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Electroluminescent properties of lanthanide pentafluorophenolates

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Lanthanide pentafluorophenolates Ln(OC\textsubscript{5}F\textsubscript{3})(L\textsubscript{x})(Ln = Pr, Nd, Sm, Eu, Dy, Ho, Er, Yb; L = 1,10-phenanthroline, 2,2'2bipyridine; x = 1, 2) are used as emissive layers in organic light emitting devices (OLEDs). Single-layer ITO/Ln(OC\textsubscript{5}F\textsubscript{3})(L\textsubscript{x})/Yb devices reveal no electroluminescence (EL) with the exception of Tb-derivatives-based OLEDs. Bilayer ITO/TPD/ Ln(OC\textsubscript{5}F\textsubscript{3})(L\textsubscript{x})/Yb samples exhibit a broad band emission peaked at 580 nm assigned to electroplex at the TPD/complex interface. Besides, the spectra of the devices based on Pr, Nd, Sm, Eu, Er, Tb and Yb derivatives contain the characteristic narrow bands of f-f transitions. Terbium-based bilayer OLEDs exhibit unusual changes in the EL spectra with increased applied voltage. The emission color of the devices tunes from orange to green. The possible nature of the phenomenon is discussed.

1. Introduction

It is known that the luminescence properties of organic complexes of non-transition, d-transition and lanthanide metals in a great extent depend on the nature of the ligands. Analysis of literature data reveals that virtually all of metal-organic luminophores contain bi- or polydentate ligands which provide an extended conjugated chains (necessary for luminescence) and sufficient stability of the material. In the case of organolanthanide emitters the ligands play a role of sensitizers, excitation energy from which is transferred to emissive f sublevels of the metal ion increasing their luminescence. Fluorescent complexes with monodentate anionic ligands are rare. The group of luminophores containing one monodentate ligand is limited by O-quinolinolinate-aryloxides of aluminum (or gallium) of the type Al(q)\textsubscript{2} (OR)\textsubscript{3} and similar N,N-bis(3,5-di-tert-butyl-salycilidene)-ethylene-diamine derivatives (\textit{tBu})\textsubscript{3}Al(OC\textsubscript{6}H\textsubscript{4} p-R)\textsubscript{3}. Photo- or electroexcitation of the complexes results in blue or green-blue emission of moderate efficiency. The luminescent properties of the complexes with two monodentate ligands were not studied. A set of metal-organic luminophores with three non-chelating anionic OC\textsubscript{5}F\textsubscript{3} and SC\textsubscript{5}F\textsubscript{5} ligands has been prepared and studied by Brennan, Riman and coauthors. The complexes (DME)\textsubscript{2}Ln(OC\textsubscript{5}F\textsubscript{3})\textsubscript{3} (Ln = Nd, Er, Tm)\textsuperscript{20} and (DME)\textsubscript{2}Ln(SC\textsubscript{5}F\textsubscript{5})\textsubscript{3} (Ln = Nd\textsuperscript{19}, Er\textsuperscript{17}, Tm\textsuperscript{16}) showed significantly greater metal-centered NIR PL efficiency than that of previously reported non-fluorinated compounds. High quantum yield of metal-centered PL was observed for the clusters containing thiophenolate and selenophenolate ligands (THF)\textsubscript{3}Nd\textsubscript{5}O\textsubscript{2}Se\textsubscript{3} (SePh)\textsubscript{19}, (THF)\textsubscript{3}Tm\textsubscript{5}Se\textsubscript{3} (SC\textsubscript{5}F\textsubscript{5})\textsubscript{2}, (DME)\textsubscript{3}Al(C\textsubscript{6}F\textsubscript{5})(SC\textsubscript{5}F\textsubscript{5})\textsubscript{2}. Both, the absence of C-H bonds in the anionic ligand and the low Ln-S or Ln-Se phonon energies are thought to minimize competitive vibrational relaxation pathways, and in turn result in the formation of exceptionally emissive molecules. As far as we know, the EL properties of the complexes with exclusively non-chelating anionic ligands have never been studied. To close this gap, we synthesized a series of lanthanide pentafluorophenolates and tested them as emissive materials in single- and bilayer OLED devices.

