Metal–organic complexation in the marine environment

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We discuss the voltammetric methods that are used to assess metal–organic complexation in seawater. These consist of titration methods using anodic stripping voltammetry (ASV) and cathodic stripping voltammetry competitive ligand experiments (CSV-CLE). These approaches and a kinetic approach using CSV-CLE give similar information on the amount of excess ligand to metal in a sample and the conditional metal ligand stability constant for the excess ligand bound to the metal. CSV-CLE data using different ligands to measure Fe(III) organic complexes are similar. All these methods give conditional stability constants for which the side reaction coefficient for the metal can be corrected but not that for the ligand. Another approach, pseudovoltammetry, provides information on the actual metal–ligand complex(es) in a sample by doing ASV experiments where the deposition potential is varied more negatively in order to destroy the metal–ligand complex. This latter approach gives concentration information on each actual ligand bound to the metal as well as the thermodynamic stability constant of each complex in solution when compared to known metal–ligand complexes. In this case the side reaction coefficients for the metal and ligand are corrected. Thus, this method may not give identical information to the titration methods because the excess ligand in the sample may not be identical to some of the actual ligands binding the metal in the sample.

Introduction

In the last two decades, our knowledge of trace metal speciation has grown tremendously. With the advent of trace metal clean sampling techniques and sensitive voltammetric techniques, the marine community now recognizes that metal speciation in seawater and estuarine waters is dominated by complexation with organic compounds of unknown composition and origin. Recent culture work has shown that microorganisms produce a variety of low molecular weight organic compounds that complex metals with high stability constants. These compounds have a variety of functional groups that include phosphate, carboxylic acids, amines, thiol and hydroxy groups. Specific functional groups such as hydroxamate, catecholate and -hydroxyaspartate are bidentate groups and organisms make molecules with three bidentate groups in a molecule. In addition, plant degradation products such as porphyrins are significant organic ligands that bind metals through four N atoms in a square planar arrangement. These latter multidentate molecules have very high stability constants with metals and are also kinetically inert to metal–ligand dissociation processes. For this reason, organisms generally uptake the free metal ion rather than a metal–ligand form. Thus, an understanding of metal–organism interactions requires an understanding of the amount of dissolved free ion present relative to the total dissolved metal concentration as well as the metal acquisition methods that an organism can use.

In this paper we review and compare the principal voltammetric methods, which provide evidence for metal–organic complexes. Most voltammetric work is performed with the hanging mercury drop electrode (HMDE) or the rotating disk electrode (RDE) with a thin mercury film (TMF) because these permit the measurement of metal–organic complexation at (sub)nanomolar levels directly in the solution of interest. The actual experimental methods can be broken into two broad categories and are based on the electrochemical behavior of the metal bound to an organic ligand.

The first method consists of titration experiments that measure the amount of ligand in excess to the metal in the solution and the conditional stability constant, for the excess ligand with the metal. The conditional stability constant equals the reciprocal of the side reaction coefficient for the metal can be corrected but not that for the ligand.

$$K_{cond \text{M-L}} = [ML]/([M] [L'])$$

where $M'$ and $L'$ are the concentrations of the metal and ligand that are not bound to each other. These are related to the total metal $[M]_T$ and $[L]_T$ via

$$[M'] = [M]_T - [ML] \quad \text{and} \quad [L'] = [L]_T - [ML].$$

The free metal $[M']$ plus the metal bound to other inorganic ligands equals $[M]$,

$$[M'] = [M'] + \sum MX_i$$

and the fraction of free metal in the solution without the organic ligand is given by

$$x_M = [M'] / [M'] + [M']$$

where

$$x = (1 + \sum K_{MX_i} [X])$$

This has also been expressed as the side reaction coefficient for $M'$, $x_M$, which is the reciprocal of $x_M$ or

$$x_M = [M'] / [M']$$
The conditional constant for $M'L$ is related to $M^{n+}L$ by

$$K_{\text{cond ML}} = [ML] / ([M^{n+}][L^-]) = K_{\text{cond ML}} (\Delta S_M)$$

Similar equations can be written for the organic ligand to give a thermodynamic constant,

$$K_{\text{therm}} = [ML] / ([M^{n+}][L^-]) = K_{\text{cond ML}} (\Delta S_L)$$

but in environmental samples the interactions of $H^+$, $Ca^{2+}$, and $Mg^{2+}$ with the ligand are unknown.

