Determination of Trace Amounts of Lead Using the Flotation-spectrophotometric method

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Abstract: The present study describes a simple and highly selective method for separation, preconcentration and spectrophotometric determination of extremely low concentrations of lead. It is based on flotation of a complex of Pb²⁺ ions and Alizarin yellow between aqueous and n-hexane interface at pH = 6. The proposed procedure is also applied for determination of lead in both tap water and prepared sea water samples. Beer’s Law was obeyed over the concentration range of 3.86 × 10⁻⁸ To 8.20 × 10⁻⁷ molL⁻¹ (8–170 ngmL⁻¹) with an apparent molar absorptivity of 1.33 × 10⁶ molL⁻¹ cm⁻¹ for a 100 mL aliquot of the water sample. The detection limit (n = 10) was 8.7 × 10⁻⁹ molL⁻¹ (1.0 ngmL⁻¹) and the Relative standard deviation (R.S.D), (n = 10) for 7.2 × 10⁻⁷ molL⁻¹ (150 ngmL⁻¹) of Pb (II) was 4.36%. A notable advantage of the method is that the determination of Pb (II) is free from the interference of almost all cations and ions found in the environment and waste water samples. The determination of Pb (II) in tap and synthetic seawater samples was also carried out by the present method. The results were satisfactorily comparable so that the applicability of the proposed method was confirmed to the real samples.

Keywords: flotation-extraction, lead, alizarin yellow, spectrophotometry
Introduction

The impact of the effects of trace chemical species in the environmental and industrial samples on human health has fostered development of analytical techniques capable of addressing these issues. Accurate and reliable analytical methods possessing high sensitivity and selectivity, coupled with convenience and economy, applicable to real world samples are required. Although some analytical techniques such as AAS, GFAAS, ICP-OES, ICP-MS, or polarographic analysis and atomic spectroscopy with high capability of determination of trace elements have already been used, however, due to their high cost and in some cases low sensitivity and selectivity, demands for more sensitive and less expensive methods are increasing in analytical chemistry. As a heavy metal lead, which produces several diseases, is one of the most important and widely distributed pollutants in the environment. Lead is one of the most toxic elements and has accumulative effects. It can occur as a result of industrialization in the production of pigments, anticorrosion coatings, lead smelters, alloys, and batteries, causing significant contaminations in air, dust, soil, water, sediments, food, etc. The World Health Organization (WHO) and the US Environmental Protection Agency (EPA) have set maximum levels of 50 and 10 $\mu$gL$^{-1}$ of Pb in seawater and drinking water, respectively. Due to these reasons, the determination of lead is becoming increasingly important. However, the number of reagents available for the spectrophotometric determination of lead is relatively small. The main reagents are dithiazone, diethyldithio-carbamate, 4-(2-pyridilazo) resorcinol, diphenylcarbazone, Arsenazo III, 2-(2-thiazolylazo)-p-cresol and porphyrin compounds. Although each chromogenic system has its advantages and disadvantages with respect to sensitivity, selectivity and rapidity due to using different chromogenic reagents, most of them require extraction using an organic solvent, surfactants or even fierce toxic cyanide as a masking agent to increase the sensitivity or selectivity.

Also, the determination of lead(II) has been widely done in various systems such as in human and artificial teeth blood, urine, cookies, baker’s yeast, ashes, coals, sediments, sludge, soils, freshwaters, drugs, human hair, wine, seawater and minerals, using different classical and instrumental techniques.

One of the classical noninstrumental methods of preconcentrating samples which we have applied in our study is flotation. It is a well known technique for selective separation of valuable substances from ores, minerals and etc. but today this technique is used mainly in other fields of chemical engineering and more rarely in analytical chemistry also. It is a process in which valuable minerals are separated from gangue minerals. It is a complex combination of various physical principles, such as surface chemistry, colloid chemistry, crystallography and physics. The exact manner is not well understood. Various factors influence the performance of a flotation unit including the bubble size, stator and rotor configuration, type and quantity of chemicals added and residence time.

