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Synthesis, Optical Spectroscopy, Structural, and DFT Studies on Dimeric Iodobridged Copper(I) complexes

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Abstract:

Three new iodo-bridged copper(I) complexes of the type $[Cul(PPh_3)L]_2$, where $L = Ar - \Xi - C_5H_4N$, Ar = phenyl (**C**₁), biphenyl (**C**₂) and flourenyl (**C**₃) have been synthesized *via* coordination-driven self-assembly processes. Two of Cu(I) complexes, **C**₂ and **C**₃, have been characterized by single-crystal X-ray diffraction studies. The complexes have two molecules of the P-donor ligand and two molecules of the N-donor ligand in *trans* configurations, supporting the central Cu₂I₂ unit. Absorption properties of the complexes have been investigated. Extensive DFT calculation has been carried out to delineate the influence of aromatic spacers on the optical properties and the nature of excited states. The ease of synthesis of these Cu(I) dimers and the wide range of ethynylpyridine supporting ligands that can be incorporated highlights the potential for these materials to form polymers by linking through the ethylylpyridine ligands.

Keywords: Copper(I) complexes; Ethynylpyridine; DFT studies; Self-assembly; X-ray crystal structures

1. Introduction

Transition metal complexes have attracted significant attention due to their diverse photophysical properties and applications.[1-5] The last four decades has witnessed a great upsurge in the design and development of d¹⁰ ion-based complexes such as those of copper(I) bearing homo- and heteroleptic ligands.[6, 7] Among luminescent transition metal complexes, copper(I)-based systems offer easy synthetic routes, a diverse range of structural arrangements, bright luminescence encompassing a broad range of the visible and UV spectrum, high quantum yields, etc. It has been demonstrated that neutral copper(I) complexes bearing Nand P-coordinating ligands surrounding a halogenido-bridged Cu₂X₂ (X = halogen) core are good candidates for optoelectronic applications.[8] By simple variation of the coordination sphere (i.e. the coordinating and bridging ligands) around the metal, both absorption and emission can be manipulated. For instance, it has been demonstrated that metal to ligand charge transfer (MLCT) or halide to ligand charge transfer (XLCT) emission bands can be significantly tuned by varying the π^* level of ligands.[8] Because of these features, several new copper(I) complexes have been synthesized and tested as components for photovoltaics (PVs), organic-light emitting diodes (OLEDs), sensors, etc.[7, 9] The low toxicity and the pseudo-tetrahedral geometry of the complexes has also prompted researchers to assess their biological activity.[3]

Within the class of Cu₂X₂ complexes, synthetic manipulation can readily be achieved by altering the nature of the supporting ligands, usually with donor N- or P-bridgehead atoms.[10-12] The pyridine nitrogen donor atom has a strong tendency to interact with a variety of metals and, as a result, a number of complexes bearing simple to functionalized pyridine (or higher pyridines) have been reported.[3] Not only does the electron-deficient pyridine endow structural rigidity, but a judicious choice of functionalization and macrocyclization through pyridine core can also impart helicity/chirality to the resulting complexes.[13] Bearing these features in mind, we have investigated structural and photophysical behavior of a number of acetylide-functionalized pyridinyl copper(I) complexes. Both dimeric and tetrameric copper(I) complexes bearing ethynylpyridinyl-based coordinating ligands have been investigated [5, 14, 15] For example, we demonstrated that copper(I) complexes bearing pyridine ligands connected to 4-ethynyl ferrocene show intriguing electrochemical properties. In such complexes, the ferrocene units act as an electron reservoir/sink and was detrimental to the emission properties.[15] Similarly, we recently demonstrated that dye sensitized solar cells (DSSC) based on Cu₂X₂ bridged complexes with heteroarylethynyl attached to 4position of pyridine (Chart 1) are highly emissive and show up to 2% light to energy conversion efficiency.[5] The performance of the dye was found to depend significantly on two factors: (i) nature of the aryl group attached to pyridine via π -linker; and (ii) presence/absence of an anchoring group associated with the phosphine ligands.



Chart 1: Recently reported acetylide-functionalized pyridine-based Cu(I) complexes for PV applications.

