High efficiency phosphorescent organic light-emitting diodes using carbazole-type triplet exciton blocking layer

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Device performances of green phosphorescent organic light-emitting diodes using (4,4′-N,N′-dicarbazole)biphenyl (CBP) and N,N′-dicarbazolyl-3,5-benzene (mCP) as an exciton blocking layer were investigated. CBP and mCP were introduced between hole transport layer and emitting layer to block triplet exciton quenching and efficient hole transport to emitting layer. The efficiency of green devices could be improved by more than three times by using mCP exciton blocking layer. © 2007 American Institute of Physics. [DOI: 10.1063/1.2742788]

Phosphorescent organic light-emitting diodes (PHOLEDs) are promising as light-emitting materials in OLEDs due to their merit of high efficiency. 100% internal quantum efficiency could be realized in green PHOLEDs Ref. 1 and high power efficiency of 77 lm/W was reported. There have been many studies about light emission and high efficiency in PHOLEDs. Many different device architectures were tried to improve the light-emitting efficiency of PHOLEDs. A hole blocking layer or exciton blocking layer (EBL) was introduced in PHOLEDs to block hole injection from light-emitting layer (EML) to electron transport layer and it was effective to get high efficiency in PHOLEDs. An electron blocking layer was also used in blue PHOLEDs to block electron injection from EML to hole transport layer (HTL). Fac-tris(1-phenylpyrazolato-N,C2′) iridium [Ir(pz)3] with its lowest unoccupied molecular orbital (LUMO) level of 1.7 eV was efficient as an electron blocking material. Other than these, a double EML structure was used and high quantum efficiency of 19.3% was reported. A graded doping structure was studied by our group and it also gave a long lifetime as well as high power efficiency. In this work, we studied the use of (4,4′-N, N′-dicarbazole)biphenyl (CBP) and N,N′-dicarbazolyl-3,5-benzene (mCP) as an EBL to get high efficiency in green PHOLEDs. A detailed mechanism for exciton blocking of CBP and mCP was clarified and device performances were investigated.

The standard device structure used in this experiment was indium tin oxide (150 nm)/N,N′-diphenyl-N,N′-bis-[4-(phenyl-m-tolyl-amin o)-phenyl]-biphenyl-4, 4′-diamine (60 nm)/N, N′-di-(1-naphthyl)-N, N′-diphenylbenzidine (NPB) (30 nm)/PH1: tris(2-phenylpyridine) iridium [Ir(ppy)3] (30 nm, 5% doping)/biphenoxy-bi(8-hydroxy-3-methylquinoline) aluminum (5 nm)/tris(8-hydroxyquinoline) aluminum (20 nm)/LiF (1 nm)/Al (200 nm). Three devices with different HTL structures were fabricated to investigate the effect of CBP and mCP EBLs on device performances. Device I had NPB (30 nm) as a HTL and device II had both NPB (20 nm) and CBP (10 nm) as a double layer HTL, while device III had NPB (20 nm) and mCP (10 nm) as a double layer HTL. PH1 was supplied from Merck Co. and it has a spirobifluorene-type backbone structure with high electron transport properties because of spirobifluorene units. The triplet band gap of PH1 was 2.4 eV and the highest occupied molecular orbital (HOMO) and LUMO were 5.9 and 2.8 eV, respectively. The current density–voltage–luminance characteristics of the devices were measured with Keithley 2400 source measurement unit and PR 650 spectrophotometer.

It is important to confine excitons in EML to increase the recombination efficiency of holes and electrons in OLEDs by blocking charge carriers and exciton diffusion out of EML. Hole and exciton blocking in PHOLED was effective in electron transport layer side and electron and exciton blocking in HTL side is expected to give high recombination efficiency through charge and exciton confinement inside EML. Exciton blocking materials should have good hole transport properties to transport holes from hole injection layer to EML, and it is required to have a wide triplet band gap to block exciton diffusion and quenching. In addition, it should have electron blocking function to prevent electron overflow to HTL. Therefore, CBP and mCP were chosen as an exciton blocking material in this work and their role as an EBL was investigated.