Experimental

Materials and instrumentation

All the reagents were of analytical grade obtained from Merck, Germany. Double distilled fresh water was used in all the experiments. The standard solution of 1000 $\mu$gM$^{-1}$ Pb (II) was prepared by dissolving the appropriate amount of Pb (NO$_3$)$_2$ in H$_2$O. The required volumes of this solution were used to prepare the working solutions. A sodium tetraborate buffer solution with the concentration of 0.05 M (with addition of NaOH, 1M) was prepared by mixing nearly the appropriate amounts and adjusting to pH = 6 by the addition of a few drops of HCl solution. The solution of $3.23 \times 10^{-3}$ molL$^{-1}$ Alizarin yellow GG was made by dissolving of Alizarin yellow GG in the appropriate amount of freshwater.

The pure stable Cetyltrimethylammonium bromide (CTAB) solution (0.5 $\mu$gmL$^{-1}$) was prepared by dissolving the appropriate amount in H$_2$O. The 0.1M NaCl solution was prepared by dissolving the appropriate amount of the salt in water. Pure methanol and n-hexane were also used as solvent and extraction organic phase, respectively. A Shimadzu mini UV-Vis model 1245 spectrophotometric was used for all the absorbance measurements with a 10 mm quartz cell. A Jenway model 3505pH meter was used for pH measurements.

Experimental

To a 100 ml volumetric balloon, 1 ml of a solution containing up to 10 $\mu$gM$^{-1}$, and 12 ml
3.23 × 10⁻³ molL⁻¹ Alizarin yellow GG were added and its pH was adjusted to 6 by the addition of 3 mL of the buffer solution. 1 mL NaCl (0.1M) and 7 mL CTAB solution (0.5 µg/mL⁻¹) were also added and the volume reached to 100 ml with freshwater. The solution was maintained for 12 min to form complex and then transferred to a 100 ml separating funnel. 12 mL of n-hexane was added to it. The funnel was stoppered and vigorously shaken for 90 sec, and then left to rest for 5 minutes to give a perfect floated layer in the aqueous/organic interface. The floated layer was adsorbed on the inner walls of the funnel. By slowly opening the stopcock of the funnel, the aqueous phase was released in 5 minutes. To extract the lead content, 2 mL pure methanol was added to the funnel and vigorously shaken again for few minutes. The organic phase containing only the lead–alizarin complex was separated and its absorbance was measured at 364 nm against a reagent blank prepared in the same manner.

**Results and Discussion**

**Overall procedure**
The procedure included two steps as follow:

**Step 1:** Separation and pre-concentration of Pb (II) by its flotation as a complex with Alizarin

**Step 2:** Determination, extraction and detection by the spectrophotometric method with methanol.

**Influencing parameters**
To obtain reliable results, a number of parameters influencing the steps were optimized. Some of them that related to the flotation step including the type of the organic phase, pH of the solution, concentration of alizarin, surfactant, the electrolyte solution, floatation relaxation time, and complexation time were firstly investigated and reported. The others that related to the extraction and determination of Pb (II) content are discussed as the following. Since almost all the interfering agents were eliminated via the flotation process, no troublesome was observed on the performance of the extraction procedure. There was also no inherent interference due to the presence of alizarin. In this viewpoint, selection of a suitable reagent to extract the Pb (II) content gives emphasis on the following considerations.

1. The reagent should form a very stable complex with Pb (II).
2. In viewpoint to increase the sensitivity of the determination, the molar absorptivity coefficient of the extracted complex must be inherently much greater than the ion-associate which is in turn justifying the extraction step, evidently.
3. The extraction process of Pb (II) should be carried out using a high selectivity reagent to avoid any interfering element in the extraction process.
4. To assure on quantitative extraction of Pb (II) content in one step, the gain of the extraction process must be very high.