The titration experiments include (1) anodic stripping voltammetry\(^2\) (ASV), which is useful for metals that react at the electrode directly ($Cu^{2+}$, $Zn^{2+}$, $Cd^{2+}$, $Pb^{2+}$), and (2) cathodic stripping voltammetry/competitive ligand exchange\(^8,9\) (CSV-CLE) which is useful for metals that do not react at the electrode directly but have a metal–ligand complex that does ($Fe^{3+}$, $Co^{2+}$). The CSV-CLE method depends on the measurement of a known metal–ligand complex (the competing ligand), that adsors to the mercury electrode. In addition, a kinetic CSV-CLE approach\(^10,11\) for excess ligand binding a metal has been used to measure the metal organic formation rate constant ($k_1$), dissociation rate constant ($k_2$), the half-life or residence time ($t_{1/2}$) of the complex and $K_{\text{cond ML}}$ (from $k_1/k_2$). The second type of voltammetry method involves the breakdown of the actual complex \textit{in situ} and is termed pseudovoltammetry\(^45\)–\(^48\) which is useful for metals that react at the electrode directly. This method gives information on the amount of ligand binding to a specific complex with a thermodynamic constant, $K_{\text{therm}}$, that differs from $K_{\text{cond ML}}$. $K_{\text{cond ML}}$ is corrected for the side reaction coefficient of the metal but not the ligand whereas $K_{\text{therm}}$ is corrected for the side reaction coefficients of the metal and ligand \textit{via} comparison to metal–ligand complexes of known $K_{\text{therm}}$ (chelate scale).

We describe the use of these methods for unknown ligands in seawater as well as with model ligands in UV irradiated seawater for the metals $Cu(n)$, $Zn(n)$ and $Fe(m)$. In the case of CSV-CLE, we show for known $Fe(m)$-organic complexes that the use of different ligands [1-nitroso-2-napthol, or 1N2N, and salicylaldehyde, or SAL] gives comparable $K$ and ligand concentration data.

**Experimental**

The details of the experimental procedures for ASV and pseudovoltammetry work have been previously described by our group.\(^45\)–\(^46\) Total $Zn$ and $Cu$ concentrations were performed using the method of Bruland \textit{et al.} CSV-CLE and kinetic Fe(m) measurements with IN2N were performed as we have outlined previously.\(^10\)–\(^12\) CSV-CLE experiments with SAL were performed as described by Rue and Bruland.\(^6,7\) Examples of model ligands commonly used in experiments are given in Appendices 1 and 2. Appendix 2 shows types of strong ligands (functional groups are circled) that bind to Fe(m) and which may bind to other metals.

**Results and discussion**

**Metal–ligand complexes**

Voltammetry can provide information on a ligand actually binding a metal because many metal–ligand complexes give a discrete peak or half-wave potential. In a sample, these peaks can be compared to known ligand–metal complexes in the form of a metal-chelate scale (see pseudovoltammetry below). Fig. 1A shows the voltammetric reduction of inorganic $Zn(n)$ in UV irradiated seawater ($E_{p} = -1.05$ V) and Fig. 1B shows the reduction when $Zn(n)$ is bound to two nitrotriatic acid molecules (NTA: $E_p = -1.52$ V). The reduction is more negative for the $Zn$ complex with NTA than for inorganic $Zn$ because the ligand donates electrons more strongly than simple monodentate ligands such as chloride and hydroxide. In addition, two to four atoms in one NTA molecule can bind to $Zn(n)$ and the displacement of monodentate inorganic ligands by multidentate ligands gives rise to higher stability constants \textit{via} the “chelate” effect which is an entropy driven reaction; i.e., there are more product molecules than reactant molecules for the reaction\(^2\) (generalized eqn. (1) and (2); charges omitted for simplicity).

$$M(H_2O)_6 + L\text{--}L \rightarrow M(H_2O)_6(L\text{--}L) + 2H_2O$$

where $L\text{--}L$ indicates a bidentate ligand

$$\Delta G = \Delta H - T\Delta S = -RT\ln K$$

Every ligand that reacts with a metal can have a unique reduction potential that can be used for analysis and this is the basis for both the CSV-CLE and pseudovoltammetry approaches.

