Nickel separation from human blood samples based on amine and amide functionalized magnetic graphene oxide nano structure by dispersive sonication micro solid phase extraction

Anne Trégouët*, Masoud Khaleghi Abbasabadib and Pooya Gholamic,\*  
\* Department of Chemistry, University Paris-Saclay, Saint-Aubin, Paris, France.  
\* Department of Chemistry, Iran University of Science and Technology, Tehran, Iran  
\* Nano Technology Center, Research Institute of Petroleum Industry (RIPI), P.O. Box 14665-1998, Tehran, Iran

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A B S T R A C T  
Nickel (Ni) is toxic effect on human body and must be determined in human blood samples. In this study, Ni ions separated and preconcentrated from blood samples based on magnetic Fe₃O₄-supported amine/amide-functionalized graphene oxide (Fe₃O₄@A/A-GO) nanoparticles by dispersive sonication micro solid phase extraction (DS-μ-SPE). By procedure, 10 mg of Fe₃O₄@A/A-GO was dispersed in 10 mL of human blood samples with sonication for 5.0 min and then separated from liquid phase with magnetic accessory. The Ni ions was extracted based on amine/amide covalence bonding of Fe₃O₄@A/A-GO sorbent (Ni---: NH₂). Then, the Ni ions back-extracted from Fe₃O₄@A/A-GO in low pH with nitric acid (0.2 mL, 0.3 M) which was diluted with DW up to 0.5 mL and finally, was determined by ET-AAS (peak area). The LOD, linear range (LR), enrichment factor (EF) and absorption capacity (AC) were obtained 35 ng L⁻¹, 0.15 - 7.2 μg L⁻¹, 19.8 and 131.6 mg g⁻¹, respectively. The method was validated by spiking samples.

1. Introduction  
Nickel (Ni) is one of the toxic compounds in water contamination and caused to acute and chorionic effect in human bodies[1]. Ni(II) are released into environment from waste of different chemical factories such as battery manufacturing, mining and electroplating. The air of near factories contain the significant amount of heavy metals such as nickel and caused to adverse effects in environment and humans[2]. Nickel is known to bind to specific proteins and/or amino acids in the blood serum and the placenta. Orally absorbed nickel is distributed to the kidneys, followed by the liver, brain, and heart. The harmful health effects of nickel lead to possible symptoms includes, chronic bronchitis, lung dysfunction, cancer in lung and nasal sinus[3]. Other target organs include the cardiovascular system, immune system, and the blood. In large doses (>0.5 g), some forms of nickel may be acutely toxic to humans when taken orally. Oral LD₅₀ values for rats range from 67-9000 mg Ni per kg (ATSDR).
Toxic effects of oral exposure to nickel usually involve the kidneys with some evidence from animal studies showing a possible developmental/reproductive toxicity effect (ATSDR). Normal range for Ni in healthy peoples is 0.2 µg L⁻¹ in serum and less than 3.0 µg L⁻¹ in human urine. A national health and nutritional examination survey (NHANES) of hair found mean nickel levels of 0.39 mg L⁻¹, with 10% of the population having levels less than 1.5 mg L⁻¹ [4, 5]. The various techniques were used for measurement of nickel (Ni²⁺) in human bodies such as serum and blood samples. Due to previous studies, the flame atomic absorption spectrometry (F-AAS) and electrothermal atomic absorption spectrometry (ET-AAS) were reported for analysis of heavy metals such as nickel (Ni) which was suitable for the determination in biological matrixes [6-9]. Occasionally, the atom trapping flame atomic absorption spectrometry (AT-FAAS) and fluorescence spectrometry (XRF) also was used for heavy metal determination such as nickel in biological samples [10, 11]. Also, different methods such as, polarography [12], inductively coupled plasma-optical emission spectrometry (ICP OES), inductively coupled plasma-mass spectrometry (ICP-MS) [13, 14], and spectrophotometry [15] were used for nickel analysis in human samples [16]. But as difficulty matrixes in human biological samples such as blood or serum, the sample preparation is required to separate nickel ions from liquid phases. The different sample preparation technology exist for nickel extraction from blood samples such as liquid–liquid extraction (LLE) [17], solid-phase extraction (SPE) and functionalized magnetic-SPE [18], dispersive solvent by liquid–liquid microextraction method (DLLME) [19], phase extraction based on cloud point (CPE) [20], dispersive solid phase microextraction (D-SPME) [21], ultrasound-assisted solid phase extraction (USA-SPE) [22]. Recently, the dispersive sorbent in the liquid phase was presented as micro SPE (D-µ-SPE) for separation/determination of nickel in water and biological samples [23]. Some advantages such as high extraction efficiency, simple usability, fast and low time caused to select as a favorite technique for metal extraction. The sorbent characterizations are a main factor which was effected on heavy metal extraction by the D-µ-SPE procedure. The selective of favorite nanosturactures improved recovery. The different nanosorbent such as, carbon nanotubes (CNTs), graphene / graphene oxide sheets (NG, NGO) and silica (MSN) were used for extraction of Ni in blood samples [24-26]. Between them, the NGO was reported as efficient sorbents for metal extraction due to their surface properties. In this study, a novel sorbent based on Fe₃O₄-supported amine/amide-functionalized graphene oxide (Fe₃O₄@A/A-GO) was used for separation/speciation Ni(II) from human blood samples by dispersive sonication micro solid phase extraction (DS-µ-SPE) at pH=8.0. The method was developed in blood samples by ET-AAS.