In order to find a favorable reagent for the extraction process, a number of conventional reagents were investigated. It was observed that some of them, such as Alizarin Red (s), methyl thymol blue and Xylenol orange were not able to extract Pb (II) completely in the presence of n-hexane. Alizarin yellow GG is one of the best extractants, which has been recognized as a sensitive reagent for the determination of Pb (II) in acidic media. It is capable to form primary and secondary complexes with Pb (II) in acidic and alkaline media, respectively. However, due to a higher absorptivity coefficient and solubility in polar organic phase such as methanol, the primary lead–alizarin chelate was preferred in the spectrophotometric determination. To enhance the selectivity of the extraction of the primary lead–alizarin, the process was carried out in acidic medium in the presence of excess alizarin.

**Optimization of reagent (Alizarin) concentration**
Using alizarin with a sufficient high concentration to extract all the Pb (II) content in one step was very important. The effect of alizarin concentration was studied over the concentration range of 1.29 × 10⁻⁴ to 5.17 × 10⁻⁴ molL⁻¹. The maximum absorbance occurs to alizarin concentrations above 3.23 × 10⁻⁴ molL⁻¹. Since, by increasing the concentration of Pb (II) ions, a greater amount of alizarin was required, a solution with 3.88 × 10⁻⁴ molL⁻¹ alizarin was chosen for further investigations (Fig. 1).

**Influence of pH**
The influence of pH and the volume of buffer were investigated in the range of between 1 to 9. As shown
in Figure 2 an optimum volume of borate buffer with pH = 6 has been obtained.

**Effect of surfactant concentration**

The volume of two surfactants (CTAB and CPC) with the concentration of 0.5 ppm was investigated in the range of between 0 to 10 mL. The optimum volume was 7 mL of CTAB (Fig. 3).

**Conformity with Beer’s Law and figures of merit**

Under the optimum conditions, a linear calibration curve was constructed for pb (II) determination over the range of $3.86 \times 10^{-8}$ to $8.20 \times 10^{-7}$ molL$^{-1}$ (8–170 ngmL$^{-1}$). The correlation coefficient was ($r = 0.998$), showing an acceptable linearity of the calibration curve. The apparent molar absorptivity at 364 nm was $1.33 \times 10^6$ molL$^{-1}$cm$^{-1}$ for a 100 mL aliquot of the extracted aqueous phase. The RSD obtained ($n = 10$) for $7.2 \times 10^{-7}$ molL$^{-1}$ (150 ngmL$^{-1}$) of Pb (II) was 4.36% and the detection limit, defined as the sample concentration giving a signal equal to the blank average signal (10 blank) plus three times the standard deviation of the blanks, was found to be $8.7 \times 10^{-9}$ molL$^{-1}$ (1.0 ngmL$^{-1}$).

**Effect of foreign ions**

Possible interference from various ions may be found in the complex matrix and examined by introducing them into aliquots of Pb (II) solutions with the concentration of $2.4 \times 10^{-7}$ molL$^{-1}$ (50 ngmL$^{-1}$). The tolerance limit was fixed at the maximum amount of an ion causing an error not greater than 5% in the absorbance of the extract (Table 1). Almost all of the cations and anions were tolerated at high ion/Pb (II) ratio except for Zn$^{2+}$, Cd$^{2+}$ and Al$^{3+}$ which exhibit proportionally lower tolerated limits. The interference of Cd$^{2+}$, Zn$^{2+}$

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**Figure 1.** The effect of alizarin concentration on determination of Pb (II). The pH of the solution was adjusted to 6, $C_{Pb} = 100$ ngmL$^{-1}$ in all examinations.

**Figure 2.** The effect of pH and volume of buffer on the determination of Pb (II) by the proposed method. The pH of the solution was adjusted to 6, $C_{Pb} = 100$ ngmL$^{-1}$, $C_{alizarin} = 3.88 \times 10^{-4}$ molL$^{-1}$ in all examinations.

