Title: HETEROCYCLIC COMPOUNDS AND THEIR USE IN ELECTRO-OPTICAL OR OPTO-ELECTRONIC DEVICES

Abstract: Compounds exhibiting high hole mobility and/or high glass transition temperatures are provided which are of the formula \[ \text{[Ar}^i\text{]}_m[\text{Ar}^j] \], wherein \( m \) is an integer from 1-3 and \( n \) is an integer and may be 1 or 2; \( \text{Ar}^1 \) represents a dibenzo[b,e][1,4]dioxine, phenoanthrene, thiophene, dibenz[a]anthracene or selenanthrene residue having a linkage to \( \text{Ar}^2 \) at one or two positions selected from ring positions 1-4 and 5-8 and optionally mono-, bi- or poly-substituted with \( \text{C}-\text{C}_2\text{-alkyl} \), \( \text{C}-\text{C}_2\text{-alkoxy} \), fluoro, phenyl or biphenyl which in the case of phenyl or biphenyl may be further substituted with \( \text{C}-\text{C}_2\text{-alkyl} \), \( \text{C}-\text{C}_2\text{-alkoxy} \) or fluoro; \( \text{Ar} \) represents a residue derived from an arylamine in which the aryl rings are phenyl, naphthyl or anthracenyl optionally substituted with \( \text{C}-\text{C}_2\text{-alkyl} \), \( \text{C}-\text{C}_2\text{-alkoxy} \) or fluoro, a polycyclic fused or chain aromatic ring system optionally containing nitrogen or sulphur and in a chain aromatic ring system optionally containing one or more chain oxygen or sulphur atoms, a triarylphosphine oxide or an arylsilane the rings of any of which are optionally substituted with \( \text{C}-\text{C}_2\text{-alkyl} \), \( \text{C}-\text{C}_2\text{-alkoxy} \) or fluoro. The compounds may be used in electron transport layers and may be doped with p-type dopants. They may be incorporated into OLEDs, organic photo-voltaic devices, imaging memory and thin film transistors. In further embodiments there are provided OLEDs or other devices e.g. electrostatic latent image forming members in which improved efficiency is obtained by using as electron transport layers, electron injectors, hosts and emitting (dopants) ambipolar or electron-transmitting compounds in which thiophene is bonded to aryl e.g. 1-anthracenyl-9-yl-thianthrene, 1-biphenyl-4-yl-thianthrene and 9,10-Bis(1-thianthrenyl) anthracene.
HETEROCYCLIC COMPOUNDS AND THEIR USE IN ELECTRO-OPTICAL OR OPTO-ELECTRONIC DEVICES

FIELD OF THE INVENTION

This invention relates to novel compounds and to their use in electro-optical or opto-electronic devices, *inters alia* optical light emitting devices, for example in a hole transport layer. It also relates to a second class of compounds having ambipolar properties. It further relates to novel compounds and to their use in electro-optical or opto-electronic devices, *inter alia* optical light emitting devices, for example in an electroluminescent device in the field of flat panel displays and lighting in an electron transport layer, hole blocking layer, host layer and emissive layer. The invention also relates to novel compounds which can be used as electron transporters in organic photovoltaics and semiconductors in thin film transistor devices.

BACKGROUND TO THE INVENTION

Hole transport materials

One class of hole transport materials comprises aromatic tertiary amines including at least two aromatic tertiary amine moieties (e.g. those based on biphenyl diamine or of a "starburst" configuration), of which the following are representative and of which at this time a-NPB (formula in the specific description below) is believed to be the most widely accepted and used in commercial production.

WO 2011/021803 (Duksan) discloses compounds having a thianthrene structure and their use in OLEDs. In examples, the five compounds below were synthesized.
The above compounds were tested as host materials forming part of a doped electroluminescent layer in an OLED. A layer of 10 nm copper phthalocyanine on an ITO electrode had deposited thereon 30 nm of a-NPB (also known as a-NPD) as hole transporter, a layer of one of the above thianthrene compounds or of CBP to serve as host material for Ir(ppy)$_3$ as dopant, 10 nm of aluminium biphenoxy bis(2-methyl quinolate) as hole blocker, 40 nm of aluminium quinolate as electron transporter, 0.2 nm of lithium fluoride as electron injector and aluminium as cathode. Green electroluminescence with substantially the same colour coordinates was obtained when the test thianthrene compounds were used as host as when CBP was used as host, and turn-on voltage and luminous efficiency (cd/A) ranged from slightly worse than with CBP to somewhat better. However, the fused carbazole ring structures of the Duksan compounds exhibit relatively low hole mobility so that these compounds would be expected to exhibit poor performance if used as hole transporters. It should be mentioned that CBP, which also has carbazole rings linked directly to an extended
aromatic system, has relatively low hole mobility and is also better as a host material in an electroluminescent layer than as a hole transport layer material. It is unsurprising, therefore, that Duksan employs a conventional hole transporter and does not employ any of the thianthrene compounds for that purpose.

Functionalised thianthrenes alleged to have hole transport properties and alleged to be blue emitters are disclosed by Šwist et al., ARKIVOC 2012 (iii), 193-209 (2012). Suzuki coupling of thianthren-2-yl-2-boronic acid with a variety of brominated aromatic amino compounds having C₄H₉, C₁₂H₂₅ or C₁₆H₃₃ substituents gave oils or, in one instance, a solid of low melting point. Stille coupling of 2,8-dibromothianthrene with (Bu)₃Sn- derivatives of thiophene, oxazole, furan and pyridine gave 2,8-bis(2-oxazolyl) thianthrene which was an oil, 2,8-bis(2-thienyl) thianthrene m.p. 176-179°C, 2,8-bis(2-furanyl)thianthrene m.p. 204-206°C and 2,8-bis(2-pyridyl)thianthrene (m.p. 113-114°C, all of which are undesirably low for device applications. Although the compounds were investigated by cyclic voltammetry, DPV spectroscopy, UV-Visible spectroscopy and fluorescence - were alleged to have band gaps in a range appropriate for semiconductors, no hole mobility measurements were made, and the compounds were not tested in OLEDs or other practical devices. They are alleged to be castable into uniform films, but this would not be a property shared by those compounds which are oils. There is no disclosure or suggestion that the materials reported, or any of them, should be used as hole transport layers in OLEDs as opposed to alternative devices such as organic photovoltaic devices, or that any of them give better properties in a hole transport layer of an OLED than established materials e.g. a-NPB and no reason to suppose that this is the case.

US 8012606 (Takahiro et al, Nippon Steel) discloses heterocyclic compounds represented by the general formulæ

\[
\begin{array}{c}
\text{Ar}_1 \\
\text{Ar}_2 \\
\text{N} \\
\text{O} \\
\text{O} \\
\text{Ar}_3 \\
\text{N} \\
\text{Ar}_4 \\
r
\end{array}
\]
wherein: R represents a hydrogen atom, a substituted or unsubstituted alkyl group, or a substituted or unsubstituted aryl group, Ari, Ar₂, Ar₃, and Ar₄ denote independently a substituted or unsubstituted aryl group or Ari, Ar₂ together with the nitrogen atom bonded thereto or Ar₃, Ar₄ together with the nitrogen atom bonded thereto may form a nitrogen-containing hetero ring (e.g. N-carbazolyl, N-phenoxazinyl, N-phenothiazinyl), and m and n are independently 1 or 2. Examples disclose compounds as described above serving as components of the light-emissive layer of an OLED and in one instance as a hole transport layer. The compounds are alleged when applied to an organic EL device to enables the device to be driven at low voltage. When used as a host material, electrons and holes are alleged to move in a well-balanced way to form a wide range of emission of light and attain high luminous efficiency. Furthermore, the heterocyclic compounds have a high triplet energy which is important in an electroluminescent device utilizing phosphorescence. Hence, when used as a host material or an electron-transporting material for a phosphorescent device, the energy of the triplet excited state of a phosphorescent dopant can be confined efficiently and phosphorescence can be obtained at high efficiency. In addition to these good electrical properties, the said heterocyclic compounds are alleged to be stable when formed into thin film. An organic EL device comprising the heterocyclic compound of this invention in its organic layer efficiently emits light of high brightness at low voltage and shows excellent durability and it is applicable to flat panel displays Representative compounds include:

2,7-bis(phenylamino)dibenzodioxin,

2,7-bis(9-carbazolyl)dibenzodioxin,
2,7-bis(N-3-biphenylyl-N-phenylamino)dibenzodioxin,
2,7-bis(N-l-naphthyl-N-phenylamino)dibenzodioxin and
2,7-bis(9-carbazolyl) thianthrene

5 Light emission

Materials that emit light when an electric current is passed through them are well
known and used in a wide range of display applications. Devices which are based on
inorganic semiconductor systems are widely used. However these suffer from the
disadvantages of high energy consumption, high cost of manufacture, low quantum
efficiency and the inability to make flat panel displays. Organic polymers have been
proposed as useful in electroluminescent devices, but it is not possible to obtain pure
colours; they are expensive to make and have a relatively low efficiency. Another
electroluminescent compound which has been proposed is aluminium quinolate, but it
requires dopants to be used to obtain a range of colours and has a relatively low
efficiency.

Patent application WO 98/58037 describes a range of transition metal and
lanthanide complexes which can be used in electroluminescent devices which have
00/26323, WO 00/32719, WO 00/32717, WO 00/32718 and WO 00/44851 describe
electroluminescent complexes, structures and devices using rare earth chelates. US
Patent 5128587 discloses an electroluminescent device which consists of an
organometallic complex of rare earth elements of the lanthanide series sandwiched
between a transparent electrode of high work function and a second electrode of low
work function, with a hole conducting layer interposed between the electroluminescent
layer and the transparent high work function electrode, and an electron conducting layer
interposed between the electroluminescent layer and the electron injecting low work
function cathode. The hole conducting layer and the electron conducting layer are
required to improve the working and the efficiency of the device. The hole transporting
layer serves to transport holes and to block the electrons, thus preventing electrons from
moving into the electrode without recombining with holes. The recombination of
carriers therefore mainly takes place in the emissive layer.
In order to enhance the performance of electroluminescent organometallic complexes the electroluminescent organometallic complex can be mixed with a host material and we have now devised an improved electroluminescent material using a metal quinolate as the host material.

Electron transport materials

Kulkarni et al, Chem. Mater. 2004, 16, 4556-4573 (the contents of which are incorporated herein by reference) have reviewed the literature concerning electron transport materials (ETMs) used to enhance the performance of organic light-emitting diodes (OLEDs). In addition to a large number of organic materials, they discuss metal chelates including aluminium quinolate, which they explain remains the most widely studied metal chelate owing to its superior properties such as high EA (— 3.0 eV; measured by the present applicants as - 2.9 eV) and IP (~ -5.95 eV; measured by the present applicants as about - 5.7 eV), good thermal stability (Tg -172 °C) and ready deposition of pinhole-free thin films by vacuum evaporation. Aluminium quinolate remains a preferred material both for use as a host to be doped with various fluorescent materials to provide an electroluminescent layer and for use as an electron transport layer. For a hole transporter or electron transporter to work effectively in a phosphorescent device, the triplet levels of the respective materials should be higher than the triplet level of the phosphorescent emitter.

SUMMARY OF THE INVENTION

A problem with which the invention is concerned is to provide further, or in the alternative improved, compounds having good hole transport properties when used in OLEDs and other electro-optic or opto-electronic devices. A further problem with which invention is concerned is to provide OLEDs of improved performance and lifetime e.g. by the linkage of nitrogen to a tricyclic ring structure not directly but via an aryl spacer group..

In one aspect the invention provides a compound of the formula \([Ar^1]_m[Ar^2]_n\) wherein:

\(m\) is an integer from 1-3 and \(n\) is an integer and may be 1 or 2;
Ar\(^{1}\) represents a dibenzo[b,e][1,4]dioxine, phenoaxthine, thianthrene, dibenzo[b,e],4-thielenene, selenanthrene or phenothiazinyl residue having a linkage to Ar\(^{2}\) at one or two positions selected from ring positions 1-4, 2,7 and 6-9 and in the case of phenothiazine derivatives optionally at the ring nitrogen atom and optionally mono-, bi- or poly- substituted with Ci-C\(_{4}\)-alkyl-, Ci-C\(_{4}\)-alkoxy-, fluoro, phenyl or biphenyl which in the case of phenyl or biphenyl may be further substituted with Ci-C\(_{4}\)-alkyl-, Ci-C\(_{4}\)-alkoxy- or fluoro;

Ar\(^{2}\) represents a residue derived from an arylamine in which the aryl rings are phenyl, naphthyl or anthracenyl optionally substituted with Ci-C\(_{4}\)-alkyl-, Ci-C\(_{4}\)-alkoxy- or fluoro and in which there is aryl between nitrogen and Ar\(^{1}\), a polycyclic fused or chain aromatic ring system optionally containing nitrogen or sulphur and in a chain aromatic ring system optionally containing one or more chain oxygen or sulphur atoms, a triarylphosphine oxide or an arylsilane the rings of any of which are optionally substituted with Ci-C\(_{4}\)-alkyl-, Ci-C\(_{4}\)-alkoxy- or fluoro.

In such compounds, in many embodiments Ar\(^{1}\) represents thianthrene. In embodiments Ar\(^{1}\) is linked to Ar\(^{2}\) at the 1-position, at the 2-position, the 1- and 8-positions or the 2- and 7-positions.

In many embodiments Ar\(^{2}\) represents a diarylamine residue the nitrogen of which is linked to Ar\(^{1}\), and whose rings are optionally substituted Ci-C\(_{4}\)-alkyl-, C\(_{1}\)-C\(_{4}\)-alkoxy- or fluoro, phenyl, naphthyl, anthracenyl or phenylpyridyl. In other embodiments Ar\(^{2}\) represents a triarylamine residue whose rings are optionally substituted Ci-C\(_{4}\)-alkyl-, Ci-C\(_{4}\)-alkoxy- or fluoro. In the diarylamine series e.g. diphenylamine, N-phenyl-1-naphthylamino, N-phenyl-2-naphthylamino or in the triarylamine series e.g. triphenylamine the rings may optionally be substituted with Ci-C\(_{4}\)-alkyl-, Ci-C\(_{4}\)-alkoxy- or fluoro. In the triarylamine series m may be 1, 2 or 3. In further embodiments Ar\(^{2}\) represents carbazole, spiro-bicarbazole or dibenzothiophene residue optionally ring-substituted with Ci-C\(_{4}\)-alkyl-, Ci-C\(_{4}\)-alkoxy- or fluoro.

Embodiments of the present compounds exhibit a surprisingly favourable combination of hole mobility and high glass transition temperature or melting point, and the compounds also find utility in a hole injection layer and/or in an electron blocking layer. Since they are in general small molecules, many of them are purifiable by sublimation, which is desirable for the production of compounds of the purity required
for OLEDs and other device applications. Embodiments of these compounds exhibit ambipolarity i.e they can be doped to form either electron or hole transport layers depending upon whether they are doped with p-type or n-type dopants. Such molecules are sought-after by device manufacturers because in some embodiments the number of different materials in different layers that have to be used is reduced.

HOMO and LUMO levels compared to α-NPB are shown below.

In a further aspect the invention provides a hole transport material comprising a heterotricyclic compound as defined above and a p-dopant which is an acceptor-type organic molecule. In some embodiments the dopant is present in an amount such that when the material is deposited to form a layer the dopant contributes about 10-40% to the layer thickness, e.g. about 33%. Other materials that may be present in the composition, in embodiments in minor amounts e.g. <50 wt% based on the weight of the heterotricyclic compound, include known organic hole transport agents e.g. an aromatic tertiary amine of biphenyl or starburst configuration, in some embodiments a-NPB.

In a further aspect the invention provides an optical light-emitting diode having first and second electrodes and between said electrodes a layer comprising a compound as defined above or a material as defined above. The layer may be a hole transport layer or a hole injection layer. The device may include an emissive layer comprising a fluorescent emitter, a phosphorescent emitter, an ion fluorescent (rare earth based) emitter or an optical light-emissive quantum dots. A hole injection layer may be
The device may form part of a flat panel display or a lighting panel. Other end-uses may include organic photovoltaic devices, imaging members for forming an electrostatic latent image, organic thin film transistors and dye-sensitised solar cells.

A further problem with which the invention is concerned is to provide compounds which may be used in OLEDs as electron transport materials, electron injectors, hosts, hole blockers and emissive materials and in some embodiments materials emitting in the blue region of the electromagnetic spectrum.

