NHC-Based Iron Sensitizers for DSSCs

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Abstract: Nanostructured dye-sensitized solar cells (DSSCs) are promising photovoltaic devices because of their low cost and transparency. Ruthenium polypyridine complexes have long been considered as lead sensitizers for DSSCs, allowing them to reach up to 11% conversion efficiency. However, ruthenium suffers from serious drawbacks potentially limiting its widespread applicability, mainly related to its potential toxicity and scarcity. This has motivated continuous research efforts to develop valuable alternatives from cheap earth-abundant metals, and among them, iron is particularly attractive. Making iron complexes applicable in DSSCs is highly challenging due to an ultrafast deactivation of the metal–ligand charge-transfer (MLCT) states into metal-centered (MC) states, leading to inefficient injection into TiO₂. In this review, we present our latest developments in the field using Fe(II)-based photosensitizers bearing N-heterocyclic carbene (NHC) ligands, and their use in DSSCs. Special attention is paid to synthesis, photophysical, electrochemical, and computational characterization.

Keywords: iron complexes; NHC ligands; excited states; photophysics

1. Introduction

Replacement of fossil fuels is one of the major concerns mankind is currently facing. To keep fulfilling an always-increasing energy demand, along with a reduction of greenhouse gas emissions, renewable energies are undoubtedly the most appealing alternative for a long-term sustainable solution for our society. Therefore, huge investments have been made during the last decades to develop commercially viable technologies to convert these renewable energies. Among these, the sun is a clean, secure energy source that can supply in just one day the whole world’s energy demand of an entire year in just one day. In consequence, photovoltaic (PV) devices based on light harvesting and its conversion into electricity have been actively investigated. As an alternative to typical solar cells based on expensive, highly pure semiconductor materials, nanostructured dye-sensitized solar cells (DSSCs) are promising photovoltaic devices because of their low-cost fabrication process and
partial transparency [1–3]. Their principle is to collect photons using a dye, also called a sensitizer, linked to a photoanode (coated with nanocrystalline TiO$_2$ semiconductor (SC)). The light-induced dye excitation is followed by a fast electron injection into the SC. A redox electrolyte subsequently regenerates the dye’s ground state. The oxidized form of the mediator is finally regenerated by reduction at the cathode, thus forming a current source for electrical circuits. Ruthenium polypyridine complexes have long been considered as lead sensitizers due to their ideal photophysical and structural properties [4–6], allowing the attainment of up to 11% conversion efficiency [7–9]. Very high electron injection quantum yields [10–12] are at the origin of the remarkable success of these sensitizers. Indeed, they exhibit long-lived metal—ligand charge-transfer (MLCT) states that allow the electron injection into the sensitized semiconductor conduction band (CB). Electron injection is also further favored by an efficient interfacial electron-hole separation and by their peculiar adsorption geometry [13]. These same characteristics minimize unwanted side-processes such as detrimental charge recombination between the photoanode and the sensitizer. However, ruthenium is a scarce, potentially toxic, and expensive metal, limiting the real-world industrial development. This has motivated the quest for valuable alternatives, and among them, earth-abundant, low-cost, and environmentally friendly d-block metals such as iron [14–16] and copper [17,18] are particularly attractive. Despite an intense MLCT absorption, however, conventional Fe(II) polypyridyl complexes are, unfortunately, characterized by an ultrafast (ca. 100 fs) deactivation of the MLCT manifold to low-lying metal-centered (MC) states, proceeding via the triplet $^3$MC and ultimately populating the quintuplet $^5$T$_2$ state [13,19–22], hence impeding electron injection into the sensitized semiconductor.

The difference in the involved excited-state relaxation pathways between Ru– and Fe–polypyridine complexes is shown in Figure 1 [23]. In the former case, the metastable MLCT states are populated with a relatively long lifetime due to the absence of competitive relaxation pathways. When they are linked to a photoanode, since the MLCT state energy is high enough, electron injection into TiO$_2$ is favored (Figure 1a). As for Fe–polypyridyl complexes, on the other hand, the MC states lie at lower energy than the MLCT ones [24]. This results in the presence of efficient photophysical channels that allow the deactivation of the optically excited $^{1,3}$MLCT states (Figure 1b). From a light-to-energy conversion point of view, this means that the useful photon energy is available during a much too short time lapse. In other words, intramolecular relaxation rates are much larger than those of functional photochemical processes.

![Energetic diagrams of Ruthenium polypyridine complexes and Iron polypyridine complexes](image)

**Figure 1.** Energetic diagrams of: (a) Ruthenium polypyridine complexes; (b) Iron polypyridine complexes.

Therefore, the main challenge for further development of Fe–polypyridine complexes applicable in photochemical energy conversion processes is to enhance the lifetime of the $^{1,3}$MLCT.
manifold by precisely tuning the MLCT and MC potential energy surfaces, and most importantly, their relative positions.

As a first approximation, one can consider that the efficiency of the MLCT to MC relaxation is strongly dependent on the field strength brought by the ligands around the iron center. Indeed, the stronger the ligand-field splitting, the more separated the pseudo $t_{2g}$ and $e_{g}$ orbitals will be and hence the stronger the MC states’ destabilization. On the other hand, the increase of the π-conjugation of the ligand, and the presence of electron-acceptor group, will generally lower the energy of the MLCT states.

Recently, as shown by Wärnmark and coworkers, N-heterocyclic carbenes (NHC) have been proven to promote notable improvements of $^3$MLCT lifetimes, thanks to strong σ-donating effects inducing the destabilization of MC states [25–28]. While no DSSCs have been built from the newly designed complexes, this work has fostered the revival of research towards the development of iron-based sensitizers for DSSCs, which has been almost asleep since the pioneering works by Ferrere using the Fe(dcbpy)$_2$(CN)$_2$ dye [14].

In a different approach, using high-spin ground state halogen-substituted Fe(terpy)$_2$ complexes, it was shown that the $^5T_2$ to $^5D$MLCT transition can lead to $\approx16$ ps excited-state lifetimes, which presents a more than 100-fold increase with respect to traditional Fe–polypyridine complexes [29].

This review is devoted to our group’s latest developments in the field of NHC-based Fe(II) photosensitizers and their use in DSSCs. We focus on the synthesis of both homo- and heteroleptic complexes, discuss their photophysical, electrochemical, and computational characterization, and finally, we describe the performance of these complexes as photosensitizers in DSSCs.

2. Design of Sensitizers

When designing a potentially efficient iron-based sensitizer, one has to carefully take into account all the operating parameters of the DSSC. In this regard, the energetic levels of the $1,3^2$MLCT manifolds have to be adequately positioned to allow the injection of the photogenerated electron into the CB of the semiconductor (SC). As previously stated, the difference between first-row Fe(II) and second-row Ru(II) transition metals stems from the MC energy level, which is much lower for the former. Therefore, although having appropriate MLCT states, favorable electron transfer reactions into the SC typically only take place in Ru(II) complexes, since MLCT states from Fe(II) complexes undergo an ultrafast depopulation towards the MC states (Figure 1). In order to reverse the position of both MLCT and MC states in Fe(II) complexes and thus their interstate relaxation processes, special attention has to be paid to the electronic environment of the metal center through the modification of the metal–ligand interaction. To reach this challenging goal, the introduction of strong-field ligands is a powerful tool to raise the energy of the MC levels.

