

Mixed-ligand complex formation equilibria of Cu^{II} with biguanide in presence of glycine as the auxiliary ligand

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Abstract. Equilibrium study on the mixed ligand complex formation of Cu^{II} with biguanide(Bg) and glycine (HG), indicated the formation of the complexes: Cu(Bg)²⁺, Cu(Bg)₂²⁺, Cu(Bg-H)(Bg)⁺, Cu(Bg-H)₂, Cu(Bg)(OH)⁺, Cu(Bg-H)(OH); Cu(G)⁺, Cu(G)(OH), Cu(G)₂; Cu(G)(Bg)⁺, Cu(G)(Bg-H); (G)Cu(Bg)Cu(G)²⁺, (G)Cu(Bg-H)Cu(G)⁺, and (G)Cu(Bg-2H)Cu(G). From the deprotonation constants of coordinated biguanide (Bg) in the complexes Cu(Bg)(OH)⁺, Cu(Bg-H)(Bg)⁺ and Cu(G)(Bg)⁺, the Lewis basicities of the coordinated ligand species (Bg-H)⁻, OH⁻ and glycinate (G⁻) were found to be of the order: (Bg-H)⁻ >> OH⁻ > G⁻. Bridging (N¹-N⁴, N²-N³) tetradentate mode of coordination by biguanide species Bg, (Bg-H)⁻ and (Bg-2H)²⁻ was indicated from the occurrence of biguanide-bridged dinuclear mixed ligand complexes (G)Cu(Bg)Cu(G)²⁺, (G)Cu(Bg-H)Cu(G)⁺, (G)Cu(Bg-2H)Cu(G) in the complexation equilibria.

Keywords. Copper^{II}-biguanide; glycinate; ternary complex; formation constants.

1. Introduction

Biguanide or guanylguanidine (Bg) is an extensively studied (N,N) donor bidentate ligand. Transition metal complexes of biguanide, and its various N-substituted derivatives and related ligands have been investigated by many workers.^{1,2} Most of these studies were aimed at isolation of transition metal complexes with these ligands and their characterization and structure elucidation by various physicochemical techniques. Modes of metal ion coordination by biguanide were extensively investigated by many workers and (N,N) bidentate chelation by the N² and N⁴ atoms were established to be the most acceptable mode of coordination.³⁻⁷ Protonation-deprotonation equilibria of biguanide and substituted biguanides were also investigated by several workers.⁷⁻¹¹ Stepwise formation and decomposition of Cu^{II} and Ni^{II} binary complexes with biguanide and some substituted biguanides were studied by pH metric, polarographic and spectrophotometric methods⁸⁻¹⁴ and the order of relative stabilities of such complexes were established. Stability constant of Ag^{III}-ethylenedibiguanide complex was determined by pH metric and potentiometric methods.¹⁵ Instabi-

lity constants of Co^{III} and Cr^{III} tris(biguanide) and tris(phenylbiguanide) complexes were determined by potentiometric investigation on the decomposition of the corresponding complexes with acid.¹⁶⁻¹⁸ Literature survey revealed the absence of any report of a systematic equilibrium study on the mixed ligand complex formation of metal ions with biguanide and related derivatives. In view of the importance of biguanide, substituted biguanides and their metal derivatives as antimalarial and antidiabetic agents,^{19,20} as well as antimicrobial, antifungal and ovicidal activities of certain metal biguanide complexes,²¹ it was considered worthwhile to study the mixed ligand complex formation equilibria of biological metal ions with biguanide and related ligands in the presence of typical biomolecules, such as, amino acids and small peptides and other biomolecules as auxiliary ligands, to throw some light upon the molecular mechanism of actions of biguanide derivatives as drugs. Results of such investigation would provide useful clues to the design of new bio-inorganic drugs with greater selectivity and specificity at the same time lower toxicity.

In this paper we describe the results of a combined pH-metric and spectrophotometric equilibrium study on the mixed ligand complex formation of Cu^{II} with biguanide (Bg) in presence of the simplest amino

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Table 1. Formation constants (b_{pqrs}) of binary and mixed ligand Cu^{II} complexes with biguanide (Bg) and glycinate (G⁻) in aqueous solution, ionic strength, $I = 0.1 \text{ mol dm}^{-3}$ (NaNO₃) at $25 \pm 1^\circ\text{C}$.

