Separation of trace antimony and arsenic prior to hydride generation atomic absorption spectrometric determination

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Abstract

A separation method utilizing a synthetic zeolite (mordenite) was developed in order to eliminate the gas phase interference of Sb(III) on As(III) during quartz furnace hydride generation atomic absorption spectrometric (HGAAS) determination. The efficiency of the proposed separation method in the reduction of suppression effects of transition metal ions on As(III) signal was also investigated. Among the volatile hydride-forming elements and their different oxidation states tested (Sb(III), Sb(V), Se(IV), Se(VI), Te(IV), and Te(VI)), only Sb(III) was found to have a signal depression effect even at low (μg L⁻¹) concentrations under the experimental conditions employed. It has been shown that mordenite adsorbs Sb(III) quantitatively, even at a concentration of 1000 μg L⁻¹, at pHs greater than two, and also, it reduces the initial concentrations of the transition metal ions to lower levels which can be tolerated in many studies. The adsorption of Sb(III) on mordenite follows the Freundlich isotherm and is endothermic in nature.

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1. Introduction

Arsenic and antimony compounds are quite widespread in the environment. They have many industrial applications; they are used in semiconductors, in steel industry to reduce the ductility of the product and to increase brittleness, in semiconductors as an additive, in agriculture, and in glass manufacturing. Antimony is used in alloys and batteries. Also they have similar chemical and toxicological properties and they cause cancer in trivalent state [1].

Toxicity, reactivity and bioavailability of the elements depend on their chemical forms. This necessitates speciation analysis in which different chemical and physical forms of an analyte existing in a sample are determined. Speciation analysis mostly involves two steps, separation and determination; and one of the key issues of the speciation analysis is to preserve the composition of the sample during sampling, storage and pretreatment [2]. Sensitivity of the method should also be considered as the analytes of interest are usually at trace concentrations. Review articles on speciation of various elements can be found elsewhere [3].

There are many methods for the determination of antimony and arsenic in various matrices. Hydride generation atomic absorption spectrometry (HGAAS) is probably the most popular technique for the determination of arsenic and antimony, either directly or after a suitable pretreatment step. An important advantage of HGAAS is the separation of the analyte from the matrix. This facilitates reduction of spectral interferences and also preconcentration of the analyte. On the other hand, serious non-spectral interferences may arise in the liquid phase during hydride formation and transfer from solution (liquid phase interferences) or the interferents can influence the analyte in the gas phase (gas phase interferences) [4]. Liquid phase interferences are caused by changes in the rate of hydride release from the liquid phase or by a decreased efficiency of hydride release. Gas phase interferences
ences are apparently due to volatile species, most often by hydrides. Many studies have been done in order to reduce the interferences in HGAAS, especially caused by transition metal ions and volatile hydride-forming elements. Welz and Schubert-Jacobs [5] studied the influence of acid and sodium borohydride concentrations to reduce the interference of transition metal ions in arsenic and selenium determination and concluded that higher acid–lower sodium borohydride concentrations improve the interference-free determination. Brindle and Le [6] used thiol-containing compounds and amino acids for the enhancement of signals. Moreover, reducing agents such as L-cysteine [6,7], and thiourea [7] were also used for the reduction of transition metal ion interferences. Campos et al. [8] suggested p-cresollic acid and L-cysteine mixture to eliminate Cu and Ni interferences and tolerable concentrations were found up to 4000 and 1000 mg L\(^{-1}\) of nickel and copper, respectively. Another important interfering group is volatile hydride-forming elements that have mutual interferences during determination [4]. Welz and Melcher [9] removed the effect of Se(IV) on As(III) and As(V) adding copper to the sample solution. The authors explained the mechanism in a way that, selenium hydride, which is formed first, reacts with copper ions in the solution to form insoluble selenides (CuSe), whereas copper arsenide (Cu5As2) is soluble in acids. Therefore, the evolution of selenium hydride from solution, and its interference on arsenic signal, is prevented. On the other hand, graphite tube atomizers at temperatures >2000 °C was suggested by Dittrich and Mandy [10]. The mechanism was explained as the formation of diatomic molecules between analyte and matrix elements such as AsSb. The effect of KI on the interferences by Bi, Se and Te on Sb signal and the interferences of Sb, As, Bi, Te, and Sn on the Se signal using oxygen and a mixture of ozone were examined by Barth et al. [11]. In a study by Ay and Henden [12], a separation procedure using a GC column was applied to eliminate the inter-element interference of arsenic and antimony. In the GC column, arsine is eluted first and separated from stibine. They suggest taking stibine outside the atomizer for arsenic determination. In the next study by Erdem and Henden [13], a pyrex adsorption U-tube trap containing glass wool was placed between the drying tube and quartz tube atomizer and the mutual interference between As and Sb was investigated by selective heat decomposition of arsine and stibine.

