## **40. Models for Copper-Dioxygen Complexes: the Chemistry of Copper(I1) with Some Planar Tridentate Nitrogen Ligands**

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The solution chemistry of **Cu(I1)** with a series of five planar tridentate nitrogen ligands, 2,6-bis(benzimidazol-2-y1)pyridine (bzimpy, **l), 2,6-bis(l-methylbenzimidazol-2-yl)pyridine** (mbzimpy, **2) 2,6-bis(benzothiazol-2-y1)**  pyridine (bzthpy, **3), 2,6-bis(benzoxazol-2-yl)pyridine** (bzoxpy, **4),** and 2,2',6',2"-terpyridyl (terpy, **5)** is reported. Electronic and EPR spectra are consistent with the complexes  $\text{[CuL]}^{2+}$  having essentially tetragonal structure in solution, with the fourth coordination site in the plane of the ligand occupied by solvent. bzthpy and bzoxpy show smaller ligand-field splittings than bzimpy, mbzimpy, and terpy, and are easily decomplexed from the copper. Substitution of the coordinated solvent molecule in the plane of the ligand is observed with  $Cl^-$  and  $OH^-$  (provided that the ligand has no acidic protons) for all ligands except terpy. The reaction between  $[Cu(mbzimpy)]^{2+}$  and imidazole has been studied by potentiometric titration in MeCN/H<sub>2</sub>O 1:1 and shows strong binding of the imidazole in the plane ( $log K = 4.5$  at  $25^{\circ}$ ), and also the formation of an imidazolate-bridged dinuclear species.

Introduction. - There is currently considerable interest in the chemistry of the oxygen transporting protein oxyhemocyanin [ 11 and related systems such as tyrosinase **[2]** and the laccases **[3]** which possess an active site of type 111 where the dioxygen moiety is thought to be bound to **Cu(I1)** as in **I** below:



This structure has been proposed on the basis of the extended X-ray absorption fine structure **(EXAFS)** measurements [4], electronic **[5],** and resonance *Raman* **[6]** spectroscopy. However, some discrepancies between EXAFS results have led to the suggestion that three histidines may be bound to Cu(1I) in the oxy form of hemocyanin **[7],** and a recent crystal-structure determination of deoxyhemocyanin shows the Cu(1) to be bound to three histidines in the deoxy form [8]. There is, however, general agreement as to the formulation of these complexes as  $\mu$ -peroxodicopper(II) complexes, and a synthetic complex of this type has recently been characterised by X-ray crystallography [9].

In the search for possible precursors to stable copper-dioxygen complexes, our attention was attracted to planar tridentate nitrogen ligands, since such ligands show similari-

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ties to the known natural systems, and, in view of the tendency of Cu(I1) to display a distorted octahedral geometry with four short equatorial and two long axial bonds [lo], Cu(I1) complexed by a planar tridentate system is likely to show a strong affinity for ligands in the fourth equatorial binding site, and, under these conditions, the binding of 0, in the form of peroxide at this site might be envisaged [l 11.

The following criteria were established for the choice of ligand: *i)* the ligand should be constrained to be planar; *ii)* the coordinating N-atoms should possess some unsaturated character in order to allow the stabilisation of Cu(1) [12]; *iii*) it should be possible to modify the ligands slightly in order to vary the ligand field strength, and thereby influence the copper redox potential. The ligands shown below, **2,6-bis(benzimidazol-2-yl)pyridine**  (bzimpy, **l), 2,6-bis(l-methylbenzimidazol-2-yl)pyridine** (mbzimpy, **2),** 2,6-bis(benzothiazol-2-y1)pyridine (bzthpy, **3),** and **2,6-bis(benzoxazol-2-yl)pyridine** (bzoxpy, **4)** satisfy these criteria and may readily be synthesised in high yield using the modified *Philips*  reaction [13].



**A** complex of bzimpy with Ni(I1) [14] and a few complexes of bzthpy with Mn(II), Fe(II), Co(II), Ni(II) [15], and Cu(II) [16] have been reported in the literature, but no extensive studies have been made. During the preparation of this manuscript, however, we were informed of a very recent study reporting the preparation of some complexes of bzimpy and mbzimpy with  $Cu(II)$  and  $Zn(II)$ , and the crystal structures of  $[Cu(bzimpy)(MeCN)](ClO<sub>4</sub>)<sub>2</sub>$  and  $[Cu(mbzimpy)<sub>2</sub>(ClO<sub>4</sub>)<sub>2</sub>]\cdot H<sub>2</sub>O$  [17], and a paper reporting the chemistry of bzimpy, mbzimpy, and bzthpy with  $Fe(II)$  and  $Fe(III)$  has also been published [18].

In this paper, we report on the solution chemistry of ligands **1-4** with Cu(I1) and compare it with the well known tridentate N ligand 2,2',6',2"-terpyridyl (terpy, **5)** in order to: *i)* compare the bonding properties of the five ligands, *ii)* establish the coordination of the Cu(I1) in solution, and *iii)* investigate to what extent these complexes do bind another ligand in the plane of the three N-atoms. We consider a thorough understanding of the properties of the  $[CuL]^{2+}$  species to be essential for the assessment of the likely stability of a Cu(I1)-peroxo complex. The chemistry of the most favorable of these ligands, mbzimpy, with Cu(I) will be reported later [19].

**Experimental.** ~ Unless otherwise stated, solvents and starting materials were purchased from *FIuka* (Switzerland) and used without further purification.

*Preparution qfligunds.* bzimpy **(1)** and bzthpy **(3)** were prepared according to the procedure described in [13], and this method was also applied to the synthesis of bzoxpy **(4)** obtained in good yield *(5* 1 %). The yields are greatly increased, if mechanical stirring is applied during the reaction. All ligands were fully characterized by their IR, <sup>1</sup>H-NMR, and EI-MS  $(M^+$ : bzimpy, 311; bzthpy, 345; bzoxpy, 313). mbzimpy **(2)** was prepared under N<sub>2</sub> using *Schlenek* techniques. Tetramethylurea (TMU) and pentane were previously distilled from NaH under N,. Compound **1** (4.0 **g,** 12.9 mmol) was dissolved in 30 ml of dry TMU. NaH (oil dispersion 58 %; 1.62 g, 38.8 mmol) was washed with  $5 \times 10$  ml of dry pentane and then suspended in 30 ml of dry TMU. This suspension was added to the well-stirred soln. of 1 at 0°. The mixture was allowed to stand at r.t., then heated to 35° for 1 h. The mixture was cooled to  $0^\circ$ , 5.51 g (38.8 mmol) of MeI added, heated to  $40^\circ$  for 3 h, then hydrolysed with H<sub>2</sub>O, the pH adjusted to 13 with 1M NaOH, and 3.86 g (11.4 mmol) of white 2 were separated by filtration and dried at  $10^{-2}$  Torr/60° for 12 h (yield 89%). The product was characterized by its IR ( $\tilde{v}$ (NH) has disappeared and three new bands appear at 2910, 2840, and 1415 cm<sup>-1</sup> characteristic for the Me groups), <sup>1</sup>H-NMR (in CDCI<sub>3</sub>, a *singlet* at 4.25 ppm (CH<sub>3</sub>)), and its EI-MS (339 *(M')).* 

