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1 2	An Electrochemical Sensor for the Detection of P-Anisidine through Electrochemistry
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7 Abstract

8 In this study, a sensitive electrochemical voltammetry method for the analysis of p-anisidine

9 (pA) using a carbon paste electrochemical (CPE) modified with a porous material, such as

¹⁰ natural phosphate (NP), was proposed. p-anisidine strongly adsorbed on a electrode surface

and prepared electrode NP-CPE provides easy methods for electrochemical quantitative

¹² electro-p-anisidine. Operational parameters have been optimized, and performance

¹³ voltammetric stripping was investigated by cyclic voltammetry (CV). The current intensity

¹⁴ peaks are very linear, with good sensitivity NP- CPE.

15

Index terms — modified electrodes; cyclic voltametry; natural phosphate; moringa oleifera p-anisidine; carée
 wave voltammetry.

18 1 Introduction

19 adsorbed p-anisidine was assessed using cyclic voltammetric (CV) and square wave voltammetry (SQW).

20 **2 II.**

²¹ 3 Experimental a) Reagents

Potassium nitrate was dissolved into Bidistilled deionized water (BDW) to form 1mg.L -1 stock solutions. Working standards for calibration were prepared by diluting the primary stock solution with BDW. Carbon paste was supplied from (Carbon, Lorraine, ref. ??900, ??rench). All chemicals were of analytical grade and used without further purification.

²⁶ 4 b) Electrodes preparation

Firstly, the carbon-paste electrode was prepared according the following procedure [13]. The carbon-paste electrode was prepared by mixing the graphite powder with paraffin oil used as a binder.

The mixture was grinding in a mortar agate and then a portion of the resulting composite material was housed in PTFE cylinder. The geometric surface area of the working electrode was 0.1256cm 2. A bare of carbon vitreous inserted into carbon paste provided the electrical contact, and then the Phosphate natural film is electrodeposited onto carbon paste electrode. The deposit of Phosphate natural on carbon paste electrode

³³ surfaces was processed at 20 V. The current was maintained by a galvanostat with a function generator.

³⁴ 5 c) Apparatus

Electrochemical experiments were performed using a voltalab potentiostat (model PGSTAT 100, Eco Chemie
B.V., Ultrecht, The Netherlands) driven by the general purpose electrochemical systems data processing software
(voltalab master 4 software).

All the electrochemical experiments were performed in a standard one-compartment three reference electrode was SCE and the counter electrode was platinum. All electrode potentials were referred to this

 $_{40}$ reference electrode. The working electrode was natural phosphate modified carbon paste electrode (NP-CPE).

41 **6 III.**

42 7 Results and Discussion

⁴³ 8 a) Characterization of prepared electrodes

The surface structure of NP-CPE surface was observed using scanning electron microscopy (Fig. ??). The film layer of NP was formed on the surface of ara-anisidine (p-anisidine), is the most toxic [1] of the three isomers of anisidine and causes damage to the blood when ingested orally, by inhalation or skin contact. If heated strongly it can release highly toxic fumes of oxides Trogen lev- [2][3][4]. Melting and boiling point 243°C P-anisidine (PA) is colorless crystal with point 57.2 it is an important intermediate for the synthesis of dyes, medicine and fragrances, pigments, and other chemical compounds [5][6][7].

P-Anisidine reacts with secondary oxidation products such as aldehydes and ketones in the fats and oils to form products which absorb at 350 nm of light wavelength; therefore, it is used as an official method for detection them by the American Oil Company Chemistsy [8][9]. It is particularly good at detecting unsaturated aldehydes, which are those most likely to generations at unacceptable flavors, making it especially useful in food quality testing [10]. Traditional preparation of p-Anisidine uses iron powder or sodium sulfide as reducing agent [11], which reproduces a large quantity of waste and results in the problem of serious environmental pollution. [12]

56 In this article, we describe the electrochemical analysis of p-anisidine on a modified clay carbon paste electrode.

57 The electrochemical characterization of carbon paste electrode; it was not disintegrated or detached from the

 $_{\rm 58}$ $\,$ surface when immersed in the electrolytic solution (0.1M Na 2 SO 4). The treatment described previously gives

59 compact particle fractions between 100 and 400µm rich in phosphate. Natural phosphate treaty has the following

60 chemical composition: CaO (54.12%), P 2 O 5 (34.24%), F (3.37%), SiO 2 (2.42%), SO 3 (2.21%), CO 2 (1.13%),

⁶¹ Na 2 O (0.92%), MgO (0.68%), Al 2 O 3 (0.46%), Fe 2 O 3 (0.36%), K 2 O (0.04%) and order of several ppm ⁶² metals.