**Figure 3.** The effect of surfactant (CTAB and CPC) concentration on determination of Pb (II). The pH of the solution was adjusted to 6, $C_{Pb} = 100$ ngmL$^{-1}$, $C_{alizarin} = 3.88 \times 10^{-4}$ molL$^{-1}$ in all examinations.

**Table 1.** Tolerance limits for diverse ions in $2.41 \times 10^{-7}$ mol L$^{-1}$ (50 ng mL$^{-1}$) Pb (II) in a 100 mL solution of seawater sample.

| Mole ratio of interfering ion to Pb (II) | Ions |
|----------------------------------------|------|
| 10000                                  | Na$^+$, K$^+$, NH$^+_4$, Ca$^{2+}$, Ba$^{2+}$, Co$^{3+}$, Cu$^{2+}$, Fe$^{2+}$, Mn$^{2+}$, Al$^{3+}$, Ni$^{2+}$, Cr$^{3+}$, Cu$^{2+}$, Fe$^{3+}$, F$^-$, Cl$^-$, Br$^-$, SCN$^-$, ClO$_4^-$, SO$_4^{2-}$, Zn$^{2+}$, CH$_3$COO$^-$, CO$_3^{2-}$, C$_2$O$_4^{2-}$, MoO$_4^{2-}$, Hg$^{2+}$, HPO$_4^{2-}$ |
| 1500                                   | Ag$^+$, Sr$^{2+}$, WO$_4^{2-}$, Mg$^{2+}$, Cd$^{2+}$ |
| 100                                    | Fe$^{3+}$, Th$^{4+}$, Bi$^{3+}$, Cr$^{3+}$ |
| 30                                     | Zn$^{2+}$, Cd$^{2+}$, Al$^{3+}$ |

**Note:** *Tolerated after masking with cyanide—in which 5 ml of 0.1 M solution was added to the solution before the flotation process.*
and Ag⁺ was attributed to the formation of a more stable complex with masking reagent, cyanideion. This interfering complex can be floated similarly at the critical concentrations and more. During the extraction of lead content the complex can also be extracted with alizarin in the same manner. The tolerance limit of Zn²⁺, Cd²⁺ and Cu²⁺ were sufficiently enhanced by the addition of 0.1 molL⁻¹ cyanide-ions as a masking agent, but Al³⁺ could be neither masked nor separated with none of the conventional agents. Fortunately, the concentration of Al³⁺ was very low in usual real samples and Al³⁺ was not a serious interferrent on the determination of Pb (II) in practical analysis of the real samples.

Applications to real samples
The proposed method was applied for the determination of Pb (II) in several water samples including a tap water and a synthetic laboratory sea water sample. The synthetic sea water composition was prepared based on the procedure of Arancibia et al.³ Along with the samples, several known amounts of Pb (II) were spiked to examine the reliability of the method. In treatment with the first two samples, aliquots of 100 ml were directly employed, and then the prepared solutions were subjected to the proposed method. The spiked amounts demonstrate that the proposed method exhibit a good reliability. Since the concentration of Cl⁻ is essentially very high in sea-water, to eliminate its interference in treatment with the flotation step, the iodide concentration was increased up to five times of the normal case. The slopes of the calibration graphs prepared for the water samples were found almost identical with that of the standard addition plot. Hence; the standard curve method with R.S.D. of 4.36% was used in the determination of various samples (Table 2).

| Test no. | Samples          | Lead added (ngmL⁻¹) | Measured (ngmL⁻¹) | Recovery (%) |
|----------|------------------|--------------------|------------------|--------------|
| This method (n = 7) | Tap water | 20.00 | 23.05 ± 0.14 | 115.3 |
| 1        | 30.00           | 28.73 ± 0.16 | 95.8             |
|          | 40.00           | 39.12 ± 0.23 | 97.8             |
| 2        | Sea water       | 40.00           | 38.11 ± 0.14 | 95.3 |
|          | 60.00           | 57.09 ± 0.19 | 95.1             |

Conclusions
The developed method offers a good sensitivity and selectivity for the determination of Pb (II) in the range of 8–170 ngmL⁻¹ in various environmental samples. The important feasibility of the method is its simplicity in treatment with the aqueous samples. Results were found not to be significantly different from standard addition. In comparison with the conventionally alizarin extraction method, the detection limit, sensitivity, and selectivity were enhanced considerably due to benefit a high preconcentration factor (100) utilizing the flotation process. The recovery yield and RSD of the measured data denoted that the proposed method can be satisfactorily applied to the determination of trace Pb (II) in real samples.