2. Results and discussions

2.1. Synthesis

Among the lanthanide complexes with pentafluorophenolate ligands the promising as emitters in the OLED devices could be simplest tris(phenolates) Ln(OC\textsubscript{5}F\textsubscript{3})\textsubscript{3} because they do not contain the ancillary groups with luminescence quenching C-H, N-H or O-H fragments. However, low thermal stability and high hygroscopicity of these compounds\textsuperscript{21} prevent the preparation of thin films by vacuum sublimation or spin-coating methods. In the case of vacuum deposition the complexes explode on the evaporator, while the films prepared from solutions by spin-coating are quickly destroyed because of water absorption. Both of these drawbacks of the tris(phenolates) are caused by open coordination sphere of the metal ion (vide infra). To solve the problem we synthesized the phenolates containing 1,10-phenanthroline (Ln(OC\textsubscript{5}F\textsubscript{3})(phen) – 1(Ln), Tb(OC\textsubscript{5}F\textsubscript{3})(phen)\textsubscript{2} – 2) or 2,2'-bipyridine (Tb(OC\textsubscript{5}F\textsubscript{3})(bpy) – 3, Tb(OC\textsubscript{5}F\textsubscript{3})(bpy)\textsubscript{2} – 4) as neutral ancillary ligands. As expected, the introduction of additional groups in the coordination sphere of the lanthanides significantly increased thermal and hygroscopic stability of the complexes.\textsuperscript{21}
3.560 Å) indicate an existence of some intermolecular π-π interactions.

Our attempts to prepare other phenolates in the crystalline form suitable for X-ray analysis failed. The structures some of the complexes were determined using Quantum chemical modeling. According to these data the complex I(Tb) exists in the form of dimer with two \( \text{C}_6\text{F}_5\text{O} \) bridges (phen)\( \text{Tb} \)\( \text{O} \)(\( \text{OC}_6\text{F}_5 \))\( \text{Tb} \)\( \text{O} \)(\( \text{OC}_6\text{F}_5 \))(phen) (Fig. S1). Energy of this molecule is 19 kcal/mol lower than that of system of two monomers \( \text{Tb} \)\( \text{O} \)(\( \text{OC}_6\text{F}_5 \))(phen) (Fig. S2(a)) and 34 kcal/mol lower than that of the dimer with bridging phen ligands (Fig. S3). The phenolates I(Pr, Nd, Sm, Yb) are isoostructural to I(Tb) (Table S1).

The DFT calculations revealed as well that in three coordinated phenolates \( \text{Ln} \)\( \text{O} \)(\( \text{C}_6\text{F}_5 \)) there are weak intermolecular F→\( \text{Ln} \) interactions in axial directions. These interactions persist in dimer form with four \( \text{C}_6\text{F}_5\text{O} \) bridges (Fig. S4) which is formed with energy gain ca. 40 kcal/mol. Presumably, such an interaction causes low thermal stability of the perfluorinated phenolates free of ancillary neutral ligands because facilitates the formation of strong Ln-F bonds. Coordination to the metal center of 1,10-phenanthroline or 2,2'-bipyridine molecules in mononuclear complexes \( \text{Ln} \)\( \text{O} \)(\( \text{C}_6\text{F}_5 \)) leads to large energy release about 45 and 25 kcal/mol for the first and the second neutral ligand, respectively, and suppresses the dimerization. Since the coordination sphere of the metal ion turned out filled the thermal and hygroscopic stability.