Figure 1 shows current density–voltage–luminance characteristics of green devices with CBP and mCP EBL compared with standard devices. The current densities of device I with NPB as a HTL and device II with NPB and CBP double layer as a HTL were similar, while device III with NPB and mCP as a HTL was rather low compared with that of devices I and II. The similar current density in devices I and II is due to similar energy barrier between organic layers. Energy level diagrams of devices II and III are schematically described as an inset in Fig. 1. The energy barrier for hole injection from NPB to CBP is 0.4 eV and there is no energy barrier for hole injection from CBP to PH1. Device I also has 0.4 eV HOMO difference between NPB and PH1, and the hole injection barrier between HTL and EML in device I is...
the same as that of device II. Therefore, hole injection in devices I and II can be similar, leading to similar current density in the device. Compared with device II, device III has a large energy barrier of 0.6 eV between NPB and mCP, resulting in low current density even though there is no energy barrier between mCP and PH1. Luminance-voltage characteristics of three devices are also shown in Fig. 1. The luminance of device II with CBP EBL was higher than that of other devices, and device III showed the lowest luminance value at the same driving voltage. Considering similar current densities of devices I and II, the high luminance of device II compared with device I indicates that exciton formation and light emission are efficient in device II.

Quantum efficiency–luminance of three devices was plotted in Fig. 2 based on current density and luminance of devices. Device III with mCP as an EBL showed the best quantum efficiency value of 14% and device II with CBP EBL also exhibited high efficiency of 11.9% compared with 3.4% of device I at 1000 cd/m². As expected from energy level diagram in Fig. 1, mCP can play a role of an electron blocking layer as well as an EBL. Electrons can be blocked by mCP and triplet exciton quenching can be reduced by mCP because of the wide triplet band gap of mCP (2.9 eV), resulting in an efficient recombination of holes and electrons inside the emitting layer. Even though NPB is efficient as an electron blocking layer due to the high LUMO energy barrier of 0.4 eV between NPB and PH1, the triplet energy level of NPB is only 2.3 eV, which is not high enough for exciton blocking from EML because the triplet energy level of host and dopant materials is 2.4 eV. Compared with NPB, mCP has a triplet energy level of 2.9 eV (Ref. 5) and it can block triplet exciton quenching. CBP can also act as an EBL due to a wide triplet band gap of 2.6 eV even though it is not so efficient as mCP as an electron blocking layer. The exciton blocking function of mCP was confirmed from photoluminescence (PL) spectra. Figure 3 shows PL spectra of standard device and device with carbazole-type EBL. PL intensity was high in the device with CBP and mCP EBL, indirectly proving that CBP and mCP reduce triplet exciton quenching by NPB. Even though mCP has a wider triplet band gap than CBP, PL intensity was quite similar to each other. This result indicates that mCP and CBP show similar performances as a triplet EBL. Considering similar exciton blocking effect but different efficiency in CBP and mCP devices, the high efficiency in mCP device might be due to good hole-electron balance in emitting layer which originated from less hole flow in mCP device.

Figure 4 shows electroluminescence spectra of three devices. Green emission with peak position of 515 nm was observed in all devices which originated from Ir(ppy)₃ triplet emission. In addition to green emission, device I also showed another broad peak at 458 nm which is assigned to NPB emission, while devices II and III exhibited no emission in blue wavelength range. The NPB emission in device I may be due to hole accumulation at the interface between NPB and PH1 and electron overflow from PH1 to NPB. In general, holes are strongly trapped by dopant materials in Ir(ppy)₃ doped devices and holes can be accumulated between NPB and PH1 because hole transport through Ir(ppy)₃ is limited. The accumulated holes can recombine with some electrons injected from EML inside NPB, giving rise to NPB...
emission. Compared with device I with blue emission, no blue emission in devices II and III is due to the reduced hole accumulation at the interface between HTL and EML. Holes are injected from CBP or mCP layer and can be transported through PH1 host materials. Therefore, hole accumulation can be greatly reduced and most of the holes can recombine with electrons in EML, leading to efficient green emission. No blue emission in PH1 device with CBP EBL supports this explanation. The LUMO level of CBP is only 2.6 eV and electrons can be efficiently injected from PH1 to the CBP and NPB layer, which will induce NPB blue emission through recombination with accumulated holes at the interface. However, NPB emission was not observed in the CBP device, which supports the fact that efficient hole transport dominates light emission in PH1 device with CBP or mCP EBL. It can be inferred from electroluminescence spectra that the recombination zone of holes and electrons was shifted from HTL side to EML by EBL, considering the disappearance of blue emission in devices II and III.

In summary, the use of CBP and mCP as an excition blocking layer in green PHOLEDs was effective to get high luminance efficiency, and quantum efficiency was enhanced from 3.4% to 14.0% by mCP EBL. Both CBP and mCP were efficient as an EBL, and there was no emission from NPB in the device with CBP and mCP EBL.

![Graph of electroluminescence spectra](image)

**FIG. 4.** Electroluminescence spectra of green phosphorescent devices with different hole transport layers.