Complex species	p	q	r	s	$\log b_{pqrs}$
H ₂ Bg ²⁺	0	1	0	-2	14.20
HBg ⁺	0	1	0	-1	10.90
H ₂ G ⁺	0	0	1	-2	11.87
HG	0	0	1	-1	09.61
Cu(OH) ⁺	1	0	0	1	-6.29
Cu(OH) ₂	1	0	0	2	-13.05
Cu(G) ⁺	1	0	1	0	08.23
Cu(G)(OH)	1	0	1	1	01.11
Cu(G) ₂	1	0	2	1	15.19
Cu(Bg) ²⁺	1	1	0	0	06.74
Cu(Bg)(OH) ⁺	1	1	0	1	01.70
Cu(Bg-H)(OH)	1	1	0	2	-5.18
Cu(Bg) ₂ ²⁺	1	2	0	0	12.51
Cu(Bg)(Bg-H) ⁺	1	2	0	1	05.99
Cu(Bg-H) ₂	1	2	0	2	-3.34
Cu(G)(Bg) ⁺	1	1	1	0	16.84
Cu(G)(Bg-H)	1	1	1	1	10.74
G(Cu)(Bg)Cu(G) ²⁺	2	1	2	0	31.24
G(Cu)(Bg-H)Cu(G) ⁺	2	1	2	1	25.36
G(Cu)(Bg-2H)Cu(G)	2	1	2	2	19.48

Deprotonation constants of coordinated biguanide (Bg)

$\log K^{\text{H}}_{\text{Cu(Bg)(Bg-H)}}$	$\log K^{\text{H}}_{\text{Cu(Bg)(OH)}}$	$\log K^{\text{H}}_{\text{Cu(G)(Bg)}}$
-16.09	-6.88	-6.10

Limits of error: ± 0.02

where, p , q , r and s are stoichiometric numbers. p , q and r may be positive integers or zero. s is a negative integer for a protonated species, zero for a neutral species and a positive integer for a deprotonated or a hydroxo species. In dilute solution, for binary 1 : 1, 1 : 5 Cu^{II} : ligand systems, p is 1 and the values of q and r depend upon the molar proportions of the metal : ligand. For 1 : 1 Cu^{II} : ligand systems, q or r does not exceed 1, but for the 1 : 5 Cu^{II} : ligand systems, higher values may be possible. For the ternary 1 : 1 : 1 Cu^{II} : Bg : HG system, $p = q = r = 1$ and for the 2 : 1 : 2 Cu^{II} : Bg : HG system, $p = 2$, $q = 1$, $r = 2$. The stoichiometries of the possible complexes occurring in all the binary and ternary systems are presented in the combined table 1, which also includes the computer-refined values of $\log b_{pqrs}$ and some other relevant constants of the binary and mixed ligand complexes. As the pH range of complex formation equilibria are overlapping with the hydrolytic equilibria of the Cu(aq)²⁺ ions, formation of the hydroxo species, Cu(OH)⁺ and Cu(OH)₂ have also

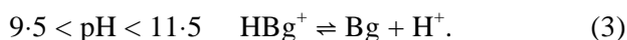
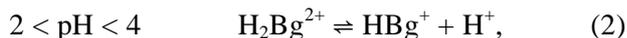
been considered in calculating the formation constants. However, all the pH-titration data have been collected at pH values prior to the appearance of any turbidity.

Although (N, O⁻) bidentate mode of coordination of glycinate ligand in its metal complexes is well known, models involving methyl amine-like only N-coordinated glycine, acetate-like only O⁻-coordinated glycinate and bridging (N, O⁻) coordinated glycinate containing mixed ligand complex species, viz. (Bg)Cu(H₂NCH₂COOH), (Bg)Cu(O⁻OCCH₂NH₃⁺) and (Bg)Cu(H₂NCH₂COO⁻)Cu(Bg) respectively, have been also considered in the calculation in addition to the expected complexes involving (N, O⁻) bidentate coordinated glycinate ligand species.²⁶ However, in all these cases the standard deviations are large and the percentage concentrations of the corresponding complex species are negligibly very small and are not even identifiable on the speciation curves (figures 2–4). Hence, these complexes have been excluded in the final calculation of the formulation constants.

