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Chromium (III) ion selective electrode based on di(benzylamino)glyoxime

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## ABSTRACT

A new poly/viny/chlorido) membrane sensor for CP<sup>2</sup> ions based on di (benzylamino)glyoxime as an ionophore was prepared. The electrode has a linear dynamic mage 1 to 1<sup>2</sup>/<sub>2</sub> to 1<sup>2</sup> mol 1<sup>2</sup>/<sub>2</sub> with a Nemstain slope of 20.3 ± 0.5 mV per decede and a detection limit of 2×10<sup>2</sup>. It has a first response time of <15 a and can be used for at least 3 months without any considerable divergence in potential. The proposed electrode revealed good sensitivity for Cq(III) over a wide variety of mental ions and could be used in a plf range of 2×5.5. The proposed sensor was used for the determination of CP<sup>2</sup> ions in real samples such as test leaves, coffee and cance powder by direct potentionerty.

Keywords: Ion-selective electrode: Chromium(III); Di(benzylamino)glyoxime: Poly(vinyl chloride) membrane

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### INTRODUCT

The increasing use of ion sensors in the fields of environmental, agricultural and medicinal analysis is forcing analytical chemies to develop new sensors for the fast, accurate, reproducible and selective determination of various species. Among heavy metals, trivalent metals have received less attention in spite of their widespread occurrence in food products, sea and fresh water, plants and animals [1].

A literature survey revealed that a large number of ISEs based on PVC membrane number of ISEs based on PVC membrane ware reported for many inorganic ions [2], but very little was published on PVC based trivalent ions. The first report on Cr(III) was in 1990. [3] in which a PVC based chromium-wire electrode was made by incorporating videous control of Cr(SCN), ion-pair studied, and the electrode gave a full response in the Cr(SCN), ion-contentation range of 10°-10° mol I° and has the Nernstian slop of 58 mV men rOC.

In 1987 a Cr(III) selective electrode with PVC membrane based on 8-quinlinechromium dithizonate was built that was a precipitate based selective electrode [5]. Singh et al. used a Cr(III)-selective electrode based on the macrocyclic compound for electrode, was prepared, based on 4methylaminoazobenzene with a linear concentration range of 1.7×10<sup>-6</sup>-1.0×10<sup>-2</sup> mol I and a limit of detection of 8.0×10<sup>-7</sup> mol I<sup>-1</sup> [7]. A Nernstian slope of 19.5 mV per decade and pH range of 3.0-5.5 were reported for this electrode. The selectivity coefficient of some metal ions such as Ag(I) and Mn(II) are small enough to be considered (about 10°1) for PVC-based Cr(III)-selective electrode based on glyoxal bis(2-hydroxyanil) has which has a linear range of 3.0×10° -1.0×10° mol 1°, with a Nemstian slope of 19.8±0.5 mV [8]. However these sensors [3-8] suffer from the disadvantages of significant interferences by some cations, deviation from Nemstian behavior, small linear range and narrow pH range.

In this paper, we describe the construction and evaluation of a plasticized PVC membrane chromium(III) - selective electrode based on di (benzylamino) glyoxime (DBAG) as neutral carrier (Fig. 1).

Fig. 1. Structural representation of the DBAG used as ionophore in the chromium(III)-selective electrode.

### Experimental

## 1. Reagents and materials

Reagent grade of dibuthyl phthalate (DBP), acetophenon (AP), dioctyl phthalate (DOP), tetrahydrofuran (THF), potassium tertakis(p-chlorophenylborate) (KTpCIPB), oleie acidi(DA), high relative molecular weight PVC and nitrate or chloride salts of all cations obtained (from Merck or Fluks) were used as received. The ionophore DBAG was synthesized. Double distilled deionized water used

#### 2. EMF measurements

Potentials were measured with a Corning ion analyzer pH/m meer against a double junction saturated calome leterrode (SCE) at constant temperature (25-6). The chamben of the SCE was filled with a potassium choloride solution. A sitversitive chloride electrode containing 3 M solution of KCI was used as the internal reference electrode. The electrode cell assembly of the following type was used:

