2,7(3,6)-Diaryl(arylamino)-substituted Carbazoles as Components of OLEDs: A Review of the Last Decade

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Abstract: Organic light emitting diode (OLED) is a new, promising technology in the field of lighting and display applications due to the advantages offered by its organic electroactive derivatives over inorganic materials. OLEDs have prompted a great deal of investigations within academia as well as in industry because of their potential applications. The electroactive layers of OLEDs can be fabricated from low molecular weight derivatives by vapor deposition or from polymers by spin coating from their solution. Among the low-molar-mass compounds under investigation in this field, carbazole-based materials have been studied at length for their useful chemical and electronic characteristics. The carbazole is an electron-rich heterocyclic compound, whose structure can be easily modified by rather simple reactions in order to obtain 2,7(3,6)-diaryl(arylamino)-substituted carbazoles. The substituted derivatives are widely used for the formation of OLEDs due to their good charge carrier injection and transfer characteristics, electroluminescence, thermally activated delayed fluorescence, improved thermal and morphological stability as well as their thin film forming characteristics. On the other hand, relatively high triplet energies of some substituted carbazole-based compounds make them useful components as host materials even for wide bandgap triplet emitters. The present review focuses on 2,7(3,6)-diaryl(arylamino)-substituted carbazoles, which were described in the last decade and were applied as charge-transporting layers, fluorescent and phosphorescent emitters as well as host materials for OLED devices.

Keywords: substituted carbazole; amorphous material; charge transporting layer; emitter; host material; organic light emitting diode

1. Introduction

Since the first discovery of organic electroluminescent compounds, huge attention has been devoted to the creation of new materials and optimized multilayer device architectures for viable and practical organic light emitting diodes (OLEDs), which would demonstrate low-driving voltage, high brightness, full-color emission, a long lifetime, and the easy formation of flexible thin-film devices. The commercial benefit of OLED devices is seen in the emergence of digital cameras, mobile cell phones as well as display technologies; however, there is still a strong demand to considerably improve the performance and lifetime of the devices for lighting technologies [1–5].

Formerly, structurally and chemically well-defined, carbazole-based compounds and their characteristics were investigated due to their properties of biological activity [6–8]. However, in the last three decades, the carbazole rings or substituted carbazole-containing compounds have been widely investigated in the field of organic optoelectronics, particularly in OLED technologies [9–12]. The substituted carbazole-based derivatives are widely used as components of the devices due to their good charge injection and transport characteristics in the layers of the materials [13], electroluminescence [14], thermally activated delayed fluorescence [15], improved thermal and morphological stability as well as their homogenous thin film formation properties [16]. In addition, the rather high triplet
energies of some aryl(arylamino)-substituted derivatives make them useful components, acting as host materials for the phosphorescent triplet emitters in electrophosphorescent OLED technologies [17].

The 9H-carbazole core as electron donor (D) can be easily substituted at the 9,2,7 (3,6)-positions by using different aromatic fragments. Donor–acceptor (D-A) derivatives (9-arylcarbazoles), asymmetric 9,2(3)-diarylcarbazoles as well as symmetric 2,7(3,6)-diarylcarbazoles could be synthesized by using various arylation or amination reactions. D-D fragments containing derivatives are usually used as hole-transporting layer materials or fluorescent emitters [18–20]. The D-A and A-D-A structures that have carbazoles are more suitable as host materials for phosphorescent triplet emitters, thermally activated delayed fluorescence (TADF)-based emitters as well as in other less investigated fields such as exciplex components, intramolecular hosts, dopants, etc. [21]. D-A structure derivatives and asymmetric 9,2(3)-diaryl(diarylamino)carbazoles are rather widely described in earlier review articles [22,23]. In the present review, we analyze and present only symmetric 2,7(3,6)-diaryl(arylamino)-substituted carbazoles, which were developed and described in last decade and were applied as charge-transporting layers, fluorescent and phosphorescent emitters as well as host materials for the OLED devices.

2. Synthesis of 2,7(3,6)-Diiodo(dibromo)carbazoles and 2,7(3,6)-Diaryl(diarylamino)-substituted Objective Carbazoles

Both the 3,6-Diiodo-9H-carbazoles and 3,6(bromo)-9H-carbazoles are intermediate materials for the preparation of objective carbazole-based derivatives for OLEDs. Structures of the halogenated derivatives are shown in Scheme 1. The bromination reaction of 9H-carbazole with N-bromosuccinimide (NBS) [24] or its Tucker iodination reaction [25] correspondingly yields 3,6-dibromocarbazole or 3,6-diiodocarbazole. On the other hand, 2,7-dibromocarbazole is obtained by a two-step synthesis, as shown in the Scheme 1 [26]. The starting material 4,4′-dibromo-1,1′-biphenyl is firstly nitrified. The obtained 4,4′-dibromo-2-nitro-1,1′-biphenyl is then reacted with triethylphosphate to get 2,7-dibromo-9H-carbazole. The nitrogen atom of the halogenated carbazoles can be then functionalized by different alkyl or aryl groups in order to obtain the key starting materials: 3,6-diiodo-9-alkyl(aryl)carbazoles (36DICr), 3,6-diiodo-9-arylcarbazoles (36DI9ArCr), 3,6-dibromo-9-alkyl(aryl)carbazoles (36DBrCr), 3,6-dibromo-9-arylcarbazoles (36DBr9ArCr), 2,7-dibromo-9-alkyl(aryl)carbazoles (27DBrCr), and 2,7-dibromo-9-arylcarbazoles (27DBr9ArCr), which are shown in the Scheme 1. The alkylation reactions with bromo or iodo alkanes under basic conditions are rather simple and widely described in the literature [27–29]. The di-halogenated 9-arylcarbazoles are prepared by Ullmann or Buchwald–Hartwig reactions [30–32].

