INTRODUCTION

The toxic effects of mercury are well known. Mercury may enter a human body by inhalation of mercury vapor [mainly in the form of Hg(0)], drinking water (mainly as inorganic mercury, Hg^{2+}), and/or by the consumption of fish and fish products (mainly as methyl-mercury, CH_{3}Hg^{2+} in the diet).^{1}

Mercury and its compounds even at low concentrations are known to be a very dangerous neurotoxin for living organisms. Cold vapor atomic absorption spectrometry (CVAAS) with the typical detection limit of 0.05 mg L^{-1} level due to its advantages including simplicity, high sensitivity, and freedom from interference, is one of the most selective, sensitive, and popular techniques for mercury(II) determination in a wide variety of samples. The determination of traces of mercury ion at the sub ng mL^{-1} level even using sensitive CVAAS usually requires a prior separation and/or preconcentration step in order to improve method sensitivity with separation of target compounds as well as minimization of the matrix effects. Selective separation of a given metal ion from a complex mixture of other ionic species is of increasing importance in industrial and analytical chemistry and separation sciences. In most cases, before detection, various interfering species must be removed and/or the species of interest must be enriched.

Numerous extraction and separation methods for mercury determination have been developed. Liquid membrane transport mediated with a mobile carrier has been proposed as a promising technology for separation and purification of various metal ions. Liquid membranes (liquid phases), existing in either supported or unsupported form, serve as selective barriers between liquid or gas phases and have shown great potential and application in separations.

Selective transport of cationic substrates by membrane carriers is of great importance in chemistry, biology, and separation sciences. Compared with conventional separation processes such as liquid–liquid extraction, membrane techniques are characterized by the technical simplicity and high efficiency in separating or enriching material from gaseous or liquid mixtures. Also, these methods reduce the solvent inventory requirements and also allows the use of expensive and highly selective extractant which otherwise would be uneconomic in solvent extractions.

Bulk liquid membrane (BLM) is one of the simple, lowest and efficient types of liquid membranes. In this technique similar to liquid membrane configurations, viz ion transport across membranes combine the extraction, diffusion, and back extraction of analytes are particularly drawing maximum attention. BLM constitute the cheapest separation techniques because of their relatively small inventory and low capital cost.

In a BLM, a relatively thick layer of immiscible fluid is used to separate the source and receiving phase. There is no means of support for the membrane phase and it is kept apart from the external phases only by means of its immiscibility. A recent development in liquid membranes is the incorporation of selective carriers within the liquid membrane phase which selectivity and efficiency via chemical reaction facilitate the transport of a specific compound across the membrane.

Carrier mediated transport through liquid membrane is well known as one of the most powerful tools for such concentration, separation and recovery of target compound. The selective transport of metal ions across a membrane is known to play an essential role in many biological processes. There has been a growing interest in the transport of metal ions mediated by receptor molecules where the carrier operates selective across artificial or biological membranes.

In this regard, there has been considerable interest in the use of proton ionizable carriers for the transport of metal cations between two aqueous solutions through an organic membrane. These pH-regulated ligands described until now present the fact that transfer of the counter-anion from the aqueous phase into the organic medium is avoided. Such chelating agents do not involve concomitant transfer of one or more aqueous phase anions into the organic medium. This factor was of immense importance to potential applications in which hard aqueous phase anions such as chloride, nitrate and sulfate would be involved.

Transport of mercury through liquid membranes using various carriers such as polynatriadiene, triisobutylphosphite (TBP), \(^{10}\) 18-crown-6 and 21-crown-7, bis (di (2-ethylhexyloxy) thiophosphoryl) disulfide \(^{11}\) have been carried out. The present paper describes the use of isopropyl 2-[(isoproxycarbothioyl)disulfanyl]ethane thioate (IIDE) as a selective neutral carrier for Hg^{2+} ion transport through...
a bulk liquid membrane. For this purpose the influence of effective parameters such as stirring rate, concentration of HCl in the source phase, concentration of NaSCN in the receiving phase, concentration of IIDE in the membrane and contact time was optimized.

