Redox behavior of Biologically Important Cu(II)-Valine Complexes at Different pH by Electron Transfer Process

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Abstract: In human and animal bodies trace amount of transition metals such as Iron (Fe), Copper (Cu), Manganese (Mn) etc. are present. Fe is the most essential metal ion usage in hemoglobin for oxygen transport. Cu is one of the biologically important chemical species that functions as a co-factor in metalloenzymes and metalloproteins. Mn is also an important microelement for plants and adult human body. Electrochemical studies of such metal ions in presence of ligands have been carried out in order to understand the mode of interaction and electron transfer mechanism of transition metals complexes with amino acids or proteins. The electrochemical behavior of metal Cu(II) with Val has been studied by Cyclic voltammetry (CV), Chronamperometry (CA), UV-Vis and FTIR spectroscopic techniques. This research was carried out to studied the electrochemical behavior of metals Cu(II) with Valine (Val) by Cyclic voltammetry (CV), Chronamperometry (CA), UV-Vis and FTIR spectroscopic techniques. Electrochemical and spectroscopic techniques reveal the interaction of Cu(II), with the ligands (Val) that may be formed of metal-ligand complexes. Electrochemical studies have been carried out in variation of metal ion concentration, ligand concentration, buffer solution of different pH and different scan rate at different electrodes. In this study all the experiment is carried out only in aqueous buffer solution; KCl or other conventional supporting electrolytes are not used in the solution. The effect of pH of Cu(II)-Val was studied by varying pH from 3.0 to 9.0. The maximum peak current was obtained at pH 4. This shows that the electrochemical oxidation of Cu(II)-is facilitated in acid media and consequently the rate of electron transfer is quicker. The slopes of the plots of E_p against pH of Cu-Val metal ligand system was determined graphically as anodic peak 21-31 mV/pH at 0.1V/s, which is close to the theoretical value of 30 mV which indicates that the oxidation of all the studied metal-ligand systems proceeded via the $2e^{-}/2H^{+}$ processes. During the bulk electrolysis the electrosynthesized metal-ligand products were generated. UV-Vis and FTIR techniques were also used to confirm the interaction of Cu(II) with Val. Keywords: Cyclic voltammetry, pH effect, Cu(II)-Valine complex, Chronamperometry (CA), UV-Vis and FTIR.

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I. Introduction

Transition metal complexes are seen at the active sites of a great variety of oxidation-reduction active proteins owing to their capacity of accessible in more than one stable oxidation state illustration them to function in detoxification, transport, catalysis and storage in biological processes [1]. Due to the physiological importance of metal-ligand complexes, there has been much attempt taken to know their redox nature.

The rate of electron transfer of the complexes depends upon the degree of charge localization, that is, the coupling between the mixed-valence centers. Coupling depends on several factors, including distance between mixed-valence centers and the environment surrounding each center. Robin and Day developed three broad categories describing the degree of interaction between mixed-valence centers. Class I compounds are described as having localized valences, where there is little or no communication between the mixed-valence centers, with an activation energy barrier to electron transfer between the centers. Class III compounds are described as having completely delocalized charge, with no barrier to coupling between mixed-valence centers [1].

Electrochemical behavior of polymetallic clusters, such as oligomers of ferrocene, often include cyclic voltammograms with multiple one-electron couples, the separation of which depends upon the degree of

coupling between metal centers [2, 3]. These complexes are unique in that it is the metal center of the complex that acts as the bridge between redox-active ligands, rather than ligands acting as a bridge between metal centers. Some of these complexes exhibit properties desirable in Cu(II)-Val, with multiple reversible redox couples within a narrow range of potential. These may supply or accept multiple electrons to catalyze redox reactions. One advantage of ligand-based redox chemistry is that it does not rely on oxidative additon/reductive elimination to transfer electrons, so the chemistry it facilitates does not have to be on substances that ligate with the metal. Furthermore, the narrow potential range of the redox couples mitigates the problem of chemicallyactive intermediates by allowing multiple electrons to be transferred nearly simultaneously [2,3].

