Improved Current Efficiency in Organic Light-Emitting Devices with Co-Doped Host Layer

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Abstract—A co-doping host layer 4,4’-N,N’-dicarbazole-biphenyl (CBP) and tris-(8-hydroxyquinoline)aluminum (Alq), was incorporated tris (2-Phenylpyridine) iridium [Ir(ppy)3] doped into host as the emitting layer (EML) based organic light-emitting device. Such a structure helps to increase the energy transfer from host to guest resulting in an improved current efficiency. The highest luminous efficiency of the device with 6:4 of CBP: Alq achieved 11.9 cd/A, which is nearly 4.3 times that of the device with 8:2 of CBP: Alq, and the power efficiency achieved 4.69 lm/W, which is nearly 6.7 times that of the device with 8:2 of CBP: Alq.

Keywords—organic light emitting devices; co-doping host layer; current efficiency; power efficiency

I. INTRODUCTION

Organic light-emitting devices (OLEDs) would be probably promising ways to offer the satisfying new display technologies. In the past decade, the architecture of the OLEDs was studied for better power efficiency and device lifetime [1–13]. These emitting layer (EML) structures can be categorized as the uniformly mixed (UM) layer [8,9], step-wise graded mixed (SGM) layer [12] and continuously graded mixed (CGM) layer [9,11,12]. The common feature between these EMLs is that these are basically bipolar transport layers. The main difference between these three EML structures lies in the spatial distribution of the electron-and hole-transport materials. The EML of UM-OLED consists of a constant mixed composition, which is also known as the mixed host (MH) OLED [13]. For SGM-OLED, the EML consists of multilayers with different transport material (HTM): electron transport material (ETM) compositions or ratios, hence it is coiled as a graded layer. For example, when Tris-(8-hydroxyquinoline)aluminum (Alq) acts as host; tris-(2-Phenylpyridine) iridium [Ir(ppy)3] was selected as the green guest; 4,7-diphenyl-1,10-phenanthroline (BPhen) acts as the electron transporting layer; LiF as an electron injecting layer, and Al as cathode electrode.

The routine cleaning procedure, including ultrasonication in acetone, Decon90, and rinsing in deionized water, was firstly carried out to clean ITO glass. The deposition was carried out in a high vacuum condition of about 5×10⁻⁴Pa. Except for host materials, all other materials were deposited simultaneously for the four samples (the same evaporation process). This avoids the uncertainties in comparing devices fabricated with different evaporation processes. Electroluminescence (EL) spectra of the fabricated devices were measured with a PR655 Spectra Scan spectrometer. The luminance–current density–voltage characteristics were recorded simultaneously with the measurement of the EL spectra by combining the spectrometer with a Keithley model 2400 programmable voltage–current source. All measurements were carried out at room temperature under ambient atmosphere without any encapsulation of the OLEDs.

nearly 4 times that of the device with 8:2 of CBP: Alq, and the power efficiency achieved 4.69 lm/W, which is nearly 6.7 times that of the device with 8:2 of CBP: Alq, respectively.

II. EXPERIMENTAL DETAILS

Figure I shows the molecular structure of the materials used in this work, the structure of device. The structure of the device is ITO/m-MTDATA (40 nm)/ NPB (10 nm)/ CBP x%: Alq y%:Ir(ppy)3 8% (20 nm)/ BPhen (50 nm)/ LiF(1 nm)/ Al (150nm) were fabricated, where x: y = 8:2,6:4,4:6,2:8 and the corresponding devices are named A, B, C and D, respectively. In our devices, 4,4’,4”-tris[N-(3-methylphenyl)-N-phenylamin}triphenylamine (m-MTDATA) and N, N’-di(naphthalene-1-yl)-N,N’-diphenyl-benzidine (NPB) are used as hole-injecting layer and hole transport layer; 4,4’-N,N’-dicarbazole-biphenyl (CBP) and tris-(8-hydroxyquinoline)aluminum (Alq) acts as host; tris-(2-Phenylpyridine) iridium [Ir(ppy)3] was selected as the green guest; 4,7-diphenyl-1,10-phenanthroline (BPhen) acts as the electron transporting layer; LiF as an electron injecting layer, and Al as cathode electrode.

