Methoxydiphenylamine-substituted fluorene derivatives as hole transporting materials: role of molecular interaction on device photovoltaic performance

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The molecular structure of the hole transporting material (HTM) play an important role in hole extraction in a perovskite solar cells. It has a significant influence on the molecular planarity, energy level, and charge transport properties. Understanding the relationship between the chemical structure of the HTM’s and perovskite solar cells (PSCs) performance is crucial for the continued development of the efficient organic charge transporting materials. Using molecular engineering approach we have constructed a series of the hole transporting materials with strategically placed aliphatic substituents to investigate the relationship between the chemical structure of the HTMs and the photovoltaic performance. PSCs employing the investigated HTMs demonstrate power conversion efficiency values in the range of 9% to 16.8% highlighting the importance of the optimal molecular structure. An inappropriately placed side group could compromise the device performance. Due to the ease of synthesis and moieties employed in its construction, it offers a wide range of possible structural modifications. This class of molecules has a great potential for structural optimization in order to realize simple and efficient small molecule based HTMs for perovskite solar cells application.

Hybrid organic–inorganic perovskite materials have shown great potential for use not only in photovoltaics but also in lasers, plasmonics, light-emitting diodes, tandems with silicon, photodetectors and sensors¹–⁴. Recently, perovskite solar cells (PSCs) have attracted considerable attention due to extremely large and rapid performance progress. In 2009, Kojima et al. demonstrated that methylammonium lead iodide (MAPbI₃) perovskite can work as a solar cell material with 3.8% power conversion efficiency (PCE)⁵. Since then published values leaped to 21.6%⁶ in only five years with a currently certified record PCE of 22.1%⁷. Such values are already exceeding commercialized polycrystalline silicon solar cells and rapidly approach crystalline silicon solar cells with record efficiency around 25.6%⁸.

In the PSC perovskite light absorber layer, either with or without mesoporous scaffold, is positioned between the electron and hole transport layers. By precisely manipulating charge carriers along the entire pathway from the absorber to both electrodes, an increase in the power conversion efficiency could be achieved. Although research is conducted on each layer, the biggest progress has been made in the area of perovskite film processing and relevant material design. Several reports show stability, performance and reproducibility improvements for the perovskite materials using mixtures of multiple cations, i.e. methylammonium, formamidinium and Cs,

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Instead of single methylammonium\textsuperscript{8,10}. Tuning the halide composition, by using an increased amount of bromide instead of iodide, has also been reported to improve thermal and humidity stability\textsuperscript{11}.

Although a number of new hole transporting materials (HTM) has been developed and investigated\textsuperscript{12}, the field is still dominated by costly 2,2',7,7'-tetrakis(N,N-di-p-methoxyphenylamine)-9,9'-spirobifluorene (Spiro-OMeTAD) and even more expensive poly[bis(4-phenyl)(2,4,6-trimethylphenyl)amine] (PTAA)\textsuperscript{13–15}. The main factors contributing to the high cost of these materials are multistep synthesis procedures, expensive reagents and costly purification methods used\textsuperscript{16}. For example, sublimation-grade Spiro-OMeTAD is needed in order to obtain high-performance devices. A significant effort has been made to develop less costly and easier to synthesize HTMs with performance comparable or better than that of Spiro-OMeTAD or PTAA\textsuperscript{17–21}. One promising avenue for simplification of the HTM synthesis procedure is substitution of the difficult to obtain 9,9'-spirobifluorene core with a simpler alternative. Methoxydiphenylamine-substituted fluorene and triphenylamine derivatives have demonstrated their effectiveness as HTMs for number of applications and they form a basis of two most effective HTMs used for PSC construction, Spiro-OMeTAD and PTAA\textsuperscript{8,13,18,22}. Therefore, it's appropriate to test the combination of the two as a new class of HTMs for application in PSCs. Additionally, it was demonstrated that structure of the HTMs has a significant influence on the molecular planarity, energy level and charge transport properties\textsuperscript{23}. Understanding the relationship between the chemical structure of the HTMs and the photovoltaic performance is crucial in continued development of the more efficient organic charge transporting materials and is imperatively needed\textsuperscript{24}. Numerous investigations are being carried out aimed at enhancement of hole drift mobility; however, until now only a few tangible recommendations have been made with regard to molecular structure modifications\textsuperscript{25–27}.