**ASV titrations**

We first discuss the titration approach for the measurement of metal–organic ligand complexes for metals that react directly at the Hg electrode (ASV experiment). In titration experiments, metal is added to an unknown sample and the inorganic form of the metal (e.g., Fig 1A for inorganic Zn indicates that the deposition potential should be more negative than $-1.1$ V) is analyzed \textit{via} deposition experiments for possible reaction at the Hg electrode. More than 95–99% of the metal is normally bound to an unknown organic compound(s), which is in excess to the metal in the sample. Fig. 2A shows that the inorganic Zn reduction peak from a Delaware Bay sample is suppressed until the excess ligand has been titrated by the addition of inorganic Zn. Linearization of the titration data is typically performed by use of the Langmuir or Ruzic transformation\(^35\)–\(^41\) [eqn. (3)] or the Scatchard transformation\(^59\) [eqn. (4)]. For the Langmuir linearization (Fig. 2B),

$$\frac{[M]}{[ML]} = \frac{[M]}{C_L} + \frac{\Delta S_M}{(K_{\text{cond ML}} C_L)}$$

a plot of $[M]/[ML]$ \textit{vs}. $[M]$ yields a straight line with slope $C_L$ from which $K_{\text{cond ML}}$ (the conditional stability constant uncorrected for the side reaction coefficient of the ligand) can be evaluated from the intercept. Note that $M_L = [M] = [ML]$, $[M]$ is the labile or inorganic $M$, and $\Delta S_M$ is the side reaction coefficient of the metal ion ($\Delta S_M$ for the divalent cations of the
first transition series in seawater is usually <0.2 log \( K \) units.\(^{30,51}\)

In Fig. 2B, the linearization plot for data in Fig. 2A shows that there is a single straight line showing only one complex with a \( C_L = 36.1 \) nM and a log \( K_{\text{cond ML}} = 9.03 \).

The Scatchard transformation is given by eqn. (4) and shown in Fig. 2C for the data in Fig. 2A. A plot of \([ML]/[M]\) vs. \([ML]\) gives a slope which is \( K_{\text{cond ML}} \) and \( [C_L] \) is the \( x \)-intercept for the regression line. In this linearization, two separate slopes are noted with a total ligand content of 37.1 nM. These data suggest that two ligands or ligand classes may be present in the sample. By convention, \( L_1 \) is the stronger ligand with a higher log \( K_{\text{cond ML}} \) of 9.13 (concentration = 33.7 nM) and \( L_2 \) is the weaker ligand with a smaller log \( K_{\text{cond ML}} \) of 8.89 (concentration is 37.1 – 33.7 = 3.4 nM). The Scatchard transformation is usually the better of the linearization methods for determining separate ligand classes especially when the log \( K_{\text{cond ML}} \) data are similar. More recently, non-linear methods\(^{42} \) have been gaining popularity.

It is important to reiterate that the \( K_{\text{cond ML}} \) data cannot be corrected for the side reaction coefficient of the unknown ligand in samples. Bruland\(^2 \) showed that the Zn-EDTA complex has a log \( K_{\text{cond ML}} = 7.9 \) in UV seawater but log \( K_{\text{cond ML}} > 11 \) in 0.1 M KCl of the same pH. The difference in these constants is due to the interaction of Ca and Mg in seawater with the carboxylic acid functional groups of EDTA. However, the actual thermodynamic constant for Zn–EDTA is log \( K_{\text{therm}} = 16.3 \). The fact that a log \( K_{\text{cond ML}} > 11 \) is calculated indicates that there is a titration window for these types of ASV titration experiments. The titration window depends on the concentration of the unknown ligand and the metal. In general, there is a window of about six log \( K \) units for these types of titrations.

**CSV-CLE titrations**

Any known metal–organic complex, which gives a voltammetry signal, can be used to study the interactions of that metal with an unknown ligand(s) in a sample. In this example, the known ligand is a competitive ligand, one competing for the metal in a sample. This approach must be used for metals such as Fe(n)\(^{3,12} \) and Co(n)\(^{23} \) that do not react directly at the mercury electrode. Several studies have also used this approach for metals\(^{40,41,44,46} \) such as Zn and Cu, which can be measured at the electrode. Comparison of the ASV and CSV-CLE methods\(^{42} \) for these metals shows similar \( [C_L] \) and \( K_{\text{cond ML}} \) data.

In the CSV-CLE case, metal in increasing concentration (from zero added metal) is added to a series of electrochemical cells containing the sample with the same amount of a competitive ligand. After analyzing each electrochemical cell, a plot similar to Fig. 2A results. Linearization of the data is given in eqn. (5), which is identical to eqn. (4)

\[
\frac{[M]}{[ML]} = \frac{K_{\text{cond ML}} [C_L] - K_{\text{cond ML}} [ML]}{[ML]} \quad (4)
\]

\[
\frac{[M]}{[ML]} = \frac{[C_L] + (2K_{\text{cond ML}} + 2K_{\text{cond ML}})[ML]}{K_{\text{cond ML}}[CL]} \quad (5)
\]