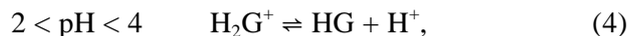
3. Results and discussion

3.1 Proton-ligand equilibria with biguanide (Bg)

In weakly acidic solution (pH ~ 2), biguanide (Bg) exists in its dipronated form, H_2Bg^{2+} , and with rise of pH, undergoes successive deprotonation to form the monoprotonated (HBg^+) and neutral (Bg) species, providing two well-separated buffer regions.¹

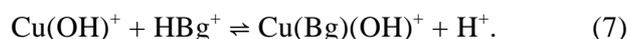
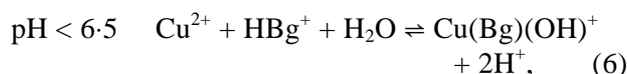


In the pH range 2–4, glycine (HG) exists in its monoprotonated form (H_2G^+), which with rise of pH also provides two well-separated buffer regions²⁶ due to successive deprotonation of the $-\text{COOH}$ group (equilibrium (4)) and the NH_3^+ group (equilibrium (5)).



3.2 Binary Cu^{II} : Bg equilibria

Complex formation equilibria in both (1:1) and (1:5) Cu^{II} :Bg systems start above pH 5 (figures 2a,b), where, Cu^{II} exists as $\text{Cu}(\text{aq})^{2+}$, $\text{Cu}(\text{OH})^+$ and $\text{Cu}(\text{OH})_2$. The predominant complex species at pH < 6.5 in the 1:1 Cu^{II} :Bg system is the ternary hydroxo complex, $\text{Cu}(\text{Bg})(\text{OH})^+$, formed out of Cu^{2+} , $\text{Cu}(\text{OH})^+$ and HBg^+ according to equilibria (6) and (7) as the speciation curves (figure 2a) imply:



With rise of pH, the 1:1 Cu^{II} :Bg system shows another buffer region of one mole of H^+ per mole of

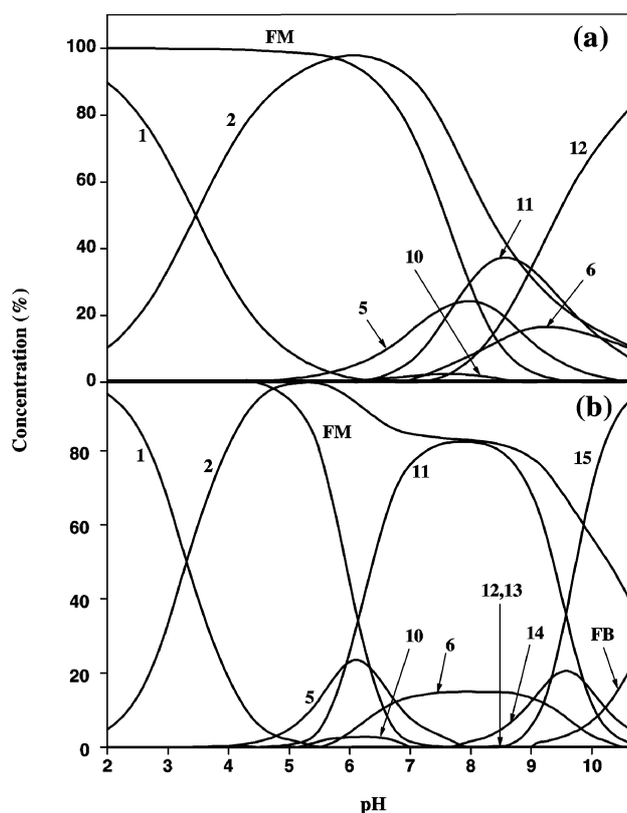


Figure 2. Speciation curves of (a) 1:1 and (b) 1:5 Cu^{II} :Bg systems: 1, H_2Bg^{2+} ; 2, HBg^+ ; 5, $\text{Cu}(\text{OH})^+$; 6, $\text{Cu}(\text{OH})_2$; 10, $\text{Cu}(\text{Bg})^{2+}$; 11, $\text{Cu}(\text{Bg})(\text{OH})^+$; 12, $\text{Cu}(\text{Bg}-\text{H})(\text{OH})$; 13, $\text{Cu}(\text{Bg})_2^{2+}$; 14, $\text{Cu}(\text{Bg}-\text{H})(\text{Bg})^+$; 15, $\text{Cu}(\text{Bg}-\text{H})_2$; FM = Free Cu^{II} , FB = Free Bg.