The key starting di-halogenated compounds 36DICr, 36DI9ArCr, 36DBrCr, 36DBr9ArCr, 27DBrCr, and 27DBr9ArCr can then be used in the following amination or C-C coupling reactions to obtain the target compounds: 2,7-diarylamino-9-alkyl(aryl)carbazoles (27DNCr and 27DNArCr), 3,6-diarylamino-9-alkyl(aryl)carbazoles (36DNCr and 36DNArCr), 2,7-diarylamino-9-alkyl (aryl)carbazoles (27DCr and 27DArCr), or 3,6-diarylamino-9-alkyl(aryl)carbazoles (36DCr and 36DArCr) (Scheme 2). Most of the objective carbazole-based derivatives, which are used later as electroactive components for the production of OLED devices, are obtained from the di-halogenated carbazoles by the Ullmann, Suzuki, Stille, or Buchwald–Hartwig reactions.
Scheme 1. Synthesis of starting compounds: 3,6-diiodo(dibromo)-9-alkyl(aryl)carbazoles and 2,7-dibromo-9-alkyl(aryl)carbazoles.

Scheme 2. Synthetic pathway of the 2,7(3,6)-diaryl(arylamino)-substituted objective 9-alkyl(aryl)carbazoles.

The Ullmann reaction is a coupling reaction between aryl halides and aromatic amines or aromatic heterocyclic compounds in the presence of copper-based catalysts. The mechanism of the Ullmann reaction was extensively studied for many years. Complications arise because the reactions are often heterogeneous, especially those catalyzed with metallic copper. Suzuki cross-coupling procedures, developed by Nobel laureate Akira Suzuki, are among the most widely investigated reactions in the formation of carbon–carbon bonds in aromatic compounds. These reactions were generally catalyzed by inorganic catalysts, such as soluble palladium (Pd) complexes having various ligands, and more recently, also in aqueous media. The Buchwald–Hartwig amination procedure is a chemical reaction used in organic chemistry for the formation of carbon–nitrogen bonds via palladium-catalyzed coupling reactions of aromatic amines with aryl halides. Other reactions as the Stille coupling, Diels Alder, and Friedel–Craft have also been used to obtain the target carbazole compounds, but in rarer cases. In the general...
case, diarylamino carbazoles (27DNCr, 27DNArCr, 36DNCr, and 36DNArCr) are usually obtained by Ullmann or Buchwald–Hartwig reactions. On the other hand, diaryl carbazoles (27DCr, 27DArCr, 36DCr, and 36DArCr) are usually prepared by Suzuki or Stille reactions, as presented in Scheme 2.

3. Diaryl(diarylamino)-substituted Carbazoles as Charge-Transporting Layer Materials for OLEDs

Structures of the diaryl-substituted compounds (Cx), which were used to form the hole-transporting layers (HTLs) in OLED devices, are shown in Scheme 3. The target compounds C1–C14 [44–48] and C17–C26 [49–54] were obtained under the conditions of the Suzuki reaction. Compounds C15–C16 were generated by the Diels–Alder reaction between conjugated diene and substituted alkene, forming the substituted cyclohexene fragments [55]. Authors of the research studied properties of the derivatives and used these materials for the formation of HTLs in OLED devices. Thermal properties were examined for compounds C1–C13, C17–C19, C21–C22, and C24–C26. The reported material C22 demonstrated the highest thermal stability in this group, with a very high thermal decomposition temperature (T_d) of 575 °C, as well as the highest glass transition temperature (T_g) of 260 °C. The values of ionization potentials (I_p) for compounds C9–C11, C17–C20, and C23 were 5.65 eV, 5.55 eV, 5.8 eV, 5.38 eV, 5.42 eV, 5.19 eV, and 5.50 eV, respectively, and confirmed suitable hole-injecting properties for thin layers of many of the materials. HOMO and LUMO levels of the materials C1–C8, C13–C17, C19–C22, and C24–C26 are different and depend on the nature of the substituents. The HOMO level of the diarylcarbazoles varied between −4.93 and −6.02 eV, and the LUMO level was in a broad range between −0.87 and −2.93 eV due to different electron-withdrawing or donating substituents at the carbazole core.

Scheme 3. Structures of 3,6-diarylcrazoles used for hole-transporting films in OLEDs.
Positive charge drift mobility ($\mu_h$) in the thin layers of the derivatives C12, C17–C19, C21–C22, and C25–C26 were reported. The compounds demonstrated rather high hole-drift mobility in their amorphous films ranging from $5 \times 10^{-5}$ cm$^2$/V$^{-1}$s$^{-1}$ to $1.5 \times 10^{-4}$ cm$^2$/V$^{-1}$s$^{-1}$ at high electrical fields. The charge-injecting/transporting properties of these materials confirmed that they are suitable hole-transporting layer materials in OLEDs.

