pH-Responsive Emission of Novel Water-Soluble Polymeric Iridium(III) Complexes

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Abstract: The synthesis and characterization of water-soluble copolymers containing N,N-dimethylacrylamide (DMAM) and a vinylic monomer containing an Iridium(III), Ir(III), complex substituted with the quinoline-based unit 2-(pyridin-2-ylo)-6-styrene-4-phenylquinoline (VQPy) as ligand are reported. These copolymers were prepared through pre- or post-polymerization complexation of Ir(III) with the VQPy units. The first methodology led to copolymer P1 having fully complexed VQPy units, whereas the latter methodology allowed the preparation of terpolymers containing free and Ir(III)-complexed VQPy units (copolymer P2). The optical properties of the copolymers were studied in detail through UV-Vis and photoluminescence spectroscopy in aqueous solution. It is shown that the metal-to-ligand charge transfer (MLCT) emission is prevailing in the case of P1, regardless of pH. In contrast, in the case of terpolymer P2 the MLCT emission of the Ir(III) complex is combined with the pH-responsive emission of free VQPy units, leading to characteristic pH-responsive color changes under UV illumination in the acidic pH region.

Keywords: Iridium polymeric complexes; water-soluble copolymers; water-soluble metallocomplexes; pH-sensors; quinoline-pyridine ligand; photoluminescence

1. Introduction

Transition-metal complexes are well-known for their ability to achieve high-efficiency phosphorescence because they can harvest both singlet and triplet excitons. Motivated by this property, cyclometalated complexes based on metal ions such as Ru(II), Rh(I), Pt(II), and Ir(III) have been largely investigated [1–4]. Among these, iridium(III) complexes have attracted great attention because of their special optical properties [5–7], namely the ability to harvest both singlet and triplet exciton, their high photoluminescence efficiency and high quantum yields, wide range of colors emitted etc. Thus, Ir(III) complexes are promising materials for a wide variety of applications, including among others, organic light emitting diodes (OLEDs) [8–11], bioprobes [12–18], catalysts [19–23], optical chemical sensors [24–31].

Ir(III) complexes are most commonly soluble in organic solvents rather than in aqueous solutions. In contrast, the development of water-soluble luminescent Ir(III) complexes is valuable for biological and biochemical applications such as bioprobes for cell imaging or their use as indicators to determine parameters such as pH, temperature, or oxygen. To this end, the ligands may be functionalized with hydrophilic groups [13,14,32–37] or the counter ions of cationic cyclometalated Ir(III) complexes may be adequately chosen to sustain water-solubility [14,38–40]. Alternatively, the covalent attachment of Ir(III) complexes onto a polymeric chain, using water-soluble polymers such as poly(N-isopropylacrylamide), poly(N-vinylpyrrolidone), and polyethylene glycols, has been also applied [41–50].

The development of pH sensors is of great interest, since they are used in many different areas, such as environmental monitoring, biology, and food industry. For our current...
approach, we take advantage of our experience about the functionalization of polymeric or carbon materials with pH-sensitive luminescent units based on quinoline derivatives. Due to its weak basic character, the protonation/deprotonation of the quinoline group results in the change of the emitted color from blue to green upon decreasing pH [51–54]. The vinyl-functionalization of quinoline derivatives and subsequent copolymerization allowed us to tune these pH-responsive emission properties [55] or to combine them with additional functionalities, such as thermosensitivity [56], selective surfactant sensing [57], as well as encapsulation abilities for hydrophobic magnetic nanoparticles [58].

Recently, in our group, organo-soluble polymeric Ir(III) complexes were successfully prepared through “post-polymerization” or “pre-polymerization” complexation, aiming at OLED applications [59,60]. Our target in the present work is the combination of the aforementioned pH-responsive properties of quinoline derivatives with the optical properties of Ir(III) complexes into a water-soluble polymeric nanostructure. To this end, a vinlyc monomer containing a pyridine-quinoline-based ligand [61], namely 2-(pyridin-2-ylo)-6-styrene-4-phenylquinoline (VQPy), was explored as the polymerizable ligand of Ir(III) ions. Based on this ligand, highly luminescent water-soluble polymeric materials have been developed by introducing the iridium(III) complex (PPy)2Ir(VQPy) (PPy: phenylpyridine) as signaling units along a nonionic hydrophilic N,N-dimethylacrylamide, DMAM, chain (see Scheme 1), through free radical polymerization in organic solvents. While “pre-polymerization” complexation assures that all VQPy units are complexed, “post-polymerization” complexation offers the possibility to control the fraction of complexed VQPy units and to tune thus the optical properties of this metallocomplex nanomaterial.