However, the design of a sensitizer for a DSSC must also take into account the global electron transfer process from the sensitizer’s excited state, and thus, a π-accepting site (to assist the extraction of the excited electron density towards the semiconductor) also has to be introduced into the coordination sphere. Moreover, the creation of π-deficiency is also expected to contribute to lowering the MLCT level. Thus, pyridyl-NHC ligands appear to be ideal candidates by fulfilling the requirements of π-acceptor (pyridine) and σ-donor (NHC) features. Importantly, the obtained iron complex must absorb a wide portion of the solar spectrum, especially in the visible region. Furthermore, the contrasting electronic character between the donor NHC units and the acceptor pyridine rings would definitely favor the electron transfer directionality, a critical feature for the performance of the sensitizer under DSSC conditions.

In order to prevent isomeric problems, Wärnmark and coworkers initially described the synthesis of tridentate bis-imidazolium pyridine (L1) as the ligand precursor [25], and the subsequent preparation of homoleptic complex C1 (Figure 2a). The authors also prepared the terpyridine (tpy) counterpart, namely [Fe(tpy)$_3$]$^{2+}$, for comparison reasons (Figure 2a). The stronger σ-donating character of the pyridylcarbene ligand with respect to tpy was nicely evidenced by the lower oxidation potential of the
complex, which was assigned to Fe(II)/Fe(III) oxidation. Concerning the optical features, the absorption spectra of both C1 and [Fe(tpy)2]3+ is shown in Figure 2b. As can be observed, a notable difference of the MLCT band position was obtained, being 92 nm blue-shifted for C1 with respect to the one of [Fe(tpy)2]3+. Nevertheless, an exceptional increase of the excited-state lifetime of C1 was achieved. In fact, complex C1 attained a 9-ps MLCT lifetime, while that of [Fe(tpy)2]3+ was 145 fs only [30]. Despite the lower covering of the 500–800 nm window, which is where the solar spectral irradiance is maximum, this result opened up the door for this class of complexes to be employed in PV applications.

![Figure 2](image.png)

**Figure 2.** (a) Structure of the ligand precursor L1 and complexes C1 and Fe(tpy)2; (b) Comparison of the absorption spectra of C1 and Fe(tpy)2 complexes in acetonitrile.

Our group has investigated several routes to improve the 3MLCT lifetime of Fe–NHC carbenes by chemically modifying the different areas of the ligands. We also examined the ways to shift back the absorption spectra towards longer wavelengths in homoleptic complexes, mostly by a stabilization of both singlet and triplet MLCT manifolds.

### 2.1. NHC-Based Homoleptic Complexes

#### 2.1.1. Effects of Carboxylic Groups

We first decided to tune the MLCT state levels by introducing electron-withdrawing groups onto the ligands as in C2 (Scheme 1) [30]. The interest of investigating the effect of carboxylic groups was two-fold: (i) the electron-withdrawing effect was expected to stabilize the MLCT states and thus decrease the MLCT transition gap and red-shift the absorption band; (ii) carboxylic groups also allow further grafting of the sensitizer onto the semiconductor surface.

![Scheme 1](image.png)

**Scheme 1.** Synthesis of complex C2.
In Figure 3, the electronic absorption spectra of pyridine-bis-NHC Fe(II) complexes C1 and C2 are collected. For both complexes, three main bands can be identified. The intense band near 320 nm corresponds to ligand-based $\pi$-$\pi^*$ transitions. At 360–380 nm, another band is observed that seems to be little affected by the nature of the ligand. On the contrary, the lower energy bands, in the 450–600 nm range, are significantly modified in terms of absorption maxima shifts and band shape. These two bands can be assigned to MLCT transitions. Indeed, the bands at 360–380 nm can be regarded as the charge transfer to NHC moieties [$\text{d}^\text{NHC} \rightarrow \pi^*(\text{NHC})$], while the lowest-energy ones represent transfer to the central pyridine [$\text{d}^\text{Fe} \rightarrow \pi^*(\text{py})$]. As a result, the functionalization of the central pyridine nucleus with a carboxylic group in C2 promotes a dramatic red-shift of this band (vide infra), whose absorption maximum is shifted from 460 to 520 nm from C1 to C2, respectively. The cyclic voltammograms of C1 and C2 confirm these results. The decrease of the electrochemical bandgaps ($\Delta E = E_{\text{ox}} - E_{\text{red}}$) from 2.75 eV to 2.20 eV going from C1 to C2 indicates that a lower energy is required to operate the metal–ligand charge transfer in C2.

![Figure 3. Comparison of electronic absorption of spectra of complexes C1 and C2 in air-equilibrated acetonitrile solution at room temperature.](image-url)

The excited-state lifetimes of the complexes were studied by femtosecond transient absorption spectroscopy (TAS) with $\approx$80 fs time resolution and broadband white-light probing in the 300–700 nm range, in order to access the full spectra of photoinduced ground state bleach (GSB) and excited-state absorption (ESA). Details are given in ref. [28] and in its Supporting information. A typical set $\Delta A$ (differential absorbance) data is displayed for C2 in Figure 4a, which shows these spectra for different delay times from −0.2 to 64 ps. They reveal GSB as negative signals at 400 and 505 nm and positive ESA bands peaking in the near-UV (320–360 nm), the blue (420–450), and in the red part of the spectrum (>580 nm). These three ESA bands are characteristic of the $^3\text{MLCT}$ state in Fe(II) complexes [24,31]. A detailed analysis of the spectral dynamics allows us to establish the following photoinduced reaction scenario [30]:

$$
\begin{align*}
^1\text{MLCT}^+ & \underset{<\text{IRF}}{\rightarrow} \begin{cases}
^1\text{MLCT} \\
^3\text{MLCT}^+
\end{cases} \underset{\tau_1}{\rightarrow} ^3\text{MCLT} \underset{\tau_2}{\rightarrow} S_0
\end{align*}
$$

(1)

Photoexcitation leads to population of a vibrationally excited $^1\text{MLCT}^+$ manifold, which relaxes within the experimental time resolution (“$<\text{IRF}$”) into a mixture of the vibrationally relaxed $^1\text{MLCT}$ and the vibrationally excited $^3\text{MLCT}^+$ triplet [23]. The fast changes of the ESA spectrum, especially for $\lambda > 580$ nm, reflects a second, slower state of intra- and intermolecular relaxation, occurring on the timescale $\tau_1$, which is in the 80–200 fs time range, and leads into the vibrationally relaxed $^3\text{MLCT}$ state. Indeed, for delay times longer than 0.3 ps, the spectral shape of the ESA bands is time-invariant.
Their amplitude decays with the time constant $\tau_2$, which is the $^3$MLCT lifetime (Figure 4b). For C1, we confirm the $^3$MLCT lifetime of $9.0 \pm 1.0$ ps [24] and find $\tau_2 = 16.5 \pm 1.0$ ps [28]. The two COOH groups on the central pyridine moiety almost double the excited-state lifetime.

There is no indication for the occurrence of the quintuplet $^5T_2$ state, since the latter is characterized by a 20 nm narrow ESA at 355 nm for Fe(II)–polypyridine complexes [28]. In a recent work on [Fe(2,2′-bipyridine)$_2$(CN)$_2$] complexes, the relaxation of $^3$MLCT into metal-centered triplet states $^3$MC with several-picosecond lifetimes was revealed [32,33]. This should lead, as for the case of the $^3T_2$ state in Fe(2,2′-bipyridine)$_3$, to a GSB recovery retarded with respect to the $^3$MLCT ESA decay. As we do not find any indication for this—both kinetics perfectly overlap for C1 and C2 (Figure 4b)—we conclude that a possible population of $^3$MC has to have an order-of-magnitude shorter lifetime than for $^3$MLCT, and therefore a negligible concentration that goes unnoticed in the TAS data.

