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2,2’-Dipyridylamines: More Than Just Sister Members of the Bipyridine Family. Applications and Achievements in Homogeneous Catalysis and Photoluminescent Materials

S. Wang, C. Bruneau, J.-L. Renaud, S. Gaillard, C. Fischmeister

University of Rennes. UMR CNRS 6226, Institut des Sciences Chimiques de Rennes
Université de Rennes 1 263, avenue du général Leclerc, 35000 Rennes (France)
E-mail: cedric.fischmeister@univ-rennes1.fr; christian.bruneau@univ-rennes1.fr

Normandie Univ., LCMT, ENSICAEN, UNICAEN, CNRS,
6 boulevard du Maréchal Juin, 14000 Caen (France)
E-mail: sylvain.gaillard@ensicaen.fr

Abstract

2,2’-Dipyridylamine (dpa) and related compounds belong to the family of polydentate nitrogen ligands. More than a century has passed since its first report but new complexes and applications have been emerging in the recent years owing to the versatility of dpa–based architectures. This review aims to present and highlight the main achievements attained with dpa-containing metal complexes in the domains of homogeneous catalysis and luminescent materials.

1. Introduction

Chelating ligands play a pivotal role in coordination and organometallic chemistry with multiple applications in various domains. Beyond the ubiquitous bis-pyridine, bis-imine and related ligands, bis-oxazoline and NaNac ligands (β-diketiminate), 2,2’-dipyridylamine (dpa)-based ligands have been much less explored. If the first report of dpa synthesis using sodium amide goes back to 1914, it is only in the second half of the 20th century that this class of compounds was more actively investigated, in particular in coordination chemistry owing to the multiple coordination mode they allow. Surprisingly, despite the rapid growth of N’N-chelating ligands in catalysis and material sciences, it is only recently that dpa ligands have attracted increasing interest in these domains. The facilitated synthesis of dpa-derivatives through Pd-catalysed amination reactions and the modularity of this ligand undoubtedly contributed to their increasing interest (Figure 1). The present review aims to present the recent developments and applications of organometallic complexes incorporating dipyridylamine ligands L1 in catalysis and material sciences with an emphasis on luminescent materials. When possible, complexes coordinated to the well-known bipyridine, bpy-L2, or 1,10-phenantroline, phen-L3, ligands will be compared to highlight the differences between these two classes of ligands (Figure 1).

![Figure 1 Dipyridylamine-based ligands by Pd-catalysed amination](image-url)
2. Homogeneous Catalysis

2.1 Reduction reactions

Reduction and oxidation reactions are major transformations in organic chemistry and industrial processes. Recently, these reactions have seen renewed interest with the growing development of sustainable resources and energies.\(^8\)

In 1999, the electroreductive coupling of aromatic halides was reported in ethanol using a catalyst \textit{in situ} generated from NiBr\(_2\), 3H\(_2\)O and dpa L\(_1\).\(^9\) A number of biaryl compounds were obtained in yields higher than those obtained when the catalyst was prepared with 2,2’-bipyridine (bpy) L\(_2\) as ligand. In particular, several compounds not accessible with NiBr\(_2\)(L\(_2\)) were synthesized in modest to good yields with NiBr\(_2\)(L\(_1\)) (Scheme 1, a). Similarly, the same catalyst was utilized for the conjugate addition of arylbromides on activated olefins in ethanol (Scheme 1, b).

![Scheme 1 Nickel-catalysed electrosyntheses](image)

Another example of electrocatalytic reduction was reported in 2009 by Wong.\(^10\) A pyrrole-tagged dpa ligand L\(_4\) was reacted with a rhenium precursor leading to the Re(I) complex C\(_1\) shown in Figure 2. This monomer was easily electropolymerized onto a glassy carbon electrode in acetonitrile. This electrode was found active for the electroreduction of CO\(_2\) to CO as demonstrated by cyclic voltammetry measurements performed under an argon atmosphere and under a CO\(_2\) atmosphere.

![Figure 2 Pyrrole-tagged rhenium complex C\(_1\)](image)
More recently, the electrochemical reduction of CO\textsubscript{2} was investigated with metal carbonyl complexes of tungsten and molybdenum bearing bipyridine and dpa ligands (Figure 3).\textsuperscript{11} Interestingly, catalyst \textbf{C2} outperformed the bpy-based catalysts \textbf{C3} in preparative scale reduction of CO\textsubscript{2} into CO. Mechanistic investigation also revealed a major difference between the bpy- and dpa-based catalysts. Indeed, contrary to bpy-ligand, experimental and theoretical results suggested a non-innocent behaviour of the dpa ligand with CO formation mainly localized on the dpa-ligand rather than on the metal centre.

![Figure 3](image_url)  
\textbf{Figure 3} Catalyst for electroreduction of CO\textsubscript{2} into CO

The imidazolium-tagged dpa ligand \textbf{L6} was easily prepared and used as stabiliser of Pd nanoparticles for hydrogenation of olefins under biphasic conditions (Figure 4).\textsuperscript{12}

![Figure 4](image_url)  
\textbf{Figure 4} Water-soluble dpa-based ligand

The Pd nanoparticles were easily \textit{in situ} prepared by mixing Pd(OAc)\textsubscript{2}, the tagged-dpa ligand \textbf{L6} and an ionic liquid in acetone followed by reduction of Pd(II) with dihydrogen. This catalyst could perform quantitatively the hydrogenation of cyclohexene, styrene and ethyl acrylate at 35 °C under 1 bar of dihydrogen pressure. After separation, the catalyst was reused two times without yield lessening. Water compatibility is also accessible using ionic complexes. In 2010, a series of cationic ruthenium complexes were prepared and implemented in the aqueous phase transfer hydrogenation of ketones (Figure 5).\textsuperscript{13}

![Figure 5](image_url)  
\textbf{Figure 5} Cationic ruthenium-dpa complexes
Several ketones were efficiently reduced at 65 °C in water with C6 and HCO₂H/HCOONa (Figure 6). The recyclability of the catalyst was investigated showing a rapid decrease of conversion after the second run.

\[
\text{Ar} \quad \text{R} \quad \xrightarrow{\text{C6 (0.05 mol%), HCOOH/HCOONa (5 eq./eq.)}} \quad \text{H}_2\text{O}, \ 65\ °\text{C}, \ 24\ h \quad \text{Ar} \quad \text{R}
\]

Figure 6 Aqueous phase transfer hydrogenation of ketones

Simultaneously, Pandey and coworkers reported the synthesis and characterization of a series of cationic Ru-dpa complexes with various coordination spheres consisting of 2,2'-dipyridylamine L1 or di-2-pyridylbenzylamine (dpb) L7, p-cymene and benzene as arene, Cl, N₃, CN, SCN as X ligand and PF₆ or BF₄ as counter-ion (Figure 7).¹⁴ The chloro-complexes were evaluated in the transfer hydrogenation of acetophenone and benzophenone at 82 °C using isopropanol as hydrogen source. As generally observed with ruthenium complexes in catalysis, the \( \eta^6 \)-p-cymene complex displayed better performances than the \( \eta^6 \)-benzene derivative.

\[
\begin{align*}
\text{[(η}^6\text{-arene})\text{RuCl}_2\text{]}_2 \\
\text{arene} = \text{Ar} \text{PhCH}_2\text{Me, C}_6\text{H}_6 \\
\text{R}' = \text{H (dpa) L1} \\
\text{R}' = \text{benzyl (dpb) L7} \\
\end{align*}
\]

Figure 7 Transfer hydrogenation of ketones

The group of Salmain reported a series of studies in bioinorganic chemistry involving organometallic complexes incorporating dpa ligands. The structural modularity of the ligand was first used for the synthesis of ruthenium complexes C12-C14 bearing a thiol-reactive maleimide function aimed at inactivating the cysteine endoproteinase papain by S-alkylation of the cysteine active site (Figure 8).¹⁵ These complexes and related rhodium complexes were later evaluated in the catalytic reduction of NAD⁺
into NADH. The results demonstrated the great superiority of Rh vs Ru and revealed that catalysts bearing the N-alkylated dpa ligands were not as efficient as bispyridine or phenanthroline–derived complexes of different structures.

