Fused thiophenes: an overview of the computational investigations

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Abstract: Computational investigations of fused thiophenes including not only thienothiophene, dithienothiophene and thienoacene but also other thiophene-fused (hetero)-aromatic rings have been surveyed. The effect of the methods and basis sets applied on promising optoelectronic materials were elaborated. This brief review suggests the best method for the fused thiophenes to be B3LYP functional. ©2016 ACG Publications. All rights reserved.

Keywords: Computation; DFT; fused thiophenes; optoelectronic. © 2017 ACG Publications. All rights reserved.

1. Introduction

Conjugated organic compounds have been the focus of scientists due to their promising use in organic thin film transistors (TFT)1−5 and solar cells.6,7 They have less processing time and cost through their deposition from solution providing a fast and large-area fabrication.8 Fused thiophenes provide a better conjugation in the ground state and are utilized for tuning the band gap of organic materials owing to their rigid structures with an extended π-conjugation. Moreover, they enhance the intermolecular interactions in thin films which influence the molecular arrangement predominantly.9,10

Fused thiophenes comprize thienothiophenes (TTs), possessing two annulated thiophene rings, dithienothiophenes (DTTs), formed by fusing another thiophene ring to TTs, and thienoacenes having more than three thiophene units.9,10 Thiophenes were also annulated to other aromatic units, such as pyrrole, furan, benzene, pyridine, etc.

To shed light on the optoelectronic properties of organic materials, which were aimed to be used in TFT, organic light emitting diodes (OLEDs) and solar cells, their computational investigations using density functional theory (DFT) and time dependent-DFT (TD-DFT) were conducted. Estimated electronic and structure characteristics of the most appropriate candidates were compared with those collected from experimental studies to open a new avenue for the construction of highly efficient optoelectronic materials.

In this brief review, computational investigations performed to unveil the optoelectronic properties of oligomers and polymers possessing fused thiophenes will be addressed.

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2. Thiophene-Based Fused Systems

Recently, annulated thiophene systems have attracted remarkable interest, and they have been investigated theoretically and experimentally, owing to their various practical applications. Numerous publications reporting diverse aspects of the chemistry of fused thiophenes unequivocally confirm the growing interest in their material chemistry.\textsuperscript{11}

Fused thiophenes, such as thienothiophenes (TTs), possess more rigid structures than \(\alpha\)-oligothiophenes, containing two or more thiophene rings linked through a single bond. They have attracted considerable attention as they have excellent electrochemical and optical properties.\textsuperscript{12}

Oligothiophene and fused thiophene based conjugated materials could have highly \(\pi\)-conjugated electrical conductance with promising potential applications in organic light-emitting diodes (OLEDs),\textsuperscript{13-17} organic photovoltaic cells (OPVs)\textsuperscript{18-23} organic field-effect transistors (OFETs),\textsuperscript{24-27} and dye-sensitized solar cells (DSSCs).\textsuperscript{28}

Fused thiophene-based materials have emerged as one of the most promising alternatives, and led to high charge carrier mobilities exceeding 1 cm\(^2\) V\(^{-1}\) s\(^{-1}\) together with excellent stability and processability.\textsuperscript{29,30}

2.1. Structural and Electronic Properties of Fused Thiophene Based Systems and Their Application Areas

Annulation of thiophenes limits the rotational disorders between thiophene rings in favor of creating better conjugation,\textsuperscript{31} which, as a result, provides conjugated systems with low band gaps. Their stable and electron rich structures make them important building blocks for the construction of OPVs and OTFTs.\textsuperscript{32,33} DTT and its isomers demonstrated various intermolecular interactions including weak hydrogen bonding, \(\pi\)–\(\pi\) stacking, C–H interactions, and S–S interactions.\textsuperscript{34} In case of the dimer of the fused thiophene, \(\alpha,\alpha'\)-bis(dithieno[3,2-\(b\):2',3'-\(d\)]thiophene) (BDT) was found to have an unusual \(\pi\)-stacking rendering a high mobility up to 0.05 cm\(^2\) V\(^{-1}\) s\(^{-1}\).\textsuperscript{35} These endowed features allow it to be utilized in organic electronic devices.\textsuperscript{36}

Organic acceptor and donor molecules based on DTT were used for the preparation of different cation radical salts and charge-transfer complexes.\textsuperscript{37,38}