That problem is solved, according to a further aspect of the invention, by compounds in which there is employed a compound in which thianthrene is bonded to polycyclic aryl, or a composition comprising a compound as aforesaid in admixture with a p-type dopant. Examples include the materials ETS-1, ETS-2 and ETS-3 described in the examples together with other materials discussed in relation to the electron transport layer of an OLED. The exemplified compounds, in particular, exhibit a surprisingly favourable combination of electron mobility and high glass transition temperature. Also, these compounds show intense fluorescence as thin films, powders and in solution, particularly in the blue and green region of the visible spectrum.

In a further aspect, the invention provides an electro-optical or opto-electronic device having a layer comprising of compounds described above and further discussed in relation to the electron transport layer. Such devices include OLED Displays, OLED Lighting and also e.g. organic thin film transistors, organic phototransistors, organic photovoltaic cells, organic photodetectors, electronic storage devices based on bistable organic molecules and photoconductive imaging members for creating electrostatic latent images.
In a yet further aspect the invention provides an optical light emitting diode device having a first electrode, a layer comprising any of the compounds in Table 1 or its derivatives and a second electrode. The layer is in an embodiment of an electron transport layer (either neat or doped with low work function metals or metal complexes) or an electron injecting layer or host or emissive layer (host or dopant along with another dopant or host respectively).

The same layer can also act as a hole blocking layer.

It will be appreciated that OLEDs according to the invention may have the aforesaid thianthrene compounds in both the hole and in the electron transport layers or in the hole transport layer, in the electroluminescent layer and in the electron transport layer.

In a further aspect there is provided an OLED having a compound having one or two thianthrene moieties linked to conjugated or aromatic hydrocarbon other than alkyl-substituted fluorine. The OLED may incorporate a compound which is of one of the formulae set out below

(wherein n is an integer and refers to a polyphenyl chain or fused rings),
yl-2-(4-(thianthren-9-yl)phenyl)-l H -benzo[d]imidazole,

wherein X and Y are both S, the rings optionally being substituted e.g. with methyl.

The compound may be 1-anthracenyl-9-yl-thianthrene, 1-biphenyl-4-yl-thianthrene or 9,10-Bis (1-thianthrenyl) anthracene or a mixture thereof. It may be mixed with a low work function metal complex or may be doped with a fluorescent dopant or may be doped with a phosphorescent dopant or may be doped with a rare earth chelate. For example, the compound may be mixed with either lithium quinolinolate (LiQ) or its derivatives or Lithium Schiff base complexes from 1 to 99% by mass, e.g. 10 to 90% by mass, e.g. 20-90% by mass, commonly 30-80% by mass.

The compound may be a component of an emissive layer e.g. a host or it may be comprised in an electron transport layer and/or in an electron injection layer.

In one embodiment there is provided an optical light emitting diode device having a first electrode, a layer comprising 1-anthracenyl-9-yl-thianthrene, 1-biphenyl-4-yl-thianthrene or 9,10-bis (1-thianthrenyl) anthracene and a second electrode and optionally a low work function material.

The above OLEDs may have any of the following features:
(a) the emissive layer is composed of a fluorescent emitter;
(b) emissive layer is composed of a phosphorescent emitter;
(c) the emissive layer is composed of ion fluorescent (rare earth based emitters);
(d) a hole injection layer comprising CuPC, ZnTpTP, 2-TNATA, MoO$_3$, MoO$_x$, WO$_3$, WO$_x$ or NiO$_x$
(e) the hole transporting layer is doped or mixed with an electron acceptor
(f) the hole transporting layer is doped or mixed with an electron acceptor such as TCNQ or F$_4$TCNQ;
(g) the hole transporter is mixed with another hole transporter;
(h) the electron transporter is mixed with a Li, K, Cs compounds or any other compounds of low work function metals or materials;
(i) the electron transporter is mixed with a rare earth metal or its complexes;
(j) the electron transporter is mixed with Lithium Quinolinoates (Liq) or Lithium Schiff Bases;
(k) the electron transporter is mixed with another electron transporter or electron injector;
(l) the electroluminescent layer comprises a metal complex;
(m) the electroluminescent layer comprises zirconium or hafnium quinolate as host material doped with a dopant;
(n) the electroluminescent layer comprises aluminium quinolate as the host material doped with a dopant.
(o) the electroluminescent layer comprises an aromatic tertiary amine as host material doped with a dopant;
(p) the electroluminescent layer comprises a light emitting material which is a metal or metalloid complex;
(q) the electroluminescent layer comprises a luminescent material a metal quinolate, an iridium, ruthenium, osmium, rhodium, iridium, palladium or platinum complex, a boron complex or a rare earth complex;

(r) the electroluminescent layer comprises as electroluminescent material lithium quinolate or aluminium quinolate;

(s) the electroluminescent layer comprises a light-emitting conjugated polymer or copolymer or a dendrimer;

(t) a hole transport layer comprising a-NPB.

The above compounds may be incorporated into other opto-electronic or electro-optic devices as described below.

**BRIEF DESCRIPTION OF THE DRAWINGS**

The accompanying Figs 1-16 show data obtained with practical green and red OLED devices incorporating compounds or materials of the invention.
DESCRIPTION OF PREFERRED FEATURES

Cell structure

The OLEDs of the invention are useful *inter alia* in flat panel displays and typically comprise an anode and a cathode between which is sandwiched a multiplicity of thin layers including an electroluminescent layer, electron injection and/or transport layer(s), hole injection and/or transport layer(s) and optionally ancillary layers. The layers are typically built up by successive vacuum vapour deposition operations, although it may be convenient to form one or more of the layers e.g. the hole injection and hole transport layers by other methods e.g. spin coating or ink jet printing.

A typical device comprises a transparent substrate (examples include glass, plastics (PET, PEN etc.), metals or alloys, semiconductors (organic or inorganic)) on which are successively formed an anode layer, a hole injector (buffer) layer, a hole transport layer, an electroluminescent layer, an electron transport layer, an electron injection layer and a cathode layer which may in turn be laminated to a second transparent substrate. Top emitting OLED’s are also possible in which an aluminium or other metallic substrate carries an ITO layer, a hole injection layer, a hole transport layer, an electroluminescent layer, an electron transport layer, an electron injection layer and an ITO or other transparent cathode, light being emitted through the cathode. A further possibility is an inverted OLED in which a cathode of aluminium or aluminium alloyed with a low work function metal carries successively an electron injection layer, an electron transport layer, an electroluminescent layer, a hole transport layer, a hole injection layer and an ITO or other transparent conductive anode, emission of light being through the anode. If desired a hole blocking layer may be inserted e.g. between the electroluminescent layer and the electron transport layer.

OLEDs of the invention include small molecule OLEDs, polymer light emitting diodes (p-OLEDs), OLEDs that emit light by fluorescence, OLEDs that emit light by phosphorescence (PHOLEDs) and OLEDs that emit light by ion fluorescence (rare earth complexes) and include single-colour or multi-colour active or passive matrix displays.
Anode

In many embodiments the anode is formed by a layer of tin oxide or indium tin oxide coated onto glass or other transparent substrate. Other materials that may be used include antimony tin oxide, aluminium zinc oxide and indium zinc oxide. Other anode materials also include conducting polymers (example: poly(thiophene)(s), poly(aniline)(s) and poly(pyrrole)(s)). If desired a modified anode may be produced e.g. by subsequently treating the ITO surface with oxygen plasma, and then conditioned as a modified anode by decomposing CHF$_3$ gas in a plasma treatment chamber to deposit a ~1-nm-thick layer of CF$_X$. In active matrix embodiments the anode may be a high work function metal or alloy e.g. gold or platinum or may be crystalline, polycrystalline, continuous grain or amorphous silicon which may be p-doped or metal oxides such Mo, W and Ni oxides.. Cells in which the anode is of doped or un-doped graphene are also within the invention.

Hole injection and hole transport layers

Commonly OLEDs have distinct hole injection and hole transport layers. The applicant has considered the properties that are desirable for the material of a hole injection layer and for the material of a hole transport layer of an OLED, and these are set out in the table below:
<table>
<thead>
<tr>
<th>Property</th>
<th>Hole Injectors (HI’s)</th>
<th>Hole Transporters (HT’s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LUMO Level</td>
<td>Not critical</td>
<td>Critical</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$E_{\text{LUMO}} \geq -2.6 \text{ eV}$</td>
</tr>
<tr>
<td></td>
<td>(i.e., $</td>
<td>E_{\text{LUMO}}</td>
</tr>
<tr>
<td>HOMO Level</td>
<td>$E_{\text{HOMO}} \leq -5.2 \text{ eV}$</td>
<td>$E_{\text{HOMO}} \leq -5.7 \text{ eV}$</td>
</tr>
<tr>
<td></td>
<td>(i.e., $</td>
<td>E_{\text{HOMO}}</td>
</tr>
<tr>
<td>Hole mobility ($\mu$)</td>
<td>As high as possible</td>
<td>As high as possible, preferably greater than</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$1 \times 10^{-5} \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$</td>
</tr>
<tr>
<td>Capable of acting as a</td>
<td>Yes</td>
<td>Not a pre-requisite as it is normally deposited on the</td>
</tr>
<tr>
<td>planarization layer for the</td>
<td></td>
<td>anode</td>
</tr>
<tr>
<td>anode</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Interface to the anode</td>
<td>Must act as an “anchoring” agent with good adhesion.</td>
<td>Capable of forming amorphous films.</td>
</tr>
<tr>
<td></td>
<td>Must improve the film properties and</td>
<td></td>
</tr>
<tr>
<td></td>
<td>subsequent organic layers.</td>
<td></td>
</tr>
<tr>
<td>Thermal stability</td>
<td>High $T_g$ (glass transition point), $T_m$ (melting point) and $T_d$ (decomposition point)</td>
<td>High $T_g$ (glass transition point), $T_m$ (melting point) and $T_d$ (decomposition point)</td>
</tr>
<tr>
<td></td>
<td>$T_g &gt; 150^\circ \text{C}, T_m &gt; 250^\circ \text{C}, T_d &gt; 350^\circ \text{C} (\text{under nitrogen})$</td>
<td>$T_g &gt; 120^\circ \text{C}, T_m &gt; 250^\circ \text{C}, T_d &gt; 250^\circ \text{C} (\text{under nitrogen})$</td>
</tr>
<tr>
<td>Doping</td>
<td>N. A</td>
<td>Capable of being doped with acceptor molecules to enhance the conductivity.</td>
</tr>
<tr>
<td>Typical thickness</td>
<td>1 nm- 100 nm</td>
<td>20 -200 nm</td>
</tr>
</tbody>
</table>
It will be appreciated that many of the above characteristics are desirable rather than essential. For example, although Tg (where such a transition occurs or is detectable) should be as high as possible, many current hole injection materials have Tg of 100-120°C, or in some instances lower e.g. about 90°C.

**Hole injection materials**

A single layer may be provided between the anode and the electroluminescent material, but in many embodiments there are at least two layers one of which is a hole injection layer (buffer layer) and the other of which is a hole transport layer, the two layer structure offering in some embodiments improved stability and device life, see US-A-4720432 (VanSlyke et al, Kodak). The hole injection layer may serve to improve the film formation properties of subsequent organic layers and to facilitate the injection of holes into the hole transport layer.

Suitable materials for the hole injection layer which may be of thickness e.g. 0.1-200 nm depending on material and cell type include hole-injecting porphyrinic compounds - see US-A-43 56429 (Tang, Eastman Kodak) e.g. zinc phthalocyanine, copper phthalocyanine and ZnTpTP, whose formula is set out below:

![ZnTpTP](image)

Particularly good device efficiencies, turn/on voltages and/or lifetimes may be obtained where the hole injection layer is ZnTpTP. A further material that may be used is hexacyanohexaazatriphenylene (CHATP) of structure:
The hole injection layer may also be a fluorocarbon-based conductive polymer formed by plasma polymerization of a fluorocarbon gas - see US-A-6208075 (Hung et al; Eastman Kodak), a triarylamine polymer - see EP-A-0891121 (Inoue et al, TDK Corporation) or a phenylenediamine derivative - see EP-A-1029909 (Kawamura et al, TDK Corporation) or the materials described in US-A-6436559 (Ueno, Canon) and 6720573 (Se-Hwan, LG Chemical Co., Ltd.). It may also be a solution-processable hole injecting polymer e.g. PEDOT/PSS (poly-3,4-ethylenedioxythiophene doped with poly(styrenesulfonate) or a block copolymer of PEDOT and a flexible polymer such as a polyether, polysiloxane, polyester, or polyacrylate. Poly(aniline)(s) and Poly(pyrrole)(s) are also part of this invention. Methods of applying such materials include solution processing methods, e.g. spin coating, printing through a mask and ink jet printing e.g. of a relatively dilute solution where thin hole injection layers are desired.

Other hole injectors include oxides of Mo, W and Ni.

**Hole-transport materials**

Hole transport layers which may be used are in some embodiments preferably of thickness 10 to 200 nm, e.g. 20-200nm.

The hole transport compounds used herein include compounds having at least one dibenzo[£,e][1,4]dioxine, phenoxanthine, thianthrene, dibenzo[b,e]1,4-thiaselenane, selenanethrene or phenothiazine residue per molecule. In some embodiments the dibenzo[£,e][1,4]dioxine, phenoxanthine, thianthrene, dibenzo[b,e]1,4-thiaselenane or selenanethrene residue serves as scaffold and may have one or two ring substituents which are aryl or heteroaryl. Linkage may be at any ring position or in the case of compounds having two ring aryl or heteroaryl substituents they may be at any pair of stereochemically compatible ring positions e.g. the 1-, 2-, 1,9- or 2,7-positions of the tricyclic ring. Said residue or residues may be optionally mono-, bi- or poly- substituted.
with \( \text{C}_i-\text{C}_4\)-alkyl-, \( \text{C}_i-\text{C}_4\)-alkoxy-, fluoro, phenyl or biphenyl which in the case of phenyl or biphenyl substituents may be themselves further substituted with \( \text{C}_i-\text{G}_j\)-alkyl- (preferably methyl), \( \text{C}_i-\text{C}_4\)-alkoxy- (preferably methoxy) or fluoro. Alkyl substituents include methyl, ethyl, \(-\)-propyl, \(\parallel\)-propyl and \(\perp\)-butyl. Alkoxy substituents include methoxy, ethoxy, \(\langle\langle\)-propoxy, \(\langle\langle\parallel\)-propoxy and \(\langle\langle\perp\)-butoxy. The residue or residues may also include an aryl linking group attached at one end as a substituent to the dibenzo[3,e][1,4]dioxine, phenoxythine, thianthrene, dibenzo[b,e][1,4]-thiaselenane or selenanthrene residue e.g. 1,4-phenylene optionally substituted by \(\text{C}_1-\text{C}_4\)-alkyl- (preferably methyl), \(\text{C}_i-\text{C}_4\)-alkoxy- (preferably methoxy) or fluoro. The hole transport material may, for example, include one, two or three dibenzo[3,e][1,4]dioxine, phenoxythine, thianthrene, dibenzo[b,e][1,4]-thiaselenane or selenanthrene residues per molecule, with two such residues at present being believed an optimum.

The following numbering system is used for the thianthrene ring and the other related ring structures mentioned herein:

A sub-genus of the above hole transport compounds has a tricyclic structure attached to a triarylamine or triaryl phosphine oxide having a single nitrogen or phosphorus atom and includes compounds having a favourable combination of melting points, phase transition temperatures and hole mobilities for use in electronic devices.

A broad genus of compounds has one or more nitrogen, phosphorus or silicon atoms bonded to dibenzo[b,e][1,4]dioxide, phenoxythine, thianthrene, dibenzo[b,e][1,4]-thiaselenane, selenanthrene or phenothiazine via aryl. Such compounds may be of the formula:
wherein:

\( Z \) is N, P=0 or Si;

\( X \) and \( Y \) which may be the same or different each represent O, S or Se, or \( X \) may represent NH and \( Y \) may represent O, S or Se;

the groups \( R \) which may be the same or different independently represent hydrogen or one or more ring substituents selected from methyl, methoxy, ethyl, ethoxy, aryl (e.g. phenyl) and fluoro;

\( m \) is 1-3 when \( Z \) is N or P=0 or may be 1-4 when \( Z \) is Si;

\( n \) is 0 or 1;

\( p \) is 0, 1 or 2;

\( q \) and \( r \) are 0 or 1,

\( s \) is 1 when \( Z \) is N or P=0 or is 2 when \( Z \) is Si;

the dotted linkage when present indicates that the two phenyl rings form part of a \( 9H \)-carbazol-9-yl residue

the solid line indicates a linkage between the tricyclic ring directly, by phenyl or by biphenyl; and

linkages to the tricyclic ring may be at its 1- or 2-position or may be at its 2, 7 or 2.8-positions or when \( X \) is NH may be in the 10-position.