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Disclosure
This manuscript has been read and approved by all authors. This paper is unique and is not under consideration by any other publication and has not been published elsewhere. The authors and peer reviewers of this paper report no conflicts of interest. The authors confirm that they have permission to reproduce any copyrighted material.

References
1. Dessuy MB, Vale MGR, Souza AS, Ferreira S’LC, Welz B, Katskov DA. Method development for the determination of lead in wine using electrothermal atomic absorption spectrometry comparing platform and filter furnace atomizers and different chemical modifiers. Talanta. 2008;74:1321–9.
2. Chena J, Xiao S, Wu X, Fang K, Liu W. Determination of lead in water samples by graphite furnace atomic absorption spectrometry after cloud point extraction. Talanta. 2005;67:992–6.
3. Arancibia V, Nagles E, Comego S. Determination of lead in the presence of morin-5'–sulfonic acid and sodium dodecyl sulfate by adsorptive stripping voltammetry. Talanta. 2008. doi:10.1016/j.talanta.2009.06.046.
4. Li Z, Zhu Z, Chen Y, Hsu CG, Pan J. Spectrophotometric determination of lead in biological samples with dibromo-p-methyl-methylsulphonazo. Talanta. 1999;48:511–6.
5. Nakamura T, Kusata T, Matsumoto H, Sato J. Atomic absorption spectrometric determination of Cadmium and Lead in human and artificial teeth by direct atomization technique. Anal Biochem. 1995;226:256–62.
6. Yang CC, Kumar AS, Zen JM. Precise blood lead analysis using a combined internal standard and standard addition approach with disposable screen-printed electrodes, Anal Biochem. 2005;338:278–83.

7. Viksna A, Lindgren ES. Determination of lead and cadmium in whole blood of mothers and their babies. Anal Chim Acta. 1997;355:307–11.

8. Gmignberg, de Campos RC. Iridium as permanent modifier in the determination of lead in whole blood and urine by electrothermal atomic absorption spectrometry. Spectrochim Acta Part B. 2001;56:1831–43.

9. Parsons PJ, Slavin W. Electrothermal atomization atomic absorption spectrometry for the determination of lead in urine: results of an inter laboratory study. Spectrochim Acta Part B. 1999;54:853–64.

10. Aćar O, Košić E, rker ART. Determination of lead in cookies by electrothermal atomic absorption spectrometry with various chemical modifiers. Food Chem. 2000;71:117–22.

11. Skountzou P, Soupmiô M, Bekatorô A, et al. Lead(II) uptake during baker’s yeast production by aerobic fermentation of molasses. Proc Biochem. 2003;38:1479–82.

12. Lima ÉC, Brasil JL, Santos AHDP. Evaluation of Rh, Ir, Cu, W–Rh, W–Ir and W–Ru as permanent modifiers for the determination of lead in ashes, coals, sediments, sludges, soils, and freshwaters by electrothermal atomic absorption spectrometry. Anal Chim Acta. 2003;484:233–42.

13. Ahsan S, Kaneko S, Ohta K, et al. Electrothermal atomic absorption spectrometric determination of lead in calcium drug samples by direct atomization technique. Anal Chim Acta. 1998;362:279–84.

14. Kubovi J, Haniikovî V , Janos P. Extraction-spectrometric determination of lead and cadmium in human hair by atomic absorption spectrometric procedures after solid phase extraction. Anal Chim Acta. 1997;337:329–34.