⁶³ 9 b) Voltammetric characteristic of p-anisidine

The determination of p-anisidine at the NP-CPE was performed by using cyclic (CV) and square wave 64 voltammetry (SQW). The results are presented in Figures 2 and 3 show the CV and the SQW measurements, 65 respectively. Two oxidation peaks were observed at NP-CPE towards the positive sweep direction, the first one 66 around ?0. 2 V and the second at approximately 0.15 V versus SCE, scanning in the negative sense brings up 67 a cathodic peak at -0.4V. Fig. 3 shows the square wave voltammograms obtained in 0.1 mol L ?1 Na 2 SO 68 4 for unmodified and modified carbon paste electrode. When the NP-CPE was dipped into the accumulation 69 medium containing p-anisidine followed by square wave voltammetry two well defined peaks appeared. Under 70 these selected conditions, the peak current increased linearly with the p-anisidine concentration in solution using 71 a preconcentration time of 10 min (Figs. 4 and 5). The linear dynamic range was comprised between 0.8 and 72 2.4 mM in terms of the relationship between p-anisidine concentration and the oxidation peak current (Fig. ??). 73 . The relationship can be described in the following linear regression equation in the mentioned concentration 74 range: IP1 = 0,105[P-A] + 0,438 R² = 0,98375

76 10 Global

77 11 Conclusion

A new chemically modified carbon paste electrode has been developed with natural phosphte for the determination
of p-anisidine at trace levels by square wave and cyclic voltammetry. The electrode offers attractive properties
such as simplicity of electrode preparation. Also, there is no leaching of the electrode because of the low solubility

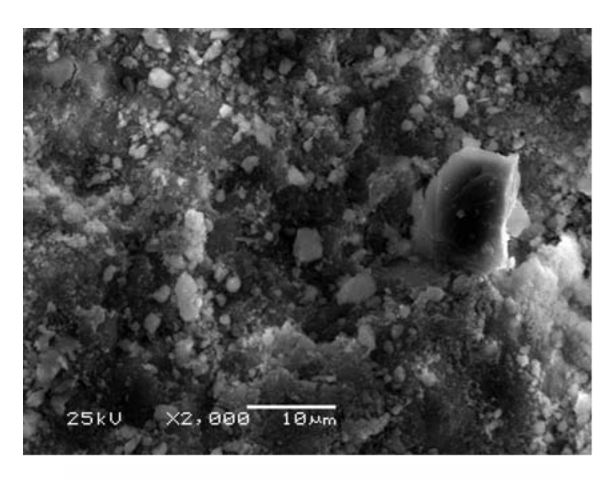
81 of the NP in aqueous solution.

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Figure 1:

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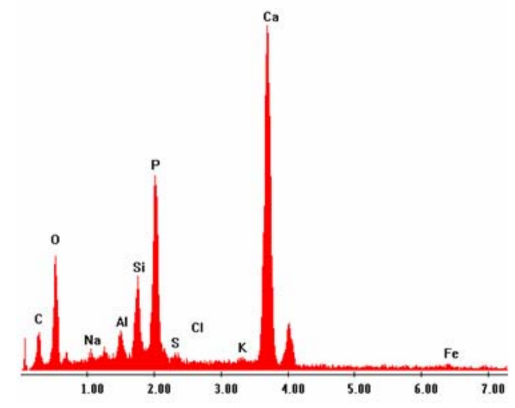


Figure 2: 2 2016 CFigure 1 :

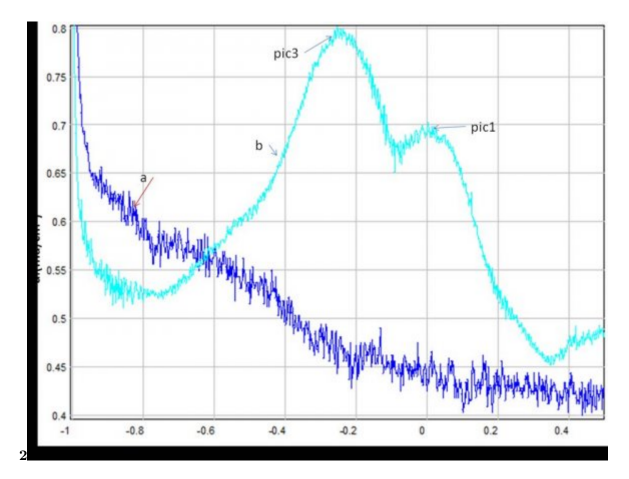


Figure 3: Figure 2 :

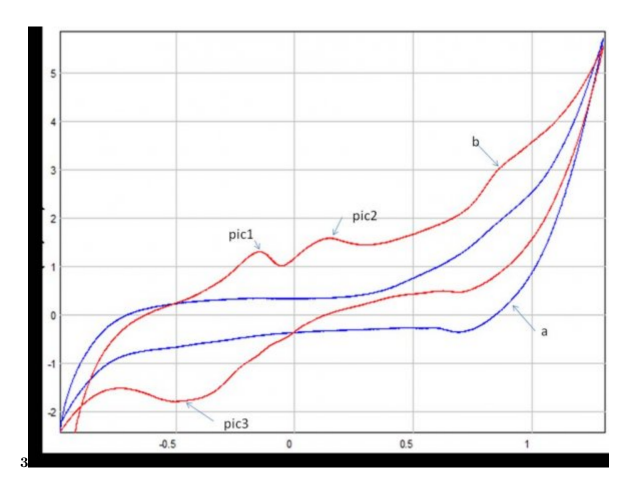


Figure 4: Figure 3 :