Acenes are polycyclic aromatic hydrocarbons consisting of linearly fused benzene rings. Although acenes containing five or more fused aromatic rings have been long known as promising materials for organic electronic devices,\textsuperscript{39} their instabilities against light and air hamper their use in material chemistry.\textsuperscript{40,41} Unlike acenes, thiienoacenes containing annulated thiophene rings possess higher charge mobilities and stabilities under ambient conditions, owing to their favorable electronic properties. For example, dithieno[2,3-\(d\):2',3'-\(d\)]thieno[3,2-\(b\):4,5-\(b\)]dithiophene (pentathiienoacene, PTA) has a molecular shape and crystal packing geometry similar to those of pentacene, whereas the larger band gap (up to 3.29 eV) of PTA compared to that of pentacene (1.85 eV) indicates the stability of PTA toward visible light.\textsuperscript{42}

2.2. Computational Studies on Thiophene Based Fused Systems

Although various synthetic methods are available in the literature, theoretical studies for the formation mechanism of oligo- and fused-thiophenes are rather rare. In addition, experimental and computational investigations on their electronic transport properties still need extensive effort.\textsuperscript{33,44}

Based on thorough theoretical works, the donor–acceptor (D–A) strategy among all approaches has been the most useful one applied in material chemistry such as in search of green electrochromic polymers and chromophores containing DTTs.\textsuperscript{35,45,46} Electrochemical and spectroscopic properties of the neutral and doped forms of D–\(\pi\)–A chromophores possessing DTTs were investigated by means of FT–Raman and Vis–NIR spectroscopies combined with DFT methods. These chromophores are used as electron relay \(\pi\)-centers to improve the electronic properties of \(\pi\)-conjugated molecules.\textsuperscript{45,46}
Dithieno[3,2-b:2′,3′-d]thiophene (1) and dithieno[2,3-b:3′,2′-d]thiophene (2), obtained through the annulation of bithiophenes by bridging the α,α′ and β,β′ positions with a sulfur atom had quite different LUMO characters and, therefore, significant difference between their HOMO-LUMO gaps was observed. While the former with a good conjugation had a gap of 4.51 eV, the latter one possessing a cross-conjugated system was calculated to have 5.27 eV of HOMO–LUMO gap (Figure 1, left). Moreover, HOMO–LUMO gaps and spin densities were also computed for diaryl-substituted dithieno[2,3-b:3′,2′-d]-thiophenes (3a–3d) to shed light on the electropolymerization of cross-conjugated DTTs. Taskiran Cankaya et al. performed the DFT calculations on 3a–3d bearing p-substituted phenyl units at 3- and 4-positions at UB3LYP/6-31+G(d) level (Figure 1, middle). Although the cross-conjugated DTTs had enough spin densities on α-carbons, which are best places for an electro-polymerization based on computations, all attempts for their electrochemical polymerizations ended up with failure. This result was explained by almost zero spin densities on related α-carbon atoms in dimer, which prevented the polymerization. (Figure 1, right).

Figure 1. (Left) Chemical structures of dithieno[3,2-b:2′,3′-d]thiophene (conjugated DTT, 1) and dithieno[2,3-b:3′,2′-d]-thiophene (cross-conjugated DTT, 2), and their HOMO and LUMO energy levels in eV. (Middle) Diaryl-substituted cross-conjugated DTTs 3a–3d and the range of spin densities on α-carbons. (Right) Dimer of 3a and the spin densities on α-carbons required for electropolymerization.

Comprehensive computations on dimers were performed by Li et al. at B3LYP/6-31G(d,p) level to unveil the effects of structural symmetry and linking mode on the electronic structures of the dimers, which made of accurate description of the ground-state electronic properties and electronic couplings between adjacent molecules (Figure 2, above). The predicted FMO plots and orbital energy levels of the dimers are displayed in