Agarwal fabricated the single-layer spin-coated OLEDs, in which the derivatives C1-C8 were sandwiched between indium tin oxide (ITO) or ITO/PEDOT:PSS [poly(3,4-ethylenedioxythiophene): poly(styrenesulfonate)] as the anode and a calcium (Ca) cathode, which was protected by a thick layer of Al. Multilayer OLEDs with the materials C1–C8 were also fabricated. The structure of the devices was ITO/F$_2$TCNQ (2,3,5,6-tetrafluoro-7,7′,8,8′-tetracyano-p-quinodimethane)/TPD (N,N′-(bis(3-methylphenyl)-1,1′-biphenyl-4,4′-diamine))/CBP (N,N′-dicarbazoloyl-4,4′-biphenyl)/C3 or C4/BCP (4,7-diphenyl-1,10-phenanthroline)/LiF/Al. Brightness as high as 900–1000 cd/m$^2$ was achieved in the multilayer OLED devices using the HTLs of C3 or C4. Green electro-phosphorescent devices ITO/PEDOT:PSS/C9 or C11/CBP/Ir(ppy)$_3$ (tris(2-phenylpyridine(ridium(III)))/TPBi (2,2′, 2″-(1,3,5-benzinetriyl)-tris(1-phenyl-1-H-benzimidazole))/LiF/Al were also presented. The device with C11 demonstrated the best results, with a turn-on voltage of 5 V, a maximum brightness of 9800 cd/m$^2$, and a maximum current efficiency of 22.5 cd/A. C12 was tested as an electron-confining HTL for phosphorescent OLEDs with a red, green, or blue emitting layer. The power efficiency of the studied red device was increased from 8.5 to 13.5 lm/W, an increment of 59%, and the maximum luminance was enhanced from 13,000 to 19,000 cd/m$^2$, an increment of 46%, as compared with an analogous device using commercial HTL material. For the blue device using C12, the power efficiency was increased from 6.9 to 8.9 lm/W, an increment of 29%, and the maximum brightness was enhanced from 9000 to 11,000 cd m$^{-2}$, an increment of 22%. C13 as an HTL was investigated by Kochapradist et al. Tris(8-hydroxyquinolinato)aluminum (Alq3) emitter-based green OLEDs, with the structure of ITO/PEDOT:PSS/C13/Alq3/LiF:Al, were fabricated. The material C13 showed excellent hole-transporting properties for the green device—a luminance (current) efficiency exceeding 5 cd/A was achieved. Lee and co-workers studied the hole-transporting properties of C14 and formed the device with ITO / CuPc (copper(II)phthalocyanine)/C14/Alq3/LiF/Al. For an objective comparison of electroluminescent properties, a reference device was fabricated and NPB was used as the HTL material. The device with the compound C14 showed a high power efficiency of 1.67 lm/W and a current efficiency of 6.12 cd/A, which were higher as compared with the NPB-based device. OLEDs with the structures ITO/2-TNATA (4,4′,4′′-tris[N-(2-naphthyl)-N-phenylamino]triphenylamine)/C15 or C16/Alq3/LiF/Al were fabricated by Park and co-workers. The devices showed pure performance, with their luminance efficiency exceeding only 3 cd/A and a power efficiency exceeding 1 lm/W.

In order to identify the charge-transporting properties of C17, multilayer OLED devices were fabricated by Kim et al. The green device ITO/CuPc/C17/Alq3/LiF /Al showed a current efficiency of 5.16 cd/A and a power efficiency of 2.35 lm/W. Braveenth and co-authors studied the properties of the compounds C18–C19 as hole-transporting materials. Device configuration was ITO/DNTPD (4,4′-bis[N-[4-[N,N-bis(3-methylbenz-1-phenylamino)biphenyl]]/C18 or C19 /Bebz$_2$ (bis[10-hydroxybenzo[h]quinolino]beryllium): Ir(mphmq)$_2$(tmq) (bis[2,4-dimethyl-6-(4-methyl-2-quinolinolyl)phenyl][2,2,6,6-tetramethyl-3,5-heptanedionate]/Bebz$_2$/LiF /Al. A C18-based phosphorescent device exhibited higher maximum current efficiency (24.6 cd/A) and higher maximum external quantum efficiency (23.2%) as compared with a C18-based OLED. In the double-layer device, ITO/C20/Alq3/LiF/Al, in which C20 was used as the hole-transporting material, a yellowish-green color arising from the Alq3 was observed. It was only reported that the luminance at the applied voltage of 10 V was about 9600 cd/m$^2$. Devices with the structures of ITO/C21/TPBi/LiF/Al and ITO/NPB/C21/TPBi /LiF/Al were fabricated to evaluate the properties of C21 as a
hole-transporting light-emitting layer. A quantum efficiency of 2.21% was obtained for the C21-based device having an additional NPB layer.

Kumar and co-workers investigated the hole-transporting properties of C25 and C26 for fluorescent OLEDs ITO/PEDOT:PSS/NPB/C25 or C26/Alq3/LiF/Al, as well as for phosphorescent devices ITO/PEDOT:PSS/NPB/C25 or C26/Ir(ppy)3/CBP/TPBi/LiF/Al. At 1000 cd/m², the fluorescent green device with C26 showed a current efficiency of 4.0 cd/A, which was about 135% higher than that of the typical HTM NPB-based device. Green phosphorescent devices with C25 showed a high current efficiency of 58.4 cd/A (power efficiency 54.8 lm/W and EQE 16.1%), while 45.1 cd/A (power efficiency 40.8 lm/W and EQE 12.5%) was measured in the C26-containing device.

Diarylamino-substituted carbazoles, which were used as HTLs in OLEDs, are shown in Scheme 4. The target compounds C27–C42 were all obtained by Ullmann amination reactions [56–62]. C30–C42 demonstrated sufficient and high thermal stability, with the thermal degradation temperatures (T_d) ranging from 263 °C to 490 °C. The derivatives C30–C42 are also suitable for glass formation, having a T_g in the range of 42–217 °C. The Ip of the compounds C30–C31 and C37–C42 were reported and demonstrated the values of 5.80 eV, 5.81 eV, 5.50 eV, 5.24 eV, 5.43 eV, 5.67 eV, 5.28 eV, and 5.34 eV, respectively. HOMO and LUMO energy levels were reported for the materials C27–C30. The values of HOMO/LUMO for the derivatives C27–C30 were −4.92/−1.91 eV, −4.92/−2.24 eV, −4.92/−2.33 eV, and −4.83/−1.87 eV, respectively.

Scheme 4. Structures of diarylamino-substituted carbazoles used for HTL formation.