2.3. Electroluminescence

The complexes \( \text{Ln} \)\( \text{O} \)(\( \text{C}_6\text{F}_5 \))(phen), \( \text{Ln} \)\( \text{O} \)(\( \text{C}_6\text{F}_5 \))(phen)_2, \( \text{Ln} \)\( \text{O} \)(\( \text{C}_6\text{F}_5 \))(bpy), \( \text{Ln} \)\( \text{O} \)(\( \text{C}_6\text{F}_5 \))(bpy)_2 can be sublimed in vacuum or spin casted from solutions to give amorphous stable in air stout entire films. That feature allowed to use them as emissive layers in OLEDs and investigate their EL properties. We have found that the EL properties of all the obtained compounds are similar except the Tb derivatives. In the single-layer devices the tested complexes showed no EL and relatively low charge-transporting properties judging by current-voltage curves (Fig. S8). In contrast, the devices based on Tb-complexes displayed intense bands of \( ^2\text{D}_j \rightarrow ^7\text{F}_j \) (j=6-3) transitions characteristic of Tb\(^{3+} \) ion (Fig. S9).

It is known that the insertion of the hole-transporting layer (HTL) between the anode and the emissive layer as well as the hole-blocking layer (HBL) between the emissive layer and cathode as a rule significantly increases efficiency of the OLEDs. In the case of the devices based on lanthanide perfluoro phenolates insertion of HTL also leads to improvement their performance but to a different extent depending on the metal. In OLEDs based on \( \text{I} \)\( \text{Pr}, \text{Nd}, \text{Sm}, \text{Eu}, \text{Dy}, \text{Ho}, \text{Er}, \text{Yb} \) insertion of TPD layer between ITO and phenolate layer resulted in appearance of broad peak at 580 nm (Table 1, Fig. S10), which can be assigned to emission of the electroluminescence at TPD/complex interfaces. This assumption is confirmed by photoluminescence (PL) spectra of double layer \( \text{I} \)\( \text{Ln} \)/TPD and blend samples \( \text{I} \)\( \text{Ln} \)/TPD containing the broad band of TPD at about 420 nm and in the cases of Sm, Dy and Tb derivatives the narrow bands of f-f transitions characteristic of the respective Ln\(^{3+} \) cation (Fig. S11). The intensity of the band at 580 nm in the EL spectra varies slightly and the maxima remains virtually the same on going from...
one lanthanide to another. In the EL spectrum of device ITO/TPD/I(Sm)/Yb the band of the electroplex at 580 nm is overlapped with the band of metal-centered emission at 600 nm (\(^2G_5\rightarrow^4H_{5,2}\) transition) and manifests as a shoulder. Unlike the enhanced PL intensity of the similar lanthanide pentafluorophenolates reported by Brennan and coworkers,\(^{20}\) the EL of the bilayer OLEDs based on the obtained compounds turned out comparable or even lower than those of devices with most conventional non-fluorinated organo-lanthanide luminophores.\(^{6,8}\) Extremally poor performance characteristics were found for the diode based on I(Eu) which exhibited very weak bands at 580 nm and 615 nm (\(^5D_0\rightarrow^7F_2\) transition). In contrast, the device with I(Yb) showed high luminescence efficiency. Weak EL activity of Eu complexes and enhanced characteristics of Yb analogs we observed earlier with substituted phenolates\(^{21}\) and naphthalates\(^{22}\) of these metals. It was assumed that great difference in the EL properties of Eu and Yb derivatives is caused by the difference stability of Ln\(^{3+}\) ions and energy levels of excited [Ln\(^{3+}\)*] ions.\(^{23}\) It is possible that the same reasons are responsible for the difference in EL efficiency of Eu and Yb pentafluorophenolates. Insertion of HBL (Bath, 4,7-biphenyl-1,10-phenanthroline, 20 nm) in the ITO/TPD/I(Tb)/Yb device between I(Tb) and Yb-cathode did not enhance the brightness but a little increased the operating voltage due to the increased resistance of the sample.