Figure 2 (below). Dimerization of α- and β-DTTs resulted in higher HOMOs and lower LUMOs, and consequently smaller HOMO–LUMO gaps for molecules 4 and 5, compared to their monomers. Further decrease in HOMO–LUMO gaps was achieved through incorporation of a vinylene bridge (4–6), possessing relatively smaller HOMO–LUMO gaps with respect to 4–6. The effect of substituents on dimers was probed by attachment of phenyl (8b) and 2-thienyl (8c) to α-positions of dimer 8a. The total density of states (DOS) and projected density of states (PDOS) of three dimers 8a–8c indicated that the sulfur atoms partially take part in the formation of both HOMOs and LUMOs, while the phenyl and 2-thienyl groups are not involved at all. Owing to the dihedral angles between phenyl or thienyl groups and the central π systems about 29.2° and 25.2°, respectively, the substituted phenyl or thienyl groups (8b and 8c) cannot extend the conjugation of π-system efficiently and only partly take place in the charge transport, whereas the sulfur atoms directly involved in the charge carrier transport (8a). The molecular geometries, the frontier molecular orbitals, molecular ionization potentials (IPs), electron affinities (EAs) and reorganization energies were proved to be influenced by the structural symmetry and linking mode in dimerization.
The relationship between geometric structures and electronic properties of fused thiophenes was investigated in depth by Park et al.\(^\text{49}\) (Figure 3, left). Molecular structures of all \(\alpha\)-([n]TA (thienoacenes)) and \(\beta\)-annulated ([n]TH (thienohelicenes)) thiophenes were optimized using B3LYP/6-31G(d) method, which indicated that there is a close linear relationship between HOMO–LUMO gaps \((E_g)\) and 1/Nc (Nc: number of C atoms) (Figure 3, right). The intercepts of the graphs provided the band gaps of 2.235 and 4.152 eV for TAs and THs, respectively. The slopes showed that the conjugation length affecting the HOMO–LUMO gaps \((E_g)\) is smaller in the TH system compared to that of TA (8.87 vs. 17.12). Based on the combined computational and experimental studies on oligothienoacenes with five and seven rings, Osuna et al.\(^\text{50}\) reported that these fused oligothiophenes have a larger band gap than most semiconductors currently used in the fabrication of organic field-effect transistors (OFETs). Conjugational and optical properties of pentathienoacene (5TA) and heptathienoacene (7TA) (Figure 3), analyzed by using Raman and UV-vis-NIR absorption spectroscopies, were interpreted by DFT and TD-DFT quantum chemical calculations at the B3LYP/6-31G(d,p) level, owing to the fact that DFT is well suited to model extended \(\pi\)-conjugated systems.\(^\text{51}\) It was reported that singlet excited states of 5TA and 7TA, experimentally measured at 355 nm (5TA) and 396 nm (7TA), are closely related to the calculated electronic transitions of 5TA at 349 \((f = 0.754)\) and 7TA at 397 nm \((f = 1.202)\). These results pointed out the high accuracy of TD-DFT approach for predicting the experimental wavelengths. On the other hand, DFT calculations allowed identification of the vibrational modes, associated with each of the strongest Raman features, selectively enhanced by the

**Figure 2.** (Above) Dimers of DTTs 4–9 and dihedral angles for 4–6. (Below) Their HOMO and LUMO energy levels and surfaces estimated at B3LYP/6-31G(d,p) level. "Reprinted (adapted) with permission from Li, P.; Cui, Y.; Song, C.; Zhang, H. RSC Adv., \textbf{2015}, \textit{5}, 50212–50222. Copyright 2015 The Royal Society of Chemistry (RSC)."
presence of effective π-conjugation. DFT calculations indicated totally symmetric collective vibrations mostly located on the sulfur linkages, responsible for these Raman features.

Figure 3. (Left) Chemical structures of [n]thienoacenes (TA) and [n]thienohelicenes (TH). (Right) HOMO-LUMO gaps ($E_g$) of [n]TA (a) and [n]TH (b) oligomers as a function of 1/Nc, where Nc is the number of carbon atoms along the chain. "Reprinted (adapted) with permission from Ra, C. S.; Yim, S.; Park, G. Bull. Korean Chem. Soc. 2008, 29, 891–893. Copyright 2008 Publications of the Korean Chemical Society (KCS)." 49