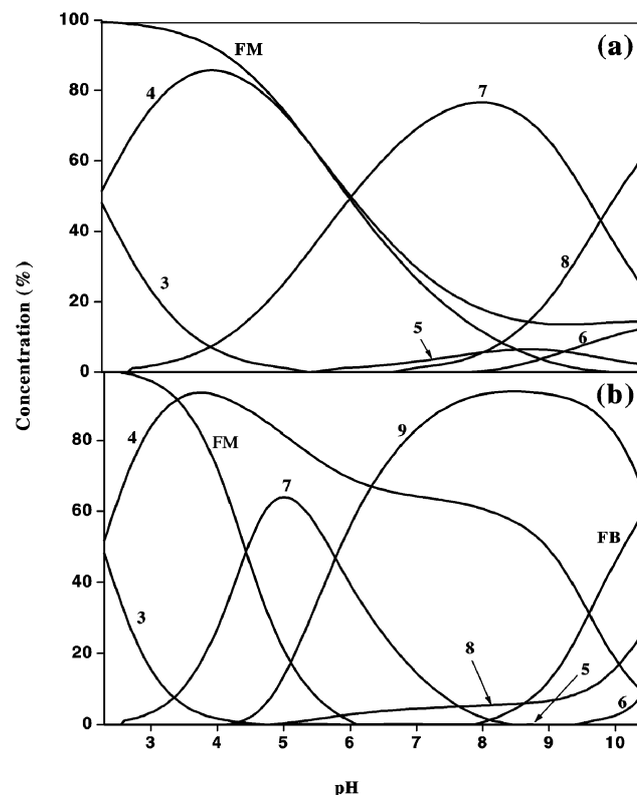
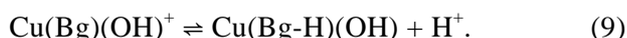
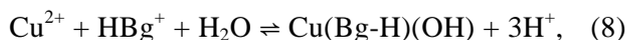


Figure 3. Speciation curves of (a) (1:1) and (b) (1:5) Cu^{II} :HG systems: 3, H_2G^+ ; 4, HG; 5, $\text{Cu}(\text{OH})^+$; 6, $\text{Cu}(\text{OH})_2$; 7, $\text{Cu}(\text{G})^+$; 8, $\text{Cu}(\text{G})(\text{OH})$; 9, $\text{Cu}(\text{G})_2$; FM = Free Cu^{II} , FB = Free G^- .

Cu^{II}, obviously due to the formation of biguanide (N² or N⁴) deprotonated ternary hydroxo complex, Cu(Bg-H)(OH), according to equilibria (8) and (9), the latter involves deprotonation of a coordinated biguanide molecule:



Titration cannot be continued beyond pH 8 due to appearance of turbidity, possibly because of precipitation of Cu(Bg-H)(OH) and/or Cu(OH)₂. Deprotonation constant of coordinated biguanide ligand in the complex, Cu(Bg)(OH)⁺, defined according to

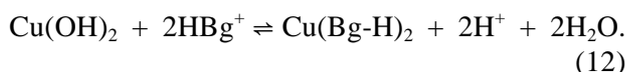
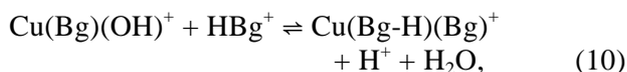
$$K_{\text{Cu}(\text{Bg})(\text{OH})}^{\text{H}} = \frac{[\text{Cu}(\text{Bg-H})(\text{OH})][\text{H}]}{[\text{Cu}(\text{Bg})(\text{OH})]}, \quad (9a)$$

may be calculated using the relation,

$$\log K_{\text{Cu}(\text{Bg})(\text{OH})}^{\text{H}} = \log \mathbf{b}_{1102} - \log \mathbf{b}_{1101}, \quad (9b)$$

where \mathbf{b}_{1101} and \mathbf{b}_{1102} are the formation constants of the complexes, Cu(Bg)(OH)⁺ and Cu(Bg-H)(OH) respectively. Charges are not shown in the mathematical expressions for simplicity.

