Exploration of Nonlinear Optical Properties for the First Theoretical Framework of Non-Fullerene DTS(FBTTh2)2-Based Derivatives

Muhammad Usman Khan,* Shabbir Hussain, Muhammad Adnan Asghar, Khurram Shahzad Munawar, Rasheed Ahmad Khera, Muhammad Imran, Mohamed M. Ibrahim, Mahmoud M. Hessien, and Gaber A. M. Mersal

ABSTRACT: Organic compounds having significant nonlinear optical (NLO) applications are being employed in the optoelectronics field. In the current work, a series of non-fullerene acceptor (NFA) based compounds are designed by modifying the acceptors with different substituents using DTS(FBTTh2)2R1 as a reference compound. To study the NLO responses to the tuning of various acceptors, DFT and TD-DFT based parameters were calculated at the M06 level along with the 6-31G(d,p) basis set. The designed compounds (MSTD2–MSTD7) showed smaller values of the energy gap in comparison to the reference compound. The energy gaps of the title compounds were linked to global reactivity insights; MSTD7 provided a lower band gap, with smaller and larger quantities for hardness and softness characteristics, respectively. Further, UV–vis analyses were performed for all of the designed compounds, displaying wavelengths red-shifted from that of DTS(FBTTh2)2R1. The intraelectron transfer (ICT) process and stability of the title compounds were explored via frontier molecular orbital (FMO) and natural bond orbital (NBO) studies, respectively. Out of all the designed compounds, the highest value of linear polarizability ($\alpha$) of 3.485 $\times$ 10$^{-22}$ esu, first hyperpolarizability ($\beta_{total}$) of 13.44 $\times$ 10$^{-32}$ esu and second-order hyperpolarizability ($\gamma$) of 3.66 $\times$ 10$^{-31}$ esu were exhibited by MSTD7. In short, all of the designed compounds exhibited promising NLO properties because of their low charge transport resistance. These NLO properties may be useful for experimental researchers to uncover NLO materials for modern applications.

INTRODUCTION

Nonlinear optically active chromophores have received more consideration in view of their importance in many technological applications in recent years. Organic-based scaffolds have a broad band spectrum, nonlinearity, and fast response time features. They are also extensively adapted for the storage of optical data, processing of images, and optical switching applications.1–11 A comparative study between the organic and inorganic nonlinear chromophores reveals that organic systems have many preferable properties such as a short response time, large molecular plasticity, outstanding processing performance, and high nonlinear polarization rate.5–12 To achieve the required nonlinear optical characteristics, researchers have concentrated on synthesizing organic molecules with certain geometries and electronic molecular characteristics.13–15

The delocalization of $\pi$ electrons in the primary framework causes polarization in organic NLO based materials.16–18 In an organic molecule, electrons flow toward the donor via a $\pi$ linker to the acceptor, consequently leading to intermolecular charge transfer (ICT).19,20 Indeed, changing the molecular structure to generate high-performance nonlinear optical materials is a smart method.21–23 As a result, organic compounds with a large conjugated delocalized $\pi$ electronic density may be employed as the preferred species for second- and third-order nonlinear optics materials.24–26 It has also been observed from the data reported earlier that D (donor) and A (acceptor) moieties are responsible for providing the required ground-state charge asymmetry.27–29

Among the different series of delocalized electronic conjugated frameworks, fullerene and its conjugated derivatives are good candidates as has been reported by their NLO output.30–33 Fullerenes exhibit a reasonable NLO response due to $\pi$ conjugation a general charge delocalization.33,34 Despite the fact that fullerene acceptors have been the most popular materials for two decades, the restricted electrical characteristics and moderate absorbing capacity of fullerene...
derivatives in the visible region has influenced the evolution of organic photovoltaic devices.35

In the past decade, non-fullerene acceptors (NFAs) have received a great deal of attention, primarily for research into the improvement of organic solar cells. To a large extent, in this area they have taken the spotlight away from fullerene derivatives. The organic acceptors, which replace the typical fullerene acceptors present in the photoactive layers of standard organic solar cells, have various advantages, including light absorption, donor−acceptor combination variety, and massive acceptor material manufacturing.36 For this, a great number of NFA species have been studied in the past decade for an exploration of their electro- and photoelectrochemical properties due to the strong electron transfer abilities of such derivatives.37 On the basis these fascinating features of π-conjugated NF-based compounds, we selected NF-based compounds in this work to explore their NLO properties from a theoretical perspective. To our knowledge, NLO research on the title NF-based molecule and its developed compounds has yet to be published. Hence, to overcome the research gap, calculations based on density functional theory (DFT) have been carried out to explore NLO properties. The NFA molecule DTS(FBTTh2)2R1 has been taken as a reference, and different NF-based acceptor compounds have been designed via coupling of various acceptors and π-conjugated linkers. The current work is extremely important for estimating the NLO applications of organic compounds. In addition to this, it helps to realize the consequences of the donor, various acceptors, and π linkers of non-fullerene-based compounds. In order to understand the NLO properties, DFT and TD-DFT calculations were performed to calculate the natural bonding orbitals (NBOs), global reactivity parameters (GRPs), frontier molecular orbitals (FMOs), densities of states (DOSs), and UV−vis absorption spectra. NFA molecules are considered to play a substantial role in the arena of NLO.