It is known that O, N and S atoms take part a key role in the coordination of metals at the active place of many metallobiomolecules [4]. O, N, S sites having metal complexes was widely examined because they have antibacterial, anticancer, industrial, antifungal, herbicidal and antiviral applications [5-10]. They act as models for biologically important species and have applications in biomimetic catalytic reactions. Chelating ligands containing N, S and O donor atoms provide wide biological activity and are of particular attention as of the diversity of behavior in which they are bonded to metal ions. It is recognized that the existence of metal ions bonded to biologically active compounds may improve their activities [8-10]. The complexes of Fe(III), Co(II), Mn (II), Cu(II) and Ni(II) ions with amines have been synthesized and analysis using diverse physico-chemical process, such as conductance measurements, elemental analysis, and UV-Vis, FTIR and NMR spectra. [11]. In order to understand the issue of interaction of Cu(II) with Val at different pH, an experimental aspect of redox interaction in terms of electrochemical current potential curves analysis is necessary. To the best of our knowledge, Cyclic voltammetry (CV), Chronamperometry (CA), UV-Vis and FTIR spectroscopic techniques and differential pulse voltammetric studies of Cu(II) with Val at different pH, at different electrodes, different scan rate and variation concentration of ligands has not been analyzed.

II. Experimental Method

The electrochemical behavior of Cu (II)-Val at various pH has been investigated using controlled potential coulometry(CPC), Amperometry and cyclic voltammetry at glassy carbon (GC) or platinum electrode (Pt) or gold electrode. Details of the instrumentation are given in the following sections. The thoroughly cleaned glass pieces were dried in electric oven. The smaller pieces of apparatus were dried in an electric oven and stored in a desiccator, while larger pieces of apparatus were used directly from the oven.

III. Results And Discussion

Electrochemical behavior of Cu(II)-L-Valine System:

The electrochemical behavior of Cu (II) with L-Valine (Val) system in aqueous buffer solution was carried out by cyclic voltammetry (CV). Cyclic voltammogram of Cu (II) ion in presence of Val at 0.1V/s at pH 4 is shown in Figure 1 (red line). Electrochemical experiments are performed in only aqueous buffer solution, KCl or other conventional electrolytes are not used as a supporting electrolyte. Buffer solution contains electrolyte that act as a supporting electrolyte. It is seen from the Figure, the anodic peak arises at 0.06V and the corresponding cathodic peak at -0.3V. In contrast, Cu (II) appears (green line) one anodic peak at 0.09V and two cathodic peaks at -0.28V and -0.1V in the identical experimental state. Pure Val (blue line) displays electroinactivity in the similar investigational condition (Figure 1). It is observed from the Figure, the peak point of the voltammogram of Cu-Val is altered negatively as regards of only Cu (II). On the adding of Val into Cu (II) the anodic oxidation peak moves left to 0.03V and reduction peak also shifts negatively 0.07V (Figure 1). The peak current also reduces considerably as regards with of free Cu (II) in the similar investigational conditions. This activity attributed the formation of Cu-Val complex. Similar voltammetric properties of Cu (II) and Mn (II) with ascorbic acid were stated [12-14].





The CV of Cu-Val in buffer solution (pH 4) at different scan rates is shown in Figure 2. The peak current of the anodic and cathodic peak rises with the increasing of scan rate (Table 1). The cathodic peaks are moved in the direction of left and the anodic peak is altered slightly to the right direction with increasing in scan rate. Figure 3 shows plot of the anodic and cathodic net peak currents of Cu-Val (second cycle) versus the square-root of the scan rates where the net current is measured by the scan-stopped method [15]. The almost proportionality of the redox peak current indicate that the peak current of the species is controlled by diffusion process. The CV of uncoordinated 2mM Cu (II) in aqueous solution at different scan rates is shown in Figure 4. Figure 5 demonstrates plots of the anodic and the cathodic net peak currents versus the square-root of the scan rates (Table 2). The proportionality of the anodic and the cathodic net peaks also attributed that the peak current of the species at each redox reaction is governed by diffusion process.