The ternary hydroxo complex, Cu(Bg)(OH)⁺, is also the predominant Cu^{II} bearing species in the (1:5) Cu^{II}:Bg system in the pH range 5.5–7.5. As the speciation curves (figure 2b) imply, it is formed out of Cu(OH)⁺, mainly according to equilibrium (7) in addition to equilibrium (6). Above pH 7.5, this ternary hydroxo complex, Cu(Bg)(OH)⁺, and the dihydroxo species, Cu(OH)₂ gradually disappear due to the formation of the (1:2) Cu^{II}:Bg complexes, Cu(Bg-H)(Bg)⁺ and Cu(Bg-H)₂ according to equilibria (10–12) as the speciation curves (figure 2b) imply,



The Cu(Bg)₂²⁺ complex is formed in negligibly such small amounts that it cannot even be identified in the speciation curves. The biguanide deprotonated complex, Cu(Bg-H)₂, is the major Cu^{II} species at

pH > 9.5. Titration cannot be continued above pH 10, as the solution becomes turbid, possibly due to precipitation of the Cu(Bg-H)₂ complex. The deprotonation constant, $K_{\text{Cu}(\text{Bg})(\text{Bg-H})}^{\text{H}}$, of coordinated biguanide (Bg) in the 1:2 complex, Cu(Bg-H)(Bg)⁺, (equilibrium (11)) may be calculated using the relation,

$$\log K_{\text{Cu}(\text{Bg})(\text{Bg-H})}^{\text{H}} = \log \mathbf{b}_{1202} - \log \mathbf{b}_{1201} + \log \mathbf{b}_{1002} + \log \mathbf{b}_{1001}, \quad (11a)$$

where, \mathbf{b}_{1201} and \mathbf{b}_{1202} are the formation constants of the complexes, Cu(Bg-H)(Bg)⁺ and Cu(Bg-H)₂, and \mathbf{b}_{1001} and \mathbf{b}_{1002} are the first and second step hydrolysis constants of Cu(aq)²⁺ ion.

3.3 Binary Cu^{II}:HG equilibria

Complex formation of Cu^{II} with glycine in the 1:1 Cu^{II}:HG system starts at pH < 3.5. The major Cu^{II} species in the pH range (3.5–7) is the 1:1 binary complex, Cu(G)⁺, formed according to the equilibria (13) as the speciation curves (figure 3a) imply:



The hydroxo complexes, Cu(OH)⁺ and Cu(OH)₂ are practically non existent in this pH range. At pH > 5.5, the solution, containing the Cu(G)⁺ complex, shows another buffer region corresponding to one mole of H⁺ per mole of Cu²⁺, obviously due to deprotonation of a coordinated water molecule of the Cu(G)⁺ complex to form the ternary hydroxo complex, Cu(G)(OH) according to,



With increase of pH, the percentage of the dihydroxo species, Cu(OH)₂, increases gradually and the solution becomes turbid, possibly due to precipitation of either or both of Cu(OH)₂ and Cu(G)(OH). In the 1:5 Cu^{II}:HG system, the major Cu^{II} species are Cu(G)⁺ and Cu(G)₂. The binary and ternary hydroxo species are practically non existent (figure 3b).

3.4 1:1:1 Cu^{II}:Bg:HG ternary equilibria

Complex formation in the ternary 1:1:1 Cu^{II}:Bg:HG system (figure 4a) starts around pH ~ 2.5 with the formation of the Cu(G)⁺ complex (equilibrium (13)), which passes through a concentration maximum (~40%) around pH 5. Except traces

(<2%) of $\text{Cu}(\text{OH})^+$ and $\text{Cu}(\text{G})(\text{OH})$, no other binary or ternary hydroxo species are identifiable in the speciation curves above pH 4.5. The ternary complexes, $\text{Cu}(\text{G})(\text{Bg})^+$ and $\text{Cu}(\text{G})(\text{Bg-H})$, appear around pH ~ 4 and 4.5 according to equilibria (15) and (16) respectively,



The biguanide deprotonated ternary complex, $\text{Cu}(\text{G})(\text{Bg-H})$, appears as the major Cu^{II} containing species at pH > 6.5. Deprotonation constant, $K^{\text{H}}_{\text{Cu}(\text{G})(\text{Bg})}$, of coordinated biguanide (Bg) in the mixed ligand complex, $\text{Cu}(\text{G})(\text{Bg})^+$, (equilibrium (16)), may be calculated using the relation,

$$\log K^{\text{H}}_{\text{Cu}(\text{G})(\text{Bg})} = \log b_{1111} - \log b_{1110}, \quad (16a)$$

