Electrochemical Chelation of Lead by NDA Modified Carbon Paste Electrode
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Abstract
The analytical performance of developed DNA film screen printed biosensors designed for the detection of metals was evaluated. The modified surface exhibited an affinity to chelating metal ions in solution, forming complexes. Square wave voltammetry combined with a pre-concentrating process and standard additions were employed for trace analysis. Finally, cyclic voltammetry technique was used to characterize the developed DNA film surface behavior in presence of heavy metal.

Keywords: DNA; Modified electrode; Biosensor; SQW; Cyclic voltammetry

Introduction
A biosensor is defined by the international Union of Pure and Applied chemistry (IUPAC) as a self-contained integrated device that is capable of providing specific quantitative or semi-quantitative analytical information using a biological recognition element (bio chemical receptor), which is retained in contact direct spatial with a transduction element [1].

The DNA-based biosensor is a system that incorporates immobilized DNA as molecular recognition element in the biological active layer on the surface, and measures specific binding processes with DNA mainly using electrochemical, optical and piezoelectric transducers. The fact that the DNA sequences are unique to each organism means that any self-replicating biological organism can be distinguished. The DNA-based biosensor is also a complementary tool for the study of biomolecular interaction mechanisms of compounds with DNA, enabling the screening and evaluation of the effect caused to DNA by health hazardous compounds and oxidizing substances. There are hundreds of compounds which bind and interact with DNA, such as toxic organic molecules or heavy metals. Exposure to toxic chemicals is the cause of many human cancers; these carcinogens act by chemically damaging the DNA. Thus it is very important to explain the factors that determine affinity and selectivity in binding molecules to DNA, identify these chemicals and plan to minimize the impact of human exposure to these products.

Electrochemical techniques have the advantage in DNA biosensor design of having a rapid response time, being quantitative, sensitive, suitable for automation, cost effective, disposable, enabling in situ generation of reactive intermediates and detection of DNA damage and solving analytical problems in a wide range of contexts in order to be commercially viable. Comprehensive descriptions of research on DNA and DNA sensing [2-10] show the great possibilities of using electrochemical transduction in DNA diagnostics.

Carbon paste electrodes chemically modified are endowed with many good qualities, such as ease of handling and applicability to anodic oxidations [11]. Different modifiers for carbon paste have been reported in the last years for the electrochemical stripping analysis of heavy metals.

Recently, methods involving radical–based mechanisms have been developed for the modification of carbon surfaces with covalently attached layers of organic species. The deposition of mono or multilayers at the carbon paste surfaces serves the purpose of electrode.

Functionalization for analytical purposes, like electro analysis of heavy metals [12,13].

The aim of this investigation is to highlight the potential for using DNA systems for the fabrication of electrochemical biosensors. The prepared electrodes were used for the chemical preconcentration of different heavy metals at trace levels. The DNA modified carbon paste electrode (DNA-CPE) can preconcentrate heavy metals from aqueous solution to the surface of the modified CPE by forming complexes with these ions and greatly increasing the sensitivity of its determination. The experimental data show that some of the DNA-CPEs have high sensitivity, adequate selectivity and reproducibility, and a wide operative linear range of concentrations for the determination of heavy metals in water samples. The DNA-CPEs have the advantages of simple methods of synthesis and purification and their low solubility in water.

Experimental
Reagents and chemicals
All chemicals were of the highest quality. Graphite powder (spectroscopic grade RWB, Ringsdorf-Werke GmbH, Bonn-Bad Godesberg, Germany) was obtained from Aldrich and was used without further purification. PbSO4 was obtained from Merck chemicals. Deionised water was used to prepare all solution. DNA systems were extracted from plants.

Apparatus
Electrochemical experiments were performed using a voltalab potentiostat (model PGSTAT 100, Eco Chemie B. V., Utrecht, 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-electrode cell. The reference electrode was

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SCE and the counter electrode was platinum. All electrode potentials were referred to this reference electrode. The working electrode was DNA-CPE.

**Electrode preparation**

The carbon paste unmodified was prepared by adding paraffin oil to carbon powder. DNA-modified carbon paste electrodes (DNA-CPEs) were prepared by immobilizing the DNA system by soaking the preformed carbon paste electrode in a solution containing the DNA solution. The effect of the contact time of the CPE electrode with the DNA solution was studied.

**Results**

**Cyclic voltammetric (CV) experiments**

Preliminary CV experiments were performed to study the behavior at carbon paste electrode (CPE) and DNA-CPE.

The electrodes were first immersed in an electrochemical cell containing buffer solution (pH 7.2). Comparing CPE and DNA-CPE voltammograms (Figure 1), we find that the shape has changed which corresponds to a change of the surface morphology.

The electrochemical behavior with DNA-CPE prepared at low contact times illustrates the great difficulty for the transition electrons from the electrode surface. Cyclic voltammogram obtained with a carbon paste electrode did not show any redox peak. The deposit of the DNA onto CPE surface requires about 48 hours (Figure 1). The developed DNA films modified carbon paste electrodes were soaked, respectively, in solutions containing different concentrations of studied heavy metals for about 10 minutes, removed, then rinsed with pure water. The electrodes were then, respectively, transferred into the electrochemical cell only containing supporting electrolyte solution for cyclic voltammetry and square wave voltammetry. Figure 2 shows CVs for Pb (II) solutions obtained at different CPE modified with DNA system. The cathodic scan produces the reduced species of lead, which is deposited on the electrode. This deposition favors the oxidation process and hence the current peak of this signal (i.e., the anodic peak). The peak current is more important in the case of DNA-CPE electrode prepared with a contact time of 24 h. The oxidation peak of lead, complexed with DNA film and CPE surface, in the cathodic scan varies as follows:

DNA-CPE cont. time 24h ≥ DNA-CPE cont. time 18h ≥ DNA-CPE cont. time 5h ≥ CPE

1. The deposition of DNA on the carbon paste electrode surface improves its sensitivity;
2. The contact time is crucial in the process of filing the DNA.

Figure 4 show the SQWVs curves recorded in buffer solution containing different concentrations of lead, at DNA-CPE electrode (contact time 24 h). The peak current increases linearly with the concentration of Pb(II) (Figure 5).

The calculated standard deviation was used in the determination of the detection limit (DL, 3 × Std) and the quantification limit (QL, 10 × Std). From these values, the detection was 1.23.10-5 mol/L. This value remains correct compared to that found with a CPE electrode modified by organic molecules [14].

The Figure 6 shows the optical microscopic images, recorded respectively, for CPE (a) and DNA-CPE (b) electrode. We note that the DNA film forms a thin layer that covers the all carbon surface.

**Conclusion**

In conclusion, it was possible demonstrating the potentiality of the proposed electrodes for chelating heavy metals. Such a biosensor is characterized by a higher sensitivity and reproducibility than those of the unmodified carbon paste electrode. Electrochemical research on DNA is of great relevance to explain many biological mechanisms. The DNA-modified electrode is a very good model for simulating the
nucleic acid interaction with cell membranes, potential environmental carcinogenic compounds and to clarify the mechanisms of action of drugs used as heavy metals.

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