Zr, Hf, Nb, Ta, W</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table I.4.1 Comparison between previous works

Detection on limit
A Journal Review: Extraction of Trace Elements in Seawater using EDTA – Modified Rice Husk Ash Solid Phase Extraction Resin

(A) The detection limit of Zr, based on three standard deviations of the blank, is 0.21 pmol/kg and Hf is 0.03 pmol/kg while for (B) the ICP-MS instruments detection limit (IDL) for each analyte will calculate from three times the standard deviation for ten determinations of 0.5 M HNO₃ with the resulting detection limit of 0.009–0.15 pmol kg⁻¹.

Recovery

<table>
<thead>
<tr>
<th>Element</th>
<th>Recovery (%)</th>
<th>(\text{Cu} )</th>
<th>(\text{Ni} )</th>
<th>(\text{Co} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.5 mL HNO₃ (0.5 mol L⁻¹)</td>
<td>94.7 ± 2</td>
<td>92.7 ± 4</td>
<td>94.2 ± 4</td>
<td></td>
</tr>
<tr>
<td>4.5 mL HCl (0.5 mol L⁻¹)</td>
<td>82.3 ± 4</td>
<td>81.7 ± 5</td>
<td>85.8 ± 5</td>
<td></td>
</tr>
<tr>
<td>4.5 mL H₂SO₄ (0.5 mol L⁻¹)</td>
<td>42.3 ± 2</td>
<td>45.5 ± 2</td>
<td>50.7 ± 4</td>
<td></td>
</tr>
<tr>
<td>4.5 mL CH₃COOH (0.5 mol L⁻¹)</td>
<td>24.6 ± 4</td>
<td>41.4 ± 3</td>
<td>46.5 ± 2</td>
<td></td>
</tr>
<tr>
<td>4.5 mL Na₂SO₄ (0.5 mol L⁻¹)</td>
<td>29.1 ± 5</td>
<td>30.8 ± 2</td>
<td>32.2 ± 2</td>
<td></td>
</tr>
<tr>
<td>4.5 mL EDTA (0.5 mol L⁻¹)</td>
<td>46.1 ± 3</td>
<td>57.1 ± 5</td>
<td>48.2 ± 3</td>
<td></td>
</tr>
<tr>
<td>4.5 mL HNO₃ (1 mol L⁻¹)</td>
<td>94.5 ± 4</td>
<td>96.5 ± 2</td>
<td>94.8 ± 4</td>
<td></td>
</tr>
<tr>
<td>4.5 mL HNO₃ (2 mol L⁻¹)</td>
<td>100.5 ± 4</td>
<td>98.0 ± 2</td>
<td>102 ± 5</td>
<td></td>
</tr>
<tr>
<td>6 mL HNO₃ (1 mol L⁻¹)</td>
<td>90.0 ± 2</td>
<td>101.0 ± 5</td>
<td>101.8 ± 4</td>
<td></td>
</tr>
<tr>
<td>6 mL HNO₃ (2 mol L⁻¹)</td>
<td>100 ± 4</td>
<td>101.0 ± 5</td>
<td>102.8 ± 4</td>
<td></td>
</tr>
<tr>
<td>4.5 mL HCl (1 mol L⁻¹)</td>
<td>84.4 ± 4</td>
<td>80.7 ± 4</td>
<td>85.0 ± 3</td>
<td></td>
</tr>
<tr>
<td>4.5 mL HCl (2 mol L⁻¹)</td>
<td>90.1 ± 5</td>
<td>85.4 ± 2</td>
<td>84.0 ± 2</td>
<td></td>
</tr>
<tr>
<td>6 mL HCl (1 mol L⁻¹)</td>
<td>90.5 ± 3</td>
<td>89.2 ± 2</td>
<td>86.4 ± 3</td>
<td></td>
</tr>
<tr>
<td>6 mL HCl (2 mol L⁻¹)</td>
<td>80.7 ± 2</td>
<td>78.7 ± 3</td>
<td>71.0 ± 5</td>
<td></td>
</tr>
</tbody>
</table>

Recovery

II. Characterization of Rice Husk Ash as sorbent [60,61]

1. Efficiency

<table>
<thead>
<tr>
<th>Adsorbent</th>
<th>mg Bi/g adsorbent</th>
<th>mg Cr/g adsorbent</th>
</tr>
</thead>
<tbody>
<tr>
<td>URH (untreated rice with husk)</td>
<td>0.47</td>
<td>0.12</td>
</tr>
<tr>
<td>RH (rice husk)</td>
<td>2.94</td>
<td>0.50</td>
</tr>
<tr>
<td>RHA-300 (rice husk treated at 300°C)</td>
<td>18.34</td>
<td>19.09</td>
</tr>
<tr>
<td>RHA-600 (rice husk treated at 600°C)</td>
<td>4.90</td>
<td>6.49</td>
</tr>
</tbody>
</table>