EXPERIMENTAL

Reagents

A stock solution of Hg (II) was prepared by dissolving an appropriate and accurately amount of mercuric chloride of analytical regent grade in 1% (v/v) HCl solution. The working standard solution was prepared by further diluting the stock solution with doubly distilled deionized water. SnCl₂ and Nitrate salts of all other cations used were of the analytical grade and used without further purification. Doubly distilled deionized water was used throughout. Extra pure chloroform purchased from Merck was used as organic phase. (Merck E. Dermstant Germany)

Synthesis of IIDE

Iodine (1 mmol) in CH₂Cl₂ (10 mL) was added to a stirred solution of potassium o-isopropyl(dithiocarbamate (1 mmol) in CH₂Cl₂ (10 mL) and stirred for 1 h. The reaction mixture was washed with 10% (w/v) aqueous Na₂SO₄ (1×10 mL) and H₂O (2×10 mL). The organic layer was dried over MgSO₄ and evaporated under reduced pressure. More purification was achieved by re-crystallization in hexane, so that, pale yellow crystals of IIDE were obtained in 90% yield (0.24 g). The structure and purity of IIDE was confirmed by elemental analysis, NMR and IR spectroscopy. 1H NMR (CDCl₃). δ (ppm): 1.43 (t, 12H, CH₃), 5.63 (m, 2H), 7.94 × 10⁻⁵ (s), 690.5 (m). The structure of IIDE is presented in Scheme 1.

Scheme 1. Structure of IIDE

Apparatus

The determination of mercury content was carried out with a Shimadzu AA-670 atomic absorption spectrometer equipped with an Hg-hollow cathode lamp (HCL) and an on-line cold vapor generation system using SnCl₂. The wavelength was set at 253.7 nm (resonance line) with the spectral bandwidth of 0.5 nm. A long path quartz cell (2 cm i.d., 10 cm long) connected to the spectrometer was used as a detection system. The determination of all other cations was carried out with a Shimadzu AA-670 atomic absorption spectrometer under recommended conditions for each metal ion. A digital pH meter, Metrohm model 632 equipped with a combined glass calomel electrode was used for the pH adjustments. A bulk type liquid membrane cell (Figure 1) was used in this study. All conductance measurements were carried out with a Metrohm 712 conductivity meter with a dip-type conductivity cell made of platinum black.

Procedure

All transport experiments were carried out at ambient temperature. A cylindrical glass cell (Figure 1) (inside diameter, 4.0 cm) holding a glass tube (inside diameter, 2.0 cm) which separates the two aqueous phases, was used. The inner aqueous phase (source phase, SP) contained mercuric chloride (5 mL, 2.0×10⁻⁵ M) and 0.04 M hydrochloric acid, while the outer aqueous phase (receiving phase, RP) contained sodium thiocyanate (10 mL, 1.6 M). The membranes phase (MP) is compromized of 20 mL of 6.0×10⁻⁵ M IIDE in chloroform below these aqueous phases and bridged the two aqueous phases. The organic layer was magnetically stirred by a Teflon-coated magnetic bar (2.0 × 5 cm, diameter). Under these conditions, while is the mixing process perfect, the interfaces between the organic membrane and the two aqueous phases remained flat and well defined. Determination of the mercury ion concentration in both aqueous phases was carried out by CV-AAS. A similar transport experiment was carried out in the absence of the carrier for reference.