Peak current ratios (I_{pa}/I_{pc}) of the corresponding peak of Cu-Val are higher than one (Table 1). It suggests that redox reactions of Cu-Val are not reversible. Peak potential difference (ΔE) of the corresponding peak is > 0.1 V. This is much higher than one electron transfer reaction. Sometimes higher values of ΔE are showed owing to uncompensated solution viscosity and the diffusion of the species is not linear [16]. The value of ΔE increases with the increasing of scan rate. This attributed that there is a restriction owing to charge transfer kinetics [17]. When current passage through the cell containing electrolytes need additional driving force to overcome the viscosity (solution resistance) of the ions to shift towards the electrode. The corresponding peak potential difference (ΔE) of Cu(II)- Val is lower than Cu(II) in the similar investigational condition.

pH effect of Cu-Valine system

CV of Cu(II)-Val in aqueous buffer solution (without additional supporting electrolyte) of various pH (3.0, 4.0, 5.0, 6.0 and 7.0) at 0.1V/s scan rate is shown in Figure 6. The peak point of the redox species is strongly influenced by pH and it is shifted negatively with the increase of pH. The peak current versus pH is plotted as shown in Figure 7. It is observed from the Figure, the oxidation peak current rises with the increasing of pH upto pH 4 then the peak current decreased (Table 3). At pH 7 the CV is shown electro-inactive. The highest peak current is found at pH 4 (Table 3). At elevated pH (pH >7) the Cu(II)-Val complexes are nearly electroinactive. This ascribed that the redox reaction of Cu(II)-Val systems is favored in acid media (pH<7). Therefore the rate of electron transfer is faster in acid media and the optimum condition is found to be pH 4.

The coordination sites of the ligands are most active at pH 4 that may be facilitated for the formation of Cu(II)Val complex. With the increase of pH (pH>7), the coordination sites of the ligands may be deactivated, therefore the anodic and cathodic peak disappeared.

The plot of oxidation peak potential, E_p values versus pH is shown in Figure 8. The slope of the plot has been measured graphically as the anodic peak 31mV/pH at scan rate 0.1V/s. This value is very close to the value for a two-electron transfer process [13]. This ascribed that the redox reaction of the Cu(II)-Val carried on by means of the 2e processes. One step two electron transfer reaction may be occurred in the solution of without conventional supporting electrolyte as: Cu(II)-Val +2e \leftrightarrow Cu(0) + Val



Figure 2: CV of 2 mM Cu(II) with 40 mM Val (1:20) of GC electrode in buffer solution of pH 4 at scan rate 0.05 V/s to 0.5 V/s (2nd cycle).



Figure 3: Plots of peak current (I_p) versus square root of scan rate $(v^{1/2})$ of 2 mM Cu(II) with 40 mM Val of GC electrode in buffer solution (pH 4).



Figure 4: CV of 2 mM Cu(II) of GC electrode in buffer solution of pH 4 (without KCl solution) at scan rate 0.05 V/s to 0.5 V/s (2^{nd} cycle).



Figure 6: Cyclic voltammogram 2 mM Cu(II) with 40mM L-Valine of GC electrode in different pH (3, 4, 5, 6 and 7) (without KCl solution) at scan rate 0.1 V/s.



Figure 5: Plots of peak current (I_p) versus square root of scan rate $(v^{1/2})$ of 2 mM Cu(II) of GC electrode in buffer solution (pH 4) (2nd cycle).



Figure 7: Plots of peak current vs pH (3, 4, 5 and 6) of 2 mM Cu(II) with 40 mM L-Valine of GC electrode at scan rate 0.1 V/s (2nd cycle).



Figure 8: Plots of peak potential vs pH (3, 4, 5, 6 and 7) of 2 mM Cu(II) with 40 mM L-Valine of GC electrode at scan rate 0.1 V/s (2nd cycle).

