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Highly sensitive and efficient voltammetric determination of ascorbic acid in food and pharmaceutical samples from aqueous solutions based on nanostructure carbon paste electrode as a sensor



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1. Introduction

Ascorbic acid is a naturally organic compound and as a water-soluble vitamin has been widely applied in large quantities to food products, drinks, animal feed, pharmaceutical formulations and cosmetics due to its valuable properties such as pH regulating and antioxidant features [1,2].

Ascorbic acid participates as a key component in biological metabolism such as reducing agent in various metabolic pathways, synthesis and maintenance of collagen, blood vessels, cartilage, bones and tendons and reacts with reactive oxygen species or free radicals and probably reducing cholesterol level [3,4]. Furthermore, ascorbic acid as a vital nutrient is commonly used in therapeutical fields such as improving immunity, preventing and healing of catarrh, infertility, skin disorders, amelioration of injuries and burns, cancer, aids and clinical diagnostic applications [5,6]. Besides, in food processing industries ascorbic acid is used as an antioxidant to prevent changes in color, taste and odor of products [7].

Therefore, due to biological importance of ascorbic acid, its accurate determination in pharmaceutical, clinical and food industries samples is

ABSTRACT

A square wave voltammetric method for the trace analysis of ascorbic acid was developed in this study. Carbon paste electrode was modified with NiO nanoparticle and 1-butyl-3-methylimidazolium tetrafluoroborate as a binder. Electro-oxidation behavior of ascorbic acid on the modified electrode was studied, which indicated that the nanostructure modified electrode could efficiently promote electrocatalytic oxidation of ascorbic acid. A fast, selective, high sensitive and simple electrochemical strategy was then developed for trace analysis of ascorbic acid using the constructed electrode. The catalytic oxidation signal exhibited a wide linear range from 0.08 to 380.0 µM toward the concentration of ascorbic acid with a sensitivity of 0.0158 µA/µM, and the limit of detection was as low as 0.04 µM. The suggested sensor was also used for quantitative determination of ascorbic acid in food and pharmaceutical samples.

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greatly demanded. Until now numerous techniques have been reported for quantitative determination of ascorbic acid including high performance liquid chromatography [8,9], spectrophotometry methods [10, 11], fluorometry methods [12,13], solid phase analysis [14,15] and chemiluminescence methods [16]. However, all mentioned methods have major disadvantages including high cost, laborious in sample preparation and time consuming, requiring large infrastructure back up and expert knowledge, background interference for fluorometry and destroying of sample for chromatography. Recently electrochemical techniques provide accurate and rapid tools with high sensitivity for routine and reliable determination of ascorbic acid in various matrices [17-20].

Among all chemically modified electrodes, carbon paste electrodes (CPEs) received high attention due to its ease of application and regeneration, cheapness, stable response and very low ohmic resistance. Modified CPEs overcome on large over potential required for oxidation of electroactive compounds by modification of electrode surface using nanostructure materials and high conductive binder to increase the conductivity of the electrode [21]. Nanostructure materials received considerable attention due to their distinctive and unique behavior with outstanding electrical, chemical, mechanical and structural properties that make them a very attractive material for large range of applications in pharmaceutical, biological and industrial procedure [22-25]. Metal nanoparticles have been used commonly in electrochemical techniques owing to its high catalytic activity in chemical reactions and high surface area for increasing current density [26]. Since metal

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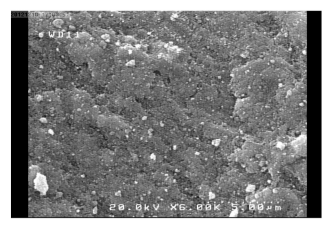


Fig. 1. SEM image of synthesized NiO nanoparticles.

nanoparticles revealed biocompatible properties, they could be applied in fabrication of electrochemical sensors to determine electroactive compound in pharmaceutical and food samples. Furthermore, using ionic liquids as a high conductive binder with specific characteristics such as good chemical and thermal stability, wide electrochemical windows and high ionic conductivity greatly benefited the fabricated electrode [27–29].