As can be seen there are many studies on the reduction of HGAAS interferences during determination. In the present study, a different approach is proposed for the elimination/reduction of interferences on As signal prior to HGAAS determination. For the separation of As(III) and Sb(III) before hydride generation a synthetic zeolite (mordenite) was used. It has also been shown that the proposed separation method has the ability to reduce the extent of transition metal interferences.

2. Experimental

2.1. Reagents and materials

All reagents were of analytical-reagent grade. Ultra-pure water (18 MΩ) was used throughout the study. Glassware and plastic containers were cleaned by soaking in 10% (v/v) nitric acid and rinsed with distilled water prior to use. The As(III) stock standard solution was prepared from As2O3 (Fischer) by dissolving in about 10% (v/v) H2SO4. The solution was then neutralized with 0.1 M NaOH and was diluted to the required volume by ultra pure water (a similar dissolution procedure as explained in [9]). The Sb(III) stock standard solution from C4H4KO7Sb (Sigma) by dissolution in ultra pure water. Standards with lower concentrations were prepared daily from their stock standard solutions. Sodium borohydride solution (1%, m/v) was prepared from fine granulated product (Merck) and was stabilized by the addition of 0.1% (m/v) NaOH (Merck). A synthetic zeolite, mordenite (Zeolyst International) was used in the sorption experiments. Mordenite used had a SiO2/Al2O3 mole ratio of 20, Na2O of 0.08% (m/m), has a surface area of 500 m2/g and its nominal cation form was ammonium.

2.2. Instrumentation and apparatus

An atomic absorption spectrometer, Thermo Elemental Solar M6 Series with an air-acetylene burner was used in all the measurements. Antimony and arsenic determinations were performed by hydride generation atomic absorption spectrometry using their hollow cathode lamps at the wavelengths of 217.6 and 193.7 nm, respectively. A deuterium hollow cathode lamp was used for background correction in all determinations. In HGAAS, the quartz tube atomizer was 10 cm long, 8 mm in internal diameter and 10 mm in external diameter with a 4 mm bore inlet tube fused at the middle for sample introduction. Air-acetylene flame was used for heating the quartz tube externally and nitrogen was used as the carrier gas. The hydride generation system coupled with segmented flow injection unit (FI90) was used in all measurements for antimony and arsenic determinations. A schematic representation of the hydride generation apparatus used is shown in Fig. 1. It is a continuous hydride generation system segmented with air bubbles. Operating parameters for the HGAAS system are given in Table 1. In the determination of transition metal ions, flame atomic absorption spectrometry

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carrier gas (N2) flow rate (ml min(^{-1}))</td>
<td>200</td>
</tr>
<tr>
<td>HCl concentration (M)</td>
<td>0.12</td>
</tr>
<tr>
<td>HCl flow rate (ml min(^{-1}))</td>
<td>6.1</td>
</tr>
<tr>
<td>NaBH4 concentration (1% (m/v)) stabilized with</td>
<td>0.1% (m/v) NaOH</td>
</tr>
<tr>
<td>NaBH4 flow rate (ml min(^{-1}))</td>
<td>3.0</td>
</tr>
</tbody>
</table>
Solid and solution phases were separated. The supernatant solution was centrifuged and then placed on a reciprocating shaker for 15 min. After the shaking period, the mixture was centrifuged and the remaining in the solution. In addition, the supernatant solutions were analyzed to determine the concentrations of metals remaining in the solution. In addition, the supernatant solutions were spiked with 25.0 μg l⁻¹ As(III) in order to see whether the remaining concentrations of the transition metal ions had any interference on As(III) signal.