A group of the above compounds has \( n=1 \), \( p=1 \) or 2 and \( m \) is 1, 2 or 2, the linkages between triaryl and \( Z \) being by phenyl or biphenyl. Another group has \( m=1 \), \( n=1 \) or 2 and \( p=0 \) or 1.

**Triarylamines and related phosphine oxides and silane derivatives**

A particular sub-group of compounds incorporating a single triarylamine residue may be of the formula:
wherein:

m is 1, 2 or 3, the dotted lines correspondingly representing bonds for each phenothiazine residue to one of the phenyl or naphthyl rings;

n is 1 or 2;

p, q and r are independently 0 or 1;

X and Y which may be the same or different each represent O, S or Se, or X may represent NH and Y may represent O, S or Se;

Z represents N or P=0;

the groups R which may be the same or different independently represent hydrogen or ring substituents selected from methyl, methoxy, ethyl, ethoxy, aryl (e.g. phenyl) and fluoro; and

linkages may be at the 1- or 2-position of the tricyclic ring or rings or may be at the 2, 7 or 2,8-positions or when X is NH may be in the 10-position, and may be α- or β- to the phenyl or naphthyl rings to which they are attached.

In one group of these compounds m is 1 and n is 1, 2 or 3. In another group of compounds m is 1 and n is 1 or 2.

For example, the scaffold may be triphenylamine with one, two or three of the phenyl rings substituted with a thianthrenyl residue of the type described above. The thiazine residue may be attached to the phenyl ring at any sterically available position, commonly the 4-position as in many of the compounds shown below.

In compounds where X is NH and Y is O, S or Se, attachment may be between the nitrogen atom and a phenyl ring at any sterically available position, most usually the 4-position.

The compound \(N\)-4-(thianthren-1-yl)tri phenylamine (also phenyl-\(N\)-(4-(thianthren-7-yl)phenyl)benzenamine) can be made by brominating thianthrene under reflux in acetic acid to give 2-bromothianthrene which is reacted with aqueous acidic
trimethyl borate to give thianthren-2-yl-2-boronic acid which is a widely useful intermediate for Suzuki coupling reactions to provide thianthren-2-yl compounds herein. The above intermediate is Suzuki coupled (Pd(0)/K$_2$CO$_3$) with N-(4-bromophenyl)-N-phenylbenzenamine to give the desired product. Likewise the compound N,N-bis(4-bromophenyl)benzenamine can be Suzuki coupled with thianthren-2-yl-2-boronic acid to give 4,4''-di-(thianthren-1-yl)triphenylamine (also phenyl-bis-4-thianthren-2-yl)phenylamine) of structure

![Chemical Structure](image)

The compound 4-(thianthren-1-yl)-4',4''-dimethoxytriphenylamine (also 4-methoxy-N-(4-methoxyphenyl)-N-(4-(thianthren-9-yl)phenyl)benzenamine) may be made as follows:

![Chemical Reaction](image)

N-4-(2,8-dimethoxythianthren-1-yl)triphenylamine (also 4-(2,8-dimethoxy-thianthren-9-yl)phenyl)-N-phenylbenzenamine) may be made by the following sequence of reactions:
4-(N-Phenyl-N-w-tolylamino)-4'-(thianthren-1-yl)biphenyl may also be made by Suzuki coupling as shown:

See also the following compounds:

wherein X and Y are as defined above.

Compounds within this sub-genus include compounds of name and/or structural formula below.

4-(thianthren-1-yl)triphenylamine (HTS-1);
4,4'-di-(thianthren-1-yl)triphenylamine (HTS-4);
4,4',4''-tri-(thianthren-1-yl)triphenylamine (HTS-2);
4-(thianthren-2-yl)triphenylamine;
4,4'-di-(thianthren-2-yl)triphenylamine;
4,4',4''-tri-(thianthren-2-yl)triphenylamine;
4-(1OH-phenothiazin-10-yl)triphenylamine (HTS-5);
4,4'-di-(1OH-phenothiazin-10-yl)triphenylamine;
4,4',4''-tri-(1OH-phenothiazin-10-yl)triphenylamine;
2,7-bis-(4-diphenylaminophenyl)thianthrene
2,7-bis-(4-diphenylaminophenyl)-3,8-dimethylthianthrene
3-(thianthren-2-yl)triphenylamine;
N-(4-thianthren-1-ylphenyl)-di-p-tolylamine;
4-(thianthren-1-yl)-4',4''-dimethoxytriphenylamine;
4-(2,8-dimethoxythianthren-1-yl)triphenylamine;
1,9-di-(4-(diphenylamino)phenyl)thianthrene;
4-(thianthren-1-yl)triphenylphosphine oxide;
N-phenyl-N-(1-(thianthren-1-yl)naphthalen-4-yl)naphthalen-1-amine;
N-(naphthalen-1-yl)-N-(4-(thianthren-1-yl)phenyl)naphthalen-1-amine;
N-(naphthalen-1-yl)-N-(1-(thianthren-1-yl)naphthalen-4-yl)naphthalen-1-amine;
4-(N-Phenyl-N-m-tolylamino)-4'-(thianthren-1-yl)biphenyl;

Triphenylphosphine oxide compounds

Compounds containing triphenyl phosphine e.g. 4-(thianthren-1-yl)triphenylphosphine oxide, 4-(thianthren-2-yl)triphenylphosphine oxide and 4-(10H-phenothiazin-10-yl)triphenylphosphine oxide and may be generically represented as indicated below:
Similar to the above sub-genus are compounds where an above defined tricyclic structure is attached to tetraphenylsilane to give a compound of formula III of which examples in the thianthrene and appear below:

Ring substitution with methyl, methoxy or fluoro is also possible, and the molecules could contain 2, 3 or 4 thianthrene residues per molecule.

Compounds containing aminobiphenyl, diaminobiphenyl, 2,7-diaminofluorene, 2,7-dimaminospirobifluorene and 2,7,2',7'tetraaminospirobifluorene residues

Hole transport materials of the above type may, for example be of the formula
wherein X and Y independently are O, S or Se, R is as defined above and Ri may be phenyl which may be substituted with methyl, methoxy or fluoro (e.g. to give w-tolyl or p-tolyl) or the groups R may together form part of a spirobifluorene group. In formulae VI and VII, it will be understood that the groups R on the biphenyl rings may form fluorine or spirobifluorene.

Examples of such compounds include:
Examples of preparative methods are shown below:

```
H N—C6H4—N=H + C8H7S2Br →
```

and
The following structures (which may be varied along the general lines indicated above e.g. with ring substitutions and change of S for O, Se or NH) are also analogous:
Compounds having one, two or three thianthrene moieties linked to conjugated or aromatic hydrocarbon other than alkyl-substituted fluorene

wherein R, X and Y is as defined in claim 1 and in embodiments X and Y are both S.
Compounds in this class include the following:

2-(naphthalen-1-yl)thianthrene
1-(phenanthren-9-yl)thianthrene,
1-(10-phenylanthracen-9-yl)thianthrene,
1-(4-(thiophen-2-yl)phenyl)thianthrene
1-(9,10-di(naphthalen-2-yl)anthracen-6-yl)thianthrene
10-(4-(9H-carbazol-9-yl)phenyl)-1OH-phenothiazine
2-(10H-phenothiazinyl)dibenzothiophene
9,10-di(thianthren-yl)anthracene
1,3,5-tri(thianthren-1-yl)benzene
1,3,5-tris-(4-(thianthren-1-yl)phenyl)benzene
(4(9-(4-(thianthren-1-yl)phenyl)-9H-carbazole
9-(4-(thianthren-1-yl)benzyl)-9H-carbazole
1-phenyl-2-(4-(thianthren-9-yl)phenyl)-1H-benzo[d]imidazole
2-phenyl-1-(4-(thianthren-1-yl)phenyl)ethanone
1-(9,9-di-p-tolyl-9H-fluoren-2-yl)thianthrene
1-(2-thianthren-1-yl)-9,9-di-p-tolyl-9H-fluoren-7-yl)thianthrene
1-(9,9-dipropyl-2-(thianthren-1-yl)-9H-fluoren-7-yl)thianthrene,
1-(4-(9-(4-(thianthren-1-yl)phenyl)-9H-fluoren-9-yl)phenyl)thianthrene
1-(4-(9-(4-(thianthren-1-yl)phenyl)-9H-fluoren-9-yl)phenyl)thianthrene

The compound 9-(4-(thianthren-1-yl)phenyl)-9H-carbazole can be prepared by reacting 9H-carbazole with 1-bromo-4-iodobenzene under reflux in NMP in the presence of Cul and K$_2$CO$_3$ to give 9-(4-bromophenyl)-9H-carbazole which in turn is Suzuki coupled with thianthren-1-yl-l-boronic acid (a generally useful intermediate for thianthren-1-yl compounds herein) to give the desired product.

Dopants for the hole transport layer
The hole transport layer may comprise any of the above mentioned compounds which are sublimable small molecules and which may be p-doped e.g. with an acceptor-type organic molecule e.g. tetracyanoquinodimethane or tetrafluorotetracyanoquinodimethane (F₄TCNQ). The amount of dopant may be such as to contribute 10-40% e.g. about 33% to layer thickness. In addition to one of the above mentioned thianthrene compounds and optionally p-type dopant, there may be present additionally one or more known hole transport compounds which may also be sublimable small molecules. For example, aromatic tertiary amines provide a class of preferred hole-transport materials, e.g. aromatic tertiary amines having at least two aromatic tertiary amine moieties (e.g. those based on biphenyl diamine or of a "starburst" configuration). For example, aromatic amines may be used of the general formulae (i) - (vii) below:

(i) 

(ii) 

(iii)
wherein the groups R in any of the formulae in (i) to (vi) can be the same or different and are selected from hydrogen; substituted and unsubstituted aliphatic groups; substituted and unsubstituted aromatic, heterocyclic and polycyclic ring structures; halogens; and thiophenyl groups; and wherein in formula (i) the methyl groups may be replaced by C₁-C₄ alkyl or monocyclic or polycyclic aryl or heteroraryl which may be further substituted e.g. with alkyl, aryl or arylamino.

Further hole transport materials comprise

wherein the groups R₁-R₄ when appearing in either of the above formulae can be the same or different and are selected from hydrogen; substituted and unsubstituted
aliphatic groups; substituted and unsubstituted aromatic, heterocyclic and polycyclic ring structures; halogens; and thiophenyl groups.

The following are representative of suitable aromatic tertiary amines:

\[
\begin{align*}
\text{TPD} & \quad T_g \, (^{\circ}\text{C}) \quad 61 \\
& \quad \mu h \, (\text{cm}^2 \, \text{V}^{-1} \, \text{s}^{-1}) \quad 1 \times 10^{-3}
\end{align*}
\]

\[
\begin{align*}
\text{a-NPB} & \quad T_g \, (^{\circ}\text{C}) \quad 98 \\
& \quad \mu h \, (\text{cm}^2 \, \text{V}^{-1} \, \text{s}^{-1}) \quad 1 \times 10^{-4}
\end{align*}
\]

A further possible material is 4,4',4''-tris(carbazolyl)-triphenylamine (TCTA) which is a hole transport material with a wider band gap than a-NPB and which can in some embodiments assist in confining excitation to the emissive layer.
Further possible materials are spiro-linked molecules which are aromatic amines e.g. spiro-TAD (2,2',7,7'-tetrakis-(diphenylamino)-spiro-9,9'-bifluorene).

A further class of small molecule hole transport materials is disclosed in WO 2006/061594 (Kathirgamanathan et al) and is based on diamino dianthracenes. Typical compounds include:

- 9-(10-(N-(naphthalen-1-yl)-N-phenylamino)anthracen-9-yl)-N-(naphthalen-1-yl)-N-phenylanthracen-10-amine;
- 9-(10-(N-biphenyl-N-2-w-tolylamino)anthracen-9-yl)-N-biphenyl-N-2-w-tolylamino-anthracen-10-amine; and
- 9-(10-(N-phenyl-N-m-tolylamino)anthracen-9-yl)-N-phenyl-N-m-tolylanthracen-10-amine.

Hole transport layers which may be used are preferably of thickness 10 to 200 nm.

15 Electroluminescent materials

In principle any electroluminescent material may be used, including molecular solids which may be fluorescent dyes e.g. perylene dyes, metal complexes e.g. Alq₃, Ir(III)L₃, rare earth chelates e.g. Tb(III) complexes, dendrimers and oligomers e.g. sexithiophene, or polymeric (conjugated or otherwise) emissive materials. The electroluminescent layer may comprise as luminescent material a metal quinolate, iridium, ruthenium, osmium, rhodium, palladium or platinum complex, a boron complex or a rare earth complex.

One preferred class of electroluminescent materials comprises host materials doped with dyes which may be fluorescent, phosphorescent or ion-phosphorescent (rare earth). The term "electroluminescent device" includes electrophosphorescent devices.

Preferably the host is doped with a minor amount of a fluorescent material as a dopant, preferably in an amount of 0.01 to 25% by weight of the doped mixture. As discussed in US-A-4769292 (Tang et al, Kodak), the contents of which are included by reference, the presence of the fluorescent material permits a choice from amongst a wide latitude of wavelengths of light emission. In particular, as disclosed in US-A-4769292 by blending with the organo metallic complex a minor amount of a fluorescent
material capable of emitting light in response to hole-electron recombination, the hue of the light emitted from the luminescent zone, can be modified. In theory, if a host material and a fluorescent material could be found for blending which have exactly the same affinity for hole-electron recombination, each material should emit light upon injection of holes and electrons in the luminescent zone. The perceived hue of light emission would be the visual integration of both emissions. However, since imposing such a balance of host material and fluorescent materials is limiting, it is preferred to choose the fluorescent material so that it provides the favoured sites for light emission. When only a small proportion of fluorescent material providing favoured sites for light emission is present, peak intensity wavelength emissions typical of the host material can be entirely eliminated in favour of a new peak intensity wavelength emission attributable to the fluorescent material.

While the minimum proportion of fluorescent material sufficient to achieve this effect varies, in no instance is it necessary to employ more than about 10 mole percent fluorescent material, based of host material and seldom is it necessary to employ more than 1 mole percent of the fluorescent material. On the other hand, limiting the fluorescent material present to extremely small amounts, typically less than about $10^{-3}$ mole percent, based on the host material, can result in retaining emission at wavelengths characteristic of the host material. Thus, by choosing the proportion of a fluorescent material capable of providing favoured sites for light emission, either a full or partial shifting of emission wavelengths can be realized. This allows the spectral emissions of the EL devices to be selected and balanced to suit the application to be served. In the case of fluorescent dyes, typical amounts are 0.01 to 5 wt%, for example 2-3 wt%. In the case of phosphorescent dyes typical amounts are 0.1 to 15 wt%. In the case of ion phosphorescent materials typical amounts are 0.01-25 wt% or up to 100 wt%.

Choosing fluorescent materials capable of providing favoured sites for light emission, necessarily involves relating the properties of the fluorescent material to those of the host material. The host can be viewed as a collector for injected holes and electrons with the fluorescent material providing the molecular sites for light emission. One important relationship for choosing a fluorescent material capable of modifying the hue of light emission when present in the host is a comparison of the reduction potentials of the two materials. The fluorescent materials demonstrated to shift the
wavelength of light emission have exhibited a less negative reduction potential than that of the host. Reduction potentials, measured in electron volts, have been widely reported in the literature along with varied techniques for their measurement. Since it is a comparison of reduction potentials rather than their absolute values which is desired, it is apparent that any accepted technique for reduction potential measurement can be employed, provided both the fluorescent and host reduction potentials are similarly measured. A preferred oxidation and reduction potential measurement techniques is reported by R. J. Cox, Photographic Sensitivity, Academic Press, 1973, Chapter 15.

A second important relationship for choosing a fluorescent material capable of modifying the hue of light emission when present in the host is a comparison of the band-gap potentials of the two materials. The fluorescent materials demonstrated to shift the wavelength of light emission have exhibited a lower band gap potential than that of the host. The band gap potential of a molecule is taken as the potential difference in electron volts (eV) separating its ground state and first singlet state. Band gap potentials and techniques for their measurement have been widely reported in the literature. The band gap potentials herein reported are those measured in electron volts (eV) at an absorption wavelength which is bathochromic to the absorption peak and of a magnitude one tenth that of the magnitude of the absorption peak. Since it is a comparison of band gap potentials rather than their absolute values which is desired, it is apparent that any accepted technique for band gap measurement can be employed, provided both the fluorescent and host band gaps are similarly measured. One illustrative measurement technique is disclosed by F. Gutman and L. E. Lyons, Organic Semiconductors, Wiley, 1967, Chapter 5.