In this report a new type of HTMs (Fig. 1), based on methoxydiphenylamine-substituted fluorene and triphenylamine fragments is reported. Using molecular engineering approach, i.e. influencing conformation and packing of the molecules in the film via changing angle of rotation between fluorene and triphenylamine moieties with the help of additional methyl groups or by placing different aliphatic substituents at the para positions of the triphenylamine fragments, we have constructed a series of HTMs in order to investigate the relationship between the chemical structure of the HTMs and the photovoltaic performance. PCEs in the range of 9% to 16.8% was measured utilizing these HTMs, highlighting the importance of the optimal chemical structure, as an inappropriately placed methyl group could compromise device performance.

## Results

Synthesis of new type of hole transporting molecules involves condensation of 2,7-dibromofluorene with corresponding formyltriphenylamine, followed by a palladium-catalysed C–N cross coupling reaction with 4,4'-dimethoxydiphenylamine (Fig. 2). More detailed information on the synthesis can be found in the Supporting Information.

Thermogravimetric analysis (TGA) reveals very good thermal stability of the investigated HTMs, the fluorene derivatives start to decompose at temperatures above 410 °C (Fig. 3a, Table 1 and Supplementary Figs S1–S4).

During differential scanning calorimetry (DSC) scans it was determined that investigated materials are fully amorphous. Only the glass to liquid transition is detected and it is in the range between 85 and 109 °C (Fig. 3b, Table 1 and Supplementary Figs S5–S8). This is desired in order to form homogeneous HTM films in SC devices. $T_g$ of HTM1–3 are similar to that of Spiro-OMeTAD (126 °C), while presence of aliphatic substituents in para positions of the triphenylamine fragment in HTM4 and HTM5 reduce the glass transition temperature by ~20 °C.

The UV-vis absorption bands of the investigated HTM1–5 are shown in Fig. 4a. All studied derivatives are based on the same principal structure, i.e. fluorene core with attached 4,4'-dimethoxydiphenylamine fragments connected with a triphenylamine unit; therefore, their UV-vis spectra have significant similarities.

Presence of a fluorene fragment with attached 4,4'-dimethoxydiphenylamine moieties, the same as in Spiro-OMeTAD, results in a comparable UV-Vis absorption spectra. The π–π* absorption bands for HTM1–5 are observed at around 383 nm, indicating a similar sized π–conjugated system as that of Spiro-OMeTAD. Presence of the triphenylamine fragment results in the increase in size of the π–conjugated system and an additional absorption shoulder at 430 nm is observed\textsuperscript{28}. Position of the methyl group in the triphenylamine segment has a noticeable influence on the intensity of the shoulder. Methyl group in the meta position of the triphenylamine fragment in HTM3 causes it to be more twisted out of the plain. As a consequence shoulder at 430 nm is less intense, compared with other investigated HTMs, indicating that methyl group at this position restricts π–electron conjugation in the molecule and limits the ability of the π–electrons of triphenylamine moiety to join the common system. On the other hand, alkyl groups at para positions of the triphenylamine moiety (HTM4, HTM5) have a positive effect on the intensity of the absorption band at 430 nm. UV-Vis-NIR absorbance spectra, upon chemical oxidation of HTM4 and Spiro-OMeTAD by the silver bis(trifluoromethanesulfonyl)imide (AgTFSI)\textsuperscript{29}, are shown in the Fig. 4b.

Similarly as Spiro(TFSI)\textsubscript{2}, chemically oxidized HTM4TFSI demonstrated absorption bands at around 460 nm, 650 nm and 1470 nm indicating formation of the oxidized species. The lower intensity of the HTM4TFSI absorption maxima could be attributed to the fact that the transition is accompanied by a smaller change in the electronic charge distribution upon excitation as HTM1–5 have one less methoxydiphenylamine-substituted fluorene fragment available for oxidation.

Cyclic voltammetry (CV) was used to determine the ground-state oxidation potentials of the HTMs (Table 2). Quasi-reversible oxidation signals are observed for all investigated materials (Supplementary Figs S12–16) and differences in the CV results are quite small, although, there are some slight variations in the energy levels. A slight increase of $E_{HOMO}$ compared with HTM1, is observed for HTM2, HTM4, HTM5 due to presence of additional alkyl groups and stronger interactions between methoxydiphenylamine-substituted fluorene and triphenylamine fragments. HTM3 on the other hand, doesn’t follow a similar pattern as methyl group in the meta position.
of the triphenylamine moiety restricts π–electron conjugation in the molecule, these results correlate well with the UV-vis absorption data.

