The short device lifetimes of blue PhOLEDs: insights into the photostability of blue Ir(III) complexes

Denis Jacquemin and Daniel Escudero

Using Ir(III) complexes as dopants in phosphorescent organic light-emitting diodes (PhOLEDs) is the most successful strategy to attain long-lasting and highly-performant electroluminescent devices. Whilst highly efficient blue PhOLEDs are accessible, their limited operational lifetimes still restrain their practical use in lighting and displays. These short lifetimes are directly related to the low intrinsic photostability of blue Ir(III) complexes. This work uses first principles to unveil the mechanisms of degradation of blue phosphors arising from energetically hot excited states (≈6.0 eV), and to propose a strategy for improving the stability of blue phosphors.

Introduction

Phosphorescent-based OLEDs (PhOLEDs), the so-called second generation of OLEDs, are the most widespread electroluminescent devices since they can attain low power consumption and record external quantum efficiencies. Electroluminescence in PhOLEDs takes place from the triplet excited states of the organometallic complexes, i.e., typically Ir(III) and Pt(II) complexes, which enables the conversion of nearly 100% of the excitons created into photons. Besides high efficiencies, it is essential that the devices possess long-lived operational stability. In contrast to red or green PhOLEDs, the fabrication of a long-lasting blue PhOLED has not been achieved yet, which has consequently limited the large-scale commercialization of PhOLEDs. For both lighting and display applications, operational lifetimes of $T_{95} > 10000$ h are mandatory (where $T_X$ is the time under constant current operation in which the luminance decreases to $X\%$ of its initial value). To illustrate the size of the gap to be bridged, state-of-the-art blue and greenish-blue PhOLEDs are characterized by $T_{80}$ of <10 h and <160 h, respectively, values clearly insufficient for practical applications. Degradation in PhOLEDs manifests itself through the formation of dark spots, the occurrence of a catastrophic failure and/or the decrease of the overall luminance during continuous operation. Whilst the dark spots and the catastrophic failure can be minimized by an optimal device fabrication, the long-term decrease of the overall luminance during prolonged operation is the major obstacle for the commercial exploitation of PhOLEDs, especially for the blue ones. Intrinsic chemical degradation of both host and dopant molecules is responsible of harming device’s operational lifetimes.

The elucidation of the mechanisms of degradation products as well as of the related mechanisms poses, however, significant experimental challenges due to (i) the limited amount of defects on an aged PhOLED stack, (ii) the difficult isolation of the degradation products from the organic semiconductor layers, and (iii) the intricate competition between multiple non-desired reactions (including exciton and charge carrier induced decomposition), that leads to a broad variety of chemical defects. Despite these inconveniences, multiple analytical techniques that provide insights into the chemical degradation of thin organic semiconductor films are nowadays available. One of the most powerful methods is the matrix assisted laser desorption/ionization time-of-flight mass spectroscopy (MALDI-TOF-MS) method, which allows trace analysis of organic layers or even entire devices. This technique enabled the identification of relevant degradation products for host materials as well as for phosphorescent dopants. This knowledge has paved the way to the development of host-dopant matching design principles and has therefore significantly contributed to the improvement of the operational stability of PhOLEDs materials. Nevertheless, in the majority of the cases, the actual degradation mechanisms remain unknown and thus, quantum chemical investigations can shed light into the structural features, nature of the electronically excited states and excited-state reactivity of host and dopant materials. However, to date, these investigations are scarce and limited to very specific cases. The present work unveils, from first principles, relevant mechanisms of degradation of state-of-the-art blue Ir(III) phosphors due to photodechelation reactions.
Beyond disclosing these mechanisms and providing paths for improving the design of Ir-based phosphors, this contribution illustrates how quantum chemical modeling may largely impact the field of blue Ir(III) dopants.