Similar sorption experiments were performed with some other volatile hydride-forming elements; namely Se(IV), Se(VI), Te(IV), Te(VI), Sb(III), and Sb(V). The concentrations of both Sb(III) and As(III) was examined separately in the pH range 2.0–10.0, and also in 1.0 and 4.0 M HCl. The concentrations of metal ions in the supernatant solutions were determined by FAAS and the percent sorption of transition metal ions on mordenite was calculated for each pH. The effect of initial concentrations of the transition metal ions on sorption on mordenite was investigated for the concentrations 1.0, 10.0, 100.0, and 1000.0 mg l⁻¹. After the usual shaking and centrifugation steps, the supernatant solutions were analyzed to determine the concentrations of metals remaining in the solution. In addition, the supernatant solutions were spiked with 25.0 μg l⁻¹ As(III) in order to see whether the remaining concentrations of the transition metal ions had any interference on As(III) signal.

The interference effect of several transition metal ions on As(III) signal in HGAAS measurements was investigated in such a way that various concentrations of the metals were prepared and spiked with As(III). The change in HGAAS signals of As(III) as a function of metal concentration was obtained. For this purpose, 30.0 ml of the separate transition metal ion solutions with the concentrations ranging from 0.1 to 5.0 mg l⁻¹, were spiked with 25.0 μg l⁻¹ As(III). The solutions were analyzed with HGAAS system for As(III). In a separate study, in order to investigate the interference effect of other volatile hydride-forming elements on As(III) signal, a similar strategy was followed with Se(IV), Se(VI), Te(IV), Te(VI), Sb(III), and Sb(V). The concentrations of both Sb(III) and As(III) were measured with HGAAS system.

Sorption characteristics of mordenite towards the selected transition metal ions and hydride-forming elements were investigated through batch process. The general strategy followed in the batch process was as follows: 30.0 ml of the related solution was prepared and 0.3 g of mordenite was added into this solution. The mixture was shaken for 1–2 min manually and then placed on a reciprocating shaker for 15 min. After the shaking period the mixture was centrifuged and the solid and solution phases were separated. The supernatant solution was analyzed by HGAAS for As(III) and Sb(III) and FAAS for the transition metals.

The effect of solution pH on the sorption of transition metal ions on mordenite was examined in the pH range 2.0–8.0, for a transition metal concentration of 5.0 mg l⁻¹. In the adjustment of pH, NH₃ (0.1–1.0 M) and HCl (0.1–1.0 M) were used. After the above-mentioned batch process, the concentrations of metal ions in the supernatant solutions were determined by FAAS and the percent sorption of transition metal ions on mordenite was calculated for each pH.

The effect of initial concentrations of the transition metal ions on sorption on mordenite was investigated for the concentrations 1.0, 10.0, 100.0, and 1000.0 mg l⁻¹. After the usual shaking and centrifugation steps, the supernatant solutions were analyzed to determine the concentrations of metals remaining in the solution. In addition, the supernatant solutions were spiked with 25.0 μg l⁻¹ As(III) in order to see whether the remaining concentrations of the transition metal ions had any interference on As(III) signal.

Sorption studies with mordenite

Sorption characteristics of mordenite towards the selected transition metal ions and hydride-forming elements were investigated through batch process. The general strategy followed in the batch process was as follows: 30.0 ml of the related solution was prepared and 0.3 g of mordenite was added into this solution. The mixture was shaken for 1–2 min manually and then placed on a reciprocating shaker for 15 min. After the shaking period the mixture was centrifuged and the solid and solution phases were separated. The supernatant solution was analyzed by HGAAS for As(III) and Sb(III) and FAAS for the transition metals.