### 2.4. Electroluminescence of devices based on Tb-complexes

Dramatic difference in EL behavior was found for Tb derivatives. First, in contrast to the phenolates of other lanthanides, the compounds I(Tb), 2, 3 and 4 exhibit the intense metal-centered EL not only in the devices containing hole-transporting layer but also in the single-layer OLEDs. Shape of EL spectra of single-layer devices was virtually the same as that of bilayer samples at high voltage while the luminescence intensity (50 cd/m\(^2\)) and other characteristics (current efficiency 1.12 cd/A, power efficiency 0.3 lm/W, EQE 1.8%) were even higher (Fig. S2). Note, that triplet level of the ligand (20408 cm\(^{-1}\), determined from low temperature PL spectra of I(Gd)) matches well with the hypersensitive \(^5D_1\) level of Tb\(^{3+}\) ion (20457 cm\(^{-1}\)) providing complete excitation energy transfer from the ligands to the metal. The second, the EL spectra of bilayer devices based on Tb-complexes with TPD or NPD as a HTL vary greatly with voltage increasing. The luminance appeared at 4V as a single broad band centered at 580 nm. Its intensity increased with increasing voltage up to 16 V but then gradually reduced and virtually vanished at 20 V. Simultaneously, at 14 V four characteristic narrow bands arising from Tb\(^{3+}\) ion at 491, 546, 586 and 622 nm (\(^5D_4\rightarrow^7F_{j=6-3}\)) appeared on a background of the first band. The intensity of these bands run up with voltage increasing and reached maximum at 20 V as it is shown in Fig. 2a for device ITO/TPD/I(Tb)/Yb. Change of EL spectra is accompanied by change of the luminescence color from orange (CIE, x 0.53; y 0.43) at 10 V to green (CIE, x 0.34; y 0.46) at 20 V (Fig. 2b). The same dependence was found when HTL was N,N'-biphenyl-N,N'-bis(1-naphthyl)-1,1'-biphenyl-4,4'-diamine (NPD) (Table 2). However, the phenomenon was not observed when 4,4'-dicarbazolyl-1,1'-biphenyl (CBP) was used as HTL.

The bilayer OLEDs with phenolates of other lanthanides also revealed dependence of EL spectra on applied voltage but to a much lesser degree. No strong enhancement of f-f emission at high voltages was observed but electroluminescence still tends to decay. Since the ligand composition, structure and energy levels of frontier orbitals, determined from the data of cyclic voltamperometry and DFT calculations (Table S2), are the same or very similar the only explanation for the phenomenon

Table 1 Performance characteristics of the devices of ITO/TPD/Ln-complex/Yb (L = OC\(_3\)F\(_3\))

<table>
<thead>
<tr>
<th>Complex</th>
<th>(V_0) [V](^{\text{a}})</th>
<th>(\lambda_{\text{max}}) [nm]</th>
<th>(I_{\text{max}})(^{\text{b}}) [(\mu)W/cm(^2)]</th>
<th>(\eta_p) [cd/A]</th>
<th>(\eta_p) [lm/W, **mW/W]</th>
<th>EQE(^{\text{c}}) [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pr(L)(phen)</td>
<td>5</td>
<td>580</td>
<td>9 (13)</td>
<td>0.09</td>
<td>0.03</td>
<td>0.09</td>
</tr>
<tr>
<td>Nd(L)(phen)</td>
<td>7.5</td>
<td>580</td>
<td>5 (20)</td>
<td>0.01</td>
<td>&lt;0.01</td>
<td>0.14</td>
</tr>
<tr>
<td>Sm(L)(phen)</td>
<td>5</td>
<td>566</td>
<td>40 (12)</td>
<td>0.35</td>
<td>0.1</td>
<td>1.1</td>
</tr>
<tr>
<td>Eu(L)(phen)</td>
<td>10</td>
<td>580</td>
<td>&lt;1 (19)</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Dy(L)(phen)</td>
<td>5</td>
<td>580</td>
<td>25 (17.5)</td>
<td>0.26</td>
<td>0.07</td>
<td>0.21</td>
</tr>
<tr>
<td>Ho(L)(phen)</td>
<td>7</td>
<td>580</td>
<td>25 (18.5)</td>
<td>0.3</td>
<td>0.08</td>
<td>0.24</td>
</tr>
<tr>
<td>Er(L)(phen)</td>
<td>6</td>
<td>580</td>
<td>8 (15)</td>
<td>0.07</td>
<td>0.02</td>
<td>0.06</td>
</tr>
<tr>
<td>Yb(L)(phen)</td>
<td>6</td>
<td>580</td>
<td>29 (18.5)</td>
<td>0.31</td>
<td>0.08</td>
<td>1.2</td>
</tr>
<tr>
<td>Yb(phen)</td>
<td>979</td>
<td>90* (18.5)</td>
<td>-</td>
<td>0.8**</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^{a}\)Voltage at 0.1 mA/cm\(^2\); \(^{b}\)values in parentheses are the voltage at which the luminance was obtained; \(^{c}\)value at 12 V.

