1996

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BecotteHaigh, P; Tyson, JF; Denoyer, E; and Hinds, MW, "Determination of arsenic in gold by flow injection inductively coupled plasma mass spectrometry with matrix removal by reductive precipitation" (1996). SPECTROCHIMICA ACTA PART B-ATOMIC SPECTROSCOPY. 1078.
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Determination of arsenic in gold by flow injection inductively coupled plasma mass spectrometry with matrix removal by reductive precipitation

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Received 19 March 1996; accepted 22 April 1996

Abstract

Arsenic was determined in gold by flow injection hydride generation inductively coupled plasma-mass spectrometry following a batch mode reductive precipitation removal of the interfering gold matrix. A solution of potassium iodide, L-ascorbic acid, and hydrochloric acid was used as the reductant. The recovery of gold by precipitation and filtration was 99 ± 3%. The detection limit for arsenic in gold was 55 ng g⁻¹ in the solid. The concentration of arsenic that was determined in the Royal Canadian Mint gold sample FAU-10 was 29.7 µg g⁻¹ in the solid; this value was indistinguishable, with 95% confidence, from values determined at the Royal Canadian Mint by graphite furnace atomic absorption spectrometry and by inductively coupled plasma–mass spectrometry. The standard deviation for four replicate determinations of the arsenic in FAU-10 was 0.972 µg g⁻¹ in the solid.

Keywords: Arsenic; Flow injection; Hydride generation; ICP-MS; Matrix removal; Reductive precipitation

1. Introduction

The determination of the common impurities in gold is important in assessing the value, physical properties, and point of origin of a sample of gold [1–3]. Arsenic is often present at ppm concentrations in gold. Digestion and dilution of the sample further reduce the concentration of arsenic to be determined, creating a need for a very sensitive analytical method. A method for this determination should also be precise so that sample homogeneity may be determined [1]. In addition, the method should require a small sample size in order to minimize the cost.

Electrothermal atomic absorption spectrometry (ET–AAS) with batch anion-exchange removal of the gold matrix [4], and laser ablation inductively coupled plasma mass spectrometry (LA–ICP–MS) [1,3] have both been used to determine trace elements in gold samples. Some of the advantages of the LA–ICP–MS method are: solid sampling, microsampling, sensitivity, and multi-element capability. The main drawbacks to LA–ICP–MS are the lack of suitable calibration and certified reference materials, and poorer precision relative to solution analysis [5].

A method for trace element determinations in gold using direct nebulization into the ICP–MS would
potentially suffer from several interferences. The conventional aqua regia gold digestion procedure introduces Cl\textsuperscript-, which produces the polyatomic interference \( ^{40}\text{Ar}^3\text{Cl}^+ \) on mono-isotopic \( ^{75}\text{As}^+ \). The total dissolved solids should not exceed 0.1-0.5\% m/m in the diluted sample or deposition of material on the sample and skimmer cones will cause drift and sensitivity loss problems [5-7]. Dilution of the matrix to reduce the total dissolved solids also dilutes the arsenic. Gold introduced could also suppress the analyte signal by its space charge effect [8-10].

The application of flow injection (FI) techniques to atomic spectroscopy in order to reduce interferences is well documented [11-17]. Several workers have applied FI to ICP-MS [5,7,18-21]. Sample and reagent consumption, as well as the total amount of dissolved solids that enter the ICP-MS are reduced with FI techniques.

Hydride generation (HG) is often used to increase the transport efficiency and therefore the sensitivity for certain vapor forming elements, such as As [18,20,22]. Another advantage of HG is that the analyte is removed from interfering matrix constituents. A depressive interference upon the hydride signal is often encountered, however, because of the adsorption of the hydride on finely divided transition metal particles that may be precipitated by the reluctant [23,24]. Preliminary experiments had shown this to be the case for the determination of As in gold [25], despite the successful use of the kinetic discrimination with a FI system for the determination of As in nickel [26].

2. Experimental

2.1. Instrumentation

An Elan 5000 ICP mass spectrometer (Perkin-Elmer SCIEX, Norwalk, CT, USA) equipped with a FIAS-200 (Bodenseewerk Perkin-Elmer, Überlingen, Germany) was used for this work. The ICP-MS operating conditions are summarized in Table 1. The operating conditions (nebulizer flow and ion optics settings) were optimized by pumping a 1 ppb (m/m) As(III) standard through the carrier line, and generating the hydride continuously. The sensitivity observed during this optimization was more than adequate for the analysis, so that the reagent concentrations and manifold parameters did not have to be optimized further. The FIAS operating program is summarized in Table 2.