Ag-AgC1 | KCI (3 M) | internal solution.  $1\times10^{13}$  M  $Cr(NO_3)_3$  | PVC membrane | test solution | Hg-Hg<sub>2</sub>Cl<sub>3</sub>,KCI(saturated),

## 3. Electrode preparation

The general procedure to prepare the PVC membrane was similar to our previous work 191. First, we mixed thoroughly 5 mg of Impophore DBAG, 30 mg of ryndered PVC. and 65 mg of plastisizer AP in a 2 cm diameter glass dish. The mixture was then dissolved in 5 ml of THF. The solvent was evaporated slowly until an oily concentrated mixture was obtained. A Pyrex tube was dipped into the mixture for ~10 s so that a membrane was formed. The tube was then pulled out of the The tube was filled with internal filling solution (1.0×10-1 mol I1 CnNO-h). The electrode was finally conditioned for 24 h in a 1.0×10°3 mol 1°1 solution of Cr(NO<sub>3</sub>). A allver/silver chloride electrode was used as an Internal reference electrode.

## Results and discussion

1. Effect of membrane composition

The DBAG as a carrier was found to be highly responsive to Cr(III) with respect to several other metal ions. Therefore, we studied in detail the performance of the plasticized PVC membrane containing this ionophore for

In order to test the performance of the nembrane, various operation parameters response neluding selectivity. ensitivity, life time, linear range, the nfluence of pH and the membrane Omposition on the response of the electrode vere investigated.

Optimization of the membrane ingredients

Number Composition(%) — (±0.5 mV per decade) (mol 1<sup>-1</sup>) PVC Plasticizer Ionophore Additive 60(AP) 65(AP) 1×10°-1×10° 5(KTpCIPB) 30

depend on the nature of ionophore used, but significantly on the membrane and additives used [9.11-13].

Thus, the influences of the membrane composition, the nature and amount of plasticizers on the potential response of the summarized in Table 1.

It is seen that, among the different incorporating 30% PVC, 65% AP ionophore DBAG shows the best sensitivity over a wide concentration range of 1×10%the presence of additive in the membrane slightly decreases the linear range of the

#### 2. Calibration curve and statistical data

The Potential responses of various ionselective membranes based on DBAG are cations tested, the largest sensitivity was obtained for Cr3+ ion. The e.m.f. vs. -log a<sub>Cr3+</sub> of the PVC membrane based on DBAG. a rectilinear range from 1×10-6-1×10-1 M. The slope of the calibrations curve was 20.3±0.5 mV per decade of Cr3+ concentration. The limit of detection, as determined from the intersection of the two extrapolated segments of the calibration graph, was 7.7×10<sup>-7</sup> M.

The standard deviation of sixteen replicate measurements is ±0.5 mV. The prepared electrode could be used for more than 3 months without any measurable change in potentials.

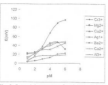


Fig. 2. Potential response of various metal ionselective electrodes based on DBAG. Composition of membrane: membrane ingredients, 30%PVC, 65%AP, 5% DBAG; internal solution, 1,0×10<sup>3</sup>mol 1<sup>3</sup> Cr(NO<sub>3</sub>)s.

#### 3. Influence of pl

The pH dependence of the potentials of the proposed electron for different concentrations of chromium(III) ion were tested over the pH range 1-7 and the results are depicted in Fig. 3. The pH was adjusted with nitric acid or sodium hydroxide solution. As I can be seen, the potential response remains almost constant the potential response remains almost constant the working pH range of the electrode. At pH higher than 5.5, the potential of electrode decreases due to the formation of chromium hydroxides in solution. The variation of

potentials in pH<2 could be related to protonation of the ligand DBAG in the membrane phase, which results in a loss of its ability to complex with Cr(III) ions, and probably response of the electrode to H<sup>\*</sup> ions in high acidic media.