2. Experimental

2.1. Instrumental

The graphite furnace atomic absorption spectrophotometer (GF-AAS, 932 GBC, Aus) was used for nickel determination in blood samples with the Avanta software. The linear range of 3.0-150 µg L⁻¹ (peak Area, Abs=1.91) was selected for Ni in optimized light. The current and wavelength of HCL lamp was adjusted 3.0 mA and 228.8 nm, respectively. The auto-sampler of spectrophotometer (Pal 3000) was used for micro injection (1µL) of sample volumes to furnace tube after adjusted injector. The pH of the sample was digitally calculated by Metrohm pH meter (Swiss). Fourier transform infrared (FT-IR) spectra were recorded from KBr pellets using a Perkin Elmer Spectrum 65 FT-IR spectrophotometer. Powder X-ray diffraction (XRD) was conducted on a Panalytical X’Pert PRO X-ray diffractometer. Scanning electron microscopy (SEM) images were obtained using a Tescan Mira-3 Field Emission Gun Scanning Electron Microscope (SEM). Magnetic
property of the catalyst sample was measured with a vibrating sample magnetometer (VSM) model LBKFB, Meghnatis Daghigh Kavir Co, Iran, at room temperature.

2.2. Reagents

Chemicals including natural flake graphite (325 mesh, 99.95%), were purchased from Merck chemical company and used as received. The standard stock solutions (1000 mg L\(^{-1}\)) of Ni (II), acetone, acetate, HNO\(_3\), NaOH, HCl and other reagents were purchased from Merck (Darmstadt, Germany). Ultra-pure deionized water (DW) purchased from Milli-Q plus water purification system (USA). The standard and experimental solutions of Ni\(^{2+}\) (0.1, 0.2.0.5, 1.0, 3, 5.9 and 7.0 μg L\(^{-1}\)) were prepared daily by appropriate dilution of the stock solutions with DW. The pH of samples were used with appropriate buffer solutions including sodium acetate for pH 3.5–5.6, sodium phosphate for pH of 5.8–8.0, and ammonium chloride for pH 8–10. All the laboratory glassware and plastic tubes were cleaned by 5% (v/v) HNO\(_3\) for at least 12 h and then washed many times with DW and dried in oven prior to use. All reagents for synthesis of Fe\(_3\)O\(_4\)@A/A-GO prepared by RIPI Company.