To examine the effect of replacing heterocyclic spacers by carbocyclic spacers (such as phenyl, biphenyl and fluorene), we now report the synthesis, the structures and photophysical and DFT studies on three new iodo-bridged copper(I) complexes supported by ethylylpyridine-based ligands. The complexes were prepared *via* self-assembly in the solution phase under mild conditions.

2. Experimental

2.1 General

All chemicals and reagents, except where stated otherwise, were obtained commercially and used without further purification. Unless otherwise stated, syntheses were performed under a dry argon atmosphere. Solvents were pre-dried and distilled before use by standard procedures.[16] NMR spectra were recorded on a Bruker AM-400 spectrometer in CDCl₃. ¹H-NMR spectra were referenced to solvent resonances and ³¹P{1H} NMR spectra were referenced to external phosphoric acid. IR spectra were recorded directly on the sample as attenuated total reflectance (ATR) on Diamond using Cary 630 FT-IR spectrometer. UV-vis spectra were recorded with an Agilent Cary 5000 UV-visible spectrophotometer using a quartz cuvette with a 1 cm path length. Column chromatography was performed on either Kieselgel 60 (230 – 400 mesh) or celite.

2.2 General protocol for the synthesis

2.2.1 Synthesis of ligands (L₁-L₃) and complexes (C₁-C₃)

The intermediates and final products were obtained *via* established cross-coupling reactions.[5, 15, 17] Briefly, 4-trimethylsililyethynylpyridine was obtained by reacting an equimolar quantity of 4-iodopyridine (2.0 mmol) with trimethylsilylethyne (TMSE, 2.0 mmol) in THF/ⁱPrNH₂ using Pd(II)/Cu(I) catalysts. (**Scheme 1**) The crude product obtained was purified by silica column chromatography using hexane/dichloromethane eluant systems. This was followed by a protodesilylation reaction under basic condition (aq. KOH in MeOH/THF) and the 4-ethynylpyridine was separated by silica column chromatography using pure dichloromethane as eluant. In the next phase of the reaction, respective aryl halides (2.0 mmol) and 4-ethynylpyridine (2.0 mmol) were coupled *via* a Sonogashira cross-coupling reaction. Final products were purified using an alumina column and obtained as tan to light brown solid in quantitative yields.

Phenylethynylpyridine (L₁): IR (v_{max}) cm⁻¹: 2199 (-C≡C-), 1598 (C-N), ¹H-NMR (700 MHz, CDCl₃): δ (ppm) 8.45 (d, 2H, J = 6.4, H-py), 7.98 (d, 2H, J = 5.8, H-py), 7.35 (d, 2H, J = 6.2, H-Ph), 6.70-6.68 (m, 3H, H-Ph). ESI-MS: m/z 179.9 (M+). C₁₃H₉N Anal. Calc.: C, 87.12; H 5.06; N, 7.82%. Found: C, 86.97; H, 4.94; N, 7.77%.

Biphenylethynylpyridine (L₂): IR (v_{max}) cm⁻¹: 2208 (-C≡C-), 1605 (C-N), ¹H-NMR (700 MHz, CDCl₃): δ(ppm) 8.28 (d, 2H, J = 5.95 Hz, H-py), 7.68 (d, 2H, J = 4.55 Hz , H-py), 7.62-7.60 (m, 2H , H-biph), 7.46 (dd, 4H, J = 5.53, 13.51 Hz, H-biph), 7.38-7.35 (m, 3H,H-biph). ESI-MS: m/z 256.1 (M+). C₁₉H₁₃N Anal. Calc.: C, 89.38; H, 5.13; N, 5.49 %. Found: C, 8.98; H, 4.83; N, 5.02%.

Fluorenylethynylpyridine (L₃): IR (v_{max}) cm⁻¹: 2205 (-C=C-), 1608 (C-N), ¹H-NMR (700 MHz, CDCl₃): δ(ppm) 8.61 (d, 2H, J = 5.74 Hz, H-py), 8.27 (d, 2H, J = 5.95 Hz, H-py), 7.81-7.79 (m, 2H, H-fluorene), 7.78 (s, 1H, H-fluorene), 7.73 (m, 2H, H-fluorene), 7.59-7.57 (m, 2H, H-fluorene), 3.93 (s, 2H). ESI-MS: m/z 268.1 (M+). C₂₀H₁₃N Anal. Calc.: C, 89.86; H, 4.90; N, 5.24%. Found: C, 88.26; H, 4.49; N, 4.94%.