![Scheme 1. Synthesis of P(DMAM-co-[(PPy)2Ir(VQPy)(PF6)]) P1, through direct free radical polymerization of DMAM with [(PPy)2Ir(VQPy)(PF6)].](image)

2. Materials and Methods

2.1. Materials

The monomer 2-(pyridin-2-ylo)-6-styrene-4-phenylquinoline (VQPy) was synthesized according to literature procedures [62], through a Friedländer reaction followed by a Suzuki cross-coupling reaction. The monomer N,N-dimethylacrylamide (DMAM), the reagent ammonium hexafluorophosphate (NH4PF6), the solvent chloroform (CHCl3) were purchased from Sigma-Aldrich (Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany). The reagents 2-phenylpyridine (PPy), IrCl3·H2O and the solvent 2-ethoxyethanol were purchased from Alfa Aesar (Thermo Fisher (Kandel) GmbH Postfach, Karlsruhe, Germany). The initiator α’–α’ azobisisobutionitrile (AIBN), the reagents for the preparation of the buffer solutions, trisodium citrate dihydrate, and hydrochloric acid, the solvents tetrahydrofuran (THF), diethyl ether (Et2O), and methanol were purchased from Merck (Merck KGaA, Darmstadt, Germany). Ultra-pure water was obtained by means of a SG Water laboratory unit.
2.2. Instrumentation

$^1$H-NMR spectra were obtained on a Bruker Advance DPX 400.13 MHz spectrometer (Bruker BioSpin GmbH, Magnet Division, Karlsruhe, Germany) at 25 °C using deuterated CDCl$_3$ or d$_6$-DMSO. Chemical shifts (δ) are reported in units, parts per million (ppm) downfield from TMS.

Attenuated total reflectance Fourier transform infrared spectra (ATR-FTIR) were recorded on a “Bruker Optics’ Alpha-P Diamond ATR Spectrometer of Bruker Optics GmbH” (Ettlingen, Germany).

A single-quadrupole Quattro micro mass spectrometer (ACQUITY SQ Detector) equipped with an electrospray ionization (ESI) interface was used for analytical detection. ESI-MS was operated in positive mode under the following operating parameters: capillary voltage 3.5 kV, cone voltage 30 V, source temperature 140 °C, desolvation temperature 250 °C, desolvation gas (nitrogen) 500 L/hour, and cone gas (nitrogen) 50 L/h. All data were acquired and processed using Masslynx 4.1 software (Waters Corp., Milford, MA, USA).

Size-exclusion chromatography (SEC) measurements were carried out using a Polymer Lab chromatographer (Agilent Technologies, Santa Clara, CA, USA) equipped with two PLgel 5 μm mixed columns and a UV detector using CHCl$_3$ as the eluent at a flow rate of 1 mL min$^{-1}$ at 25 °C; and calibrated versus polystyrene standards.

The UV–Vis spectra were recorded using a Hitachi-Science and Technology-U-1800 UV–Vis Spectrophotometer in the 190–800 nm wavelength range (Hitachi High-Technologies Europe GmbH, Mannheim, Germany).

Photoluminescence (PL) spectra were recorded using a Perkin–Elmer LS45 luminescence spectrometer (Waltham, MA, USA), using quartz glass cuvettes (optical path = 1 cm). For the photoluminescence study, the excitation and emission slits were 10 nm, while the excitation wavelengths were 350 nm or 400 nm.