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Similar sorption experiments were performed with some other volatile hydride-forming elements; namely Se(IV), Se(VI), Te(IV), Te(VI), Sb(III), and Sb(V). The concentrations of all the species were 40.0 μg l⁻¹.

Based on the results of the interference study related with hydride-forming elements, a special attention was given to Sb(III) since it showed the most suppressive interference on As(III) signal. The effect of solution pH on the sorption of Sb(III) and As(III) was examined separately in the pH range 2.0–10.0, and also in 1.0 and 4.0 M HCl. The concentrations of both Sb(III) and As(III) were 40.0 μg l⁻¹.

In addition to the pH experiments, the equilibrium sorption isotherm studies were conducted in order to better clarify the sorption of Sb on mordenite. These studies were also performed through batch process. The accurately weighed amounts of mordenite (0.300 or 1.000 g) were added into 30.0 ml of solutions containing the specified concentrations of Sb(III) (ranging from 5.0 to 500.0 mg l⁻¹) and the mixtures were shaken in a thermostated water bath (30 or 50 °C) for an hour. At the end of the shaking period the solid and solution phases were separated through centrifugation and the concentrations of Sb in the supernatant solutions were determined by HGAAS or FAAS. The sorbed amount of Sb per unit mass of mordenite was calculated from the mass balance.

3. Results and discussion

3.1. Interference studies

3.1.1. Effect of transition metal ions on As(III) signal

As explained in Section 2.3, the change in As(III) signal as a function of transition metal concentrations ranging from 0.1 to 5.0 mg l⁻¹ was investigated and the results are shown in Fig. 2. As can be seen from the figure, the highest signal depression was observed for Ni²⁺ and Cu²⁺ whereas the As(III) signal was not affected dramatically for...
Ag⁺ and Pb²⁺. These findings are in accordance with some studies in literature [14]. Higher concentrations (>5.0 mg l⁻¹) were not tried because of the reduction of metal ions to elemental state with the injection of NaBH₄ and thereby increasing the possibility of damaging and clogging of the pump tubings and narrow connection points of the SFI–HGAAS.

3.1.2. Effect of volatile hydride-forming elements on As(III) signal

Volatile hydride-forming elements can cause serious gas phase interferences in HGAAS system. These interferences can take place either during the transfer of the hydride from the sample solution to the atomizer, or in the atomizer [4]. In order to test whether or not Se(IV), Te(IV) or Sb(III) had a suppression effect on As(III) signal, a 25.0 µg l⁻¹ As(III) standard in separate solutions of potential interferents; namely, 250.0 µg l⁻¹ Sb(III), Se(IV) and Te(IV), in different HCl concentrations (0.12–3.0 M) was prepared. The relative As(III) signals are shown in Fig. 3 (a). As can be seen from the figure, Sb(III) exhibited almost a constant signal suppression (∼60%) at all HCl concentrations studied whereas no signal suppression was observed from Se(IV) or Te(IV) in 12 M HCl. But, immediately after this HCl concentration (e.g. in 0.24 M HCl), a dramatic decrease in As(III) signal was obtained with Se(IV), almost 80% suppression, and about 20% decrease with Te(IV). These results can show the strong dependence of hydride generation reaction on sample acidity.