2.2.2 Synthesis of Cu(I) complexes (C₁-C₃)

Cu(I) complexes were synthesized following the protocol reported in literature.[5, 15, 17] Complexes C_1 - C_3 were obtained by stirring a mixture of the appropriate arylethynylpyridine ligands (1.0 mmol), CuI (1.0 mmol) and triphenylphosphine (1.0 mmol) in an equimolar mixture in degassed dichloromethane. (Scheme 2) After stirring for 24 h at room temperature, the mixture was filtered, and the filtrate was concentrated under reduced pressure to produce neutral dimeric complexes C_1 - C_3 . All complexes are stable to light and air and were fully characterized. Crystals suitable for X-ray analysis were obtained by slow diffusion of dichloromethane (DCM) in hexane.

Complex [L1(Cul)₂(PPh₃)₂] (**C**₁): IR: 2206 cm⁻¹ v(C=C), 1488 cm⁻¹ v(C=N). ¹H-NMR (700 MHz, CDCl₃): δ (ppm) 8.70 (d, *J* = 5.98 Hz, 4H, H-py), 7.91 (d, *J* = 6.3 Hz, 8 H, H-py and PPh₃), 7.71–7.42 (m, 36 H, H-Ph and PPh₃). ³¹PNMR (121.53 MHz, CDCl₃): δ = 29.18 (s, PPh₃) ppm. ESI-MS: m/z 1261.9 (M+). C₆₂H₄₈Cu₂I₂N₂P₂ Anal. Calc.: C, 58.92; H, 3.83; N, 2.22%. Found: C, 59.55; H, 4.01; N, 2.17%

Complex [L2(Cul)₂(PPh₃)₂] (**C**₂): IR: 2210 cm⁻¹ v(C=C), 1489 cm⁻¹ v(C=N). ¹H-NMR (700 MHz, CDCl₃): δ (ppm) 8.65 (d, *J* = 4.48 Hz, 4 H, H-py), 7.62 (d, *J* = 10.5 Hz, 12 H, H-py and PPh₃), 7.55–7.34 (m, 40 H,

H-biph and PPh₃). ³¹PNMR (121.53 MHz, CDCl₃): δ = 29.22 (s, PPh₃) ppm. ESI-MS: m/z 1413.7 (M+). C₇₄H₅₆Cu₂l₂N₂P₂: Anal. Calc.: C, 62.76; H, 3.99; N, 1.98%. Found: C, 64.88; H, 4.11; N, 2.01%.

Complex [L3(Cul)₂(PPh₃)₂] (**C**₃): IR: 2208 cm⁻¹ v(C=C), 1493 cm⁻¹ v(C=N). ¹H-NMR (700 MHz, CDCl₃): δ (ppm) 8.64 (m, 4 H, H-py), 7.81–7.79 (m, 4 H, H-py), 7.74-7.57 (m, 30 H, H-flourene and PPh₃), 7.52-7.34 (m, 14 H, H-flourene and PPh₃), 3.94 (s, 2H). ³¹PNMR (121.53 MHz, CDCl₃): δ = 29.31 (s, PPh₃) ppm. ESI-MS: m/z 1440.2 (M+). C₇₆H₅₆Cu₂I₂N₂P₂ Anal. Calc.: C, 63.38; H, 3.92; N, 1.95%. Found: C, 64.88; H, 4.19; N, 2.01 %.