Table 1: Peak potential (E_p) , corresponding peak potential difference (ΔE), peak current Ip (μA) and corresponding peak current ratio (I_{pa}/I_{pc}) of 2mM Cu (II) with 40mM L-Valine (1:20) of GC electrode in buffer solution of pH 4 without conventional supporting electrolyte (SE) at scan rate 0.05 V/s to 0.5 V/s (2nd cycle).

ion of pH 4 without conventional supporting electrolyte (SE) at scan rate 0.05 V/s to 0.5 V/s (2^{14}					$0.5 \text{ V/s} (2^{14} \text{ cy})$	
v/Vs^{-1}	$E_{\rm pa1}/{ m V}$	$E_{\rm pc1}/{\rm V}$	$\Delta E = Epc1 \sim Epa1$	$I_{pa1}/\Delta A$	$I_{pc1}/\Delta A$	Ipa1/ Ipc1
0.05	0.05	-0.32	0.37	47.69	-25.35	1.88
0.10	0.05	-0.33	0.38	180.26	-35.73	5.04
0.15	0.05	-0.33	0.38	217.29	-46.13	4.71
0.20	0.06	-0.34	0.40	243.47	-57.46	4.23
0.25	0.06	-0.34	0.40	269.51	-61.83	4.35
0.30	0.08	-0.34	0.42	309.04	-64.98	4.75
0.35	0.08	-0.36	0.44	327.23	-66.97	4.88
0.40	0.10	-0.36	0.46	342.16	-69.18	4.94
0.45	0.10	-0.39	0.49	350.17	-66.30	5.28
0.50	0.10	-0.39	0.49	358.18	-63.36	5.65

pН	Peak Current, $I_{pa}(\mu A)$	Peak potential, E _{pa} (V)
3	179.230	0.05
4	231.310	0.04
5	197.270	0.03
6	131.130	-0.05

Table 2: Peak Current I_p (μ A) and peak potential E_p (V) of 2 mM Cu (II) with 40 mM L-Valine of GC electrode without conventional SE at scan rate 0.1 V/s in different pH media (2nd cycle)

UV-Vis spectra Analysis of Cu (II) and Valine:

The electronic spectrum of the Cu (II)-Val complex and only Val are shown in Figure 9. In this Figure, it is seen that Cu (II)-Val adduct shows a broad absorption spectrum at 264nm (red line). Uncoordinated Val shows no absorption peak. The wavelength shift of Cu(II)-Val complexes from that of pure Cu (II) and pure Val ascribed the formation of complexes [18].



Figure 9: UV-Vis spectra of 0.2 mM L-Valine (blue line) and 0.2 mM Cu(II)-Valine (red line) in buffer solution of pH 4 (without KCl solution).

FTIR Spectral ana Valis of Cu (II)-Valine complex:

The FTIR spectrum of the vibrational modes of the Cu(II)-Val complex and pure Val were shown in Figure 10.The spectrum of Cu(II)-Val is different from that of pure Val specially in finger print region. The bending vibration of Cu (II)-Val decreases enormously with respect to pure Val suggesting that the formation of complexes of Cu(II) with Val [19].



Figure 10: FTIR spectra of L-Valine (blue line) and Cu(II)-Valine (red line). Product Cu(II)-Valine extracted from bulk electroValis of Cu(II) with L-Valine in buffer solution of pH 4.

IV. Conclusion

Cyclic voltammetry (CV), UV-Vis and FT-IR techniques were used to investigate the interaction behavior of biologically important metal ions such as Cu(II), with L-Valine, in aqueous solution with various pH, and scan rates. Electrochemical and spectroscopic techniques reveal the interaction of metal (Cu,) with the ligands (Val) that may be formed of metal-ligand complexes. L-Valine, is electro-inactive ligand. The voltammogram of Cu(II)-ligand system is irreversible. From the voltammogram studies, it is seen that Cu (II)

interacts with L-Valine. The peak current of metal-ligand complexes increases with the decrease of pH indicating that at lower pH the metal-ligand complexes are highly electro-active. The electrochemical oxidation of the studied metal ligand complexes are facilitated in acid media. The slope of peak potential, $E_{\rm p}$ against pH of studied metal-ligands varies linearly and the slopes of the plot are close to 30 mV which indicates that the oxidation of the metal-ligand complexes proceeded via the 2e⁻/2H⁺ processes. The peak current of the complexes at each redox reaction is controlled by diffusion process. The optimum condition of elelectrochemical reaction such as pH, composition of metal:ligands, scan rate are determined. The spectra of the studied pure metals and ligands were different from metal-ligand complexes indicating the complexation of metals with ligands. The behavior of electrode reaction of all the studied systems is electron transfer, chemical reaction and electron transfer (ECE) type.

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