2.3. Synthesis of Compounds

[(PPy)$_2$Ir(VQPy)(PF$_6$)] complex. The complex was synthesized in two steps [63,64]. First, the chloro-bridged dinuclear cyclometalated iridium(III) [(PPy)$_2$IrCl$_2$]$_2$ was synthesized: 2-phenylpyridine (PPy) (127 μL, 0.880 mmol) was mixed with IrCl$_3$$^\cdot$H$_2$O (0.106 g, 0.360 mmol) in a mixture of 2-ethoxyethanol and H$_2$O (3/1 v/v) to reach a concentration in IrCl$_3$$^\cdot$H$_2$O of ~0.04 M. The mixture was degassed by multiple vacuum and N$_2$ purging cycles and was heated in an oil bath set at 120 °C for 24 h under N$_2$. The yellow precipitate formed during the reaction was filtered, washed with hot methanol, and dried under vacuum at 40 °C overnight. In the second step, VQPy (0.045 g, 0.120 mmol) and 2.3 mL 2-ethoxyethanol were added to 0.050 g (0.050 mmol) of the yellow precipitate, to reach again a concentration of ~0.04 M. The mixture was degassed by multiple vacuum and N$_2$ purging cycles and was heated in an oil bath set at 120 °C for 24 h under N$_2$, leading to a red solution. The reaction mixture was cooled to room temperature and was extracted several times with Et$_2$O and water. After the addition of a solution of NH$_4$PF$_6$ (10 eq in respect to the dinuclear [(PPy)$_2$IrCl$_2$]$_2$ complex) in the aqueous phase, a red solid was precipitated. The suspension was cooled to 0 °C for 1 h, filtered and washed with H$_2$O. The red solid was dried under vacuum at 60 °C overnight.

Synthesis of P(DMAM-co-[(PPy)$_2$Ir(VQPy)(PF$_6$)]) copolymers. Two different synthetic procedures were followed as depicted in Schemes 1 and 2. Both of them were based on the free radical copolymerization of DMAM with VQPy, either complexed with Ir(III) (Scheme 1) or free (Scheme 2).
Direct copolymerization of DMAM with [(PPy)$_2$Ir(VQPy)(PF$_6$)] (Scheme 1). THF was degassed by bubbling with N$_2$ to remove the dissolved oxygen. The monomers DMAM (0.4458 g, 4.5 mmol) and [(PPy)$_2$Ir(VQPy)(PF$_6$)] (0.0200 g, 0.023 mmol), as well as the initiator AIBN (2% mol over the total monomers quantity), were added in the solvent and the mixture was stirred at 70 °C for 2 days. The reaction mixture was added in Et$_2$O under stirring, leading to the precipitation of a slightly orange solid. The mixture was filtered, and the resulting product $P_1$ was washed several times with cold Et$_2$O and recovered after drying in vacuum at 40 °C. Mn = 10.000 Da, Polydispersity Index = 2.1.

Copolymerization of DMAM with VQPy and subsequent introduction of Ir(III) complex (Scheme 2). According to the second methodology, the synthesis of the P(DMAM-co-VQPy) through free radical polymerization of DMAM with VQPy preceded the complexation reaction. The monomers DMAM (1.5 mL 14.800 mmol) and VQPy (0.300 g 0.780 mmol) and the initiator AIBN (0.0513 g, 0.3124 mmol) were added in freshly distilled and deoxygenated THF (18 mL) and the mixture was stirred at 70 °C for 2 days. The polymer was precipitated in Et$_2$O and the white solid was filtered, washed with cold Et$_2$O, and dried under vacuum. Mn = 15.000 Da. Polydispersity Index = 1.8.

In the second step, the copolymer P(DMAM-co-VQPy) (0.4595 g, 0.2300 meq of VQPy) was mixed with the chloro-bridge dinuclear [(PPy)$_2$IrCl$_2$] complex (0.050 g, 0.093 mmol of Ir) in 2-ethoxyethanol (2.3 mL, the ethoxyethanol/Ir molar ratio was about 4) at 130 °C for 24 h under N$_2$. The reaction mixture was cooled to room temperature and 3 mL deionized H$_2$O was added. Then the biphasic solution was extracted with H$_2$O and Et$_2$O and the collected water phase was heated at 70 °C for 30 min. Then NH$_4$PF$_6$ was added (10 eq in respect to the dinuclear [(PPy)$_2$IrCl$_2$] complex) without however leading to the precipitation of the polymeric complex. Thus, the polymer $P_2$ was purified through dialysis (membrane.
3. Results
3.1. Synthesis and Characterization of Compounds