The charge-transporting properties of the derivatives C27–C28 and C31–C42 were studied with the time-of-flight technique. The μ_h values ranged from 1.2 × 10⁻⁸ to 2 × 10⁻³ cm²·V⁻¹·s⁻¹ at high electric fields. The μ_h in the films of C32 and C40 dispersed in bisphenol Z polycarbonate (PC-Z) reached 10⁻³ cm²·V⁻¹·s⁻¹ at high electrical fields. The charge injection/transport characteristics of some from the derivatives C27–C28 and C31–C42 showed that they are suitable as hole-transporting layers for OLED devices.

Shen and co-authors investigated two types of devices. The structure of device I was ITO/C27–C30/TPBi/Mg:Ag, and that of device II was ITO/C27–C30/(Alq3)/Mg:Ag. In type I, the derivatives functioned as both hole-transporting and emitting materials. In the type II devices, light was emitted from either the disubstituted carbazole layers or from Alq3. Device II with C30 reached a maximum external quantum efficiency of 1.3%. 
4. Diaryl(arylamino)-substituted Carbazoles as Host Materials for PhOLEDs

The structures of the diaryl-substituted carbazoles, which were used as host materials for PhOLED devices, are shown in Scheme 5. The compounds 36DICr, 36DBrCr, and 36DBr9ArCr were firstly obtained for their synthesis as key starting materials, as shown in Scheme 1. The objective compounds H1–H4 [63–65], H6–H7 [66], H10 [67], and H12–H14 [68,69] were obtained during the Suzuki reaction. The diphosphate oxide-based material H5 was obtained by the oxidation of the corresponding diphosphate, which was prepared by a lithium–halogen exchange reaction between the di-brominated aromatic bridge and n-butyl lithium, followed by a reaction with chlorodiphenylphosphine [70]. Compound H8 was obtained when 9-phenyl-9-carbazole-3,6-dicarboxylic acid reacted with SOCl2, and then the resulting compound was treated with trimethylamine and 2-aminodiphenylamine. Derivative H9 resulted from the reaction of 9-phenyl-3,6-di(2H-tetrazol-5-yl)-9H-carbazole with the o-xylene solution containing benzoyl chloride [71]. H11 was synthesized by the coupling reaction of chlorodiphenylphosphine with 3,6-dibromo-N-phenylcarbazole [72]. The reaction of 9-(4-tert-butylphenyl)carbazole with triphenylmethanol in the presence of Eaton’s reagent yielded the objective derivative H15 [73].

Scheme 5. Chemical structures of diarylcarbazoles used as host materials for PhOLEDs.

The destruction temperatures (Td) of the compounds H1–H4, H6–H10, and H12–H15 were reported. The compound H7 demonstrated a rather low thermal destruction temperature of 186 °C, while the temperatures of the other compounds reached 300 °C. All the compounds H1–H15 were obtained as amorphous substances and could form amorphous glasses with Tg values ranging from 30 °C to 269 °C. The HOMO and LUMO energy levels of H1–H15 were reported. The HOMO/LUMO energies of the derivatives H1–H15 were different and depended on substituents at the 3,6 positions of the carbazole core. The HOMO level was between −5.21 and −6.48 eV, and the LUMO level varied between −1.90 and −2.83 eV. The values of µh in the thin films of the prepared materials H3
and H15 were investigated. The amorphous layers of these materials showed very similar results, and the charge drift mobility reached $5 \times 10^{-4} \text{ cm}^2\cdot\text{V}^{-1}\cdot\text{s}^{-1}$ at high electrical fields.

Kim et al. investigated the phosphorescent devices ITO/NPD/H1 or H2: Ir(ppy)$_3$/Alq$_3$/LiF/Al. A maximum brightness of 4165 cd/m$^2$, low current efficiency of 1.58 cd/A, and also a low external quantum efficiency of 0.59% were achieved in the group of the most efficient device with the host H1 [74]. The PhOLEDs ITO/PEDOT/H3:Ir(ppy)$_3$/TPBi/LiF/Al containing the green emitter Ir(ppy)$_3$ in hosts H3 using wet and dry processes as well as dry process, with the incorporation of an additional TAPC hole-transporting layer, were fabricated by Jou and co-workers. The efficiency of the best device was reported as 11.6 lm/W (21 cd/A). The host material H4 was used in a solution processable multilayer device ITO/PEDOT/H4:Ir(ppy)$_3$/TPBi/LiF/Al. Its maximum photometric efficiency was 35.5 cd/A, and the maximum EQE exceeded 10.6%. Sapochak et al. investigated the blue devices ITO/TCTA/H5: Flrpic/DPDT (2,8-bis(diphenylphosphoryl) dibenzob[d]thiophene)/LiF/Al and obtained an EQE of 7% at a voltage of about 7 V [75]. The simple-structured blue PhOLEDs with hosts H6 and H7 and Flrpic dopant correspondingly reached a maximum efficiency of 43.9 cd/A and 46.1 cd/A. Sky blue PhOLEDs were fabricated using either Flrpic or (DFPPM)$_2$Irpic as phosphorescent emitters co-evaporated with the hosts H8 or H9 in the devices. The host material H9 gave excellent OLED characteristics, with an EQE of 17.7% for green and 20.6% for the red device. Jiang et al. investigated the host H10 and reported that the device with Flrpic as a dopant reached a maximum current efficiency of 7.1 cd/A. Ahn and co-workers formed the device ITO/HATCN (1,4,5,8,9,11-hexaazatriphenylehexacarbonitrile)/TAPC/(3,5-di(carbazol-9-yl)-N,N-diphenylaniline) DCDDP/H11:TSPO1 (diphenyl-4-tribenylylphenyl-phosphine oxide)/TPBi/LiF/Al and found that the multilayer PhOLED shows a very high external quantum efficiency of 27.8% [76]. A green PhOLED with the structure ITO/2-TNATA/NPB/H12/TPBi/LiF/Al was fabricated by Song et al. The device showed that its maximum luminescence efficiency could exceed 29 cd/A and its EQE could reach 8.68%. The double-layer solution-processed blue PhOLEDs ITO/PEDOT:PSS/PVK/H13 or H14: Flrpic/Ba/Al were fabricated. The best results yielded a device with an H14 host-maximal current efficiency of 4.16 cd/A. Tsai et al. created the device ITO/PEDT:PSS/DPAS (2,2′-bis(N,N-diphenylamine)-9,9′-spirobifluorene)/TCTA/H15:Flrpic/TAZ (3-(4-biphenylyl)-4-phenyl-5-(4-tert-butylphenyl)-1,2,4-triazole)/LiF/Al. The efficiency of the H15-based device reached 7% at 100 cd/m$^2$.