The degradation pathways of Ir(III) phosphors upon PhOLED operation root on parasitic annihilation processes between excited states (i.e., exciton-polaron and/or exciton-exciton) that lead to the formation of hot excited states (that is, highly energetic excited states at \( \equiv 6.0 \) eV above the ground-state). Among all possible mechanisms, triplet–triplet annihilation (TTA) is of uttermost relevance for PhOLEDs. TTA processes in PhOLEDs are thought to be mainly due to dopant:dopant aggregation in the emissive layer (EML) and occur via a diffusion-based Dexter transfer mechanism. In a TTA process (see Scheme 1a), according to spin statistics, one can generate 1/9 of hot singlet excited states \( (S_n) \) per 3/9 and 5/9 of hot triplet \( (T_n) \) and quintet \( (Q_n) \) states, respectively. The quintet excited states are, according to our calculations, energetically not reachable for Ir(III) complexes (exemplarily, the first quintet state for the prototypical compound 1 (see Chart 1) is located 6.11 eV above the S\(_0\) energy at the Franck–Condon point) and, therefore, they do not contribute to the global deactivation dynamics. Hence, due to the nonradiative quenching of at least half of the initial \( T_1 \) states, TTA leads to an effective quenching of photoluminescence in PhOLEDs (also known as roll-off effect)\(^7\) and to the formation of hot \( T_n \) states, since the 1/9 of \( S_n \) also lead to \( T_n \) states after efficient intersystem crossings (ISC) processes. Indeed, ISC processes occur typically with near-infrared singlet yield for Ir(III) complexes.\(^8\) Having in mind that the \( T_1 \) energy for a typical deep-blue emitter is ca. 3 eV, hot \( T_n \) states of up to 6 eV can be generated through TTA (the energetic requirement for TTA is \( E(T_n) \leq 2E(T_1) \)). The Jablonski diagram including the relevant excited states and processes is presented in Scheme 1b. While the vast majority of hot states will rapidly decay back nonradiatively\(^9\) (i.e., \( T_n \rightarrow T_1 \), process 3 in Scheme 1b), those which are in a reactive excited vibrational state may lead to dissociative states \( (D_m \), process 4 in Scheme 1b) and eventually yield the degradation of the emissive complexes. Typical bond dissociation energies of weak bonds in phosphors (see the discussion below) are well below the energy of the hot states but they are higher in energy than \( T_1 \). Therefore, it is clear that TTA is the major source of chemical degradation in PhOLEDs. We underline that often unexpected degradation events happen, since it is common that excited state reactivity greatly differs from its ground state counterpart and additionally, some groups in specific environments are more susceptible to e.g., nucleophilic substitution reactions. One typical example is the lability of the C–F bonds of the \((4,6\text{-difluorophenyl})\text{pyridinato} \) units upon PhOLED operation,\(^9\) which can not be explained in terms of the ground-state thermodynamics of the \( \text{C(sp}^3\text{)} \)-fluorine bond, which is one of the strongest known single bonds. In short, if one aims to minimize the generation of degradation products in PhOLEDs, two possible ways are available: reducing TTA or controlling the excited state reactivity by phosphor design strategies. Efficient ways to reduce TTA processes include: (i) controlling the dopant:dopant aggregation at the EML, and (ii) attaining shorter triplet excited state lifetimes. Both approaches are challenging to implement in PhOLEDs, since obtaining triplet lifetimes of Ir(III) complexes shorter than a few \( \mu \)s is difficult and controlling aggregation on the vacuum deposited EML is hardly achievable.

In this contribution we focus on disentangling the excited state reactivity for blue Ir(III) phosphors and we explain, for the first time, why blue Ir(III) dopants are more prone to degradation than the green and red ones. We next provide some phosphor design principles to limit the formation of degradation products.