![Figure 8 Complexes used in the reduction of NAD$^+$ using sodium formate as hydride source](image)

The modularity of the dpa ligand was further extended to the synthesis of artificial metalloenzymes derived from bovine $\beta$-lactoglobulin. A series of complexes bearing a dpa ligand N-alkylated with saturated and unsaturated fatty acids was prepared and characterized (Figure 9). These complexes were initially evaluated in the transfer hydrogenation of trifluoroacetophenone (TFACP) where rhodium catalysts yet surpassed their ruthenium analogs. The supramolecular assembling of these complexes within $\beta$-lactoglobulin was realised in EtOH or DMSO at pH 7.5 and the association was evidenced by circular dichroism and fluorescence spectroscopy. The resulting supramolecular catalysts were evaluated in the asymmetric transfer hydrogenation of trifluoroacetophenone with formate. Most complexes retained their original activity but the enantiomeric excess reached a low maximum value of 32% for the (R)-enantiomer.

![Figure 9 Artificial metalloenzyme for the asymmetric transfer hydrogenation of TFACP](image)

The potential of dpa ligands in transition metal-catalyzed reduction was extended to the valorization of bio-sourced compounds such as levulinic acid. A series of dpa-based ruthenium and iridium complexes was evaluated in the transfer hydrogenation and hydrogenation of levulinic acid (Figure 10). In the M-Cl series, transfer hydrogenation was performed under neat conditions in the presence of triethylamine. Both ruthenium and iridium complexes were shown competent for this transformation but iridium catalysts, in particular C19, were slightly more efficient as they could perform the reduction of levulinic acid at 120 °C when a temperature of 150 °C was necessary with the most efficient ruthenium catalyst C18. Nevertheless, recycling of the ruthenium catalyst C18 was achieved in six consecutive cycles without loss of activity. Much better results were obtained with zwitterionic complexes C22-C27 bearing...
a OSO₃ ligand (Figure 10). In particular, iridium catalyst C25 displayed high performances in aqueous phase transfer hydrogenation using formic acid (TON 9000) and more significantly in hydrogenation under 5 bar of hydrogen pressure at 130 °C, for which the highest TON was reported (TON 174000). Of note, it was demonstrated that transfer hydrogenation proceeded essentially by hydrogenation resulting from fast and efficient dehydrogenation of formic acid into CO₂ and H₂.

![Reduction of levulinic acid into γ-valerolactone (GVL)](image)

The zwitterionic iridium-complex C25 was further engaged in the reductive amination of levulinic acid into pyrrolidone derivatives (Scheme 2).

![Reductive amination of levulinic acid](image)

A broad range of pyrrolidone derivatives was efficiently synthesized from aliphatic and aromatic amines. In particular, sterically hindered amines could be employed leading to new bulky derivatives as shown in Figure 11. Isoindolinone derivatives were also efficiently prepared by reductive amination of 2-formylbenzoic acid. Here too bulky derivatives including new compounds were also prepared in high yields (Figure 11).
The hydrogenation of another biosourced compound, namely 5-hydroxymethylfurfural (HMF), into 1-hydroxyhexane-2,5-dione (HHD) was reported by de Vries (Figure 12). Among five catalysts evaluated, the iridium-dpa C19 furnished the highest yield of 76%. Notably, the well-known and efficient Ru-pincer catalyst C28 led to high conversion but poor selectivity for the desired product isolated in only 5%.

**Figure 11** Biosourced pyrrolidone and isoindolinone derivatives

**Figure 12** Reduction of HMF into HHD

### 2.2 Oxidation reactions

Beside reduction reactions, oxidation transformations involving dpa-ligands have received less attention. Early reports made use of dpa and dpa derivatives in ligand screening. In 1995, Griffith reported a study on the ruthenium- and osmium-catalysed oxidation of alkenes, alkanes and alcohols. Catalysts *in situ* generated from RuCl₃ and various bidentate N,N-ligands were evaluated in the epoxidation of cyclooctene using NaIO₄ as oxidant. A modest 50% yield was obtained with RuCl₃-dpa when 72% was obtained with RuCl₃-bpy. Dpa ligands were implemented in the synthesis of supramolecular polynuclear metal catalysts for oxidation of 3,5-di-tert-butylcatechol (DTBC). In an example, the copper-catalyzed oxidation of DTBC in a pure oxygen atmosphere was performed at 25 °C.
All the dpa-based catalysts displayed moderate conversion of ca. 45-50% with selectivities for 3,5-di-tert-butylbenzoquinone higher than 90%. In another example, eight dpa ligands were anchored onto a silsesquioxane cage. Three of these dpa could be coordinated to a molybdenum species leading to a silsesquioxane “T8-dpa-[Mo(η\(^3\)-C\(_3\)H\(_5\))Br(CO)]” supramolecular architecture. This compound was fully characterized and evaluated in the epoxidation of styrene and cyclooctene using tert-butyl hydroperoxide (TBHP) as oxidant, but it exhibited low reactivity. In another study, [Cu(II)(dpa)(H\(_2\)O)]\(^{2+}\) was advantageously used for the selective delignification of oxygen-prebleached Kraft pulp (OPK) using H\(_2\)O\(_2\). When compared to ligand-free CuCl\(_2\) bleaching, the Cu(II)(dpa) catalyst led to a 12-fold increase of the delignification selectivity. This improvement was assumed to result from the measured decrease of hydroxyl radical in reaction catalysed by Cu(II)(dpa) as compared to CuCl\(_2\).

In 2004, Bolm reported a study aimed at using dpa-ligands in stereoselective copper-catalyzed allylic oxidation of cyclohexene (Scheme 3). A series of C1- and C2-symmetric ligands were prepared by Pd-catalyzed amination reactions. Among the numerous ligands prepared, the C1-symmetric ligand L\(_8\) exhibited the best performances but the ee remained low (17%).

\[ \text{PhCO}_3\text{Bu} + \text{Cu(OTf)}_{2}/\text{ligand} \xrightarrow{(5 \text{ mol\% each})} \text{PhNHNNH}_2 \xrightarrow{(6 \text{ mol\%})} \text{PhNOCPh} \]

\( L_8 \), yield = 91\%, ee = 17\%

\( L_9 \), yield = 84\%, ee = 11\%

Scheme 3 Copper-catalysed allylic oxidation of cyclohexene

The straightforward functionalisation of the bridging NH of dpa was used for the immobilisation of a Pd-catalyst in water. A PEG-tagged dpa ligand was prepared and coordinated to Pd(OAc)_2. The resulting catalyst C\(_{29}\) was used for the aqueous phase aerobic oxidation of styrene resulting in the selective formation of benzaldehyde in 82\% yield along with acetophenone, benzoic acid and other unidentified compounds (Scheme 4). This catalyst led to better performances than other palladium catalysts coordinated with other N-ligands. It is noteworthy that the catalyst could be reused up to 7 times without alteration of its performances.
The catalytic properties of the \([\text{Cp}^*\text{Ir}(\text{dpa})\text{Cl}]\text{Cl}\) complex \(\text{C19}\) (Figure 10) were investigated in the very challenging and important water oxidation reaction driven by Cerium Ammonium Nitrate (CAN).\(^{28}\) Not only did this catalyst perform better than the bipyridine-based catalyst \([\text{Cp}^*\text{Ir}(\text{bpy})\text{Cl}]\text{Cl}\) but it exhibited a peculiar behaviour as CAN was always consumed faster than \(\text{O}_2\) evolution. An extreme situation was found with a CAN/Cat ratio of 80 for which CAN was almost fully consumed before \(\text{O}_2\) evolution began. The authors assumed that the bridging NH moiety of the dpa ligand might be responsible for the easier oxidative degradation of \(\text{C19}\) leading to more active catalytic sites as reported by Fukuzumi.\(^{29}\) Based on kinetic and DFT studies, the authors proposed an alternative pathway involving the formation and transfer of an \(\text{O}-\text{O}\) bond containing moiety from Ir to Ce species followed by oxygen release.