After this finding, effect of interferent concentration on As(III) signal was investigated in two different HCl concentrations. For this purpose, 25.0 µg l⁻¹ As(III) standard was prepared in varying concentrations (25.0–1000 µg l⁻¹) of Sb(III), Se(IV) and Te(IV) in 0.12 and 1.2 M HCl. The respective figures are shown in Fig. 3 (b) and (c). As seen in Fig. 3 (b), in 0.12 M HCl, Sb(III) suppressed the As(III) signal whereas no signal suppression was observed from Se(IV) and Te(IV). On the other hand, 1.2 M HCl created a more convenient environment for Se(IV) and Te(IV) to exhibit their interference effect on As(III) signal (Fig. 3 (c)). Se(IV) was more effective and even no As(III) signal was obtained in presence of 500 µg l⁻¹ Se(IV) in 1.2 M HCl. In addition, the pH of the reaction mixture after mixing the sample (in different HCl concentrations) and the reducing agent (1.0% (m/v) NaBH₄ in 0.1% (m/v) NaOH) was measured in the SFI–HGAAS system used. Initial concentrations of HCl and the pH of the mixtures were as follows; 0.012 M (pH 9.8), 0.060 M (pH 9.0), 0.12 M (pH 6.6), 0.18 M (pH 1.4), 0.24 M (pH 0.4), 0.60 M (pH 0.5), and 1.2 M (pH 0.1). These results are in accordance with the results mentioned earlier by Belcher et al. [15]. According to the results, H₂Se and H₂Te could possibly not escape from the solution when the initial concentration of HCl is 0.12 M (after mixing sample and reducing agent, pH of the solution was 6.6). On the other hand, when the initial HCl concentration was 0.24 M, pH of the solution changed to 1.0 after mixing, and therefore H₂Se and H₂Te could escape from the solution and therefore cause gas phase interference.
3.2. Sorption studies

3.2.1. Sorption of transition metal ions

Use of natural and synthetic zeolites including mordenite for the sorption of metal ions has been shown by many researchers [16–18]. In the present study, the initial experiment was planned to reveal the change in the percent sorption of the transition metal ions on mordenite versus the pH of the solution. It was found that any pH between 4.0 and 8.0 can be used for the quantitative sorption of all four transition metal ions. An interesting observation in this figure is that Cu$^{2+}$ and Ni$^{2+}$ showed a very similar behavior during sorption as seen in interference studies given in Fig. 2.

After determining the appropriate pH values, the specified amount of mordenite (0.3 g) was added into separate solutions of the four metal ions with different initial concentrations of 1.0, 10.0, 100.0, and 1000.0 mg l$^{-1}$. The concentrations of the metal ions that remained in the solution after sorption are given in Table 2. Also given in the table, the percent suppression of the As(III) signal when the supernatant solutions were spiked with 25.0 μg l$^{-1}$ As(III). As can clearly be seen from the table, when the initial concentration of the metals is 1000.0 mg l$^{-1}$, mordenite cannot decrease their concentration to a tolerable level. The percent suppression effect of the remaining concentration of metal ions was not examined due to the possibility that such high concentrations could have caused some uncontrollable problems such as the formation of dispersed metals in the solution and the clogging of connection points in the SFI–HGAAS system. In the case of Ag$^+$ and Ni$^{2+}$, even an initial concentration of 100.0 mg l$^{-1}$ did not cause any suppression since the percent sorption was still 98 and 100, respectively. The concentration of Ag$^+$ remained in the solution was 2.0 mg l$^{-1}$ and this relatively low concentration caused only a 4 ± 3% suppression in As(III) signal. This finding supports also the observations given in Fig. 2.

The concentrations of Cu$^{2+}$ and Ni$^{2+}$ remained in the solution after 100.0 mg l$^{-1}$ initial concentration of the metal ions were mixed with mordenite were 10 ± 1 and 19 ± 1 mg l$^{-1}$, respectively. These concentrations of Cu$^{2+}$ and Ni$^{2+}$ were still very effective in suppression of As(III) signal. In both cases, the injection of NaBH$_4$ caused the metal ions to be converted to elemental state and be dispersed in the solution, which possibly adsorbed AsH$_3$ as soon as it was formed. With the initial concentrations of 1.0 and 10.0 mg l$^{-1}$, mordenite decreased the concentrations of the metal ions below 0.04 mg l$^{-1}$, and this and lower concentrations of the metals had a very slight suppression on As(III) signal only in the case of Ni$^{2+}$. These results indicate clearly the potential of mordenite in reducing the interference effect of transition metal ions on As(III) signal.