*Preparation of Perchlorate Salts of the Complexes of Cu(II) with* **14.** For **1** and **2,** 3.0 mmol of ligand were dissolved in 50 ml of EtOH at 50° and then a soln. of 1.11 g (3.0 mmol) of  $\left[Cu(H<sub>2</sub>O)<sub>6</sub>\right](ClO<sub>4</sub>)$ , in the same solvent was added. The complex of 2 precipitated immediately and was recrystallized from DMF/EtOH to give 1.97 g (2.63) mmol) of deep green [Cu(mbzimpy)](ClO<sub>4</sub>)<sub>2</sub> 2 DMF (6; yield 88%). The complex with 1 was crystallized by addition of dioxane. We obtained 1.51 g (2.40 mmol) of light green [Cu(bzimpy)](ClO<sub>4</sub>)<sub>2</sub>·3 H<sub>2</sub>O (7; yield 80%). The same procedure was applied for **3** and **4,** but their insolubility in almost all solvents required working with a two-phase system. A suspension of 1.6 mmol of ligand in MeCN at *50"* was mixed with a stoichiometric quantity of  $[Cu(H<sub>2</sub>O)<sub>6</sub>](ClO<sub>4</sub>)$ , and the mixture was stirred, until a clear deep green soln. was obtained. The complex was then crystallized by slow evaporation *in uacuo.* We obtained 0.66 **g** (0.98 mmol) of pale green [Cu(bzoxpy)]-  $(CIO_4)_2 \cdot H_2O \cdot 2$  MeCN **(8**; yield 61%) and 0.68 g (1.09 mmol) of green  $[Cu(bzthpy)](ClO_4)_2 \cdot H_2O$  **(9**; yield 68%). These four perchlorate salts were characterized by their IR and gave satisfactory elemental analysis (C, H, Cu, N).

*Preparation of Chloro-Perchlorate Salts of the Cu(II) Complexes of* 24. For ligand 2,0.2 **g** (0.27 mmol) **of** *6*  was dissolved in 8 ml of DMF, and 0.045 g (0.27 mmol) of Et<sub>4</sub>NCl in a minimum of the same solvent was added. The product was precipitated by addition of 10 ml of Et<sub>2</sub>O, separated by filtration, and washed with two portions of CH<sub>2</sub>Cl<sub>2</sub> (to extract Et<sub>4</sub>NClO<sub>4</sub>). After recrystallization in DMF/Et<sub>2</sub>O, we obtained 0.11 g (0.197 mmol) of light green  $\text{[Cu(mbzimpy)Cl}(CIO_4) \cdot \text{H}_2\text{O}$  (10; yield 73%).  $\tilde{v}(Cu-CI) = 320 \text{ cm}^{-1}$ . For 3 and 4, the same procedure was applied with MeCN instead of DMF, yielding products which crystallized directly from the mixture without addition of Et<sub>2</sub>O. We obtained 73% of green [Cu(bzoxpy)Cl](ClO<sub>4</sub>) (11) and 75% of yellowish green  $[Cu(bzthpy)Cl]$ (ClO<sub>4</sub>) (12).  $\tilde{v}$ (Cu–Cl) was observed at 330 and 315 cm<sup>-1</sup> for 11 and 12, respectively. These three Cu(I1) chloro-perchlorate salts were characterized by IR and gave satisfactory elemental analyses.

*Caution;* perchlorate salts with ore. ligands are potentially explosive and should be handled with the necessary precautions [20].

*Preparation of Dichloro Salts of the Cu(II) Complexes of* **1** *and* **2.** A soln. of 0.31 I g (1 mmol) of **1** in EtOH (20 ml) was mixed with 0.17 g (1 mmol) of CuCl<sub>2</sub>:  $2 H_2O$  in the same solvent. The resulting precipitate was separated by filtration and recrystallized from 300 ml hot DMF. The formula [Cu(bzimpy)]CI, DMF **(13)** was established by IR, elemental analysis, and an X-ray crystal-structure determination [21]. The light green complex [Cu(mbzimpy)]Cl<sub>2</sub>. H<sub>2</sub>O (14) was obtained in the same way by using MeCN instead of EtOH and was recrystallized from MeCN/Et<sub>2</sub>O (yield 83%). This complex was characterized by its **IR** ( $\tilde{v}$ (Cu-Cl) of 295 and 270 cm<sup>-1</sup>) and gave a satisfactory elemental analysis.

We attempted to synthesize the Cu(I1) dichloro complexes of **3** and **4,** but have not succeeded in obtaining reproducible compositions for the substances isolated (IR and elemental analysis differed from one batch to another).

*Preparation of [Cu(mbzimpy) (imidazolate) Cu(mbzimpy)](ClO<sub>4</sub>)<sub>3</sub> · H<sub>2</sub>O (15). To a soln. of 0.3 g (0.4 mmol)* of  $[Cu(mbzimp)](ClO<sub>4</sub>)<sub>2</sub>·2 DMF (6)$  in 18 ml of MeCN, 1 ml (0.2 mmol) of a 0.2M soln. of imidazole in MeCN and 1 ml (0.2 mmol) of a 0.2 $\mu$  soln. of Et<sub>3</sub>N in MeCN were added slowly. The soln. rapidly turned turquoise; 30 ml of MeCN/EtOH 1:1 were added and, after standing at  $-25^{\circ}$  for 3 h, the small turquoise crystals were separated by filtration. The filtrate was concentrated under vacuum, and further addition of MeCN/EtOH and refrigeration gave a second crop of crystals. After drying *in uacuo,* we obtained 0.185 g (0.156 mmol, 78 *YO)* of blue green crystals which gave a correct analysis for  $\left[\text{Cu(mbzimby)}\right](\text{imidazolate})\text{Cu(mbzimpy)}\left(\text{ClO}_4\right)_3 \cdot \text{H}_2\text{O}$  (15).