### Results and discussion

Chart 1 gathers the homoleptic Ir(III) complexes studied here, including the green \( \text{fac-Ir} \text{(ppy)}_3 \) (where ppy = phenylpyridine) and \( \text{fac-Ir} \text{(mpy)}_3 \) (where mpy = methylphenylpyridine) complexes (1 and 2, respectively) and the blue \( \text{fac-Ir} \text{(ppz)}_3 \) (where ppz = phenylpyrazole) and \( \text{fac-Ir} \text{(pmb)}_3 \) (where pmb = 1-
phenyl-3-methylbenzimidazolyl) complexes (3 and 4, respectively). These complexes were selected due to the availability of experimental degradation studies and to unravel the intrinsic differences in the excited-state reactivities of blue and non-blue complexes. Experimentally, the blue complex 3 undergoes dissociation of a ppz ligand and secondly it coordinates a host molecule when the device operates at moderate to high current levels. As a result, an operational lifetime of $T_\text{95} < 1$ h is obtained for this device. Oppositely, the green complex 1, under the same conditions, is much more photostable and its electroluminescence performances remain unchanged. Despite the intrinsic photostability of 1, another group has reported some minor trace amounts of degradation products that involve the dissociation of the ppy cyclometallating ligand. For complex 2, the photodegradation products of aged devices have not been experimentally characterized, but when 2 is photo-degraded in inert solution it undergoes enhanced degradation processes as compared with 1,22 likely involving ligand dissociation as well. Finally, the use of N-heterocyclic carbene (NHC) ligands is beneficial not only to attain highly efficient deep-blue Ir(m) emitters but also to potentially increase their intrinsic photostabilities. Blue PhOLED devices using tris(bidentate) NHC Ir(m) complexes, such as e.g., complex 4, are known to present increased device lifetimes compared to other blue Ir(m) emitters (e.g., complex 3).23 Although degradation is still observed in these devices, the actual degradation products have not been indentified to date. The first important criterion that should be met to avoid ligand dissociation is that the reaction from the hot excited states should be thermodynamically unfeasible. In Table 1 are listed the computed Gibbs free energies for the homolytic ligand dissociation reactions of complexes 1–4 (see computational details below). For the green complex 1, a $\Delta G_{\text{react}}$ of 5.01 eV is obtained. This value is nearly isoenergetic to the energy of a green hot state (which is ca. $2E(T_1) \equiv 5.0$ eV) and therefore, ppy dissociation may occur only in exceptional occasions (as the minor traces experimentally detected corroborate).22 Complex 2 possesses a smaller $\Delta G_{\text{react}}$ value than 1 (4.80 eV), and thus it is more prone to ligand dissociation, in agreement with experimental evidences. For complex 2, another possible degradation mechanism may arise from the dissociation of a methyl group (see reaction 2’ in Table 1). Although this dissociation reaction is thermodynamically more feasible and might compete to ppy dissociation, it is likely that the latter process dominates due to the lability of the ligand dissociation in the excited states of Ir(m) complexes (see below). In the case of the blue complexes, their $\Delta G_{\text{react}}$ values for the photodechelation reactions (reactions 3 and 4 in Table 1) are well below the energy of a blue hot state (which is ca. $2E(T_1) \equiv 6.0$ eV) and thus, their ligand dissociation reaction are in principle energetically feasible. In the case of 3 the $\Delta G_{\text{react}}$ value is 4.2 eV only, partly explaining the extremely fast device degradation observed experimentally.20 Additionally, for 4 we explored the possibility of the self-fragmentation reaction of the pmb ligand through the $N$-C bond rupture (see Chart 1). This reaction is also thermodynamically feasible, see 4’ in Table 1 ($\Delta G_{\text{react}} = 4.6$ eV).

While the thermodynamic criterion, i.e., $\Delta G_{\text{react}} > 2E(T_1)$, should always be met to completely avoid ligand dissociation, the excited state kinetic does play an important role in modulating the degree of degradation for complexes that do not meet such criterion. As mentioned above, the vast majority of hot $T_n$ states rapidly decay back nonradiatively recovering the $T_1$ geometry. The $T_1$ state is usually the $^3\text{MLCT}$ state and displays a pseudo-octahedral coordination environment (see Scheme 2b). Therefore, it seems very unlikely that ligand dissociation (which involves the breaking of two bonds, i.e., $\text{Ir}^{-1}\text{-C}$ and $\text{Ir}^{-1}\text{-N}$) will take place at the $^3\text{MLCT}$ structure. In this regard, spectroscopic24 and computational studies25–27 confirmed the active role of metal centred triplet excited states ($^3\text{MC}$, see Scheme 2b) in the excited state dynamics of Ir(m) complexes, especially for blue emitters. Importantly, the $^3\text{MC}$ state can be thermally populated from the $^3\text{MLCT}$ state. In Scheme 2a, these pathways are represented. The pseudo-octahedral Ir(m) complexes at their $T_1$ geometry need to surpass a barrier (TS in Scheme 2a) to attain the $^3\text{MC}$ state geometry. This state usually displays a trigonal bipyramidal structure, where one iridium-heteroatom single bond is