■ COMPUTATIONAL STUDY

The current work was completed employing the Gaussian 09 program38 at the M06 level of theory and the 6-31G(d,p) basis set. In dichloromethane (DCM) solvent, the DFT calculations were carried out to determine the data for FMO, NBO, the density of state (DOS), absorption spectra, and NLO properties of fullerene-free molecules having a D−π−D−A−π−A configuration. An FMO analysis was performed to determine the band gap which enables an estimation of the least amount of energy needed for the transition from the HOMO to the LUMO. An NBO analysis was employed to determine the hyperconjugative interactions and intramolecular charge transfers. The DOS calculations were used to determine the scattering of energy states. UV−visible absorption spectra were obtained to investigate electronic transitions. The dipole moment (μ), linear polarizability ⟨α⟩, first-hyperpolarizability (β), and second-hyperpolarizability (γ) values were calculated by employing eqs 1−4.39,40

\[
\mu = (\mu_x^2 + \mu_y^2 + \mu_z^2)^{1/2}
\]

(1)

\[
\langle\alpha\rangle = 1/3(\alpha_{xx} + \alpha_{yy} + \alpha_{zz})
\]

(2)

\[
\beta_{\text{total}} = (\beta_x^2 + \beta_y^2 + \beta_z^2)^{1/2}
\]

(3)

Tensors of β in three directions are described in eqs S1−S3 in the Supporting Information.

\[
\gamma_{\text{tot}} = (\gamma_x^2 + \gamma_y^2 + \gamma_z^2)^{1/2}
\]

(4)

The results were obtained by utilizing Gauss View,41 Avogadro,42 and ChemCraft43 programs from the output files.

■ RESULTS AND DISCUSSION

The present work focuses on the NLO characteristics of non-fullerene-based organic scaffolds. The no-fullerene-based reference compound DTS(FBTTh2)2R1 has a D−π−A−D−A−π−D type configuration that contains three fragments: donors, π spacers, and acceptors. The branches C8H17 (n-octyl) and C6H13 (n-hexyl) in DTS(FBTTh2)2R1 have been altered with CH3 (methyl) at the donor part to overcome the computational cost, as shown in Figure 1.
The study of frontier molecular orbitals (FMOs) is extremely significant for an understanding of the optical and electric characteristics of species, and it is also utilized to investigate their chemical stability. The highest occupied molecular orbitals are referred to as HOMOs, while the lowest unoccupied molecular orbitals are referred to as LUMOs, which are very significant FMOs. An FMO analysis enables us to determine significant quantum chemistry parameters, including electronic properties, chemical stabilities, electron transfer properties, and reactivities, of the investigated compounds. The energy difference \( E_{\text{gap}} = E_{\text{LUMO}} - E_{\text{HOMO}} \) of compounds is directly associated with their kinetic stability and chemical reactivity. The \( E_{\text{gap}} \) values also control factors, including electronegativity, electrophilicity, delicate quality, hardness, reactivity, and softness. The lower the value of energy difference (\( \Delta E \)), the higher the value of polarizability, resulting in a magnificient NLO value. Table 1 gives the \( E_{\text{gap}} \) values for the compounds under investigation.

| compound       | \( E_{\text{HOMO}} \) | \( E_{\text{LUMO}} \) | \( \Delta E \) |
|---------------|----------------|----------------|----------|
| DTS(FBTTh)\(_2\)\(R_1\) | -5.119 | 2.788 | 3.331 |
| MSTD2         | -5.238 | -3.403 | 1.835 |
| MSTD3         | -5.248 | -3.528 | 1.720 |
| MSTD4         | -5.240 | -3.463 | 1.777 |
| MSTD5         | -5.234 | -3.416 | 1.818 |
| MSTD6         | -5.265 | -3.774 | 1.491 |
| MSTD7         | -5.250 | -3.668 | 1.582 |

It is concluded from the data presented in Table 1 that DTS(FBTTh)\(_2\)\(R_1\) has a theoretically estimated energy difference (\( \Delta E \)) value, i.e., 2.331 eV, greater than the \( E_{\text{gap}} \) values obtained in the designed molecules. The donors, acceptors, \( \pi \) spacers, and endchain acceptors are involved in the configuration of the designed compounds, which are the reasons for the reduction of their energy difference (\( \Delta E \)) values. The \( E_{\text{gap}} \) value for MSTD2 decreased to 1.835 eV because of the structural modification with \( 7\)-(2,2′-bithiophen-5-yl)-4-(4,4-dimethyl-6-(4-((9-methyl-9H-carbazol-2-yl)[2,2′-bithiophen]-5-yl)phenyl)-4H-silolo[3,2-b:4,5-b′]dithiophen-2-yl)-5-fluorobenzo[c][1,2,5]thiadiazole as donors and \( \pi \) spacers and 3-dinitromethylen-1-one (DNMI) as an acceptor unit. In MSTD3 and MSTD4, \( E_{\text{gap}} \) values further decreased to 1.720 and 1.777 eV by the introduction of two and one chloro species on the acceptor unit DNMI, respectively, in which two chloro groups were found to be more effective in decreasing the band gap in comparison to one chloro group. This \( E_{\text{gap}} \) value in MSTD5 slightly increased to...
1.818 eV due to the introduction of one fluoro group on the acceptor part (DNMI). This increase in the $E_{\text{gap}}$ value was perhaps due to less enhancement of the resonance effect induced due to a fluoro group as compared to a chloro group. As the fluoro and chloro groups are electron donating due to resonance effects, the compounds MSTD3, MSTD4, and MSTD5 have decreases in $E_{\text{gap}}$ values by enhancement of resonance and electron withdrawal toward the acceptor region.59,60

Furthermore, MSTD6 and MSTD7 were also found to have smaller $E_{\text{gap}}$ values in comparison to DTS(FBTTh$_2$)$_2$R1 and MSTD1–MSTD5 due to the presence of nitro groups in the DNMI acceptor. The presence of two and three nitro groups in MSTD6 and MSTD7, respectively, reduced the $E_{\text{gap}}$ values to 1.491 and 1.582 eV. The nitro group has a strongly electron attracting character, and its electron-attracting ability is a consequence of a strong negative inductive activity.61 The observed lowest $E_{\text{gap}}$ value of MSTD6 among all designed compounds is because of the extensive conjugation involved in the biphenyl rings and the end-capped acceptor part comprising nitro units.