except for \( 2K_{\text{ML}} \), which is the side reaction coefficient for the metal with the competitive ligand. Much work has been performed to understand Fe(n) speciation in seawater. For Fe(n), the \( K_{\text{cond Felab}} \) of a Fe–natural ligand complex and total natural ligand concentration \( [C_L] \) can be calculated from the intercept and slope of a \([Fe_{\text{lab}}]/[Fe]\) vs. \([Fe_{\text{lab}}]\) plot. \([Fe_{\text{lab}}]\) is that metal that can bind with the competitive ligand and is obtained from the CSV Fe peak current, \( i_p \), and the sensitivity, \( S \) (slope of a standard curve in UV seawater); i.e. \([Fe_{\text{lab}}] = i_pS = [Fe^{3+}] \times (K_{\text{Fe}^{3+}} + K_{\text{Fe}^{3+}[B]} + K_{\text{Fe}^{3+}[C]} + K_{\text{Fe}^{3+}[D]}) \) and \([Fe_{\text{lab}}] = [Fe_{\text{lab}}] + [Fe_{\text{lab}}] \).

The \( K_{\text{Fe}^{3+}} \) is the \( x \)-coefficient for all inorganic species of Fe\(^{3+} \) (10\(^{-4} \) at pH = 7; 10\(^{-10} \) at pH = 8)\(^{44} \) and \( 2K_{\text{ML}} \) is the side reaction coefficient for Fe(n)L competitive ligand complexes. For Fe(n) with 1N2N,\(^3,12,52 \) the \( 2K_{\text{Fe}^{3+}[B]} \) is about 10\(^{13.04} \) at pH = 7 and 8. For salicylaldoxime,\(^3 \) the \( 2K_{\text{Fe}^{3+}} \) is about 10\(^{2} \) at pH = 8. The window for determination of \( K_{\text{cond Felab}} \) is smaller that the ASV method (about two log units) but can be varied by changing the ligand concentration. The \( K_{\text{Fe}^{3+}} \) for salicylaldoxime indicates that the \( K_{\text{cond Felab}} \) calculated is dependent on the accuracy of \( K_{\text{Fe}^{3+}} \) used. Byrne et al.\(^53 \) have estimated a value of 10\(^{15.5} \) so \( K_{\text{cond Felab}} \) can vary 1.5 log units based on the \( K_{\text{Fe}^{3+}} \) used.

Fig. 3 shows the log \( K_{\text{cond Felab}} \), for several model ligands in UV seawater determined by CSV-CLE titrations with the two competitive ligands (1N2N and SAL). In these calculations\(^4 \) an \( K_{\text{Fe}^{3+}} \) of 10\(^{10.0} \) at pH = 8 was used. The data show that the log \( K_{\text{cond Felab}} \) data are similar—usually within one log \( K \) unit—when using either competitive ligand. The use of 1N2N at pH = 7, where the Fe1N2N voltammetric peak is most sensitive, does not compromise the data. The main reason for this is the high \( K_{\text{Fe}^{3+}[B]} \) when compared to the \( K_{\text{Fe}^{3+}} \) of Fe(n) at these pH values. The vertical lines in Fig. 3A and 3B show the range of reported Fe(n)L log \( K \) values from the world’s oceans. Fig. 3 also shows that the model ligands binding Fe(n) give similar log \( K_{\text{cond Felab}} \) data regardless of their structure. This will be discussed below.

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Kinetic approach

This approach has been used to assess the rate constants for formation and dissociation of Fe(m)L complexes. The approach is briefly described but is detailed elsewhere. 10–12, 55 Excess Fe’ is added to a sample without any competitive ligand so that the excess Fe’ can bind to the organic ligand(s) in seawater (eqn. (6)). Aliquots of this solution are measured over time at the pH of the sample after addition of a competitive ligand to the aliquot. The k_f (rate of formation of FeL) is determined from this experiment. The excess ligand binding to Fe’ is determined by kinetic analysis of the time course.

Fe’ + L → FeL

(6)

The k_d and t_{1/2} are determined by recovering Fe’ in FeL by adding a competitive ligand such as 1N2N to an equilibrated sample (eqn. (7)). This is monitored over time.