The structures of the diarylamino-substituted carbazoles, which were used as host materials for PhOLED devices, are shown in Scheme 6. The objective compounds H16–H17 [77,78], H22–H23 [79], and H24 [80] were obtained by Ullmann reaction. The compounds H18–H21 [81] and H25 [57] were prepared by the Buchwald–Hartwig amination procedure. TGA confirmed that the compounds are highly and thermally stable. The $T_d$ values of the derivatives H16–H23 were 419 °C, 384 °C, 364 °C, 389 °C, 399 °C, 417 °C, 335 °C, and 404 °C, respectively. The $T_g$ values of the materials H16–H23 ranged from 81 °C to 210 °C. $I_p$ was reported for some of the derivatives. For example, the phenothiazine-substituted compound H16 had a lower $I_p$ of 5.25 eV as compared with the carbazole-based compound H17 (5.57 eV). The described HOMO and LUMO energies of the compounds H17 and H19–H25 had different values and depended considerably on arylamino fragments. The energy levels of these derivatives ranged from $-4.86$ to $-5.62$ eV for HOMO and from $-1.78$ to $-2.74$ eV for LUMO.
Bagdziu纳斯等 fabricated three PhOLEDs in which H16 was used as host material. The structures of the devices were ITO/H16:Ir(ppy)$_3$/Bphen (4,7-diphenyl-1,10-phenanthroline)/Ca:Al or ITO/m-MTDATA/H16 :Ir(ppy)$_3$/Bphen/Ca:Al or ITO/m-MTDATA/H16 :Ir(piq)$_3$(acac)/Bphen/Ca:Al. The maximum power and EQE up to 47.5/29.6 lm/W and 20.0/10.5% for green or red PhOLED devices, respectively, were reported in the most effective device. The structure ITO/PEDOT:PSS/H17: Ir(Fppy)$_3$/LiF/Al was fabricated by Pudzs et al. [82]; however, the un-optimized OLED reached very low efficiencies of only 0.07 lm/W and 0.45 cd/A. The reported optimized phosphorescent device with the host H20 material ITO/PEDOT:PSS/TAPC/ H20:Flrpic/TmPyPB/LiF/Al [83] had a maximum current efficiency of 36 cd/A at 200 cd/m$^2$. A device stack with the carbazole-based host H22 doped with an orange phosphor tris(2-phenylquinoline)iridium (III) [Ir(2-phen)$_3$] demonstrated improved efficiency (7.4% and 16 lm/W). Most importantly, the superior stability of the device using H22 enabled a lifetime well above 10,000 h at a practical luminance of 1000 cd/m$^2$ [84]. PhOLED devices ITO/PEDOT:PSS/TAPC/H24 or H25: Ir(ppy)$_3$ /BmPyPb/LiF/Al were fabricated by Park and Lee. The green device with H24 showed the best performance, with a maximal EQE of 21.3%.

5. Diaryl(arylamino)-substituted Carbazoles as Fluorescent Emitters of OLEDs

Structures of the 2,7-diaryl- or 3,6-diaryl-substituted carbazoles, which were used as emitters (E) in the fluorescent OLED devices, are shown in Scheme 7. All the target diaryl-substituted compounds E1–E13 [85–93] and E15–E20 [94–97], except for E14, were obtained during the Suzuki reaction of the di-halogenated carbazoles with the corresponding boronic acids or their esters. The objective material E14 was synthesized by fusing triphenylethylene and one dimesitylboron group moiety into a system, with one attached to the carbazole fragment [98].
Scheme 7. Structures of 2,7(3,6)-diarylcarbazole-based emitters.