The ascending order of $E_{\text{gap}}$ of the title compounds is MSTD6 < MSTD7 < MSTD3 < MSTD4 < MSTD5 < MSTD2 < DTS(FBTTh$_2$)$_2$R1. Overall, MSTD2–MSTD7 have small energy gaps in comparison to DTS(FBTTh$_2$)$_2$R1. This decrease in band gap can be attributed to the presence of electronegative species (fluoro, chloro, and nitro groups) on the acceptor units, which can effectively remove electronic charge density from other portions, lowering the band gap (Table 1). The potential of the designed compounds to perform intramolecular charge transfer (ICT) from donor to acceptor units through $\pi$ linkers is further effectively found in frontier molecular orbital studies as coupled with the $E_{\text{gap}}$ values.

The $E_{\text{gap}}$ values define the ICT experience from the donor to the acceptor component, which is aided by the $\pi$ spacer, which gives a deep insight into the related functional associations of the NLO structures.62,63 The contour surfaces of FMOs are utilized to define the transfer of electronic charges, as shown in Figure 2. In the reference compound (DTS(FBTTh$_2$)$_2$R1), the highest occupied molecular orbital electronic density is mostly present over the entire molecule, donors, acceptors, and $\pi$ spacers, while the LUMO is mostly located on acceptors and $\pi$ spacers and to some extent on the donor unit. In MSTD2–MSTD5, the electronic density of the LUMO is only found on the acceptor unit and partially on the second $\pi$ linker. The electronic population in the HOMO is found mostly on 4-(6-(4-(2,2′-bithiophen)-5-yl)-4,4-dimethyl-4H-silolo[3,2-b:4,5-b′][dithiophen-2-yl])-5-fluorobenzo[c][1,2,5]thiadiazole (on the first $\pi$ spacer, second donor, and first acceptor). In addition to this, in MSTD6 and MSTD7 the electronic density of the LUMO is located perfectly over the various terminal acceptor moieties, while in the HOMO, the charge density is largely positioned over 4-(6-(4-(2,2′-bithiophen)-5-yl)-4,4-dimethyl-

Figure 2. HOMO–LUMO structures of the reference DTS(FBTTh$_2$)$_2$R1 and its derivatives.
4H-silolo[3,2-b:4,5-b’]dithiophen-2-yl)-5-fluorobenzo[ε]-[1,2,5]thiadiazole (on the first π-spacer, second donor, and first acceptor). The NLO response is assumed to be dependent on charge transfer from the donor to the acceptor via the π spacer. MSTD2–MSTD7 may be considered as suitable NLO candidates as a result of this effect.

**Global Reactivity Parameters.** The HOMO–LUMO energy gap may be used to describe the reactivity and stability of the designed compounds using global reactivity parameters (GRP). The energy gap of HOMOs (\(E_{\text{gap}} = E_{\text{LUMO}} - E_{\text{HOMO}}\)) may be used to calculate global reactivity parameters such as ionization potential (IP), electron affinity (EA), electronegativity (\(\chi\)),66,67 global hardness (\(\eta\)),66 electrophilicity index (\(\omega\)), chemical potential (\(\mu\)), and global softness (\(\sigma\)).65

The investigated compounds have lower global softness values (\(\sigma\)) in comparison to their chemical hardness (\(\eta\)) and electrophilicity index (\(\omega\)) values. The highest \(\sigma\) value is exhibited by MSTD7, i.e., 15.4371 \(E_h\) whereas DTS-(FBTTh2)2R1 has the lowest value of 11.6777 \(E_h\). In all examined compounds, the increasing trend of global softness (\(\sigma\)) values is reported as DTS(FBTTh2)2R1 < MSTD5 < MSTD6 < MSTD7 < DTS(FBTTh2)2R1. The highest value of chemical hardness (\(\eta\)) is shown by DTS(FBTTh2)2R1 (0.0428 \(E_h\)) which decreases to 0.0324 \(E_h\) in MSTD7. The descending order of chemical hardness is represented as MSTD7 < MSTD3 < MSTD2 < MSTD6 < MSTD5 < MSTD4 < DTS(FBTTh2)2R1.

**Natural Bond Orbital (NBO) Analysis.** An NBO analysis helps in the detection of hydrogen bonds, originating from hyperconjugated interactions in the designed molecules.\(^{32,66}\) It provides a true image for investigating the intramolecular delocalization and the shifting of electronic charge density from the donor to the acceptor region of the system in D–π–A architectures.\(^{36,66}\) With a larger value of the energy of stabilization, the interaction between the donor and acceptor moieties is very significant.\(^{70}\) Equation 12 may be used to represent the stabilization energy formula using a second-order perturbation approach:

\[
E^{(2)} = q^2 F_{ij}^2 / (\epsilon_j - \epsilon_i)
\]

where \(E^{(2)}\) describes the stabilization energy, \(q\) represents the occupancy of the donor orbital, \(F_{ij}\) shows the off-diagonal NBO Fock matrix elements, and \(\epsilon_j\) and \(\epsilon_i\) signify the diagonal elements.\(^{72}\) For the discovery of intra-atomic and subatomic holding and connection among bonds, an NBO examination is an effective procedure and a helpful premise for researching conjugative collaboration or charge movement in subatomic frameworks.\(^{70}\) An NBO analysis for DTS(FBTTh2)2R1 and MSTD2–MSTD7 has been carried out and elaborated in Tables S10–S16 in the Supporting Information, and some selected values are given in Table 3.