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III. RESULTS AND DISCUSSION

Four devices with different ratios of the co-doped host layer were fabricated and tested for this study. Figure 2(a) and Figure 2(b) shows the current density–voltage characteristics and the luminance–voltage characteristics of the devices. The current density of the device is determined mainly by the interfacial energy barrier and charge mobility of the organic materials. A low interfacial energy barrier and high carrier mobility usually give low driving voltage. In particular, interfacial energy barrier influences not only on the driving voltage but also the turn-on voltage of the devices. In other words, the turn-on voltage is closely related to the interfacial properties of the device, while the driving voltage mainly depends on the charge mobility of the material. It is essential to confine the electrons and holes in the light-emitting layer to get high performance. Under the forward bias conditions, the current increased super linearly with increasing applied voltage. It is found that the current density of device A-D at 4V achieves 0.4711mA/cm², 0.5275mA/cm², 0.7045mA/cm², and 6.1288 mA/cm², respectively. The luminance of the devices also showed the same tendency as the current density. The luminance of the device A-D achieves 4.41cd/m², 3.90cd/m², 2.30cd/m², and 21.88 cd/m², respectively. Due to the Alq can blocking hole and more exciton confinement within the EML. It is thought that accumulation of holes occurs at the EML. Therefore, the electric field of EML of OLED with the co-doped host layer is considered to be enhanced compared with the OLED without ones, thus the EL properties are improved. It is also considered that the improvement occurs by a good balance between the injected electrons and holes as Alq owns high electron mobility. It’s well known that co-doped host layer confine holes and electrons effectively in the EML. The improvement seems to be caused by a good balance between the injected electrons and holes when the co-doped host layer with proper doping ratio. The luminance of devices A-D is 4089 cd/m², 10950 cd/m², 15390 cd/m² and 16370 cd/m² at 15V, respectively.
EL spectra of the devices shows main emission peak at 516 nm originating from Irppy3. The spectral variation was found to be different structure of the devices due to the BAlq as electron transport layer can better excitons blocking noticeably affected the performances of the OLEDs fabricated. Figure 3(b) shows CIE coordinates of devices A-D at different voltage. The CIE coordinates were location in green region and essentially voltage-independent.

Figure 4(a) and figure (b) show the current efficiency-voltage and the power efficiency-voltage of the devices. It is obvious that the device B with CBP: Alq (6:4) as co-doped host layer has the highest current efficiency (11.94cd/A) that is about 4.3 times higher than that of the device with CBP: Alq
The highest power efficiency (4.69 lm/W) that is about 6.7 times higher than that of the device with CBP: Alq (8:2) (0.70 lm/W). Table 1 lists the performance of the devices. We can see that the device B with CBP: Alq (6:4) shows the maximum current efficiency and maximum power efficiency. It indicates that the holes and electrons are more balanced in the OLED with an optimum co-doped host layer of around 6:4 than in another three devices. Our results suggest that the optimum ratio of co-doped host layer improve the balance of hole and electron resulting in the improvement of the efficiency. Usually, the mobility of electrons is much lower than holes in common organic materials. This gives rise to an accumulation of excess that of holes at the EML interface. Either increasing the number of electrons or decreasing the number of holes may improve the balance. Obviously, the presence of CPB: Alq can decrease the number of holes, while the number of electrons can be confined in emitting layer. Such a structure helps to increase the energy transfer from host to guest resulting in an improved current efficiency.

The phenomena observed experimentally are consistent with such a speculation, which can be discussed by follows. The luminous efficiency was improved by co-doped host layer method. The host of the emitter layer was composed of CBP and Alq dopant. The Alq dopant did not emit in the emitter layer because their band gaps were wider than that of Ir(ppy)3, and their HOMO and LUMO levels were outside those of Ir(ppy)3. The role of the Alq dopant layer was to supply carriers and excitons must be well confined in emitting layer. The highest luminous efficiency of the device with 8:2 of CBP: Alq, and the power efficiency achieved 4.69 lm/W, which is nearly 6.7 times that of the device with 8:2 of CBP: Alq.

IV. CONCLUSION

In summary, to achieve the high efficient OLEDs, both carriers and excitons must be well confined within the emissive layer. The approach discussed here is to efficiently force carrier recombination and exciton confinement in the EML by the use of co-doped host layer method. By carefully controlling the ratio of co-doped host materials, it may be possible to balanced holes and electrons at EML/ETL interfaces. The result is a simplified structure for high efficiency devices, we have demonstrated a high-efficiency OLEDs with CBP: Alq as co-doped host layer. The highest luminous efficiency of the device with 6:4 of CBP: Alq achieved 11.9 cd/A, which is nearly 4.3 times that of the device with 8:2 of CBP: Alq, and the power efficiency achieved 4.69 lm/W, which is nearly 6.7 times that of the device with 8:2 of CBP: Alq.

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REFERENCES