The thermal properties of some of the emitters were investigated [99]. It was described that the compounds E1–E7, E9–E10, E14, and E17–E18 exhibited rather high thermal stability, with a $T_d$ value ranging from 283 to 603 °C. Meanwhile, the $T_d$ values of the compounds E15 and E16 were lower and reached 219 °C and 188 °C, respectively. It was mentioned that the combination of fragments of carbazole with various conjugated aromatic fragments is favorable for the formation of homogeneous thin films with high $T_g$ temperature values. The $T_g$ of E1, E3–E6, E9–E10, E14, E17, and E18 were 105, 138, 65, 138, 77, 70, 211, 88, 170, and 172 °C, respectively. The values of $I_p$ for the derivatives E1–E2 and E19–E20 were 5.70 eV, 5.56 eV, 5.17 eV, and 5.30 eV, respectively. It was evident that the attached fragments at the 2,7- and 3,6-positions of the carbazole core provided an effective tool for the HOMO and LUMO of the compounds. The HOMO and LUMO of E2–E7 and E9–E20 ranged from $-4.43$ to $-5.66$ and from $-2.61$ to $-3.01$ eV, respectively. Hole drift mobilities in the amorphous layer of the compounds E17–E18 were reported, and the values varied from $1 \times 10^{-5}$ to $3 \times 10^{-8}$ cm$^2$·V$^{-1}$·s$^{-1}$ at a high electric field.
The device ITO/PEDOT:PSS/CBP: E2/TPBi/LiF/Al was fabricated by Konidena and co-workers. This OLED reached a power efficiency of only 0.5 lm/W and a current efficiency of 1.1 cd/A. The same structure device just using the emitter E5 reached a power efficiency of 3.9 lm/W and a current efficiency of 4.9 cd/A, while the same device with E7 reached a power efficiency of 4.12 lm/W and a current efficiency of 9.52 cd/A. Dang and co-workers experimented with the device ITO/HATCN (dipyrazino[2,3-f’:2’,3’-h]quinoxaline-2,3,6,7,10,11hexacarbonitrile)/NPB/TCTA/E3 or E4/TPBi/LiF/Al. The material E4, with a much balanced molecular conjugation and a twisted molecular conformation, exhibited much better performance in non-doped OLEDs with maximum photometric efficiency, power efficiency, and EQE values of 7.38 cd/A, 6.81 lm/W, and 3.0%, respectively. The simple-structured ITO/E8/TPBi/Mg:Al demonstrated a power efficiency of 2.55 lm/W and a current efficiency of 4.75 cd/A. OLEDs with the structure ITO/PEDOT:PSS/E9 or E10/NPB/CPB/LiF/Al were characterized by a power efficiency of 1.68 or 2.67 lm/W and a current efficiency of 3.55 or 5.89 cd/A, respectively. Feng et al. studied the properties of the device ITO/MoO3/TAPC/TCTA/E11/TPBi/LiF/Al. The designed emitter showed a bright violet light at a wavelength of 400 nm, with a current efficiency of 0.65 cd/A and an EQE of about 2%. To examine the electroluminescent properties of the emitters E12 and E13, researchers fabricated a device by solution processing with the following configuration: ITO/PEDOT:PSS/ NPB/E12 or E13/BCP/Alq3/LiF/Al. The white light emission from the device was stable in the range from 5 V up to 20 V. The maximal luminance values were found to be 20 cd/m² and 110 cd/m², respectively, for the E12 and E13-based devices. Multilayer, non-doped OLEDs, with a configuration of ITO/HATCN /NPB/E14, E15 or E16/TPBi/LiF/Al, were fabricated. They exhibited good performance with low turn-on voltages of 4.2 V, 6.2 V, and 6.0 V, high maximal luminance values of 13,930 cd/m², 21,054 cd/m², and 4376 cd/m², and maximum current efficiency values of 4.74 cd/A, 3.34 cd/A, and 2.34 cd/A, respectively. A three-layer PhOLED ITO/PEDOT:PSS/E17:[(pbi)2Ir(acac)]/TPBi/LiF/Al was fabricated by Hung and co-authors. The maximum value for energetic efficiency of 62 lm/W and 62 cd/A for current efficiency were achieved from the system at a brightness of 120 cd/m². The structure PEDOT:PSS/TCTA/E18 Ir(ppy)3/PQ2Ir(acac)/TPBi/CsF/Al was also fabricated. Hybrid white emitting devices with E18 showed the peak external quantum efficiency exceeding 10% and a power efficiency of 14.8 lm/W, at a luminance of 500 cd/m². Skorka et al. investigated two devices: ITO/PEDOT:PSS/PVK: PBD: E19/Ca/Ag and ITO/PEDOT:PSS/PVK: PBD E19/LiF/Al. The current efficiency measured at ca. 9 V was about 0.55 cd/A for the device with the Ca/Ag cathode, and about 0.70 cd/A for that with the LiF/Al cathode.

The 2,7-di(arylamino)- or 3,6-di(arylamino) carbazoles, which were used as fluorescent emitters (Ex) in the OLED devices, are shown in Scheme 8. The compounds 27DBr9ArCr, 36DBr9ArCr, or 36D19ArCr were firstly obtained as starting materials for the synthesis, as shown in Scheme 1. The target compounds E21 [100,101] and E23 [102] were obtained during the Buchwald–Hartwig amination reaction, and the materials E22 [79], E24 [103], and E25 [104] were obtained during the Ullmann reaction of the starting derivatives.

It was reported that the compounds E21–E25 have high thermal stability, with temperatures of decomposition at 436, 440, 387, 285, and 413 °C, respectively. The derivatives E21, E24, and E25 are also suitable for glass formation, with Tg values at 88, 110, and 119 °C. HOMO and LUMO energy levels were described for all the molecules of E21–E25. The HOMO/LUMO values of the molecules were −5.49/−2.13 eV, −5.29/−1.89 eV, −5.25/−2.21 eV, −5.57/−3.26 eV, and −4.94/−2.13 eV, respectively. The electric field dependencies on positive charge drift mobility were characterized in the layers of the material E21. The derivative E21 containing two carbazolyl fragments connected to a central carbazole core demonstrated the hole mobility of 2 × 10⁻³ cm²·V⁻¹·s⁻¹ at a high electric field.
Scheme 8. Structure of diarylaminosubstituted carbazoles as fluorescent emitters for OLED devices.

Luka et al. investigated the properties of a UV OLED structure containing an indium-free transparent anode (ZnMgO:Al) and 2,7-di(9-carbazolyl)-9-(2-ethylhexyl)carbazole E21 as a light-emitting layer. The obtained device was characterized by a high turn-on voltage of ca. 9 V. The driving voltage required for the device to operate at a current of 20 mA/cm² was 10.5 V [105]. Li and co-authors studied the properties of E22 and E23, and suggested that the new compounds have potential applications as blue light emitters in the field of OLEDs. With the emitter E24, the device ITO/HAT-CN/NPB/TAPC/PPT/E24/PPT/TPBi/LiF/Al was described. The OLED displayed excellent device performance, with a power current efficiency of 15.26 cd/A, a power efficiency of 8.14 lm/W, a maximum external quantum efficiency of 13.33%, and a maximal brightness of 1526 cd/m². A single-layer OLED with the structure ITO/Cul/E25/Ca:Al and the bilayer device ITO/Cul/E25/Bphen/Ca:Al were investigated by Deksnys and co-authors. The bilayer structure had a high photometric efficiency of 10.5 cd/A, a maximum luminance of 27,000 cd/m² (at 15 V), and an EQE of ca. 3.3%.