Results cited in Table 3 indicate that the maximum value of a probable transition was observed in DTS(FBTTh2)2R1: i.e., π(C11–C12) → π*(C13–N20) at 26.73 kcal/mol has been detected to be the most reliable transition in directing the

| Compound | IP (eV) | EA (eV) | \(\chi\) (eV) | \(\eta\) (eV) | \(\mu\) (eV) | \(\sigma\) (eV) |
|----------|---------|---------|---------------|-------------|-------------|-------------|
| DTS(FBTTh2)2R1 | 1.8811 | 0.1025 | 0.1453 | 0.0428 | -0.1453 | 0.2465 | 11.6775 |
| MSTD2 | 0.1753 | 0.1056 | 0.1405 | 0.0349 | -0.1405 | 0.2828 | 14.3339 |
| MSTD3 | 0.1792 | 0.1042 | 0.1417 | 0.0337 | -0.1417 | 0.2677 | 13.3295 |
| MSTD4 | 0.1819 | 0.1033 | 0.1423 | 0.0389 | -0.1423 | 0.2600 | 12.8473 |
| MSTD5 | 0.1794 | 0.1038 | 0.1416 | 0.0378 | -0.1416 | 0.2652 | 13.2288 |
| MSTD6 | 0.1845 | 0.1143 | 0.1494 | 0.0351 | -0.1494 | 0.3181 | 14.2489 |
| MSTD7 | 0.1866 | 0.1218 | 0.1542 | 0.0324 | -0.1541 | 0.3669 | 15.4371 |

MSTD7 has the highest IP value among the proposed compounds, 0.1866 \(E_h\) (Table 2), indicating that the donor moiety efficiently distributes the electronic charge density to the acceptor, while MSTD2 has the lowest IP value of 0.1753 \(E_h\). Overall, the ascending order of the ionization potential is MSTD2 < MSTD3 < MSTD5 < MSTD4 < MSTD6 < MSTD7 < DTS(FBTTh2)2R1. The highest value of chemical hardness (\(\eta\)) is shown by DTS(FBTTh2)2R1 (0.0428 \(E_h\)) which decreases to 0.0324 \(E_h\) in MSTD7. The descending order of chemical hardness is represented as MSTD7 < MSTD3 < MSTD2 < MSTD6 < MSTD5 < MSTD4 < DTS(FBTTh2)2R1.

All values are given in hartree (\(E_h\)) units.
maximum stabilization energy and a powerful association of donor (π) and acceptor (π*) units.

With a stabilization energy value of 0.68 kcal/mol, the transition from π(C5−C6) to π*(C5−C6) was termed as the least stable. Similarly, transitions such as σ(N20−S51) → σ*(C12−C13) and σ(C7−C8) → σ*(C6−S9) have larger and smaller stabilization energy values of 8.32 and 0.50 kcal/mol, respectively. A weak contact between an electron donor (σ) and acceptor (π*) causes the outputs of these stabilization energies.

When the phenomenon of resonance is taken into account, electronic interactions such as LP2(S9) → π*(C5−C6) with the highest value of the energy of stabilization, 27.05 kcal/mol, respectively. A weak contact between an electron donor (σ) and acceptor (π*) causes the outputs of these stabilization energies.

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In compound MSTD2, the transition $\pi(C1-C2) \rightarrow \pi^*(C83-C113)$ has the highest stabilization energy of 26.69 kcal/mol while the transition $\pi(N115-O117) \rightarrow \pi^*(C90-C91)$ has the lowest energy of 0.53 kcal/mol. Hence, $\sigma(C1-C2) \rightarrow \sigma^*(C36-C37)$ and $\sigma(N115-O117) \rightarrow \sigma^*(C29-S32)$ transitions in MSTD2 had the largest stabilization energy of 8.35 kcal/mol and the lowest stabilization energy of 0.51 kcal/mol, respectively. As indicated in Table 3, the LP2(S9) $\rightarrow \sigma^*(N115-O117)$ transition due to resonance has an energy of stabilization of 177.19 kcal/mol, whereas the LP(1)(S9) $\rightarrow \pi^*(N115-O116)$ transition in MSTD2 has a stabilization energy of 21.34 kcal/mol.

The compound MSTD3 showed the transition $\pi(C60-C61) \rightarrow \pi^*(C83-C113)$ had the maximum value of the stabilization energy (27.99 kcal/mol) and the $\pi(N115-O117)$

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→ $\pi^*(C88-C89)$ transition had the lowest value of stabilization energy (0.66 kcal/mol).

For the $\sigma(C26-C27) \rightarrow \pi^*(C29-S32)$ transition, MSTD3 has the lowest stabilization energy value of 0.51 kcal/mol due to the weaker degree of association between $\sigma$(donor) and $\pi^*$(acceptor). In addition to this, LP(3)(O116) → $\pi^*(N115-O117)$ and LP2(O87) → $\pi^*(C77-C86)$ also have high and low stabilization energies of 177.93 and 21.89 kcal/mol, respectively.