2.3 X-ray Crystal Structure Determination

Single-crystal X-ray diffraction experiments were performed at 150(2) K on an Oxford Diffraction Gemini A Ultra CCD diffractometer monochromatic Mo-K α radiation ($\lambda = 0.71073$ Å). The sample temperature was controlled using an Oxford Diffraction Cryojet apparatus; CrysAlis Pro was used for the collection of frames of data, indexing reflections and determining lattice parameters.[18] Structures were solved by direct methods using SIR-92[19] and refined by full-matric least-squares, on F^2 , using SHELXL2014[20] within the OLEX2 suite of programs.[21] A multi-scan absorption correction was applied in all cases. The hydrogen atoms were generated geometrically and refined with common isotropic thermal parameters. Crystallographic data for the complexes is presented in **Table 1**. In the case of **C**₃, the fluorene rings showed significant disorder that could not be modelled effectively. As a last resort the unit was assigned isotropic displacement parameters and the carbon atoms were assigned full occupancies. While the refinement was stable with this model several significant residual electron density peaks in the vicinity of the ligand remained. The cif files for the two structures have been deposited with the CCDC: *CCDC 1898577-1898578*

2.4 Computational studies

2.5 Computational details

All DFT calculations were performed using the Gaussian 09[22] package with the Lee–Yang–Parr correlation function (B3LYP).[23] For copper and iodide, the Hay and Wadt basis-set LANL2DZ pseudopotential.[24, 25] was used. The 6-311+G(d).[26] basis-set was employed for N and P, while the

6-31G(d,p)[27] basis-set was operated for the atomic orbital of C, H to calculate all the optimized structures.TD-DFT calculations were performed using the same functional and the polarizable continuum model, CPCM with dichloromethane (ε = 8.93) as the surrounding solvent considering bulk solvent effects.[28] TD-DFT-calculated electronic spectra were depicted by using GaussSum,[29] All the molecular orbitals and spin-density diagrams were plotted using the Chemcraft Visualization program.[30]

3. Result & Discussion

3.1 Synthesis and Spectroscopic Characterization

The ligands L_1-L_3 and their corresponding complexes C_1-C_3 have been synthesized by adapting previously reported procedure.[5] Briefly, arylhalides (4-lodophenyl, 2-bromobiphenyl or 2-iodofluorenyl) were cross-coupled with TMSA using Sonogashira coupling reaction followed by protodesilylation giving the terminal alkynes (Scheme 1). The deprotected ligands obtained were then coupled with 4-iodopyridine to produce ligands L_1-L_3 as yellow-brown powders in 60-78% yield. Analytical data (IR, NMR and MS) confirmed the formulation of the ligands. For example, TMSA protection of 4-iodopyridine was confirmed by the observed $-C \equiv C$ - stretching frequency (~ 2165 cm⁻¹) while deprotection was confirmed by the presence of the $-C \equiv H$ stretching frequency (~ 3223 cm⁻¹) in addition to the $-C \equiv C$ - band. Moreover, the $v_{(C \equiv C)}$ value of 4-ethynylpyridine was found to be lower than those of the 4-(trimethylsilylethynyl)pyridine due to the fact that terminal alkynes (HC $\equiv C$ -R) have lower $v_{C \equiv C}$ stretching frequencies than their protected counterparts RC $\equiv CR$.[31] The complexes C_1-C_3 were obtained by swift solution phase self-assembled coordination. The reaction of the ligand (L_1-L_3), with copper iodide and triphenyl phosphine (1:1:1 ratio), in dry dichloromethane, under an argon atmosphere produced the complexes C_1-C_3 . In addition to the standard analytical techniques, single crystal X-ray diffraction studies provided the full three-dimensional molecular and crystal structures of the three complexes.

3.2 Structural characterization

The crystal structures of the complexes: C_2 and C_3 were determined and are shown in Figures 1 and 2. Crystal data is summarized in **Table 1** and bond parameters are listed in **Tables ST1-ST4** (*supplementary information*). Single crystals of the complexes were obtained by slow diffusion of hexane into a dichloromethane solution of each complex.



Figure 1. The crystal structure of the dimeric complex C₂. Hydrogen atoms are omitted for clarity.

