CHARACTERIZATION OF TRACE METAL SPECIES AND MEASUREMENT OF TRACE METAL STABILITY CONSTANTS BY ELECTROCHEMICAL TECHNIQUES

R. Ernst,* H. E. Allen† and K. H. Mancy

Environmental Chemistry Laboratory, Department of Environmental and Industrial Health, School of Public Health, The University of Michigan, Ann Arbor, MI 48104, U.S.A.

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Abstract—Differential pulse polarography and differential pulse anodic stripping voltammetry were used to determine copper, lead, cadmium and zinc complexes of relevance to environmental conditions. The dependency of peak current on pH and alkalinity was interpreted in terms of the variation in the metal species present. Stability constants of copper and lead complexes with carbonate were determined from the magnitude of the shift of peak potential using the method of Lingane. Inclusion of the transfer coefficient, calculated from the peak half-width for electrochemically irreversible systems, was used to determine the stability constant. By using both electroanalytical techniques, the investigator validates the correctness of his results without needing to rely on prior determinations of the stability constant for verification.

INTRODUCTION

Present awareness of the importance of trace metals in aquatic systems has prompted increased activity in both the development of trace analytical techniques and the study of the physicochemical behavior of low levels of metals. Although the majority of water quality standards specify, and environmental studies have measured only the total concentration of a trace metal, it has become apparent that the chemical form of the metal must be known to permit an accurate interpretation of both its biological effects and geochemical reactions. Prediction of trace metal environmental levels, forms, and transformations requires the development of mechanistic models which are based on equilibrium constants of metal complexes.

Methods for the determination of stability constants have been reviewed by Martell and Calvin (1952), Rossotti and Rossotti (1961) and Beck (1970). These books deal with many analytical procedures and their underlying theoretical bases. However, the examples used are almost exclusively simple systems generally containing a single metal and a single well-characterized ligand, both present at levels much higher than those in the environment. Application of these techniques to the study of environmentally significant complexation reactions has been carried out primarily by soil chemists concerned with metal–humic acid and metal–fulvic acid reactions. The techniques which have received the most use are potentiometric titration, Job’s continuous variation technique and ion-exchange equilibrium (Schnitzer and Khan, 1972; Stevenson and Arakani, 1972). Recent work by Cheng, Patterson and Minear (1975) has utilized the ion-exchange equilibrium method for determining the stability constant of various metals complexed with the soluble portion of sewage. Other investigators (Bradford, 1973; Stumm and Bilinski, 1973) have employed anodic stripping voltammetry for the determination of the stability constants of hydroxide and carbonate complexes of zinc and lead. These investigations were conducted to ascertain those complexes which predominate under environmental conditions.

We have investigated the use of both differential pulse polarography (DPP) and differential pulse anodic stripping voltammetry (DPASV) for the assessment of trace metal complexation. The application of these techniques to environmental measurements is presented here while the electrochemical basis for these results will be discussed elsewhere (Ernst, Allen and Mancy, 1974).

POLAROGRAPHIC DETERMINATION OF STABILITY CONSTANTS

Although polarography has been widely used for the quantitative determination of metals in the aquatic system, utilization of this technique for the determination of stability constants of these metals with naturally occurring ligands has been extremely limited. The detection limit for normal polarography does not permit analysis at the level at which metals normally occur in the environment. If samples are concentrated, shifts in chemical equilibria will result. Newer electroanalytical techniques such as pulse polarography and anodic stripping voltammetry have

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* Present address: Department of Chemistry, University of Lausanne, Lausanne, Switzerland.
† Present address: Department of Environmental Engineering, Illinois Institute of Technology, Chicago, IL 60616, U.S.A.
‡ To whom correspondence should be addressed.
sufficiently low detection limits that direct measurement of environmentally significant levels of trace metals is feasible.

The application of DPP and DPASV to the measurement of trace metals in natural waters has been discussed by Osteryoung and Osteryoung (1972) and Siegerman and O'Don (1972), respectively. For differential pulse measurements the potential waveform shown in Fig. 1(a) is applied to the electrode. Application of the pulse causes a large increase in the flow of both capacitive and faradaic currents. Since the capacitive current decays more rapidly than the faradaic current, the measured current at the end of the pulse will be almost exclusively faradaic [Fig. 1(b)]. It is due to this virtual elimination of the capacitive current that pulse techniques are able to achieve detection limits approximately two orders of magnitude lower than by the corresponding d.c. technique. In addition, the peak current ($i_p$) and the peak potential ($E_p$) can be easily determined from the nearly symmetrical peaks obtained by DPP or DPASV [Fig. 1(e)], which will facilitate measurement.