Yet in the domain of sustainable energy sources, an electron-enriched iridium complex \(\text{C30}\) was prepared and evaluated in the dehydrogenation of formic acid (Scheme 5).\(^{30}\) This catalyst achieved a TOF of 12321 at pH 1.8. Contrasting with most iridium-based catalysts which exhibited their best performances at pH 3.5-4 (i.e. pKa of formic acid), \(\text{C30}\) performed best at a pH 1.8 (i.e. no added sodium formate). It was also demonstrated that \(\text{C30}\) could maintain its performances within a concentration range going from 2 M to 8 M. Noteworthy, the catalyst could be used in pure formic acid in which a TOF of 4800 h\(^{-1}\) was reached hence demonstrating the high stability of this catalyst in acidic media. Based on the comparative study of numerous catalysts bearing diversely substituted dpa ligands, the authors proposed a mechanism involving H-bonding between formic acid and the proton of the bridging NH. Another liquid organic hydrogen carrier (LHOC), namely tetrahydroquinoline, could be efficiently dehydrogenated in water by \(\text{C30}\) leading to quinoline in 98% yield.\(^{31}\) Interestingly, the same catalyst was able to promote the reverse reaction under mild conditions of temperature and pressure (Scheme 4). This protocol was extended to numerous derivatives that could be efficiently converted in either way.
2.3 Olefin polymerisation

The nickel complexes C31-C33 (Figure 13) are highly active catalysts for the oligomerization of ethylene into C4-C6 products. These complexes required initial activation by Et₃Al₂Cl₃ in excess (Al/Ni= 600) and the reactions were performed at low temperature (12-30 °C) in dichloromethane under 0.6-1.3 bar for 1 h. With the most efficient catalysts C32 and C33, turnover numbers of about 600000 were reached with major formation of the C4-C6 fraction including more than 90% of butenes (C4-C6:C8-C16:>C16 ratio of 71:23:6 were obtained with C33). The oligomers were highly branched with a ratio of methyl protons to total alkyl protons of 0.61 for the higher boiling fraction.32

![Figure 13: Nickel(II) complexes used in ethylene oligomerization](image)

When the binuclear Ni(II) complexes C34-C38 (Figure 14) were associated with alkylaluminium reagents in ethylene oligomerization conditions, an excellent activity was obtained when Et₂AlCl was used whereas methylaluminoxane led to an inefficient catalytic system. Thus, with an Al/Ni ratio of 250, a catalytic activity leading to the production of 499-662 kg of oligomers/mol of nickel/h under 5 bar of ethylene at 30 °C was achieved corresponding to a TOF located in the range 18000-24000 h⁻¹. Remarkably, under these conditions only butenes (C4 fraction) were formed and catalyst C38 proved to be very selective affording the terminal 1-butene with 98% selectivity.33 The mononuclear cationic palladium(II) complexes C39-C43 (Figure 14) were prepared with the same 2,2'-dipyridylamine ligands and evaluated in ethylene oligomerization. Without any cocatalyst, these palladium species oligomerised ethylene at 25 °C in dichloromethane under 5 bar of ethylene. However, if the selectivity towards the formation of butenes was good, the productivity was low (TOF= 4.4 to 7.4 h⁻¹).34

![Figure 14: Nickel(II) and palladium(II) complexes used in ethylene dimerization](image)

It is noteworthy that 1-octene or 1-hexene as sole olefin were formed together with low molecular weight polyethylene wax when ethylene (40 bar) was treated at high temperature (> 80 °C) with 400 equivalents of methylaluminoxane in the presence of chromium complexes equipped with differently
substituted dipyridylamine ligands as exemplified in Scheme 6. This selectivity was ascribed to steric factors, the less hindered ligands around the metal center such as C45 producing 1-octene.

Scheme 6 Influence of steric hindrance of the dpa ligand on ethylene oligomerization

One or two dipyridylamine moieties connected to a polyethylene glycol chain in L10 and L11 were used in atom transfer radical polymerization of methyl methacrylate associated with a copper(II) halide (CuBr₂ or CuCl₂·2H₂O) as metal catalyst precursor (Figure 15). The polymerization reactions were carried out under thermo-regulated phase-transfer catalysis conditions using a water-toluene mixture as solvent. The copper-dpa catalyst was maintained in the aqueous phase at room temperature and transferred into the organic phase during the polymerization process at 90 °C. After cooling the catalyst was recovered in the aqueous phase and polymethacrylate was extracted from the organic phase.

Figure 15 N-methoxypoly(ethyleneglycol)-dipyridyl L10 and polyethyleneglycol-supported-dipyridyl Ligand (PSDL) L11

Dipyridylamines have also been supported on polynorbornene obtained by ring opening metathesis of a functionalized norbornene with a molybdenum catalyst and on coated silica. When these materials were loaded with CuBr·Me₂S or Cu(OAc)₂/Al(OiPr)₃, styrene could be polymerized in moderate yields (< 25%) following an atom transfer radical polymerization process initiated by ethyl-2-bromoisobutyrate. The cationic cobalt(III) complex [Co((L1)₂Me₂] C46 (Figure 16) proved to be an excellent catalyst for the polymerization of acrylonitrile in methanol in the presence of NaBH₄. It was shown that this polymerization did not involve a radical or an anionic mechanism but a coordination-type mechanism,
where the borohydride reduced the cobalt(III) complex into a catalytically active cobalt(I) species able to coordinate the acrylic double bond and favor successive insertions into a Co-C bond.\(^{38}\)

Figure 16 Cobalt(III) bis(L1) complex C46

Finally, the rhodium complex [Rh(cod)(L1)]PF\(_6\) was used in the polymerization of alkynes such as phenylacetylene and the functional N-propargylamine.\(^{39}\) Its catalytic properties in the formation of polyolefins from alkynes has been compared with other rhodium complexes equipped with other bidentate dinitrogen ligands such as 2,2’-bipyridine L2, 2,2’-bipyrazine and 4,4’-dimethyl-2,2’-bipyridine, and associated with other counter anions. It was shown that the nature of the ligands, counter anion as well as the presence of NaOH as cocatalyst had an influence on the fate of the polymerization. In both cases, polymerization of phenylacetylene and N-propargylamine, the [Rh(cod)(L1)]PF\(_6\) complex exhibited better performances than the corresponding [Rh(cod)(L2)]PF\(_6\).

2.4 Hydrolysis of esters

The dicationic species [Cu(L1)(H\(_2\)O)\(_2\)]\(^{2+}\) C47 was claimed as the first catalyst able to perform hydrolysis of unactivated esters such as methyl acetate at 25 °C at pH 7 with a rate constant similar to that obtained with natural esterases such as chymotrypsin.\(^{40}\) The same copper complex was also able to hydrolyze formamides under neutral pH conditions but a higher temperature was required (Scheme 7).\(^{41}\)

Scheme 7 Hydrolysis of esters and formamides in the presence of the copper(dipyridylamine) catalyst C47

Later on, the use of this type of copper catalyst was extended to more functionalized esters such as glycine ethyl ester,\(^{42}\) and other types of esters such as phosphotriesters.\(^{43}\) The carbonate protecting group of allyloxy carbonyl-coumarin has been removed upon treatment with phenylsilane in the presence of palladium-dipyridylamine catalysts such as C48 supported on polymer...
brushes (Scheme 8). This system is operating as a reservoir of catalytic species that are progressively released in solution and the influence of the thickness of the surface has been studied in detail.