Conductometric titration

In this experiment, 15 mL of the Hg²⁺ ion solution (2.0×10⁻⁵ M) was placed in the titration cell thermostated to 25.0 ± 0.1 °C by a home made thermostat and the conductance of the solution was measured. Then, several increments of IIDE solution using a precalibrated microsyringe was added. The conductance of the solution was measured after each addition. The stability constants of the resulting complex was evaluated from the computer fitting of the molar conductance–mole ratio data to the corresponding equations according our previous publication using the program KINFIT.¹¹

RESULTS AND DISCUSSION

IIDE as a sulfur containing ligand is insoluble in water. Due to the existence of four donating sulfur atoms in the flexible structure of IIDE based on the well known hard-soft acid–base theory, it was expected that the stability of its complex with Hg²⁺ ion be higher than the other ions including alkali, alkaline earth and many transition and heavy metal ions.

Conductometric investigation of complexation of Hg²⁺ ion with IIDE

In preliminary experiments in order to obtain useful information about the stoichiometry and stability of the complexation of IIDE with Hg²⁺ ion, the system was investigated conductometrically in acetonitrile solution. It is obvious from Figure 2 that a distinct inflection point at L/Hg²⁺ molar ratio of 1:1 were occurred. The obtained stability constant for Hg-IIIDE complex using KINFIT program is 7.94 × 10⁴.
The resulting positive charged complex can form an ion pair with an appropriate anion. The neutral form ion pair can be quantitatively extracted into a suitable organic solvent. Our preliminary experiments clearly revealed that the IIDE dissolves easily in chloroform and because of low immiscibility of this solvent with water and its higher density than water, we chose it for membrane phase. Thus, in this work, we were interested to investigate the transport behavior of Hg$^{2+}$ ion through a liquid membrane containing IIDE as a potential ion carrier and chloride as a companion anion. As expected, the nature of the ligand used as carrier in the organic membrane phase has a very significant effect on the efficiency and selectivity of Hg$^{2+}$ ion transport.

Primary results showed that this membrane could transport the Hg$^{2+}$ ion selectively among other metal ions. Using acidic medium in source phase uphill transport was achieved, while in basic receiving phase no transport efficiency was observed. In order to increase the efficiency and selectivity of transport, a particular agent in the receiving phase must be used. One of the interesting possibilities for this purpose was the use of thiocyanate ion. Finally the experimental variables must be optimized in order to achieve the highest efficiency and selectivity in the transport of Hg$^{2+}$ ions across the membrane in short time.

**Influence of condition of source phase on Hg$^{2+}$ ion transport**

Most chelating ligands are conjugate bases of weak acid groups and accordingly, have a very strong affinity for hydrogen ions. Therefore the pH, play an important role in the complexation of metal ions by chelation. The pH will determine the values of the conditional stability constants of the complexes of metal ions with a desired ligand. Due to the presence of a sulfur atom in the structure of IIDE it is expected that the of its complexation ability is sensitive to pH.

It was expected that the efficiency of transport at higher pH values (>2.0) due to the formation of Hg(OH)$_2$ precipitate (white precipitate) at top of the chloroform between the source and receiving phase must be acidic. The effect of the presence of 0.1 M of different acids including picric acid, HCl, HClO$_4$, HNO$_3$ and H$_2$SO$_4$ in the source phase on the transport of Hg$^{2+}$ ion was studied and it was observed that the maximum transport occurs in the presence of 0.1 M HCl. It was found that nitrate ion is a suitable counter anion to accompany the Hg$^{2+}$–IIDE complex into the organic phase and only 30% of Hg$^{2+}$ ions was transported into the receiving phase in a long time. The transport efficiency was increased to 45% in the presence of perchlorate ion (4 h) in source phase, while addition of picric acid to the source phase increase the transport efficiency of Hg$^{2+}$ but not quantitatively. In the presence of hydrochloric acid in the source the percentage of Hg$^{2+}$ ion transported to receiving phase quantitatively was increased.