The photophysical properties of C1 and C2 were also theoretically investigated using density functional theory (DFT) and the time-dependent (TD)-DFT techniques, employing the Hamprecht-Cohen-Tozer-Handy (HCTH) exchange and correlation functional [30]. The simulated absorption spectra are in good agreement with the experimental recordings. The $^3$MLCT states were further characterized at the Franck–Condon region by computing the natural transition orbitals (NTOs, see Figure 5), which provide excited-state descriptions based on only one (or at maximum, two) couples of orbitals [30], confirming the MLCT nature of the lowest-energy optical bands at visible wavelengths as being recorded experimentally (Figure 3). The excited-state decay of the triplet states was studied based on the relative energy of the $^3$MLCT and $^3$MC states at their corresponding equilibrium geometry and the energy difference ($\Delta E$) between these two structures. For C1, the $^3$MC minimum lies 0.73 eV below the $^3$MLCT equilibrium geometry, whereas both triplet states are almost degenerate ($\Delta E = 0.08$ eV) in the case of C2 (see Table 1). These significant differences actually support the faster decay of the $^3$MLCT state of C1 ($\tau_2 = 9.5 \pm 1.0$ ps) as compared to C2 ($\tau_2 \sim 16.5 \pm 1.0$ ps). The differences found in the relative MLCT and MC energies can be ascribed to the effect of the COOH group, only present in C2, in the electronic structure of the $^3$MLCT states. Thus, the electron-withdrawing character of this substituent is expected to stabilize the electron density localized over the aromatic rings, in global agreement with the observed and calculated red-shift of the MLCT absorption band. This explanation is consistent with the negligible impact of the COOH...
substitution in the 3MC energies, since the excitation in the latter states is entirely localized in the metal atom. As a result, the 3MLCT state has a lower energy in C2 as compared to C1, leading to quasi-degenerated 3MLCT and 3MC equilibrium geometries in the former complex, and hence to the important increase of excited-state lifetime.

![Figure 5](image)

**Figure 5.** Natural transition orbitals (NTOs) isodensity surface of C2 describing the 1MLCT states at the Franck–Condon region. λ is the absorption wavelength, f stands for oscillator strength, s*s refers to the weight of the NTO couples, and oNTO and vNTO stand for “occupied” and “virtual” NTO, respectively. Reprinted with permission from Ref. [30] (Copyright © 2015 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim).

**Table 1.** Energy difference (ΔE) between the 3MLCT and the 3MC states at their corresponding equilibrium structure, calculated as ΔE = E(3MLCT) − E(3MC) for the C1–C4 compounds computed with the TD-DFT theoretical method. The 3MLCT lifetimes recorded experimentally are also shown.

<table>
<thead>
<tr>
<th>Complex</th>
<th>ΔE (eV)</th>
<th>3MLCT Lifetime (ps)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>0.78</td>
<td>9.5 ± 1.0</td>
</tr>
<tr>
<td>C2</td>
<td>0.08</td>
<td>16.5 ± 1.0</td>
</tr>
<tr>
<td>C3</td>
<td>-</td>
<td>16.4 ± 0.4</td>
</tr>
<tr>
<td>C4</td>
<td>−0.12</td>
<td>26 ± 1.2</td>
</tr>
</tbody>
</table>

1 Ref. [30]; 2 Ref. [34].

This initial promising result prompted us to modify the NHC coordination site. In order to introduce additional π-accepting character on the NHC side and thus provide additional stabilization of the MLCT states via improved conjugation with the pyridines, the imidazolylidene (Im) moieties were substituted by benzimidazolylidene ones (BIm) [34]. The resulting complexes C3 and C4 were prepared following Scheme 2.

As can be seen in Figure 6a, C3 and C4 exhibited the same absorption features as C1 and C2. When comparing both sets of complexes, it is interesting to note that the bands in the 360–390 nm range related to the Fe–carbene MLCT [dπ(Fe) → τ(NHC)] transition [30] are strongly affected in energy when switching from imidazolylidene (Im) to benzimidazolylidene (BIm). Indeed, the absorption maxima for complexes C3 and C4 are blue-shifted by 20 nm compared with those of C1 and C2. This blue-shift is clearly in agreement with the electron-accepting effect of the phenyl ring in BIm, that must stabilize the 3MLCT state has a lower energy in C2 as compared to C1, leading to quasi-degenerated 3MLCT and 3MC equilibrium geometries in the former complex, and hence to the important increase of excited-state lifetime.
(0.85 V) complexes to C3 (1.04 V) and C4 (1.13 V) [34]. The more-stabilized metal orbitals in C3 and C4 concomitantly affected the energy of the Fe–pyridine MLCT [dπ(Fe) → π*(py)] bands, being also blue-shifted with respect to those of C1 and C2.

Cyclic voltammetry also shows the effects of the phenyl groups on the reduction potential, and thus the MLCT states. This substitution stabilizes the MLCT by 0.15 eV (C3 vs. C1) and 0.12 eV for C4 vs. C2, probably due to a better conjugation between the pyridine and the NHC [34]. The so-reduced 3MLCT–3MC energy gap is held responsible for the increased lifetime of C3.

Finally, as already observed for the Im-based complex C2 [34], the functionalization of the central pyridine ring with the carboxylic groups promotes a notable red-shift of the band at lowest energy in BIm-based complexes moving from 440 to 500 nm; i.e., from C3 to C4, respectively. Analogously, such a red-shift is due to the stabilization of the MLCT states by the electron-withdrawing effect of the carboxylic group on the ligand-conjugated π system, as shown by downshifts of ≈0.6 eV of the reduction potential [34].

The BIm leads to a significant lengthening of the 3MLCT lifetime (Figure 6b). We find \( \tau_2 = 16.4 \pm 0.5 \) and 26.0 ± 0.5 ps for C3 and C4, respectively. Again, addition of the two COOH groups leads to a significant enhancement of the 3MLCT lifetime [30].

The excited-state properties of C4 were further analyzed by means of DFT and TD-DFT [34]. Inspection of the NTOs computed at the Franck–Condon region confirmed the MLCT nature of the lowest-energy band peaking in the visible region (see Figure 7). The electronic excitation mainly occurs from the iron atom to the two pyridyl moieties, and to some extent, also involving the COOH substituents. This latter finding suggests a favorable charge injection into a semiconductor like TiO2. The decay of the triplet states of C4 was further assessed computing the 3MLCT and 3MC equilibrium

Figure 6. (a) Absorption spectra of complexes C1–C4 in air-equilibrated acetonitrile solution at room temperature. (b) Right: GSB- and ESA-normalized kinetic traces of the BIm-containing complexes C3 and C4 in acetonitrile.

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structures and comparing their relative energies (see Table 1). In contrast to C2, in which the 3MC lies 0.08 eV below the 3MLCT minimum, in C4, the energy order is inverted and the 3MC equilibrium structure lies 0.12 eV above the 3MLCT minimum, therefore supporting the observed longer 3MLCT lifetimes up to \(-26 \pm 1\) ps. C4 should hence be considered as the first example of a synthetically available iron complex presenting a 3MLCT/3MC inversion.