For MSTD4, the transition $\pi(C60-C61) \rightarrow \pi^*(C83-C116)$ has a maximum value of stabilization energy of 27.48 kcal/mol, while $\pi(C87-C88) \rightarrow \pi^*(C87-C88)$ has the lowest value of stabilization energy of 0.52 kcal/mol. The highest and lowest energy values, 8.34 and 0.5 kcal/mol, exhibited by MSTD4 are due to $\sigma(S50-N51) \rightarrow \pi^*(C36-C37)$ and $\sigma(C1-C2) \rightarrow \pi^*(C4-S10)$ transitions, respectively. The highest stabilization energy values (177.6 and 21.76 kcal/mol) are because of resonance, which is shown in LP(3)(O94) → $\pi^*(N92-O93)$ and LP2(O86) → $\pi^*(C77-C85)$ transitions, respectively (Table 3).

The most prominent transition for $\pi \rightarrow \pi^*$ exhibited by MSTD5 is $\pi(C60-C61) \rightarrow \pi^*(C83-C116)$ with a stabilization energy of 27.18 kcal/mol, while $\pi(C87-C88) \rightarrow \pi^*(C87-C88)$ has an energy value of 0.58 kcal/mol shown to have the lowest value of stabilization energy. Furthermore, the $\sigma(S50-N51) \rightarrow \sigma^*(C36-C37)$ transition exhibited the maximum stabilization energy, 8.34 kcal/mol; on the other hand, $\sigma(C1-C2) \rightarrow \sigma^*(C4-S10)$ with an energy value of 0.5 kcal/mol is found to have the minimum value of the stabilization energy. However, the highest 177.56 kcal/mol and the lowest 14.35 kcal/mol stabilization energy values are shown by LP(3)(O94) → $\pi^*(N92-O93)$ and LP2(O90) → $\pi^*(C88-N89)$ transitions because of resonance phenomena.

A similar trend of stabilization energies is observed in MSTD6, with the maximum probable $\pi \rightarrow \pi^*$ transition being $\pi(C60-C61) \rightarrow \pi^*(C83-C117)$ with a stability value of 30.34 kcal/mol. Furthermore, the transition $\pi(C37-C38) \rightarrow \pi^*(C37-C38)$ with an energy value of 0.62 kcal/mol is shown to have the lowest stability. The transitions $\sigma(S50-N51) \rightarrow \sigma^*(C36-C37)$ and $\sigma(C21-C22) \rightarrow \sigma^*(C20-S25)$ have the highest and lowest energy values of 8.36 and 0.51 kcal/mol, respectively, because of enervated connections between $\sigma$(donor) and $\sigma^*$(acceptor). In addition to this, other electronic transitions such as LP(3)(O105) → $\pi^*(N104-O106)$ and LP2(O102) → $\pi^*(C81-N101)$ had stabilization energies of 186.34 and 15.08 kcal/mol, respectively, due to resonance, as shown in Table 3.

The highest stabilization energy value for the $\pi(C60-C61) \rightarrow \pi^*(C83-C118)$ transition is 32.77 kcal/mol, whereas the lowest stabilization energy value for $\pi(C110-C111) \rightarrow \pi^*(C110-C111)$ transition is 0.71 kcal/mol in MSTD7. Due to the presence of weak interactions, 9.48 and 0.50 kcal/mol are found for the transitions $\sigma(C118-H119) \rightarrow \pi^*(C60-S66)$ and $\sigma(C26-C27) \rightarrow \sigma^*(C29-S32)$, respectively.

Due to resonance, the transitions LP(3)(O90) → $\pi^*(N89-O91)$ and LP2(O90) → $\pi^*(C88-N89)$ have the highest stabilization energy values of 179.75 and 14.68 kcal/mol. Tables S10–S16 demonstrate that more transitions have been investigated in DTS(FBTTh$_2$)$_2$R1 and MSTD2–MSTD7. Furthermore, the results showed that the $\pi$ electron bonding orbitals of C-C delocalized antibonding orbitals play a significant role in the ring stability and overall structural design. As a result, it is safe to say that extended conjugation is found in MSTD2–MSTD7 and that it is necessary for the stabilization of these systems and giving matchless NLO features.

**Density of State (DOS).** The distinct kinds of states inhabited by electrons at a quantized energy level, i.e., the different kinds of electronic states per unit volume per unit energy, are known as the densities of state (DOSs). An extraordinary high value of the DOS reveals that different kinds of states are accessible for energy levels. The zero value along the axis illustrates that no state is available for electron excitation. DOS calculations may be used to determine the general scattering of energy levels as a function of energy, as well as the energy gap.73 The DOS plots of DTS(FBTTh$_2$)$_2$R1 and MSTD2–MSTD7 at the M06/6-31G(d,p) level of theory are given in Figure 3.

For reference compound DTS(FBTTh$_2$)$_2$R1 the contribution by the acceptor group seems to be 43.2% to the HOMO and 14.7% for the LUMO. Here, donors contribute 44.1%, 44.2%, 44.1%, 44.4%, 43.3%, and 43.7% to the HOMO and 2.0%, 1.7%, 2.5%, 2.6%, 4.3% and 9.2% to the LUMO in MSTD2–MSTD7, respectively. The acceptor contributes 10.9%, 10.6%, 10.5%, 10.7%, 10.0%, and 11.6% to the HOMO and 76.1%, 76.4%, 75.8%, 74.9%, 77.1%, and 83.2% to the LUMO for MSTD2–MSTD7, respectively. The $\pi$ linkers contribute 45%, 45.2%, 45.3%, 44.8%, 46.7%, and 44.9% to the HOMO and 21.9%, 21.9%, 21.7%, 22.6%, 18.6%, and 16.4% for the LUMO for MSTD2–MSTD7, respectively.