Early workers observed that the d.c. polarographic half-wave potential ($E_{1/2}$) could vary as the medium was altered in chemical composition. Heyrovsky and Ilkovic (1935) explained such shifts as being a result of complexation. Lingane (1941) was the first to quantitatively treat the phenomena. For the reaction:

$$M^{n+} + jL \rightleftharpoons ML_j^{n+}$$

having the formation constant

$$\beta_j = \frac{[ML_j^{n+}]}{[M^{n+}][L]}$$

he developed the following relationship between the shift in potential and the equilibrium ligand concentration:

$$\Delta E_{1/2} = (E_{1/2}, k) - (E_{1/2}, l) = \frac{2.303RT}{nF} \log \beta_j + \frac{2.303RT}{nF} \log[L]$$

where $(E_{1/2}, k)$ is the half-wave potential of the simple, uncomplexed metal ion, $M^{n+}$; $(E_{1/2}, l)$ is the half-wave potential of the metal in the presence of the ligand concentration $[L]$. $n$ is the number of electrons exchanged in the reduction process; and $(2.303RT/nF)$ is a constant having the value of 0.0591 at 25°C.

The following assumptions have been made in the derivation of the Lingane equation.

(1) The diffusion coefficient of the complex must be the same as that of the simple metal ion.
(2) The complex which is formed must be electrochemically labile.
(3) The ligand concentration at the electrode surface must be equal to the bulk concentration.
(4) The reduction of the metal must be electrochemically reversible.
(5) A single complex must predominate over a wide range of ligand concentrations.

These conditions have been discussed in detail by Crow (1969) and Heyrovsky and Kuta (1965).

Frequently, the condition of having a predominant complex is not fulfilled. Consequently, when the shift in half-wave potential is plotted versus the logarithm of the ligand concentration, a curve results rather than the straight line predicted by the Lingane equation. The method of DeFord and Hume (1951) has been used to interpret these curves as a summation of overlapped Lingane relationships and allows the individual formation constants to be ascertained.

Many ligand–metal systems are not electrochemically reversible and therefore are not amenable to treatment by the Lingane method. However, these irreversible systems may be of considerable practical importance and it is, therefore, important to have methods capable of studying them.
The most common method used for irreversible systems is the indicator method introduced by Schwarzenbach and co-workers (Ackerman and Schwarzenbach, 1952; Schwarzenbach, Gut and Anderegg, 1954). A labile complex of the ligand with another metal is required as an indicator. It must be reversibly reduced at a more positive potential than the half-wave potential of the metal under investigation and the two complexes must have similar stability constants. The stability constant can be determined from comparison of the current measured for the reversibly reduced metal, first in the absence of ligand, then in the presence of both the ligand and the irreversibly reduced metal. As many environmentally significant ligands would be expected to form irreversibly reduced complexes with most metals, this procedure would be unusable.

The shift in half-wave potential for an indicator complex has been used by Ringbom and Eriksson (1953) and Eriksson (1953) to estimate the stability constant of irreversible systems. However, the necessity of the rapid reaction of the reduced metal with the ligand and the need for chemical equilibrium to be established almost instantaneously are very strong limitations of the technique (Crow and Westwood, 1965).

Subrahmanya (1960) used a modified version of the Lingane equation to account for the irreversibility of the electrode process. This treatment which is based on the work of Tamamushi and Tanaka (1949) results in the following equation for an irreversible reduction process:

$$\Delta E_{1/2} = (E_{1/2})_L - (E_{1/2})_k = \frac{2.303RT}{nF} \log \beta_j + \frac{2.303RT}{nF} \log [L]$$

(4)

where the transfer coefficient \( \alpha \) is the fraction of the total applied potential which favors the forward reaction.