FeL + 3(1N2N) → Fe(1N2N)_3 + L

(7)

Eqn. (7) can be broken into two eqns. (8) and (9)

FeL → Fe’ + L

(8)

Fe’ + 3(1N2N) → Fe(1N2N)_3

(9)

The k_d is evaluated using the steady state approximation for Fe’ which simplifies the kinetic expression to ln[FeL] = k_d t. The K_{cond} Fe_2L = k_d/k_f and K_{cond} Fe(III)L = K_{cond} FeL(x_Fe) where x_Fe = 10^{10}. Fig. 4 shows the log K_{cond} Fe(III)L data obtained from the kinetic approach at pH = 8 and the CSV-CLE approach for model ligands bound to Fe(III) in UV seawater. The agreement is excellent indicating that both methods give comparable results. To date the window for K_{cond} Fe(III)L using this method is log K = 18–23. In addition to the stability constant data, the kinetic data for FeL (Table 1) reflect the fast reaction rates via k_f and slow dissociation rates via k_d. The t_{1/2} and residence times for Fe(III)L complexes come directly from the k_d data (t_{1/2} × k_d = 0.693) and correlate with other estimates of iron residence times in the ocean. 56, 57

These CSV-CLE and kinetic data as well as solubility data58, 59 indicate that Fe(III) is primarily complexed by natural organic compounds in seawater.

Pseudovoltammograms and chelate scales

Metal reduced to an amalgam; e.g. ZnL + 2e^- → Zn(Hg) + L. When a metal–ligand complex is reduced to a metal amalgam, the half-wave potential of a metal complex, E_{1/2}', or the peak potential, E_p, can be directly related to the thermodynamic stability constant, K_{therm} 45–48 by eqn. (10):

E_{1/2}' = E_{1/2} - [2.303 RT ln K_{therm}]/nF

(10)

where E_{1/2} is the reduction potential of the free metal ion and n is the number of electrons involved in the process and K_{therm} = K_{ox} = [ML]/[M][L] for a 1:1 complex (for...
Table 1 Comparison of model FeL complex formation and dissociation rate constants, conditional stability constants, and Fe$^{3+}$ and Fe$^{2+}$ residence times in treated with Chelex, photo-irradiated seawater as determined using the kinetic method. Errors represent average mean ± s (standard deviation) from two separate replicates. Data taken from ref. 12.

| Model ligand                  | $k_f \times 10^5$/M$^{-1}$s$^{-1}$ | $k_d \times 10^{-4}$/s$^{-1}$ | log $K_{Fe/L}$ kinetic | log $K_{Fe^{3+}/L}$ | Fe$^{3+}$ residence time/yr | Fe$^{2+}$ residence time/yr | log $K_{therm}$ |
|------------------------------|------------------------------------|-------------------------------|------------------------|----------------------|---------------------------|---------------------------|-----------------|
| Protoporphyrin IX$^a$        | 6.2 ± 0.8                          | 0.7 ± 0.7                     | 11.9 ± 0.5             | 21.9 ± 0.5           | 0.031                     | 645                       | —               |
| Protoporphyrin IX$^b$        | 15.3 ± 0.2                         | 0.2 ± 0.9                     | 13.0 ± 0.2             | 23.0 ± 0.2           | 0.116                     | 5866                      | —               |
| Dimethyl ester$^c$           |                                    |                               |                        |                      |                           |                           | —               |
| Phaeophytin$^d$              | 12.7 ± 0.1                         | 12.3 ± 16.8                   | 11.0 ± 1.2             | 21.0 ± 1.2           | 0.002                     | 72                        | —               |
| Apoferritin$^e$              | 0.93 ± 0.3                         | 0.08 ± 0.04                   | 12.1 ± 0.1             | 22.1 ± 0.1           | 0.275                     | 820                       | —               |
| Phytic acid$^f$              | 12.8 ± 0.1                         | 0.51 ± 0.28                   | 12.4 ± 0.2             | 22.4 ± 0.2           | 0.043                     | 1820                      | —               |
| Alterobactin A$^g$           | 3.8 ± 0.8                          | 0.17 ± 0.04                   | 12.3 ± 0.4             | 22.3 ± 0.4           | 0.129                     | 1620                      | 49–53$^{18}$    |
| Alterobactin B$^h$           | 8.0 ± 0.6                          | 0.25 ± 0.02                   | 12.5 ± 0.3             | 22.5 ± 0.3           | 0.088                     | 2330                      | 43.6$^{18}$     |
| Enterobactin$^i$             | 10                                 | 15.8                          | 10.8                   | 20.8                 | 0.013                     | 46.0                      | 49.0$^{20}$     |
| Ferrichrome$^j$              | 4.6 ± 2.9                          | 0.05 ± 0.04                   | 12.9 ± 0.1             | 22.9 ± 0.1           | 0.439                     | 6700                      | 29.07$^{31}$    |
| Desferrioxamine$^k$          | 19.6 ± 10.1                        | 1.5 ± 1.8                     | 12.1 ± 0.6             | 22.1 ± 0.6           | 0.015                     | 952                       | 30.60$^{31}$    |

Fe complexing moieties for the model ligands: $^a$Porphyrin. $^b$Protein. $^c$Phosphate. $^d$β-Hydroxyaspartate/catecholate. $^e$Bis-catecholate. $^f$Tris-catecholate. $^g$Tris-Hydroxamate.