*Physical Measurements.* **UVjVIS** Spectra: in soh. with a *Perkin Elmer Lambda 5* spectrophotometer at 20' using quartz cells of 1, 0.1, and 0.01 cm path length. IR: KBr pellets, with a *Perkin Elmer IR 597* spectrophotometer. 'H-NMR: on a *Varian EM360A* (60 MHz), a *Varian XL* 200 (200 MHz), or a *Bruker WH360* (360 MHz) spectrometer. EI-MS (70 eV): *VG* 7000 *E* and *Finnigan 4000.* EPR: frozen solns. at 77 K, at 9 **GHz** on a *Bruker ER 200D-SRC.* Cyclic voltammograms were recorded using a *Tacussel PRGE-DEC* potentiostat connected to a function generator and a **XY** plotter. A 3-electrode system consisting of a stationary Pt disk working electrode, a Pt counter electrode, and a non-aqueous  $Ag/Ag^+$  reference electrode was used.  $Bu_4NClO_4$  **(0.1m)** or 0.1m  $Bu_4NPF_6$ served as inert electrolytes. Propylene carbonate  $[22]$  and MeNO<sub>2</sub>  $[23]$  were purified according to the literature. MeCN was distilled from  $P_2O_5$  and then passed through an *Alox* column (pH 9.5). All other solvents used were distilled under reduced pressure  $(10^{-2}$  Torr): DMSO from CaH<sub>2</sub>, TMU from NaH and DMF from *Alox*. The reference potential  $(E^{\circ} = +0.66 \text{ V} \text{ vs. NHE})$  was standardized against the known complex  $\text{[Ru(bipy),NCO4)}_2$  [24a].

The scan speed used was 0.2 **Vjs,** and voltammograms were analyzed according to established procedures **[24b].**  Conductimetric results were obtained with a *Metrohm EA240* cell  $(f = 0.61 \text{ cm}^{-1})$  immerged in a thermostatted vessel at 25". Conductivity measurements were made with a *Metrohm E527 Wheatstone* bridge; the conductivity was measured after each addition of the soln. of the anion from a syringe. The results were corrected for dilution. Potentiometric measurements were made in a closed cell thermostatted at  $25 \pm 0.1^{\circ}$  using a *Metrohm* combined electrode and a *Metrohm E500* digital pH-meter. The conditions for a typical titration were as follows: to 25 ml **of**  MeCN/H<sub>2</sub>O 1:1  $(v/v)$  were added 56.1 mg (7.5·10<sup>-5</sup> mol) of 6,750 µl of N-methylimidazole (0.1m in MeCN/H<sub>2</sub>O), and 750  $\mu$  of HClO<sub>4</sub> (0.1 M in MeCN/H<sub>2</sub>O), and the ionic strength was adjusted to 0.05M with KNO<sub>3</sub>. This soln. was placed in the cell and the pH electrode introduced. The soln. was titrated with a soln. of 0.02M KOH in MeCN/H<sub>2</sub>O whose ionic strength was adjusted to 0.05<sub>M</sub> with KNO<sub>3</sub>. Acid and base solns. were standardized against *Merck* titrisol solns. The measured pH values  $(\pm 0.02 \text{ units})$  were fitted to pK values using a non-linear least-squares program [25]. Elemental analyses (C, H, CI, N) were performed by Dr. *H. Eder* **of** the Microchemical Laboratory of the University **of** Geneva. Cu was determined by atomic absorption *(Pye Unicum SP9)* after acidic oxidative mineralisation of the complex.

**Results and Discussion.** - 1. *Synthesis and Properties of Ligands.* We have followed the synthetic procedure developed by *Addison* and coworkers [13] to obtain bzimpy **(1)** and bzthpy **(3),** and we have extended it to the synthesis of bzoxpy **(4).** The double methylation of **1** was easily performed by deprotonation with 2 equiv. of NaH in dry TMU followed by alkylation with Me1 to give mbzimpy **(2)** in 89 % yield. The ligands **1** and **2**  are sparingly soluble in EtOH, insoluble in H,O, but very soluble in polar aprotic solvents like DMF or DMSO. Ligands **3** and **4** are very poorly soluble in all solvents except hot CHCl<sub>3</sub>. The H-atoms of the benzo moiety of 1 give a symmetrical  $AA'BB'$ -spin system in the 'H-NMR spectrum which becomes a very complicated ABCD system for the dimethyl-substituted derivative **2. A** first-order interpretation of the 360-MHz 'H-NMR spectra (CDC1,) of **3** and **4** is fully satisfactory with two *triplets* and two *doublets* for the benzo H-atoms between *7.5* and 8.0 ppm and a AB,-spin system for the pyridine H-atoms 8.1-8.5 ppm *us.* TMS. The UV spectra of these ligands in propylene carbonate solution consist of one broad envelope of  $\pi \rightarrow \pi^*$  transitions centered around 31250 cm<sup>-1</sup> with  $\varepsilon_{\text{max}} = 30000 \text{ M}^{-1} \cdot \text{cm}^{-1}$ . The comparison with the spectrum of terpy (5) in the same solvent  $(36000 \text{ cm}^{-1}, \varepsilon_{\text{max}} = 20000 \text{ m}^{-1} \cdot \text{cm}^{-1})$  shows a red-shift for our ligands of roughly 4700 cm<sup>-1</sup>. This transition is assigned to  $\pi_1 \rightarrow \pi^*$  according to *Nakamoto* [26].

2. Properties of *Isolated Complexes with* 1–4. The proposed formulae of the Cu(II) complexes **614** were consistent with elemental analysis based on C, H, Cu, CI, and N. In all cases, we isolated complexes with one ligand per Cu(I1) and two anions. The IR spectra showed characteristic ligand vibrations ( $C=C=N$  stretching) in the 1620–1500 $cm^{-1}$  range which are shifted toward high energy (about 15-25 cm<sup>-1</sup>) upon complexation to Cu(II). The ClO<sub>4</sub> anion showed two IR bands at 625 and 1095 cm<sup>-1</sup> which are symmetric, when it was not coordinated. Only complex 11 shows a splitting of  $ClO<sub>a</sub>$ vibrations compatible with coordination to  $Cu(II)$ , which is confirmed by a new absorption at 375 cm<sup>-1</sup> attributed to the Cu $\sim$ OClO<sub>3</sub> stretching frequency. The stretching vibrations Cu-N(ligand) are rather weak and appear at *ca*. 305 cm<sup>-1</sup> for 6 and 7, and are displaced to lower energy for **8** (250 cm-') where benzoxazoles replace benzimidazoles. This is an indication of the weaker coordination of **4** to Cu(I1) compared with **1** or **2. A** square-pyramidal structure for complex **13** was determined by X-ray diffraction [21], and the powder EPR spectrum at 298 K of the analogous complex with **2, 14,** showed a  $(d_{x^2-y^2})^1$  ground state with  $g_{\parallel}=2.22 > g_{\perp}=2.10 > 2.0$  indicating a similar structure [101[271.

3. *Chemistry of the Complexes in Aprotic Solution.* 3.1.  $\Gamma$ *Cu(L)* $\Gamma$ *(ClO<sub>4</sub>)*, **(L** = **1–4**). The complexes **6** and **7** are soluble in polar aprotic solvents like DMF, DMSO, propylene carbonate, MeCN, but **4** and **3** are decomplexed from Cu(I1) in strong donor solvents (DMSO or DMF) resulting in precipitation of the poorly soluble free ligands which were isolated and characterized by IR. Complexes **8** and **9** have, therefore, only been studied in MeCN or propylene carbonate, and this last solvent was used for comparison of the different ligands.