The influence of the length of fused thiophenes on the capabilities of the charge transfer properties of a series of multifunctional molecular devices was elaborated by Fan et al.52 Based on computations involving the combination of non-equilibrium Green’s functions (NEGF)53 and first-principle DFT, it was reported that multifunctional molecular devices can be produced when suitable substituents are attached to the fused thiophenes, and the peak position of negative differential resistance (NDR) and rectification ratio can be controlled by changing the molecular length. The molecules having thiol end groups depicted in Figure 4 (left) were fully optimized at the B3LYP/6-31G(d,p) level. These thiol subunits were used in molecular devices to place the molecules between two gold electrodes (Figure 4, right). Geometry optimizations and transport calculations for two-probe systems were done conducting DFT computations combined with Non-Equilibrium Green’s Functions (NEGF/DFT) approach. The exchange-correlation potential was described by Perdew-Burke-Ernzerh (PBE) of parametrization of the generalized gradient approximation (GGA). LDOSs (Local Density of States), MPSH (Molecular Projected Self-Consistent Hamiltonian), transmission spectra, and fermi level calculations of all two-probe systems led to the following results; a) the increase in molecular conjugation length shifts the NDR peak from the higher bias to the lower bias, b) Rectifying performance can be realized by introduction of push-pull substituent groups breaking the symmetry of molecules, c) 13c can be regard as a good candidate for multifunctional molecular device owing to its excellent rectifying performance.

Figure 4. (Left) Chemical structures of the investigated molecules, (right) a representative model of the molecular junction; large yellow, medium yellow, medium gray and small white balls represent Au, S, C and H atoms, respectively. "Reprinted (adapted) with permission from Fan, J.; Gathitu, N. N.; Chang, Y.; Zhang, J. J. Chem. Phys, 2013, 138, 074307. Copyright 2013 AIP Publishing LLC." 52

Although electropolymerization of cross-conjugated DTTs failed,48 conjugated TT and DTT possessing co-monomers linked through 4,4’-dinonyl-2,2’-bithiazole were polymerized successfully electrochemically by Sezer and Ustamehmetoğlu.54,55 They investigated the electrochemical and optical properties of these co-monomers PhTT-NBT-PhTT (14), containing 4,4’-dinonyl-2,2’-bithiazole and 3-phenylthieno[3,2-b]thiophene units, and T-NBT-Ph2DTT-NBT-
T (15), possessing 4,4'-dinonyl-2,2'-bithiazole and 3,5-diphenyldithieno[3,2-b:2',3'-d]thiophene groups (Figure 5). Their polymer films, produced via electropolymerization of co-monomers in trifluoroborate–diethyl ether and acetonitrile, were characterized experimentally, and their band gaps were obtained to be 2.04 and 1.86 eV, respectively, by in situ spectro-electrochemical measurements. Molecular orbitals and electropolymerization of the co-monomers were elaborated by means of DFT level computations. While 6-31+G(d,p) basis set with MN12-SX method, one of Meta-GGA DFT functionals, was applied on neutral molecules, unrestricted MN12-SX with the same basis set was used for their radical cations. Solvent effect was also included in the calculations with integral equation formalism of the Polarizable Continuum Model (IEFPCM). The absorption maxima of the monomers NBT and Ph₂DTT were recorded as 234, 332 nm and 260, 293 nm, respectively, which were explained according to TD-DFT level calculations at (IEFPCM:DCM)-TD-MN12-SX/6-31+G(d,p) level, providing 219, 335 nm and 260, 318 nm absorptions with main contributions from HOMO to LUMO excitations. The results obtained computationally are in well agreement with the experimentally recorded absorption values. Both optical investigations and DFT level predictions provided the band gaps of NBT-Ph₂DTT-NBT (16) and 15 as 2.76 eV and 2.64 eV, respectively. For 14, while optical band gap of 2.58 eV was obtained experimentally, theoretical studies provided the band gap of 2.44 eV. Computational studies indicated a smooth electro-polymerization starting generally from α-carbons of the explored compounds.

![Figure 5. PhTT-NBT-PhTT (14) T-NBT-Ph₂DTT-NBT-T (15) and NBT-Ph₂DTT-NBT (16) comonomers investigated for electropolymerization.](image)

Buyruk et al.\textsuperscript{56} investigated the properties of donor–acceptor (D–A) copolymers possessing TT/DTT as electron donating units and 1,2,3-triazole as an acceptor. Systematic studies on their optoelectronic and electrochemical properties provided the corresponding band gaps varying from 2.2 to 2.8 eV. (TD)-(CPCM:THF)-B3LYP/6-31G(d) level computations performed on the model compounds, which unveiled the presence of an intramolecular charge transfer transition between TT/DTT units and 1,2,3-triazole group (Figure 6).