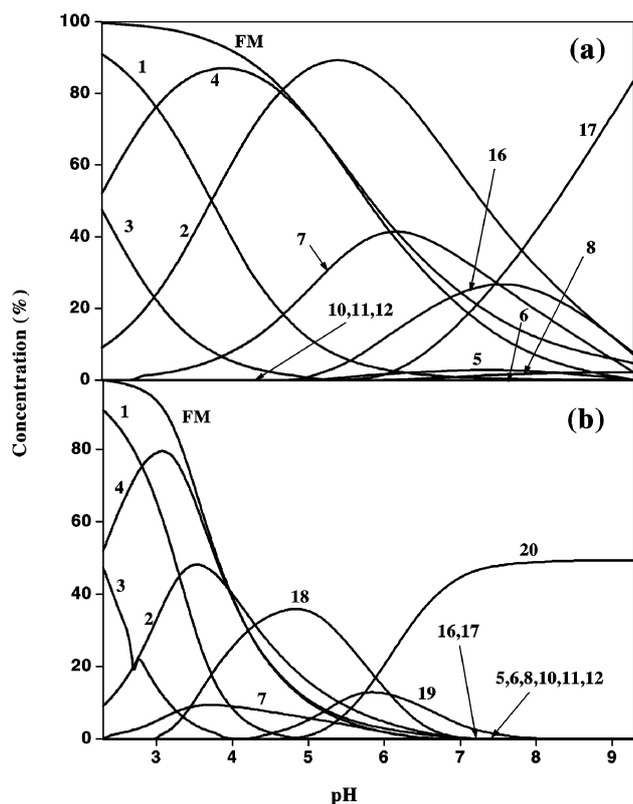


Figure 4. Speciation curves of (a) 1:1:1 and (b) 2:1:2 $\text{Cu}^{\text{II}}:\text{Bg}:\text{HG}$ systems: 1, H_2Bg^{2+} ; 2, HBg^+ ; 3, H_2G^+ ; 4, HG ; 5, $\text{Cu}(\text{OH})^+$; 6, $\text{Cu}(\text{OH})_2$; 7, $\text{Cu}(\text{G})^+$; 8, $\text{Cu}(\text{G})(\text{OH})$; 10, $\text{Cu}(\text{Bg})^{2+}$; 11, $\text{Cu}(\text{Bg})(\text{OH})^+$; 12, $\text{Cu}(\text{Bg-H})(\text{OH})$; 16, $\text{Cu}(\text{G})(\text{Bg})^+$; 17, $\text{Cu}(\text{G})(\text{Bg-H})$; 18, $\text{G}(\text{Cu})(\text{Bg})\text{Cu}(\text{G})^{2+}$; 19, $\text{G}(\text{Cu})(\text{Bg-H})\text{Cu}(\text{G})^+$; 20, $\text{G}(\text{Cu})(\text{Bg-2H})\text{Cu}(\text{G})$; FM = Free Cu^{II} .

where, b_{1110} and b_{1111} are the formation constants of the complexes, $\text{Cu}(\text{G})(\text{Bg})^+$ and $\text{Cu}(\text{G})(\text{Bg-H})$ respectively. Deprotonation constants of coordinated biguanide (Bg) in these ternary complexes are found to be of the order: $\text{Cu}(\text{G})(\text{Bg})^+ > \text{Cu}(\text{Bg})(\text{OH})^+ \gg \text{Cu}(\text{Bg})(\text{Bg-H})^+$, which indicates the Lewis basicity order of: $(\text{Bg-H})^- \gg \text{OH}^- > \text{G}^-$, when they are coordinated to Cu^{II} . This may be very useful information in relation to drug actions of biguanide derivatives and their metal complexes.