Fig. 2 EL spectra (a) and CIE coordinates (b) of ITO/TPD (20 nm)/I(Tb) (40 nm)/Yb device recorded at bias voltage from 10 to 20 V.

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observed in the diodes with terbium complexes is a felicitous correlation energies of the triplet and the resonance levels of Tb3+ ion. The change of EL spectra with increasing of operating voltage is typical for many OLEDs of various compositions. However, such dramatic alterations as in the devices with Tb pentafluorophenolates to the best of our knowledge were not realized with exclusion of air and water using Schlenk techniques. Reagent-grade chemicals were used in all experiments, and solvents were purified using standard procedures. The lanthanide pentafluorophenolates \(\text{Ln}[(\text{OC}F_5)_2]\) were prepared according to the published procedures. CONTENT of metal in the products was determined by complexometric titration. The lanthanide pentafluorophenolates \(\text{Ln}[(\text{OC}F_5)_2](\text{phen})\) \((\text{Ln} = \text{Pr}, \text{Nd}, \text{Er}), \text{Tb}[(\text{OC}F_5)_2](\text{phen})_2\) \((2)\),

<table>
<thead>
<tr>
<th>Complex</th>
<th>HTL</th>
<th>Luminance</th>
<th>(I_{\text{max}}^{b)^{[\text{cd/m}^2]}})</th>
<th>(\eta_{\text{p}}^{a)^{[\text{cd/A}]}})</th>
<th>(\eta_{\text{p}}^{a)^{[\text{lm/W}]}})</th>
<th>EQE(^{c)^{[%]}})</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{Tb}(L)^3)(\text{(phen)})</td>
<td>TPD</td>
<td>electrolplex</td>
<td>6 (14)</td>
<td>0.31</td>
<td>0.07</td>
<td>0.8</td>
</tr>
<tr>
<td></td>
<td>(\text{f-f})</td>
<td>28.5</td>
<td>0.34</td>
<td>0.06</td>
<td>0.59</td>
<td></td>
</tr>
<tr>
<td></td>
<td>NPD</td>
<td>electrolplex</td>
<td>2 (19)</td>
<td>0.05</td>
<td>&lt;0.01</td>
<td>0.13</td>
</tr>
<tr>
<td></td>
<td>(\text{f-f})</td>
<td>30 (27)</td>
<td>0.13</td>
<td>0.02</td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CBP</td>
<td>electrolplex</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>(\text{Tb}(L)^3)(\text{(bpy)})</td>
<td>TPD</td>
<td>electrolplex</td>
<td>4 (13)</td>
<td>0.35</td>
<td>0.09</td>
<td>0.9</td>
</tr>
<tr>
<td></td>
<td>(\text{f-f})</td>
<td>45 (19)</td>
<td>0.26</td>
<td>0.04</td>
<td>0.45</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(\text{f-f})</td>
<td>40 (19)</td>
<td>0.34</td>
<td>0.05</td>
<td>0.6</td>
<td></td>
</tr>
<tr>
<td>(\text{Tb}(L)^3)(\text{(bpy)})</td>
<td>TPD</td>
<td>electrolplex</td>
<td>5 (12.5)</td>
<td>0.4</td>
<td>0.1</td>
<td>1.0</td>
</tr>
<tr>
<td></td>
<td>(\text{f-f})</td>
<td>40 (20)</td>
<td>0.35</td>
<td>0.05</td>
<td>0.6</td>
<td></td>
</tr>
</tbody>
</table>