Since the slope of the d.c. polarographic wave is a measure of the reversibility of the process, measurement of the wave at \( \frac{1}{4} \) and \( \frac{3}{4} \) of its maximum value permits the determination of the value of \( \alpha \) from the relationship

$$E_{3/4} - E_{1/4} = -\frac{0.0563}{n}$$

(5)

For DPP the peak half-width was shown by Parry and Oldham (1968) to be dependent on both the amplitude of the applied pulse and the reversibility of the reduction. When using small values for the pulse amplitude the peak half-width \( (W_{1/2}) \) is given by the equation:

$$W_{1/2} = 3.52 \frac{RT}{nF}$$

(6)

from which the value of \( \alpha \) can be evaluated.

**EXPERIMENTAL**

A Princeton Applied Research polarograph Model 174 was used in the differential pulse mode. A Houston Instrument Omnigraphic Model 2000 recorder was used for recording the polarograms. For DPASV a PAR Model 172 Drop Timer was used to dislodge the mercury drops. All potentials were measured at 25°C vs a saturated calomel electrode.

The alkalinity of samples was fixed at \( 10^{-2}, 10^{-2}, \) or \( 10^{-1} \) equiv l \(^{-1} \) by adding the required amount of NaHCO\(_3\). The pH could then be controlled and adjusted as desired by equilibrating the sample with the appropriate partial pressure of CO\(_2\). The carbonate concentration at each alkalinity and pH was calculated using the equations presented by Roberts and Allen (1972).

Stock solutions, \( 10^{-2} \) M in metal, were prepared by dissolving the required amount of reagent-grade metal nitrate and adjusting to pH 3. All samples were prepared with respect to the trace metal and, in addition to the NaHCO\(_3\), 0.1 moles l \(^{-1} \) KNO\(_3\) was added as a non-reactive supporting electrolyte.

For experiments with organic ligands, in addition to the previously named reagents, the required amount of \( 10^{-3} \) M glycine or 500 mg l \(^{-1} \) humic acid (K & K Laboratories) was added.

Solutions were deaerated for 30 min prior to their initial measurement. Measurements were repeated to ensure that equilibrium had been attained. After further addition of ligand or lowering of the pH by increasing the partial pressure of CO\(_2\), generally only a short time was required to reattain equilibrium.

For DPASV the solution was mixed with a magnetic stirrer during the electrodeposition. The deposition was conducted with the instrument in the differential pulse mode so that a voltage excursion + 25 mV from the deposition potential occurred once each second. The deposition potential was 200–300 mV more cathodic than the peak potential so that the effect of the 25 mV pulses on the amount of metal deposited was small. The deposition potentials were -200 mV for Cu\(^{2+}\), -800 mV for Pb\(^{2+}\), -750 mV for Cd\(^{2+}\) and -1200 mV for Zn\(^{2+}\). Samples were electrolyzed for 100 s while the solution was stirred and for 20 s more under quiescent conditions. In the stripping step a +25 mV pulse was superimposed once each second on a voltage ramp which increased at 0.1 or 2 mVs \(^{-1} \). In the case of glycine and humic acid solutions for both DPP and DPASV, the voltage scan rate of 0.1 mV s \(^{-1} \) was used to permit measurement of the half-peak width and the peak potential with increased precision.

The same conditions, but with both the sweep rate and the pulse being cathodic, were used for DPP.

![Fig. 2. Variation of DPP current as a function of pH and alkalinity for 2.5 × 10^{-6} M copper. Current has been normalized to that for Cu\(^{2+}\).](image_url)
Fig. 3. Variation of DPASV current as a function of pH and alkalinity for 2.5 x 10^{-8} M copper. Current has been normalized to that for Cu^{2+}.

DEPENDENCE OF $i_p$ ON pH AND ALKALINITY

A pH-dependent model for the chemical speciation of copper, zinc, cadmium and lead in seawater has been developed by Zirino and Yamamoto (1972). They considered the ligands CO$_3^{2-}$, HCO$_3^-$, SO$_4^{2-}$, Cl$^-$, and OH$^-$ at the levels occurring in seawater. Within the pH range 7-9, the hydroxy and carbanato complexes and the free metals were the most important species except in the case of cadmium, where the high level of chloride in the seawater results in chloride complexes being dominant over the entire pH range considered. The increase in ASV signal resulting from lowering the pH of seawater from 8 to 5.5 was reported in support of the model.