Electronic spectral I_{max} values of 1:1:1 $\text{Cu}^{\text{II}}:\text{Bg}:\text{HG}$ mixtures show successive blue shifts with rise of pH (figure 5). At pH ~ 5 the major Cu^{II} species is the $\text{Cu}(\text{G})^+$ complex (~50%) and $\text{Cu}^{2+}(\text{aq})$ ion (~50%). $\text{Cu}(\text{G})^+$ complex is likely to take up two molecules of solvent H_2O for completing a square planar geometry around Cu^{II} . The calculated I_{max} value of $[\text{Cu}(\text{COO}^-, \text{NH}_2)(\text{H}_2\text{O})_2]$ geometry³¹ as in the complex $\text{Cu}(\text{G})(\text{H}_2\text{O})_2^{2+}$ ion, is around 716 nm, which is in close agreement with the experimental value of 723 nm. Concentration of the ternary $\text{Cu}(\text{G})(\text{Bg})^+$ complex passes through a maximum around pH ~ 6, where the concentration of $\text{Cu}(\text{aq})^{2+}$ is only ~10% and those of $\text{Cu}(\text{G})(\text{H}_2\text{O})_2^{2+}$, $\text{Cu}(\text{G})(\text{Bg})^+$ and the biguanide deprotonated $\text{Cu}(\text{G})(\text{Bg-H})$ complex are around ~25% each. The estimated I_{max} values of $\text{Cu}(\text{aq})^{2+}$, square planar $[\text{Cu}(\text{COO}^-, \text{NH}_2)(\text{H}_2\text{O})_2]$ geometry as in $\text{Cu}(\text{G})(\text{H}_2\text{O})_2^{2+}$ $[\text{Cu}(\text{COO}^-, \text{NH}_2)(\text{NH}_2, \text{NH})]$ geometry as in $\text{Cu}(\text{G})(\text{Bg})^+$ and $[\text{Cu}(\text{COO}^-$,

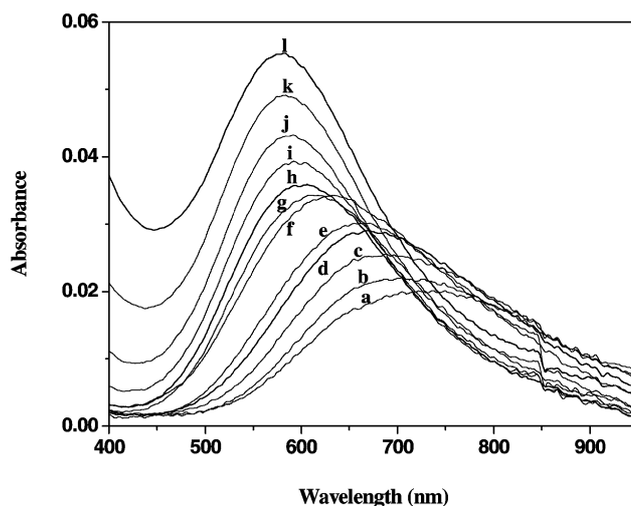
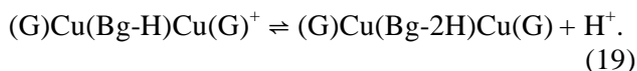
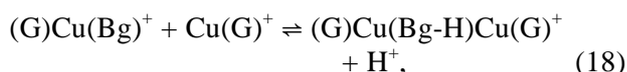


Figure 5. Electronic spectral curves of 1:1:1 $\text{Cu}^{\text{II}}:\text{Bg}:\text{HG}$ mixtures at different (pH): a (4.06), b (4.30), c (4.57), d (5.00), e (5.50), f (6.00), g (6.50), h (7.00), i (7.50), j (8.00), k (8.50), l (9.00). Ionic strength 0.1 mol dm^{-3} (NaNO_3).

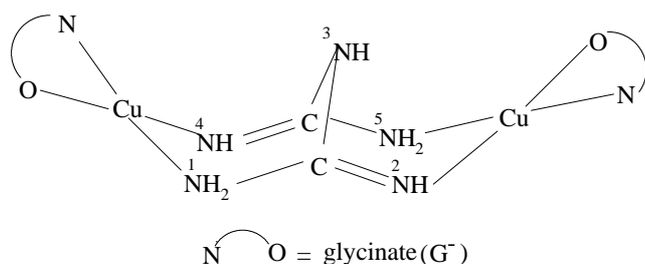
NH₂)(N⁻, NH₂)] geometry as in the Cu(G)(Bg-H), assuming a (>C=NH) moiety as equivalent to a (>C=O) moiety of a neutral peptide (-CONH) bond and (>C=N⁻) moiety as equivalent to a (-C(=O)N⁻) moiety of a deprotonated peptide bond, are around 830, 716, 645 and 577 nm respectively. The weighted average of these I_{\max} values at pH ~ 6 comes to ~646 nm. The experimental I_{\max} value of the solution at this pH is 668nm. With rise of pH, the I_{\max} value shows a further blue shift to 583 nm at pH ~ 7, indicating an increase of ligand field strength, as a coordinated neutral biguanide ligand as in the Cu(G)(Bg)⁺ complex is transformed to its monoanionic (Bg-H)⁻ species due to deprotonation at N² or N⁴ atom, to produce the complex, Cu(G)(Bg-H), according to equilibrium (16).