Fig. 5 (A) Square wave voltammogram of 100 μM Zn with 50 μM NTA and (B) pseudovoltammogram of 10 nM Zn with 500 nM NTA using anodic stripping square wave voltammetry.
weak third-ligand complex at 2.108 V (log $K_{therm}$ = 4.14 M$^{-2}$) is due to inorganic ligands and/or weak acids such as oxalate. The Zn concentration bound to each of these complexes in increasing negative potential is 1.7, 0.90 and 3.5 nM (5.7 nM combined based on the Zn peak sensitivity) whereas the total Zn concentration in the sample is 24.7 nM. Thus, 19 nM of complexed Zn compounds are still unaccounted for. This could be due to strong organic complexes (log $K_{therm}$ > 40) which have been found in natural waters that have log $K_{therm}$ > 40. These Zn–ligand complexes cannot be determined by the pseudovoltammetry approach because of sodium ion reduction, which permits an upper limit of log $K_{therm}$ = 18 for Zn.

These data are now compared with the ASV titration approach shown in Fig. 2. The latter method indicates that one complex (perhaps a second) with ligand in excess to the metal is present with a value for the conditional log $K_{cond ML}$ = 9.03.

The conditional $K_{cond ML}$ and $K_{therm}$ data are not readily comparable for Zn(II) because $K_{therm}$ data are due to the actual ligand complexes in the sample and $K_{cond ML}$ data are for the ligands in excess to the metal in seawater. The actual ligands binding Zn may or may not be the same as the excess ligands to total Zn in the sample. The log $K_{therm}$ data that are less than 9.03 are weak complexes that are not detected by both Langmuir and Scatchard linear transformations. The complex with log $K_{therm}$ = 11.45 (close to the ZnEDDA complex) may not be related to the log $K_{cond ML}$ data of 9.03 either because the actual ligand concentration binding Zn in this case via the pseudovoltammograms is smaller than the ASV titration calculation of 36.1 nM. Thus, the two methods appear to be giving information on different Zn complexes.

A similar approach has been used for Cu(II) as shown in Fig. 8. In that study, 17 known organic ligands were used to develop a chelate scale with a log $K_{therm}$ range of 12–26.5. The upper limit for this scale based on the sodium reduction wave is log $K_{therm}$ ~ 49. Interestingly, the largest log $K_{therm}$ value for a CuL ligand is smaller than the estimated CuL data from field and culture samples ($E'_{1/2}$ is more negative for the field samples) demonstrating that very strong CuL complexes can be formed. The strong CuL complex found in Martha’s Vineyard waters was matched by a ligand produced by a strain of Synechococcus. The moderately strong CuL complex in Eel’s Pond and in Martha’s Vineyard waters did not match the ligands from other cultures. The three cultures tested showed a great variability of ligands that can be produced by different organisms. The log $K_{cond Cu(II)L}$ for these complexes as determined by ASV titration ranged from 10.8 to 14.3. These conditional constants indicate that the side reaction coefficients for the ligand(s) are high and similar to what has been observed for ligands that form Fe(III) complexes, which are discussed below.

Reduction of a metal complex to a lower valency (no amalgam formation) Fe$^{3+}$ + e$^- \rightarrow$ Fe$^{2+}$. Similar chelate scale data can be obtained for metal complexes which do not decompose at the electrode to form metal amalgams. In this case, $E'_{1/2}$ is proportional to the ratio of the thermodynamic stability constants of the reduced and oxidized complexes according to eqn. (12):

$$E'_{1/2} = E_{1/2} - \frac{2.303 \, RT \, ln F}{nF} \, \log \frac{K_{red}}{K_{ox}}$$

where $K_{ox}$ and $K_{red}$ are the stability constants of Fe$^{3+}$ and Fe$^{2+}$. $[\text{Zn}]$ is the concentration of Zn in the sample. $[\text{Fe}]$ is the concentration of Fe in the sample.