The complexes **69** behave as 2 : 1 electrolytes in polar aprotic solution *(Table 1)* with molar conductivities corresponding to the highest values proposed by *Geary* **[28]** (220- 340 in MeCN and 130-170 in DMF). However, our values are consistent with the results of *Nelson et al.* [29] for  $\lceil Cu(L), \rceil (ClQ_4)$ , in MeCN (250–270  $\Omega^{-1}$  mol<sup>-1</sup>  $\cdot$  cm<sup>2</sup>, where L is a *Schiff* base derived from 2,6-diacetylpyridine whose greater size is presumably responsible for the lower value).

Complexes		Propylene carbonate	<b>DMSO</b>	Other solvents
[Cu(mbzimp)](ClO <sub>4</sub> )	(6)	83	103	196 (DMF)
$[Cu(bzimpy)](ClO4)2$	(7)	81	107	
$[Cu(bzospy)](ClO4)_{2}$	(8)	80		360 (MeCN)
$[Cu(bzthpy)](ClO4)2$	(9)	77		340 (MeCN)
[Cu(mbzimpy)Cl](ClO <sub>4</sub> )	(10)	27	53	
[Cu(bzospy)Cl](ClO <sub>4</sub> )	(11)	32		
[Cu(bzthpy)Cl](ClO <sub>4</sub> )	(12)	32		
$Bu_4NClO_4$		41	51	
$Bu_4NCl$		35	45	
		$\sim$		

Table 1. *Conductivity Data at* 25<sup>oa</sup>)

<sup>a</sup>) Molar conductivities are given in  $\Omega^{-1}$  mol<sup>-1</sup> cm<sup>2</sup> for  $10^{-3}$  m soln. at 25°. Estimated experimental errors: *\*5%.* 

Compound	Assignment			
	$\pi_1 \rightarrow \pi^*$		d-d	
bzimpy $(1)$		30 400 (30 000)		
$1 + Zn^{2+}$	28 330 (25 400)	32 360 (25 600)		
$1 + Cu^{2+}$	27780 (19500)	31850 (27000)	14470 (88)	
mbzimpy $(2)$		31 150 (29 000)		
$2 + Zn^{2+}$	28 330 (21 500)	32 570 (23 500)		
$2 + Cu^{2+}$	27470 (16000)	31 850 (23 500)	14770 (98)	
bzthpy $(3)$		30490 (28000)		
$3 + Zn^{2+}$	27 550 (28 800)	32 360 (21 200)		
$3 + Cu^{2+}$	27 100 (21 200)	31450 (14100)	13210 (77)	
$b$ zoxpy $(4)$		31 450 (30 000)		
$4 + Zn^{2+}$	28 900 (21 000)	33 445 (20000)		
$4 + Cu^{2+}$	28 090 (20 000)	33 110 (22000)	13 120 (55)	
terpy $(5)$		35970 (18800)		
$5 + Zn^{2+}$	30210 (17300)	35 340 (15000)		
$5 + Cu^{2+}$	29 326 (15000)	34970 (13500)	15080 (92)	

Table 2. *Electronic Spectra in Propylene Carbonate Solution at 20"3* 

<sup>a</sup>) Energies are given for the maximum of the band envelope in cm<sup>-1</sup> and  $\varepsilon$  (in parentheses) in  $M^{-1}$  cm<sup>-1</sup>. All values refer to 10<sup>-3</sup> M propylene carbonate solution obtained either directly from the isolated complexes for  $Cu^{2+}$  or formed *in situ* ( $Zn^{2+}$ ) by mixing stoichiometric quantities of ligand and  $[Zn(H_2O)_6](ClO_4)_2$ .





The electronic spectra of the complexes **6-9** in solution present two distinct absorption domains. Between  $12\,500$  and  $15\,500$  cm<sup>-1</sup>, we find the envelopes of d-d transitions  $(e = 50-100 \text{ M}^{-1} \cdot \text{cm}^{-1})$  and between 25 000 and 37 000 cm<sup>-1</sup> the  $\pi \rightarrow \pi^*$  transitions of the aromatic ligands  $(\varepsilon = 15000-25000 \text{ M}^{-1} \cdot \text{cm}^{-1})$ . According to *Nakamoto* [26], terpy (5) shows two bands in the UV region attributed to two  $\pi \rightarrow \pi^*$  transitions:  $\pi_1 \rightarrow \pi^*$  at *ca.* 35 700 cm<sup>-1</sup> and  $\pi_2 \rightarrow \pi^*$  at 41 700 cm<sup>-1</sup>. Upon complexation to  $\mathbb{Z}n^{2+}$  or Ni<sup>2+</sup> (or Mg<sup>2+</sup> with 2,2-bipyridyl [30]), the  $\pi_1 \rightarrow \pi^*$  is split into two components, with  $\pi_1^b \rightarrow \pi^*$  at *ca.* 30300 cm<sup>-1</sup> and  $\pi_1^2 \rightarrow \pi^*$  at 35300 cm<sup>-1</sup> for [Zn(terpy)]Cl<sub>2</sub>. These results are paralleled by the ligands 1-4 which show a similar splitting of the  $\pi_1 \rightarrow \pi^*$  transition upon complexation with  $Zn^{2+}$  or  $Cu^{2+}$  in PC *(Table 2 and Fig. I)*. We have found this splitting to be a useful confirmation of the complexation of ligands **14.** 

Comparison of the d-d transition energies in propylene carbonates shows the benzimidazole ligands **1** and **2** to have comparable ligand-field strength to terpy, while **3** and **4**  are appreciably weaker. Since the conductivity measurements suggest that the  $ClO<sub>a</sub>$ anions are not coordinated, there are two axial and one equatorial binding sites free on the Cu(II)  $(cf.$  **II)**, and solvent dependence of the spectra is expected and is, indeed, observed.



For  $[Cu(2)]^{2+}$  the d-d maxima are 13720 (DMSO) < 14100 (DMF) < 14770 (propylene carbonate)  $<$  15385 (MeCN)  $<$  15870 cm<sup>-1</sup> (MeNO<sub>2</sub>). This order corresponds to increasing ligand-field strength in the spectrochemical series  $(f(\text{MeCN}) = 1.22$ ,  $f(DMF) = 0.98$ , and  $f(DMSO) = 0.91$  [31]), and decreasing donor character as established by *Gutmann* [32]. These results indicate strong interaction of the solvent with the vacant coordination sites. Similar solvent shifts on going from propylene carbonate to DMSO have been reported for Ni(II), Cr(III), and Cu(I1) [22].