The FI manifold is shown in Fig. 1. The gas–liquid separator (GLS) (Perkin-Elmer, Norwalk, CT, USA) is the same as the one described by Welz and Schubert-Jacobs [27], except that the device was filled with glass beads to the top of the waste drain. The nebulizer was replaced with a Teflon adapter that screwed directly to the torch mount and had a 2 mm i.d. central channel. The GLS was connected directly to the Teflon torch adapter via 1 m of 1.24 mm i.d. Teflon tubing. The injection volume was 500 µl.

2.2. Reagents and sample preparation

All reagents were of analytical reagent grade unless otherwise stated. All reagents and samples were prepared with distilled, deionized water (E-Pure, Barnsted, Boston, MA). Fisher Scientific (Fair

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Table 1

<table>
<thead>
<tr>
<th>Instrumental parameters</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>t.f. power/W</td>
<td>1000</td>
</tr>
<tr>
<td>Argon gas flow:</td>
<td></td>
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<tr>
<td>Outer gas flow rate/(l min(^{-1}))</td>
<td>15</td>
</tr>
<tr>
<td>Intermediate gas flow rate/(l min(^{-1}))</td>
<td>0.80</td>
</tr>
<tr>
<td>Aerosol carrier gas flow rate/(l min(^{-1}))</td>
<td>0.9-1.1</td>
</tr>
<tr>
<td>Sample and skimmer cones</td>
<td>nickel</td>
</tr>
<tr>
<td>Mass spectrometer running pressure/Pa</td>
<td>(130 \times 10^{-6})</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Data acquisition parameters</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Mode</td>
<td>Peak hop transient</td>
</tr>
<tr>
<td>Dwell time/ms</td>
<td>40</td>
</tr>
<tr>
<td>Points per peak</td>
<td>1</td>
</tr>
<tr>
<td>Readings per replicate</td>
<td>250</td>
</tr>
<tr>
<td>Number of replicates</td>
<td>5</td>
</tr>
</tbody>
</table>
Lawn, NJ, USA) brand concentrated hydrochloric and nitric acids were used throughout this work.

The sodium tetrahydroborate solution was prepared fresh daily by dissolving sodium tetrahydroborate (Fisher Scientific) in 0.05% (m/m) sodium hydroxide (certified, Fisher Scientific), then suction filtering the solution through #1 Whatman filter paper. A 1000 ppm (m/v) stock solution of As(V) was prepared by dissolving sodium arsenate (certified, Fisher) in 10% (v/v) nitric acid. A 500 ppb (m/m) As(III) stock solution was prepared by adding 20 ml of 10% (m/v) potassium iodide (Aldrich, Milwaukee, WI), 10% L-ascorbic acid (Fisher), and 20 ml concentrated hydrochloric acid to 0.1 ml of the 1000 ppm (m/v) As(V) stock solution in a 250 ml Nalgene bottle, diluting to 200 g, and allowing the solution to stand for 45 min.

Four portions of the gold sample, each between 0.05 and 0.17 g (FAU-10, 29.4 ppm (m/m) As) supplied by the Royal Canadian Mint (RCM) were each dissolved by placing the material in a 50 ml beaker with 1 ml concentrated nitric acid and 3 ml concentrated hydrochloric acid, covering with a watchglass, and heating slowly until the resulting solution evaporated to approximately half of its original volume. Each solution was then diluted with 1% (v/v) hydrochloric acid to yield a yellow solution containing 0.05% Au (m/m). Forty gram aliquots of the 0.05% (m/m) FAU-10 sample solutions were reduced by adding 20 ml of 10% (m/v) L-ascorbic acid, 10% (m/v) KI solution, 20 ml hydrochloric acid and diluting to 200 g. An opaque brown precipitate formed in the samples upon the addition of the KI/L-ascorbic acid solution. This precipitate was removed by filtering through glass wool, dissolved in aqua regia, and diluted to 200 ml to produce a yellow solution. The precipitate immediately assumed a metallic gold appearance on addition of aqua regia. A 0.1 ml aliquot of the solution in which the precipitate was dissolved was diluted to 100 g. The filtrate was clear and colorless.