![Figure 6. Model compounds 17–20 containing TT/DTT donors and 1,2,3-triazole acceptor for DFT computations.](image)

Instead of heteroaromatic unit such as 2,2'-bithiazole, successfully incorporated into TT and DTT molecules to make the electropolymerization feasible,\textsuperscript{54,55} Turkoglu et al.\textsuperscript{57} utilized boron as a strong acceptor in their OLED molecules. They analyzed the photophysical and
electrochemical properties of novel organoboron compounds 21–23, possessing conjugated TT as a π-conjugated spacer between mesitylboron acceptor and triphenylamine (TPA) donor units (Figure 7). The ground state geometric optimizations, optical band gaps, wavelength absorption maxima and fluorescence spectra of compounds 21–23 were investigated computationally by performing (TD)–DFT calculations at B3LYP/6-31G(d) level. While the $\lambda_{\text{max}}$ values of 21–23 were calculated to be 425, 474 and 528 nm, with the incorporation of solvent effect (THF), their $\lambda_{\text{em}}$ values were estimated as 519, 544 and 582 nm, respectively. For all the organoboron compounds 21–23, experimental data (absorption maxima: 390, 411 and 454, emission maxima: 502, 520 and 545, respectively) are well in agreement with the computational results.

![Figure 7. Organoboron compounds 21–23.](image)

Sahin et al\textsuperscript{58} used heteroatom boron as an acceptor in their seven soluble polymers 24–30 connecting substituted-TT donors (Figure 8). These compounds were reported as a first example of TT-boron copolymers emitting white color. The structural and electronic states of the polymers were elaborated by performing DFT and TD-DFT calculations on dimers and trimers at B3LYP/6-31G level. Their UV-Vis absorption spectra and vertical excitation energies were predicted using the TD–DFT method with solvent effect (THF). DFT level computations indicated an intramolecular charge transfer transitions between donor TT units and boron acceptor groups. Furthermore, the estimated absorption and emission spectra of the dimers and trimers matched well with the experimental results. For example; the UV-Vis absorption maxima of 24 measured in THF were 390 and 295 nm, while excited state electronic transitions predicted at (PCM:THF)-TD-B3LYP/6-31G(d) level were calculated as 418 and 282 nm.

![Figure 8. Polymers 24 – 30 possessing TT and mesitylboron units.](image)
Figure 9. Polymers 31–36 having DTT and DTT-S,S-dioxide units connected via mesitylboron.

Figure 10. HOMO and LUMOs (isosurface value = 0.025 au) of monomers and dimers predicted at B3LYP/6-31G(d) level in gas phase. "Reprinted (adapted) with permission from Sevinis, E. B.; Sahin, C.; Cinar, M. E.; Eroglu, M. S.; Ozturk, T. Polym. Eng. Sci., 2016, 56, 1390–1398. Copyright 2016 John Wiley and Sons."
Sevinis et al.\textsuperscript{59} applied this methodology on DTT and DTT-4,4-dioxide (DTT-\(\text{S}_2\text{SO}_2\)) to achieve their copolymers with mesitylboron for potential applications in optoelectronic devices (Figure 9). Structural and electronic states of the model compounds, which were prepared as dimers, having boron acceptor units between DTT donor groups, were unveiled by conducting DFT method in order to shed more light on their optical properties (Figure 10). While HOMOs of all the molecules, except that of dimer \textit{34}, are localized on DTT donor units, LUMOs of the dimers are exclusively located on boron acceptor atoms with small contributions of DTT groups. The UV–Vis absorption spectra and vertical excitation energies of the dimers and their emission spectra were predicted at (PCM:THF)-TD-DFT level, which are well in alignment with the experimental absorption and emission values. While optical band gaps were obtained between 2.46 and 3.21 eV for \textit{31–36}, the estimated values ranged from 2.34 to 3.25 eV.