15. Bispo MS, Korn Maô, Mortea ESô, Teixeira LSG. Determination of lead in seawater by inductively coupled plasma optical emission spectrometry after separation and preconcentration with cocrystallized naphthalene alizarin. Spectrochim Acta Part B. 2002;57:2175–80.

16. Zendelesovska D, Pavlovskâ G, Cundeva K, Stafilov T. Electrothermal atomic absorption spectrometric determination of cobalt, copper, lead and nickel traces in aragonite following flotation and extraction separation. Talanta. 2001;54:139–46.

17. Koksâal J, Synesk V, Janos P. Extraction-spectrometric determination of lead in high-purity aluminium salts. Talanta. 2002;58:325–30.

18. Yaman M. The improvement of sensitivity in lead and cadmium determinations using flame atomic absorption spectrometry. Anal Biochem. 2005;339:1–8.

19. Veseleva IA, Shekhovsotsna VN. Visual determination of lead (II) by inhibition of alkaline phosphatase immobilized on polyurethane foam. Anal Chim Acta. 2000;413:95–101.

20. Li J, Lu F, Umemura T, Tsunoda K. Determination of lead by hydride generation inductively coupled plasma mass spectrometry. Anal Chim Acta. 2006;543:65–72.

21. Shams E, Abdollahi H, Yektaehz M, Hajar R. H-point standard addition method in the analysis by differential pulse anodic stripping voltammetry Simultaneous determination of lead and tin. Talanta. 2004;63:359–64.

22. Erber D, Quick L, Winter F, Cammann K. Investigations for the hydride trapping within determination of lead by in situ a graphite electrothermal atomizer for routine analysis. Talanta. 1995;42:927–36.

23. Kuramochi M, Tomioka K, Fujinami M, Oguma K. Rapid determination of lead extracted by acetic acid from glazed ceramic surfaces by flow injection on-line preconcentration and spectrophotometric detection. Talanta. 2005;68:287–91.

24. Mousavi MF, Rahmani A, Golabi SM, Shamsipur M, Sharghi H. Differential pulse anodic stripping voltammetric determination of lead(II) with a 1,4-bis(prop-2-enyl)-1,4-phenylethylenediamine modified carbon paste electrode. Talanta. 2001;55:305–12.

25. Zenki M, Minamisawa K, Yokoyama T. Clean analytical methodology for the determination of lead with Arsenazo III by cyclic flow-injection analysis. Talanta. 2005;68:281–6.

26. Sp’ev a’ckova V, Sm’id J. Determination of lead in teeth of children for monitoring purposes by electrothermal atomic absorption spectrometry. Spectrochim Acta Part B. 1999;54:865–71.

27. Rubio J, Souza ML, Smith RW. Overview of flotation as a wastewater treatment technique. Mine Eng. 2002;15:139–55.

28. Morey MS, Grano SR, Ralston J, Prestidge CA, Verity B. The electrochemistry of Pb(II) activated fahlerite in relation to flotation. Minerals Engineering. 2001;14(9):1009–17.

29. Aki MA, Ismael DS, El-Asmy AA. Precipitate flotation-separation, speciation and hydride generation atomic absorption spectrometric determination of selenium(IV) in food stuffs. Microchem J. 2006;83:61–9.

30. Aslan N, Fidan R. Optimization of Pb flotation using statistical technique and quadratic programming. Separat Purif Tech. 2008;62:160–5.

31. Bakkali A, Cotra E, Berrueta LA, Gallo B. Study of the solid-phase extraction of diclofenac sodium, indomethacin and phenylbutazone for their analysis in human urine by liquid chromatography. Journal of Chromatography B: Biomedical Sciences and Applications. 1999;729:139–45.

32. Millers benz J, Hare LG, Frym M, et al. The use of hydrophilic lipophilic balanced (HLB) copolymer SPE cartridges for the extraction of diclofenac from small volume paediatric plasma samples. Journal of Pharmaceutical and Biomedical Analysis. 2001;25(5–6):871–9.

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