### Table 1

| Complex | Reaction | $\Delta G_{\text{react}}$ (eV) |
|---------|----------|-------------------------------|
| 1       | 1: Ir(ppy)$_3$ $\rightarrow$ [Ir(ppy)$_2$][ppy] $+$ [ppy] | 5.01 (5.0) |
| 2       | 2: Ir(mppy)$_3$ $\rightarrow$ [Ir(mppy)$_2$][mppy] $+$ [mppy] | 4.80 (5.0) |
| 2’      | 2’: Ir(mppy)$_3$ $\rightarrow$ [Ir(mppy)$_2$][ppy] $+$ [-CH$_3$] | 3.93 (5.0) |
| 3       | 3: Ir(ppz)$_3$ $\rightarrow$ [Ir(ppz)$_2$] $+$ [ppz] | 4.18 (6.0) |
| 4       | 4: Ir(bmp)$_3$ $\rightarrow$ [Ir(bmp)$_2$][bmp] $+$ [bmp] | 5.77 (6.0) |
| 4’      | 4’: Ir(bmp)$_3$ $\rightarrow$ [Ir(bmp)$_2$][C$_6$H$_5$] (C$_6$NH$_7$)$_2$ | 4.60 (6.0) |
| 5       | 5: $\text{[Ir(mimf)(pzpyphF)]} \rightarrow$ [Ir(mimf)]$^-$ $+$ [pzpyphF]$^+$ | 10.00 (6.0) |
| 5’      | 5’: $\text{[Ir(mimf)(pzpyphF)]} \rightarrow$ [Ir(pzpyphF)$^-$] $+$ [mimf] | 9.06 (6.0) |

$^a$ Energetic threshold for a blue or a green hot state.
broken (see Scheme 2b). After population of the $^3$MC well, two processes might occur: (i) reversible return to the $^3$MLCT well; or (ii) irreversible radiationless recovery of the ground state ($^1$GS) geometry through the $^1$GS/$^3$MC minimum energy crossing point (MECP).\(^{28}\) The MECP geometry usually exhibits a further distorted trigonal bipyramid arrangement. This kinetic scenario can be summarized as:\(^{29}\)

$$\text{3MLCT} \rightarrow \text{3MC} \rightarrow \text{1GS}. \quad (1)$$

At room temperature (RT), $^3$MLCT $\rightarrow$ $^3$MC equilibration occurs for many Ir(III) complexes.\(^{27}\) This equilibrium is controlled by the magnitude of the activation barriers ($E_a$, $E_b$ and $E_c$ in Scheme 2a). A straightforward approach to prevent the thermal population of the $^3$MC is to maximize the $E_a$ value. In Table 2 are tabulated the computed $E_a$, $E_b$ and $E_c$ values for complexes 1–4 (see computational details). In the blue complex 3, the $^3$MC state is adiabatically located lower in energy than the emissive $^3$MLCT state. Additionally it is accessed in a barrierless manner, since no TS is found along the $^3$MLCT $\rightarrow$ $^3$MC reaction coordinate. Therefore, at RT, this nonradiative pathway is the most prominent deactivation channel for 3, and thus it leads experimentally to a complete quench of phosphorescence. Having in mind these kinetic considerations, and under steady-state conditions, TTA will predominantly take place in 3 between two $^3$MC states. Since, at the $^3$MC structure, one iridium–nitrogen single bond is already broken, the probability of reaching in the TTA process a reactive vibrational excited state is the ligand-to-metal charge transfer (3LMCT) state shown in Fig. 1b, which involves an excitation of the Ir–ppz orbital. At the $^3$MC optimized geometry this state corresponds to $T_{88}$ and its excitation energy is 5.84 eV, and thus it can be populated after TTA. The scan along the Ir–C bond cleavage reaction coordinate starting from the $^3$MC structure (see computational details below). In Fig. 1a, this scan is exemplarily presented for complex 3. One possible dissociative excited state is the ligand-to-metal charge transfer ($^1$LMCT) state shown in Fig. 1b, which involves an excitation from a $\sigma_{ppz}$ + $\pi_{Ir-ppz}$ orbital to a $d + \sigma_{Ir-ppz}$ orbital. At the $^3$MC optimized geometry this state corresponds to $T_{88}$ and its excitation energy is 5.84 eV, and thus it can be populated after TTA. The scan along the Ir–C bond cleavage reaction coordinate confirms the dissociative nature of this $^1$LMCT state, that potentially leads to ppz ligand dissociation upon relaxation on this PES. Note that in the real device situation, the “cage effect”

\[\text{Table 2: PWPB95-D3BJ activation barriers (eV) for the temperature-dependent nonradiative channels (see Scheme 1a) and 3MLCT – 3MC adiabatic energy differences for 1–4} \]