![Scheme 8](image)

**Scheme 8** Hydrolysis of carbonate by PhSiH₃ in the presence of palladium(dpa) catalysts immobilized on well-defined polymer brushes

### 2.5 Transesterification

2,2'-Dipyridylamine (L1) has been coordinated to zinc, cadmium and mercury leading to various types of mononuclear complexes and polymeric forms, which have been evaluated in transesterification reactions. Thus, when Zn(OBz)₂ (OBz= benzoate) was reacted with L1, the monomeric complex Zn(OBz)₂(L1) was formed, whereas ZnBr₂ and ZnCl₂ produced hydrogen-bonded ZnX₂(L1) polymers. On the other hand, Zn(H₂O)₂(L1)(SO₄) was obtained as a coordination polymer from Zn(SO₄) and L1. All these forms exhibited catalytic activity for the transesterification of various benzoates and acetates with methanol with a catalyst loading of 4 mol% leading to complete conversion within 2-9 days into methyl benzoate and methyl acetate, respectively. The hydrogen-bonded [ZnCl₂(L1)] heterogeneous system appeared to give the best efficiencies. Later on, it was shown that the introduction of a nitrate ligand in [Zn(NO₃)₂(L1)]₂(NO₃) provided a more efficient catalyst. The substitution of zinc by cadmium has led to a variety of cadmium complexes featuring one, two or three dpa ligands. All of them were tested in transesterification reaction of acetates and benzoates with methanol and [Cd(L1)₃][ClO₄]₂ offered the best catalytic activity. Notably, the mercury complexes Hg(OBz)₂(L1) and Hg₂(L1)₂(NO₃)₂ have also shown catalytic activity for transesterification reactions.

### 2.6 Cross coupling reactions

There are only a few examples of application of 2,2'-dipyridylamine derivatives as ligands associated with palladium in cross coupling reactions and they are mostly devoted to catalysis in water or aqueous biphasic media. Thermo-regulated systems have been reported, where the central nitrogen atom of the dpa (L12) is connected to a decyloctaethyleneglycol and coordinated to a palladium centre upon treatment with
Na$_2$PdCl$_4$ to generate the neutral C49 and dicationic palladium complex C50 (Scheme 9). These complexes are readily prepared and show different solubility properties due to the presence of the amphiphilic ligand, the neutral C49 is insoluble in water whereas the ionic C50 is soluble. Heck reaction were carried out at 110-120 °C in biphasic aqueous media with in situ generated catalysts based on Na$_2$PdCl$_4$ and an excess of ligand. The organic phase consisted of the substrates without additional solvent. The catalytic reactions took place in this organic phase and after cooling the catalyst was transferred to the water phase and the products isolated from the organic phase. Starting from iodobenzene, E-cinnamic acid was isolated in 89% yield after 20 h from ethyl acrylate, whereas trans-stilbene was recovered in more than 90% yield after 40 h from styrene. The catalyst recovered in the aqueous phase could be reused but its activity decreased probably because of accumulation of salts in this phase.

Scheme 9 Heck cross coupling with thermo-regulated catalytic systems

The copper-free Sonogashira coupling was also performed efficiently at room temperature in water with an in situ generated catalyst based on equimolar amounts of Pd(NH$_3$)$_2$Cl$_2$ as metal precursor and L1 as bidentate ligand (Scheme 10). The reaction required the presence of triethylamine as a base, other organic amines such as diisopropylamine or tributylamine being inefficient as well as the classical KOH, NaOAc, K$_3$PO$_4$, Cs$_2$CO$_3$. In this study, it was shown that the dipyridylamine ligand triggered much higher catalytic activity than other bidentate nitrogen ligands such as 2,2’-bipyridine L2, 1,10-phenanthroline L3, 2-imidazol-2-ylpyridine and 2-pyrazol-3-ylpyridine. The cross-coupling reaction of phenylacetylene and electron-enriched phenylacetylenes with various phenyl iodides gave more than 73% yield of the corresponding 1,2-diarylacetylenes whereas lower yields located in the range 59-73% were obtained with the electron-deficient 4-acetylphenylacetylene.
Scheme 10 Sonogashira coupling promoted by a L1-containing palladium catalyst

The Suzuki-Miyaura coupling was also achieved in water with palladium complexes equipped with ionic water soluble dipyridylamine ligands. These complexes were obtained by treatment of Pd(OAc)$_2$ with equimolar amounts of ligands L13-L15 in dichloromethane (Scheme 11).$^{51}$

Scheme 11 Suzuki-Miyaura coupling in water soluble palladium catalysts based on ionic dipyridylamines

Pd(OAc)$_2$(L15) was the most active catalyst for the cross coupling of aryl bromides and chlorides with arylboronic acids indicating a clear effect of the pendant ionic group of the ligands. A turnover frequency of 96000 h$^{-1}$ was reached during the coupling of 4-bromoacetophenone and phenylboronic acid at 100 °C in water in the presence of 0.001 mol% of this catalyst leading to 96% yield of 4-acetylbiphenyl within 1 h. Arylbromides containing electron-withdrawing groups in 4-position gave the cross coupling products in excellent yields in short reaction times (less than 1 h) with 0.1 mol% catalyst loading. In addition, the catalyst tolerated the presence of withdrawing and donating functional groups in the arylboronic acid partner without loss of reactivity. Starting from aryl chlorides required more drastic conditions including longer reaction times and the presence of a phase transfer agent such as tetra-$n$-butylammonium bromide. In these systems, nanoparticles with small size and narrow size distribution stabilized by the ionic dipyridylamine ligands were detected at the end of the reactions.

Ring opening copolymerization of (Z)-9-oxabicyclo[6.1.0]non-4-ene with tris(cyclooct-4-enyl-1-oxy)methylsilane with a Grubbs third generation ruthenium catalyst has been used to prepare palladium nanoparticles embedded in the pores of a polymeric support (Scheme 12).$^{52}$ This material CS1 has shown catalytic activity for the Suzuki-Miyaura cross coupling of bromobenzene derivatives with phenylboronic acid at 50 °C for 24 h in water in the presence of tBuOK as a base and tetra-$n$-butylammonium bromide as phase transfer agent. Assuming that only the surface palladium atoms participated in the catalytic reaction, turnover numbers up to 2940 were reached. On the other hand, this catalyst was much less...
efficient for the copper-free cross coupling of the same bromobenzene substrates with phenylacetylene at 80 °C leading to a moderate 36% yield, whereas iodobenzene proved to be much more reactive providing 95% yield upon reaction with 2,4,6-trimethylphenylacetylene.