These contributions show that the design of different compounds by varying different efficient acceptor moieties has a leading role in the transmittance of an electronic charge cloud in different ways. Each compound is split into three respective segments: i.e., donor, $\pi$ spacer, and acceptor. For compound MSTD2, the highest charge density of the HOMO is found to be between −7 and −12 eV and it seems to be on the $\pi$ spacer and the highest density of the LUMO is found to be −1.5 eV by the donor group, which confirms the efficient charge transfer from the donor to the acceptor group through the $\pi$ spacer. In the case of MSTD3, the highest HOMO density is found to be −7 eV, which is 45.2% on the $\pi$ spacer, while the highest LUMO density is found to be on the acceptor which makes a 76.4% charge contribution. For MSTD4 and onward, a maximum HOMO density of around −7 to −14 eV is due to the acceptor and the maximum LUMO density of around 2–4 eV is restricted by the $\pi$ spacer, which increases the electron-withdrawing nature and also extends the conjugation. All of the results from a DOS analysis strongly support the same results computed from an FMO analysis.

**UV–Vis Analysis.** UV–vis spectroscopy offers further evidence with regard to different types of transitions in the compounds, such as maximum absorption wavelength ($\lambda_{max}$), transition energy ($E$), and oscillator strength ($f_{os}$).74 These parameters are determined by TD-DFT calculations at the M06/6-31G(d,p) level in the gaseous phase and are given in Table 4, while further findings are presented in Tables S21–S27. A pictographic illustration of the absorption shifts of DTS(FBTTh$_2$)$_2$R1 and MSTD2–MSTD7 is given in Figure 4.

Table 4 shows that the energy and maximum absorption wavelengths ($\lambda_{max}$) are inversely proportional to each other.75 The calculations of absorption spectra of designed compounds (MSTD2–MSTD7) and reference compound DTS(FBTTh$_2$)$_2$R1 at M06/6-31G(d,p) level of theory were performed. The maximum absorption wavelength ($\lambda_{max}$), oscillator strength ($f_{os}$), and excitation energy, calculated in
Table 4. TD-DFT-Calculated Transition Energies (eV), Maximum Absorption Wavelengths (λ_{max} nm), Oscillator Strengths (f_{nm}) and Transition Natures of the Designed Compounds

| Compounds          | λ   | E   | f_{nm} | MO contributions                  |
|--------------------|-----|-----|--------|-----------------------------------|
| DTS(FBTTh2)_2R1    | 711.408 | 1.743 | 2.092 | H → L (94%), H1 → L+1 (5%)        |
| MSTD2              | 810.190 | 1.530 | 1.589 | H → L (83%), H1 → L (8%)          |
| MSTD3              | 859.563 | 1.442 | 1.396 | H → L (87%), H1 → L (7%)          |
| MSTD4              | 833.726 | 1.487 | 1.479 | H → L (85%), H1 → L (8%)          |
| MSTD5              | 818.264 | 1.515 | 1.592 | H → L (84%), H1 → L (8%)          |
| MSTD6              | 780.997 | 1.588 | 1.228 | H → L (27%), H1 → L+1 (52%)       |
| MSTD7              | 930.247 | 1.333 | 1.009 | H → L (91%), H1 → L (5%)          |

Figure 4. Absorption plots of DTS(FBTTh2)_2R1 and MSTD2–MSTD7.

The λ_{max} values of the compounds under investigation range from 711.408 to 930.247 nm. The maximum absorption was observed at 711.408 nm for the reference compound, whereas the absorption values of MSTD2–MSTD7 are 810.190, 859.563, 833.726, 818.264, 780.997, and 930.247 nm, respectively. MSTD7 has a λ_{max} value of 930.247 nm, and there is a small value of the energy gap (1.582 eV) from DTS(FBTTh2)_2R1 because of the presence of an end-capped electron-withdrawing acceptor group.

The increasing order of the designed molecules on the basis of λ_{max} is MSTD6 < MSTD2 < MSTD5 < MSTD4 < MSTD3 < MSTD7. It is observed that the reference molecule DTS(FBTTh2)_2R1 has the lowest λ_{max} value among the designed molecules. Experimentally, the absorption spectra of reference molecule obtained correlate with the simulated results. It is obvious from the results that electron-withdrawing end-capped acceptor moieties have a significant effect on λ_{max} values. These electron-deficient moieties are responsible for the red shift in absorption spectra. Absorption band lengths of MSTD2–MSTD7 ranges from 780.997 to 930.247 nnnm which are red-shifted relative to the reference molecule DTS(FBTTh2)_2R1. The designed compounds MSTD3 and MSTD7 have a greater red shift with the lowest excitation energies of 1.442 and 1.333 eV, respectively. Due to the small energy gap, charge transfer is feasible. The λ_{max} values of MSTD2, MSTD4, and MSTD5 are closely related, having an oscillation strength of ~1.5. The excitation energies of MSTD2–MSTD7 are 1.530, 1.442, 1.487, 1.515, 1.588, and 1.333 eV, respectively, which are lower in comparison to that of the reference compound DTS(FBTTh2)_2R1: i.e., 1.743 eV.