To provide pertinent information regarding the effect of ligands on ASV, we have compared the electrodeposition step as measured by DPP to both the electrodeposition and stripping steps as measured by DPASV. Detailed experimental investigation in which both pH and carbonate have been independently varied is not available and, therefore, interpretation of ASV signals from environmental samples in terms of species rather than labile and non-labile components is not possible (Mancy, 1972).

For DPP measurements of copper, lead, cadmium and zinc solutions a maximum signal was obtained at low pH values where the hydrated metal ion was the dominant species. At higher pH values, the signal decreased but this decrease was less in solutions of high alkalinity. Typical results for the effect of pH and alkalinity on DPP are shown in Fig. 2. The decrease in signal at higher pH indicates the formation of insoluble species such as Cu(OH)$_2$, Zn(OH)$_2$, and PbCO$_3$. Higher levels of carbonate appear to favor metal solubility due to formation of ion pairs such as CuCO$_3^-$ and PbCO$_3^-$, which favorably compete with solid species for the metals.

DPASV curves of $i_p$ vs pH were similar to those obtained for DPP. For example, the results for copper which are plotted in Fig. 3, exhibit a similar dependency of $i_p$ on pH. However, for zinc and lead this dependency is greater for DPASV than for DPP. If there were no difference between DPASV and DPP data, this would indicate that the cathodic reaction is the controlling step. Since results by DPASV and DPP differ, the signal obtained by the DPASV measurement is dependent upon both the plating and the stripping steps. Present research in this laboratory (Twork, 1974) is directed toward the elucidation of the dependency of $i_p$ upon the chemical composition of the medium used in the stripping step.

Only qualitative interpretations were based on $i_p$ vs pH diagrams. Absolute measurements of current are complicated by slow precipitation and adsorption of dissolved or colloidal materials onto the cell. The slowness of the reaction at high pH and low alkalinity is indicated in Fig. 4.

Copper

Analysis of copper curves (Figs. 2 and 3) indicates that both hydroxide and the carbonate system are important in regulating the species of copper present. The importance of carbonate ions is shown most clearly in Fig. 2, where three independent, non-supersposed curves were obtained for the three alkalinities. At any pH, the [CO$_3^{2-}$] is almost directly proportional to the total alkalinity. If only carbonate were important parallel curves displaced on the pH axis would be expected in Figs. 2 and 3. The fact that the DPP curves are not parallel indicates the contribution of the hydroxide ion in regulating copper speciation. This conclusion supports the equilibrium model of Zirino and Yamamoto (1972) who calculate that Cu$^{2+}$, CuCO$_3^-$ and Cu(OH)$_2$ are the dominant copper species in solution. Based on previous work in this laboratory (Mancy and Allen, 1974) an equilibrium distribution diagram for copper was calculated (Fig. 5), and carbonate and hydroxide are both important.
The plots of \( i_p \) vs pH for the lead system, shown in Fig. 6, are similar to those for copper. However, the three lead curves tend to be more parallel than those for copper shown in Fig. 2. We can infer, at least qualitatively, that carbonate is much more important to lead speciation than is hydroxide. For a given pH, high alkalinity is associated with high current value which is an indication of increased solubility. Increased alkalinity results in decreased diminution of signal due to lead precipitation, which cannot be \( \text{PbCO}_3 \). This conclusion contradicts the model for lead solubility in natural waters proposed by Hem and Durum (1973). They indicated that while \( \text{Pb(OH)}_2 \) was the solid phase present at high pH, solid \( \text{PbCO}_3 \) was present at lower pH. Their equilibrium solubility calculations in the pH range 5-9 show lower soluble lead concentrations at higher total carbonate levels. An explanation for the discrepancy between our experimental data and their calculation might be the omission of the \( \text{PbCO}_3 \) ion pair from the model. In the model proposed by Zirino and Yamamoto (1972), the \( \text{PbCO}_3 \) ion pair was predicted to be the dominant soluble species over the pH range considered. Recent work by Bilinski and Stumm (1973) supports the model of Zirino and Yamamoto.

**Zinc**

Zinc curves, for the three values of alkalinity, are almost superimposed and are strongly pH dependent above pH 7.5. This is shown in Fig. 7. The decrease in signal is assumed to be due to the precipitation of zinc hydroxide. In the pH range between 8 and 9, the peak potential shifts towards more cathodic values. At pH greater than 9 a second peak appears and its signal increases with increasing pH. This second peak, which occurs at a potential more cathodic than the first, results from the direct electrochemical reduction of a non-identified zinc complex at the electrode surface.