| Complex | $E_a$ | $E_b$ | $E_c$ | $\Delta E(3MC - 3MLCT)$ |
|---------|-------|-------|-------|--------------------------|
| 1       | 0.352 | 0.025 | 0.038 | 0.327                    |
| 2       | 0.365 | 0.287 | 0.016 | 0.078                    |
| 3       | 0.000 | 0.153 | 0.006 | -0.153                   |
| 4       | 0.638 | 0.562 | 0.079 | 0.076                    |

\[\text{Fig. 1: (a) TD-DFT scan of the } T_{88} \text{ state along the Ir–C bond cleavage reaction coordinate of complex 3. (b) Kohn–Sham Frontier orbitals involved in the dissociative } ^1\text{LMCT state.}\]
exerted by the host surrounding material will favour the ligand recombination to a certain extent.

Let us now propose blue phosphor design principles to avoid the formation of degradation products using the thermodynamic and kinetic criteria discussed above. The most straightforward way to prevent ligand dissociation is to attain $\Delta G_{\text{react}}$ values above the energetic threshold of a blue hot state ($i.e., >6.0$ eV). To this end, the use of bis(tridentate) Ir(III) complexes, in lieu of the more popular tris(bidentate) ones, is suited, since in the former complexes ligand dissociation involves the cleavage of three Ir-heteroatom bonds. In addition, as demonstrated herein, the use of NHC ligands in a tris(bidentate) fashion is a successful strategy to decrease the kinetic liability of the dissociative processes, as their stronger Ir-C carbon bonds lead to large $E_a$ values, consequently preventing the formation of their 3MC states. However, in view of the computed $\Delta G_{\text{react}}$ values for 4, it seems unlikely that the thermodynamic criterion can be met for any tris(bidentate) NHC Ir(III) complex. Nevertheless, we predict that there is still room to attain larger $E_a$ and $\Delta G_{\text{react}}$ values by ligand design strategies. As an optimal way to avoid degradation, we propose to combine bis(tridentate) architectures with NHC ligands. Indeed, highly efficient deep-blue neutral Ir(III) complexes have recently reported using this strategy. One representative example is the heteroleptic Ir(mimf)[pzpyph}] complex 5 shown in Chart 2 (where mimf is a pincer dicarbene ligand and pzpyph stands for a tridentate 6-pyrazolyl-2-phenylpyridine ligand). We computed the $\Delta G_{\text{react}}$ values for the two possible ligand dissociation reactions of 5 (see reactions 5 and 5’ in Table 1). Both dissociations are highly endergonic ($ca. 10$ and $9$ eV, respectively) and they are located significantly above the energetic threshold of a blue hot state. Therefore, one can anticipate that ligand dissociation cannot take place for complex 5 upon PhOLED operation. Although the presence of C-F bonds in complex 5 might be a problem for practical PhOLEDs applications, this general design approach should help improving complex stability. To our knowledge, this approach remains to be used in the PhOLEDs community.

Conclusions

In this manuscript we unveil from first principles relevant degradation mechanisms of state-of-the-art blue Ir(III) complexes upon PhOLED operation. Energetically hot states ($\approx 6.0$ eV) generated after annihilation processes between excited states are responsible of the intrinsic chemical deterioration of PhOLEDs materials. These parasitic processes compromise the large-scale commercialization of PhOLEDs, especially the blue ones. Through our quantum chemical investigations, we unravel one of the most critical degradation processes, i.e., the photodechelation reaction. We found that both kinetic and thermodynamic criteria do play an important role in the degradation processes. The larger tendency to form the 3MC states in the blue complexes (enhanced kinetic liability) stands as the ultimate reason for their reduced intrinsic photostability. In view of these results we propose some blue phosphor design strategies. In particular, combining bis(tridentate) architectures with NHC ligands looks a very promising method to prevent ligand dissociation reactions. Due to their extreme robustness, these complexes may also be very good candidates to be used as hot excited state managers in managed EML. Finally, whilst the photodechelation mechanism is found to be common for blue and non-blue complexes, it is certainly not the exclusive mechanisms of deterioration of blue dopants. Other possible chemical reactions triggered by hot states include radical reactions occurring at the chelate ligands, as shown for complexes 2 and 4, that may potentially lead to further reactions with the host material surrounding the complexes. Another typical example found in the literature is the degradation of the picolinate ligand, where the elimination of thermodynamically stable CO$_2$ takes place. Therefore, phosphor design strategies should always be accompanied by a careful choice of the ligands. Computationally-guided ligand design strategies are currently under investigation in our group.