Scheme 12 Palladium nanoparticles immobilized within small pores functionalized by dipyridylamine groups

Finally, nickel species generated in situ from equimolar amounts of NiBr₂.3H₂O and L₁ have been used for the reductive coupling of aryl halides in ethanol at room temperature under electrolysis conditions at a constant current of 0.6 A.dm⁻² (Scheme 13). Under similar conditions, the reductive cross coupling of arylbromides with electron-deficient olefins such as acrylates and ethyl maleate was also achieved in good yield, whereas acrylonitrile and ethyl methacrylate were not reactive (Scheme 13).⁵³
Dipyridylamine copper complexes have been used as catalysts for the oxidative coupling of 2,6-dimethylphenol. Depending on the nature of the catalyst precursor and the experimental conditions, 3,3',5,5'-tetramethyl-4,4'-diphenoquinone (DPQ) and poly(2,6-dimethyl-1,4-phenylenether) (PPE) could be formed (Scheme 14). Under mild conditions, e.g. 25 °C under atmospheric pressure of dioxygen in a methanol/toluene mixture as solvent, the copper complex C52 (catalyst loading: 0.4 mol%) led to the selective formation of the PPE polymer.\(^{54}\)

The oxidation of 2,6-dimethylphenol was also studied with \(\text{H}_2\text{O}_2\) as oxidant and the copper complex with the proposed structure C53 as catalyst in aqueous buffer in the range of pH 6.0-9.0 at 25 °C. It was shown that the L1 ligand provided the most efficient catalyst and that the initial rate of formation of DPQ strongly depended on the value of the pH, the maximum being located around pH 7.\(^{55}\) The introduction of surfactants in the reaction medium such as cetyl trimethylammonium bromide leading to the formation of micelles had a beneficial effect on the rate of oxidation of 2,6-dimethylphenol and favored the formation of PPE polymer.\(^{56}\)
Scheme 14 Oxidative coupling of 2,6-dimethylphenol with copper catalysts

2.7 Miscellaneous reactions

Photocatalyzed aza-Henry reaction

Among a series of cationic iridium complexes featuring two N,C-cyclometallated ligands based on benzoquinoline, 2-phenylisoquinoline, and dpa L1, the iridium complex C54 has revealed efficient catalytic properties for the photocatalyzed aza-Henry reaction involving N-phenyltetrahydroisoquinoline and nitromethane (Scheme 15).

In the presence of 1 mol% of catalyst C54, full conversion of the aromatic substrate was achieved and 69% of product resulting from selective sp3C-H bond activation/functionialization at the benzylic position of the tetrahydroisoquinoline ring was obtained at room temperature under air within 11 h. This result competes with the 79% isolated yield obtained with the more active [Ir(2-phenylpyridine)2(di-tert-butylbipyridine)][PF6] catalyst used for comparison.

Scheme 15 Aza-Henry reaction photocatalyzed by the iridium catalyst C54
**Hydroformylation of olefins**

Very early on, 2,2’-dipyridylamine L1 has been used in hydroformylation of olefins for comparison purposes with other bidentate nitrogen ligands such as 2,2’-bipyridine L2, 1,10-phenanthroline L3. For instance the hydroformylation of 1-octene at 180 °C with a pressure of CO/H₂ (1:2 ratio) of 80 bar in the presence of 3 mol% of RuO₂-L1 as catalyst precursor, led to a mixture of branched and linear nonanals and nonanols with major formation of alcohols, and a reactivity and selectivity comparable to the results obtained with bipyridine ligands. On the other hand, the catalytic species generated from Ru₃(CO)₁₂ and L1 on silica carrier was active in hydroformylation of 1-hexene at 150 °C under 50 bar of syngas and led to heptanals as major products but the productivity remained lower than when bipyryne was used.

3 Photoluminescent materials

A rapid survey of the literature shows that around 700 complexes bearing dipyridylamine derivatives as ligand are reported in the literature. These include chromium, molybdenum, manganese, rhenium, iron, ruthenium, osmium, cobalt, rhodium, iridium, nickel, palladium, platinum, copper, silver, gold, zinc, cadmium, and mercury as metal centers. Among this plethora of transition metal complexes, chromium, rhenium, ruthenium, osmium, rhodium, palladium, platinum, copper, silver, zinc, cadmium, and mercury complexes have been reported to emit or absorb in the visible region. These photophysical properties render this class of organometallic complexes interesting for ion or organic molecule detection by selective binding of cation accompanied by a luminescent response, for electroluminescent material such organic light-emitting diodes (OLEDs) or light-emitting electrochemical diodes (LECs), for nuclear detection via plastic scintillators and finally for dye-sensitized solar cells (DSSCs) by using the absorption properties in the visible region.

3.1 Ion and organic molecule detection

Regarding the ion or organic molecule detection, two strategies can be considered, (i) the fluorescence (or phosphorescence) extinction when an ion binds or interacts with the emissive complex; and (ii) the generation or strong modification of fluorescence (or phosphorescence) when an ion binds or an organic molecule interacts with a non-emissive complex. The most representative examples are based on strategy (i). However, some dpa derivatives are known to be intrinsically fluorescent and, as mentioned above, several transition metal complexes possess fluorescence properties.

As first example, a star-shaped ligand L16 (Figure 17) possessing a benzene core and 6 biphenyl-2,2’-dipyridylamine substituents has been synthesized and studied in self-assembling on graphite surface. In the presence of a copper(II) salt, ligand L16 leads to a non-emissive 2D extended structure. The quenching of the fluorescence property of L16 (emission wavelength (λ_em) = 409 nm in dichloromethane solution) was reached when a 1:2 ratio of L16:CuII was used. Unfortunately, no other cation has been studied to show selectivity for such extinction selectivity.
Simpler structures having two to three dpa units (ligands L17-L20, Figure 18) and the corresponding copper(I) complexes have also been described (Figure 19). Various copper salts (CuI, CuBr and CuCN) were investigated and, surprisingly, depending on the salt, various polymetallic structures were characterized. As copper(I) can coordinate two dpa ligands, organometallic polymers were formed adopting different conformations (trans, cis, inward- and outward-conformations), depending of the copper(I) precursor and the ligand/metal ratio (Figure 19).

Figure 17 Structure of the star-shaped ligand 16

Figure 18 Structures of poly-dpa derivatives L17-L20

Figure 19 Coordination modes of ligands L17-L20
Notably, the two pyridine units were not always coordinated to the copper(I) metal center. In some cases, each pyridine bond a different metal center leading to a 2D Metal Organic Framework (MOF).

Interestingly, a complete extinction of the blue fluorescence observed with the free ligand \( \text{L17} \) (\( \lambda_{\text{em}} = 423 \) nm (main band) and 517 nm (shoulder) in dichloromethane solution) was observed with the organometallic polymer generated with ligand \( \text{L17} \). The emission spectra of \( \text{L17} \) in the presence of different concentration of copper(I) salt showed a drastic decrease of the fluorescence until a 1:6 ratio of \( \text{L17}:\text{Cu(I)} \). In this work, no selectivity in binding metal salts was reported. Such study was demonstrated with a second family of poly-dpa ligands, namely 1,2-\( \text{L21} \), 1,3-\( \text{L22} \), 1,4-\( \text{L23} \), 2,6-bis-(2,2'-dipyridylaminomethyl)pyridine \( \text{L24} \) and 1,3,5-tris-(2',2''-dipyridylaminomethyl)benzene \( \text{L25} \) for bulk membrane transport (Figure 20).97 The coordination properties of these polydentate ligands were investigated with various silver(I) salts. The selectivity in binding metal ions depends on the nature of the fluxional core. Indeed, in the presence of a mixture of transition metal salts (Co(II), Ni(II), Cu(II), Zn(II), Cd(II), Zn(II), Ag(I) and Pb(II)), dpa derivatives \( \text{L22-L23} \) and \( \text{L25} \) bind selectively the silver(I) cation. Unfortunately, no emission of the ligands or of the resulting silver(I) complexes have been reported.

![Figure 20](image)

Figure 20 Structures of poly-dpa derivatives \( \text{L21-L25} \).

In 2000, the coordination of 2,2',2''-tripyridylamine \( \text{L26} \) and 2,2',3''-tripyridylamine \( \text{L27} \) with various Zn(II) salts has been studied (Figure 21).98

![Figure 21](image)

Figure 21 Structures of ligands \( \text{L26} \) and \( \text{L27} \)

The ligand \( \text{L26} \) acted as a chelate with ZnCl\(_2\) (1 equivalent of \( \text{L26} \)) or Zn(OTf)\(_2\) (2 equivalents of \( \text{L26} \)) whereas only the 3-substituted pyridine moiety in ligand \( \text{L27} \) bound Zn(OTf)\(_2\). The authors stated that the different mode of coordination (four ligand \( \text{L27} \) mono coordinated to the zinc metal centre) was due to some steric hindrance. Interestingly, all the corresponding Zn(II) complexes exhibited a blue emission (\( \lambda_{\text{em}} \) between 422 to 432 nm).