With host materials which are themselves capable of emitting light in the absence of the fluorescent material, it has been observed that suppression of light emission at the wavelengths of emission characteristics of the host alone and enhancement of emission at wavelengths characteristic of the fluorescent material occurs when spectral coupling of the host and fluorescent material is achieved. By "spectral coupling" it is meant that an overlap exists between the wavelengths of emission characteristic of the host alone and the wavelengths of light absorption of the fluorescent material in the absence of the host. Optimal spectral coupling occurs when the emission wavelength of the host is within ±25nm of the maximum absorption of the
fluorescent material alone. In practice advantageous spectral coupling can occur with peak emission and absorption wavelengths differing by up to 100 nm or more, depending on the width of the peaks and their hypsochromic and bathochromic slopes. Where less than optimum spectral coupling between the host and fluorescent materials is contemplated, a bathochromic as compared to a hypsochromic displacement of the fluorescent material produces more efficient results.

Useful fluorescent materials are those capable of being blended with the host and fabricated into thin films satisfying the thickness ranges described above forming the luminescent zones of the EL devices of this invention. While crystalline organometallic complexes do not lend themselves to thin film formation, the limited amounts of fluorescent materials present in the host permit the use of fluorescent materials which are alone incapable of thin film formation. Preferred fluorescent materials are those which form a common phase with the host. Fluorescent dyes constitute a preferred class of fluorescent materials, since dyes lend themselves to molecular level distribution in the host. Although any convenient technique for dispersing the fluorescent dyes in the host can be used preferred fluorescent dyes are those which can be vacuum vapour deposited along with the host materials.

One class of host materials comprises metal complexes e.g. metal quinolates such as lithium quinolate, aluminium quinolate, titanium quinolate, zirconium quinolate or hafnium quinolate which may be doped with fluorescent materials or dyes as disclosed in patent application WO 2004/058913.

In the case of quinolates e.g. aluminium quinolate:

(a) the compounds below, for example, can serve as red dopants:
the compounds below, for example can serve as green dopants:

wherein R is C\textsubscript{i} - C\textsubscript{4} alkyl, monocyclic aryl, bicyclic aryl, monocyclic heteroaryl, bicyclic heteroaryl, aralkyl or thienyl, preferably phenyl; and

c) for biphenyloxy aluminium bis-quinolate (BA\textsubscript{1}Q\textsubscript{2}), BAq2 (aluminium bis(-2-methylquinolate) or aluminium quinolate the compounds perylene and 9-(10-(N-(naphthalen-8-yI)-N-phenylamino)anthracen-9-yI)-N-(naphthalen-8-yI)-N-phenylanthracen-10-amine can serve as a blue dopants.

Another preferred class of hosts is small molecules incorporating conjugated aromatic systems with e.g. 4-10 aryl or heteroaryl rings which may bear substituents e.g. alkyl (especially methyl), alkoxy and fluoro and which may also be doped with fluorescent materials or dyes.

An example of a system of the above kind is a blue-emitting material based on the following compound (Compound H) as host
and perylene or 9-(10-(N-(naphthalen-8-yl)-N-phenylamino)anthracen-9-yl)-N-(naphthalen-8-yl)-N-phenanthracen-10-amine as dopant. Further examples of host materials which are small aromatic molecules are shown below:

2,9-Bis(2-thiophen-2-yl-vinyl)-[1,10] phenanthroline may, as explained above, may be used as host in the electroluminescent layer or may be present on its own.
Other blue emitting systems include

![Tetra-tert-Butyl Perylene](image)

![TBADN](image)

Blue-emitting materials may be based on an organic host (e.g. a conjugated aromatic compound as indicated above) and diarylamine anthracene compounds disclosed in WO 2006/090098 (Kathirgamanathan et al.) as dopants. For example, CBP may be doped with blue-emitting substituted anthracenes inter alia

9,10-bis(4-methylbenzyl)-anthracene,
9,10-bis(2,4-dimethylbenzyl)-anthracene,
9,10-bis(2,5-dimethylbenzyl)-anthracene,
1,4-bis(2,3,5,6-tetramethylbenzyl)-anthracene,
9,10-bis(4-methoxybenzyl)-anthracene,
9, 10-bis-(9H-fluoren-9-yl)-anthracene,
2,6-di-t-butylantracene,
2,6-di-t-butyl-9,10-bis-(2,5-dimethylbenzyl)-anthracene,
2,6-di-t-butyl-9, 10-bis-(naphthalene- 1-ylmethyl)-anthracene.

Further blue-emitting materials may employ TCTA as host and it may be doped with the blue phosphorescent materials set out below, see WO 2005/080526 (Kathirgamanathan et al.):

**Blue Phosphorescent Materials**

\[
\begin{align*}
\text{Ir} & \quad \text{N} & \quad \text{N} & \quad \text{Ir} & \quad \text{N} & \quad \text{N} \\
& \quad \text{N} & \quad \text{N} & \quad \text{Ir} & \quad \text{O} & \quad \text{N} & \quad \text{Ph} \\
& \quad \text{N} & \quad \text{N} & \quad \text{Ir} & \quad \text{O} & \quad \text{N} & \quad \text{Ph}
\end{align*}
\]

\[
\begin{align*}
\lambda_{\text{max}} & \quad 495\text{nm (DCM)} & \quad 493\text{nm (DCM)} & \quad 485\text{nm (DCM)}
\end{align*}
\]

\[
\begin{align*}
\text{Ir} & \quad \text{N} & \quad \text{N} & \quad \text{Ir} & \quad \text{N} & \quad \text{N} \\
& \quad \text{N} & \quad \text{N} & \quad \text{Ir} & \quad \text{O} & \quad \text{N} & \quad \text{Ph} \\
& \quad \text{N} & \quad \text{N} & \quad \text{Ir} & \quad \text{O} & \quad \text{N} & \quad \text{Ph}
\end{align*}
\]

\[
\begin{align*}
\lambda_{\text{max}} & \quad 485\text{nm (DCM)} & \quad 484\text{nm (DCM)} & \quad 483\text{nm (DCM)}
\end{align*}
\]
Examples of green phosphorescent materials that may be employed with CBP or TAZ are set out below (see WO 2005/080526):

**Green Phosphorescent Materials**
Examples of red phosphorescent materials that may be employed with CBP or TAZ are set out below (see WO 2005/080526):

**Red Phosphorescent Materials**
Further examples are:
Other examples include dendrimers such as
As further dopants, fluorescent laser dyes are recognized to be particularly useful fluorescent materials for use in the organic EL devices of this invention. Dopants which can be used include diphenylacridine, coumarins, perylene and their derivatives. Useful fluorescent dopants are disclosed in US 4769292. One class of preferred dopants is coumarins. The following are illustrative fluorescent coumarin dyes known to be useful as laser dyes:

FD-1 7-Diethylamino-4-methylcoumarin,
FD-2 4,6-Dimethyl-7-ethylaminocoumarin,
FD-3 4-Methylumbelliferone,
FD-4 3-(2’-Benzothiazolyl)-7-diethylaminocoumarin,
FD-5 3-(2’-Benzimidazolyl)-7-N,N-diethylaminocoumarin,
FD-6 7-Amino-3-phenylcoumarin,
FD-7 3-(2’-N-Methylbenzimidazolyl)-7-N,N-diethylaminocoumarin,
FD-8 7-Diethylamino-4-trifluoromethylcoumarin,
FD-9 2,3,5,6-H4-Tetrahydro-8-methylquinolazino[9,9a,l-gh]coumarin.
FD-10 Cyclopenta[c]julolindino[9,10-3]-1 H-pyran-1 1-one,
FD-11 7-Amino-4-methylcoumarin,
FD-12 7-Dimethylaminocyclopenta[c]coumarin,
FD-13 7-Amino-4-trifluoromethylcoumarin,
FD-14 7-Dimethylamino-4-trifluoromethylcoumarin,
FD-15 1,2,4,5,3H,6H, 10H-Tetrahydro-8-trifluoromethyl [1]benzopyrano[9,9a, 1-ghquinolinizin- 10-one,
FD-16 4-Methyl-7-(sulphometamino)coumarin sodium salt,
FD-17 7-Ethylamino-6-methyl-4-trifluoromethylcoumarin,
FD-18 7-Dimethylamino-4-methylcoumarin,
FD-19 1,2,4,5,3H,6H, 10H-Tetrahydro-carbethoxy [1]benzopyrano[9,9a, 1-ghjquinolinizino- 10-one,
FD-20 9-Acetyl-1,2,4,5,3H,6H,10H-tetrahydro[l]benzopyrano[9,9a,1-ghjquinolinizino-10-one,
FD-21 9-Cyano- 1,2,4,5,3H,6H, 10H-tetrahydro[ 1]benzopyrano[9,9a, 1-ghjquinolinizino- 10-one,
FD22  9-(t-Butoxycarbonyl)-1,2,4,5,3H,6H, 10H-tetrahydro[1]-benzopyrano[9,9a,1]-ghjquinolizino-10-one,
FD-23  4-Methylpiperidino[3,2-g]coumarin,
FD-24  4-Trifluoromethylpiperidino[3,2-g]coumarin,
FD-25  9-Carboxy-1,2,4,5,3H,6H, 10H-tetrahydro[1]benzopyrano[9,9a,1]-ghjquinolizino-10-one,
FD-26  N-Ethyl-4-trifluoromethylpiperidino[3,2-g].

Other dopants include salts of bis benzene sulphonic acid (require deposition by
spin-coating rather than sublimation) such as

![Chemical Structure]

and perylene and perylene derivatives and dopants. Other dopants are dyes such as the
fluorescent 4-dicyanomethylene-4H-pyrans and 4-dicyanomethylene-4H-thiopyrans,
e.g. the fluorescent dicyanomethylene pyran and thiopyran dyes. Useful fluorescent dyes
can also be selected from among known polymethine dyes, which include the cyanines,
complex cyanines and merocyanines (i.e. tri-, tetra- and poly-nuclear cyanines and
merocyanines), oxonols, hemioxonols, styryls, merostyryls, and streptocyanines. The
cyanine dyes include, joined by a methine linkage, two basic heterocyclic nuclei, such
as azolium or azinium nuclei, for example, those derived from pyridinium, quinolinium,
isoquinolinium, oxazolium, thiazolium, selenazolium, indazolium, pyrazolium,
pyrrolium, indolium, 3H-indolium, imidazolium, oxadiazolium, thiadiazolium,
benzoxazolium, benzothiazolium, benzoselenazolium, benzotellurazolium,
benzimidazolium, 3H- or 1H-benzoindolium, naphthoxazolium, naphthothiazolium,
naphthoselenazolium, naphthotellurazolium, carbazolium, pyrrolopyridinium,
phenanthrothiazolium, and acenaphthothiazolium quaternary salts. Other useful classes
of fluorescent dyes are 4-oxo-4H-benz-[d,e]anthracenes and pyrylium, thiapyrylium,
selenapyrylium, and telluropyrylium dyes.
Further blue-emitting materials are disclosed in the following patents, applications and publications, the contents of which are incorporated herein by reference:

US-A-5 141671 (Bryan, Kodak) - Aluminium chelates containing a phenolato ligand and two 8-quinolinolato ligands.

WO 00/32717 (Kathirgamanathan) - Lithium quinolate which is vacuum depositable, and other substituted quinolates of lithium where the substituents may be the same or different in the 2,3,4,5,6 and 7 positions and are selected from alky, alkoxy, aryl, aryloxy, sulphonic acids, esters, carboxylic acids, amino and amido groups or are aromatic, polycyclic or heterocyclic groups.

US 2006/0003089 (Kathirgamanathan) - Lithium quinolate made by reacting a lithium alkyl or alkoxy with 8-hydroxyquinoline in acetonitrile.


WO 03/006573 (Kathirgamanathan et al) - Metal pyrazolones.

WO 2004/084325 (Kathirgamanathan et al) - Boron complexes.

WO 2005/080526 (Kathitgamanathan et al) - Blue phosphorescent iridium-based complexes.

Ma et al., Chem. Comm. 1998, 2491-2492. Preparation and crystal structure of a tetranuclear zinc(II) compound \([\text{Zn}_4\text{O}(\text{AID})_6]\) with 7-azaindolate as a bridging ligand. Fabrication of inter alia a single-layer LED by vacuum deposition of this compound (<200 °C, 2 x 10⁻⁶ Torr) onto a glass substrate coated with indium-tin oxide to form a thin homogeneous film was reported.

Further electroluminescent materials which can be used include metal quinolates such as aluminium quinolate, lithium quinolate, titanium quinolate, zirconium quinolate, hafnium quinolate etc.

Many further electroluminescent materials that may be used are disclosed in WO 2004/050793 (pyrazolones), WO 2004/058783 (diiridium metal complexes), WO 2006/016193 (dibenzothiophenyl metal complexes) and WO 2006/024878 (thianthrene metal complexes), see also WO 2006/040593 the contents of which are incorporated herein by reference. Rare earth chelates, in particular may be employed as green and red
emitters. Furthermore, there may be used as electroluminescent materials, conducting polymers and conjugated polymers e.g. polyaniline, phenylene vinylene polymers, fluorene homopolymers and copolymers, phenylene polymers, as indicated below:

Further examples of dopants from the class of compounds based on rare earth chelates are:

\[
\text{Eu(DBM)}_3\text{.Phen}
\]

\[
\text{Eu(DBM)}_3\text{.OPNP}
\]

\[
\text{Eu(TTA)}_3\text{.Phen}
\]

\[
\text{Tb(TMHD)}_3\text{.Phen}
\]

\[
\text{Tb(TMHD)}_3\text{.OPNP}
\]
Furthermore, there may be used as electroluminescent materials conducting polymers e.g. phenylene vinylene polymers, fluorene homopolymers and copolymers, phenylene polymers.

5

**Thermally Activated Delayed Fluorescent Material Based Devices**

The present invention includes the use of the materials claimed in this invention in OLED devices where the emitter is a thermally activated delayed fluorescent (TDAF) material (For example see Scientific Reports: Hajime Nakantonai et al., *Scientific Reports, 3:2127,* DOI:10.1038/srep02127) the contents of which are incorporated herein by reference.
Conducting Polymers

PEDOT-PSS \((\sigma = 1 \, \text{S cm}^{-1})\) Polyaniline (PANI) \((\sigma = 1 - 10 \, \text{S cm}^{-1})\)
Polythiophene (PT) \((\sigma = 1 - 500 \, \text{S cm}^{-1})\) Polypyrrole (PPy) \((\sigma = 1 - 100 \, \text{S cm}^{-1})\)

Mixed host materials have also been disclosed in the literature and may be used in OLED devices according to the invention. Various references disclose additives and mixed hosts for OLEDs in an attempt to further improve properties. Jarikov et al, *J. Appl. Phys.*, **100**, 014901 (2006) discloses flat and rigid polycyclic aromatic hydrocarbons (PAHs) as LEL additives e.g. perylene.

Jarikov *et al* further report *J. Appl. Phys.*, **100**, pp. 094907-094907-7 (2006) perylene derivatives as light-emitting-layer (LEL) additives in organic light-emitting diodes (OLEDs). These molecules readily form emissive aggregates when added to the LEL. Addition of these polycyclic aromatic hydrocarbons increases the half-life \((t_{50})\) of undoped and doped OLEDs by 30-150 times e.g. in an Alq3+dibenzo[b,k]perylene mixed host. The authors yet further report in *J. Appl. Phys.*, **102**, 104908 (2007) a synergistic effect of a lifetime-extending light-emitting-layer (LEL) additive and improved electron injection and transport in organic light-emitting diodes (OLEDs). Di-(2-naphthyl)perylene (DNP) serves as a LEL additive said to extend the operating lifetime of OLEDs by over two orders of magnitude. Using 2-phenyl-9,10-di(2-
naphthyl)anthracene (PADN) as an electron-transport layer (ETL) and a separate layer of 4,7-diphenyl-1,10-phenanthroline (BPhen) as an electron-injection layer (EIL) the authors claimed to have significantly improved electron delivery into the charge recombination zone relative to traditional ETL made of tris(8-quinolinolate)aluminium (Alq). See also US-A-7 175922 (Jarikov et al) the disclosure of which is incorporated herein by reference.

J.C. Deaton et al (supra) disclose an a-NPB host with a "blue" aluminium quinolate as co-host and an iridium dopant. Very good yields were obtained with low concentrations of dopant for phosphorescent devices and it was found that the mixed host device provided increased power efficiency. It was hypothesized that the explanation was a reduction in the energy barrier to inject holes into the emissive layer by mixing the hole-transporting NPB having an ionization potential of 5.40 eV into the dominantly electron-transporting "blue" aluminium quinolate, having a higher ionization potential of 6.02 eV.