The crystal structures of the dimeric complexes C_2 and C_3 have a similar planar, parallelepiped Cu₂I₂ core and a *trans* arrangement of the two phosphine and the two L ligands. In each case the center of the Cu₂I₂ core sits on a crystallographic center of symmetry, so that the crystallographic asymmetric unit contains half of a dimeric molecule. Each Cu(I) center is in a tetrahedral coordination environment consisting of two bridging iodo ligands, one PPh₃ and the bridgehead nitrogen atom of L₂ and L₃. The unique bond distance for Cu-I, Cu-P and Cu-N are 2.6487(4), 2.2412(6) and 2.063(2) Å, respectively, in C₂ and are 2.651(8), 2.235(2) and 2.059(6) Å, respectively, in C₃. These bond distances are similar to the equivalent bond lengths in a family of previously reported [(Py)(Ph₃P)CuX]₂ adducts.[32] The angles at the copper center in the Cu₂I₂ core are obtuse, *e.g.* 111.45(1)° and 108.13(3)° in C₂ and C₃, respectively. While the angle at the iodine center is acute, **e.g.** 68.54(1)° and 71.83(3)° in C₂ and C₃, respectively. The Cu-Cu separations in the dimers: C₂ (3.0196(5) Å) and C₃ (3.1394(5) Å), are too long for a formal Cu-Cu bonding interaction; however, it remains similar to reported analogues.



Figure 2. The crystal structure of the dimeric complex C₃. Hydrogen atoms are omitted for clarity.

Table 1. Crystallographic data for C2-C3

	C ₂	C ₃
Empirical formula	$C_{74} H_{56} Cu_2 I_2 N_2 P_2$	$C_{76} H_{56} Cu_2 I_2 N_2 P_2$

Formula weight	1416.02	1440.04	
Т(К)	150(2)	150(2)	
λ (Å)	0.71073	0.71073	
Crystal system	Monoclinic	Monoclinic	
Space group	P21/n	P2 ₁ /n	
а	9.43370(10)	9.3125(2)	
b	22.5428(3)	22.1452(4)	
c	14.2331(2)	15.2590(3)	
α	90°.	90	
β	98.562(1)	101.061(2)	
γ	90	90	
V (Å ³)	2993.11(7)	3088.36(11)	
Z	2	2	
ρ _{calc.} Mg m ⁻³	1.571	1.549	
μ (Mo-K _α) (mm ⁻¹)	1.842	1.787	
F(000)	1416	1440	
Crystal size (mm)	0.60 x 0.40 x 0.40	0.2 x 0.2 x 0.04	
θ range (deg)	2.834 to 31.192	2.971 to 29.689	
Reflections collected	79295	40941	
Independent reflections [R(int)]	9128 [0.0375]	7906 [0.0385]	
Completeness to theta = 25.242°	99.90%	99.80%	
Max. and min. transmission	1.00 and 0.730	1.000 and 0.9209	
Data / restraints / parameters	9128 / 0 / 370	7906 / 0 / 302	
Goodness-of-fit on F ²	1.133	1.034	
Final R ₁ , wR ₂ indices [I>2σ(I)]	0.0305, 0.0584	0.0719, 0.1870	
R indices (all data)	0.0454, 0.0647	0.0927, 0.2013	
Largest diff. peak, hole (e.Å ⁻³)	1.101, -0.516	3.528, -1.267	

3.3 Optical Spectroscopy

The electronic spectra of complexes C_1-C_3 were measured in dichloromethane at room temperature (**Figure 3a**). The onset of absorption for all ethylpyridinyl-based ligands and their corresponding Cu complexes were red-shifted compared to ligand precursor 4-(trimethylsilylethynyl)pyridine ($\lambda_{max} = 286$ nm). The complex incorporating the fluorenylethynylpyridine ligand (C_3) showed red-shifted band compared to phenyl and biphenylethnylpyridine Cu(I) complexes (C_1 and C_2). Complex C_1 showed a strong absorption band at 300 nm (λ_{max}) whereas complexes C_2 and C_3 showed absorptions peaking at 310 and 334 nm, respectively. This can be ascribed to the increasing π -conjugation length leading to a lower energy for the complex and absorption at longer wavelengths. The absorption peaks in the UV spectrum ($\lambda_{max} < 400$ nm) can be attributed to the $\pi \rightarrow \pi^*$ transition associated with the arylethynylpyridine group. **Figure 3b** shows the effect of attaching different aromatic spacers to 4-ethynylpyridine ligands on the absorption maxima. As expected, the λ_{max} values move to the red (≥ 10 nm) upon the incorporation of heterocyclic spacers.[5] Comparison of these data with our previous results [5] clearly indicate that the presence of a heavy atom