The **EPR** spectra of frozen propylene-carbonate solutions show pseudo-axial features with  $g_1 > g_2 > 2.0$  *(Table 3)* which are typical of a  $(d_{x_1}^2, d_x^2)$  ground state [10]. This implies that planarity of ligands is maintained in solution with the three coplanar heterocyclic

Complexes	$\mathbf{g}_{\parallel}$	$g_{\perp}$	$A_{\parallel}$	$E_{\rm p/2}$ [v]	$E_{\rm pc} - E_{\rm pa}$ [mV]
6	2.24	2.03	182	$+0.38$	
7	2.26	2.04	172	$+0.42$	1000
8	2.29	2.06	151	$+0.85$	500
	2.35	2.06	149	$+0.76$	500
9	2.28	2.06	142	$+0.73$	600
				$+0.63$	700
$6 + 1$ equiv. $Cl^{-}$	2.23	2.03		$+0.37$	180/800
		2.06			
$7 + 1$ equiv. $Cl^-$	2.23	2.03	169	$+0.27$	380
		2.06			
$8 + 1$ equiv. Cl <sup>-</sup>	2.25	2.05	153	$+0.69$	120/600
$9 + 1$ equiv. $Cl^-$	2.26	2.05	140	$+0.58$	140/700

Table 3. *EPR Parameters* (in frozen propylene carbonate solution at 77 K) *and Electrochemical Reduction Potentials* (in propylene carbonate solution)")

<sup>a</sup>) EPR parameters: estimated errors of  $\pm 0.01$  on g<sub>ll</sub>,  $\pm 0.01$  on g<sub>L</sub>, and  $\pm 3 \cdot 10^{-4}$  cm<sup>-1</sup> for A<sub>ll</sub> which is given in  $10^{-4}$  cm<sup>-1</sup>. Electrochemical potentials for Cu(II)/Cu(I) are given *vs*. NHE for the half reduction peak at 20°. Estimated error on  $E_{p/2}$  is  $\pm 0.01$  V.



 $in 5 \cdot 10^{-3}$ M frozen propylene carbonate solution at 77 K

N-atoms bound to Cu(II) destabilising the  $d_{x-y^2}$  orbital. It has been shown that, for any given Cu(II) coordination compound, its position on a plot of  $g_{\parallel}$  against  $A_{\parallel}$  is governed by *a)* the nature of the donor atoms in the tetragonal plane, with increasing  $g_{\parallel}$  values  $S \le N \le O$ , *b*) the formal charge carried by the four donor atoms plus the metal, and *c*) the dihedral angle [33]. The values for the four complexes  $6-9$  lie near to a N<sub>4</sub> donor set on the plot and are fully compatible with a  $\left[\text{Cu(N)}\right]$  (solvent)]<sup>2+</sup> unit in pseudo-tetragonal plane. The existence of such a unit is confirmed by the isolation of  $[Cu(1)(MeCN)](ClO<sub>4</sub>)<sub>2</sub>$ from EtOH/MeCN [17]. Hyperfine structure is visible for 6 in the  $g_{\perp}$  domain due to N superhyperfine coupling  $(A \approx 14.10^{-4} \text{ cm}^{-1})$ , confirming the coordination of three Natoms to Cu(I1) *(Fig.* 2). Complex **8** (with the ligand bzoxpy **(4))** shows two axial species, one with  $g_{\parallel} = 2.35$  and the other with  $g_{\parallel} = 2.29$  which have been tentatively assigned to two differently solvated pseudo-axial species as proposed by *Reed et al.* [34] in a similar case, and complex **9** (ligand bzthpy **(3))** shows a similar effect in MeCN solution. There is a slight solvent dependence of the **EPR** parameters.

Cyclic voltammetry studies of complexes **69** show highly irreversible Cu(II)/Cu(I) redox waves, and *Table 3* reports the half-peak reduction potentials  $E_{p/2}$  in propylene carbonate solution. The apparent ease of reduction of the bzoxpy **(4)** and bzthpy **(3)**  complexes **8** and **9,** respectively, is probably not significant, since we have evidence that **4**  and 3 dissociate from Cu(I) [19], and, in propylene carbonate solution, the Cu<sub>solv</sub>,  $Cu_{solv}^+$ ,  $Cu_{solv}^+$ couple is even more positive  $(E_{p/2} = +1.03 \text{ V})$ ; furthermore, **8** and **9** show two reduction peaks. For complexes **6** and **7,** the reduction peaks are strongly solvent dependent: for *6*  the potentials *(vs. NHE)* vary in the order  $+0.25$  (DMSO)  $< +0.29$  (DMF)  $< +0.38$  (PC)  $< +0.60$  V (MeNO<sub>2</sub>). This order correlates with decreasing solvent donor strength [32]. If *Addison's* formula [12] for  $E^{\circ}$  is applied for Cu(II) with three heterocyclic N donors, the predicted value is  $+0.33$  V, in reasonable agreement with our results given that the donor number for H,O (18) is intermediate between DMF (26.6) and propylene carbonate (15.1).

3.2. *Reactivity of*  $\int Cu(L)/\langle ClO_4 \rangle$ , (6–9) *with Chloride*. To determine whether the strong binding at the fourth equatorial site noted in the crystal structure [21] of  $[Cu(1)]C$ , is also observed in solution, we have investigated the reactivity of the complexes  $\left[\mathrm{Cu}(L)\right]^{2+}$  $(L = 1-4)$  with Cl<sup>-</sup> and OH<sup>-</sup> ions in  $10^{-3}$  M solution, following the reactions by conductivity and UV/VIS spectroscopy. Conductimetric titrations have previously been used to show that  $[Cu(terpy)(OH<sub>2</sub>)]<sup>2+</sup>$  binds two Cl<sup>-</sup> in MeNO<sub>2</sub> solution [35]. The results for OH<sup>-</sup> are discussed in the following section.