As a result of the above explanation, it can be inferred that the newly proposed molecules have better optoelectronic characteristics in comparison to the reference molecule. It is believed that all of the newly designed compounds have good excitation energies, oscillator strengths, and absorptions. Thus, MSTD2–MSTD7 can be used as significant non-fullerene acceptor moieties in photovoltaic devices.

Nonlinear Optical (NLO) Properties. The nonlinear optical (NLO) response of organic compounds with adequate electronic communication across their various moieties is rather good.76–77 In research communities, several experimental and computational scientists have focused on the fields of material science, physics, and chemistry.85,86 Herein, Table 5 shows the computed μ_{total}, ⟨α⟩, β_{total} and ⟨γ⟩ values of the designed compounds.

Table 5. Dipole Moment, Average Polarizability, First Hyperpolarizability, Second-Order Hyperpolarizability, and Major Contributing Tensors (esu)

| compound          | μ_{total} × 10^{-27} | ⟨α⟩ × 10^{-22} | β_{total} × 10^{-27} | ⟨γ⟩ × 10^{-31} |
|--------------------|----------------------|----------------|----------------------|----------------|
| DTS(FBTTh2)_2R1    | 0.3                  | 2.123          | 0.008118             | 0.403          |
| MSTD2              | 5.26                 | 3.098          | 6.763                | 1.45           |
| MSTD3              | 6.13                 | 3.277          | 8.730                | 2.02           |
| MSTD4              | 5.20                 | 3.184          | 7.684                | 1.71           |
| MSTD5              | 4.92                 | 3.131          | 7.221                | 1.57           |
| MSTD6              | 8.72                 | 3.485          | 13.44                | 3.66           |
| MSTD7              | 6.39                 | 3.296          | 10.39                | 2.67           |

Urea was employed as a reference material for a comparison of the initial hyperpolarizability and dipole moment values of MSTD2–MSTD7, and they were found to be 5.26, 6.13, 5.20, 4.92, 8.72, and 6.39 D, respectively, according to a review of the literature. It is found that the value of dipole moment of all designed compounds are greater relative to urea (1.373 D).87 The dipole moment of MSTD6 is the highest (8.72 D) among all of the title compounds. The overall decreasing order of values of dipole moments is MSTD6 > MSTD7 > MSTD3 > MSTD2 > MSTD4 > MSTD5 > DTS(FBTTh2)_2R1. The decreasing order of average linear polarizability values of the title compounds was observed to be MSTD6 > MSTD7 > MSTD5 > MSTD4 > MSTD3 > DTS(FBTTh2)_2R1. The average polarizability was found to be the greatest. An extraordinary increment in its value was found, i.e., 3.485 × 10^{-22} esu. This was because of the presence of three nitro groups that enriched the electron density near the acceptor unit and enhanced the electron-withdrawing ability. The β
factor generally increases with an increase in the strength of the substituents such as fluoro, chloro, and cyano groups attached to the acceptor unit 3-dinitromethyleneindan-1-one (DNMI), which affects the nonlinearity of a compound. Furthermore, as the substitutions take place, the contribution of the conjugated system is extended and $\beta$ is more dominant. The decreasing order of $\beta_{\text{total}}$ value for all designed compounds was as follows: MSTD6 > MSTD7 > MSTD3 > MSTD4 > MSTD5 > MSTD2 > DTS(FBTTh2)2R1. Additionally, MSTD6 has highest value of $\beta_{\text{total}}$ ($13.34 \times 10^{-27}$ esu) in comparison to the other designed compounds. In addition to this, a relative study was also carried out by using urea as a standard compound, whose $\beta_{\text{total}}$ value is found to be $3.714 \times 10^{-31}$ esu. The first-hyperpolarizability values of MSTD2–MSTD7 are $8.33 \times 10^{-25}$, $1.07 \times 10^{-24}$, $9.46 \times 10^{-25}$, $8.89 \times 10^{-25}$, $1.65 \times 10^{-25}$, and $1.27 \times 10^{-25}$ times greater, respectively, in comparison to the reference compound DTS(FBTTh2)2R1 and $1.82 \times 10^4$, $2.35 \times 10^5$, $2.06 \times 10^5$, $1.94 \times 10^5$, $3.61 \times 10^5$, and $2.79 \times 10^5$.

Figure 5. TDM graphs of DTS(FBTTh2)2R1 and MSTD2–MSTD7.
times greater, respectively, in comparison to the $\beta_{\text{total}}$ value of urea. Briefly, for MSTD2–MSTD7 because of the incorporation of stronger electron-withdrawing species in the acceptor part, the NLO response exposure is higher, which might minimize the HOMO–LUMO energy gap, increase the ICT between the orbitals, and make the molecule more polarized. The reverse order of the $E_{\text{gap}}$ between the HOMO and LUMO orbitals revealed the second-order hyperpolarizability $\langle \gamma \rangle$ values of all proposed compounds. MSTD6 has the highest value of $\langle \gamma \rangle$ (3.66 $\times$ 10$^{-31}$ esu). The following is the descending order of all of the created compounds: MSTD6 $>$ MSTD7 $>$ MSTD3 $>$ MSTD4 $>$ STD5 $>$ MSTD2 $>$ DTS(FBTTh$_2$)$_2$R1. The current research provides insights into the fascinating NLO data of the aforementioned π-conjugated push–pull organic compounds for modern NLO applications. In addition, due to their high NLO response, the designed compounds can be used as targets for further research.