The vinylic complex [(PPy)₂Ir(VQPy)(PF₆)] was synthesized in two steps, involving initially the preparation of the chloro-bridged dinuclear cyclometalated iridium(III) complex [(PPy)₂IrCl₂] and the subsequent reaction of this complex with the vinylic monomer VQPy. The final complex was characterized through ¹H NMR and MS-ESI. The detailed attribution of the peaks observed in the ¹H NMR spectrum is shown in Figure 1. As seen, the characteristic peaks of the vinylic bond and the peaks attributed to the aromatic groups of the complex are clearly detected. In addition, the verification of the synthesis was also based on the determination of the molecular weight of the complex through MS-ESI (Figure 1, inset). The main product of the sample with a molecular weight of 885.56 g/mol is revealed in the MS-ESI spectrum. Having in mind that the expected molecular weight of the complex is 886.62 g/mol, this suggests that the desired complex has been successfully synthesized.

![Figure 1. ¹H NMR spectrum of complex [(PPy)₂Ir(VQPy)(PF₆)] in DMSO-d6. The inset shows the respective MS-ESI spectrum.](image)

For the incorporation of this complex into the water-soluble poly(N,N-dimethylacrylamide) chain, two alternative methodologies were followed. According to the first one, namely pre-polymerization Ir(III) complexation, the target copolymer P(DMAM-co-[(PPy)₂Ir(VQPy)(PF₆)]) was synthesized via free radical copolymerization of the monomer DMAM with the vinyl-modified complex [(PPy)₂Ir(VQPy)(PF₆)] (Scheme 1, product P1). The feed molar content of the Ir(III) complex was adjusted at 0.5 mol% for this product.

The second methodology, namely the post-polymerization Ir(III) complexation, is illustrated in Scheme 2. For this route, as a first step, the copolymer P(DMAM-co-VQPy) was initially synthesized through free radical copolymerization in an organic solvent. The molar feed composition of VQPy units was adjusted at ~5 mol%. The second step was the complexation of VQPy units of the copolymer with Ir using the chloro-bridged dinuclear complex [(PPy)₂IrCl₂]. The molar ratio of [(PPy)₂IrCl₂] over VQPy units of the copolymer (based on feed composition) was 0.2. As a consequence, the final product is expected...
to be a terpolymer, namely P(DMAM-co-VQPy-co-[(PPy)2Ir(VQPy)(PF6)]) (product P2), containing comparable contents of complexed and free VQPy units.

The products P1 and P2 were characterized by 1H NMR and ATR-IR. As seen in Figure 2, where the 1H NMR spectrum of product P1 is shown as an example, weak peaks are observed in the region 7–9 ppm, as a consequence of the aromatic content owing to the [(PPy)2Ir(VQPy)(PF6)] units of this copolymer. In addition, the peaks in the 2.5–3 ppm region are attributed to the methyl groups of the DMAM units, while the methylene and methine protons of the polymeric backbone are found in the 1–2.5 ppm region.

The ATR spectra of products P1 and P2 are compared in Figure 2b with the respective spectra of the copolymer P(DMAM-co-VQPy) and the vinylic Ir(III) complex [(PPy)2Ir(VQPy)(PF6)]. In the spectrum of the monomer complex, the peaks at ~840 cm\(^{-1}\) and ~550 cm\(^{-1}\) are attributed to the existence of hexafluorophosphate. As it can be observed,
the same peaks appear at the spectrum of the copolymer P(DMAM-co-[(PPy)2Ir(VQPy)(PF6)]) but do not exist at the spectrum of the copolymer P(DMAM-co-VQPy). Finally, the peak at the 1610 cm⁻¹ is attributed to quinoline VQPy double bond (C=C and C=N) of the aromatic rings. The peaks at ~2900 cm⁻¹ of the copolymers’ spectra are attributed to the main polymer backbone C-H bonds.