2.3. Standards

Standards were prepared by dissolving and pre-reducing/filtering 0.082 g high purity gold (99.999%, RCM) in the same manner as the samples, and adding appropriate amounts of the 500 ppb (m/m) As(III) standard before the final dilution, to produce 0, 0.99, 2.5 and 5.0 ppb (m/m) As(III) standards. This procedure was followed in an attempt to match the sample matrix composition. These calibration standards were used for the determination of As in the samples. A set of calibration standards was prepared by spiking 10% hydrochloric acid with the 500 ppb (m/m) As(III) solution to produce 0, 1.0, 2.0, 4.0 and 6.0 ppb (m/m) As(III) standards for comparison with:

<table>
<thead>
<tr>
<th>Step</th>
<th>Time/s</th>
<th>Pump 1/(rev min⁻¹)</th>
<th>Pump 2/(rev min⁻¹)</th>
<th>Valve position</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-sample</td>
<td>20</td>
<td>120</td>
<td>120</td>
<td>Rinse/fill loop</td>
</tr>
<tr>
<td>1</td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>Fill loop</td>
</tr>
<tr>
<td>2</td>
<td>15</td>
<td>120</td>
<td>0</td>
<td>Inject</td>
</tr>
</tbody>
</table>
Table 3
Results (in µg/g) for the determination of As in FAU-10

<table>
<thead>
<tr>
<th>Method</th>
<th>Concentration determined</th>
<th>Standard Dev.</th>
<th>Replicates</th>
</tr>
</thead>
<tbody>
<tr>
<td>FI–HG–ICP–MS with pre-reduction</td>
<td>29.7</td>
<td>0.972</td>
<td>4</td>
</tr>
<tr>
<td>ICP–MS</td>
<td>28.3</td>
<td>1.26</td>
<td>8</td>
</tr>
<tr>
<td>GFAAS*</td>
<td>30.4</td>
<td>1.10</td>
<td>10</td>
</tr>
</tbody>
</table>

* Data obtained by the Royal Canadian Mint.

the matrix matched standards. Another set of calibration standards (0, 0.053, 0.026, 0.13, 0.26, 0.93 and 3.5 ppb (m/m) As(III) was subsequently prepared by pre-reducing and spiking gold solutions to investigate the calibration response at lower concentrations, and to calculate a detection limit. A 10 ppb (m/m) gold solution was prepared as a single point calibration for the determination of the gold recovery by the precipitation. The gold determination was performed by FI ICP–MS with a single-line manifold. The gold content in a pre-reduced/filtered sample was also determined by this method.

3. Results and discussion

When a sample containing gold was injected a black precipitate formed downstream of the confluence point at which the sodium tetrahydroborate solution was merged. Once this precipitate formed subsequent injections of samples and standards gave no signal. It was hoped that this gold interference would be eliminated by separation of the gold by precipitation during sample pretreatment. However, it was considered possible that arsenic might coprecipitate with the gold, leading to inaccurate determinations.

3.1. Gold recovery

The concentration of gold in the solution in which the precipitate was dissolved would be 500 ppb (m/m) if all of the gold was recovered by the precipitation/filtration/dissolution procedure mentioned above. The gold content of the solution was 492 ± 16 ppb (m/m). This value corresponds to 99 ± 3% recovery (95% confidence). The gold content of a pre-reduced/filtered sample solution was determined to be 5.1 ± 0.8 ppb (m/m, 95% confidence). Gold was shown to be completely removed from the matrix by a batch mode reduction prior to analysis.

3.2. Determination of arsenic

The results for the determination of arsenic in FAU-10 gold by FI–HG–ICP–MS are compared with the values obtained by the RCM by ICP–MS and graphite furnace atomic absorption spectrometry (GFAAS) in Table 3. The values determined in this work are indistinguishable from the values determined by the RCM, with 95% confidence. Therefore, arsenic was not lost during the gold precipitation.

The sensitivity of the matrix matched standards calibration was 200 kilo counts per second (kcps) per ppb As(III), as compared to 180 kcps per ppb As(III) for the non-matrix matched standards. The difference between these values may be due to the instrumental drift that was observed. The correlation coefficient for each of these calibrations was greater than 0.9999. Approximately 60 injections per hour were possible with the FIAS program used.

The calibration that was performed using the lower concentration matrix-matched standards had a correlation coefficient of 0.999. The detection limit for this method, based upon 3 standard deviations of the blank response, was 55 ppb (m/m) As in the solid sample. Since the RCM requires that the method detect arsenic at concentrations down to 1 ppm (m/m) in the solid, this method would be adequate for their purposes.

Acknowledgements

The provision and maintenance of the Perkin-Elmer Sciex Elan 5000 and FIAS 200 instruments at the
Amherst campus by Perkin-Elmer are gratefully acknowledged.

References