The transport efficiency of Hg$^{2+}$ ion was also found to be dependent on the concentration of HCl in the SP (Figure 3). Maximum transport occurs at HCl concentration range of 0.01-0.1 M, while further transport studies of Hg$^{2+}$ ion was carried out at 0.04 M HCl. At higher HCl concentrations there was a decrease in the percentage of transport of Hg$^{2+}$ ion, probably due to the competition of H$^+$ with Hg$^{2+}$ ion for binding to IDE and competition of chloride ion with IIDE for complexation with Hg$^{2+}$ ion. The efficiency of transport decreases at lower HCl concentrations, probably due to hydrolysis of Hg$^{2+}$ ion.

**Effect of ligand concentration**

The preferential binding of sulfur containing ligands towards Hg$^{2+}$ ion has been reported in the literature. The influence of the concentration of IIDE in the organic phase on the transport efficiency of Hg$^{2+}$ ion was studied. It was seen that the percentage of transport of Hg$^{2+}$ ion increases with an increase in IIDE concentration in the organic phase. Maximum transport occurs at concentration range of 0.5 mM-0.04 M IIDE. A further excess of the carrier had no considerable effect on the transport efficiency. Thus, a concentration of 6.0 mM of IIDE was adopted for further studies. This is most probably due to the fixed stoichiometry of the resultant 1:1 carrier– Hg$^{2+}$ ion complex.

**Effect of receiving phase agent**

The nature and composition of the receiving phase was found to have a dramatic influence on the Hg$^{2+}$ ion transport. Preliminary experiments revealed that the nature and composition of the receiving phase could have a significant effect on the efficiency and selectivity of transport. Among different receiving agents used such as hydrochloric acid, hydrobromic acid, bromide, thiosulfate, sulfuric acid, nitric acid, EDTA and thiocyanate, SCN$^-$ ion with increased complexing ability towards Hg$^{2+}$ ion acts as the most suitable receiver for the release of cation from the membrane phase into the receiving phase.

The influence of the concentration of KCSCN in the receiving phase on the transport efficiency of Hg$^{2+}$ ion was also investigated and the results are shown in Figure 4. As is obvious, while only 20% Hg$^{2+}$ ion transport occurs in the absence of the receiving agent, the transport of Hg$^{2+}$ ion increases sharply with increasing the concentration of KSCN. Quantitative transport of Hg$^{2+}$ ion occurs at a 1.6 M of the receiving agent.

The Hg$^{2+}$ ions form a complex with a deprotonated ionophore to be extracted from the source phase into the membrane phase. The
thiocyanate plays an essential role in the metal ion releasing process in the receiving phase via the formation of a ternary ligand–metal–thiocyanate complex which has been reported by other researchers.\(^{20-22}\) This co-operation probably assists the selective releasing of Hg\(^{2+}\) ion to the receiving phase, which lead to increase in the percentage of transported Hg\(^{2+}\) ion. Such a ternary complex has already been recognized as being an important transient species in biological transports.\(^{23}\)

The mechanism of transport in the presence of this carrier could be related to the rapid complex formation and exchange of ions at the membrane interface. In the membrane and receiving phase interface exchange of ligand with thiocyanate for formation of Hg (SCN)\(^2^-\) take place. Based on the overall stability constant (Log \(B_1\)) of Hg\(^{2+}\) ion complexes with respect to SCN, and Cl as (Hg(SCN))\(^2^-\), HgCl\(^+\):20.9 and 15.1,\(^{24}\) one may notice that this is consistent with the selectivity order of described procedure. As is known use of thiocyanate ion as a receiving agent effectively avoids the formation of Hg(OH)\(_2\)(s) precipitate. Consequently, this compound was used as a receiving phase agent in our later transport experiments.

**Influence of stirring rate**

The effect of stirring speed on Hg\(^{2+}\) ion transport through bulk liquid membrane was investigated. The effect of stirring rate of organic phase in the range of 50–500 rpm on the Hg\(^{2+}\) ion transport efficiency was also studied. The results revealed that replicate transport occur in the stirring rate about 300 rpm. The experimental data indicate that beyond 300 rpm the flux of Hg\(^{2+}\) ions through the bulk liquid membrane becomes independent of the stirring rate. In higher rates more than 400 rpm, some mixing of source phase and receiving phase will occur. In receiving phase of 300 rpm the thickness of the aqueous diffusion films reaches a constant limiting value. Consequently, all subsequent experiments were carried out at a stirring rate of 300 rpm in the source and receiving phase solutions.