### Table 2

<table>
<thead>
<tr>
<th>Initial concentration (mg l$^{-1}$)</th>
<th>Concentration in effluent (mg l$^{-1}$)</th>
<th>% Sorption</th>
<th>% Suppression in As signal</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Cu$^{2+}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.01 ± 0</td>
<td>99 ± 0</td>
<td>2 ± 2</td>
</tr>
<tr>
<td>10</td>
<td>0.04 ± 0</td>
<td>100 ± 0</td>
<td>4 ± 2</td>
</tr>
<tr>
<td>100</td>
<td>10 ± 1</td>
<td>90 ± 1</td>
<td>45 ± 4</td>
</tr>
<tr>
<td>1000</td>
<td>778 ± 33</td>
<td>22 ± 3</td>
<td>NA</td>
</tr>
<tr>
<td>(b) Ni$^{2+}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>ND</td>
<td>100</td>
<td>NS</td>
</tr>
<tr>
<td>10</td>
<td>0.03 ± 0</td>
<td>100 ± 0</td>
<td>1 ± 1</td>
</tr>
<tr>
<td>100</td>
<td>19 ± 0</td>
<td>81 ± 1</td>
<td>96 ± 1</td>
</tr>
<tr>
<td>1000</td>
<td>822 ± 5</td>
<td>18 ± 1</td>
<td>NA</td>
</tr>
<tr>
<td>(c) Ag$^+$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.023 ± 0</td>
<td>98 ± 1</td>
<td>NS</td>
</tr>
<tr>
<td>10</td>
<td>0.04 ± 0</td>
<td>99 ± 0</td>
<td>NS</td>
</tr>
<tr>
<td>100</td>
<td>2 ± 0</td>
<td>98 ± 0</td>
<td>4 ± 3</td>
</tr>
<tr>
<td>1000</td>
<td>362 ± 7</td>
<td>64 ± 1</td>
<td>NA</td>
</tr>
<tr>
<td>(d) Pb$^{2+}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>ND</td>
<td>100</td>
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<tr>
<td>10</td>
<td>ND</td>
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<td>NS</td>
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<tr>
<td>100</td>
<td>0.4 ± 0</td>
<td>100 ± 0</td>
<td>NS</td>
</tr>
<tr>
<td>1000</td>
<td>301 ± 12</td>
<td>69 ± 0</td>
<td>NA</td>
</tr>
</tbody>
</table>

NA: not analyzed; ND: not detected; NS: no suppression.

3.2.2. Sorption of volatile hydride-forming elements

The sorption of different oxidation states of Se, Te, and Sb on mordenite was investigated as explained in Section 2.4. At the end of the usual sorption procedure, the lower oxidation states of the volatile hydride-forming elements; namely, Sb(III), Se(IV) and Te(IV) were determined by SFI–HGAAS directly after bringing it to the required acidity for hydride generation step. On the other hand, a prereduction step was applied to measure the higher oxidation states (Sb(V), Se(VI) and Te(VI)). Treatment of the supernatant solution with 2% (m/v) L-cysteine was sufficient for Sb(V) to Sb(III) reduction [19] whereas heating the solution in 6M HCl served as an efficient prereduction step for Se(VII) [20] and Te(VI) [21].

Among the elements and their oxidation states examined, mordenite showed selectivity only to Sb(III) (95 ± 2%)
whereas percentage uptake values towards Te(IV), Te(VI), Se(IV), Se(VI), Sb(III), and Sb(V) were 19 ± 2, 4 ± 1, 9 ± 8, 1 ± 1, and 2 ± 2, respectively. Except Sb(III), none of the volatile hydride-forming elements investigated had a suppression effect on As(III) signal, under the experimental conditions employed (Fig. 3(b)).

