Synthesis of nonlinear optical chromophores containing electronexcessive and -deficient heterocyclic bridges. The auxiliary donor– acceptor e**ff**ects

Ching-Fong Shu* and Yuh-Kai Wang

Department of Applied Chemistry, National Chiao T ung University 1001 Ta-Hsueh Road, Hsin-Chu, Taiwan, 30035, R.O.C.

electron donor (acceptor) group and π is a conjugated bridge which was allowed to react with BuⁿLi yielding 2-chloro-5possessing large molecular second-order nonlinear optical lithiothiazole. Adding N,N-diethylformamide to the 2-chloro- (NLO) responses and good thermal stability, are currently of 5-lithiothiazole solution and quenching with water yielded 2 interest because of their applicability to electro-optic devices.¹ diethylamino-5-formylthiazole 6^{10} Wittig–Horner conden-
In such molecules, the donor and acceptor substituents provide sation of diethyl 2-thienylme In such molecules, the donor and acceptor substituents provide sation of diethyl 2-thienylmethylphosphonate with the thiazole the requisite ground-state charge asymmetry, whereas the π - aldehyde derivative 6 produced c conjugation system provides a pathway for the redistribution of electric charges under the influence of electric fields. Benzene rings are widely used in combination with polyenes in π conjugated bridges because of their thermal and oxidative stability, synthetic availability and substituent positional selectivity. However, the barrier resulting from the aromatic delocalization energy of the benzene ring leads to diminution of β values.2 Synthetic studies have demonstrated that replacing the benzene ring of a chromophore π bridge with easily delocalizable five-membered heteroaromatic rings, such as thiazole and thiophene, results in enhanced molecular hyperpolarizability.3 Recently, computational studies have suggested that heteroaromatic rings play a subtle role in influencing the second-order NLO response properties of donor–acceptor compounds.4 While the aromaticity of heteroaromatics affects electronic transmissions between donor and acceptor substituents, the electron-excessive or electron-deficient nature of the heterocyclic ring systems also plays a major role in determining the overall electron-donating and -accepting ability of the substituents: electron-excessive heterocycles act as auxiliary donors and electron-deficient heterocycles act as auxiliary acceptors.^{4b} Thus, attaching a strong donor to an electronexcessive heteroaromatic, such as thiophene,⁵ and a strong electron acceptor to an electron-deficient heteroaromatic, such as thiazole,⁵ will yield chromophores with significantly enhanced NLO responses. Contrarily, reversing the architecture of the heterocyclic rings in the π -conjugated system will lead to chromophores with diminished NLO responses. **Synthesis of nonlinear optical chromophores containing electron-

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> In this report, we present novel NLO chromophores with π -conjugating moieties that contain thiazole and thiophene, with an ethylene segment in between, and show that the location of the heterocyclic rings indeed has a pronounced effect on the energy of charge-transfer (CT) transition of the chromophores, as was predicted in previous theoretical calculations.⁴ The synthetic routes used to prepare the NLO chromophores $5c-d$ and $7c-d$ containing thiazole and thioscheme 1 Reagents: i, BuⁿLi, N-formylmorpholine; ii, NaBH₄; iii, CCl₄, chromophores 5c–d and 7c–d containing thiazole and thio-

> pph₃; iv, PPh₃; v, 2-diethylamino-5-formylthiophene, Bu^tOK;

> phene are shown in phene are shown in Scheme 1. The thiazole parts of compounds vi, BuⁿLi, N,N-diethylformamide; vii, diethyl 2-thienylmethylphos-5 were synthesized starting from 2-bromothiazole, which was

> treated with BuⁿLi at -78 °C, and the halogen–lithium Push-pull substituted nonlinear optical chromophores with exchange product was quenched with N-formylmorpholine to thiazole and thiophene rings and interposed ethylene units as produce 2-formylthiazole 1.⁶ Reduction of t produce 2-formylthiazole 1.⁶ Reduction of the formyl group of π -conjugated bridges were synthesized. The effects of the π and π an nature and location of the heterocycles on the energy of the transformed into 2-chloromethylthiazole 3 upon chlorination charge transfer transition for the chromophores are discussed. with carbon tetrachloride–triphenylpho with carbon tetrachloride–triphenylphosphine. The reaction of chloromethyl derivative 3 with triphenylphosphine yielded the corresponding phosphonium salt 4.7 Wittig reaction of 2 diethylamino-5-formylthiophene8 with the phosphonium salt 4 in Bu^tOK–benzene led to compound 5a. The thiazole por-Dipolar chromophores of the form D- π -A, where D (A) is an tions of compounds 7 were prepared from 2-chlorothiazole,⁹ aldehyde derivative 6 produced compound 7a. Compounds 5a