In the current work a great attempted have been done to fabricate a novel modified carbon ionic liquid paste electrode using NiO nanoparticles and 1-butyl-3-methylimidazolium tetrafluoroborate (BMITFB) as a binder. The electrochemical behavior of ascorbic acid was investigated and the obtained results demonstrate the advantage of BMITFB/NiO/NPs/CPE in compare to the bare carbon paste electrode in terms of higher sensitivity and reliability. The well fabricated BMITFB/NiO/NPs/CPE revealed an extraordinarily low background current, an extensive operating potential window, convenient modification, reproducibility, renewability and low cost. The analytical performance of the fabricated electrochemical sensor was evaluated by determination of ascorbic acid in liposome dope with ascorbic acid and food samples.

2. Materials and methods

2.1. Chemical and reagents

Ascorbic acid, NaOH, mineral oil, methanol and graphite powder were obtained from Merck. 1-butyl-3-methylimidazolium tetrafluoroborate was purchased from Sigma-Aldrich. Phosphate buffer solution (PBS) with various pH was prepared by mixing the stock solution of 0.1 M H_3PO_4 . The doubly distilled water was used in all solution preparations. All of the other chemicals were purchased in analytical grade from Merck.

2.2. Apparatus

Voltammetric measurements were performed on a Sama-500 electrochemical workstation (Isfahan, Iran). A three-electrode system was employed with a modified or unmodified carbon paste electrode as working electrode, a Pt wire as counter electrode, and a Ag/AgCl/KCl_{sat} electrode as reference electrode.

Microstructure and surface morphology of nanoparticles were identified by a scanning electron microscope (SEM, Philips).

2.3. Preparation of BMITB/NiO/NPs/CPE

BMITB/NiO/NPs/CPE was prepared by mixing 0.2 g of BMITB, 0.8 g of paraffin, 0.1 g of nanoparticles and 0.9 g of graphite powder. Then the mixture was mixed well for 1 h until a uniformly wetted paste was obtained. A portion of the paste was filled firmly into a glass tube as

described above to prepare BMITB/NiO/NPs/CPE. When necessary, a new surface of BMITB/NiO/NPs/CPE was obtained by pushing an excess of the paste out of the tube and polishing it on a white paper.

2.4. Real sample preparation

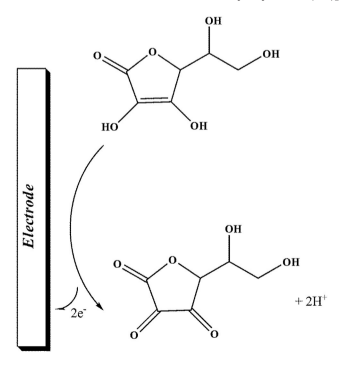
Vegetable and fruit juices were obtained using a mechanical squeezer. The juices obtained were filtered into a beaker and acidified (pH = 2) using citric acid. A 2.0-mL portion of the filtrate was added to the phosphate buffer solution pH = 7.0 in voltammetric cell. Standard addition method was used for determination of ascorbic acid in the juice samples.

Ten tablets of ascorbic acid powdered in mortar and then dissolved in 100 mL water with ultrasonication. Then, 1.0 mL of the solution plus 9.0 mL of the buffer (pH 7.0) was used for the analysis with standard addition method.

Liposomes containing ascorbic acid were prepared using film hydration method. Lecithine and cholesterol were dissolved in chloroform solvent with three different compositions of 70:30, 60:40 and 50:50, respectively. Rotary evaporator was used to evaporate the solvent, until an oily concentrated mixture was obtained and then known amount of ascorbic acid which is dissolved in di-water, was added to the mixture and for 30 min rotary evaporating process was continued until milky color suspensions containing liposomes were formed. To prevent the oxidation of Lecithine the procedure was carried out under inert atmosphere. To determine the amount of ascorbic acid loading in all formulations, the non-entrapped compounds were separated from encapsulated ones by centrifuge. The supernatant phase which contains the non-trapped ascorbic acid was separated. Standard addition method was used to determine the percentage of non-trapped ascorbic acid using proposed sensor.

2.5. Synthesis of NiO nanoparticles

To prepare the nanoparticles (NiO), 0.25 M aqueous solution of NiO(NO₃)₂· $6H_2O$ and a 0.5 M aqueous solution of NaOH was prepared in distilled water. The NiO(NO₃)₂· $6H_2O$ solutions were added drop wise (slowly for 3.0 h) to the above solution under high-speed stirring. The beaker was sealed at this condition for 3 h. The precipitated Ni(OH)₂



Scheme 1. Eelectrochemical mechanism for electro-oxidation of ascorbic acid.

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