6. Diary(arylamino)-substituted Carbazoles as TADF Emitters for OLEDs

It should be mentioned that only 3,6-diaryl-substituted carbazoles are still reported as TADF compounds suitable for emitting layers (Tx in Scheme 9). The starting materials (36DBrCr and 36DBr9ArCr) for the synthesis of the emitters as objective derivatives were prepared, as shown in Scheme 1. The target compound T1 was obtained during the Suzuki reaction [106]. The compounds T2–T3 were obtained by nucleophilic substitution [107,108]. Other compounds, T4–T5, were obtained during the Buchwald–Hartwig reaction [109].

The Td of compounds T1–T3 were very high and reached 178 °C, 242 °C, and 287 °C, respectively. These destruction temperatures were lower than those of compounds T4–T5, where they exceeded 500 °C. The Ip of the studied derivative T2 in solid-state was reported as 5.89 eV. The values of HOMO/LUMO for compounds T1, T4, and T5 were −5.77/−2.40 eV, −5.63/−2.64 eV and −5.62/−2.67 eV. Only the charge transporting properties for layers of compounds T2 were described. They exhibited a bipolar charge transport with balanced hole and electron mobility exceeding \(10^{-4} \text{ cm}^2\cdot\text{V}^{-1}\cdot\text{s}^{-1}\) at higher than \(3 \times 10^5\) V/cm electric fields.
Scheme 9. Structure of 3,6-diarylcarbazoles as TADF emitters for OLED devices.

The literature reported the TADF-based device ITO/HATCN/NPB/T1/TPBi/LiF/Al, which showed green-bluish light with a 5 V turn-on voltage, a maximum luminance of 8264 cd/m², and a maximum photometric efficiency of 3.96 cd/A. While the blue device ITO/MoO₃/TCTA/mCP/T2/TSP01/TPBi/Ca/Al demonstrated a turn-on voltage of 4.4 V, a maximum brightness of 3100 cd/m², and a photometric efficiency of 5.4 cd/A. Multilayer blue TADF OLEDs with a configuration of ITO/HATCN/TAPC/DCDPA/DBFPO:T4 or T5/DBFPO/TPBi/LiF/Al were fabricated by Karthik et al. These blue devices showed maximal EQE/photometric efficiency/luminance values of 30.7%/46.7cd/A/18,160 cd/m², and 29.1%/36.4 cd/A/11,690 cd/m² for the T4 and T5-based emitters, respectively.

The 3,6-Di(arylamino)carbazoles, which were used as the TADF emitters for OLEDs, are shown in Scheme 10. The target compounds T6–T12 were obtained during the Ullmann amination reaction of the corresponding diiodo or dibromo derivatives shown in Scheme 1 [110–112]. The compounds T13–T14 were synthesized during the Buchwald–Hartwig reactions of the starting materials [113].

Scheme 10. Structures of 3,6-di(arylamino)-substituted carbazoles as TADF emitters for OLED devices.
Thermal stability was presented for the conjugates $T_6$–$T_9$ and $T_{11}$–$T_{14}$. These derivatives have rather high stability, with $T_d$ values from 291 °C to 530 °C. The highest $T_d$ value, which exceeded 530 °C, was mentioned for derivative $T_{12}$. DSC measurements showed that the conjugates $T_{11}$–$T_{14}$ formed amorphous films, with the $T_g$ ranging from 86 °C to 155 °C. The derivatives $T_{13}$–$T_{14}$ containing carbazolyl or diphenylamino fragments had the highest $T_g$ values, reaching 155 °C. The HOMO and LUMO energies of $T_{11}$–$T_{14}$ were presented to be $-5.60/-2.63$ eV, $-5.53/-2.58$, $-5.06/-2.31$ eV, and $-5.17/-2.24$ eV, respectively.

In order to evaluate the potential of the mentioned TADF emitters for OLEDs, authors fabricated devices by using the synthesized TADF materials. Compounds $T_6$–$T_9$ were tested in TADF-based devices of the following structures: ITO/MoO$_3$/one of $T_6$–$T_9$/TCz1:Frlpic/TSP01/TPBi/Ca:Al. The most promising OLED reached a current efficiency of 46.3 cd/A, a power efficiency of 33.2 lm/W, a maximum external quantum efficiency of 20.5%, and a brightness exceeding 3300 cd/m$^2$. Devices with $T_{11}$–$T_{12}$ were fabricated by using the following configuration: ITO/NPB/TAPC/mCBP/$T_{11}$ or $T_{12}$/PPT/TmPyPb/LiF/Al. The OLEDs using $T_{11}$ and $T_{12}$ showed very different maximum EQEs of 9.4 and 23.9%, respectively. Meanwhile, current efficiencies and power efficiencies of the devices were 16.3 and 56.5 cd/A with $T_{11}$, and 14.6 and 50.6 lm/W with $T_{12}$, respectively. In order to study the light-emitting performance of compounds $T_{13}$–$T_{14}$, several devices were designed and fabricated by Zhao et al. with the TADF emitters. Device A was ITO/TAPC/$T_{14}$/TmPyP/LiF/Al and device B was ITO/TAPC/$T_{13}$/TmPyP/Al; both were fabricated using different concentrations of doping ranging from 3 to 30 wt % of the emitters. The OLED B with 15 wt % of the dopant $T_{13}$ displayed excellent device performance, with a current efficiency of 72.1 cd/A, a power efficiency of 61.5 lm/W, a maximum external quantum efficiency of 22.5%, and a brightness exceeding 42,700 cd/m$^2$.