<table>
<thead>
<tr>
<th>(\lambda) (nm) (state number)</th>
<th>(f)</th>
<th>s*</th>
<th>oNTO</th>
<th>vNTO</th>
</tr>
</thead>
<tbody>
<tr>
<td>532 (#5)</td>
<td>0.3</td>
<td>0.570</td>
<td>0.413</td>
<td></td>
</tr>
<tr>
<td>433 (#9)</td>
<td>0.09</td>
<td>0.818</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Figure 7.** NTO isodensity surface of C4 describing the 1MLCT states at the Franck–Condon region. \(\lambda\) is the absorption wavelength, \(f\) stands for oscillator strength, \(s^*\) refers to the weight of the NTO couples, and oNTO and vNTO stand for “occupied” and “virtual” NTO, respectively. Reproduced from Ref. [34] with permission from the PCCP Owner Societies.

The effect of the number of carboxylic groups on photophysical properties was also examined. To investigate this, complex C5 [35] was prepared bearing the COOH group on only one ligand (see Scheme 3).

**Scheme 3.** Preparation of monocarboxylic complex C5.

In contrast to C1 and C2, absorption spectra of complex C5 exhibit one additional band in the visible region, that is, at 440 nm, that has no correspondence with those of the former (Figure 8a). This band can be nicely assigned to the charge-transfer transition between the metal and the pyridine ring without the carboxylic group. The lowest-energy band at 510 nm is attributed to a MLCT transition...
We find an intermediate lifetime for \( C_5 \) wavelength is adjusted to be slightly longer than the absorption maximum for all complexes. As for (a) lowering of the MLCT energies due to electron-accepting groups (\( 1 \)). The absorption spectrum indicates a 45–50 meV upshift of the iron(II) chloride under basic conditions (see Scheme 4).

The excited-state lifetimes of \( C_1, C_2, \) and \( C_5 \) in acetonitrile are reported in Figure 8b. The excitation wavelength is adjusted to be slightly longer than the absorption maximum for all complexes. As for the other complexes, \( C_5 \) follows the sequential excited-state relaxation scenario (see Equation (1)). We find an intermediate lifetime for \( C_5 \), which is \( \tau_2 = 14.0 \pm 1.0 \) ps, i.e., slightly shorter than that of \( C_2 \). Excitation is selective in the low-energy transition related to the COOH-substituted NHC ligand. The absorption spectrum indicates a 45–50 meV upshift of the \( ^1 \)MLCT transition in \( C_5 \) with respect to \( C_2 \), and the \( ^3 \)MLCT level can be expected to follow the same trend, so that the energy difference \( \Delta E \) increases. \( C_5 \) thus follows the same trend as the other complexes: \( \tau_2 \) shortens for an increasing value of \( \Delta E \) (Table 1).

Taken together, complexes \( C_1 \) to \( C_5 \) underscore the action of two effects induced by ligand design: (a) lowering of the MLCT energies due to electron-accepting groups (\( C_2, C_4, \) and \( C_5 \)) and (b) an additional downshift of the MLCT states, most probably due to an increased \( \pi \)-back-donation (\( C_3 \) vs. \( C_1 \)).

2.1.2. Effects of Alkyl Substituents

Wärnmark et al. [25] reported that the introduction of a bulky t-butyl group on the carbenic part enhances the Fe–C(NHC) bond length in the ground state reducing the \( \sigma \)-donor strength, and thus leading to an SCO-type behavior with a 0.3 ps lifetime for the \( ^3 \)MLCT. From the excited-state relaxation perspective, it is well known that the elongation of the Fe–ligand distance is a critical step in the spin cross-over process, and consequently, also for the MLCT/MC deactivation [27]. This is in line with the simple interpretation of the iron-based MC states as involving the population of the \( \epsilon_g \) (i.e., antibonding orbitals). In this context, the modification of the alkyl substituents on the carbene moiety may be a way to counteract the deactivation processes, and hence, increase MLCT lifetimes. We have examined the effect of the chain length of the alkyl substituent by switching from methyl to a linear hexyl chain. Complexes \( C_8 \) and \( C_9 \) have been prepared by reaction of the appropriate imidazolium salts with iron(II) chloride under basic conditions (see Scheme 4).
As expected, the absorption spectra of C1 and C8 are almost identical (Figure 9), both for the pyridine- and the imidazole-related transitions. Indeed, the alkyl side chains are not electronically conjugated with either of the two moieties. Comparison of C3 and C9, however, reveals a splitting of the pyridine transition, by the appearance of a shoulder in the spectrum of C9 at 410 nm and broadening of the low-energy tail. This indicates some interaction of the hexyl chain with the pyridine moiety. In any case, the 3MLCT lifetimes were determined as for the other complexes by excitation of the pyridine- and the imidazole-related transitions. Indeed, the alkyl side chains are not electronically conjugated with either of the two moieties. Comparison of C3 and C9, however, reveals a splitting of the pyridine transition, by the appearance of a shoulder in the spectrum of C9 at 410 nm and broadening of the low-energy tail. This indicates some interaction of the hexyl chain with the pyridine moiety. In any case, the 3MLCT lifetimes were determined as for the other complexes by excitation of the lowest-energy Fe–pyridine transition. Figure 9b shows the effect of the side-chain substitution for C8, which displays a longer \( \tau_2 \) than C1 (12.0 ± 1.0 ps). A similar 20% increase of the 3MLCT lifetime is observed for C9, \( \tau_2 = 19.5 \pm 1.0 \text{ ps} \), in comparison with C3. The trend goes in the right direction, but it is unclear whether these relatively small improvements are due to the postulated increase of the Fe–ligand bond rigidity or shielding effects against solvent quenching, since we are lacking structural data for these compounds.

![Scheme 4. Synthesis of C8 and C9.](image)

**Figure 9.** Effect of replacing the methyl by a hexyl side chain on the carbene moiety (Scheme 4): (a) Normalized ground-state absorption spectra; (b) Normalized kinetic traces of the GSB and ESA of C1 and C8 in acetonitrile.
2.1.3. Effects of NHC Counts

From the previous examples, it clearly appears that the strong \( \sigma \)-donating effect brought by the NHCs, along with varying contributions of \( \pi \)-accepting effects, play a fundamental role in the modulation of the MLCT lifetimes. Hence, we decided to investigate the influence of the number of carbenic moieties on the iron complexes’ photophysical properties (unpublished results). Thus, complex C10, bearing only two carbene–Fe bonds, and not four as in the above examples, was prepared from a bipyridine–imidazolylidene (bpy–Im) ligand (Scheme 5).

The UV-vis spectrum of C10 is depicted in Figure 10a together with those of C1. Although possessing intense high-energy \( \pi-\pi^* \) bands, the transitions within the visible range are less intense for C10 than for C1. At ca. 400 nm, the band corresponding to the Fe–NHC MLCT transition can be identified, which is comparable with those of C1. However, the lowest-energy band associated with the Fe–bpy MLCT transition is considerably red-shifted up to 565 nm. This result can be explained in terms of low-lying \( \pi^* \) states of the bpy unit, together with less stabilized metal orbitals as evidenced by cyclic voltammetry. The oxidation potential associated to the redox couple Fe(II)/Fe(III) (0.84 V/SCE) is slightly more positive than for C1 bearing four NHC moieties. On the negative side, two reversible waves were observed at \(-1.35\) and \(-1.62\) V/SCE. In contrast, for C1, the first injection of an electron in the ligand is dramatically more negative (\( E_{\text{red1}} = -1.95\) V/SCE). This comparison suggests that the injected electron is more easily delocalized in the bipyridine moiety than in the central pyridine bearing the two carbene groups.