Later, the coordination behavior of the 2,2',3''-tripyridylamine ligand \( \text{L27} \) has been studied with different silver(I) and zinc(II) metal salts.99 Various dinuclear complexes or 1D organometallic polymers were obtained with silver depending on the nature of the anion (CF\(_3\)CO\(_2\), TfO\(^-\), PF\(_6\) or ClO\(_4\)\(^-\)). Silver complexes are in a general manner di or tri-coordinated to two or three ligands \( \text{L27} \). Unlike zinc which was able to bind 3-substituted pyridines, ligand \( \text{L27} \) bind nonspecifically silver metal centre by the 2- or 3-substituted pyridine rings. The emissions of these silver complexes were observed between 366 to 368 nm in acetonitrile solution when ligand \( \text{L27} \) was emitting at 361 nm. The silver complexes exhibited also lower photoluminescent quantum yield (\( \phi_{\text{em}} \) between 0.16 to 0.18 for silver complexes vs. 0.38 for
ligand L27). This property allowed the silver ion detection through the analysis of the decrease of the fluorescence intensity. Good correlation with the silver concentration (up to 4 mM) was obtained. Noteworthy, no decrease of fluorescence was observed in the presence of zinc(II) salts.

For organic molecule detection, Wang, Brown and co-workers reported the synthesis and photophysical properties of a zinc(II) complex in which the star-shaped ligand L28 held a 1,3,6-tris-(4'-dpa)phenylbenzene core L28 (Figure 22).88i

In the presence of three equivalents of ZnCl₂, Ligand L28 led to the formation of the complex [(ZnCl₂)₃ L28]. This complex exhibited a broad fluorescence with an emission wavelength λₑₘ at 417 nm in solution and 432 nm in the solid state. Then, complex [(ZnCl₂)₃ L28] has been introduced in an optical sensor and twelve different solvents (500 ppm) were selected to measure the decrease of the fluorescence intensity of the zinc(II) complex. As a result, benzene led to an important decrease of the fluorescence intensity whereas the structurally close solvents xylene and toluene almost did not alter its fluorescence intensity. Other protic polar solvents, alkanes or halogenated solvents did not modify the fluorescence. Based on crystallographic data, the affinity of benzene with [(ZnCl₂)₃ L28] was observed through π-π interactions between aromatic ring and the core of the ligand L28. These interactions might be responsible for the fluorescence quenching.

Finally, a dpa derivative based on guanosine has also been investigated for zinc salt detection.100 The fluorescence of the ligand L29 decreased upon addition of 6 equivalents of Zn(ClO₄)₂·6H₂O or Zn[(S)-O₂CCH(Br)CH(CH₃)CH₃] without any shift of the emission wavelength, while a red-shift (from 403 nm to nearly 490 nm for λₑₘ) and a decrease of the fluorescence intensity were observed with the analogous ligand L30. This double effect rendered the detection of the zinc(II) salt easier (Figure 23).
3.2 Electro and photoluminescent materials

Fluorescent or phosphorescent transition metal complexes can also find applications in electroluminescent materials such as OLED and LEC. Both devices are dedicated for light applications and the most important difference is that LEC devices are specific of ionic transition metal complexes which lead to simpler architecture device as no injection and transport electron or hole layers are required.

In 2010, Huo et al. synthesized dpa-based ligands \( \text{L31} \) and \( \text{L32} \) bearing either a phenyl or a \((2',4'-\text{difluorophenyl})\) substituent and the corresponding cyclometallated platinum complexes \( \text{C55-C56} \) (Figure 24).

![Figure 24 Structure of ligands L31 and L32 and platinum(II) complexes C55 and C56](image)

The phosphorescent emissions of such platinum complexes have been assigned by the author to an admixture of intraligand charge transfer (ILCT) and metal to ligand charge transfer (MLCT). The \( \text{C55} \) complex exhibited a high quantum yield of 0.74 with two maxima for the emission wavelengths at 512 and 548 nm (in 2-methyltetrahydrofuran solution). Due to its highly emissive properties \((\varphi_{\text{em}})\), complex \( \text{C55} \) has been investigated in OLED technology with an optimized device configuration consisting in ITO/\( \text{CF}x\)/NBP/TCTA/(TPBI + 30 wt % TCTA)+ 4 wt % \( \text{C55}/\text{TPBi/Alq/Mg:Ag} \) (ITO: indium tin oxide; \( \text{CF}x\), NPB: \(4,4'-\text{bis[N-(1-naphthyl)-N-phenylamino]biphenyl} \); TCTA: \(4,4',4''\text{-tris(N-carbazolyl)-triphenylamine} \); TPBi: \(1,3,5\text{-tris(N-phenylbenzimidazol-2-yl)benzene} \). The best device showed bright green emission \((\lambda_{\text{em}}/\text{CIE coordinates } (x,y)): 512 \text{nm}/0.318,0.622)\) with external quantum efficiency \((\text{EQE})\) of 14.7% and current efficiency of 50.0 \(\text{Cd.A}^{-1}\) at a current density of 0.01 mA.cm\(^{-2}\). The highest luminance \(L\) of 3698 Cd.m\(^{-2}\) was observed at a current density of 10 mA.m\(^{-2}\) (at 8.64 V) but also led to a decrease of the \(\text{EQE}\) to 10.6%.

In direct line with the previous study, the unsymmetrical dpa derivatives \( \text{L33-L35} \) were prepared and the corresponding platinum(II) complexes evaluated in the preparation of new OLED devices (Figure 25).
The three platinum(II) complexes C57-C59 (Figure 25) were green emitters with λ_{em} ranging from 504 to 513 nm with a shoulder centered approximately at 535 nm (in degassed dichloromethane). Complexes with ligands L33 and L34 exhibited photoluminescent quantum yields of 0.27, when C59 complex displayed a quantum yield of 0.47 (in degassed dichloromethane relative to fac-[Ir(ppy)$_3$], $\phi_{em}$ = 0.4). The OLED devices possessed the following architecture ITO/HAT-CN/TAPC/TCTA/mCP: Pt(II) (10 wt %)/TmPyPB/Liq/Al (HAT-CN: 1,4,5,8,9,11-hexaaazatriphenylene-hexacarbonitrile; TAPC: 1,1-[4-[N,N-di(p-tolyl)-amino]phenyl]cyclohexane; mCP: 1,3-bis(N-carbazolyl)benzene; TmPyPB: 1,3,5-tri[3-pyridyl]phen-3-yl]benzene; Liq: 8-hydroxyquinolinolato lithium). The emissions of the devices were slightly red-shifted compared to the emissions in solution. Indeed, C57 emitted greenish yellow light (λ_{em}/CEI coordinates (x,y): 548 nm/0.44,0.53) with EQE of 11.8 %, current efficiency of 38.8 Cd.A$^{-1}$ and power efficiency of 20.2 lm.W$^{-1}$ at brightness of 1000 Cd.m$^{-2}$. C58 and C59 complexes showed yellowish-green emission (λ_{em}/CEI coordinates (x,y): 540 nm for both complexes/0.36,0.58 and 0.32,0.61, respectively) with EQE of 11.5 and 11.9, current efficiency of 38.5 and 42.4 Cd.A$^{-1}$, and power efficiency of 23.7 and 29.4 20.2 lm.W$^{-1}$ at brightness of 1000 Cd.m$^{-2}$, respectively.