7. Concluding Remarks

Recent developments on 2,7(3,6)-diaryl(arylamino)-substituted carbazole-based electroactive derivatives are presented here with a short description of their preparation, physical properties, and the characteristics of organic light emitting diodes using the materials. The reviewed derivatives had different functions in the OLED devices, including hole transport in the thin layers of the materials, electroluminescence or thermally activated delayed fluorescence in emitting layers as well as host functions for phosphorescent dopants. Some of the derivatives function very effectively as positive charge-transporting layer compounds, enhancing quantum efficiencies, lowering driving voltages, and increasing the life time of the OLEDs. Positive charge drift mobility in the thin films of the most effective derivatives can exceed $2 \times 10^{-3}$ cm$^2$.V$^{-1}$.s$^{-1}$ at high electrical fields.

Among various carbazole-based host derivatives, the diaryl-substituted conjugates are very effective as hosts for the blue (EQE > 27%), green (EQE > 210%), and red (EQE > 20%) phosphorescent organic light emitting diodes. As derivatives for the light emitting layer, the diaryl(arylamino)-substituted carbazoles cover the broad range of emitted lights, from the color blue to green, through the substitution and introduction of different aromatic fragments into the carbazole core at the 2,7(3,6) positions. For example, by using the thermally activated delayed fluorescence (TADF) function of the carbazole-based emitters, blue devices showed a maximum EQE efficiency that exceeded 30%.

Therefore, the low molar mass 2,7(3,6)-dialyl(arylamino)-substituted carbazoles are promising as charge-transporting layer derivatives, emitters, and hosts for various configurations of OLED devices, and further research in the field of new carbazole-based electroactive materials is actively ongoing to improve the characteristics of future OLED devices.

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Abbreviations

NBC  N-bromosuccinimide
36DI Cr  3,6-diiodo-9-alkylcarbazoles
36DI9 Ar Cr  3,6-diiodo-9-arylcarbazoles
36DBr Cr  3,6-dibromo-9-alkylcarbazoles
36DBr9 Ar Cr  3,6-dibromo-9-arylcarbazoles
27DBr Cr  2,7-dibromo-9-alkylcarbazoles
27DBr9 Ar Cr  2,7-dibromo-9-arylcarbazoles
27DNCr  2,7-diarylamino-9-alkylcarbazoles
27DNAr Cr  2,7-diarylamino-9-arylcarbazoles
36DNCr  3,6-diarylamino-9-alkylcarbazoles
36DNAr Cr  3,6-diarylamino-9-arylcarbazoles
27DCr  2,7-diaryl-9-alkylcarbazoles
27Dar Cr  2,7-diaryl-9-arylcarbazoles
36DCr  3,6-diaryl-9-alkylcarbazoles
36Dar Cr  3,6-diaryl-9-arylcarbazoles
Cx  charge transporting compound
HTL  hole transporting layer
T_d  thermal decomposition temperature
T_g  glass transition temperature
I_p  ionization potential
HOMO  the highest occupied molecular orbital
LUMO  the lowest unoccupied molecular orbital
µ_h  hole drift mobility
ITO  indium tin oxide
PEDOT:PSS  poly(3,4-ethylene-dioxythiophene): poly(styrene-sulfonate)
F4TCNQ  2,3,5,6-tetrafluoro-7,7',8,8'-tetracyano-p-quinodimethane
TPD  N,N'-bis(3-methylphenyl)-1,1'-biphenyl-4,4'-diamine
CBP  N,N'-dicarbazolyl-4,4'-biphenyl
BCP  4,7-diphenyl-1,10-phenanthroline
Ir(ppy)_3  tris(2-phenylpyrindine)(iridium(III))
TPBi  2,2',2''-(1,3,3-benzinetriyl)-tris(1-phenyl-1-H-benzimidazole)
Alq3  tris(8-hydroxyquinolinato)aluminum
CuPc  copper(II)phthalocyanine
2-TNATA  4',4'',4'''-tris[N-(2-naphthyl)-N-phenylamino]-triphenylamine
DNTPD  4,4'-bis[N-[4-[N,N,N-bis(3-methylphenyl)aminophenyl]-N-phenylamino]biphenyl
Bebq2  bis[10-hydroxybenzo[h]quinolinato]beryllium
Ir(mphmq)_{2}(tmd)  bis[2,4-dimethyl-6-(4-methyl-2-quinolinyl)phenyl][2,2,6,6-tetramethyl-3,5-heptanediionate
PC-Z  bisphenol Z polycarbonate
HATCN  1,4,5,8,9,11-hexaazatriphenylehexacarbonitride
DCDPA  3,5-di(carbazol-9-yl)-N,N-diphenylaniline
TSPO1  diphenyl-4-triphenylsilylphenyl-phosphine oxide
PhOLED  Phosphorescent organic light-emitting diodes
TAZ  3-(4-biphenylyl)-4-phenyl-5-(4-tert-butylphenyl)-1,2,4-triazole
TGA  thermogravimetric analysis
DSC  differential scanning calorimetry
Bphen  4,7-diphenyl-1,10-phenanthroline
Ir(2-phq)_{3}  tris(2-phenylquinoline)iridium(III)
HATCN  dipyrizarino[2,3-f:2',3'-h]quinoxaline-2,3,6,7,10,11hexacarbonitride
EQE  external quantum efficiency
TADF  thermally activated delayed fluorescence
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