![Scheme 5. Synthesis of C10.](image)

![Figure 10. Photophysics of the two-carbene compound C10. (a) Comparison of the extinction coefficients of compounds C1 (black) and C10 (blue) in acetonitrile. (b) Transient absorption spectra under 515 nm excitation of C10 in acetonitrile. Data in the 490–530 nm region show excessive noise due to pump laser scatter and are replaced by a straight line.](image)
From the TAS experiments, it turns out that C10 is a photoinduced ultrafast spin crossover complex (SCO) since the population of $^5T_2$ occurs within $\approx 0.5$ ps. This is clearly observed in the time-resolved differential $\Delta A(t)$ spectra in acetonitrile upon excitation at 515 nm (Figure 10b), by the ultrafast sub-100-fs decay of the MLCT ESA in the 400–600 nm range and the concomitant rise of the narrow photoinduced absorption (PIA) at 330 nm characteristic of metal-centered $^5T_2$ for time delays $<0.5$ ps [25]. The energy of this PIA is almost identical to that of $^5T_2$ in Fe(bpy)$_3$, meaning that it is situated only 0.5–0.6 eV above $S_0$ [36]. We recall here that $^5T_2$ is thus too low in energy to serve for electron injection in the CB of TiO$_2$. At later times, only the latter PIA remains and the visible part of the spectrum is dominated by GSB. Both signals decay with a 100 ps lifetime, indicating electronic relaxation from $^5T_2$ back to the ground state. A detailed global fitting analysis of the data is performed, providing very good agreement with the experimental kinetic traces (Figure 11).

![Figure 11](image.png)

**Figure 11.** Comparison of the normalized kinetic traces of C1 and C10 probed at 400 nm (ESA, red) and 545 nm (GSB, blue).

It reveals a sequential electronic relaxation scenario, which can be summarized as:

$$1MLCT^+ \xrightarrow{<\text{IRF}} 3MLCT^+ \xrightarrow{120 \text{ fs}} 5T_2 \xrightarrow{0.5 \text{ ps}} 5T_2 \xrightarrow{100 \text{ ps}} S_0$$

(2)

As for the above complex, the photoexcited $^1$MLCT relaxes vibrationally and into the $^3$MLCT manifold on a non-resolved ultrafast timescale (<IRF), giving rise to the early broad ESA in the visible part of the spectrum. From there, formation of the vibrationally excited quintuplet state is observed with a 120 fs lifetime, slightly slower but in agreement with the sub-100 fs lifetimes reported for Fe(2,2'-bipyridine)$_3$ [24]. Small spectral shifts occur on a 0.5 ps timescale (not shown), which are consistent with vibrational cooling. The $^5T_2$ state then recombines into the singlet ground state relatively fast, within 100 ps.

These observations complete the TAS data reported by Bauer and coworkers for the same complex [37], since they probe the characteristic near-UV absorption of $^5T_2$. Here, the initial relaxation scenario and formation time is analyzed in more detail. The same authors studied also a heteroleptic compound with tridentate ligands, bearing three carbene bonds (compound 3 in Ref. [37]). The additional carbene bond leads to a longer 3.6 ps lifetime of the $^3$MLCT state, limited by the relaxation into an unidentified MC state, which is relatively short-lived (32 ps) [37]. This work therefore demonstrates that for tridentate ligands made of pyridine and imidazole moieties,
a minimum of four carbene bonds is needed in order to avoid the $^3$MLCT-quenching SCO process. The question remains whether an increased $\pi$-back-donating character, as implemented with the Blm ligand, or increased bite angle could sufficiently increase the $^3$MLCT lifetime in a three-carbene heteroleptic complex.

2.2. Heteroleptic Complexes

Taking into account the excellent absorption of $[\text{Fe(tpy)}]^{2+}$ in the 500–600 nm window, the mixing of tpy and py–bis(Im) ligands was firstly investigated for the possibility of combining good spectral coverage and enhanced lifetime, respectively. Different terpyridine ligands were proposed, such as pytpy, since it offered an additional electronic tuning via its protonation [38]. The ligands were prepared according to the literature (see details in ref. [38]), and the complexes were synthesized following the strategy depicted in Scheme 6.

![Scheme 6. Synthesis of complex C11 and its protonated form C11-H.](image)

The absorption spectra of homoleptic C8 and heteroleptic C11 are shown in Figure 12a. As can be observed, replacement of one py–bis(Im) ligand in C9 by a tpy ligand in C11 results in a noticeable bathochromic shift of 60 nm of the two MLCT transition bands in the visible domain, offering a wide absorption domain with nearly constant intensity in the 500–580 nm range. Furthermore, the protonation of the pyridine substituent in C11-H, and the concomitant increase of electron-accepting character, had a marked effect on the absorption spectrum (see Figure 12b). Indeed, the 560 nm band in C11 was red-shifted up to 590 nm due to protonation, extending the visible light harvesting capacity of C11-H from 450 to 650 nm. Importantly, the carbene-containing complex was found to be stable under acidic conditions, which validated our strategy to shift the absorption towards longer wavelengths upon replacing one py–bis(Im) ligand by a pytpy ligand.

Cyclic voltammetry also clearly showed the influence of the pytpy ligand. In fact, while for the homoleptic iron complex C8, the reversible Fe(II)/Fe(III) oxidation process occurred at 0.41V (vs. SCE), C11 led to an oxidation potential of 0.54 V. The protonated form C11-H exhibited a reversible Fe(II) oxidation at a slightly more positive potential ($\Delta\text{E}_{1/2} = 0.02$ V) as a result of the formation of the positive charge on the pyridine nitrogen that increased the withdrawing effect on the central metal.

From a theoretical standpoint, the NTOs of the alkyl-substituted C8, C11, and C11-H complexes were computed to understand the differences in the absorption properties observed experimentally [38]. Figure 13 shows the NTO isodensity surfaces for the optically active states of those compounds at the Franck-Condon region. The homoleptic C8 complex shows a bright $^1$MLCT state at 449 nm, in which the two oNTOs (i.e., the hole orbitals) are mainly centered on the iron atom, even though non-negligible electron density is also found delocalized over the ligand moieties. This implies that part of the electron density of the aromatic rings is also redistributed after excitation. The NTO analysis also revealed that such electronic redistribution takes place over the two ligands due to symmetry reasons. In contrast, the heteroleptic complex C11 has two bright $^1$MLCT states in the visible region (457 and 546 nm),
in agreement with the experimental data, where these transitions are observed at 505 and 560 nm. In these low-energy MLCT absorptions, the electron density is excited from the metal to the pytpy ligand, whereas the alkyl-substituted ligand does not participate in the process. The red-shift of the optical bands shown in Figure 12b and observed in the experiments due to the protonation of the pyridine ring in C11-H is also supported by the theoretical calculations. In this latter complex, three 1MLCT states are predicted in the visible region (at 439, 653, and 717 nm). The two lowest-energy states may be described via the electronic density redistribution from the iron atom to the protonated pyridine ring of the pytpy ligand. The red-shift is thus rationalized and ascribed to the electron-withdrawing effect of the additional proton, which stabilizes the electron density distributed over the π* orbital in the 1MLCT excited state.

![Figure 12](image-url)

**Figure 12.** Comparison of absorption spectra in air-equilibrated acetonitrile solution at room temperature: (a) C8 and C11; (b) C11 and C11-H. Figure 12b reprinted with permission from Ref. [38] (Copyright © 2014 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim).