**Transition Density Matrices (TDMs) and Exciton Binding Energy ($E_b$).** The transition density matrix (TDM) is mainly used to elucidate the electronic charge transfer in the designed compounds MSTD2–MSTD7 and the reference compound DTS(FBTTh$_2$)$_2$R1. TDM analysis is achieved by using the data obtained by the transference of charges from the donor to acceptor moiety via a π linker.90 The TDMs of investigated compounds were computed at the M06 level of theory and the 6-31G(d,p) basis set. The TDM diagrams of all the investigated compounds show the nature of transitions in these compounds. The needed atoms are divided into sections according to their contribution: i.e., donor (D), π spacer (π), and acceptor (A). Hydrogen atoms have been excluded due to their low susceptibility in effective charge transfer phenomena. All of the results for DTS(FBTTh$_2$)$_2$R1 and MSTD2–MSTD7 are shown in Figure 5.

Figure 5 shows that the electron density of all molecules is mostly on the diagonal of the donor as well as on π linkers. A diagonal charge transfer is identified in all non-fullerene-based compounds that have been studied. The π linker acts as a bridge and assists in the relocation of electrons from the donor toward the non-fullerene acceptor units. The diagonal charge is efficiently transferred through a π bridge from the donor to acceptor region, which transfers the charge without or with less charge trapping.

The binding energy ($E_b$) is the difference between electrical ($E_{\text{el}}$) and optical ($E_{\text{opt}}$) band gap energies and is an important parameter for an evaluation of the optoelectronic properties of the studied compounds. Equation 13 is used for the theoretical calculation of the binding energy of reference and designed compounds.

$$E_b = E_{\text{el}} - E_{\text{opt}}$$  

In eq 13, $E_b$ is the energy gap, $E_{\text{el}}$ is the band gap, and $E_{\text{opt}}$ is the first excitation energy.$^2$ All calculated values of binding energy are given in Table 6.

The binding energy of the reference compound was found to be 0.588 eV. The binding energy values of compounds MSTD2–MSTD7 are 0.305, 0.278, 0.290, 0.303, −0.097, and 0.249, respectively. The binding energy values are noted to be in ascending order as MSTD6 $<$ MSTD7 $<$ MSTD3 $<$ MSTD4 $<$ STD5 $<$ MSTD2 $<$ DTS(FBTTh$_2$)$_2$R1. All of the designed MSTD2–MSTD7 molecules give lower values of $E_b$ (eV) in comparison to the reference compound DTS(FBTTh$_2$)$_2$R1. The NF-based designed compounds show promising binding energy values and so may also be used in a vast range of NLO applications.

**CONCLUSION**

Herein, six novel organic molecules (MSTD2–MSTD7) were theoretically designed from DTS(FBTTh$_2$)$_2$R1 by the modification of the different substituted acceptors. The influence of different acceptors is explored for nonlinear amplitudes. FMO results revealed that all of the designed compounds have a small band gap in the range of 1.491–1.835 eV relative to DTS(FBTTh$_2$)$_2$R1 (2.331 eV). Global reactivity parameters were associated with a lower band gap having lower values of hardness with higher values of softness. Moreover, all of the developed compounds had higher $\lambda_{\text{max}}$ values and lower transition energies in the UV–vis region in comparison to the reference molecule (711.408 nm). The development of charge separation between the donor (D) and acceptor (A) species in the molecules was shown by an NBO analysis. Subsequently, this charge separation could be the reason for the great NLO response because of charge transfer from D to A. Interestingly, unusually large $\langle \alpha \rangle$, $\beta_{\text{total}}$, and $\langle \gamma \rangle$ values computed to be $3.485 \times 10^{-22}$, $13.44 \times 10^{-27}$, and $3.66 \times 10^{-31}$ esu, respectively, were found for MSTD6. The first-hyperpolarizability values of MSTD2–MSTD7 are $8.33 \times 10^{-25}$, $1.07 \times 10^{-24}$, $9.46 \times 10^{-25}$, $8.89 \times 10^{-25}$, $1.65 \times 10^{-25}$, and $1.27 \times 10^{-25}$ times greater than that of the reference compound DTS(FBTTh$_2$)$_2$R1 and 1.82 $\times 10^4$, $2.35 \times 10^4$, $2.06 \times 10^4$, $1.94 \times 10^4$, $3.61 \times 10^4$, and $2.79 \times 10^4$ times greater than the $\beta_{\text{total}}$ value of urea. Similarly, MSTD6 has the highest value of $\langle \gamma \rangle$ ($3.66 \times 10^{-31}$ esu). The following is the descending order of all of the created compounds: MSTD6 $>$ MSTD7 $>$ MSTD3 $>$ MSTD4 $>$ STD5 $>$ MSTD2 $>$ DTS(FBTTh$_2$)$_2$R1. We expect that our research will aid in the development of organic compounds with the desired characteristics for improving optical device performance.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.2c01474.

Cartesian coordinates, calculated energies ($E$) and energy gaps ($\Delta E$), global reactivity parameters, natural bond orbital (NBO) analysis, dipole polarizabilities, dipole moments, computed first hyperpolarizabilities ($\beta_{\text{total}}$), second hyperpolarizabilities, UV–vis data (wavelengths, excitation energies and oscillator strengths), and structures of DTS(FBTTh$_2$)$_2$R1 and its derivatives (PDF)
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Complete contact information is available at: https://pubs.acs.org/10.1021/acsomega.2c01474

Notes
The authors declare no competing financial interest.

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