Different sorption behavior of mordenite towards Sb(III) necessitated a further investigation of the case. For this purpose, effect of solution pH and HCl concentration on the sorption of As(III), As(V), Sb(III), and Sb(V) on mordenite were examined more closely. The results are shown in Fig. 4. It can be said that mordenite shows high adsorption selectivity for Sb(III) among the hydride-forming elements investigated may also lead to such applications as separation and preconcentration of Sb(III) from solutions containing the species with similar chemical properties like As(III).

Another set of experiments was also planned in order to reveal some of physicochemical aspects of Sb(III) sorption on mordenite as explained in Section 2.4.

### 3.2.3. Sorption isotherm and thermodynamic relations in Sb uptake by mordenite

The sorption experiments of Sb(III) were performed in the batch process. The equilibrium relationship between the amount of Sb(III) adsorbed per unit mass of mordenite ([C]s) and the residual Sb(III) concentration ([C]l) in solution phase were expressed using adsorption isotherms. The Sb(III) concentrations ranged from 5.00 to 500 mg l−1 and two different amounts of solid (0.3 and 1 g) and temperatures (30 and 50 °C) were examined. The applicability of the Freundlich sorption isotherm was tested under these specified conditions.

The general expression of Freundlich isotherm is given as:

\[
[C]_s = k [C]_l^n \quad \text{(1)}
\]

where \([C]_s\) is the equilibrium concentration of the sorbed species on the solid phase (meq g−1); \([C]_l\) the equilibrium concentration of the sorbate in the liquid phase (meq ml−1); \(k\) the constant related to sorption affinity; \(n\) the constant related to linearity of the sorption curve.

This expression can be linearized as:

\[
\log[C]_s = \log k - n \log[C]_l \quad \text{(2)}
\]

Thus, plotting \(\log[C]_s\) versus \(\log[C]_l\) would lead to linear behavior with ‘\(n\)’ being the slope and ‘\(k\)’ the intercept of the isotherms. Freundlich isotherm model allows for several kinds of adsorption sites on the solid, each kind having a different heat of adsorption. The Freundlich isotherm represents well the data at low and intermediate concentrations and is a good model for heterogeneous surfaces. When the value of Freundlich constant ‘\(n\)’ is equal to unity, Freundlich equation becomes linear and the Freundlich constant ‘\(k\)’ becomes equivalent to the distribution ratio, \(R_d\) which is an empirical constant usually used in the quantification of the sorption process [22].

As can be seen from Fig. 5, the adsorption data for Sb on mordenite follows the Freundlich isotherm. The characteristic Freundlich constants are given in Table 3.

In batch adsorption processes, the adsorption reaction can be written as:

\[
C_1 + y \rightarrow C_y \quad \text{(3)}
\]

where \(C_1\) and \(C_y\) stands, respectively, for the equilibrium concentration of the sorbate species in the liquid and on the solid sorbent, and \(y\) refers to the concentration of the sorption sites on the sorbent. The importance of \(y\) stems from the fact that sorption sites can act as the limiting reactant when the sorption reaction takes place at higher loadings of the sorbate, the thing that leads to a plateau of saturation beyond a certain high initial concentration. For dilute solutions, the fractional coverage of the sorption sites can be assumed to be small enough to justify approximating the activity (or concentration) of the sorption sites to be constant, and the activity coefficients of the sorbate species in liquid solution and at the solid to be equal to unity. Under these approximations, the equilibrium constant of the above reaction becomes equivalent to the distribution ratio, \(R_d\) (ml g−1), an empirical constant that is valid at a certain concentration for a particular set of experimental conditions.