On titration with Bu<sub>4</sub>NCl, solutions of  $6-9$  in propylene carbonate show a change in conductivity with a more or less well resolved end point at  $1$  equiv. of  $Cl^-$  per Cu. The conductivity of the solution is consistent with that of an equimolar mixture of  $Bu<sub>4</sub>NCIO<sub>4</sub>$ and another 1:l electrolyte; similar results are obtained for **6** and **7** in DMSO. The electronic spectra also show end points at 1 equiv. per Cu. The d-d band is red-shifted by 300-900 cm-' *(Table 4),* and there **is** a rise in absorption around 38400 cm-' which we associate with  $Cl^- \rightarrow Cu(II)$  charge transfer (an analogous band has been observed at  $36\,500\,\mathrm{cm}^{-1}$  for CuCl<sup>+</sup> in propylene carbonate [22]). The complexes  $\text{[Cu(L)Cl]}^+$  have been isolated as C10; salts for ligands **24,** and their spectra and conductivities are identical with those determined by titration. The spectra of the monochloro complexes are less sensitive to change in solvent than the diperchlorato salts, the variation for [Cu-  $(m\bar{b}zimpy)$ Cl]<sup>+</sup> (10) being 13 550 (DMSO) < 13 717 (DMF) < 14 085 cm<sup>-1</sup> (propylene carbonate), a range of 535 cm<sup>-1</sup>, while, in the absence of coordinated  $CI^-$ , the variation was 1050 cm<sup>-1</sup>. EPR spectra of the monochloro complexes show a  $(d_{x^2-y^2})^1$  ground state, but for  $[Cu(bzimpy)Cl]^+$  and  $[Cu(mbzimpy)Cl]^+$  the  $g_{\perp}$  signal is slightly split into two components g,, and g,, as described by *Siddiqui* and *Shepherd* [36] for tetragonal complexes such as  $[Cu(cyclam)]^{2+}$ . The values of  $g_{\parallel}$  decrease upon Cl<sup>-</sup> coordination, presumably as a result of 'core-charge' diminution [27] [33], but A<sub>l</sub> is virtually unchanged. The addition of the C1- lowers the reduction potential by 50-1 50 mV *(Table 3),* probably as a result of the decrease in the charge of the complex [37], the systems remaining irreversible.

The behaviour on adding more than 1 equiv. of  $Cl^-$  is extremely varied. In propylene carbonate solution the bzimpy **(1)** and mbzimpy (2) complexes **6** and **7,** respectively, show a second conductimetric end-point after the addition of 2 equiv. per Cu, and [Cu(L)]CI, precipitates, but in **DMSO** the spectra are unchanged by the addition of excess CI-, suggesting that no further complexation occurs. This is confirmed by the observation that [Cu(2)]CI2.H,O **(14)** dissolves in **DMSO** to give a 1:l electrolyte (conductivity 53  $\Omega^{-1}$ ·mol<sup>-1</sup>·cm<sup>-1</sup>) expected for  $[Cu(2)Cl]^+ + Cl^-$ . The solutions of the complexes with bzoxpy **(4)** and bzthpy **(3)** in propylene carbonate change from green to yellow on addition of a second equiv. of Cl<sup>-</sup>: the d-d band falls to 11 560 cm<sup>-1</sup>, a new band appears at 21 280 cm<sup>-1</sup> ( $\varepsilon = 340$  M<sup>-1</sup> · cm<sup>-1</sup>), and the  $\pi_1^{a,b} \rightarrow \pi^*$  bands coalesce to the single broad band observed for the free ligand. When 4 equiv. of  $Cl^-$  are added, the 21 280-cm<sup>-1</sup> band shifts to 24630 cm<sup>-1</sup>; these values are typical for the  $Cl^- \rightarrow Cu(II)$  charge transfer of CuCl, and  $CuCl<sub>4</sub><sup>2</sup>$ , respectively [22]. If the complexes with **4** and **3** are titrated with Bu<sub>4</sub>NBr, then the solutions become violet, and the characteristic [38]  $Br^- \rightarrow Cu(II)$  charge-transfer bands of  $[CuBr_1]$ <sup>-</sup> and  $[CuBr_4]$ <sup>2-</sup> are observed at 18870 cm<sup>-1</sup> and 15385 cm<sup>-1</sup>, respectively, confirming the formation of halide complexes, and the displacement of the heterocyclic ligand. These observations presumably explain our failure to isolate  $[Cu(L)Cl]$ ,  $(L = 3, 4)$ *(Scheme 2).* 





For  $L = 3$ , 4, *Reaction 2* in *Scheme 2* is replaced by

$$
[CuLCl]^{+} + Cl^{-} \rightarrow L + CuCl_{2}
$$
 (3)

To compare ligands **14** with terpy **(5),** we have carried out similar experiments with this ligand. The behaviour of **5,** however, is surprisingly complex. It has been known for a long time that **5** is very rapidly exchanged in Cu(I1) complexes [39]. *Harris* and *Lockyer*  [35] have reported that green  $[Cu(5)_2]$ <sup>2+</sup> is stable in presence of halide ions in aqueous solution, but instantaneous rearrangement to the mono complex  $\text{[Cu (terpy)]X}$ , occurs in MeNO,. In propylene carbonate, we have established by spectrophotometric titrations of **5** with Cu(H<sub>2</sub>O)<sub>6</sub>(ClO<sub>4</sub>), that two species,  $[Cu(terpy)]^{2+}$  and  $[Cu(terpy)]^{2+}$ , are present in solution. The complex with Cu/ligand ratio 1 :2 possesses a weak characteristic band at 21 370 cm<sup>-1</sup>  $(\epsilon = 155 \text{ M}^{-1} \cdot \text{cm}^{-1})$  absent in the 1:1 complex. Titrations with NBu,X  $(X = Cl<sup>-</sup> or Br<sup>-</sup>)$  of  $[Cu(terpy)]<sup>2+</sup>$  formed *in situ* in propylene carbonate solution show the sudden appearance of the 21 370-cm<sup>-1</sup> band (with an intensity corresponding to half of the Cu(II) present) for 1.0 or more  $X^-$  per Cu(II). In a donor solvent such as DMSO, the characteristic band of the 2 : 1 complex is always present indicating that **1** : **1** complex is not formed under these conditions. Thus, we propose that terpy **(5)** is rapidly exchanged in the presence of good donors (DMSO,  $Cl^-, Br^-, OH^-$ ).

$$
[Cu(II)(5)(S)]^{2+} + X^- \rightarrow [Cu(II)(5)X]^+ + S \tag{4}
$$

$$
[Cu(II)(5)(X)]^{+} \rightarrow 1/2[Cu(II)(5)2]2+ + 1/2 CuX2
$$
 (5)

This behaviour is similar to that of  $[{\rm Mn}(5)X_2]$  which dissociates rapidly in aqueous solution to give  $[{\rm Mn}(5)_2]$ <sup>2+</sup> [39]. Despite the close similarity of the crystal structures of [Cu(l)Cl,] [21] and [Cu(5)Cl,] [40], **5** behaves very differently in solution, from the ligands **1-4.** 

3.3. *Reactivity of*  $\lceil Cu(L)/(ClO_4), (6-9) \rceil$  *with OH<sup>-</sup>. We have used the same methods* to investigate the reactivity of the basic nucleophile OH- toward complexes **6-9.** Conductimetric titrations in propylene carbonate (and in DMSO for  $6, 7$ ) with NBu<sub>4</sub>OH (0.1 $\mu$  in i-PrOH/MeOH) give two end-points after 1 and 2 OH<sup>-</sup> per Cu(II); the conductivities after addition of 1 equiv. of OH- suggest the formation of a 1 : 1 electrolyte. Complexes *6,* 

Complexes	$l$ equiv. $Cl^-$	1 equiv. $OH^-$	
o	14085 (97)	14600 (81)	
	13890 (95)	16 180 (88)	
	12820 (84)	14025 (55)	
	12300 (103)	13850 (60)	