2.3. Synthesis of graphene oxide (GO)

Graphite black powder was oxidized to GO following the modified Hummers method in the several steps [27-29]. Fore pre oxidation of graphite powder, 250 mL of H\(_2\)SO\(_4\) was added to 5 g of graphite powder and the resulting mixture was stirred for 24 h. After 24 h, 30 g of KMnO\(_4\) was added to the mixture stirring for 72 h at 50 °C. Next, a solution of 45 mL of H\(_2\)O\(_2\) (30%) in 400 mL of deionized water was added to the mixture after which the brown color of the mixture turned into bright yellow. The GO dark solution was centrifuged and washed with deionized water and 10% HCl solution, and dried at 65 °C.

2.4. Preparation of nanomagnetic Fe\(_3\)O\(_4\) supported Amine/Amide-functionalized graphene oxide (Fe\(_3\)O\(_4\)@A/A-GO)

2.4.1 Acyl-chlorination of graphene oxide

In the first step, thionyl chloride 60 mL was added GO (0.3 g) and stirred in at 70°C for 24 h. After acyl-chlorination reaction, the Acyl-chlorination of GO was washed with THF four times and the precipitated dried at 65°C [30].

2.4.1 Functionalization of graphene oxide with Ammonia

A total of 1 g of Ammonia was added to Acyl-chlorinated GO (0.3 g). Then, the mixture was refluxed for 72 h at 120°C under argon condition. Finally, the mixture reaction washed with DI water. A dark powder of Amine/Amide-functionalized graphene oxide was obtained [31].

2.4.2 Nano magnetization of Amine/Amide-functionalized graphene oxide

The Fe\(_3\)O\(_4\)-supported Amine/Amide-functionalized graphene oxide (Fe\(_3\)O\(_4\)@A/A-GO) nanoparticles were synthesized by co-precipitation of FeCl\(_2\)-4H\(_2\)O and FeCl\(_3\)-6H\(_2\)O, in the presence of Amine/Amide-grafted graphene oxide. First, a solution of FeCl\(_2\)-4H\(_2\)O and FeCl\(_3\)-6H\(_2\)O was prepared with a molar ratio of 2:1. The weight ratio of GO to FeCl\(_3\) in the nano composite was 1 per 20 (mGO: mFeCl\(_3\) = 1:20). Afterward, 10 mg of Amine/Amide-grafted graphene oxide in 15 mL of DI water was ultrasonicated for 20 min. 12.5 mL solution of FeCl\(_2\)-4H\(_2\)O (125 mg) and FeCl\(_3\)-6H\(_2\)O (200 mg) in deionized water (10 mL) was added to the reaction mixture at room temperature. In order to raise pH value to 11, an aqua solution of 30% ammonia was added in the reaction mixture at 70 °C. Then, the reaction mixture was allowed to cool to room temperature and Fe\(_3\)O\(_4\)@A/A-GO washed five times with DI H\(_2\)O dried at 70 °C [32].

2.5 Analytical Procedure

In proposed procedure, 10 mL of blood and
standard solutions was prepared by buffer solution. The whole blood samples diluted with DW (1:1) before procedure. By DS-μ-SPE procedure, the pH of the standard solution containing 0.2 - 7.0 μg L⁻¹ of nickel was adjusted up to 8 and then 0.01 g of Fe₃O₄@A/A-GO as adsorbent was added to samples. The standard / blood samples was shaked for 5.0 min at room temperature and Ni ions physically adsorbed on surface of Fe₃O₄@A/A-GO and chemically extracted by amine and amide covalence bonding at pH=8. Then, the Fe₃O₄@A/A-GO separated from liquid phase by magnetic accessory. Finally, the Ni ions back-extraction with nitric acid (0.2 M) from Fe₃O₄@A/A-GO and after dilution up to 0.5 mL with DW was determined by ET-AAS. In optimized conditions, the recovery of physically and chemically adsorption by Fe₃O₄@A/A-GO was almost obtained 25.3% and 97.8%, respectively. The extraction efficiency of proposed method based on Fe₃O₄@A/A-GO was calculated by equation 1. The C_a is the stock concentrations of nickel and C_b is the remain concentration of Ni(II) after procedure (n=10, Eq. 1).