US-A-6392250 (Aziz et al, the disclosure of which is incorporated herein by reference.) discloses organic light emitting devices comprising a mixed region comprising a mixture of a hole transport material e.g. an aromatic tertiary amine, an electron transport material e.g. a quinolate and a dopant material. For example N,N'-di-l-naphthyl-N,N'-diphenyl-1',l'-biphenyl-l,l'-biphenyl-4,4'-diamine (NPB), and tris (8-hydroxyquinoline) aluminium (Alq₃) may be used as the hole transport material and the electron transport material, respectively and N,N'-dimethylquinacridone (DMQ), 5,6,1 1,12-tetraphenylnaphacene (Rubrene), and Nile-red dye (available from Aldrich Chemicals of Milwaukee, Wis.) may be used as dopants.

US 2002/0074935 (Kwong et al) also discloses devices with an emissive layer containing PtOEP or bis(benzo[4,5-c][1,2,5]benzothienyl-pyridinato-NAC)Iridium(III) (acetylacetonate) as a dopant and equal proportions of NPB and Alq as host materials. It is explained that the mixed host electroluminescent mixed layer serves to substantially reduce the accumulation of charge that is normally present at the heterojunction interface of heterostructure devices, thereby reducing organic material decomposition and enhancing device stability and efficiency.
In US 2004/0155238 (Thompson et al.) a light emitting layer of the OLED device contains a wide band gap inert host matrix in combination with a charge carrying material and a phosphorescent emitter. The charge carrying compound can transport holes or electrons, and it is selected so that charge carrying material and phosphorescent emitter transport charges of opposite polarity.

M. Furugori et al. in US 2003/0141809 disclose phosphorescent devices where a host material is mixed with another hole- or electron transporting material in the light emitting layer. The document discloses that devices utilizing plural host compounds show higher current and higher efficiencies at a given voltage.

T. Igarashi et al. in WO 2004/062324 disclose phosphorescent devices with the light emitting layer containing at least one electron transporting compound, at least one hole transporting compound and a phosphorescent dopant.

WO 2006/076092 (Kondakova et al., the contents of which are also incorporated herein by reference) discloses OLED device comprising a cathode, an anode, and located therebetween a light emitting layer (LEL) comprising at least one hole transporting co-host e.g. an aromatic tertiary amine such as 4,4’-Bis[N-(l-naphthyl)-N-phenylamino]biphenyl (NPB), 4,4’-Bis[N-(l -naphthyl)-N-(2-naphthyl)amino]biphenyl (TNB), 4,4’-Bis[N-(3-methylphenyl)-N-phenylamino]-biphenyl (TPD), 4,4’-Bis-diphenylamino-terphenyl or 2,6,2’,6’-tetramethyl-N,N,N’,N’ -tetraphenyl-benzoine and at least one electron transporting co-host e.g. a substituted 1,2,4-triazole such as 3-phenyl-4-(l-naphthyl)-5-phenyl-1,2,4-triazole or a substituted 1,3,5-triazine such as 2,4,6-tris(diphenylamino)-1,3,5-triazine, 2,4,6-tricarbazolo-1,3 ,5-triazine, 2,4,6-tris(N-phenyl-2-naphthylamino)- 1,3,5-triazine, 2,4,6-tris(N-phenyl- 1-naphthylamino)- 1,3,5-triazine and 4,4’,6,6’-tetraphenyl-2,2’-bi-l,3,5-triazine together with a phosphorescent emitter, wherein the triplet energy of each of the co-host materials is greater than the triplet energy of the phosphorescent emitter, and further containing an exciton blocking layer comprising a hole transporting material with triplet energy greater or equal to 2.5 eV adjacent the emitting layer on the anode side, which may be a substituted triarylamine e.g. 4,4’,4”-tris[(3-methylphenyl)phenylamino]triphenylamine (MTDATA), 4,4’,4”-tris(N,N-diphenyl-amino)triphenylamine (TDATA), N,N-bis[2,5-dimethyl-4-[(3-methylphenyl)-phenylamino]phenyl]-2,5-dimethyl-N’-(3-methylphenyl)-N’-phenyl-
1,4-benzenediamine. The devices are said to exhibit improved efficiency and reduced drive voltage.

US-A-7045952 (Lu, Universal Display Corporation) discloses an organic light emissive device comprising an emissive region disposed between and electrically connected to an anode and a cathode, wherein the emissive region comprises (i) a first single-host emissive layer, comprising a first host material, and (ii) a mixed-host emissive layer in direct contact with the first single-host emissive layer, wherein the mixed-host emissive layer comprises the first host material, and a second host material, and wherein the first single-host emissive layer and the mixed-host emissive layer each further comprise a phosphorescent emissive material.

It is believed that cells of the present kind will also operate where the light-emitting layer is based on electroluminescent quantum dots (Nanoco Group PLC) of diameter 1-20 nm, e.g. based on CdSe quantum dot cores (emission 480-640 nm) or CdS quantum dot cores (emission 380-480 nm) and cadmium-free electroluminescent quantum dots (see US7867557 Pickett; US 7588828 Mushtaq and 2011/0070443 O’Brien, the disclosures of which are incorporated herein by reference). WO 2011/0601180 (Kalzas, QD Vision, Inc) discloses the use of CdSe/CdZnS core-shell particles in a cell having inter alia a zinc oxide electron transport layer and an organic (NPB) hole transport layer. Displays based on quantum dots are expected to exhibit brighter images, lower power consumption, improved colour purity and longer life. Significantly, only a single material is needed to generate a full colour display the emission being tuneable by particle size. Because of the narrow emission bands, a display based on quantum dots are expected to have a wider colour gamut and better colour saturation than current display technologies. Quantum dot LEDs are also expected to be useful in solid state lighting. The present techniques of hole injection, hole transport, electron injection and electron transport are believed to be applicable to a quantum dot electroluminescent layer in the same way as for a current fluorescent or phosphorescent electroluminescent layer.
**Electron Transport material**

Known electron transport materials may be used, including, for example, quinolates.

Aluminium quinolate is thermally and morphologically stable to be evaporated into thin films, easily synthesized and purified and is widely used despite its problems of relatively low mobility, bandgap and tendency to ashing during sublimation. As disclosed in patent application WO 2008/0781 14, improved electron transport materials consist of or comprise zirconium or hafnium quinolate, zirconium quinolate being preferred for many embodiments.

Zirconium quinolate has a particularly advantageous combination of properties for use as an electron transport material and which identify it as being a significant improvement on aluminium quinolate for use as an electron transport material. It has high electron mobility. Its melting point (388°C) is lower than that of aluminium quinolate (414°C). It can be purified by sublimation and unlike aluminium quinolate it resublimes without residue, so that it is even easier to use than aluminium quinolate. Its lowest unoccupied molecular orbital (LUMO) is at - 2.9 eV and its highest occupied molecular orbital (HOMO) is at - 5.6 eV, similar to the values of aluminium quinolate. Furthermore, unexpectedly, it has been found that when incorporated into a charge transport layer it slows loss of luminance of an OLED device at a given current with increase of the time for which the device has been operative (i.e. increases device lifetime), or increases the light output for a given applied voltage, the current efficiency for a given luminance and/or the power efficiency for a given luminance. Embodiments of cells in which the electron transport material is zirconium quinolate can exhibit reduced turn-on voltage and up to four times the lifetime of similar cells in which the electron transport material is zirconium quinolate. It is compatible with aluminium quinolate when aluminium quinolate is used as host in the electroluminescent layer of an OLED, and can therefore be employed by many OLED manufacturers with only small changes to their technology and equipment. It also forms a good electrical and mechanical interface with inorganic electron injection layers e.g. a LiF layer where there is a low likelihood of failure by delamination. Of course zirconium quinolate can be used both as host in the electroluminescent layer and as electron transfer layer. The properties of hafnium quinolate are generally similar to those of zirconium quinolate.
Zirconium or hafnium quinolate may be the totality, or substantially the totality of the electron transport layer. It may be a mixture of co-deposited materials which is predominantly zirconium quinolate. The zirconium or hafnium may be doped as described in GB 06 14847.2 filed 26 July 2006, the contents of which are incorporated herein by reference. Suitable dopants include fluorescent or phosphorescent dyes or ion fluorescent materials e.g. as described above in relation to the electroluminescent layer, e.g. in amounts of 0.01-25 wt% based on the weight of the doped mixture. Other dopants include metals which can provide high brightness at low voltage. Additionally or alternatively, the zirconium or hafnium quinolate may be used in admixture with another electron transport material. Such materials may include complexes of metals in the trivalent or pentavalent state which should further increase electron mobility and hence conductivity. The zirconium and hafnium quinolate may be mixed with a quinolate of a metal of group 1, 2, 3, 13 or 14 of the periodic table, e.g. lithium quinolate or zinc quinolate. Preferably the zirconium or hafnium quinolate comprises at least 30 wt% of the electron transport layer, more preferably at least 50 wt%.

Other examples of electron transporters are shown below:

![QAPPI](image1)

![PyPySiPyPy](image2)
In embodiments of the invention there may be used ambipolar or electron
transporting materials either alone or as a composition co-deposited with an n-type
dopant e.g lithium quinolate or another lithium complex mentioned herein.

Compounds that may be used for this purpose have having one or two
thianthrene moieties linked to conjugated or aromatic hydrocarbon other than alkyl-
substituted fluorine.
Structures of this type (which may be ring-substituted with groups R as defined above but which do not include compounds containing fluorine alkyl-substituted at the 5-membered ring) are set out below.

(ambipolar)

(ambipolar)

(ambipolar)

(ambipolar)
(wherein \( n \) is an integer and refers to a polyphenyl chain or fused rings)

The compound 1-phenyl-2-(4-(thianthren-9-yl)phenyl)-1H-benzo[d]imidazole may be made as follows:
wherein X and Y are as defined above; e.g. X and Y are both S.

The compound may be 1-anthracenyl-9-yl-thianthrene, 1-biphenyl-4-yl-thianthrene or 9,10-Bis (1-thianthrenyl) anthrace or a mixture thereof. It may be mixed with a low work function metal complex or may be doped with a fluorescent dopant or may be doped with a phosphorescent dopant or may be doped with a rare earth chelate. For example, the compound may be mixed with either lithium quinolinolate (LiQ) or Lithium Schiff base complexes from 1 to 99% by mass, e.g. 10 to 90% by mass, e.g. 20-90% by mass, commonly 30-80% by mass. An aspect of the invention therefore comprises any of the compounds described above in combination with an n-type dopant which may be e.g. any of the lithium compounds described below for the electron injection layer.

**Electron injection material**

Any known electron injection material may be used, LiF being typical. Other possibilities include BaF$_2$, CaF$_2$, CsF, MgF$_2$ and KF.

The electron injection layer is deposited direct onto the cathode and may also comprise a compound of the formula

```latex
\text{wherein X and Y are as defined above; e.g. X and Y are both S.}
```

```latex
\text{ambipolar}
```

```latex
\text{wherein X and Y are as defined above; e.g. X and Y are both S.}
```
R_i is a 1-5 ring aryl (including polycyclic aryl or aryl-substituted polycyclic aryl), aralkyl or heteroaryl group which may be substituted with one or more C_1-C_4 alkyl or alkoxy substituents; and

R_2 and R_3 together form a 1-5 ring aryl (including polycyclic or aryl-substituted polycyclic aryl), aralkyl or heteroaryl group which may be substituted with one or more C_1-C_4 alkyl or alkoxy substituents. A compound of the above formula may be used alone or in combination with another electron injection material e.g. a quinolate such as lithium or zirconium quinolate. The Schiff base preferably comprises at least 30 wt% of the electron injection layer, more preferably at least 50 wt%.

In the formula set out above, R_i may be polycyclic aryl e.g. naphthyl, anthracenyl, tetracenyl, pentacenyl or a perylene or pyrene compound or may have up to 5 aromatic rings arranged in a chain e.g. biphenyl. It is preferably phenyl or substituted phenyl. R_2 and R_3 together may form the same groups as R_i and are preferably phenyl or substituted phenyl. Where substituents are present they may be methyl, ethyl, propyl or butyl, including t-butyl substituted, or may be methoxy, ethoxy, propoxy or butoxy including t-butoxy substituted. Particular compounds include

Cathode

In many embodiments, aluminium is used as the cathode either on its own or alloyed with elements such as magnesium or silver, although in some embodiments other cathode materials e.g. calcium may be employed. In an embodiment the cathode may comprise a first layer of alloy e.g. Li-Ag, Mg-Ag or Al-Mg closer to the electron injection or electron transport layer and a second layer of pure aluminium further from the electron injection or electron transport layer. Cells in which graphene serves as cathode are also within the invention.

How the invention may be put into effect will now be described with reference to the following examples.
Example 1.

4-(Thianthren-1-yl)triphenylamine (also diphenyl-(4-thianthren-1-yl-phenyl)-amine; (HTS-1)

To a mixture of 4-Bromotriphenyl amine (2.0 g; 0.0062 mole), 1-thianthrenylboronic acid (1.9 g; 0.0074 mole), tetrakis(triphenylphosphine) palladium (0.36 g; 0.00030 mole) in ethyleneglycoldimethyl ether (30 ml) was added potassium carbonate (4.3 g; 0.060 mole) in water (20 ml). The reaction mixture was magnetically stirred, refluxed under nitrogen atmosphere for 20 hours, allowed to cool and then filtered through a short suction column of silica gel and a pad of Hyflosuper gel. The solvent removed from the filtrate, the residue dissolved in dichloromethane and extracted with water. The organic phase was dried over anhydrous magnesium sulphate and solvent removed to give a light green residue. This was again dissolved in dichloromethane adsorbed onto silica gel and then subjected to flash column chromatography over silica gel eluting with dichloromethane. The fractions containing the product were collected together and solvent removed using a rotary evaporator to give a residue which was dissolved in diethyl ether. Addition of small amounts of petroleum ether and cooling in ice water bath gave a clear solid which was dried under vacuum at 80°C, yield 1.95g (69%). The clear solid was further purified by sublimation to give a colourless glassy solid, 1.3 g (67%), showing light bluish-purple fluorescence under UV.

Found: C 77.95, H 4.69, N 2.88, S 14.44%.
C_{30}H_{21}NS_{2}, requires C 78.40, H 4.61, N 3.05 and 13.94%.
UV: $\lambda_{\text{max}}$(CH$_2$Cl$_2$/nm ($\epsilon$/dm$^3$ mol$^{-1}$ cm$^{-1}$) 313 (22,964) and 261(30,886).

$\lambda_{\text{max}}$(Thinfilm)/nm: 317, 266 and 203 nm.

Optical band gap: 3.3 eV.

FL: $\lambda_{\text{max}}$/nm(CH$_2$Cl$_2$/em: 417; ex/nm: 350; $\lambda_{\text{max}}$/nm(Powder)/em: 409, ex/nm: 350; $\lambda_{\text{max}}$/nm(Thinfilm)/em: 400, ex/nm: 330.

TGA/°C (% weight loss): 314 (1) and 345(5).

**Example 2.**

4,4',4''-tri-(thianthren-1-yl)triphenylamine (also tris-(4-thianthren-1-y1-phenyl)-amine; HTS-2)

![Reaction Scheme]

To a mixture of tris(4-bromotriphenylamine) (2.0 g; 0.00415 mole), 1-thianthrenylboronic acid (3.6 g; 0.037 mole), tetrakis(triphenylphosphine) palladium (0.72 g; 0.0062 mole) in ethyleneglycoldimethyl ether (70 ml) was added potassium carbonate (3.0 g; 0.022 mole) in water (50 ml). The reaction mixture was magnetically stirred and refluxed under nitrogen atmosphere for 20 hours, allowed to cool and the solvent removed under reduced pressure. The residue was dissolved in dichloromethane and extracted with brine (The product was not completely soluble in dichloromethane). The organic phase was washed with water, dried over anhydrous magnesium sulphate and solvent removed to give a green solid. To the solid methanol was added, stirred for 6h and filtered off under suction. The product was washed with diethyl ether and dried under vacuum at 70 °C. Crude yield 3.15 g.
Example 3

9-(4-Bromo-phenyl)-9H-carbazole

A mixture of carbazole, 95% (5g; 0.03 mole), 4-iodo-bromobenzene, 98% (16.9 g; 0.06 mole), copper(I) iodide, 98% (2.8g; 0.015 mole) and potassium carbonate (8.3 g; 0.06 mole) in 1-methyl-2-pyrrolidinone, 99+% (60 ml) was refluxed under a nitrogen atmosphere for 20 h. After 10 minutes the reaction mixture turned blue-green in colour. The solvent was removed under reduced pressure and the residue was dissolved in 1M HCl and extracted with dichloromethane. The red dichloromethane solution was washed thoroughly with brine, water dried over anhydrous magnesium sulphate and solvent removed to give a red crystalline solid which was purified by column chromatography over silica gel (eluent CH₂Cl₂). The fractions containing the product were collected together and solvent removed to give a red solid. Trituration with petroleum ether gave an off-white solid which was suction filtered, washed with diethyl ether and dried under vacuum at 75 °C. The filtrate was evaporated and trituated with methanol-diethyl ether to give small amounts of further product. Yield 7.2 g (75 %). M.p 132 °C (DSC, onset).