From the previous section, it appears that the main absorption band of C11 in the 480 to 600 nm range does only imply MLCT transitions to the pytpy moiety and not to the NHC ligand. Transient absorption spectroscopy reveals, however, that both transitions lead to two clearly different relaxation pathways. TAS was carried out with pump pulses at 480 nm for this complex, which leads to excitation of both transitions, in particular the high-energy one at 505 nm, favoring charge transfer to the peripheral pyridine moiety (Figure 13). However, the high-energy tail of the lower energy 1MLCT transition at 560 nm is also excited. The data (Figure 14) can be best described by two independent pathways, one of them following Equation (2), i.e., ultrafast SCO, and the other one a generalized version of Equation (1), avoiding SCO. Indeed, the sub-100 fs spectra are very similar to those of C10, with the ultrafast decay of the ESA in the 400–470 nm range, indicating MLCT decay and a sharpening of the 340 nm ESA for time delays <0.2 ps, attributed to the population of the 5T2 state. Interestingly, the inset in Figure 14b shows an overdamped wave packet oscillation with a 330-fs period, probed at the 5T2 ESA wavelength. A similar relaxation-induced vibrational wave packet was reported for Fe(bpy)3 [24], but with a shorter period, which is consistent with the smaller ligands in these complexes.
<table>
<thead>
<tr>
<th>Complex</th>
<th>λ (nm)</th>
<th>oNTO</th>
<th>vNTO</th>
</tr>
</thead>
<tbody>
<tr>
<td>C8</td>
<td>449 (0.25)</td>
<td><img src="image" alt="C8 NTO isodensity surface" /></td>
<td><img src="image" alt="C8 NTO isodensity surface" /></td>
</tr>
<tr>
<td>C11</td>
<td>457 (0.15)</td>
<td><img src="image" alt="C11 NTO isodensity surface" /></td>
<td><img src="image" alt="C11 NTO isodensity surface" /></td>
</tr>
<tr>
<td>C11-H</td>
<td>546 (0.18)</td>
<td><img src="image" alt="C11-H NTO isodensity surface" /></td>
<td><img src="image" alt="C11-H NTO isodensity surface" /></td>
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<tr>
<td></td>
<td>439 (0.09)</td>
<td><img src="image" alt="C11-H NTO isodensity surface" /></td>
<td><img src="image" alt="C11-H NTO isodensity surface" /></td>
</tr>
<tr>
<td></td>
<td>653 (0.19)</td>
<td><img src="image" alt="C11-H NTO isodensity surface" /></td>
<td><img src="image" alt="C11-H NTO isodensity surface" /></td>
</tr>
</tbody>
</table>

**Figure 13.** NTO isodensity surface of C8, C11, and C11-H species describing the 1MLCT states at the Franck-Condon region. λ is the absorption wavelength, f stands for oscillator strength, and oNTO and vNTO stand for “occupied” and “virtual” NTO, respectively. Reprinted with permission from Ref. [38] (Copyright © 2014 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim, Germany).
The latter, however, is not very useful for light-harvesting, since irrespective of the nature of the excited state, 2.7 ps is probably too short a lifetime. In any case, the effect of the additional pyridine is interesting in that it leads to an SCO-avoiding pathway.

In Figure 15a, the Incident-Photon-to-Current-Efficiency (IPCE) plots of C10, however, an additional 2.7 ps decay time is observed both in the ESA spectra >600 nm and in the GSB recovery at all bleach wavelengths (Figure 14b). It is absent in the $^5T_2$ ESA at 340 nm, stressing the interpretation of two independent relaxation pathways, with the second one accessed from the 505-nm excitation not leading to SCO. Notably, both transitions at 505 and 560 nm share the same $t_{2g}$ ground state, since the GSB spectrum in the 470–600 nm range is identical to the ground-state absorption (Figure 12a), so that both the 2.7 ps and the slow 55 ps decay of $^5T_2$ are found in the GSB recovery. The electronic nature associated with the 2.7 ps lifetime state is not clear. Excitation at 580 nm reduces its amplitude slightly relative to the GSB, indicating that it is related to the excited state reached by the 505 nm transition. In addition to the >600 nm ESA, it has also some ESA in the 400–470 nm range, since the GSB is almost canceled. While $^5T_2$ does not have long-wavelength ESA [28], it has recently been shown, for [Fe(bpy)(CN)$_4$)$_2$ in different solvents, that both $^3MC$ and $^3MLCT$ states show this kind of feature [32]. In any case, the effect of the additional pyridine is interesting in that it leads to an SCO-avoiding pathway. The latter, however, is not very useful for light-harvesting, since irrespective of the nature of the excited state, 2.7 ps is probably too short a lifetime.

In summary, the introduction of the pytpy ligand is beneficial for extending the absorption bandwidth, but the presence of the NHC counterpart is not sufficient to avoid SCO. An interesting coexistence of two independent pathways is found: involving either ultrafast SCO or non-SCO MLCT decay.

The preceding section has shown that a concerted combination of TD-DFT modeling and ultrafast spectroscopy is able to identify excitation wavelength-dependent relaxation pathways. Interestingly, C11 is an example where SCO-producing and avoiding-excited-state pathways coexist, depending on the localization of the charge promoted by the selected MLCT transition. While femtosecond transient absorption spectroscopy excels in time resolution, it remains a challenge to differentiate $^3MLCT$ and $^3MC$ states from the structure of the TAS spectra.

3. Photovoltaic Properties of Iron–NHC Complexes

3.1. Homoleptic Complexes

To evaluate the photovoltaic performances of these promising new iron–NHC complexes, DSSC cells were assembled using these compounds as the sensitizers. At a first stage, the photovoltaic...
properties for the homoleptic complexes C2 and C4 were investigated, since they both bear carboxylic functionalities acting as grafting groups for the TiO2 substrate [30,35].

In Figure 15a, the Incident-Photon-to-Current-Efficiency (IPCE) plots of C2 and C4 are collected. As it can be observed, the former led to a modest, though appreciable, IPCE value (about 2.3% at 500 nm). As for C4, however, only negligible electron injection was detected. In any case, it is important to bear in mind that even if modest for practical applications, those results represent the first evidence of a photocurrent generated using iron–NHC complexes. Concerning the measured photocurrent densities, DSSC cells sensitized with C2 exhibited values three times higher than those with C4 (0.41 vs. 0.12 mA cm\(^{-2}\)), along with a 20% increment in open circuit voltage (V\(_{oc}\)). Despite their good optical and photophysical properties, both the poor performance of the C4 complex and the extremely low photocurrents measured for C2 point to a nonoptimal interfacial charge separation process, possibly associated with a fast recombination mechanism occurring in the pico- and nanosecond timescale. These suggestions suppose that the relatively high injection yield reported by Wärnmark [26] for C2 grafted on TiO2 is valid in our systems. Indeed, and as compared to the typical 50 ps time window required for complete charge injection of the Ru N719 dye [39], the excited-state lifetimes of C2 and C4 in acetonitrile (16 and 26 ps, respectively), should afford a PCE (Photon Conversion Efficiency) > 1%. In addition, as shown in Table 2, the surface dye load value for C2 is even larger than that of the N719 reference dye, which shows state-of-the-art PCE properties, allowing us to leave out any problems related to an ineffective TiO2 sensitization. Strikingly, the IPCE spectrum of C4 does not show any sensitization activity of the MLCT transitions of this complex (Figure 15a).