The solid-state ionization potential ($I_p$), was measured by photoelectron spectroscopy in air (PESA) method in the films of the undoped HTMs (Supplementary Figs S9 and S10 and Table 2). The structure/energy level dependency is less clearly defined in tightly packed films compared with the solvated molecules. Measured $I_p$ values are slightly lower than the HOMO levels found in the CV experiments and the difference may arise from different measurement methods and conditions (solution in CV and solid film in the photoemission method) used. Overall from the ionization potential data presented in the Table 2 it can be stated that $I_p$ values of the HTMs containing alkyl substituents in the triphenylamine fragment are lower as compared to the parent derivative HTM1. The most noticeable decrease in $I_p$ is observed for HTM4 with two methyl groups in para positions of the triphenylamine moiety. PESA and CV measurements reveal that solid-state $I_p$ and oxidation potentials in solution of the investigated compounds are similar to those of the Spiro-OMeTAD, as all of them share the same methoxydiphenylamine-substituted fluorene main fragment.

Charge-transporting properties of the investigated HTMS were determined by the xerographic time-of-flight (XTOF) technique. Supplementary Fig. S17a demonstrates representative dU/dt hole-transients for the thin film of HTM3. It exhibits a dispersive hole-transport; which, along with the strong electric-field mobility dependence, suggests the trap–dominant charge transport in this material. Nevertheless, well-defined transit times ($t_t$) established from the intersection points of two asymptotes of double-logarithmic plots provided hole-drift mobility at respective applied fields. Similar performance was also established for HTM2, HTM4, and HTM5. Hole transport

Figure 1. Structures of the investigated hole transporting materials HTM1–5.
of HTM1 is Gaussian and transit time is defined in linear plot of transients (Supplementary Fig. S17b). Examples of mobility field dependencies are given in the Supplementary Fig. S11. In all cases investigated mobility \( \mu \) may be well approximated by the formula:

\[
\mu = \mu_0 \exp(\alpha \sqrt{E})
\]

here \( \mu_0 \) is the zero field mobility, \( \alpha \) is Pool-Frenkel parameter and \( E \) is electric field strength. Such mobility dependence is explainable by terms of the Borsenberger and Weiss\(^{30}\), and Borsenberger \textit{et al.}\(^{31}\) disorder formalism. The mobility defining parameters \( \mu_0 \) and \( \alpha \) values as well as the mobility value at the 6.4 × 10⁵ V cm⁻¹ field are given in Table 2.

As seen from the results, synthesized HTMs demonstrated competitive charge mobility. XTOF measurements of the films prepared from HTM1, HTM2, HTM4, HTM5 indicate that the hole-drift mobility is ~10⁻⁵ cm² V⁻¹ s⁻¹ at weak electric fields and ~3 × 10⁻⁴ cm² V⁻¹ s⁻¹ at a field strength of 6.4 × 10⁵ V cm⁻¹. The hole mobility of these HTMs is comparable to that of Spiro-OMeTAD, while HTM3 demonstrates several orders of magnitude lower results. Most likely methyl group in the meta position of the triphenylamine fragment causes it to be more twisted out of the plane which prevents tight packing of the molecules and makes it more difficult for the charge to hop between sites.