3. Thiophene-Based Fused Heteroacenes

3.1. Structural and Electronic Properties of Thiophene-Based Fused Heteroacenes and Their Applications

One of the focus areas for the development of new materials has been continuing to be acenes and fused heteroacenes. Recent synthetic efforts have been devoted to the development of acene-based materials having better chemical stability and solution processability.\textsuperscript{60} When chemically unstable large polyacenes such as pentacene, hexacene, heptacene, etc. are transformed to thiophene-based fused heteroacenes \textit{via} thiophene rings, new modified structures simultaneously exhibit good stability and high charge carrier mobility.\textsuperscript{61,62} Thiophene based heteroacenes, compared to carbon analogue polyacenes, are expected to be more stable in air, more soluble, provide high transistor performance and good packing in solid state. In addition, they are easily modified at thiophene units.\textsuperscript{63,64}

Heteroaromatic oligomers widely used in organic-based electronic devices have received remarkable attention owing to their unique photophysical properties and their good stabilities against photo-degradation and oxidation processes.\textsuperscript{65–68} Although the synthesis of benzenannulated fused oligothiophenes, derivatives of heteroacenes, is not straightforward, they have interesting properties including their strong intermolecular interactions. They also exhibit strong performances as p-channel organic semiconductors.\textsuperscript{69,70} New thienoacene-based organic semiconductors possessing benzothiophene showed very high mobility exceeding 10 cm\(^2\) s\(^{-1}\) V\(^{-1}\), which could be expected to expand horizons of organic semiconductors.\textsuperscript{71}

Thiophene-based fused heteroacenes containing different heteroatoms such as –siloene,\textsuperscript{72,73} -borole,\textsuperscript{74,75} and -phosphole\textsuperscript{76,77} have also received considerable attention as key components of organic materials for construction of optical and electronic devices.

Some of heteroacenes derivatized from acenes and thiophene rings such as tetraceno[2,3-\(b\)\textit{]}benzo[\(d\)\textit{]}thiophene (TBT, \textit{38}), 2,5 di[thieno[3,2-\(b\)]\textit{]}benzothiophenic-2-\(\text{H}\)-thiophene (DBTBT, \textit{39})\textsuperscript{79}, bis[1]benzothieno[2,3-\(d\):2',3'-\(d'\)]benzo[1,2-\(b\):4,5-\(b'\)]dithiophene (BBTBDT, \textit{40})\textsuperscript{69} bis(naphtho[2,3-\(b\)\textit{]}thieno[2,3-\(d\):2',3'-\(d'\)]benzo[1,2-\(b\):4,5-\(b'\)]dithiophene (BNTBDT, \textit{41})\textsuperscript{80} and dibenzo[\(d,d'\)]thieno[3,2-\(b\):4,5-\(b'\)]dithiophene (DBTBDT, \textit{42})\textsuperscript{82}, are shown in Figure 11. Thienoacene co-oligomers and thiophene-fused acenes are very appealing for maximizing the overlap of \(\pi\)-orbitals by reducing the free rotation around the single bonds in thiophene oligomers, which results in a densely packed crystal structure with face-to-face \(\pi\)-stacking motifs. Recently, naphthalene, anthracene, tetracene and fluorene, etc., have been successfully introduced into oligothiophene \textit{via} co-oligomerization. They were used as excellent stable semiconductor materials with relatively high band gaps and low energy level HOMOs.
3.2. Computational Studies on Thiophene-Based Fused Heteroacenes

In parallel with the recent experimental work on thiophene based fused heteroacenes, theoretical efforts have indeed started to become a major source of valuable information that complements experimental studies, thereby contributes to the understanding of the molecular electronic structures as well as the optoelectronic properties of the materials.

Sokolov et al demonstrated that novel derivatives of the dinaphtho[2,3-b:2',3'-f]thieno-[3,2-b]thiophene with high hole mobility and air stability can be used as a high performance semiconductor (Figure 12). Geometries of the neutral ground state and radical cations were optimized at B3LYP/6–31G(d,p) level. To shed light on the crystal structure, which is an important feature for OFET applications, properties of the isolated molecules in a unit cell were calculated by Perdew–Burke–Ernzerhof parametrization with the double numeric plus polarization basis (PBE/DNP) and B3LYP/6–311+G(d,p) methods. The screening procedures were performed for compounds 44–50, which are potential derivatives of compound 43, such as (i) computational screening for small reorganization energy possessing molecules, (ii) optimization of crystal structure, (iii) calculation on charge transport providing relative mobilities, (iv) picking up the potential molecules having the highest mobility, (v) realizing the synthesis and (vi) fabrication of the device and investigations of its properties. According to computational results, 2 was a potential candidate owing to its higher transfer integral. Therefore, its synthesis and characterization were performed for single-crystal OFET devices, which provided saturation region hole mobilities as high as 12.3 cm² V⁻¹ s⁻¹ and a linear region mobility of 16.0 cm² V⁻¹ s⁻¹.