**Cadmium**

Cadmium undergoes minimal complexation or precipitation in the media tested. There was less than 10% decrease in peak current at any pH or alkalinity investigated. The results from the model of Zirino and Yamamoto (1972) are not directly comparable with our experimental results since chloride, which was included in their model, forms strong cadmium complexes. In this model, neglecting the chloro-complexes, the hydrated cadmium ion predominates until...
Fig. 7. Variation of DPP current as a function of pH and alkalinity for $2.5 \times 10^{-6}$ M zinc. Current has been normalized to that for Zn$^{2+}$.

pH 9, at which point the concentration of the ion pair CdCO$_3^2$ equals that of Cd$^{2+}$.

DEPENDENCE OF $E_p$ ON INORGANIC LIGAND CONCENTRATION

The data obtained from DPP and DPASV were further analyzed to explain the observed electrochemical results. When initially recording the voltammograms, it was obvious that the peak potential varied as the pH was adjusted. This effect was pronounced at higher values of pH while at low pH values the potential was invariant. In addition, the peak potential at high values of pH was different for the various alkalinities.

Lead

When the peak potential was plotted vs the pH, a regular pattern, such as that obtained for the lead carbonate system (Fig. 8), could be observed. The aforementioned effects appeared to be both regular and systematic. Since different curves were obtained for the different alkalinities, it was apparent that the shift in peak potential was not due solely to hydroxyl ion, but must also have been dependent upon carbonate.

The data for DPP presented in Fig. 8 have been replotted in Fig. 9 to show the peak potential as a function of carbonate concentration rather than as a function of pH. The linear dependence indicates that a single carbonate complex of lead predominates from pH 6.5 to 9.0 for all alkalinities studied. For DPASV a linear regression with the same slope as DPP is obtained but the intercept of the line is different. The difference is in agreement with prediction (Ernst et al., 1974).

Acid solutions, which contained no bicarbonate, were analyzed by DPP and DPASV to determine the value of the peak potential for the free metal ion. These values which were $-375$ and $-405$ mV, respectively, were used in determining the shift in peak potential, $\Delta E_p$, resulting from complexation. The Lingane equation was used to calculate the number of ligands and the stability constant of the complex.

To determine the stability constant for the lead-carbonate system, a line was fitted by inspection to the data shown in Fig. 9. At low carbonate concentrations insufficient ligand was present to cause appreciable complex formation. Thus, the slope in this region approaches zero. At high concentrations the points again deviate from the line, but this is due to an increase in slope indicating that a higher order complex becomes predominant. The slope and intercept of the line shown in Fig. 9 were computed by a least squares linear regression.

Both DPP and DPASV produced similar results for the ligand number as well as the stability constant (Table 1). The ratio of carbonate to lead was determined to be 1.07 from the DPP data and 1.06 from the DPASV data. Values for the logarithm of the stability constant were 6.2 by DPP ($\pm 0.2$ S.E.) and 6.3 by DPASV ($\pm 0.4$ S.E.).

The method of Lingane is applicable only to reversible, diffusion controlled systems (Crow, 1969). If the system were not reversible, the same results could be
Measurement of trace metal stability

Table 1. Summary of results for stability constant determinations by DPP and DPASV