It is well known that quality of the prepared films can have significant impact on the results of mobility measurements\(^{32}\). Every molecule has different film forming properties, therefore in order to mitigate film quality influence on mobility results we have also performed XTOF measurements with HTMs dispersed in polycarbonate (PC) polymeric matrix which minimizes HTM molecule dependent film quality variations (Fig. 5 and Table 2). Naturally, absolute mobility values are lower due to the presence of large portion of nonconductive polymer; however structure/properties relation is much better expressed in this case. Methyl group in the meta position of the triphenylamine fragment in HTM3 causes significant negative changes in the molecule’s conformation which translates into lowest charge mobility. While mobility in HTM4, HTM5, with alkyl groups at para positions of the triphenylamine moiety, is up to two orders of magnitude higher due to better optimized structure. XTOF measurement results in the HTM:polycarbonate mixtures also correlate well with the UV-Vis spectroscopy data.
The new semiconductors HTM1–5 were tested as HTMs in perovskite solar cells using a device stack of fluoro-doped tin oxide (FTO)/compact TiO2/mesoporous TiO2/perovskite/HTM/Au. Position of the methyl groups in the molecule had a noticeable impact on the performance of the semiconductors in the PSC. Arrangement of the molecules into conformations less favourable for the charge transport in HTM3 had a visible negative impact on charge mobility. Expectedly, decreased mobility had a negative effect on the performance of the HTM in the PSC as the PCE was only 9.15% (Fig. 6 and Supplementary Table S1). Change of the methyl group position from meta to ortho (HTM2) or its removal (HTM1) increases the mobility and also performance of the HTM in the PSC, PCE jumps to ~14–15%. The best results were obtained with structures containing aliphatic substituents in the para position of the triphenylamine moiety (HTM4, HTM5) which incidentally also demonstrated highest mobility in HTM:polycarbonate mixtures. PCE of 16.8% under AM 1.5 G illumination was recorded in the PSC with HTM4 (Table S1). The open-circuit voltage (Voc) was determined to be 1052 mV, current density (Jsc) 21.3 mA cm⁻² and fill factor 0.75 (Fig. 6). Similar PCE (16.5%) was also showed by HTM5. The best device prepared following the same device fabrication procedure, but using Spiro-OMeTAD as hole transporter, exhibited PCE of 17.88%.

**Conclusion**

In conclusion, a new group of small-molecule hole transporting materials, based on methoxydiphenylamine-substituted fluorene and triphenylamine fragments, is demonstrated. These HTMs are synthesized in two steps from commercially available materials. Relationship between the chemical structure of

| Compound | Tm (°C) | Tg° (°C) | Tdec (°C) | Absmax b (nm) | ε (M⁻¹ cm⁻¹) |
|----------|---------|---------|-----------|---------------|--------------|
| HTM1     | —       | 108     | 416       | 382           | 5.5 × 10⁴     |
| HTM2     | —       | 104     | 419       | 383           | 5.8 × 10⁴     |
| HTM3     | —       | 104     | 418       | 383           | 5.5 × 10⁴     |
| HTM4     | —       | 89      | 413       | 383           | 5.2 × 10⁴     |
| HTM5     | —       | 85      | 413       | 385           | 5.4 × 10⁴     |
| Spiro    | 245     | 126     | 449       | 387           | 6.9 × 10⁴     |

Table 1. Thermal and optical properties of the Spiro-OMeTAD, HTM1–5. *Determined by DSC: scan rate, 10 K min⁻¹; N₂ atmosphere; second run. bMeasured in 10⁻⁴ M THF solution.
the HTMs and the photovoltaic performance has been investigated using molecules with strategically placed aliphatic substituents. It was found that the structure of the HTMs play an important role in their hole extraction in PSCs and can have a significant influence on the molecular planarity, charge transport properties and device characteristics. We have observed that aliphatic substituents in meta position of the triphenylamine fragment cause it to be more twisted out of the plain and severely reduce charge transport properties of the HTM and overall device characteristics of the perovskite solar cells. In general, altering the structure of the phenyl ring connecting the fluorene and triphenylamine moieties produces undesirable conformational changes in the molecule and reduces its overall performance of the HTM in the PSC. On the other hand, positive effect on the performance of the hole

Figure 4. UV–vis absorption spectra of HTM1–5, Spiro-OMeTAD (a) and oxidized Spiro(TFSI)$_2$, HTM4TFSI (b) in THF (c = 10$^{-4}$M).

Table 2. Electrochemical characteristics, $I_p$, hole mobility for Spiro-OMeTAD, HTM1–5. $^*$CV measurements were carried out at a glassy carbon electrode in dichloromethane solutions containing 0.1 M tetrabutylammonium hexafluorophosphate as electrolyte and Pt wire as the reference electrode. Each measurement was calibrated with ferrocene (Fc). Conversion factors: ferrocene in DCM vs SCE 0.4634, SCE vs SHE: 0.244 35, SHE vs. vacuum: 4.43 36. $^*$Ionization potential was measured by the photoemission in air method from films. $^*$Mobility value at zero field strength. $^*$Mobility value at 6.4 × 10$^5$ V cm$^{-1}$ field strength.
transporting materials is observed after substitution at para position of the triphenylamine moiety. The overall efficiency of the investigated HTM in perovskite-based solar cells was in a range of 9% to 16.8% demonstrating the importance of the optimal molecular structure. Due to the ease of synthesis and moieties employed in its construction, it offers a wide range of possible structural modifications. The reported class of molecules has a great potential for further structure optimization in order to realize simple and efficient small molecular HTMs applied in perovskite solar cells.