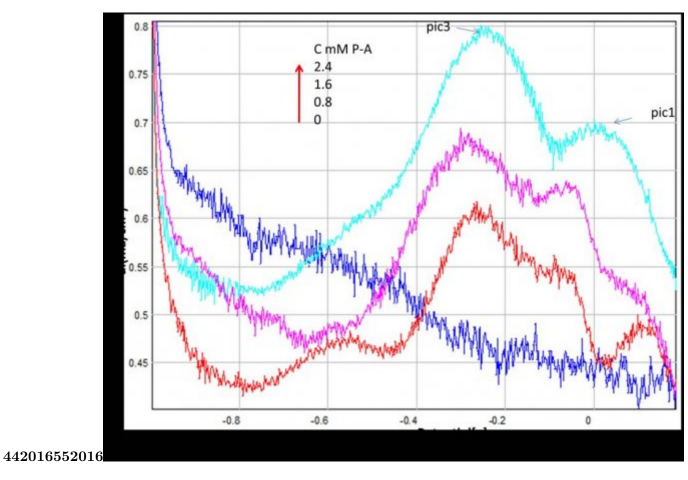


Figure 5: Figure 4 : 4 2016 CFigure 5 : 5 2016 C©

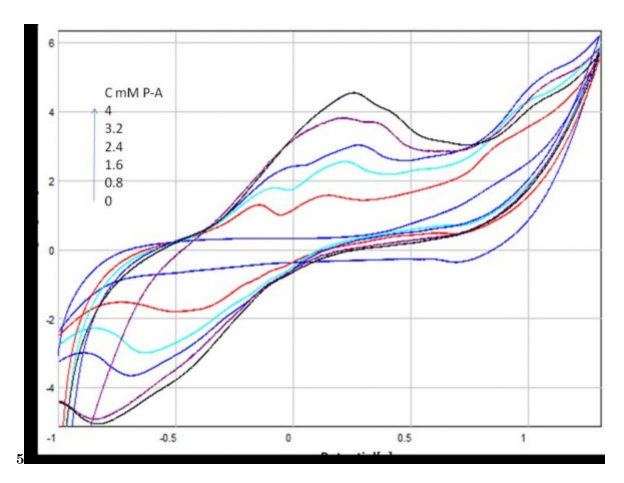


Figure 6: Figure 5 :

- 82 [Pei-Kun ()], Tang Pei-Kun. 2002. China: Chemical Industry Press.
- 83 [Liaoning Chemical Industry ()], Liaoning Chemical Industry 2002. 31 (6) p. .
- 84 [Fox ()] 'Determination of nitrite'. J B Fox . CRC Critical Reviews in Analytical Chemistry 1985. 15 p. .
- [Lijinsky and Epstein ()] 'Nitrosamines as environmental carcinogens'. W Lijinsky , S S Epstein . Nature 1970.
 225 p. .
- [Wang Gui-Lin and Wei ()] 'study of side reaction in the synthesis of p-anisidine'. Wang Gui-Lin , Wei . J]
 Zhejiang chemical industry 1996. (4) p. .
- [Wen-Nan ()] 'Synthesis of p anisidine'. Zhang Wen-Nan . Fine and Specialty Chemicals, 2001. 13 p. .
- ⁹⁰ [Zhang Jian-Hua ()] 'Synthesis of p-anisidine by liquid phase hydrogenation method'. Zhang Jian-Hua . J] China
 ⁹¹ Chlro-Alkali 2003. 5 p. .
- ⁹² [Shu-Qing and Chong ()] 'Technology study of preparation of p-anisidine from p-nitroanisole'. Wang Shu-Qing ,
 ⁹³ Gao Chong . J. Zhejiang Chemical Industry 1996. 27 (4) p. .
- ⁹⁴ [Usher ()] 'Telling; Analysis of nitrate and nitrite in foodstuffs'. C D Usher , GM . Journal of the Science of Food
 ⁹⁵ and Agriculture 1975. 26 p. .
- 96 [Walters ()] 'The exposure of humans to nitrite'. C L Walters . Oncology 1980. 37 p. .
- ⁹⁷ [Foreman and Goodhead ()] 'The formation and analysis of n-nitrosamines'. J K Foreman , K J Goodhead .
 ⁹⁸ Journal of the Science of Food and Agriculture 1975. 26 p. .
- ⁹⁹ [Valery Hambate Gomdje ()] 'Thérèse Rosie Lauriane Ngono, Salah Eddine El quoatli, Rachida Najih, Abdelilah
 ¹⁰⁰ Chtaini'. Valery Hambate Gomdje . Acta Technica Corviniensis 2013. 6 p. .
- 101 [Wolf and Wasserman ()] I A Wolf, A E Wasserman. Nitrates, nitrites, and nitrosamines, 1972. 177 p. .
- 102 [Ya-Jing] Zhu Rui-fen; Study on the Synthesis of p-methoxy aniline from p-nitrochlorobenzene, Zhang Ya-Jing.