Electron rich benzodithiophene, coupled with electron deficient TT isomer, namely thieno[3,4-b]thiophene via a single bond, was also considered in material chemistry. Borges et al reported structural and electronic properties of the ground and lowest excited states of poly-(thieno[3,4-b]thiophenebenzodithiophene) (51) by help of high-level theoretical investigations (Figure 13). Although TDDFT is an efficient method for computing the electronic spectra of such large π-conjugated polymers, its excited charge transfer (CT) state description was considered to be poor. To amend this feature, the algebraic diagrammatic construction method to second order, ADC(2), was introduced, which is a combination of ab initio wave-function-based polarization propagator method with resolution of identity (RI) method. ADC(2) was used to predict data for excited state energies, oscillator strengths and bond length alternation (BLA) analysis. The results
obtained from ADC(2) were compared with TD-DFT outcomes at B3LYP, CAM-B3LYP, BHandHLYP, PBE and LC-wPBE levels. With regard to excitation energies, CAM-B3LYP and BHandHLYP provided the best matching with the results estimated using ADC(2).

Simone et al. investigated a series of fused thiophene–pyrrole ring systems, so-called S,N heteroacenes (52–58, SNn, n = 6–12, 14, Figure 14) by means of DFT methods at different exchange–correlation functionals. Frequencies and molecular geometries of SN6–SN10 (52–57) were calculated at the M06, B3LYP-D3, PBE03, wB97XD, CAM-B3LYP and OPBE exchange–correlation functionals with all the valence 6–31G(d) basis set. Percentage errors are listed in Table 1. The electronic spectra of neutral, radical cation and dication species have been obtained using TDDFT. According to computational results; (i) absolute error of excitation energies in all exchange–correlation functionals, except PBE03, increased with the molecular size, (ii) percentage errors obtained from wB97XD, OPBE and CAMB3LYP calculations were higher than the others, (iii) M06, PBE0 and B3LYP-D3 reproduced the experimental data very well with percentage errors varying from 0.26 to 2.90. This excellent agreement was constant along all the considered molecules, (iv) the ionization potentials decreased along the series reaching the lower value for 57, (v) N-alkylated substituents in the synthesized molecules did not affect significantly the absorption properties.

Fused thiophenes comprising thienothiophene, dithienothiophene, thienoacene, and other thiophene based fused heteroacenes have been the interest of scientists owing to their promising properties in material chemistry. The experimental studies including their design, synthesis, characterization and determinations of their properties have been performed extensively.

Table 1. Percentage errors obtained from TD-DFT methods with 6-31G(d) basis set compared to experimental absorption wavelengths (nm).

|       | M06  | B3LYP-D3 | PBE03 | OPBE | wB97XD | CAM-B3LYP | Exp  
|-------|------|----------|-------|------|--------|------------|-------
| 52 (SN6) | 0.26 | 0.26     | 2.90  | 6.33 | 9.76   | 8.18       | 379   |
| 53 (SN7) | 1.28 | 1.02     | 2.55  | 9.20 | 11.5   | 9.46       | 391   |
| 54 (SN7_1) | 1.01 | 2.02     | 1.77  | 10.1 | 10.6   | 8.86       | 395   |
| 55 (SN8) | 0.96 | 2.17     | 1.45  | 10.9 | 10.6   | 8.95       | 413   |
| 56 (SN9) | 2.14 | 2.38     | 1.66  | 12.6 | 12.4   | 10.0       | 420   |
| 57 (SN10) | 1.60 | 2.75     | 1.15  | 13.1 | 12.4   | 9.86       | 436   |

4. Conclusion

Fused thiophenes comprising thienothiophene, dithienothiophene, thienoacene, and other thiophene based fused heteroacenes have been the interest of scientists owing to their promising properties in material chemistry. The experimental studies including their design, synthesis, characterization and determinations of their properties have been performed extensively.
Nevertheless, to shed more light on their characteristics and open a new avenue for their proper use, computational investigations have been conducted. Especially, DFT, which is known to be an efficient and relatively cheap method compared ab initio tools, has been selected in numerous research fields to unveil and provide back up for experimental outcomes. Among DFT methods B3LYP functional is generally employed due to the good agreement between predicted and the experimental results.

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