3.5 2 : 1 : 2 Cu^{II} : Bg : HG ternary equilibria

In the 2 : 1 : 2 Cu^{II} : Bg : HG system, an additional 1.5 mole of H⁺ per mole of Cu^{II} is released over those in the 1 : 1 : 1 system (figure 1). This indicates the release of at least 2 H⁺/Cu^{II} after the formation of the Cu(G)(Bg)⁺ complex. The only source of such proton release, of course, may be the formation of dinuclear mixed ligand complexes, (G)Cu(Bg)Cu(G)²⁺, (G)Cu(Bg-H)Cu(G), (G)Cu(Bg-2H)Cu(G), according to equilibria (17–19) as the speciation curves (figure 4b) imply:



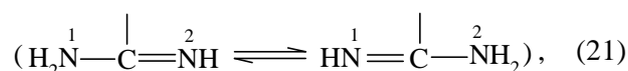
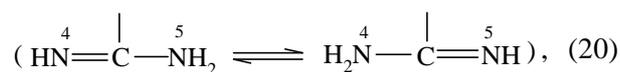
These 2 : 1 : 2 Cu^{II} : Bg : HG dinuclear complexes may be formed if biguanide in its neutral, monoan-



Scheme 2.

ionic and even in dianionic forms, Bg, (Bg-H)⁻ and (Bg-2H)²⁻ respectively, can act as a bridging ligand, that can exist in a puckered boat like conformation (scheme 2), providing two pairs of chelating (N,N-donor sites, viz., (N¹ and N⁴) and (N² and N⁵) for bridging two square planar complex Cu(G)(H₂O)₂⁺ ions, displacing the coordinated H₂O ligands from this complex.

Due to the possibility of tautomeric equilibria of the types (20) and (21) below,



within the coordinated biguanide molecule, the N⁴-C-N⁵ and N¹-C-N² sequences may form two separate triangular inclined planes, e.g. N⁴-C(-N³)-N⁵ and N¹-C(-N³)-N², meeting at the common point at N³ (scheme 2). Such a conformation may function as a bridging quadridentate ligand providing two sets of bidentate (N,N) chelating sites to the Cu^{II} atoms of two different Cu(G)⁺ complexes, as described above, thereby forming the dinuclear complex, (G)Cu(Bg)Cu(G)²⁺. With rise of pH above 4.5 the biguanide deprotonated dinuclear mixed ligand complexes appear in the equilibrium through release of protons from the coordinated =N²H and =N⁴H sites, (18) and (19).

4. Conclusions

Equilibrium study of the complex formation of Cu^{II} with biguanide (Bg) in the absence and in the presence of the simplest amino acid glycine (HG) in different molar proportions of the metal ion and the ligands revealed that biguanide may coordinate Cu^{II} as a neutral (N,N) and anionic (N,N⁻) bidentate chelating ligand in mononuclear binary and ternary complexes, using its N¹ and N⁴ or N² or N⁵ atoms. In forming 2 : 1 : 2 Cu^{II} : Bg : HG dinuclear mixed ligand complexes, the biguanide ligand species may exist in a puckered boat-like conformation, serving as neutral, monoanionic and dianionic bridging tetradentate ligand, chelating one Cu^{II} ion with N¹ and N⁴ atoms and the other Cu^{II} ion with N² and N⁵ atoms. Deprotonation constants of Cu(Bg)(OH)⁺, Cu(Bg)(G)⁺, Cu(Bg)(Bg-H)⁺ complexes indicate the

Lewis basicity order of $(\text{Bg-H})^- \gg \text{OH}^- > \text{G}^-$ for these coordinated ligand species.

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