\(^{a}\)Voltage at 0.1 mA/cm\(^2\); \(^{b}\)values in parentheses are the voltage at which the luminance was obtained; \(^{c}\)value at maximum luminance.

The change of EL spectra with increasing of operating voltage is typical for many OLEDs of various compositions. However, such dramatic alterations as in the devices with Tb pentafluorophenolates to the best of our knowledge were observed for the first time. In most cases with bilayer and multilayer non-doped devices change of EL spectra upon increasing of applied voltage is explained by the offset of the electron-hole recombination zone from one emissive layers to another due to differences of the energy barriers for the injection of electrons and holes into these layers.\(^{25-29}\) In the devices with the doped emissive layers or with blend luminophores the change of voltage-controlled luminance color is caused by shift of the emission from matrix to dopant or from one component of a blend to another.\(^{30}\) The shift of the luminance color from green-yellow to almost white upon increasing the bias voltage in the device based on Pt(acac)(ppy-R) (acac-acetylacetone, ppy-R – substituted phenylpyridyl) complex was explained by the change of the relative triplet-singlet intensity ratio.\(^{31,32}\) The dependence of EL spectra on bias voltage for the device with Sm(TTA)\(3^+(\text{TPPO})_2\) emissive layer (TTA - thenoyltrifluoroacetonate, TPPO - triphenylphosphine oxide) Reyes and coworkers explained by different intensity of the metal-centered and ligand-centered emission at different voltages.\(^{33}\) The change of visible EL spectra of bilayer device TPD/Ho(DBM)_4(Bath) (DBM - dibenzoylmethanato, Bath - bathophenanthroline) at various voltages the authors attributed to luminescence of interfacial exciplex.\(^{34}\) The emission intensity of the exciplex showed a tendency to saturation at high driving voltage, while the emissions of the Ho\(^{3+}\) ions kept increasing resulting in change of the luminescence of the Ho\(^{3+}\) ions. Varying of emission colour of bilayer device NPD/Zn(4-TimBTZ)\(_2\) (4-TimBTZ - 2-(4-trifluoromethyl-2-hydroxyphenyl)benzothiazole) the authors explained by electrolyte formation in the interface.\(^{35}\) The intensity of the electroluminescence was found to decrease progressively with applied voltage. We suppose that peculiarities of EL of devices based on Tb pentafluorophenolates is also caused by the formation of electroluminescence at TPD/Tb-complex interface and relatively low hole-