**Table 4.** *d-d Muximum Energies for Complexes Formed in Propylene Curbonate Solution at 20"')* 

**8,** and **9** show essentially similar behaviour, the electronic spectra showing a slight shift on coordination of 1 OH- *(Table 4)* ; if more than 1 equiv. of OH- is added, the splitting of the ligand  $\pi^{a,b}\rightarrow\pi^*$  band disappears, and, on standing, precipitation of free ligand and poorly soluble Cu(OH), species is observed. For **9,** the precipitation of the free ligand was confirmed by IR spectroscopy. These precipitates are not soluble in excess hydroxide. EPR spectra of frozen propylene carbonate solutions of *6* and **8** with 1 equiv. of OH-

show monomeric species with  $(d_{x^2-y^2})$  ground state similar to the monochloro complexes  $(g_{\parallel} = 2.23$  for 6, 2.25 for 8). Cyclic voltammograms of the 1:1 complexes show irreversible Cu(II)/Cu(I) couples with the reduction potentials slightly lowered as observed for the monochloro complexes. We conclude that a monohydroxo complex similar to the chloro complex is formed initially, and then that the ligands **2, 4,** and **3** are displaced by excess OH-.

$$
[CuL(S)]^{2+} + HO^{-} \rightarrow [CuLOH]^{2+} + S
$$
 (6)

$$
[CuLOH]^{+} + HO^{-} \rightarrow L + Cu(OH)_{2}
$$
 (7)

The behaviour of **7,** with the ligand **1,** is, however, quite different: although the conductivity measurements suggest formation of a  $1:1$  electrolyte after addition of 1 equiv. of OH-, and precipitation occurs on adding more than 1 equiv., the precipitate redissolves completely when 3 equiv. of  $OH^-$  per Cu(II) have been added. The spectra also show a significant rise in the d-d transition energy (by  $1710 \text{ cm}^{-1}$  to  $16180 \text{ cm}^{-1}$  after addition of 1 OH<sup>-</sup>, to 16475 cm<sup>-1</sup> after addition of 3 OH<sup>-</sup>), whereas the complex with the

Scheme 3. *Reactivity of*  $\left[Cu(bzimp)\right]^{2+}$  *(7) with OH<sup>-</sup> in Aprotic (propylene carbonate or DMSO) <i>Solution*  $(S = solvent)$ 



similar ligand **2** shows no such sharp rise. We explain this difference by the presence of the two weakly acidic protons of the benzimidazoles, which may be removed by base: this would be expected to increase the donor power of the ligand, and raise the d-d transition energy. To confirm this hypothesis, we have titrated solutions of **6** and **7** in DMF solution with the weak, sterically hindered base Et,N. Complex **6** shows no change in spectrum, showing that  $Et<sub>3</sub>N$  does not coordinate to the Cu(II), but with 7 the spectrum obtained after addition of 1 equiv. of  $Et<sub>1</sub>N$  is identical to that found after addition of 1 equiv. of  $OH^-$ , as would be expected, if the first reaction is indeed deprotonation of the ligand. We propose, therefore, that the reaction of  $\left[\text{Cu}(1)\right]^{2+}$  with OH<sup>-</sup> involves two deprotonations of the ligand **1** to give an insoluble neutral complex of H, bzimpy, followed by redissolution of an anionic complex  $[Cu(H,bzimpy)(OH)]$ . Deprotonation of 1 coordinated to Fe(II) and Fe(III) has previously been observed [17] [41]. It is worth noting that  $OH^$ does not displace the deprotonated form of 1 from the Cu(I1).

With terpy (5) in propylene carbonate solution, the electronic spectrum after the addition of 1 equiv. of OH<sup>-</sup> shows the formation of  $[Cu(5)]^{2+}$  with the formation of a fine precipitate, suggesting that reactions similar to *Eqns. 4* and *5* are also observed for OH<sup>-</sup>.

4. *Equilibrium Studies of Binding to*  $\left[\frac{Cu(mbzimp)}{l^2}\right]$ *.* To obtain a quantitative estimate of the strength with which additional ligands may be bound to the  $Cu(II)$  in these complexes, we have carried out a number of potentiometric titrations of solutions of  $[Cu(mbzimpy)]^{2+}$  (hereafter referred to as CuL) in 1:1 MeCN/H<sub>2</sub>O. This solvent was chosen for reasons of solubility. The solutions (at  $25^{\circ}$  and with an ionic strength of 0.05 $\mu$ ) were titrated with KOH, and the resulting pH/(volume of base) curves analyzed by a non-linear least-squares program [25] to obtain apparent  $pK<sub>a</sub>$  values (determined as the pH value at half neutralisation). Each system investigated was studied at several concentrations of CuL, and the quoted constants are the averages obtained from several titrations.

For solutions of CuL alone, we observe the titration of two protons, the first in the pH range 5.5 to 10, and the second between pH 10 and 12. The completion of the second deprotonation is accompanied by precipitation of Cu(OH), and free ligand. The titration curve in the range where no precipitation is observed is fitted within experimental error by two constants corresponding to the reactions

[CuLOH<sub>2</sub>]<sup>2+</sup> 
$$
\rightleftharpoons
$$
 [CuLOH]<sup>+</sup> + H<sup>+</sup>, pK<sub>8</sub> = 7.80(5) (8)

[CuLOH]<sup>+</sup> + H<sub>2</sub>O 
$$
\rightleftharpoons
$$
 Cu(OH)<sub>2</sub>  $\downarrow$  + L $\downarrow$  + H<sup>+</sup>, pK<sub>9</sub> = 13.3(1) (9)

As expected, the apparent  $pK_a$  for *Reaction 8* is independent of the total Cu(II) concentration (and excludes the formation of a hydroxo-bridged dimer), whereas that of *Reaction 9*  increases with the initial concentration of CuL. The dependence of the apparent  $pK_n$ values upon total CuL concentration has been used as confirmation of the stoichiometry of all the reactions studied. The observed  $pK$ , of H<sub>2</sub>O coordinated to  $[CuL]<sup>2+</sup>$  is in the range observed for Cu<sub>aa</sub> (7.3–8.0) [42] and slightly below the values found for [Cu(terpy)- $(OH<sub>2</sub>)$ <sup>2+</sup> (8.3 [43]) and  $[Cu(pip)(OH<sub>2</sub>)]^{2+}$  (8.4 [44], pip = 2-{[2-(2-pyridyl)ethylimino]methyl} pyridine).