such as sulfur play important role in shifting the λ_{max} values in dimeric iodo-bridged Cu(I) complexes. As mentioned before, Cu(I) complexes are well known for their emissive properties in both the solid and solution states.[8] In the past, we have demonstrated that the emission properties of ethynylpyridine ligands-based Cu(I) complexes in DCM solution is mainly governed by the electron withdrawing/donating nature of the spacer group. Moreover, there is only little impact of the alkynyl unit (especially for mono-ynes) on the emission wavelength.[5] Therefore, a trend similar to absorption is expected on the emission properties. We are currently investigating the solid-state as well as solution luminescence, electrochemical, thermal and photovoltaic properties of carbocyclic and heterocyclic spacer incorporated ethynylpyridine ligands-based Cu(I) complexes. The results will be communicated separately.





4. DFT Calculations

The structures of complexes C_1 - C_3 , bearing three different carbocylic groups (with varying conjugation lengths) attached to the ethynylpyridine ligands were confirmed by DFT calculations employing the B3LYP level of theory (see computational details in the experimental section). Geometry-optimization of complexes C_2 and C_3 was performed using the X-ray crystallographic coordinates and the complex C_1 model was optimized using the same hybrid level of theory. The minima were verified *via* computed vibration frequencies. The DFT calculated bond-lengths and bond-angles of the optimized structures are (Figure 4 and Table ST5, supplementary file) in good agreement with those obtained experimentally (Table ST1-4, supplementary file). The time-dependent DFT (TD-DFT) calculations were executed on these optimized geometries to analyze the photophysical properties (Figure 5). Other frontier molecular orbitals involved in electronic transitions in complexes C_1 — C_3 , as obtained by TD-DFT calculation are shown in Figure SF1-SF3 (supplementary information).



Figure 4.



Figure 5. TD-DFT-Calculated electronic spectra for the Cu(I) complexes (a) C₁, (b) C₂, and (c) C₃ using the B3LYP level of theory. Each plot shows the simulated absorption profile (green), with the bars marked by blue lines representing oscillator strength. Isosurface plots of the highest-occupied and lowest-unoccupied molecular orbitals (HOMOs/LUMOs) for each system are shown to the right of the simulated spectra.

To obtain deeper insight of the electronic transitions for all three complexes, time-dependent (TD-DFT) calculations have been performed. Here we semi-quantitatively explain the effect of extended conjugation embedded within alkynyl group attached to the pyridyl ligands. The calculated and experimental UV-vis spectra are depicted in **Figure 6**. To analyze the spectroscopic activity of these complexes, we have mainly considered the transition in each case with maximum oscillator strengths, *f*, among all the transitions with excitation wavelengths in the 300–400 nm region. From TD-DFT assignments it was revealed that all transitions in this region are largely due to charge-transfer (CT) mainly from the iodide ligands towards the pyridyl rings within the terminally conjugated alkynyl groups. Along with this, there is a substantial contribution of MLCT involving the metal \rightarrow substituted pyridyl rings. **Tables 2** and **ST6-ST7** (*supplementary information*) lists all excitation energies, wavelengths, oscillator strengths, and an assignment of each state in terms of contributions from individual orbital transitions.



Figure 6. TD-DFT simulated (a) and absorption (b) electronic spectra of complexes C₁-C₃ in DCM

Excitation energy (eV)	λ (nm)	f	Transition	Character
2.780	445	0.074	α/β-H[~58%L]→ α/β- L[~80%L](92%)	Charge transfer (CT) mainly from iodide towards the pyridyl rings within the terminally conjugated alkynyl groups. A significant contribution of MLCT is also observed involving metal \rightarrow substituted pyridyl rings.
3.756	330	0.310	α/β-H-3[~57%M]→ α/β- L+1[~89%L](86%) α/β-H-5[~63%L]→ α/β- L+1[~89%L](36%)	Combination of MLCT involving metal → substituted pyridyl rings with subsequent CT from iodide towards pyridyl rings within the terminally conjugated alkynyl groups Charge transfer (CT) mainly from iodide towards the pyridyl rings within the terminally conjugated alkynyl groups. An appreciable contribution of MLCT is also observed here involving metal → substituted pyridyl rings.
3.8144	325	1.62	α/β-H-8[~50%L]→	Combination of MLCT involving metal → substituted pyridyl rings with subsequent CT

Table 2. TD-DFT calculated electronic transitions of C_1 using B3LYP level of theory.