Recovery of extraction=(C_a-C_b)/C_a×100 (Eq.1)

3. Results and Discussion

3.1. Characterization of the nano Fe₃O₄@A/A-GO

The Fe₃O₄-supported Amine/Amide-functionalized graphene oxide nanomagnetic (Fe₃O₄@A/A-GO) was synthesized according to the synthetic route shown in Figure 1. Firstly, graphite was oxidized to GO by the modified Hummers method [27-29]. Afterward, the GO was amination and amidation with Ammonia according to our previously reported procedure [30, 31]. Finally, the resulted Amine/Amide-functionalized graphene oxide (Fe₃O₄@A/A-GO) was nano-magnetized by co-precipitation of ferrous (Fe²⁺) and ferric (Fe³⁺) ions in the presence of A/A-GO to afford the target adsorbent Fe₃O₄@A/A-GO [32].

Figure 2 shows the FT-IR spectra of GO and Fe₃O₄@A/A-GO. The broad peak in the range between 2600-3500 cm⁻¹ in the IR spectra of these compounds is related to the (O-H stretching)
vibration of carboxylic and enolic functionalities [35]. The peaks at 3420, 1719, 1621, and 1060 cm\(^{-1}\) shown in the spectra of GO and Fe\(_3\)O\(_4\)@A/A-GO are ascribed to the (C-O stretching), (C=C stretching), (C=O stretching), and (O-H stretching) respectively [33, 34]. Also, the absorption bands at 3360, 3181, 1630, and 1234 cm\(^{-1}\) shown in the spectra of GO and amination GO are ascribed to the stretching bands \(\nu(\text{O-H})\), \(\nu(\text{N-H})\), \(\nu(\text{C}=\text{O})\), and \(\nu(\text{C-N})\) respectively. In the spectrum of Fe\(_3\)O\(_4\)@A/A-GO, the peaks observed at around 628 and 583 cm\(^{-1}\) are related to the Fe–O stretching vibration [37-39]. These results prove that the successful Amination and amidation of GO and synthesis of Fe\(_3\)O\(_4\)@A/A-GO.

The XRD patterns of GO, and Fe\(_3\)O\(_4\)@A/A-GO are demonstrated in Figure 3 (a,b). GO has the two main peaks at \(2\theta = 11.5^\circ\), 42.58\(^\circ\) are related to the diffraction planes of (002) and (100) respectively [40-42]. As shown in Figure 3 (b), the peaks at 2\(\theta = 24^\circ\) and 42.58\(^\circ\) It is evident from the XRD pattern of Fe\(_3\)O\(_4\)@A/A-GO that was proved the presence of GO [42]. The main peaks at 2\(\theta = 30.12, 35.45, 43.07, 56.97, 62.47\) in XRD pattern of Fe\(_3\)O\(_4\)@A/A-GO are in good accordance with the standard XRD data of magnetite Fe\(_3\)O\(_4\). The main diffraction peaks at
$2\theta^o = 62.47, 56.97, 43.07, 35.45, 30.12$ are related to the reflection planes of cubic spinel crystal structure of Fe$_3$O$_4$ at (440), (511), (400), (311), (220) respectively [43-50].

The SEM image of GO shows that graphene oxide nano sheets consists of randomly accumulated and wrinkled thin sheets (Fig. 4a). Also, the SEM image of Fe$_3$O$_4$@A/A-GO shows in Figure 4(b) that the average diameter of Fe$_3$O$_4$ nanoparticles was about 35.43 nm and indicate a regularly spherical morphology on A/A-GO [51-52].