Table II.1 Comparison of adsorption capacity of URH, RH, RHA-300 and RHA-600

The loading capacity of the URH, RH, RHA-300 and RHA-600 will perform by the batch method as describe in experimental section. The results show that RHA-300 has the highest capacity of all. The maximum amount of bismuth and chromium sorb on the RHA300 is higher than many collectors, which are successfully use for the preconcentration of trace elements in water samples. This means that the loading capacity of RHA-300 is large enough to collect heavy metals from a large volume of water samples. Therefore, all the column experiments will use RHA-300.
2. **Particle Size**

![Figure II.2 Comparison of particle size to % retention](image)

Adsorption studies for different particle sizes of RHA-300 and RHA-600 (diameters of 38–53, 53–106, 106–150, 150–300, 300–600 and 600–850mm) and Bi and Cr concentration of 5.0mg/L reveal that adsorption decrease from 99–98% to 91–84% with increasing particle size. Higher adsorption obtain for smaller particle size can be attribute to larger surface area of smaller particles per unit masses of the both RHAs.

The effect of particle size on the retention of Bi and Cr on RHA-300 and RHA-600 (RHA-300 and RHA-600: 0.5g; pH=6; shaking time: 30min; sample volume: 25mL).

3. **Temperature**

![Figure II.2 Comparison of pH to % retention](image)

The properties of silica in RHA are strongly affect from thermal conditions. At elevated temperatures silica concentration in Rice Husk Ash increases and the colors of the ash samples are black-brown and nearly white at 300 and 600 deg C, respectively. When RHA will heat, carbon will remove and the percentage of carbon reduce gradually which cause the silicon dioxide to be increased relatively. On the other hand, the pH of the solution containing Rice Husk Ash increase at elevated pretreatment temperatures.
The effect of pH on the retention of bismuth (Untreated Rice Husk: 5.0g; Rice Husk: 1.0g; RHA heated at 300 deg C (RHA-300) and RHA heated at 600 deg C (RHA-600): 0.5g; sample volume: 25mL; shaking time: 48hours for URH and RH, 30min for RHA-300 and RHA-600).

Discussion

(Lutfi, 2007)

The researchers compared their methods from the existing methods that are widely use determination of trace elements namely Zr, Hf, Nb, Ta, and W with the latter having highly accessible studies. Data shows that the perform method is more precise and gives lower procedural blank. And finally, it covers lower detection limit, giving it an advantage to outperform existing or other widely use methods for determination of the said trace metals.

The method is use for determining Zr, Hf, Nb, Ta, and W's concentration (in molality) on NASS-5, a seawater reference material that is collect in the North Atlantic at a known depth and accurate location. The method provided information value that can be use for future studies that several laboratories and analysts can use to develop and refine method to determine the mentioned trace metals in seawater.

(Sohrin, 2008)

The researchers found that the methodology used has many advantages compared to other methods that were used in determining the specified trace elements. High selectivity, 100 4% recovery rate of trace metals, lower detection limit of the method, and its ability to remove almost 100% of alkali and alkaline earth metals are just part of those that justifies its efficiency.

The method proposed and used was then tested to NASS-5 and CASS-4, seawater reference materials, in order to validate its precision and accuracy. Data gathered has shown that the method and technique provided excellent agreement and possess similarities with the certified values of the seawater reference materials. The results given by the analysis seems to give oceanographically consistent data, thus making it viable for an actual trace metal determination method for seawater samples from other sources.

Conclusion

GSJ: Volume 7, Issue 1, January 2019
ISSN 2320-9186
The chelating resin with ethylenediaminetriacetic acid functional groups, like in Sohrin’s experiment has excellent selectivity for trace metals. They applied this resin to preconcentration of Al, Mn, Fe, Co, Ni, Cu, Zn, Cd, and Pb in seawater for determination by using ICP-MS. Through the construction of closed preconcentration systems and optimization of the procedure, such advantages were realized: the recovery of trace metals was 100 ± 4% and >99.9% of alkali and alkaline earth metals were removed. However, the concentrations for Fe, Zn, and Cd in surface water in the open ocean would be lower than the overall detection limits. Moreover, the preconcentration was said to be performed in a non-clean room laboratory and took only 2 hours for one cycle. This method will be very useful to elucidate the stoichiometry and systematic behavior of trace metals in the oceans. Also, certain resins gave satisfactory performance in tracing elements for other studies.

The researcher’s sorbent, rice husk, gave an overwhelming info in extracting trace metals. Its components that were discussed are truly a help in our environmental problems. Certain parameters such as pH, temperature, particle size, etc. are important in conducting experiments. That’s why many studies are conducted to prove such parameters.

References


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[33] Ajward, N., Rajendra, J., & Perera, V., Elemental Analysis of Rice Husk Ash obtained from different varieties of rice grown in Polonnaruwa District.


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