9-(4-Thianthren-l-yl-phenyl)-9H-carbazole (HTS-003)
To mixture of 9-(4-Bromophenyl)-9H-carbazole (95.0 g; 0.0155 mole) and tetrakis(triphenylphosphine) palladium (0.9 g; 0.00078 mole) in dimethoxyethane (70 ml) was added thianthrene-1-boronic acid (4.4 g; 0.017 mole), followed by potassium carbonate (10.7 g; 0.078 mole) in water (50 ml). The reaction mixture was magnetically stirred and refluxed under nitrogen for 20 h during which it became green in colour. The solvent was removed and the residue was dissolved in dichloromethane (250 ml) and extracted with water. The organic phase was dried over anhydrous magnesium sulphate and solvent removed to give a dark green solid. Methanol and petroleum ether was added to the solid and the solution was magnetically stirred for 15 minutes, suction filtered and washed with diethyl ether to give an off-white solid, 5.25g. The product was purified by sublimation at 240 °C (2.5 x 10-6 torr) to give a white crystalline solid, 4.1 g (78%).

M.p 233 °C, Tg 92 °C.

Elemental analysis:

Found, C 78.70, H 4.12, N 3.07, S 14.27.
C30H19NS2 requires C 78.74, H 4.18, N 3.06 and S 14.02 %.

Example 4

4,4’-di-(thianthren-2-yl)triphenylamine (phenyl-bis(4-thianthren-1-yl-phenyl)-
amine; HTS-004)

To a mixture of 4,4'-dibromotriphenylamine (5.0 g; 0.0124 mole), 1-thianthrenylboronic acid (7.1 g; 0.0273 mole), tetrakis(triphenyl phosphine) palladium (1.4 g; 0.0012 mole) in ethyleneglycoldimethyl ether (60 ml) was added potassium
carbonate (17.2 g; 0.124 mole) in water (40 ml). The reaction mixture was magnetically
stirred and refluxed under a nitrogen atmosphere for 20 hours, allowed to cool and the
solvent was removed under reduced pressure. The residue was dissolved in
dichloromethane and extracted with water. The organic phase was dried over
anhydrous magnesium sulphate and solvent removed to give a light green residue which
was again dissolved in dichloromethane, adsorbed onto silica gel and then subjected to
flash column chromatography over silica gel eluting with dichloromethane. The
fractions containing the product were collected together and solvent removed using a
rotary evaporator to give a residue which was triturated with methanol and addition of
small amounts of petroleum ether. Cooling in an ice water bath gave a pale yellow solid
which was dried under vacuum at 80°C, yield 7.4g (88 %). It was further purified twice
by sublimation to give a light yellow glassy solid, 2.4 g (33 %), showing light bluish
fluorescence under UV. DSC did not show any melting peak, but it showed a T_g at
122 °C.

**Example 5**

**Device Structure**

A pre-etched ITO coated glass piece (10 x 10 cm²) is used. The device is
fabricated by sequentially forming layers on the ITO, by vacuum evaporation using a
Solciet Machine, ULVAC Ltd. Chigasaki, Japan. The active area of each pixel is 3mm
by 3mm. The coated electrodes are encapsulated in an inert atmosphere (nitrogen) with
UV-curable adhesive using a glass back plate. Electroluminescence studies are
performed with the ITO electrode always connected to the positive terminal. The
current vs. voltage studies are carried out on a computer controlled Keithly 2400 source meter.

Devices with green emitters are formed by the method described above consisting of an anode layer, buffer layer, hole transport layer, electroluminescent layer (DPQA doped metal complex), electron transport layer, electron injection layer and cathode layer, film thicknesses being in nm:

Device A

ITO (100 Ohms per square)/ZnTp TP (E9363, 20 nm) /Compound (HTS-1)(100 nm)/Alq₃:DPQA (40:0.1 nm)/ Alq₃ (20 nm)/Liq (6 nm)/Al

wherein DPQA is diphenyl quinacridone of formula:

![DPQA](image)

Device B

ITO (100 Ohms per square)/ZnTp TP (E9363, 20 nm) / a-NPB (100 nm)/Alq₃:DPQA (40:0.1 nm)/ Alq₃ (20 nm)/Liq (6 nm)/Al

wherein DPQA is diphenyl quinacridone.

Compared to where a-NPB (Device B) is used, devices according to the invention exhibit lower operating voltages and greater stability for devices (Device A) with HTS-1 as shown in Figures 1-6.

Similar results are obtained with the red phosphorescent device shown in Figs 7-10, the dopant in the electroluminescent layer being tris(l-phenyl-isoquinolinato-C2,N)iridium of formula:
EXAMPLE 6

4-(10H-phenothiazin-10-yl)triphenylamine (N-(4-(10H-phenothiazin-10-yl)phenyl)-
N-phenylbenzenamine; HTS-5)

To a solution of 4-bromotriphenyl amine (5.0 g; 0.0154 mole) in toluene (40 ml)
was added palladium acetate (0.35 g; 0.00156 mole). After magnetically stirring for 5
min, phenothiazine (3.4 g; 0.017 mole) in toluene (20 ml) was added followed by tert-
butyl phosphine (6.3 ml; 0.0031 mole). Finally sodium tert-butoxide (4.45 g; 0.046
mole) and toluene (20 ml) were added. The reaction mixture was magnetically stirred
and refluxed under nitrogen. After, one hour reflux the reaction mixture became dark
and viscous and therefore further toluene (50 ml) was added to the reaction mixture
which was then refluxed for a further sixteen hours. Solvent was removed under
reduced pressure and the residual oil was cooled to give a crystalline solid which was
filtered off, washed with methanol, diethyl ether and petroleum ether, and dried under
vacuum at 75°C. The solid (yield 4.1g) exhibited a bluish fluorescence. The filtrate was
evaporated; residue dissolved in small amounts of dichloromethane and passed through
a column of silica gel, after which the column was eluted in dichloromethane. Fractions
containing the product were evaporated and to the residue petroleum ether (40-60°C) was added to give an off-white solid. It was separately dried under vacuum. Yield 1.23g. Tic examination showed the presence of some phenothiazine. The crystalline solid was sublimed to give colourless to light pink solid.

The compound melts into a liquid and then sublimes. M,p 187 °C; Tg 67 °C.

Found: C, 81.29; H, 4.97; N, 6.41, and S 71.12.  
C₃₀H₂₂N₂S requires C, 81.42; H, 5.01; N, 6.33 and S 7.24 %.

UV: Xmax/nm (λ) (CH₂Cl₂); 307(28,712), and 259(46,691);
UV: Xmax/nm (thin film); 310 and 262; Band gap: 3.49 eV

FL: Xmax/nm(s) (CH₂Cl₂) 448 (excitation wavelength: 330 nm),
Xmax/nm(s) (powder) 443 (excitation wavelength: 330 nm);
Xmax/nm(s) (thin film), 449, (excitation wavelength: 330 nm).

CV: electrolyte (l00mM-Tetrabutylammonium tetrafluoroborate), analyte (1 mM), solvent-dichloromethane: HOMO: -5.5 eV and LUMO: -2.0 eV calculated from optical absorption edge of the thin film.

In this compound the HOMO level is similar to that of a-NBP but the LUMO level of 2.0 eV is low compared to that of other compounds, the LUMO level of a-NBP, for example, being 2.4 eV. For that reason in addition to potential use as a hole transport layer, the above compound may also find use as part or all of a host material in an electroluminescent layer or as part or all of an electron transport layer.

Further compounds which are analogs of the other compounds whose synthesis is reported herein or whose structures are suggested herein, and which can be made in similar manner, include

![Chemical structure images]
tris(4-(1OH-phenothiazin-10-yl)phenyl)amine

10-(4-(9H-carbazol-9-yl)phenyl)-1OH-phenothiazine

N-(4-(1OH-phenothiazin-10-yl)phenyl)-4-(1OH-phenothiazin-10-yl)-N-phenylbenzenamine

2.8-bis(1OH-phenothiazinyl)dibenzothiophene

bis [3-(phenylvinyl-1OH-phenothiazinyl)]-4,4'-biphenyl

The present phenothiazinyl compounds may be substituted on the phenothiazinyl ring(s) e.g. with phenyl, naphthyl which may be further substituted or with methyl or methoxy on phenyl rings where present.
EXAMPLE 7

HTS-7

To a mixture of 1-thianthrenylboronic acid (0.62 g; 0.0024 mole), N-(4'-iodophenyl-4-yl)-N-(m-tolyl)aniline (1.0g; 0.0022 mole) and tetrakis(triphenylphosphine) palladium (0.125g; 108 mmol) in 1,2-dimethoxy ethanol (30 ml) was added potassium carbonate (1.5g; 0.01 mole) in water (15 ml). The reaction mixture was magnetically stirred and refluxed under nitrogen for 18 hours. Solvent was removed from the reaction mixture and the residue was extracted with dichloromethane and water. The organic phase was washed with water, dried over anhydrous magnesium sulphate and solvent removed to give an oil which was purified by column chromatography using dichloromethane-petroleum ether (40-60°C) (85:15) to give light orange fractions. Eluents containing the product were evaporated to give a viscous solid which was recrystallised from methanol-petroleum ether (40-60°C) to give a yellow solid which was filtered off and dried under vacuum at 75 °C, to give 1.0 g (83 %) of the required product which was further purified by sublimation at 300°C (1.6 x 10⁻³ torr) to give a light yellow glassy solid. No melting peak was observed on DSC, but Tg was at 84 °C. It exhibited intense bluish violet fluorescence under a UV lamp.

Found: C, 81.24; H, 4.53; N, 2.71.

C₃₇H₂₇N₂S₂ requires C, 80.84; H, 4.95; N, 2.55 %.

UV: Xmax/nm (λ) (CH₂Cl₂); 344(29,362), 262(38,576) and 212(20,834);
UV: Xmax/nm (thin film); 349 and 266; Band gap: 3.15 eV
FL: Xmax/nm (ε) (CH₂Cl₂); 441 (excitation wavelength: 350 nm),
λ_{max}/nm (8) (powder) 423 (excitation wavelength: 350 nm);
X_{max}/nm (8) (thin film), 428, (excitation wavelength: 350 nm).

CV: electrolyte (100 mM-Tetrabutylammonium tetrafluoroborate), analyte (1 mM), solvent-dichloromethane: HOMO: -5.7 eV and LUMO: -2.54 eV Calculated from optical absorption edge of the thin film;

TGA/°C (% weight loss): 382 (1) and 430(5).

EXAMPLE 8

2,8-Bis(l-thianthrenyl)dibenzothiophene [HTS-8]

To a solution of 2,8-Dibromodibenzothiophene (1.0 g; 2.9 mmol) in ethyleneglycol dimethyl ether (50 ml) at 60°C was added 1-thianthreneboronic acid (1.6 g; 6.1 mmol) and tetrakis(triphenylphosphine)palladium (0.34 g; 0.29 mmol) followed by a solution of potassium carbonate (3.2 g; 23 mmol) in water (20 ml). The reaction mixture slowly darkened and became greenish in colour. It mixture was magnetically stirred and refluxed under nitrogen for 20 hours, after which solvent was removed from the cooled reaction mixture under reduced pressure. The residue was dissolved in dichloromethane (150 ml), extracted with water and then brine, after which the organic phase was dried over anhydrous magnesium sulphate. Removal of the solvent gave a greenish residue which was purified by column chromatography over silica gel using dichloromethane-petroleum ether (40-60 °C) (4:1) as eluent. A crude product was obtained by trituration with diethyl ether-petroleum ether (40-60 °C) to give an off white solid, (1.3 g; 73 %) and further purified by sublimation at 300°C (1.6 x 10⁻⁶ barr) to give a glassy solid, 0.48 g which did not show any melting peak on the DSC. Tg 133 °C.

Found: C, 70.10; H, 3.41; S, 26.53.

C_{36}H_{20}S_{5} requires C, 70.55; H, 3.29; S, 26.16 %.

UV: X_{max}/nm (ε) (CH₂Cl₂); 294(sh)(30,989), 263(97,473) and 230(35,934);
UV: X_{max}/nm (thin film); 303 and 268; Band gap: 3.85 eV
EXAMPLE 9

1-(3-Carboxaldehyde phenyl)-thianthrene

To a mixture of 3-bromobenzaldehyde (1.4 g; 7.57 mmole) and tetrakis(triphenylphosphine) palladium (0.5 g; 0.43 mmole) in tetrahydrofuran (100 ml) was added thianthrene-1-boronic acid (2.0 g; 7.69 mmole), followed by sodium hydrogen carbonate (3.0 g; 35.7 mmole) in water (50 ml). The reaction mixture was magnetically stirred and refluxed under nitrogen for 20 h. The cooled reaction mixture was poured into water and extracted with ethyl acetate. The organic phase was dried over anhydrous magnesium sulphate and solvent removed to give a crude product, which was purified by column chromatography (toluene) to give the pure product, 1.6 g (66%).

Bis [3-(phenylvinyl-1-thianthrenyl)]-4,4'-biphenyl (HTS-9)
To a solution of biphenyl-bis(methyldiethyl phosphinate) ester (0.43g; 1.04 mmol) in dimethyl sulphoxide (20 ml) was added potassium tert-butoxide (0.34 g; 3 mmol) followed by 1-(3-carboxaldehyde phenyl)-thianthrene (0.7g; 2.19 mmol). The reaction mixture was stirred at room temperature under argon atmosphere for eighteen hours and the product suction filtered and washed with methanol and petroleum ether to give a solid 0.62 g. The product was purified by sublimation to give a yellow solid, 0.3 g, M.p 259 °C (DSC), Tg 104 °C.

Elemental analysis:
Found, C 79.69, H 4.24.

C52H34S4 requires, C 79.35, H 4.3%.

Fluorescence (powder): λmax (emission): 474 nm; excitation wavelength, 340 nm.

Fluorescence (dichloromethane): Xmax (emission): 428 and 407 nm; excitation wavelength, 340 nm.

EXAMPLE 10

Cells were made having the structures shown in Figs 11-14 and gave the electro-optic properties shown in those figures. The compound HTS-8 exhibited a favourable combination of properties with performance similar to or better than a-PB and with a relatively high glass transition temperature.

EXAMPLE 11

1-Anthracen-9-yl-thianthrene (ETS-1)
To a mixture of 9-Bromoanthracene, (Aldrich, 94%) (2.0 g; 0.0078 mole), tetrakis(triphenyl phosphine) palladium (0.45 g; 0.00039 mole) in ethylene glycol dimethyl ether (50 ml) was added 1-thianthrenylboronic acid (2.4 g; 0.0092 mole) and ethyleneglycol dimethyl ether (10 ml), followed by potassium carbonate (5.4 g; 0.039 mole) in water (25 ml). The reaction mixture was magnetically stirred and refluxed under nitrogen atmosphere for 20 hours. It was allowed to cool and the solvent removed under reduced pressure. The residue was dissolved in dichloromethane and extracted with brine. The organic phase was washed with water, dried over anhydrous magnesium sulphate and solvent removed to give a residue to which methanol was added, cooled and magnetically stirred to give a light yellow solid. The product was filtered off under suction washed with methanol, petroleum ether and dried under vacuum at 70 °C. Yield 1.65 g. TLC examination (CH₂Cl₂) showed a single spot. The product was sublimed to give a light yellow solid, M.p 220 °C.

Elemental analysis:

Found, C 79.38, H 4.10, S 16.44.

C₂₀H₂₆S₂ requires, C 79.56, H 4.11 and S 16.33 %.

EXAMPLE 12

1-Biphenyl-4-yl-thianthrene (ETS-2)

To a mixture of 4-Bromobiphenyl, (Aldrich,) (2.0 g; 0.0086 mole), tetrakis(triphenyl phosphine) palladium (0.5 g; 0.00043 mole) in ethylene glycol dimethyl ether (40 ml) was added 1-thianthrenylboronic acid (2.0 g; 0.0092 mole) {actual amount should be 2.5g; 0.0094 mole} and ethyleneglycol dimethyl ether (10 ml), followed by potassium carbonate (5.4 g; 0.039 mole) in water (25 ml). The reaction mixture was magnetically stirred and refluxed under nitrogen atmosphere for 20 hours. It was allowed to cool and the solvent removed under reduced pressure. The residue was dissolved in dichloromethane and extracted with brine. The organic phase
was washed with water, dried over anhydrous magnesium sulphate and solvent removed to give a residue to which methanol was added, cooled and magnetically stirred overnight to give a light brown solid. The product was filtered off under suction washed with methanol, diethyl ether and dried under vacuum at 70 °C to give an off-white solid, 1.69 g. The product was further purified by sublimation 200 °C \( (3 \times 10^5 \text{ torr}) \) to give a colourless solid which exhibited bluish fluorescence under UV. M.p 184°C (DSC, onset).