### Table 3

<table>
<thead>
<tr>
<th>m/V (g l−1)</th>
<th>(n)</th>
<th>(k) (meq g−1)</th>
<th>(\Delta\Delta G^\circ) (kJ mol−1)</th>
<th>(\Delta\Delta H^\circ) (kJ mol−1)</th>
<th>(\Delta\Delta S^\circ) (J mol−1 K−1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.3</td>
<td>0.32</td>
<td>0.27</td>
<td>0.20</td>
<td>0.15</td>
<td>–8.2</td>
</tr>
<tr>
<td>33.3</td>
<td>0.19</td>
<td>0.27</td>
<td>0.04</td>
<td>0.11</td>
<td>–8.8</td>
</tr>
</tbody>
</table>
conditions, and defined as:

\( R_d = \frac{[C_s]}{[C_l]} \)  (4)

under such condition, when the equilibrium is attained, the distribution ratio may be related to the change in Gibbs free energy, \( \Delta G_m \), by the following equation:

\[ \Delta G_m = -RT \ln R_d \]  (5)

Gibbs free energy can also be written in terms of enthalpy change, \( \Delta H_m \), and the entropy change, \( \Delta S_m \), as given below:

\[ \Delta G_m = \Delta H_m - T\Delta S_m \]  (6)

Here, \( \Delta H_m \) can be calculated using the relation:

\[ \Delta H_m = \frac{R(T_2) \ln \left( \frac{1}{T_1} \frac{1}{T_2} \right)}{T_2 - T_1} \]  (7)

As can be seen from Table 3, the calculated \( \Delta G_m \) values indicate that the sorption process is spontaneous in all cases. It is also obvious that spontaneity increases with increasing temperature leading to higher coverage at higher temperatures. The endothermic nature of sorption of Sb on mordenite is indicated by the positive values of the enthalpy change \( \Delta H_m \). The positive sign of the \( \Delta S_m \) values indicate that sorption leads to more randomness/mobility in the system, possibly due to the dehydration steps and/or release of a larger number of ions or molecules by mordenite upon fixation of Sb species.

As explained in Section 3.1.2, Se(IV) and Te(IV) did not cause any suppression on As(III) signal under the experimental conditions employed whereas Sb(III) had a serious suppression effect even at very low concentrations (Fig. 3(b)). Knowing this and the capability of mordenite in sorbing Sb(III), it was thought that the sample could be pretreated with mordenite which will guarantee the interference-free determination of As(III) in the presence of the above-mentioned volatile hydride-forming elements. This was confirmed as follows; 40.0μg l⁻¹ As(III) was prepared in varying concentrations of Sb(III), from 50.0 to 1000.0μg l⁻¹, and the solutions were analyzed both directly and after the proposed mordenite treatment. The results are demonstrated in Fig. 6. As can be seen from the figure, the presence of Sb(III) will affect the As(III) signal starting from very low concentrations. But, when the same solutions were processed by mordenite, the As(III) signal is regenerated and the interference of Sb(III) is reduced to a tolerable level. Even for a solution containing 1000.0μg l⁻¹ Sb(III), 87 ± 2% of the 40.0μg l⁻¹ As(III) signal is regenerated.

3.2.4. Reduction of Sb(III) interference on As(III) signal

Meanwhile, the uptake of Sb(III) species on mordenite was studied using XRD and SEM-EDX techniques. According to the XRD patterns, mordenite reflections were unaffected, in terms of peak positions, intensity, and peak shapes, thus indicating the structural stability of this mineral upon retention of Sb(III) species, under the experimental conditions employed. Moreover SEM micrographs showed no change in the morphology or crystal size of mordenite particles upon loading with Sb(III), the signals of which were observed to be equally distributed on the surface of mordenite, as implied by EDX mapping analysis.

4. Conclusion

It has been demonstrated that a pretreatment method utilizing a synthetic zeolite, mordenite, offers an efficient new route for the interference-free determination of As(III) in the presence of Sb(III) which is one of the most suppressive interferences in As(III) determination by HGAAS. Morden-
ite exhibits high selectivity towards Sb(III) and the sorption was observed to be endothermic and followed Freundlich isotherm model. Several other hydride-forming elements, such as Se(IV), Se(VI), Te(IV), Te(VI) were found not to interfere with As(III) determination with the experimental conditions employed. The proposed pretreatment method was also shown to be capable of reducing the interference effects of several transition metal ions (Ag⁺, Ni²⁺, Cu²⁺, and Pb²⁺) in the reaction medium.

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References