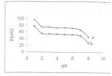


Fig. 3. Influence of pH of test solution on the potential response of Cr(III) electrode at different Cr<sup>3+</sup> concentrations: (a)1.0×10<sup>-2</sup> M and (b)1.0×10<sup>-3</sup> M.

#### 3.4. Selectivity of the electrode

The influence of interfering ions on the response behavior of ion-selective membrane electrodes is usually described in terms of selectivity coefficients,  $K_{c}^{PR}$ . The methods based on the Nicolsky-Eisenman equation for the determination of potentiometric selectivity coefficients (e.g. the fixed interference method and the mixed solution method) are among the most commonly methods used thus far [13-17].

Table 2

Selectivity coefficients (  $\kappa_{LB}^{SSM}$  ) of various interfering cations obtained using SSM.

[Cr3"]=1×10"3 M, [M8"]=1×10"3 M, pH=3

Cation	K 4,8	Cation	K 4.8
Na	1.2×10 <sup>-3</sup>	Zn <sup>2-</sup>	1.6×10 <sup>-3</sup>
Mg <sup>2</sup>	2.4×10 <sup>-3</sup>	Ni2-	7.1×10 <sup>-1</sup>
Al <sup>3</sup> - Ca <sup>2</sup> -	1.8×10-5	Li	1.8×10°
Ca <sup>2</sup>	3.2×10 <sup>-3</sup>	Ag	2.7×10 <sup>-3</sup>
CH	4.9×10 <sup>-3</sup>	Co.	5.2×10 <sup>-3</sup>
K° Pb²-	1.7×10 <sup>-3</sup>	Ba <sup>2-</sup>	3.3×10 <sup>-3</sup>
Pb2-	4.6×10 <sup>-1</sup>	Cd2-	6.7×10 <sup>-3</sup>

However, it has been shown that these methods suffer some limitations in terms of values for ions of unequal charges, non-Nemstian behavior of interfering ions, and activity dependence, of values. Thus, in this work, the recommended separate solution method (SSM) was used for selectivity measurements [18-24].

According to the SSM, the potential of a cell comprising an ions-selective electrode and a reference electrode is measured with two separates solutions, one containing the ion A at the activity at, four to  $\rm D$  ), the other one containing the ion B at the same activity  $\rm a_{\rm c}^{\rm min} \rm b_{\rm c}$  (but no A). If the measured values are E, and E\_B, respectively, the value of  $K_{\rm c}^{\rm min}$  is scalable to fix the according to

log 
$$K_{A,B}^{Pot} = (E_B - \overline{E}_A) + (1 - \frac{z_A}{z_B}) \lg a_A$$
  
which is equivalent to  $K_{A,B}^{Pot} = a_A^{-1} (1 + \frac{z_A}{z_B}) \lg a_A$ 

good detection limit. The electrode is characterized by a fast response, reproducibility, a reasonable long-term stability and low cost. The electrode also has a life time of more than three months. The proposed sensor has successfully been used as an indicator electrode to determine Cr(III) ion in some real samples.

#### 5. Application

The electrode has been used successfully for determination of Cr<sup>3+</sup> in various real samples. The experimental conditions employed and the resulting values are given in Table 2. A perusal of selectivity coefficient data presented in Table 2 indicates that the electrode is sufficiently selective over a large number of cations except Po<sup>2+</sup> and N<sup>3+</sup>.

The results (Table 3) indicate a very good agreement between the two values (obtained by AAS and the proposed sensor)

Table 3.

Results of determination of chromium in different real samples.

Sample	Average chromium concentration (ppm)		
	Proposed sensor	AAS	
Tea leaves	1.1±0.1	1.2±0.1	
Coffee	1.1±0.2	1.3±0.1	
Cacao powder	0.9±0.2	1.0±0.2	

#### 4. Conclusion

The results obtained in the present work demonstrate that the DBAG as a inonphore can be used in the development of a PVC-based ion-selective electrode. The electrode response to the Cr<sup>3+</sup> ions in a Nernstian fashion, and presents good selectivity and a

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