The ligand L36 having 6,6’-trifluoromethylpyrazolyl substituent has been coordinated to an osmium(II) metal center associated with phosphine ligands (Figure 26). The two synthesized complexes [Os(L36)(PMePh$_2$)$_2$] and [Os(L36)(PMe$_2$Ph)$_2$] exhibited strong red emission at λ_{em} 628 and 634 nm with $\phi_{em}$ of 0.40 and 0.27 (in degassed dichloromethane), respectively. Due to these strong deep red emissions of complexes [Os(L36)(PMePh$_2$)$_2$] and [Os(L36)(PMe$_2$Ph)$_2$], organic light emitting diodes (OLEDs) have been prepared with the following architecture ITO/NPB/TCTA/TPBi: [Os(L36)(PMePh$_2$)$_2$] or [Os(L36)(PMe$_2$Ph)$_2$] (4 vol%)/TPBi/LiF/Al (NPB: 4,4’-bis[N-(1-naphthyl)-N-phenylamino]biphenyl; TCTA: 4,4’,4”-tris(N-carbazolyl)-triphenylamine; TPBi: 1,3,5-tris(N-phenylbenzimidazol-2-yl)benzene). Additionally, to the devices prepared with complexes [Os(L36)(PMePh$_2$)$_2$] C60 and [Os(L36)(PMe$_2$Ph)$_2$] C61, a device doped with [Ir(piq)$_3$] was prepared for comparison (piq: phenyl isoquinoline). All devices prepared with [Ir(piq)$_3$], C60 and C61 exhibited a deep red emission with CIE coordinates at (0.67,0.33),
(0.63, 0.36) and (0.65, 0.35) with current density of 5 mA cm$^{-2}$, respectively. Both devices prepared with complexes $\text{C60}$ and $\text{C61}$ showed better performance than the device with $[\text{Ir(piq)}_3]$ as phosphorescent dopant. The highest current efficiency of 14.0 Cd A$^{-1}$ at 5 mA cm$^{-2}$ and best EQE of 9.8% were observed with the device prepared with complex $\text{C60}$. However, the devices prepared with Os(II) complexes required higher operation voltage than with Ir(III) complex due to the shallow HOMO level of Os(II) which may induce a large carrier injection barrier and charge trap.

The tridentate extended dpa derivatives $\text{L37}$ incorporating an additional 2-pyridylamino framework on the dpa motif has been extensively studied by the group of Heinze and the properties of the corresponding complexes have been compared with their analogous complexes bearing the tripyridine ligand $\text{L38}$ (Figure 27).$^{78c,104}$

Among these complexes, the photoluminescent ruthenium(II) complexes $\text{C62-C64}$, bearing either $\text{L37}$ or $\text{L38}$ ligand in combination with an ester-substituted terpyridine ligand, have been studied in light emitting electrochemical cell (LEC) technology (Figure 28).$^{105}$

The configuration of the material was based on the following structure ITO/PEDOT:PSS/ruthenium(II) complex/Ag (PEDOT: poly(3,4-ethylenedioxythiophene); PSS: poly(styrenesulfonate). Different devices have been prepared with or without 20 wt% PMMA (poly(methylmethacrylate) as additive with complexes $\text{C62}$ or $\text{C63}$ as phosphorescent dopant of the emissive layer. At low voltage (3V), only the device incorporating complex $\text{C63}$ without PMMA emitted in the red region ($\lambda_{em}$; CIE coordinates $(x,y)$:}
731 nm; 0.725, 0.275). At higher voltage (up to 6V), the devices with PMMA as additive and incorporating the Ru(II) complexes C62-C63 emitted in the red to near infra-red ($\lambda_{em}$/CIE coordinates ($x,y$): 733 nm/0.709-0.291 for C62 at 4V; 722 nm/0.717-0.283 for C63 at 6V; 745 nm/0.729/0.271 for C64 at 5V). The authors stated that these tridentate ligand-ruthenium complexes containing devices were the LECs possessing the longest emission wavelength. Regarding the highest performances of these LECs devices, luminance ($L$) and external quantum efficiency ($EQE$) were obtained with the device incorporating complex C63 at a voltage of 6V and was measured at 3.4 cd.m$^{-2}$ and 0.028 %.

Photoluminescent cationic copper(I) complexes bearing dpa ligand L1 in association with N-heterocyclic carbenes (NHC)$^{86}$ have also been studied in LEC technology. The cationic copper(I) complex C65 (Figure 29) exhibited a bright and blue emission with $\lambda_{em}$ at 463nm.

![Figure 29](#)

**Figure 29** Structures of complexes C65-C68

The $\lambda_{em}$ was ascribed to a MLCT and presented a large Stokes shift. The analog complexes [Cu(bpy)][IPr][PF$_6$] (C66)$^{86}$ or [Cu(phen)][IPr][OTf] (C67)$^{106}$ (Figure 29) both exhibited a weak and red-shifted $\lambda_{em}$ around 650 nm. This important difference in the $\lambda_{em}$ is due to the strong electron donating property of the central nitrogen atom in dpa ligand. Indeed, the DRX analyses of these different copper(I) complexes showed that the dpa ligand is planar in a general manner indicating that the p orbital of the nitrogen atom is parallel to the p orbital of the pyridine ring thus allowing delocalization of the electrons in the pyridine rings. This structural observation was also confirmed by DFT and TD-DFT calculations.$^{107}$

DFT calculations have also shown that the HOMO of complex C65 was located on the metal center and the LUMO mainly located on the dpa ligand. In this complex, the $\lambda_{em}$ is ruled by the nitrogen ligand, unlike the bipyridine or phenanthroline complexes. The introduction of electron withdrawing groups, stabilizing the LUMO, generated a red shift of the $\lambda_{em}$ whereas electron donor groups, destabilizing the LUMO, generated a blue shift of the $\lambda_{em}$. Taking advantage of the blue luminescence of the family of (NHC)copper(I) complexes bearing 2,2'-dipyridylamine ligands, one of the brightest complexes having photoluminescent quantum yield ($\varphi_{em}$) up to 0.86 has been applied in the fabrication of light emitting electrochemical cells (LECs).$^{108}$ Devices obtained with complex C68 were the first LEC incorporating cationic copper complexes having blue emission (with CIE coordinate of 0.23/0.28) showing figure of
merit similar to those obtained for LECs incorporating iridium complexes, at that time. Nevertheless, their lifetime $t_{1/2}$ were short with best value of 16.5 min.

In the context of nuclear detection with plastic scintillator, dpa ligands have also been used as ancillary ligands for cationic cyclometallated iridium(III) complexes (cyclometallating ligands = 2-phenylpyridine, ppy; $N$-phenylpyrazine, ppz; benzoquinoline, bzq or phenylisoquinoline, piq) (Figure 30). The photophysical properties of the corresponding complexes have been studied and compared to those of complexes incorporating a bipyridine L2 or phenanthroline L3 ligand.

Considering the absorption spectra, the profile appeared to be mainly driven by the cyclometallating ligand. Nevertheless, replacing bipyridine L2 or phenanthroline L3 ligands by dpa L1 led in most cases to a blue shift of the $\lambda_{em}$ originating from a MLCT. Some of these cationic iridium(III) complexes have been selected as dopant for the preparation of plastics. Due to solubility issue of these complexes in the styrene matrix, samples were prepared with concentration below or equal to 0.05 wt%. Among the plastics obtained, those incorporating dpa-iridium complexes exhibited emission resulting only from the styrene matrix due to the low $\phi_{em}$ of the dopant. Best results of radio- and photoluminescence were here obtained with plastic incorporating bipyridine or phenanthroline iridium(III) complexes and showed scintillation yields between 400 to 1500 ph.MeV$^{-1}$.