**EXAMPLE 13**

9, 10 - Bis (1-thianthrenyl) anthracene

To a magnetically stirred solution of 9,10-dibromoanthracene (5.0 g; 0.015 mole) in ethylene glycol dimethyl ether (100 ml), tetrakis(triphenyl phosphine)palladium (1.9 g; 0.0016 mole) was added followed by 1-thianthrenyboronic acid (8.5 g; 0.033 mole). Potassium carbonate (12.4 g; 0.090 mole) in water (50 ml) was then added and the reaction mixture was refluxed under nitrogen atmosphere for 20 hours. After 15 minutes, the reaction mixture became yellow green in colour. The solvent was removed under reduced pressure and the residue was dissolved in dichloromethane and extracted with dilute acid. (The layers were not easily separable without the addition of acid).

The organic phase was washed with water, dried over anhydrous magnesium sulphate and the solvent filtered through a pad of silica gel. After the removal of the solvent, methanol was added to the residue and stirred at room temperature overnight to give a greenish black solid which was dried under vacuum at 80°C. Yield 6.0 g
TLC examination showed a single product which was purified by sublimation to give a dark yellow solid that exhibited intense yellow fluorescence under UV. It was then further purified by double sublimation. The first sublimation gave 2.1 g of the product and the second sublimation gave 1.45 g of the product. M. p 381 °C (DSC, onset), Tg 149 °C.

Elemental analysis:

Found: C 75.34; H 3.88, and S 21.44.

C38H22S4, requires: C 75.21; H 3.65, and S 21.14 %.

UV (CH2Cl2): λmax (ε/M-1 cm-1), 259(133,155), 342(4107), 359(8929), 379(15,19) and 400(14,940). UV (Thin film): λmax (Abs): 196(1.47), 266(1.45), 364(0.16), 384(0.245) and 406(0.25), Film thickness: -60 nm.

FL(CH2Cl2) λmax (em): 431, excitation wavelength: 350 nm.

FL(Powder) λmax (em): 442, 465(sh), 508 and 540(sh).

FL(Thin film) λmax (em): 422(sh), 439 and 500(sh).

CV (CH2Cl2): electrolyte: Tetrabutylammonium tetrafluoroborate (100 mM), analyte (ImM).

Optical band gap: 2.9 ev; HOMO: -6.0 eV and LUMO: -3.1 eV.

TGA/ °C (% weight loss): 400 (1) and 433(5).

Physical Properties and Fluorescence characteristics of the compounds in Examples 11-13 are summarised in the following Table.
<table>
<thead>
<tr>
<th>Code</th>
<th>Compound</th>
<th>Melting point °C</th>
<th>Tg / °C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>DSC (onset)</td>
<td></td>
</tr>
<tr>
<td>ETS-1</td>
<td><img src="image" alt="Compound" /></td>
<td>220</td>
<td>76</td>
</tr>
<tr>
<td>ETS-2</td>
<td><img src="image" alt="Compound" /></td>
<td>184</td>
<td>-</td>
</tr>
<tr>
<td>ETS-3</td>
<td><img src="image" alt="Compound" /></td>
<td>381</td>
<td>149</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Code</th>
<th>Compound</th>
<th>HOMO/eV</th>
<th>LUMO/eV</th>
<th>Band Gap/eV</th>
</tr>
</thead>
<tbody>
<tr>
<td>ETS-1</td>
<td><img src="image" alt="Compound" /></td>
<td>-5.95</td>
<td>-2.95</td>
<td>3</td>
</tr>
<tr>
<td>ETS-2</td>
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<td>-6.05</td>
<td>-2.25</td>
<td>3.80</td>
</tr>
<tr>
<td>ETS-3</td>
<td><img src="image" alt="Compound" /></td>
<td>-6.00</td>
<td>-3.10</td>
<td>2.90</td>
</tr>
</tbody>
</table>
Table 3: Photoluminescence Data

<table>
<thead>
<tr>
<th>Code</th>
<th>Compound</th>
<th>PL (λ&lt;sub&gt;max&lt;/sub&gt;, Thin film)/nm</th>
<th>PL (λ&lt;sub&gt;max&lt;/sub&gt;, Powder)/nm</th>
<th>PL (λ&lt;sub&gt;max&lt;/sub&gt;, in CH&lt;sub&gt;2&lt;/sub&gt;Cl&lt;sub&gt;2&lt;/sub&gt;)/nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>ETS-1</td>
<td><img src="image1" alt="Compound" /></td>
<td>452</td>
<td>447</td>
<td>417</td>
</tr>
<tr>
<td>ETS-2</td>
<td><img src="image2" alt="Compound" /></td>
<td>412 and 459</td>
<td>387, 409, 435</td>
<td>356 and 370</td>
</tr>
<tr>
<td>ETS-3</td>
<td><img src="image3" alt="Compound" /></td>
<td>440 and 500 (shoulder)</td>
<td>441, 508</td>
<td>430</td>
</tr>
</tbody>
</table>

**EXAMPLE 14.**

**Device Structure**

A pre-etched ITO coated glass piece (10 x 10cm<sup>2</sup>) is used. The device is fabricated by sequentially forming layers on the ITO by vacuum evaporation using a Solciet Machine, ULVAC Ltd. Chigasaki, Japan. The active area of each pixel is 3mm by 3mm. The coated electrodes are encapsulated in an inert atmosphere (nitrogen) with UV-curable adhesive using a glass backplate. Electroluminescence studies are performed with the ITO electrode always connected to the positive terminal. The current vs. voltage studies are carried out on a computer controlled Keithly 2400 source meter.

Devices with green emitters are formed by the method described above consisting of an anode layer, buffer layer, hole transport layer, electroluminescent layer (doped metal complex), electron transport layer, electron injection layer and cathode layer, film thicknesses being in nm:
Device 1.

ITO/ZnTp TP (20 nm/a-NPB (100 nm)/Alq₃:DPQA (40 nm:0.1nm)/Alq₃ (20 nm))/Liq(3 nm))/Al

wherein DPQA is diphenylquinacridone.

Device 2.

ITO/ZnTp TP (20 nm/a-NPB (100 nm)/Alq₃:DPQA (40 nm:0.1nm)/Alq₃:Liq (10 nm:10 run))/Liq(3 nm))/Al

Device 3.

ITO/ZnTp TP (20 nm/a-NPB (100 nm)/Alq₃:DPQA (40 nm:0.1nm)/ETS-001: Liq (15 nm:5 run))/Liq(3 nm))/Al

Figures 15 and 16 compare the performance of the devices as far as the current injection characteristics of the different electron transporting layer. Device 2 has Alq₃:Liq (1:1) as this is the optimum concentration. The compounds ETS-002 and ETS-003 are expected to give similar performance to ETS-001 although the ratio of ETS-002 or 3 to LiQ dopant may vary from that for ETS-001 and may have to be determined by routine trial and error.

It will be appreciated that compounds of the invention are industrially applicable. They may find use in applications in OLED displays such as microdisplays, mobile displays and TV (2-D) and TV (3-D), in OLED lighting, in devices comprising quantum dots, in light sources for light activated printing, in OLED devices where a combination of fluorescent or phosphorescent or rare earth or quantum dot emitters are employed. Devices incorporating the present compounds may be made by vacuum thermal evaporation, solution based printing (spin coating, inkjet printing), OVPD (organic vapour phase deposition) etc. OLED devices incorporating compounds of the invention may employ a combination of fluorescent or phosphorescent or rare earth or quantum dot emitters are employed. The invention also covers a mixture of hole transporters including at least one compound according to the invention and hole transporters according to the invention doped with acceptors.

Alternative names for many of the above compounds, generated automatically with ChemDraw, and which are formally correct but may be less easy to understand, appear below:

N-phenyl-N-(4-(thianthren-9-yl)phenyl)benzenamine (HTS-1),
N-phenyl-N-(3-(thianthren-2-yl)phenyl)benzenamine
N-phenyl-4-(thianthren-1-yl)-N-(4-(thianthren-9-yl)phenyl)benzenamine (HTS-4)
N-phenyl-N-(4-(thianthren-2-yl)phenyl)-4-(thianthren-3-yl)benzenamine,
4-(thianthren-1-yl)-N-(4-(thianthren-1-yl)phenyl)-N-(4-(thianthren-9-yl)phenyl)benzenamine (HTS-2),
N-(4-(thianthren-2-yl)phenyl)-4-(thianthren-3-yl)-N-(4-(thianthren-8-yl)phenyl)benzenamine
4-methoxy-N-(4-methoxyphenyl)-N-(4-(thianthren-9-yl)phenyl)benzenamine
N-(4-(2,8-dimethoxythianthren-9-yl)phenyl)-N-phenylbenzenamine
N-phenyl-N-(1-(thianthren-9-yl)naphthalen-4-yl)Naphthalen-1-amine
N-(naphthalen-1-yl)-N-(4-(thianthren-9-yl)phenyl)naphthalen-1-amine
N-(naphthalen-1-yl)-N-(1-(thianthren-9-yl)naphthalen-4-yl)naphthalen-1-amine
N,N-bis(4-tert-butylphenyl)thianthren-1-amine
2-(naphthalen-1-yl)thianthrene,
1-(phenanthren-9-yl)thianthrene,
1-(10-phenylanthracen-9-yl)thianthrene,
N-(4-methoxyphenyl)-N-phenylthianthren-2-amine,
N-phenyl-N-m-tolylthianthren-2-amine,
N2,N2,N7,N7-tetraphenylthianthren-2,7-diamine,
N2,N2,N7,N7-tetrakis(p-tolyl)thianthren-2,7-diamine,
N2,N7-diphenyl-N2,N7-di(m-tolyl)thianthren-2,7-diamine,
N2,N7-tetrakis(4-methoxyphenyl)thianthren-2,7-diamine,
N2,N7-bis(4-methoxyphenyl)-N2,N7-diphenylthianthren-2,7-diamine,
9-(4-(thianthren-1-yl)phenyl)-9H-carbazole
9-(4-(thianthren-1-yl)benzyl)-9H-carbazole
9-phenyl-3,6-di(thianthren-1-yl)-9H-carbazole
4-methyl-N-(4-(thianthren-9-yl)phenyl)-N-p-tolylbenzenamine
N-(naphthalen-3-yl)-N-phenylthianthren-2-amine
1-(3,5-di(thianthren-1-yl)phenyl)thianthrene
1-(9,10-di(naphthalen-2-yl)anthracen-6-yl)thianthrene
1-phenyl-2-(4-(thianthren-9-yl)phenyl)-lH-benzo[d]imidazole
2-phenyl-1-(4-(thianthren-1-yl)phenyl)ethanone,
4-(thianthren-1-yl)-4'-diphenylamino-1,1-biphenyl,
1-(3,5-di(thianthren-1-yl)phenyl)thianthrene,
1-(9,9-dip-tolyl-9H-fluoren-2-yl)thianthrene,
1-(4-(thiophen-2-yl)phenyl)thianthrene,
1,3,5-tris-[4-(thianthren-1-yl)-1-phenyl]benzene,
1-(2-(thianthren-1-yl)-9,9-dip-tolyl-9H-fluoren-7-yl)thianthrene,
1-(9,9-dipropyl-2-(thianthren-1-yl)-9H-fluoren-7-yl)thianthrene,
1-(4-(9-(4-(thianthren-1-yl)phenyl)-9H-fluoren-9-yl)phenyl)thianthrene,
1-(10-(thianthren-1-yl)anthracen-9-yl)thianthrene
2,7-di(thianthren-1-yl)-9,9'-spiro-bifluorene,
N-(4-(1-(4-(diphenylamino)phenyl)thianthren-9-yl)phenyl)-N-phenylbenzenamine
N1,N4-di(naphthalen-1-yl)-N1,N4-di(thianthren-1-yl)benzene-1,4-diamine
N-(4-(10H-phenothiazin-10-yl)phenyl)-N-phenylbenzenamine (HTS-5)
tris(4-(10H-phenothiazin-10-yl)phenyl)amine
10-(4-(9H-carbazol-9-yl)phenyl)-1OH-phenothiazine
N-(4-(1OH-phenothiazin-10-yl)phenyl)-4-(1OH-phenothiazin-10-yl)-N-phenylbenzenamine
2.8-bis(1OH-phenothiazinyl)dbenzothiophene
bis [3-(phenylvinyl-1OH-phenothiazinyl)]-4,4'-biphenyl
2.8-bis(l-thianthrenyl)dbenzothiophene [HTS-8]
bis [3-(phenylvinyl-l-thianthrenyl)]-4,4'-biphenyl (HTS-9)
CLAIMS

1. A compound of the formula $[\text{Ar}^1]_m[\text{Ar}^2]_n$ wherein:
   $m$ is an integer from 1-3 and $n$ is an integer and may be 1 or 2;

   $\text{Ar}^1$ represents a dibenzo[b,e][1,4]dioxine, phenoanthine, thianthrene, 
dibenzo[b,e]l,4-thiaselenane, selenanthrene or phenothiazine residue having a linkage 
to $\text{Ar}^2$ one or two positions selected from ring positions 1-4 and 6-9 and optionally
mono-, bi- or poly- substituted with $\text{Cl}_1$-$\text{C}_4$-alkyl-, $\text{Cl}_1$-$\text{C}_4$-alkoxy-, fluoro, phenyl, 
biphenyl, naphthyl, antharyl, pyridyl or thienyl which in the case of phenyl, biphenyl
naphthyl, antharyl, pyridyl or thienyl may be further substituted with $\text{Cl}_1$-$\text{C}_4$-alkyl-, $\text{Cl}_1$-$\text{C}_4$-alkoxy-
or fluoro;

   $\text{Ar}^2$ represents a residue derived from an arylamine in which the aryl rings are 
phenyl, naphthyl or anthracenyl optionally substituted with $\text{Cl}_1$-$\text{C}_4$-alkyl-, $\text{Cl}_1$-$\text{C}_4$-alkoxy-
or fluoro and in which there is aryl between nitrogen and $\text{Ar}^1$, a polycyclic fused or
chain aromatic ring system optionally containing nitrogen or sulphur and in a chain
aromatic ring system optionally containing one or more chain oxygen or sulphur atoms,
a triarylpshosphate oxide or an arylsilane the rings of any of which are optionally
substituted with $\text{Cl}_1$-$\text{C}_4$-alkyl-, $\text{Cl}_1$-$\text{C}_4$-alkoxy- or fluoro.

2. A compound according to claim 1, of the formula:

   \[
   \begin{array}{c}
   \text{Z is } N, P=0 \text{ or Si;}
   \\
   \text{X and Y which may be the same or different each represent O, S or Se, or X}
   \\
   \text{may represent NH and Y may represent O, S or Se;}
   \end{array}
   \]

   \[
   \begin{array}{c}
   \text{wherein:}
   \\
   \text{m and n}
   \end{array}
   \]
the groups R which may be the same or different independently represent hydrogen or one or more ring substituents selected from methyl, methoxy, ethyl, ethoxy, aryl (e.g. phenyl) and fluoro;

m is 1-3 when Z is N or P=0 or may be 1-4 when Z is Si;

n is 0 or 1;

p is 0, 1 or 2;

q and r are 0 or 1.

s is 1 when Z is N or P=0 or is 2 when Z is Si;

the dotted linkage when present indicates that the two phenyl rings form part of a 9H-carbazol9-yl residue

the solid line indicates a linkage between the tricyclic ring directly, by phenyl or by biphenyl; and

linkages to the tricyclic ring may be at its 1- or 2-position or may be at its 2, 7 or 2,8-positions or when X is NH may be in the 10-position.

3. A compound according to claim 1, of the formula:

\[
\text{II}
\]

wherein:

m is 1, 2 or 3, the dotted lines correspondingly representing bonds for each phenothiazine residue to one of the phenyl or naphthyl rings;

n is 1 or 2;

p, q and r are independently 0 or 1;

X and Y which may be the same or different each represent O, S or Se, or X may represent NH and Y may represent O, S or Se;

Z represents N or P=0;
the groups R which may be the same or different independently represent hydrogen or ring substituents selected from methyl, methoxy, ethyl, ethoxy, aryl (e.g. phenyl) and fluoro; and

linkages may be at the 1- or 2-position of the tricyclic ring or rings or may be at the 2, 7 or 2,8-positions or when X is NH may be in the 10-position, and may be o- or p- to the phenyl or naphthyl rings to which they are attached.