<table>
<thead>
<tr>
<th>Metal</th>
<th>Ligand</th>
<th>Technique</th>
<th>Number of observations</th>
<th>Transfer coefficient*</th>
<th>Ligand number*</th>
<th>Logarithm of stability constant*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pb</td>
<td>CO$_3^-$</td>
<td>DPP</td>
<td>18</td>
<td>Reversible</td>
<td>1.07 ± 0.05</td>
<td>6.2 ± 0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>DPASV</td>
<td>14</td>
<td>Reversible</td>
<td>1.06 ± 0.10</td>
<td>6.3 ± 0.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>DPP</td>
<td>19</td>
<td>Reversible</td>
<td>0.93 ± 0.04</td>
<td>5.7 ± 0.2</td>
</tr>
<tr>
<td>Cu</td>
<td>CO$_3^-$</td>
<td>DPP</td>
<td>10</td>
<td>Reversible</td>
<td>0.99 ± 0.04</td>
<td>6.1 ± 0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>DPASV</td>
<td>12</td>
<td>Reversible</td>
<td>1.63 ± 0.09</td>
<td>8.1 ± 0.3</td>
</tr>
<tr>
<td>Cu</td>
<td>CO$_3^-$</td>
<td>DPP</td>
<td>8</td>
<td>Reversible</td>
<td>1.55 ± 0.07</td>
<td>8.2 ± 0.2</td>
</tr>
<tr>
<td>Cu</td>
<td>Glycine</td>
<td>DPASV</td>
<td>7</td>
<td>0.93</td>
<td>1.28 ± 0.04</td>
<td>6.4 ± 0.1</td>
</tr>
<tr>
<td>Cu</td>
<td>Humic acid</td>
<td>DPASV</td>
<td>5</td>
<td>0.50</td>
<td>1.69 ± 0.04</td>
<td>10.2 ± 0.2</td>
</tr>
<tr>
<td>Pb</td>
<td>Humic acid</td>
<td>DPASV</td>
<td>6</td>
<td>0.76</td>
<td>1.14 ± 0.06</td>
<td>7.8 ± 0.4</td>
</tr>
<tr>
<td>Pb</td>
<td>Humic acid</td>
<td>DPP</td>
<td>6</td>
<td>0.61</td>
<td>0.66 ± 0.03</td>
<td>4.0 ± 0.1</td>
</tr>
<tr>
<td>Pb</td>
<td>Humic acid</td>
<td>DPASV</td>
<td>8</td>
<td>0.76</td>
<td>2.48 ± 0.11</td>
<td>14.8 ± 0.5</td>
</tr>
<tr>
<td>Pb</td>
<td>Humic acid</td>
<td>DPP</td>
<td>4</td>
<td>0.61</td>
<td>1.08 ± 0.01</td>
<td>6.1 ± 0.1</td>
</tr>
</tbody>
</table>

* The error in the estimation of the transfer coefficient was not estimated. This error would be propagated into the error for both $j$ and $\beta$. Stability constants and ligand numbers for glycine and humic acid complexes are for pH 6.8.

† The half-peak width varied and, therefore, the transfer coefficient was not a constant. No estimates of $j$ or $\beta$ were made.

obtained by DPP and DPASV using the Lingane equation only if both the anodic and cathodic processes were irreversible to the same extent—clearly an extremely unlikely circumstance. When the same result is calculated by both DPP and DPASV it must be assumed that not only is the value of the equilibrium constant correct, but also the electrochemical processes and complexation are rapid. Furthermore, DPP peaks approximated the theoretical half-peak width of 45.5 mV for a reversible reaction (Parry and Osteryoung, 1965), which further substantiates the reversibility of the reactions.

In applying the Lingane equation to the determination of stability constants by ASV, Bradford (1973) noted the requirement that the reaction must be rapid. Because his experimental value was in good agreement with published results, he concluded both that the reaction was fast and that his value for the stability constant was correct. In the present study no such comparison with literature data is necessary due to the previously discussed ability to validate results by comparing DPP to DPASV and by measuring half-peak width. Such a capability becomes most important when a stability constant for the complex being studied has not been reported previously.

In developing their pH dependent model for the chemical speciation of copper, zinc, cadmium and lead in seawater, Zirino and Yamamoto (1972) could find no stability constants reported for the lead carbonate ion pair. Based on the method of Garrels and Christ (1965) of estimating stability constants from solubility products and electronegativities, they estimated the logarithm of the stability constant for PbCO$_3$ to be 7.5. In a preliminary research report, Bilinski and Stumm (1973) reported 6.4 as the logarithm of the stability constant for the ion pair. This was based on ASV measurements, but with no confirmation of the result or test that the reaction was fast. The present measurements of the stability constant therefore validate that of Bilinski and Stumm.

Copper

Data for the variation of copper peak potential with pH and total alkalinity showed a similar pattern for that of lead (Fig. 8). A strong dependence on pH with separate curves obtained for each of the alkalinites investigated was apparent. Therefore the potentials for both the DPP and DPASV copper peaks were plotted (Fig. 10) as a function of the carbonate ion concentration. Two separate linear segments are discernable for both the DPP and DPASV data.