Methods

Photovoltaic device preparation. Chemically etched FTO glass (Nippon Sheet Glass) was sequentially cleaned by sonication in a 2% Helmanex solution, acetone and ethanol for 30 min each, followed by a 15 min UV-ozone treatment. To form a 30 nm thick TiO₂ blocking layer, diluted titanium diisopropoxide bis(acetylatedone) (TAA) solution (Sigma-Aldrich) in isopropanol was sprayed at 450 °C. For the 200 nm mesoporous TiO₂ layer, mesoporous-TiO₂ layers were made by spin-coating a commercially available TiO₂ paste (Dyesol 30NRD). Substrates were baked at 500 °C for 30 min. Then, Li-doping of mesoporous TiO₂ is treated by spin coating a 0.1 M solution of Li-TFSI in acetonitrile at 3000 rpm for 10 s followed by another sintering at 500 °C for 20 min before the deposition of the perovskite layer. Mixed-perovskite precursor was prepared by mixing 1.15 m PbI₂, 1.10 m FAI, 0.2 m PbBr₂, 0.2 m MABr in a mixed solvent of DMF:DMSO = 4:1 (volume ratio). Perovskite solutions are successively spin-coated in the glovebox as follows: first, 2000 rpm for 10 s with a ramp-up of 200 rpm s⁻¹; second, 6000 rpm for 30 s with a ramp-up of 2000 rpm s⁻¹. Chlorobenzene (CB, 100 μl) was dropped on the spinning substrate during the second spin-coating step 20 s before the end of the procedure. Films were annealed at 100 °C for 90 min. The hole-transporting materials were applied from a 60 mM solution in chlorobenzene. All HTMs were doped with bis(trifluoromethylsulfonylimide lithium salt (Li-TFSI, Sigma-Aldrich), tris(2-(1H-pyrazol-1-yl)-4-tert-butylpyridine)-cobalt(III) tris(bis(trifluoromethylsulfonyl)imide) (FK209, Dynamo) and 4-tert-Butylpyridine (TBP, Sigma-Aldrich). The molar ratio of additives for the HTMs were: 0.5, 0.03 and 3.3 for Li-TFSI, FK209 and TBP respectively. The HTM solutions were spin-coated onto the perovskite layers at 4000 rpm for 30 s. The gold electrodes were deposited by thermal evaporation of 80 nm gold in high vacuum conditions.

Figure 5. Electric-field dependencies of the hole drift mobilities in charge-transport layers of HTM1–5 dispersed in polycarbonate polymeric matrix.

Figure 6. Current (J)-voltage (V) curves of the solar cells with HTM1–5, and Spiro-OMeTAD (control) recorded under AM 1.5 conditions (100 mW/cm²).
Photovoltaic device testing. The solar cells were measured using a 450 W xenon light source (Oriel). The spectral mismatch between AM1.5 G and the simulated illumination was reduced by the use of a Schott K113 Tempax filter (Präzisions Glas & Optik GmbH). The light intensity was calibrated with a Si photodiode equipped with an IR-cutoff filter (KG3, Schott), and it was recorded during each measurement. Current–voltage characteristics of the cells were obtained by applying an external voltage bias while measuring the current response with a digital source meter (Keithley 2400). The voltage scan rate were 5 and 10 mV s⁻¹ and no device preconditioning, such as light soaking or forward voltage bias applied for long time, was applied before starting the measurement. The starting voltage was determined as the potential at which the cells furnish 1 mA in forward bias, no equilibra-

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Acknowledgements
The authors acknowledge funding from the European Union Seventh Framework Programme [FP7/2007–2013] under grant agreement no 604032 of the MESO project. We thank E. Kamauska for his help with ionization potential measurements.

Author Contributions
V.G., M.K.N. conceived the initial idea and supervised the research. S.A. contributed to design, discussion, and improvement of all experiments. R.T. conducted the synthesis and structural characterization of the compounds. P.S., M.S. performed fabrication and characterisation of solar cells. M.D., T.M. carried out the DSC, TGA, CV, UV-Vis-NIR lights absorption measurements. J.N., V.J. carried out the ionization potential and mobility measurements. All authors contributed in writing this paper.

Additional Information
Supplementary information accompanies this paper at doi:10.1038/s41598-017-00271-z

Competing Interests: The authors declare that they have no competing interests.

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