**Fig. 6** A plot of $E'_{1/2}$ from pseudovoltammograms vs. log $K_{therm}$ for Zn–organic complexes dissolved in seawater. 1 ~ oxalic acid; 2 ~ CTP; 3 ~ ethylenediamine; 4 ~ glycine; 5 ~ 8-hydroxyquinoline; 6 ~iminobis(methylene phosphonic acid); 7 ~EDDA; 8 ~NTA as Zn(NTA)$_2$. The numbers 9–11 refer to unknown complexes in Delaware Bay waters (see Fig. 7).
Fe^{2+}L, respectively. If the $K_{red}$ values for all complexes are similar as shown for Fe(III) then the $K_{red}$ term can be incorporated into the intercept and the equation simplifies to eqn. (13):

$$E_{1/2} = \left[ E_{1/2} + \left( 2.303 \frac{RT}{nF} \right) \log K_{red} \right]$$

$$E_{1/2} = \left[ E_{1/2} + \left( 2.303 \frac{RT}{nF} \right) \log K_{ox} \right]$$

For this case the electrode processes are reversible (checked by CV or SWV) because the complex does not dissociate or become destroyed, and $E_{1/2}$ is independent of the ligand concentration (check by titrating the metal with ligand until no further change in $E_{1/2}$ is observed).

A chelate scale for Fe(III) has been developed using seven natural ligands (Table 2 and Fig. 9) which react with Fe(III) to form complexes spanning 20 log $K_{therm}$ units. For this example we discuss the binding properties with regard to eqn. (2). The first 3 complexes contain CDTA or NTA with or without a catechol. The low $E_p$ and log $K$ reflect that carboxylic acids do not bind (lower $D_H$) with Fe(III) like the other complexes which contain only catechol functional groups. Fe(cate)$_{2}^{2-}$ is stronger than these but weaker that the tris-catechol complexes. The nitro-catechol binds more weakly in the tris complex to Fe(III) than catechol because the nitro group is an electron withdrawing group. The enterobactin complex with Fe(III) shows the stronger binding effect of one molecule with three catechol groups than three separate catechol ligands. This is related to entropic effects via the ‘chelate’ effect. For example, Fe(cate)$_{3}^{3-}$ and Fe(ent)$_{3}^{3-}$ have the following reactivity [eqn. (14a) and (14b)] based on Fe(III).

$$\text{Fe(H}_2\text{O)}_{6}^{3+} + 3\text{cat}^{2-} \rightarrow \text{Fe(cate)}_{3}^{3-} + 6\text{H}_2\text{O}$$

$$\text{Fe(H}_2\text{O)}_{6}^{3+} + \text{ent}^{6-} \rightarrow \text{Fe(ent)}_{3}^{3-} + 6\text{H}_2\text{O}$$

The larger log $K_{therm}$ reflects that $\Delta G$ is controlled by $\Delta S$ in

Table 2 Electrochemical and stability constant data for Fe(III) complexes with selected ‘model’ ligands and natural siderophores. Measurements were made in 5 mM Bistris buffer adjusted to 0.1 M ionic strength with NaCl.

| Complex* | pH | $E_{p}/V$ vs. SCE | Log $K_{therm}$ ($I = 0.1, 25^\circ C$) |
|-----------|----|-------------------|------------------------------------------|
| Model ligands |
| 1. [FeCDTA]$^{2+}$ | 7 | -0.145 | 30.0 |
| 2. [FeNTA$tiron$]$^{2+}$ | 7 | -0.182 | 37.1 |
| 3. [FeNTA$catechol$]$^{2+}$ | 7 | -0.211 | 32.9 |
| 4. [Fe(cate)]$^{2+}$ | 7 | -0.354 | 34.7 |
| 5. [Fe($4N$cat)$_3$]$^{3+}$ | 7 | -0.440 | 40.0 |
| 6. [Fe(cate)$_2$]$^{3+}$ | 7 | -0.680 | 43.7 |
| 7. [Fe(ent)$_3$]$^{3+}$ | 7 | -0.924 | 49.0 |
| Siderophores |
| 8. [Fe(Alt-B)$_2$]$^{2+}$ | 6 | -0.428 | 37.6 |
| 9. [Fe(Alt-B)$_3$]$^{3+}$ | 8.2 | -0.672 | 43.6 |
| 10. Fe-PCC7002 No. 1 | 7 | -0.445 | 38.1 |
| 11. Fe-PCC7002 No. 3 | 7 | -0.618 | 42.3 |
| 12. Decapeptides mefp1 | 7 | -0.510 | 39.6 |
| 13. mefp1 | 7 | -0.542 | 41.6 |

*CDTA = cis-1,2-cyclohexylenedinitrilotetraacetate, NTA = nitrilotriacetate, tiron = 4,5-dihydroxy-1,3-benzenedisulfonic acid, cat = catechol, 4Ncat = 4-nitrocatechol, ent = enterobactin, Alt-B = alterobactin B, PCC7002 = Synechococcus sp. PCC7002 isolates (complex stoichiometry is not known for eqn.(10) and (11)). $\pm 10$ mV. See Table 1 of ref. 47 for references.