4. A compound according to claim 1 which either

(a) has one, two or three thianthrene or 10H-phenothiazin-10-yl moieties linked to conjugated or aromatic hydrocarbon other than alkyl-substituted fluorine or

(b) is of any of the formulae:
(wherein \( n \) is an integer and refers to a polyphenyl chain or fused rings)
wherein R, X and Y is as defined in claim 1.

5. A compound according to claim 1, 2 or 3, wherein m is 1.

6. Any of the following compounds:
   4-(thianthren-1-yl)triphenylamine (HTS-1);
   4,4’-di-(thianthren-1-yl)triphenylamine (HTS-4);
   4,4’,4”-tri-(thianthren-1-yl)triphenylamine (HTS-2);
   4-(thianthren-2-yl)triphenylamine;
   4,4’-di-(thianthren-2-yl)triphenylamine;
   4,4’,4”-tri-(thianthren-2-yl)triphenylamine;
   4-(1OH-phenothiazin-10-yl)triphenylamine (HTS-5);
   4,4’-di-(1OH-phenothiazin-10-yl)triphenylamine;
   4,4’,4”-tri-(1OH-phenothiazin-10-yl)triphenylamine;
   2,7-bis-(4-diphenylaminophenyl)thianthrene
   2,7-bis-(4-diphenylaminophenyl)-3,8-dimethylthianthrene
3-(thianthren-2-yl)triphenylamine;
N-(4-thianthren-1-ylphenyl)-di-p-tolylamine;
4-(thianthren-1-yl)-4',4''-dimethoxytriphenylamine;
4-(2,8-dimethoxythianthren-1-yl)triphenylamine;
1,9-di-(4-(diphenylamino)phenyl)thianthrene;
N-phenyl-N-(1-(thianthren-1-yl)naphthalen-4-yl)naphthalen-1-amine;
N-(naphthalen-1-yl)-N-(4-(thianthren-1-yl)phenyl)naphthalen-1-amine;
N-(naphthalen-1-yl)-N-(1-(thianthren-1-yl)naphthalen-4-yl)naphthalen-1-amine;
4-(N-Phenyl-N-m-tolylamino)-4'-(thianthren-1-yl)biphenyl;

7. Any of the following compounds:
4-(thianthren-1-yl)triphenylphosphine oxide
4-(thianthren-2-yl)triphenylphosphine oxide and
4-(1OH-phenothiazin-10-yl)triphenylphosphine oxide

8. Any of the following compounds
9. A compound according to claim 1, which is a mono or diarylamino compounds attached to a linear 5,10-triheterocycle.

10. Any of the following compounds:
    - N,N'-diphenylthianthrene-2-amine
    - N-phenyl-N'-w-tolylthianthren-2-amine,
    - N-(4-methoxyphenyl)-N'-phenylthianthren-2-amine,
    - N-(naphthalen-3-yl)-N'-phenylthianthren-2-amine
    - N,N'-bis(4-t-butylphenyl)thianthren-1-amine
    - N,N',N',N'-tetraphenylthianthrene-2,7-diamine,
    - N,N',N',N'-tetrakis(p-tolyl)thianthrene-2,7-diamine,
    - N,N'-diphenyl-N,N'-di(w-tolyl)thianthren-2,7-diamine,
    - N,N'-diphenyl-N,N'-di(l-naphthyl)thianthren-2,7-diamine
    - N,N',N',N'-tetrakis(4-methoxyphenyl)thianthrene-2,7-diamine,
    - N,N'-bis(4-methoxyphenyl)-N,N'-diphenylthianthren-2,7-diamine,
    - 2,7-bis-(4-diphenylaminophenyl)thianthren
    - 2,7-bis-(4-diphenylaminophenyl)-3,8-dimethylthianthren

11. A compound according to claim 1, containing an aminobiphenyl, diaminobiphenyl, 2,7-diaminofluorene, 2,7-dimaninospirobifluorene and 2,7,2', 7'tetraaminospirobifluorene residue.

12. Any of the following compounds:
    - 4-(N-phenyl-N-m-tolylamino)-4'-(thianthren-9-yl)biphenyl
bis [3-(phenylvinyl-10H-phenothiazinyl)]-4,4'-biphenyl
13. A compound according to claim 1, having one, two or three thianthrene moieties linked to conjugated or aromatic hydrocarbon other than alkyl-substituted fluorine.

14. Any of the following compounds:
   2-(naphthalen-1-yl)thianthrene
   1-(phenanthren-9-yl)thianthrene,
   1-(10-phenylanthracen-9-yl)thianthrene,
   1-(4-(thiophen-2-yl)phenyl)thianthrene
   1-(9, 10-di(naphthalen-2-yl)anthracen-6-yl)thianthrene
   10-(4-(9H-carbazol-9-yl)phenyl)-1OH-phenothiazine
   2-(10H-phenothiazinyl)dibenzothiophene
   9, 10-di(thianthren-yl)anthracene
   1,3,5-tri(thianthren-1-yl)benzene
   1,3,5-tris-(4-(thianthren-1-yl)phenyl)benzene(4(9-(4-(thianthren-1-yl)phenyl)-9H-carbazole
   9-(4-(thianthren-1-yl)benzyl)-9H-carbazole
   1-phenyl-2-(4-(thianthren-9-yl)phenyl)-IH-benzo[d]imidazole
   2-phenyl-1-(4-(thianthren-1-yl)phenyl)ethanone
   1-(9,9-di-p-tolyl-9H-fluoren-2-yl)thianthrene
The compound of claim 1, wherein one or more of the following applies:

(a) $\text{Ar}^1$ represents thianthrene;
(b) $\text{Ar}^1$ is linked to $\text{Ar}^2$ at the 1-position;
(c) $\text{Ar}^1$ is linked to $\text{Ar}^2$ at the 2-position;
(d) $\text{Ar}^1$ is linked to $\text{Ar}^2$ at the 1- and 6 or 1-and 9-positions;
(e) $\text{Ar}^1$ is linked to $\text{Ar}^2$ at the 2- and 7-positions;
(f) $\text{Ar}^2$ represents a diarylamine residue the nitrogen of which is linked to $\text{Ar}^1$, and whose rings are optionally substituted $\text{C}_1$-$\text{C}_4$-alkyl-, $\text{C}_1$-$\text{C}_4$-alkoxy- or fluoro;
(g) $\text{Ar}^2$ represents a triarylamine residue;
(h) $n.m$ is 1;
(i) $m$ is 2;
(j) 3;
(k) $\text{Ar}^2$ represents carbazole, spiro-bicarbazole or dibenzothiazole residue optionally ring-substituted with $\text{C}_1$-$\text{C}_4$-alkyl-, $\text{C}_1$-$\text{C}_4$-alkoxy- or fluoro.

16. A hole transport material comprising a compound of any preceding claim and a p-dopant which is an acceptor-type organic molecule.

17. The material of claim 16, wherein the dopant is present in an amount such that when the material is deposited to form a layer the dopant contributes about 10-40% to layer thickness.
18. The material of claim 17, wherein the dopant is present in an amount such that when the material is deposited to form a layer the dopant contributes about 33% to layer thickness.

19. The material of any of claims 16-18, wherein the dopant comprises tetracyanoquinodimethane or tetrafluorotetracyano-quinodimethane.

20. The material of any of claims 16-19, comprising a compound of any of the general formulae (i)-(vii) below:

(i)

(ii)

(iii)
wherein the groups R in any of the formulae in (i) to (vii) can be the same or different and are selected from hydrogen; substituted and unsubstituted aliphatic groups; substituted and unsubstituted aromatic, heterocyclic and polycyclic ring structures; halogens; and thiophenyl groups; and wherein
in formula (i) the methyl groups may be replaced by C\textsubscript{1}-C\textsubscript{4} alkyl or monocyclic or polyclic aryl or heteroraryl which may be further substituted e.g. with alkyl, aryl or arylamino, or of the formula viii or ix

wherein the groups R\textsubscript{1}-R\textsubscript{4} when appearing in either of the above formulae can be the same or different and are selected from hydrogen; substituted and unsubstituted aliphatic groups; substituted and unsubstituted aromatic, heterocyclic and polycyclic ring structures; halogens; and thiophenyl groups.

21. The material of claim 16, comprising a tertiary amine which is a-NPB or β-NBP.

22. An optical light-emitting diode having first and second electrodes and between said electrodes a layer comprising a compound as claimed in any of claims 1-15 or a material as claimed in any of claims 16-21.

23. The device of claim 22, wherein said layer is a hole transport layer.

24. The device of claim 22, wherein said layer is a hole injection layer.

25. The device of claim 22, 23 or 24, having an emissive layer comprising a fluorescent emitter.

26. The device of claim 22, 23 or 24, having an emissive layer comprising a phosphorescent emitter.

27. The device of claim 22, 23 or 24, having an emissive layer comprising ion fluorescent (rare earth based) emitters.
28. The device of claim 22, 23 or 24, having an emissive layer comprising quantum dots.

29. The device of claim 22, 23 or 24, having an emissive layer comprising thermally activated fluorescent (TADF) materials.

30. The device of any of claims 22-29, having a hole injection layer comprising CuPC, ZnT2P, 2-TNATA or

![Chemical Structure]

31. The device of any of claims 22-30, which forms part of a flat panel display.

32. The device of any of claims 22-30, which forms part of a lighting panel.

33. An organic photovoltaic device including a compound as claimed in any of claims 1-15 or a material as claimed in any of claims 16-21.

34. An imaging member for forming an electrostatic latent image device including a compound as claimed in any of claims 1-15 or a material as claimed in any of claims 16-21.

35. An organic thin film transistor comprising a compound as claimed in any of claims 1-15 or a material as claimed in any of claims 16-21.

36. A dye sensitised solar cell device comprising a compound as claimed in any of claims 1-15 or a material as claimed in any of claims 16-21.
37. A printed device comprising a compound as claimed in any of claims 1-15 or a material as claimed in any of claims 16-21.

38. A quantum dot based electroluminescent device comprising a compound as claimed in any of claims 1-16 or a material as claimed in any of claims 16-21.

39. An OLED device used as a light source to print conductive, resistive, dielectric, piezoelectric or pyroelectric films or lines or grids comprising a compound as claimed in any of claims 1-19 or a material as claimed in any of claims 16-21.

40. An OLED lighting panel comprising a compound as claimed in any of claims 1-15 or a material as claimed in any of claims 16-21.

41. An electro-optic or opto-electronic device having a compound having one or two thianthrene moieties linked to conjugated or aromatic hydrocarbon other than alkyl-substituted fluorine.

42. The device of claim 41, wherein the compound is of one of the formulae set out below
(wherein \( n \) is an integer and refers to a polyphenyl chain or fused rings)
wherein X and Y are both S.

43. The device of claim 41, wherein the compound is 1-anthracenyl-9-yl-thianthrene, 1-biphenyl-4-yl-thianthrene or 9,10-Bis (1-thianthrenyl) anthranc or a mixture thereof.

44. The device of claim 41, 42 or 43, wherein the compound is mixed with a low work function metal complex or is doped with a fluorescent dopant or is doped with a phosphorescent dopant or is doped with a rare earth chelate.

45. The device of any of claims 41-44, wherein the compound is mixed with either lithium quinolinolate (LiQ) or Lithium Schiff base complexes from 1 to 99% by mass.

46. The device of any of claims 41-44 wherein the compound is mixed with either lithium quinolinolate (LiQ) or lithium Schiff base complexes from 10 to 90% by mass.

47. The device of any of claims 41-44 wherein the compound is mixed with either lithium quinolinolate (LiQ) or lithium Schiff base complexes from 20 to 90% by mass.

48. The device of any of claims 41-44 wherein the compound is mixed with either lithium quinolinolate (LiQ) or lithium Schiff base complexes from 30 to 80% by mass.
49. The device of any of claims 41-48, wherein the compound is a component of an emissive layer e.g. a host.

50. The device of any of claims 41-49, wherein the compound is comprised in an electron transport layer and/or in an electron injection layer.

51. The device of any of claims 41-50, which is an OLED.

52. An optical light emitting diode device having a first electrode, a layer comprising 1-anthracenyl-9-yl-thianthrene, 1-biphenyl-4-yl-thianthrene or 9,10-bis (1-thianthrenyl) anthracene and a second electrode and optionally a low work function material.

53. The device of any of claims 41-52, having any of the following features:
   (a) the emissive layer is composed of a fluorescent emitter;
   (b) emissive layer is composed of a phosphorescent emitter;
   (c) the emissive layer is composed of ion fluorescent (rare earth based emitters);
   (d) a hole injection layer comprising CuPC, ZnTTP, 2-TNATA or
   \[
   \begin{align*}
   &\text{NC} &\text{N} &\text{N} &\text{N} &\text{N} &\text{NC} \\
   &\text{NC} &\text{C} &\text{N} &\text{C}
   \end{align*}
   \]
   (e) the hole transporting layer is doped or mixed with an electron acceptor
   (f) the hole transporting layer is doped or mixed with an electron acceptor such TCNQ or F₄TCNQ;
   (g) the hole transporter is mixed with another hole transporter;
   (h) the electron transporter is mixed with a Li, K, Cs compounds or any other compounds of low work function metals or materials;
   (i) the electron transporter is mixed with a rare earth metal or its complexes;
(j) the electron transporter is mixed with Lithium Quinolinoates (Liq) or Lithium Schiff Bases;  
(k) the electron transporter is mixed with another electron transporter or electron injector;  
(1) the electroluminescent layer comprises a metal complex;  
(m) the electroluminescent layer comprises zirconium or hafnium quinolate as host material doped with a dopant;  
(n) the electroluminescent layer comprises aluminium quinolate as the host material doped with a dopant.  
(o) the electroluminescent layer comprises an aromatic tertiary amine as host material doped with a dopant;  
(p) the electroluminescent layer comprises a light emitting material which is a metal or metalloid complex;  
(q) the electroluminescent layer comprises as luminescent material a metal quinolate, an iridium, ruthenium, osmium, rhodium, iridium, palladium or platinum complex, a boron complex or a rare earth complex;  
(r) the electroluminescent layer comprises as electroluminescent material lithium quinolate or aluminium quinolate;  
(s) the electroluminescent layer comprises a light-emitting conjugated polymer or copolymer or a dendrimer;  
(t) a hole transport layer comprising a-ΝPB.  

54. The device of any of claims 41-53, which is:  
(a) part of a flat panel display;  
(b) part of an imaging member for creation of an electrostatic latent image;  
(c) part of an OLED lighting panel  
(d) part of a photovoltaic device  
(e) part of a thin film transistor.
Fig. 1

Fig. 2
Fig. 3

Fig. 4
Fig. 5

TO/ZnTp TP (E9363, 20 nm)/HTL(100 nm)/Alq3:DPQA(40:0.1 nm)/Alq3 (20 nm)/Liq (6 nm)/Al

Fig. 6

TO/ZnTp TP (E9363, 20 nm)/HTL(100 nm)/Alq3:DPQA(40:0.1 nm)/Alq3 (20 nm)/Liq (6 nm)/Al
ITO(100Ω)/ZnTTP(20nm)/HTL(40)/α-NPB+Ir(Piq)_3 (40)/Bphen(20)/Alq_3(30)/Liq(3)/Al

Fig. 7

\[ J \, \text{mAcm}^{-2} \]

\[ \text{Voltage / V} \]

\[ 10^{-2}, 10^{-1}, 10^{0}, 10^{1}, 10^{2}, 10^{3} \]

\[ \text{Ir piq}_3 \]

\[ \text{ZnTTP} \]

ITO(100Ω)/ZnTTP(20nm)/HTL(40)/α-NPB+Ir(Piq)_3 (40)/Bphen(20)/Alq_3(30)/Liq(3)/Al

Fig. 8

\[ \text{Luminance / cd m}^{-2} \]

\[ \text{Voltage / V} \]

\[ 10^{-1}, 10^{0}, 10^{1}, 10^{2}, 10^{3} \]

\[ \text{Ir piq}_3 \]

\[ \text{ZnTTP} \]
ITO(100Ω)/E9363(20nm)/HTL(40)/a-NPB+Ir Piq (40)/Bphen(20)/Alq3(30)/E101(3)/Al

Fig. 13

Current Efficiency / cdA⁻¹

Luminance / cdm⁻²

Fig. 14

Power Efficiency / lmW⁻¹

Luminance / cdm⁻²
Fig. 15

Fig. 16