3.2. Optical Properties

The optical properties of the Ir(III)-containing copolymer obtained through both methodologies were investigated in aqueous solution. It is worthy to note that the initial complex [(PPy)2Ir(VQPy)(PF6)] is practically insoluble in water and thus, no UV-Vis absorbance is monitored when trying to study the aqueous dispersion of this material, as shown in Figure 3. In contrast, it is readily soluble in chloroform, exhibiting the characteristic properties of such Ir(III) complexes. In particular, the strong absorption band at ~300 nm, closely resembling the absorption spectrum of the free ligand VQPy [54], is attributed to the π-π* transition of the quinoline group. In addition, the complex presents weaker absorption bands in the range of 350–450 nm, which are attributed to the metal-to-ligand charge transfer (MLCT) transitions. More precisely, the absorption band at 350–400 nm is assigned to singlet metal-to-ligand charge transfer (1MLCT) transitions, whereas the weaker absorption area above 400 nm is attributed to triplet metal-to-ligand charge transfer (3MLCT) transitions [65,66]. The same absorption bands are also observed in the cases of the two copolymers. However, it should be stressed that now this behavior is monitored in aqueous solution, proving that the methodologies elaborated in the present work can effectively be used for the preparation of water-soluble polymeric Ir(III) complexes focusing on optical applications in aqueous environments.

In addition to the absorption characteristics, the copolymers maintain also the emission characteristics of the Ir(III) complexes in aqueous environment, as seen in Figure 4 where the normalized photoluminescence spectra of products P1 and P2 in water and of the initial vinylic [(PPy)2Ir(VQPy)(PF6)] complex in DMSO are compared, upon excitation at 400 nm. Under these conditions an emission peak at 610 nm is clearly detectable. This characteristic band is assigned to the strong singlet and triplet metal-to-ligand charge transfer (MLCT) transition of the Ir(III) complex. As a result, the aqueous solutions of products P1 and P2 emit red light under UV illumination as shown in the right part of Figure 4.

Figure 3. UV-Vis absorption spectra of the copolymers P(DMAM-co-[(PPy)2Ir(VQPy)(PF6)]) P1, P(DMAM-co-VQPy-co-[(PPy)2Ir(VQPy)(PF6)]) P2, in aqueous solution and of the complex [(PPy)2Ir(VQPy)(PF6)] in CHCl3 and in water.
we recorded the photoluminescence spectra of terpolymer P2 containing both Ir(III)-complexed and free VQPy units, a very interesting pH-controlled photoluminescence behavior was observed. To demonstrate this behavior, we recorded the photoluminescence spectra of terpolymer P2 in buffer solutions covering the pH range 2–8, upon excitation at 350 nm and 400 nm (Figure 5). As seen in Figure 5, the red emission peak of the Ir(III) complex at 610 nm is rather constant, regardless of pH. On the other hand, it is known that the emission properties of quinoline-based products are controlled by the acidity of the environment, as a consequence of the weak basic character of quinoline units [52,53,62]. More specifically, quinoline-based polymers can respond to the acidity of their solution due to the weak basic character of quinoline. The protonation of the quinoline unit when incorporated in a polymer, results into a small red-shift of the polymer’s absorption spectrum. The emission spectrum, on the other hand, is more evidently red-shifted (even up to 100 nm). Moreover, the emission peak of the protonated quinoline-based polymers is much broader and significantly weaker. The large shift of the emission maximum to higher wavelengths in acidic media is an indication for the formation of aggregates originating from intra- or intermolecular interactions attributed to acid-based interactions of the quinoline group leading to excimer formation [52]. Actually, quinoline homopolymers or quinoline-rich copolymers show significantly quenched emission of the protonated form of the quinoline unit [62]. On the other hand, the green emission of the protonated quinoline unit can be clearly observed when luminescence quenching is suppressed thus, not allowing the formation of excimers, e.g., when fixed onto carbon nanotubes [67] or when low fractions of quinoline units are randomly fixed onto cross-linked polymeric nanoparticles [54]. Through this latter approach pH-sensing applications have been demonstrated for water-soluble quinoline-based copolymers having low quinoline content. These quinoline-labeled water-soluble polymers exhibit sensitive pH-responsive photoluminescent properties, showing blue to green color gradual changes under UV irradiation upon decreasing pH [51,55].