Experimental section

Computational details

All calculations rely on density functional theory (DFT). The geometries of the singlet ground state ($^1$GS), the triplet $^3$MC and $^3$MLCT states as well as of the transition states (TS) were optimized for complexes 1–5 using the dispersion corrected hybrid functional B3LYP-D3 [ref. 34 and 35] in combination with the 6-31G(d,p) atomic basis set for all atoms. Relativistic effects were considered for the Ir atom by using the ECP-60-mwb pseudopotential. The Hessian was computed at the same level of theory to confirm the nature of the stationary points. The minimum energy crossing points (MECP) between the $^1$GS and the $^3$MC potential surfaces were optimized using Harvey’s algorithm, as implemented in the ORCA software; with the B3LYP-D3 functional in combination with the def2-svp basis set and the same pseudopotential for Ir. To obtain accurate relative energies for the PES, single-point calculations were performed with the ORCA software using the dispersion-corrected double-hybrid PWPB95-D3BJ functional and the def2-svp basis set (ECP-60-mwb pseudopotential for Ir). This exchange-correlation functional outperforms the majority of the available functionals for the thermodynamic properties of 5d transition metal complexes. All calculations apart from the MECP optimization and the single point PWPB95-D3BJ calculations were carried out using this strategy.
with the Gaussian09 program package. The same protocol (i.e., PWPB95-D3J/def2-svp//B3LYP-D3/6-31G(d,p)) was used to compute the $\Delta G_{\text{react}}$ values. For the Ir–3C scan, gas phase TD-B3LYP-D3 vertical triplet excitation energies were obtained for 3 starting at the $^1$MC optimal geometry. For these calculations we used the 6-31G(d,p) atomic basis set and the ECP-60-mwb pseudopotential for Ir.

**Conflicts of interest**

There are no conflicts to declare.

**Acknowledgements**

DE thanks funding from the European Union’s Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie grant agreement No. 700961. This research used resources of the CCIPL (Centre de Calcul Intensif des Pays de Loire), and of a local Troy cluster.