3. Experimental Section

3.1. General procedure

The syntheses and manipulations of the compounds described below were realized with exclusion of air and water using Schlenk techniques. Reagent-grade chemicals were used in all experiments, and solvents were purified using standard procedures and collected in a reaction vessel by condensation in vacuum. Pentafluorophenol, 2,2'-bipyridine and 1,10-phenanthroline (purchased from Aldrich) were purified and dried by sublimation under reduced pressure. Silylamides \(\text{Ln}[(\text{SiMe}_3)_2]\) were prepared according to the published procedures.\(^{37}\) CONTENT of metal in the products was determined by complexometric titration. The lanthanide pentafluorophenolates \(\text{Ln}[(\text{OC}F_5)_2](\text{phen})\) \((\text{Ln} = \text{Pr}, \text{Nd}, \text{Er}), \text{Tb}[(\text{OC}F_5)_2](\text{phen})_2\) (2),
Tb(OC\textsubscript{3}F\textsubscript{3})(bpy)\textsubscript{2} (4) were synthesized by the reactions of amides Ln[N(SiMe\textsubscript{3})\textsubscript{2}] with pentafluorophenol in benzene or toluene in the presence of 1,10-phenanthroline or 2,2\textsuperscript{′}bipyridine as described in the paper.\textsuperscript{21} Preparation and characterisation of Ln(OC\textsubscript{3}F\textsubscript{3})(phen) (H(Sm, Eu, Dy, Ho, Tm, Yb)) and Tb(OC\textsubscript{3}F\textsubscript{3})(bpy) (3) are described in the Supplementary information.

3.2. Photophysical measurements

IR spectra were recorded on a Specord M-75 spectrometer from 4000 to 450 cm\textsuperscript{-1}. The samples were prepared as Nujol mulls and as films between KBr plates. UV-vis spectra were recorded with a Perkin Elmer Lambda 25 spectrometer in a region from 200 to 1100 nm. The photoluminescence spectra were recorded from 400 to 800 nm on a spectrometer Perkin Elmer Fluorescence LS-55 (spectral resolution 1 nm) with excitation at 270 nm. The samples were prepared as the films on optical quartz.

3.3. Cyclic voltammetry

Cyclic voltammetry of the complexes was performed in a typical three-electrode cell with a platinum sheet working electrode, a platinum sheet counter electrode (bigger surface), and a resistive heaters for organic and metal layers.\textsuperscript{42} The bilayer devices ITO/HTL(20 nm)/complex(40 nm)/Yb(150 nm) (HTL = TPD, NPD, CBP) were prepared similarly. A commercial ITO on a glass substrate with 5 Ω/□ was used as the anode material (Luminescence Technology Corp.) and commercial Yb, 99.9% trace metals basis (Sigma-Aldrich) as the cathode material. The deposition rate for the TPD and metallo-complexes was 1 nm/s. The active area of the devices was 4×4 mm. The EL spectra in visible region and current-voltage-luminescence characteristics were measured with Ocean Optics USB2000 fluorimeter calibrated with Ocean Optics LS-1 CAL lamp, the computer controlled GW Instek PPE-3323 power supply and GW Instek GDM-8246 digital multimeter. The spectra and power efficiency in the NIR region was determined by Ocean Optics NIR-512 spectrometer. The device characteristics measurements were carried out under ambient conditions.

4. Conclusions

The EL properties of lanthanide pentafluorophenolates were studied for the first time. In the single-layer devices the complexes revealed no EL but insertion of HTL resulted in appearance of the metal-centered emission in the cases of Pr, Nd, Sm, Eu and Yb complexes along with the luminescence of electroluminescent formed at HTL/complex interface which is observed also for the OLEDs based on the Ho and Dy complexes.

Efficiency of the metal-centered luminescence is comparable with that of the devices based on conventional non-fluorinated organo-lanthanide luminophores unlike the enhanced PL of perfluorinated lanthanide phenolates. Unusually strong change in the EL spectra and luminescence color with increasing the applied voltage was found for the bilayer devices based on Tb-complexes. The observed phenomenon is explained by shifting of recombinant zone and respectively the emission from the interfacial HTL/complex electroluminescent compound that causes metal-centered luminescence. Due to these features, the systems may be used for the fabrication of voltage-color tunable organic LEDs. Besides, the observed tendency of lanthanide pentafluorophenolates to form electroluminescent with donor compounds suggests that these complexes may be suitable acceptor materials in design of efficient molecular photovoltaic cells. Further investigations in this direction are in progress.

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Notes and references

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