The Magnetic behavior of the Fe$_3$O$_4$@A/A-GO was recorded using a VSM. As shown in the Figure 5, the saturation magnetization of Fe$_3$O$_4$@A/A-GO was found to be 42 emu g$^{-1}$. This amount of a saturation magnetization value, the magnetized nanocomposite is expected to have considerable paramagnetism to make it magnetically separable from reaction mixture [32, 48].

### 3.2. Optimization of extraction procedure

For efficient extraction of nickel ions in human blood samples, the main parameters must be studied. The effective features such as, pH, sample volume, amount of Fe$_3$O$_4$@A/A-GO, shaking time and interferences ions must be optimized. Chemical bonding was strongly depended on the pH solutions. By procedure, the effect of pH on extraction of nickel through the amine functional group of Fe$_3$O$_4$@A/A-GO was evaluated. For this purpose, the different pH values from 1 to 11 with nickel concentration of 0.2-7 μg L$^{-1}$ as LLOQ and ULOQ was examined according to DS-μ-SPE procedure. Obviously, the maximum of extraction efficiencies for Ni (II) ions based on Fe$_3$O$_4$@A/A-GO were obtained at pH range of 8.2, and then the recoveries were decreased by increasing or decreasing of pH (Fig. 7). In optimized conditions, the effect of sample volume of blood on nickel extraction was studied between 2 - 20 mL. The results showed, the optimum extraction was achieved for 12 mL of blood sample and 15 mL of standard samples, So, 10 mL of sample volume was used for further studies. Also, the amount of Fe$_3$O$_4$@A/A-GO for nickel extraction was evaluated by DS - μ-SPE procedure. Based on experimental results, 10 mg of Fe$_3$O$_4$@A/A-GO was selected as optimum point. The sonication time for extraction of Ni(II) was studied in optimized pH. Based on previous research, kind and size of adsorbents are the most important factors for extraction and sonication time. Therefore, the effect of sonication in blood samples was examined by DS-μ-SPE procedure. The results showed, the maximum efficiency was achieved at 5 min. The concentration of Interfering ions such as Na$^+$, K$^+$, Mg$^{2+}$, Ca$^{2+}$, Cu$^{2+}$, Zn$^{2+}$, Co$^{2+}$, Al$^{3+}$, Hg$^{2+}$, SO$_3^{2-}$, I$^-$, NO$_3^-$, Cl$^-$ and F$^-$ caused less than ± 5% deviation in the recovery of Ni(II) as the tolerance limit. So, the interfering ions has no effect on the recovery efficiencies of Ni(II) in blood samples.
efficiencies for Ni (II) ions based on Fe₃O₄@A/A-GO were obtained at pH range of 8.2, and then the recoveries were decreased by increasing or decreasing of pH (Fig. 6). In optimized conditions, the effect of sample volume of blood on nickel extraction was studied between 2-20 mL. The results showed, the optimum extraction was achieved for 12 mL of blood sample and 15 mL of standard samples, So, 10 mL of sample volume was used for further studies. Also, the amount of Fe₃O₄@A/A-GO for nickel extraction was evaluated by DS-μ-SPE procedure. Based on experimental results, 10 mg of Fe₃O₄@A/A-GO was selected as optimum point. The sonication time for extraction of Ni(II) was studied in optimized pH. Based on previous research, kind and size of adsorbents are the most important factors for extraction and sonication time. Therefore, the effect of sonication in blood samples was examined by DS-μ-SPE procedure. The results showed, the maximum efficiency was achieved at 5 min. The concentration of Interfering ions such as Na⁺, K⁺, Mg²⁺, Ca²⁺, Cu²⁺, Zn²⁺, Co³⁺, Al³⁺, Hg²⁺, SO₄²⁻, I⁻, NO₃⁻, Cl⁻ and F⁻ caused less than ± 5% deviation in the recovery of Ni(II) as the tolerance limit. So, the interfering ions has no effect on the recovery efficiencies of Ni(II) in blood samples.