The two linear portions for each analytical method were analyzed by least squares regression analysis. The linear portion extending from approximately $10^{-5}$-10$^{-4}$ M carbonate ion would be included in the pH range 7.3–9.4 for a water having a total alkalinity of 0.001 equiv l$^{-1}$. Thus, this complex might be expected to be quite significant for many natural water systems since many natural waters have alkalinites and pH ranges similar to this. The complex was calculated to have 0.99 carbonate per copper by DPASV measurement and 0.93 carbonate per copper by DPP measurement (Table 1). The standard errors for both measurements were 0.04. The logarithm of the stability constant and its standard error was calculated as 6.1 ± 0.2 for DPASV and 5.7 ± 0.2 for DPP.

It should be noted that these values for the stability constant are significantly lower than the value of 6.77 for CuCO$_3$ determined by Silman (1958) and reported
It is important to note that, irrespective of the exact nature of the complex, it may be assumed that DPP and DPASV can be used to determine its stability constant since the present study has demonstrated that the same value is obtained for anodic as well as cathodic processes.

Cadmium and zinc

As previously noted, variation in the alkalinity had a smaller effect on both the cadmium and zinc analyses than it did in the copper and lead analyses. In addition, the pH was shown to influence the cadmium system to a much smaller extent than any of the others. In addition to these observations being true for the previously discussed variations of peak current with pH, they also help explain the variation of \( E_p \) with pH and alkalinity.

In the case of cadmium, a virtually invariant \( E_p \) was obtained irrespective of pH or alkalinity. Such behavior indicates that neither hydroxy nor carbonate complexes of cadmium have any appreciable existence within the range of conditions investigated.

For zinc a large potential shift occurred at the highest values of pH investigated for the \( 10^{-2} \text{ equiv L}^{-1} \) total alkalinity solution. This cathodic shift, which amounted to 150 mV when the pH was raised from 8.05 to 9.7, had the largest change above pH 9 which was the same region in which the peak current was found to increase. Although the complex was not identified, it must contain carbonate since the shift occurred only at the highest alkalinity investigated.

DEPENDENCE OF \( E_p \) ON ORGANIC LIGAND CONCENTRATION

Glycine

Metal glycinate stability constants were determined both by DPP and DPASV. In both cases the medium was 0.1 M in NaHCO\(_3\) and was maintained at pH 6.8. Although a smaller fraction of the glycine is capable of complexation at this pH than at higher pH, this pH simulates natural water conditions more closely than does a quite basic value. Similarly a solution containing carbonate was more realistic than one from which all other ligands were excluded.

Samples containing lead, cadmium or zinc at \( 2.5 \times 10^{-6} \text{ M} \) showed no shift in peak potential even when the total glycine concentration was as high as \( 4.8 \times 10^{-4} \text{ M} \). This is an indication that the stability of the metal hydroxide or carbonate complexes are greater than that of the metal glycine complexes under the experimental conditions. Under these conditions no complexation with glycine takes place; glycine is ineffective as a ligand. Although stability constants could not be computed for these complexes, this result is instructive in illustrating behavior which might occur in the investigation of ligands in environmental matrices.

A frequent problem is to predict the behavior of a constituent added to the environment. Concern has been expressed relative to the effect of adding NTA,
citrate or other replacements for phosphate in deter-
gents. Even when the stability constants for the com-
plexes of interest are known, our lack of knowledge
regarding levels of naturally occurring ligands and
their stability constants with the metals under consi-
deration prevents prediction of the effect of such addi-
tions. We are even less able to predict the effect of
additions of ligands such as algal metabolites or sew-
age whose structures are uncharacterized and whose
stability constants have not been measured. By deter-
mining whether a peak shift occurs on addition of
the ligand of interest, a rapid assessment of whether
the added ligand significantly complexes the trace
metal can be made.

The copper peak for both DPP and DPASV did
shift when glycine was added to the bicarbonate-con-
taining sample. This indicates that the copper–gly-
cinate complex is more stable than is the copper car-
bonate complex under these conditions. Besides shift-
ing the peak potential, addition of glycine caused a
decrease in the peak current. The decrease was ap-
proximately 10%, indicating that the copper–glycinate
complex is labile and that its diffusion coefficient is
similar in magnitude to both the inorganic complex
of copper initially present and to that for the hydrated
copper ion.