![Figure 15. (a) Incident-Photon-to-Current-Efficiency (IPCE) curves for DSSCs sensitized with C2 (black) and C4 (red line) using I\(_2/\)I\(_{-}\) as the mediator. (b) I–V curve and performance data of DSSCs sensitized with C2 and C4. Reproduced from Ref. [35] with permission from the PCCP Owner Societies.](image)

<table>
<thead>
<tr>
<th>Dye</th>
<th>(J_{sc}) (mA/cm(^2))</th>
<th>(V_{oc}) (mV)</th>
<th>FF</th>
<th>PCE%</th>
<th>Dye Load (µmol/cm(^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>C2</td>
<td>0.41</td>
<td>457</td>
<td>0.68</td>
<td>0.13</td>
<td>0.09</td>
</tr>
<tr>
<td>C4</td>
<td>0.12</td>
<td>368</td>
<td>0.71</td>
<td>0.03</td>
<td>0.06</td>
</tr>
<tr>
<td>N719</td>
<td>13.25</td>
<td>687</td>
<td>0.67</td>
<td>6.1</td>
<td>0.08</td>
</tr>
<tr>
<td>TiO(_2)</td>
<td>0.01</td>
<td>364</td>
<td>0.51</td>
<td>0.01</td>
<td>-</td>
</tr>
</tbody>
</table>

\(^a\) Measurements performed under AM 1.5 G irradiation (100 mW cm\(^{-2}\)); irradiated area: 0.36 cm\(^2\) on TiO\(_2\) layers (10–11 µm). Values obtained from at least three DSSCs for each dye. \(^b\) Dye load calculated from absorbance data of the sensitized electrode. Values confirmed by desorption of dyes from TiO\(_2\) under basic conditions. \(J_{sc}\): short-circuit photocurrent, FF: Fill Factor.

To gain insight into the peculiar interfacial properties of the C2- and C4-sensitized TiO2 interfaces and hence provide a fundamental understanding of the charge generation and sensitization capabilities,
we reported a quantitative analysis based on high-level DFT calculations and the modelling of the relevant interfacial electron-transfer processes [35].

The electronic structure of the C2- and C4-sensitized TiO2 interfaces can be analyzed by looking at the projected density of states (PDOS) and frontier molecular orbitals (MOs) (Figure 16), obtained by DFT calculations in an implicit acetonitrile solution with the hybrid B3LYP functional and a 6-31G* basis set. More details on the methodology adopted and the calculations performed can be found in Ref. [35]. The Fe(II)–carbene/TiO2 interfaces show the typical alignment of Ru(II)–polypyridyl/TiO2 interfaces, with the HOMOs and LUMOs lying above the semiconductor valence band (VB) and conduction band (CB), respectively [40,41]. Both dyes present HOMOs completely delocalized and significantly mixed with the TiO2 VB states (around 30–50%), as can be inferred by the tail of the TiO2 VB PDOS extending through the dye’s HOMOs PDOS. This might suggest a facile back electron recombination between the injected electrons and the oxidized chromophore. However, a closer look at the nature and alignment of the unoccupied states discloses more detailed information on the interfacial charge separation and electron injection efficiency. For both C2 and C4, the dye’s LUMO, lying at −2.63 and −2.71 eV, respectively, appears to be uncoupled with the TiO2 CB states (93% and 97% of electron density localization on the dye for C2 and C4, respectively), and more importantly, it is localized over the ligand, not anchored to the TiO2 surface (see isodensity plots in Figure 16). Contrarily, the LUMO+1 distribution is markedly broadened and strongly coupled with the CB states. The lower-energy tail of the LUMO+1 shows contributions mainly localized on the grafted moiety with strong delocalization toward the CB TiO2 states, while at higher energies, significant contributions on the non-anchored ligand still appear, particularly for C2. These results clearly indicate that even if the interfacial energy level alignment is favorable for an efficient electron injection step, the primary charge transfer associated to the lowest-energy MLCT state goes in the opposite direction with respect to the injection into the TiO2 CB. This aspect directly comes from the bidentate adsorption of the deprotonated dye onto the semiconductor surface, which removes the near degeneracy between the two lowest unoccupied orbitals localized on the two ligands. As the TiO2 electron-accepting capability is weaker than that of the opposite proton, the orbitals localized on the anchored ligand are destabilized with respect to the one bearing the free COOH group. It is important to point out, however, that the cell preparation and operational conditions might sizeably change the extent of the energy splitting between these two MLCT states. In particular, ionic additives in the electrolyte, acidic treatment of TiO2, pH, and the coexistence of different anchoring modes can restore/remove their near degeneracy. In fact, as discussed in Ref. [35], the deprotonation of the not-coordinated COOH group in C2 is sufficient to revert the LUMO/LUMO+1 topology and hence obtain again the correct charge flow toward the semiconductor.

3.2. Heteroleptic Complexes

As previously discussed on the basis of the dye/TiO2 calculations, the presence of two carboxylic groups pointing towards opposite directions on the dye was deleterious for the injection process. Consequently, one of the carboxylic groups was removed and the heteroleptic complexes C5, C6, and C7 were investigated [35]. The complexes were prepared according to Scheme 7. Apart from the presence of only one anchoring group, different donor groups, e.g., p-methoxyphenyl in C6 and p-(diphenylamino)phenyl in C7, were introduced at the imidazole moiety. The resulting donor–acceptor (COOH) molecular design would eventually (i) improve the vectorial electronic charge transfer into the SC and (ii) limit the possible recombination reactions by keeping the photogenerated hole far from the SC surface.
The relaxation scenario (1), as already discussed above for C5, inorganic interfacial charge separation properties, the photovoltaic performances of the DSSCs based on heteroleptic complexes did not overperform those from C2-sensitized ones (Table 3).

### 3.2. Heteroleptic Complexes

Importantly, it is localized over the ligand, not anchored to the TiO₂ surface (see isodensity plots in Figure 16). Contrarily, the LUMO₊₁ distribution is markedly broadened and strongly coupled with the CB states. The lower-energy tail of the LUMO₊₁ shows contributions mainly localized on the grafted moiety with strong delocalization toward the CB TiO₂ states, while at higher energies, it is important to point out, however, that the cell preparation and operational conditions might sizably change the extent of the energy splitting between these two MLCT states. In particular, ionic additives in the electrolyte, acidic treatment of TiO₂, pH, and the coexistence of different anchoring modes can restore/remove their near degeneracy. In fact, as discussed in Ref. [35], the deprotonation of the not-coordinated COOH group in (diphenylamino)phenyl in opposite direction with respect to the injection into the TiO₂ CB. This aspect directly comes from the injection step, the primary charge transfer associated to the lowest-energy MLCT state goes in the opposite direction with respect to the injection into the TiO₂ CB. This aspect directly comes from the injection step, the primary charge transfer associated to the lowest-energy MLCT state goes in the opposite direction with respect to the injection into the TiO₂ CB.

Femtosecond transient spectroscopy reveals that these four-carbene-containing complexes follow the relaxation scenario (1), as already discussed above for C5. They have very similar ³MLCT lifetimes.
(10 ps for C6 and 12 ps for C7), not above 20 ps, but still enough to expect PCE values >1%, assuming that electron injection occurs on the same timescales as for N719 [39].

The IPCE and J−V curves are presented in Figure 17.

![Figure 17. (a) IPCE curves and (b) J/V curves for DSSCs sensitized with C2, C5, C6, and C7 using I$_2$/I$^-$.](image)

From the J−V curves (Figure 17b) and PV data in Table 3, despite possessing the desired interfacial charge separation properties, the photovoltaic performances of the DSSCs based on heteroleptic complexes did not overperform those from C2-sensitized ones (Table 3).