**Effect of time**

Additionally, the concentration–time profile of Hg\(^{2+}\) ion transport was studied. Figure 5 shows a rapid rise in metal concentration in the receiving phase as well as a sharp decrease in Hg\(^{2+}\) ion concentration in the source phase during the first 120 min of transport and most of Hg\(^{2+}\) ions in source phase evacuated but residual complexes in the membrane will slowly transport to receiving phase. However, the release of Hg\(^{2+}\) ions from the membrane into the aqueous receiving phase occurs at a slower rate. It was confirmed under optimal conditions more than 97% of Hg\(^{2+}\) ions were transported from the source phase into the receiving phase after 2.5 h.

**Selectivity**

The selectivity of the membrane system in tertiary mixture was studied at optimum conditions for the transport of Hg\(^{2+}\) ion with respect to other cations which were initially present with in equimolar concentrations.

Table 1 shows the percentage of cations transported into receiving phase and remained in source phase. High stability constant of Hg\(^{2+}\) ion with IIDE and thiocyanate ion in membrane and receiving phase largely improved the method selectivity.

**Table 1. Amount of cation transported from different cation mixtures through the BLM**

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<thead>
<tr>
<th>Mixture</th>
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<th>Percentage transported</th>
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<tbody>
<tr>
<td>Hg(^{2+})</td>
<td>98</td>
<td>Hg(^{2+})</td>
<td>98</td>
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<tr>
<td>Pb(^{2+})</td>
<td>0</td>
<td>Co(^{2+})</td>
<td>0</td>
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<tr>
<td>Cd(^{2+})</td>
<td>0</td>
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<tr>
<td>Fe(^{3+})</td>
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\(^{a}\)Conditions: source solution, 5 mL of 2.0 \(\times\) 10\(^{-5}\) M Hg\(^{2+}\), 2.0 \(\times\) 10\(^{-4}\) M other cations and 0.04 M HCl in source phase 0.006 M IIDE in Membrane and 1.6 M SCN\(^-\) in receiving phase.

As can be seen from Table 1, after 2.5 h, Hg\(^{2+}\) ions were almost completely transported through the chloroform membrane from the source phase into the receiving phase. Meanwhile, most of the other metal ions used, even Cu\(^{2+}\) which is known to have affinity for this sulfur containing ligand, were hardly transferred through the membrane. The observed selectivity pattern toward Hg\(^{2+}\) ion is due to the high tendency of the cation, as a soft acid, towards donating sulfur atoms of the carrier as soft bases.

**Recommended mechanism**

The recommended mechanism for the transport of Hg\(^{2+}\) ion through bulk liquid membrane, which operated in this study, is shown schematically in Figure 6. Movement of the charged species through the hydrophobic organic membrane is accomplished by the presence of host carrier IIDE ion paired with chloride as a suitable counter
Highly selective transport of mercury(II) ion through a bulk liquid membrane

The chloride ion not only neutralizes the charged Hg\(^{2+}\)-IIDE complex but also induces a more lipophilic character to the mercury complex so that it can be readily extracted into the membrane phase. After complexation of Hg\(^{2+}\) ion with carrier on the source side of the membrane, the complex diffuses down its concentration gradient. On the receiving side of the membrane, the metal ion is released into the receiving phase via formation of a ternary complex (carrier-Hg\(^{2+}\)-SCN). Then the free carrier diffuses back across the membrane and cycle starts again. The net result is the transport of Hg\(^{2+}\) ion from the aqueous source phase to the aqueous receiving phase across the BLM.

**ACKNOWLEDGEMENT**

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**REFERENCES**