Although plots of $E_p$ vs log [glycine] are linear
for both DPP and DPASV, the two lines are not
parallel (Fig. 11). This can be best seen by comparison
of the curves in Fig. 11 to those in Fig. 10. In the
absence of ligand the DPP peak potential is anodic
to that for DPASV; in the presence of glycine, the
DPP peak is cathodic to the DPASV peak. Therefore,
the two lines must have crossed.

For both the DPP and the DPASV data sets, the
peaks are wider in the presence of glycine than in its
absence. The broadened peaks are a result of the
irreversibility of the electrochemical reaction. The
ratio of the peak width for a reversible reaction to
that for the irreversible reaction was used as the mea-
sure of the transfer coefficient $\alpha$ (Ernst et al.,
1974). Since the peaks had the same half-width at all glycine
concentrations, the degree of electrochemical reversi-

Humic acid

Addition of humic acid to copper, lead or zinc solu-
tions resulted in a decreased DPP or DPASV signal.
At high concentrations of humic acid the signal was
reduced to 10–20% of that measured in the absence
of humic acid (Fig. 12). This signal decrease did not
appear to be due to the precipitation of metal humate
which we found to precipitate at higher con-
centrations of metal.

From the value of the concentration of humic acid
at the end point of the titration curve, we calculated
the equivalent weight of humic acid. For both copper

![Fig. 11. Variation of DPP and DPASV peak potentials as a function of glycine concentration for 2.5 x 10^{-6} M copper at pH 6.8.](image-url)
and lead by both DPP and DPASV, identical values of 950 g equiv \(^{-1}\) were obtained. This is in good agreement with the values reported by Visser and Mendel (1971). This procedure offers an alternative to those presently used.

Based on the end point of the titration curve, concentrations of humic acid in excess of 1:1 stoichiometry were used for stability constant determinations. The concentration of humic acid was expressed in m-equiv \(^{-1}\) and the peak potentials were plotted vs the logarithm of these concentrations (Fig. 13). For each metal two linear segments were apparent by each technique. In no case were the electrochemical reactions reversible. However, constant half-peak widths for lead by DPP and DPASV and for copper by DPP permitted estimation of the \(x\) values given in Table 1. Copper determinations by DPASV showed that the peak half-width varied and, therefore, the data were not treated further.

The stability constants determined from the DPP and DPASV data for the lead–humic acid system were quite different. In addition, the values of the ligand number are also quite different as determined by the two techniques. Since the same equivalent weight was determined by both methods and used to calculate the excess humic acid concentration, the ligand number resulting from the excess humic acid concentration, the ligand number should have been equal to 1. Because this was not the case, and because different values for log \(\beta\) and \(j\) were obtained by the two techniques, it can be implied that the necessary conditions for the application of the formula were not met. With the exception of electrochemical reversibility, all the conditions previously listed as having to be met for the Lingane equation must also be met for the modified equation. The decrease in signal accompanying complexation may be indicative of the non-labile character of a soluble complex or to the formation of a colloidal metal humate. Therefore, the system is not truly amenable to treatment by this means of data analysis.

In addition to the obvious non-lability of the metal–humic acid complex, the concentration of humic acid at the electrode surface may not be the same as in the bulk of the solution. When humic acid was added to a solution which had not been stripped of oxygen, the peak potential for oxygen shifted cathodically. This shift may indicate that the reversibility of oxygen reduction is decreased because humic acid is adsorbed at the electrode surface. Because the concentration of humic acid in the diffusion layer is unknown, application of conventional electroanalytical techniques to the calculation of stability constants is not possible.

**Conclusions**

This study has shown that cathodic and anodic voltammetric techniques can be used to characterize trace metal interactions in aquatic environments. Metal complexation as a function of pH and alkalinity could be detected easily by changes in the peak current and this technique was used to verify existing equilibrium models.

Stability constants were determined from shifts in the peak potential using the Lingane equation. Although this method has usually been used only for measurements involving reversible electrode processes, by the inclusion of the transfer coefficient in the Lingane equation, it was possible to extend the method to irreversible systems. For both the reversible and irreversible cases the applicability of the data treatment could be ascertained by comparison of DPP and DPASV results. Obtaining the same result by cathodic and anodic measurement implies that the result is correct.

At present it would appear that accurate measurement of stability constants involving ligands adsorbed at the electrode surface is not possible. This should not, however, preclude the use of electrochemical measurements in other situations.

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