### Table 3. Photovoltaic performances of DSSCs sensitized with heteroleptic dyes

<table>
<thead>
<tr>
<th>Dye</th>
<th>$J_{SC}$ (mA·cm$^{-2}$)</th>
<th>$V_{OC}$ (mV)</th>
<th>FF</th>
<th>PCE%</th>
<th>Dye Load ($\mu$mol·cm$^{-2}$) $^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>C2</td>
<td>0.41</td>
<td>457</td>
<td>0.68</td>
<td>0.13</td>
<td>0.09</td>
</tr>
<tr>
<td>C5</td>
<td>0.33</td>
<td>400</td>
<td>0.73</td>
<td>0.10</td>
<td>0.10</td>
</tr>
<tr>
<td>C6</td>
<td>0.36</td>
<td>440</td>
<td>0.73</td>
<td>0.11</td>
<td>0.12</td>
</tr>
<tr>
<td>C7</td>
<td>0.36</td>
<td>390</td>
<td>0.71</td>
<td>0.10</td>
<td>0.07</td>
</tr>
<tr>
<td>N719</td>
<td>13.25</td>
<td>687</td>
<td>0.67</td>
<td>6.1</td>
<td>0.08</td>
</tr>
</tbody>
</table>

$^a$ Measurements performed under AM 1.5 G irradiation (100 mW·cm$^{-2}$); irradiated area: 0.36 cm$^2$ on TiO$_2$ layers (10–11 μm). Values obtained from at least three DSSCs per dye. $^b$ Dye load calculated from absorbance data of the sensitized electrode. Values confirmed by desorption of dyes from TiO$_2$ under basic conditions.

Also, in this case, it can be worthwhile to examine the PDOS of the dye/TiO$_2$ system displayed in Figure 18. As is apparent, the desired charge-transfer nature of the lowest-energy dye level is obtained, in line with the PBE0 results reported in Ref. [42] for the same complex adsorbed on a (TiO$_2$)$_{92}$ cluster model. As shown in Table 3, the PV performances, however, are still poor, and this seems to be in contradiction with the favorable energetics and topology calculated for this system.

To propose a further rationalization of the charge separation properties of the Fe(II)−NHC@TiO$_2$ interface, we computed the rates for both the electron transfer from the dye’s donor state (LUMO or LUMO+1, in this case) into the acceptor states of the TiO$_2$ (CB) and for the back-electron transfer (recombination) from the CB states to the dye’s HOMO. Table 4 reports the calculated injection probabilities, $G(\epsilon)$, obtained at the LUMO/LUMO+1 and HOMO energy for electron injection and recombination, respectively; the corresponding VB or CB DOS; and the injection rates, $k_{\text{inj}}$. 

![Table 4. Calculated injection rates, $G(\epsilon)$, obtained at the LUMO/LUMO+1 and HOMO energy for electron injection and recombination, respectively; the corresponding VB or CB DOS; and the injection rates, $k_{\text{inj}}$.](image)
Concerning the homoleptic complexes (C2 and C4), data in Table 4 indicate, as expected, that the electronic coupling between the dye’s LUMO and the semiconductor’s CB states is extremely weak, with \( \Gamma \) values of the order of \( 10^{-4} \) eV, yielding very low injection rates of about \( 10^{12} \) s\(^{-1}\). That is, the electron injection for the homoleptic dyes is foreseen to occur in the ps timescale, coinciding with the timescale for the deactivation of the \( ^3 \)MLCT state. Moreover, a quantitative difference between C2 and C4 is also predicted, with a lower injection probability for the latter, possibly arising from the more delocalized nature of the MOs induced by the BIm groups. In line with the previous PDOS analysis (Figure 16), on the other hand, an ultrafast injection mechanism is calculated for LUMO+1, with \( \Gamma \) values of the order of \( 10^{-1} \) eV and injection rates of \( 10^{15} \) s\(^{-1}\) (fs timescale).

Turning now to the heteroleptic C5 complex, an appreciable change in the electronic structure, i.e., in the diabatic LUMO nature, is obtained, resulting in lower electronic coupling with the low-energy TiO\(_2\) CB states (where the LUMO is energetically located) and larger coupling with higher-energy states (more details can be found in Ref. [35]). As a consequence, the calculated \( \Gamma(\varepsilon_{\text{LUMO}}) \) for C5,
despite the correct interfacial charge-transfer topology, is also in the range of several ps; that is, in the same timescale as the $3\text{MLCT}$ lifetimes (14 ps).

Further understanding also comes from the hole injection analysis. Even if, as shown by the data in Table 4, in our diabatic-like scheme recombination to the oxidized dye is calculated to be extremely slow, as a consequence of the negligible energetic overlap between the dye HOMO and the TiO$_2$ CB states (ms timescale), we found for C2, C5, and C4 (even if at a lower extent) a large electronic coupling with both the VB and CB states due to the above discussed high delocalization of the HOMO boosting the electronic conjugation with the TiO$_2$ MOs [35]. It is worthwhile to stress that in the complexity of the “working” dye-sensitized electrode, where heterogeneous binding and pair interactions perturb the energetics discussed so far, the actual back-recombination processes might be faster, taking place in the nanosecond and even picosecond timescale, as reported in Ref. [26].

4. Conclusions and Outlook

In this review, we have shown the impact of different molecular designs of iron(II) complexes on their photophysical and electrochemical properties. In particular, the key strategies to enlarge their light absorption range as well as to increase their excited-state lifetimes are presented. Thus, upon a judicious chemical design to adequately tune their optoelectronic properties, NHC-based iron(II) complexes can be regarded as promising sensitizers for DSSCs, which truly turns them into valuable alternatives to rare metals such as ruthenium.

In iron(II) complexes, the sufficient destabilization of low-lying MC states in order to allow the exploitation of MLCT states, such as for electron injection into the SC band, is highly challenging. In this regard, NHC coordinating units are of fundamental importance. By modifying the ligands around the iron center as stated, we succeeded in improving the excited-state lifetime by up to 26 ps, especially using the benzimidazolylidene moiety as the carbenic part. It is worth noting that TD-DFT computations revealed for the first time a $3\text{MLCT}/3\text{MC}$-level inversion.

The good absorption features and the remarkably long-lived MLCT excited states encouraged their use as dyes for DSSC. The first DSSC units assembled with a set of NHC-based iron complexes show that (i) they can effectively sensitize titanium dioxide and (ii) their MLCT band can be excited to produce a photocurrent upon solar irradiation. However, despite a priori sufficiently long MLCT lifetimes, the efficiency of the cells was found to be low.

By means of interfacial DFT calculations, we demonstrate the presence of inefficient electron coupling between the sensitizers and the SC orbitals, thus considerably decreasing the injection rate that falls in the same regime as the MLCT/MC deactivation process.

Work is in progress to design new complexes combining further-improved excited-state lifetimes and absorbances, along with appropriate electronic properties for better cell performances. Here, in addition to tuning of the ground-state properties (ligand field splitting), design strategies that influence the nonadiabatic excited-state relaxation kinetics [27] remain unexplored for the moment.

From a spectroscopic point of view, additional methods are needed to determine the location of the electronic states along the excited-state relaxation pathway. We recently showed, by ultrafast fluorescence spectroscopy, that the $3\text{MLCT}$ in Fe(II)–NHC complexes has a relatively small red-shift of (<0.4 eV) with respect to $1\text{MLCT}$ [43]. Besides, it is highly desirable to reveal metal-centered d–d transitions and thus to obtain direct evidence for the impact of chemical design on the induced ligand field splitting.

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