3.3. Validation methodology

Many methods was used for validation of methodology by SPE [53-55]. The analytical results of the developed DS-μ-SPE procedure were shown method at optimum conditions (Table 1). After sample preparation, Ni concentration in human blood and standard samples was determined by ET-AAS. The Human blood, serum and plasma as a real sample was used for determination of Ni by DS-μ-SPE procedure. The results was verified by analyzing the spiked samples with standard concentration of Ni (II) in human samples (Table 2). Based on results, a high and favorite recovery was obtained by spiking samples which confirms the accuracy of results in difficulty matrix. The

![Fig. 6. The effect of pH on nickel extraction by DS-μ-SPE procedure](image)

| Table 1. Analytical results for Ni(II) extraction based on Fe₃O₄@A/A-GO by DS-μ-SPE |
|----------------|-----------------|----------------|-----------------|-----------------|
| Element    | SV  | LR  | R²   | LOD (n = 10) | RSD (%) | EF |
| Ni(II)     | 10  | 0.15-7.2 | 0.9997 | 0.035 | 2.8%  | 19.8 |

a sample volume (mL), b Linear rang (µg L⁻¹), c Limit of detection (µg L⁻¹), d Relative standard deviation, e enrichment factor(EF),
recoveries of spiked samples demonstrated that the results was satisfactory for Ni analysis by DS-μ-SPE. In order to validate the method, the extraction efficiency for intra-day and inter day analysis was evaluated by spiking samples (Table 3).

4. Conclusions

A fast and efficient method based on Fe₃O₄@A/A-GO as adsorbant was used for preconcentration, separation of trace Ni (II) in human blood samples by DS-μ-SPE procedure. The mechanism of extraction was achieved by interaction between negative charge (-) of amine group of Fe₃O₄@A/A-GO with positive charge (+) of nickel ions in favorite pH (Ni²⁺... NH₂). After extraction, the concentration of nickel was determined by ET-AAS technique. Finally, the developed method has low ion interference, simple usage with low LOD, favorite RSD(%) values and good LR with high recoveries for Ni extraction in blood samples (>95%). Therefore, the proposed method can be considered as applied techniques for Ni separation and determination in blood samples by DS- μ-SPE coupled to ET-AAS.

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| Sample | Added(μg L⁻¹) | * DS-μ-SPE (μg L⁻¹) | Recovery (%) |
|--------|---------------|---------------------|--------------|
| Blood  |               |                     |              |
| ---    |               |                     |              |
| 3.0    | 3.65 ± 0.18   | 3.48 ± 0.15         | 97.3         |
| Plasma |               |                     |              |
| 1.5    | 1.45 ± 0.06   | 1.57 ± 0.07         | 95.3         |
| Serum  |               |                     |              |
| 3.0    | 2.92 ± 0.14   | 2.86 ± 0.12         | 94.6         |

*Mean of three determinations of samples ± confidence interval (P = 0.95, n =10)

| Sample | Added(μg L⁻¹) | Found (μg L⁻¹) | Recovery (%) |
|--------|---------------|----------------|--------------|
| Blood  |               |                |              |
| ---    |               | 2.61 ± 0.13    | 98.4         |
| 2.5    | 5.07 ± 0.24   |                |              |
| Blood B| ---           | 1.73 ± 0.08    |              |
| 2.0    | 3.76 ± 0.18   | 101.5          |              |
| Serum C| ---           | 3.06 ± 0.15    |              |
| 3.0    | 5.98 ± 0.33   | 97.3           |              |
| Serum D| ---           | 2.72 ± 0.14    |              |
| 3.0    | 5.68 ± 0.28   | 98.6           |              |
| Plasma E| ---          | 1.46 ± 0.11    |              |
| 1.5    | 3.02 ± 0.16   | 104.6          |              |
| Plasma F| ---          | 0.25 ± 0.02    |              |
| 0.25   | 0.49 ± 0.03   | 96.0           |              |

*Mean of three determinations of samples ± confidence interval (P = 0.95, n =10)
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