Scheme 1 Reagents: i. BuⁿLi, N-formylmorpholine: ii, NaBH₄; iii, CCl₄, phonate, Bu^tOK; viii, BuⁿLi, DMF; ix, BuⁿLi, TCNE; x, CH₂(CN)₂

Table 1 Electronic absorptions and thermal stabilities

$\lambda_{\rm max}/\rm nm$ in dioxane chromophore $\epsilon/10^4$ м $^{-1}$ cm $^{-1}$ $T_{\rm d}/^{\circ}{\rm C}^a$ 702 5.84 $217\,$ CN. CN ΝC 5c 590 4.18 237 CN NC 5d
622 4.97 242 CN .CN NC
7c 514 2.94 268 .CN NC
$7d$ 640^{b} 245 \sum (240^b) S, СN
8с 513 ^c NC.

"DSC, 10 deg min⁻¹. ^bRef 13. 'Ref 3(d).

of their conjugating moieties, were lithiated with BuⁿLi and that replacing a benzene ring on the donor end with an subsequently quenched with tetracyanoethylene and N,N- electron-deficient heterocycle, such as thiazole, decreases the dimethylformamide to give formyl derivatives 5b and 7b, and electron-donating ability of the dialkylamine group, and tricyanovinyl-substituted 5c and 7c. A further Knoevenagel counteracts the effects of reduced aromaticity. reaction of 5b and 7b with malononitrile afforded the dicyanov- The relative thermal stabilities of the chromophores were inyl-substituted chromophores 5d and 7d. All the compounds also studied using differential scanning calorimetry (DSC). The studied here were characterized using conventional spectro-
onset temperatures (T_a) of the chromo studied here were characterized using conventional spectro-
consistentially of the chromophore thermal decompo-
consistentially and the contracted using conventional spectro-
idea and the consistence in Table 1. Although scopic techniques.¹¹ sition exotherms are shown in Table 1. Although, as expected,

characterized by a long-wavelength CT transition that contrib- not lead to substantially lower thermal stability, we note that utes strongly to the second-order molecular hyperpolariz- chromophores with lower charge-transfer transitional energies ability.¹² All of the chromophores in this study had strong tend to possess lower thermal stabilities. bands in the visible region of the spectrum. Table 1 lists the In summary, compounds 5c,d and 7c,d represent the first positions of the absorption maxima for the chromophores in example of NLO chromophores synthesized in which the this study, and for related chromophores.^{3d,13} Chromophores conjugating moieties contain both electron-excessive hetero-5c,d, which have an electron-excessive heterocycle (thiophene) cycles (thiophene) and electron-deficient heterocycles (thiaon their donor ends, and an electron-deficient heteocycle(thia- zole).5 As predicted by theoretical calculations, the CT zole) on their acceptor ends, show pronounced bathochromic transitional energies of the chromophores depends not only shifts (ca. 80 nm) in their CT bands as compared with chromo- on the natures but also strongly on the locations of the phores 7c,d, which have reversed heterocycle substitution heterocyclic rings, which in turn may have significant effects patterns. This result is reasonable since in chromophores $5c,d$, on the molecular second-order NLO response properties.⁴ the thiophene and thiazole rings act as additional donor/ Experiments to measure the second-order molecular hyperpolacceptor groups, facilitating CT transitions. In chromophores arizabilities of these chromophores are in progress. 7c,d, the thiazole ring reduces the donation ability of the donor substituent and the thiophene ring attenuates the acceptance We are grateful to Professor Alex K.-Y. Jen, Northeastern power of the acceptor substituent, an arrangement that impedes University, for useful discussions. We also thank the National CT transitions. When compared to chromophores 8c,d, Science Council (ROC) (NSC 85-2113-M-009-003) for financhromophores 7c,d have no red-shifted CT bands, and even cial support.

Most donor–acceptor substituted NLO chromophores are the replacement of a benzene ring with a heterocyclic ring does

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