Figure 4. Photoluminescence spectra of the copolymers P1 and P2 in aqueous solution and of the monomer complex [(PPy)$_2$Ir(VQPy)(PF$_6$)] in DMSO, after excitation at 400 nm. The photographs in the right, show the aqueous solution of P1 in daylight (left photo) and under UV illumination at 365 nm (right photo).
Thus herein, in agreement with previous observations by our group for similar water-soluble copolymers bearing free quinoline-based units [51,55–58], an emission peak at ~420 nm is observed in Figure 5a upon excitation of aqueous P2 solutions at 350 nm at high pH, attributed to the unprotonated form of VQPy. Apparently, for our system this blue emission does not perturb the observed color of the solution, which remains red from pH = 8 down to pH = 4.5 (Figure 6, left), due to the emitted color of the Ir(III) complex. Upon further pH decrease, free VQPy units turn to their protonated form and an emission band, reflecting a blue-green color, centered at ~480 nm is seen, while the emission band at 420 nm gradually disappears. Apart from the band at 610 nm, the evolution of both bands is clearly observable when the system is excited at 350 nm (Figure 5a), whereas only the evolution of the green emission of VQPy is monitored upon excitation at 400 nm (Figure 5b), since the absorption of this unit at 400 nm is negligible. As a result of the combination of this green emission from VQPy with the red emission of the Ir(III) complex, a gradual color change of the solution as pH decreases below the value pH = 4.5 is seen, to finally attain a yellow-green color at pH = 2.5 (Figure 6).
we proceeded to the full complexation of the remaining free VQPy units with an excess of VQPy units. In this case, the MLCT emission of the Ir(III) complex is combined with the (VQPy) has been used in the present work for the preparation of water-soluble polymeric copolymers in aqueous solution. In fact, it is shown that the post-polymerization complexation methodology may lead to terpolymers containing both free and Ir(III)-complexed VQPy moieties.

The observed color changes are due to the terpolymer’s exact chemical structure in which the coexistence of free VQPy and iridium-complexed VQPy moieties allows the protonation of the free VQPy at acidic pH environments, therefore leading to PL emission changes and visibly clear color changes of the polymer metallocomplex solutions at the various pH values. This combination of the emission of free VQPy units and Ir(III) complexes, demonstrates that such terpolymer libraries with adequately adjusted compositions may permit the fine tuning of this pH-responsive emission in the acidic region.

Though such a detailed study will be the subject of a future work, to test the basic idea, i.e., that the color changes are the result of the terpolymer chemical structure of P2, we proceeded to the full complexation of the remaining free VQPy units with an excess of the chloro-bridge dinuclear [(PPy)2IrCl2]2 complex to obtain the copolymer P2 (m = 0) (Scheme 2 P2 in which m = 0). In this case, no free VQPy are available anymore and the red color persists within the whole pH region, even if pH is as low as 1 (Figure 6, right).

4. Conclusions

The complexation of Ir(III) with a vinylic unit containing quinoline-pyridine groups (VQPy) has been used in the present work for the preparation of water-soluble polymeric Ir(III) complex nanomaterials. Two synthetic methodologies have been elaborated, namely pre- or post-polymerization complexation of Ir(III) with VQPy.

Here, we mostly focused on the optical properties, especially emission, of the novel copolymers in aqueous solution. In fact, it is shown that the post-polymerization complexation methodology may lead to terpolymers containing both free and Ir(III)-complexed VQPy units. In this case, the MLCT emission of the Ir(III) complex is combined with the pH-responsive emission of free VQPy units, leading to characteristic pH-responsive color changes under UV illumination in the acidic pH region. Apart from the preparation of pH-responsive luminescent sensory nanomaterials, the present materials could also prove useful for other applications where water-solubility of luminescent Ir(III) complexes is needed, for example cell imaging applications. Moreover, the methodologies elaborated in the present work offer two facile and easily adaptable routes for the incorporation of Ir(III) complexes in more complicated polymeric architectures [68,69].

Figure 6. Demonstration of the pH induced color changes of aqueous solutions of copolymer P(DMAM-co-VQPy-co-[(PPy)2Ir(VQPy)(PF6)]) and of the persistent red color of the P2 copolymer bearing only Ir(III)-complexed VQPy moieties. All solutions are shown under UV illumination at 365 nm.
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