**Notes and references**

1. C. Adachi, M. A. Baldo, M. E. Thompson and S. R. Forrest, *J. Appl. Phys.*, 2001, **90**, 5048.
2. H. Yersin, A. F. Rausch, R. Czerwieniec, T. Hofbeck and T. Fischer, *Coord. Chem. Rev.*, 2001, **255**, 2622–2652.
3. M. Hack, J. J. Brown, M. S. Weaver and M. Premutico, *Lifetime OLED display, U.S.*, Patent 9257665, B2, 2016.
4. J. Zhuang, W. Li, W. Su, Y. Liu, Q. Shen, L. Liao and M. Zhou, *Org. Electron.*, 2013, **14**, 2596–2601.
5. K. P. Klubek, S.-C. Dong, L.-S. Liao, C. W. Tang and L. Rothberg, *Org. Electron.*, 2014, **15**, 3127–3136.
6. J. Zhuang, W. Li, W. Wu, M. Song, W. Su, M. Zhou and Z. Cui, *New J. Chem.*, 2015, **39**, 246–253.
7. Y. L. Kang and Y. Y. Lee, *Org. Electron.*, 2016, **32**, 109–114.
8. S. Scholz, D. Kondakov, B. Lüssem and K. Leo, *Chem. Rev.*, 2015, **115**, 8449–8503.
9. S. Schmidbauer, A. Hohenleutner and B. König, *Adv. Mater.*, 2013, **25**, 2114–2129.
10. D. Y. Kondakov, W. C. Lenhart and W. F. Nichols, *J. Appl. Phys.*, 2007, **101**, 024512.
11. M. Karas, D. Bachmann, U. Bahr and F. Hillenkamp, *Int. J. Mass Spectrom. Ion Processes*, 1987, **78**, 53–68.
12. S. Scholz, C. Corten, K. Walzer, D. Kuckling and K. Leo, *Org. Electron.*, 2007, **8**, 709–717.
13. R. Meerheim, S. Scholz, S. Olthof, G. Schwartz, S. Reineke, K. Walzer and K. Leo, *J. Appl. Phys.*, 2008, **104**, 014510.
14. G. Treboux, J. Mizukami, M. Yabe and S. J. Nakamura, *J. Photopolym. Sci. Technol.*, 2008, **21**, 347–348.
15. T. Setzer, C. Lennartz and A. Drew, *Dalton Trans.*, 2017, **46**, 7194–7209.
16. J. Lee, C. Jeong, T. Batagoda, C. Coburn, M. E. Thompson and S. T. Forrest, *Nat. Commun.*, 2017, **8**, 15566.
17. C. Murawski, K. Leo and M. C. Gather, *Adv. Mater.*, 2010, **25**, 6801–6827.
18. G. J. Hedley, A. Ruseckas and I. D. W. Samuel, *J. Phys. Chem. A*, 2009, **113**, 2–4.
19. R. Seifert, I. Rabelo de Moraes, S. Scholz, M. C. Gather, B. Lüssem and K. Leo, *Org. Electron.*, 2013, **14**, 115–123.
20. M. J. Jurow, A. Bossi, P. I. Djurovich and M. E. Thompson, *Chem. Mater.*, 2014, **26**, 6578–6584.
21. I. Rabelo de Moraes, S. Scholz, M. C. Gather, B. Lüssem and K. Leo, *Org. Electron.*, 2011, **12**, 341–347.
22. S. Schmidbauer, A. Hohenleutner and B. König, *Beilstein J. Org. Chem.*, 2013, **9**, 2088–2096.
23. J. Lee, H.-F. Chen, T. Batagoda, C. Coburn, P. I. Djurovich, M. E. Thompson and S. F. Forrest, *Nat. Mater.*, 2016, **15**, 92–98.
24. T. Sajoto, P. I. Djurovich, A. B. Tamayo, J. Oxgaard, W. A. Goddard III and M. E. Thompson, *J. Am. Chem. Soc.*, 2009, **131**, 9813–9822.
25. D. Escudero and D. Jacquemin, *Dalton Trans.*, 2015, **44**, 8346–8355.
26. X. Zhou, P. L. Burn and B. J. Powell, *Inorg. Chem.*, 2016, **55**, 5266–5273.
27. D. Escudero, *Chem. Sci.*, 2016, **7**, 1262–1267.
28. D. Escudero, E. Heuser, R. J. Meier, M. Schäferling, W. Thiel and E. Holder, *Chem.-Eur. J.*, 2013, **19**, 15639.
29. K. Tsuchiya, S. Yagai, A. Kitamura, T. Karatsu, K. Endo, J. Mizukami, S. Akiyama and M. Yabe, *Eur. J. Inorg. Chem.*, 2010, **2010**, 926–933.
30. A. J. Wilkinson, A. E. Goeta and J. A. G. Williams, *Inorg. Chem.*, 2004, **43**, 6513–6515.
31. Y. Chi, T.-K. Chang, P. Ganesan and P. Rajakannu, *Coord. Chem. Rev.*, 2017, **346**, 91–100.
32. H.-H. Kuo, Y.-T. Chen, L. R. Devereux, C.-C. Wu, M. A. Fox, C.-Y. Kuei, Y. Chi and G.-H. Lee, *Adv. Mater.*, 2017, **29**, 1702464.
33. C.-Y. Kuei, W.-L. Tsai, B. Tong, M. Jiao, W.-K. Lee, Y. Chi, C.-C. Wu, S.-H. Liu, G.-H. Lee and P.-T. Chou, *Adv. Mater.*, 2016, **28**, 2795–2800.
34. A. D. Becke, *J. Chem. Phys.*, 1993, **98**, 5648.
35. C. T. Lee, W. T. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1988, **37**, 785–789.
36. D. Andrae, U. Häusermann, M. Dolg, H. Stoll and H. Preuss, *Theor. Chim. Acta*, 1990, **77**, 123–141.
37. J. N. Harvey, M. Aschi, H. Schwarz and W. Koch, *Theor. Chem. Acc.*, 1998, **99**, 95–99.
38. F. Neece, *Orca, an ab initio, DFT and semiempirical SCF-MO package 2.8.0 R2327*, University of Bonn, Bonn, Germany, 2011.
39. L. Goerigk and S. Grimme, *J. Chem. Theory Comput.*, 2011, **7**, 291–309.
40. M. Steinmetz and S. Grimme, *ChemistryOpen*, 2013, **2**, 115–124.
41. M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery Jr, J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi,
J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, Ö. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09, Revision A.1*, Gaussian, Inc., Wallingford CT, 2009.