### 3.3 Application in Dye-Sensitized Solar Cells

Dipyridylamine metal complexes have been also studied as catalysts in photoredox chemistry thanks to their photophysical properties, and more specifically their absorption in the visible region, but also in material science and more particularly as component for dye-sensitized solar cells (DSSCs). To the best of our knowledge, the first example was reported in 2009 by Masuda and co-workers. The dpa derivative L39 was used as ancillary ligand of a ruthenium(II) complex C69. Both the tert-butyl and methoxy substituent on the $N$-phenyl-2,2'-dipyridylamine modified the HOMO and LUMO energy level and increased the lifetime performance of the device (Figure 31).
The absorption spectrum of complex **C69** showed two bands centered at 381 nm (molar extinction coefficient \( \varepsilon = 14.7 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \)) and 514 nm (\( \varepsilon = 49.4 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \)) which are assigned to MLCT bands. The ionization potential (\( E_{\text{ox}} \)) was measured at -5.1 eV, which is 0.3 eV more negative than the one of I2/I3. The excited-state ionization potential (\( E_{\text{ox}}^* \) or \( E_{\text{LUMO}}^* \)) of **C69**, obtained from zero-zero excitation energy (\( E_{\text{0-0}} \)), was approximated at -3.3 eV, which is 0.6 eV more positive than the TiO2 conduction band. Devices were prepared with complex **C69** and the so-called N719 complex **C70** (\( E_{\text{ox}} = -6.01 \text{ eV and } E_{\text{ox}}^* = -3.64 \text{ eV} \)). The device prepared with **C69** led to a short circuit current density (\( J_{\text{SC}} \)) of 17.3 mA.cm\(^{-2} \), an open-circuit photo-voltage (\( V_{\text{oc}} \)) of 640 mV and a fill factor (\( FF \)) of 0.65, giving an overall conversion efficiency (\( \eta \)) of 7.2%. Interestingly, these performances were slightly lower than the one observed under the same conditions with the ruthenium complex **C70**.

The group of Kim modified the dipyridylamine ligand by replacing the \( N \)-para-methoxy phenyl substituent by an aliphatic chain and the \( t \)Bu groups of the dpa skeleton by a (hetero)aryl conjugated framework (ligands **L40** and **L41**, Figure 32).113 The corresponding complexes **C71** and **C72**, analogs of **C69**, have been prepared and studied in solar cell devices.

**Figure 32** Structure of ligands **L40** and **L41**, and ruthenium complexes **C71** and **C72**.

Absorption properties of complexes **C71** and **C72** were as follows, maxima absorption wavelengths (\( \lambda_{\text{abs}} \)) of 418 nm (\( \varepsilon = 16.9 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \)) and 531 nm (\( \varepsilon = 10.9 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \)) for **C71** and 420 nm (\( \varepsilon = 24.4 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \)) and 531 (\( \varepsilon = 15.4 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \)) for **C72**. Note, the molar extinction coefficient of **C72** was higher than that of **C70** (\( \varepsilon = 14.9 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \) at \( \lambda_{\text{abs}} = 380 \text{ nm and } \varepsilon = 14.4 \times 10^3 \text{ M}^{-1}\text{cm}^{-1} \) at \( \lambda_{\text{abs}} = 520 \text{ nm} \)). The \( E_{\text{ox}} \) of **C71** and **C72** were measured at 0.95 and 0.92 V vs. NHE (-3.49 and 3.52 eV vacuum scale)) and \( E_{\text{ox}}^* \) approximated at -1.00 and -0.96 V vs. NHE (-5.44 and 5.40 eV vacuum scale), respectively. Solar cells devices with **C71** and **C72** were prepared and compared with device using N719. In the following conditions (0.18 cm\(^2\) working area; electrolyte: 0.6 M DMPII, 0.05 M I2, 0.1 M LiI and tert-butylpyridine in acetonitrile), the performances were \( J_{\text{SC}} \) of 17.1 and 18.89 mA.cm\(^{-2} \), \( V_{\text{oc}} \) of 660 mV (in both case), and \( FF \) of 0.69 (in both case), corresponding to \( \eta \) of 7.90 and 8.76 %, respectively. Of note,
the device prepared with C72 exhibited higher performance parameter that the one prepared with C70 (η of 8.76 vs. 8.19 % for C72 vs C70) and was attributed to the higher ε of C72 compared to C70.

Ruthenium complexes C73-C75 containing ligands L37 (Figure 27) and L42, represented in Figure 33, are related to the above-mentioned complexes C62-C64 (Figure 28). Their deep-red emissions allowed some studies in lighting devices, and their absorption properties in the UV and visible regions fitted with DSSCs applications.

Indeed, complexes C73-C75 showed several maxima absorption wavelengths assigned to MLCT bands in the visible region between 472 and 680 nm. Nevertheless, the ε calculated for these complexes were at least three-fold lower than that of C70. Even if photostability was more important for complexes C73-C75 compared to N719 complex C70, the performance parameters of DSSCs with complexes C73-C75 as dyes were twenty-fold lower than those of device prepared with C70 (η = 5.0 % for N719 vs. η = 0.26 % for complex C63 for example). These poor DSSCs performances were attributed to electron recombination with the oxidized electrolyte (I3−) at the TiO2 electrode. The positive charge on the ruthenium(II) complexes might attract the negatively charged I3− near the TiO2 surface and consequently might provoke undesired back electron transfer. To tackle this issue, positively charge Co3+/Co2+ couples, [Co(bipy)3]2+/3+ and C76, were envisaged to replace the negatively redox mediator I3−/I2−. Using complex C74 as dye, several DSSCs were prepared with electrolytes involving [Co(bipy)3]2+/3+ or C76 as positively charged redox mediator and the I3−/I2− for comparison (Figure 34).

Figure 33 Structures of ligands L42 and Ruthenium(II) complexes C73-C75

Figure 34 Structures of Co3+/Co2+ redox mediators [Co(bipy)3]2+/3+ and C76
DSSCs prepared with [Co(bipy)]$_{2}^{2+/3+}$ showed lower efficiency than devices using I$_{3}$/I$_{2}$ as redox mediator, the one elaborated with C76 increased both $I_{SC}$ and $V_{OC}$ and led to an efficiency two-fold higher. To explain the different results with the [Co(bipy)]$_{2}^{2+/3+}$ and C76 redox mediators, the authors proposed that the efficiency was correlated to the π$^{*}$ orbitals energy levels which were higher in the case of the electron rich ligand L39 than with bipyridine ligand. Importantly, the enhancement of the DSSC efficiency using C76 as redox mediator depended on the dye as shown later by the same group using another ruthenium(II) complex as dye and for which I$_{3}$/I$_{2}$ was found better than C76.$^{116}$

4 Conclusion

2,2'-Dipyridyliamine and 2,2-dipyridyliamine-based metal complexes have been implemented with success in a number of catalytic processes and light-emitting materials and devices. In this review, we have demonstrated that these ligands sometimes lead to improved performances when compared to the ubiquitous 2,2'-bipyridine-based organometallic complexes. One major difference between these two family of ligands is found in the formation of metallacycles with 5- or 6-members, but may be more importantly, the bridging nitrogen atoms in dpa is a direct and easily accessible manifold to modulate the electronic and steric properties of dpa-based ligands. Furthermore, the nitrogen atom is also an anchorage point for supported catalysis of for the design of task specific ligands. Last but not least, the presence of a bridging N-H group at the proximity of a catalytic centre may play a crucial role for substrate orientation/activation and product release through H-bonding. Altogether, these properties make dpa a ligand of high potential which should be considered and utilized with the same attention and interest as other well-stablished N,N-chelates. It is likely that the future years will see an increasing interest for this so far neglected ligands.

5 References