			α/β- L+1[~89%L](28%) α/β-H-7[~53%M]→ α/β-L [~80%L](66%)	from iodide towards the pyridyl rings within three terminally conjugated alkynyl groups. MLCT involving Metal → substituted pyridyl rings with terminally conjugated alkynyl groups and CT in substituted pyridyl rings with terminally conjugated alkynyl groups
4.013	308	0.295	α/β-H-10[~77%M]→ α/β-L[~80%L](34%)	MLCT involving metal \rightarrow substituted pyridyl rings within the terminally conjugated alkynyl groups.
			α/β-H-8[~50%L]→ α -L+1[~890%L](46%)	Combination of MLCT involving metal \rightarrow substituted pyridyl rings with subsequent CT from iodide towards the pyridyl rings within the terminally conjugated alkynyl groups.

The calculated HOMO-LUMO gaps associated with low energy transitions (~ 440-450 nm) are also given in the **Table 3** though they are not clearly visible experimentally due to the broadening of absorption bands. In complex **C**₁, the brightest transition at nearly 300 nm (experimentally) could be assigned as HOMO-5 to LUMO+1, HOMO-7 to LUMO, and HOMO-8 to LUMO+1 excitations, respectively, with oscillator strengths, f = 1.62. We also found a considerably weaker transition between the HOMO and LUMO orbitals at 445 nm (f = 0.074). A gradual red shift (**Table 3**) in the absorption edge (with maximum f) of complex **C**₂ (310 nm) and complex **C**₃ (334 nm) has been observed experimentally, as we move forward from complex **C**₁ towards complex **C**₃. This can be explained on the basis of their structural configurations. Due to the incorporation of the more delocalized functionalities into the alkynyl group attached to the pyridyl ligands in Cu(I) complexes, in going from **C**₁ to **C**₃ a noticeable bathochromic shift is observed in the spectra. Here, the computational modeling also semi-qualitatively represents the key inferences extracted from the optical spectroscopy. It summarized the importance of extended delocalization of the ligands on the frontier orbitals of complex **C**₁ and its analogues leading to red shift across the metal centers and between the aromatic ligands.

Complex	Δ(HOMO – LUMO)/ kcal/mol	λ (nm)		
		Expt.	DFT calculated	
C ₁	64.7838	300	324	
C ₂	74.0262	310	353	
C ₃	74.1203	334	375	

Table 3. The calculated HOMO-LUMO gaps associated with low energy transitions, the experimental and calculated λ values.

5. Conclusion

We have synthesized a range of $[Cul(PPh_3)L]_2$ dimeric complexes, with a Cu_2l_2 core, where L is a series of ethynyl-functionalized pyridine ligands (Ar-C=C-C₅H₄N), to include the aryl groups phenyl (C₁), biphenyl (C₂) and fluorenyl (C₃) using established coordination-driven self-assembly synthetic strategies.

Crystallographic studies confirm the expected *trans* arrangement of the L ligands across the planar parallelepiped Cu₂I₂ core. The UV/visible absorption spectra of the three ethylpyridinyl-based ligands and their corresponding Cu complexes were red-shifted compared to ligand precursor 4-(trimethylsilyl ethynyl)pyridine. In the complexes the magnitude of the red shift increases in the order $C_1 < C_2 < C_3$ which can be ascribed to the increasing π -conjugation length of the arylethynylpyridine ligand. DFT calculations confirm that the bands in the absorption spectra are a mixture of charge transfer from the iodine ligands to the functionalized pyridine groups and MLCT from the copper centers also to the functionalized pyridine ligands.

This study, coupled with our previous work, [5] indicates that these dinuclear copper (I) systems are amenable to coordination with a wide range of functionalized $Ar-C=C-C_5H_4N$ ligands, and opens up the possibility of synthesizing one-dimensional polymeric systems *via* further manipulation of the Ar groups to form functionalized linkers through covalent bond formation.

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