Fig. 8 A plot of $E_{1/2}$ from pseudovoltammograms vs. log $K_{therm}$ for Cu(II)–organic complexes dissolved in seawater. Blue symbols are from cultures and green symbols are from natural waters as indicated.

Fig. 9 A plot of $E_{p}$ from square wave voltammograms vs. log $K_{therm}$ for Fe(III)–organic complexes dissolved in 0.1 M KCl. Numbers refer to compounds in Table 2.
eqn. (14b) because the $\Delta H$ term for catechol functional groups is similar for eqn. (14a) and (14b). These data suggest that $k_d$ for the Fe(ent)$^{3+}$ containing a tris-catechol structure is smaller than for three separate catechol groups in Fe(cat)$^{3+}$. Because $K = k_d/k_f$, $K$ and $\Delta G$ increase with smaller $k_d$.

Fig. 9 also shows data for catechol ligands produced by different organisms. Mytilus edulis produces a 100 kDa foot protein (mefp1) which contains several catechol groups. It is not known how many catechol groups bind to Fe(n) in the protein but $E_0$ and log $K_{\text{therm}}$ are larger than the bis-catechol Fe(n) complexes. Tryptic digests of the foot protein produce decapeptides that react to form FeL$^2$ bis-catechol complexes, as in Fig. 10. These bind more strongly with Fe(n) than two catechols in Fe(cat)$^{3+}$ and this stronger binding appears related to interaction of the peptide chains with each other which helps to lower $k_d$. Similar results have been noted for the bis(catechol) complex of alterobactin-B from Alteromonas luteoviolacea.

These data show that the known ligands have a significant difference in log $K_{\text{therm}}$. However, these ligands and other known Fe(n) binding ligands have remarkably similar log $K_{\text{cond Fe(n)l}}$ values (Fig. 3 and 4) despite having different structures. This suggests that proton loss from the ligand and not Mg, Ca binding are important for the binding of Fe by the ligand. The side reaction coefficients for these ligands differ in such a way that when correcting for proton effects, the $K_{\text{therm}}$ data is different. The model ligands range from catecholate groups, which have 2 protons per functional group (6 total for enterobactin), to one proton per functional group for hydroxamates (3 total for desferrioxamine because there is no proton attached to the C=O group). The three proton difference for enterobactin (also alterobactin-A) vs. desferrioxamine leads to a different $K_{\text{therm}}$. In addition, porphyrins have 2 protons per functional group and all four N atoms can bind Fe. The effect of losing 6 protons from Alterobactin-A, 3 protons from desferrioxamine and 2 protons from a porphyrin lead to similar log $K_{\text{cond Fe(n)l}}$.

**Conclusions**

The ASV and CSV-CLE methods for the determination of organic–metal complexation give similar results for $K_{\text{cond ML}}$ and ligand concentration. These data relate to the ligand in excess to the metal in solution. ASV titrations have a larger $K_{\text{cond ML}}$ window than CSV-CLE, which can be varied by changing the competing ligand concentration. The excess ligand may or may not be the same as the actual ligand in the metal–ligand complex in the sample. The pseudovoltammogram method gives $K_{\text{therm}}$ and ligand concentration data on the actual complex(es) in solution within the window limited by the reduction of sodium ion. The data from the titration methods and the pseudovoltammogram data are not necessarily similar as shown for Zn. For Fe(n), the choice of competitive ligand for the CSV-CLE methods does not appear to affect the log $K_{\text{cond Fe(n)l}}$ data. The kinetic approach also agrees with the CSV-CLE methods. These similarities are due to measuring the same ligand types; i.e. excess ligand to the metal in the sample.

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**Appendix**

**Appendix 1**

![Fig. 10 Hyperchem MM+2 calculation for the FeL$^2$ complex from a decapeptide prepared from Mytilus edulis foot protein 1. The two decapeptide ligands appear to interact to stabilize the complex and prevent dissociation. The log $K_{\text{therm}}$ estimated is 40.2 vs. 34.7 for the bis(catechol) complex. The Fe atom bound to six oxygen atoms is in the upper left part of the figure.](Image)

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