Titration of CuL in the presence of 1 equiv. of 1-methylimiddzolium (Hmim+) shows three deprotonations corresponding successively to *Reactions 10, 11,* and *9* :

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$$
[CuL(OH2)]2+ + Hmin+ \rightleftharpoons [CuL(min)]2+ + H3O+, pK10 = 2.1(2)
$$
 (10)

$$
[CuL(min)]^{2+} + 2 H_2O \rightleftharpoons [CuLOH]^+ + \text{mim} + H_3O^+, pK_{11} = 12.1(2)
$$
 (11)

These assignments were confirmed by the dependence of apparent  $pK_a$  upon total Cu(II) concentration. Using the  $pK_a$  value of 6.57(2) determined for Hmim<sup>+</sup> under our conditions, we obtain for *Reaction 12* 

$$
[CuL(OH2)]2+ + \min \rightleftharpoons [LCu(\min)]2+ + H2O \log K12 = 4.5(2)
$$
 (12)

significantly greater than that found for  $[Cu(min)(pip)]^{2+}$  (log  $K = 3.3(1)$  [44]). If imidazole (Him) **is** used instead of mim, the first deprotonation corresponds to formation of the complex  $\text{[Cul(Him)]}^{2+}$  and gives a stability constant log  $K = 4.5(1)$  identical to that of mim; no difference between mim and Him was found for [Cu(pip)] [44].

The possibility of forming imidazolate-bridged dimers on further titration of imidazole complexes is now well-established [44] [45] and, on titrating a 2:1 mixture of  $[CuL]^{2+}$ and H,im+, a second deprotonation corresponding to

$$
[CuL(Him)]^{2+} + [CuL(OH2)]^{2+} \rightleftharpoons [CuL(im)CuL]^{3+} + H3O+, pK13 = 4.1(2)
$$
 (13)

is observed *(Fig. 3, a)*. The observed  $pK$ , is identical to that observed for the formation of the glycylglycine complex  $[(glygly)Cu(im)Cu(glygly)]^{-}(4.1 [45])$ . Formation of the dimer is confirmed by the correct negative dependence of the apparent  $pK_a$  on the total Cu(II) concentration, and by the isolation of [LCu(im)CuL](ClO<sub>4</sub>)<sub>3</sub>·H<sub>2</sub>O (15). The IR spectrum of this compound shows an absorption due to  $H_2O$  of crystallisation at 3450 cm<sup>-1</sup>, but no  $\tilde{v}(N-H)$  in the region of 3100-3300 cm<sup>-1</sup>. The EPR spectrum of the powder at room temperature shows a broad rhombic signal in the  $g = 2$  region and a signal corresponding to the  $\Delta m = \pm 2$  transition at g = 4.2, confirming the presence of a bridged dimer [44].



Fig. 3. a) *Typical titration curve of a 2:1 mixture of*  $[Cu(2)]^{2+}$  *and*  $H_2$ *im*<sup>+</sup>. b) *Calculated species distribution as a function of pH for the titration curve in* (a). The assignment of  $[Cu(2)OH]$ <sup>+</sup> is tentative (see text).

Solutions of **15** in MeCN at  $25^{\circ}$  give a molar conductivity of  $123 \pm 5$   $\Omega^{-1}$  cm<sup>2</sup> mol<sup>-1</sup> which is consistent with a 3:1 electrolyte *(Table 1)*. Using the constant determined above, we may determine

$$
[CuL(imH)]^{2+} + [CuLOH]^{+} \rightleftharpoons [LCu(im)CuL]^{3+} + H_{2}O, \log K_{14} = 3.7(2)
$$
 (14)

If the titration is continued after the formation of the dimer, further deprotonation is observed, leading eventually to precipitation of Cu(OH), and free L. The UV/VIS spectra

of the solutions after the addition of a 3rd equiv. of base suggest the major species in solution to be [CuLOH]'

$$
[LCu(im)CuL]^{3+} + 3 H_2O \rightleftharpoons 2 [CuLOH]^{+} + Him + H_3O^{+}
$$
 (15)

as was observed with the system  $\left[\text{Cu(pip)}\right]^2$  [44]. The dependence of the observed p $K_a$  on total Cu(I1) concentration does not, however, agree with this, and we cannot exclude the presence of  $\text{[CuL(im]}^+$  as seen with the glycylglycine complex [45]. Study of this reaction is hindered by precipitation, but the observed titration curves show that the dimer [LCu(im)CuL]<sup>3+</sup> is stable in the pH range of 7–9 *(Fig. 3, b)*. Titration of 1:1 mixtures of  $[Cul]^2$ <sup>+</sup> and H<sub>2</sub>im<sup>+</sup> show initial formation of  $[Cul(Him)]^{2+}$  followed by a deprotonation which is consistent with formation of the dimer; further titration leads to precipitation as for the 2:1 mixtures.

**Conclusions.** – On the basis of the d-d band maxima for  $\lbrack \text{CuL} \rbrack^{2+}$ ,  $\lbrack \text{CuLCl} \rbrack^{+}$ , and [CuLOH]', the ligands bzimpy **(1)** and mbzimpy **(2)** are only slightly weaker donors than terpy *(5),* whereas bzthpy **(3)** and bzoxpy **(4)** are significantly weaker. This agrees with the *Mössbauer* results for the complexes  $[FeL<sub>2</sub>]<sup>2+</sup>$ , where the complexes with 1 and 2 are low-spin and with **3** and **4** are high-spin at room temperature [41]. In solution, the EPR and UV/VIS spectra of  $[CuL]<sup>2+</sup>$  are consistent with expected tetragonal structure, and the change in the magnitude of the solvent effect on the electronic spectrum on going from [CuLI2' to [LCuCl]' shows that the *Lewis* acidity is indeed concentrated in the fourth equatorial site. From a purely structural point of view, therefore, all five ligands studied here seem to satisfy our initial requirements. **A** closer examination of their solution chemistry reveals, however, that bzthpy **(3)** and bzoxpy **(4)** are too easily decomplexed by donor solvents or other ligands to be of much use. More surprising, but equally limiting for model studies, is the observation of the ease with which  $\left[\text{Cu(terpy)}\right]^{2+}$  complexes disproportionate to give  $\left[\text{Cu}( \text{terpy})\right]^{2+}$  and  $\text{CuX}$ , *(Eqn. 5).* Of the two remaining ligands, bzimpy **(1)** suffers from the weak acidity of the benzimidazole protons which are likely to interfere with any basic peroxo complex. The most promising precursor for a copper-dioxygen complex is, therefore, the complex  $[Cu(mbzimp)]^{2+}$ ; this complex shows good thermodynamic stability, being decomplexed only by an excess of OH- at fairly high pH. The potentiometric measurements show that imidazole and 1 -methylimidazole are quite strongly bound to the fourth coordination site (log  $K = 4.5$ ), an order of magnitude greater than  $\left[\text{Cu(pip)}\right]^{2+}$ . The formation of an imidazolate-bridged dimer, although of more direct interest for modelling the active site of superoxide dismutase [44] [45], shows that dimerisation of  $\left[\text{Cu(mbzimp)}\right]^{2+}$  units is not impossible sterically, and that the formation of a peroxo-bridged dimer may be envisaged.

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