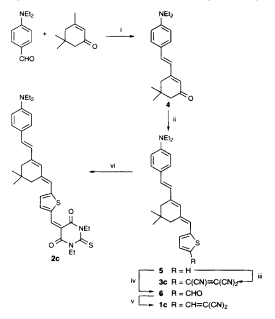
## Synthesis of second-order nonlinear optical chromophores with enhanced thermal stability and nonlinearity: a conformation-locked *trans*-polyene approach

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A facile synthesis that combines the use of an easily polarizable thiophene ring and a ring-locked triene as efficient conjugating moieties is employed to simultaneously enhance molecular nonlinearity (as high as  $13\,000 \times 10^{-48}$  esu) and thermal stability; incorporating this highly nonlinear chromophore in polyquinoline results in high electro-optic activity and long-term alignment stability at 100 °C for more than 1000 h.

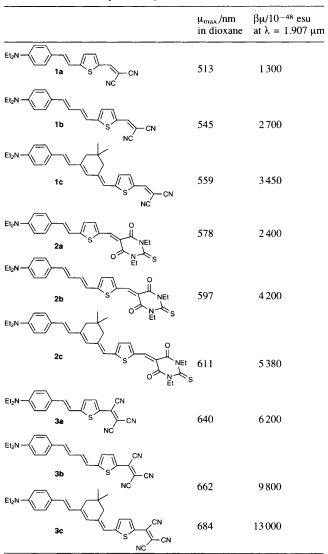
The growing interest in organic polymeric materials for electrooptic (E-O) device applications has led to the development of various classes of asymmetric  $\pi$ -conjugated molecules.<sup>1-6</sup> In poled E-O polymer systems, the alignment of the polar order is preserved either by using host polymers with high glass transition temperatures (200-300 °C),<sup>7</sup> or by cross-linking the chromophore polymer-composite.8 In either case, it poses a challenge for chemists in this field to develop highly nonlinear chromophores with thermal stabilities exceeding 200 °C in order to survive the material processing and device fabrication conditions. Recently, it has been shown that very large nonlinearities can be achieved by employing heteroaromatic rings which have lower aromatic stabilization energy upon charge separation,<sup>9-10</sup> or by using extended polyene  $\pi$ -bridge systems that have strong electron acceptors.<sup>11</sup> Incorporation of these guest chromophores into host polymer matrices and their subsequent alignment by an electric field have demonstrated very large electro-optic coefficients ( $r_{33} = 45-55 \text{ pm V}^{-1}$ ) at a wavelength of 1.3 µm.<sup>12,13</sup> These values are significantly larger



Scheme 1 Reagents: i, Bu'OK-Bu'OH; ii, diethyl 2-thienylmethylphosphonate; iii, BuLi, TCNE; iv, BuLi, DMF; v,  $CH_2(CN)_2$ : vi (from 6), diethylthiobarbituric acid

than the value of 31 pm V<sup>-1</sup> for commercially available LiNbO<sub>3</sub>. However, the long-term alignment stability of these E-O polymer systems was limited due to the low inherent thermal stability (<175 °C) of these chromophores that prevents them from being processed and poled at higher temperatures. We have recently developed a facile approach to the synthesis of NLO chromophores with both enhanced nonlinearity and thermal stability. This synthetic method combines the advantages of using a thiophene ring and a triene as efficient conjugating moieties for easier charge separation, and the use of

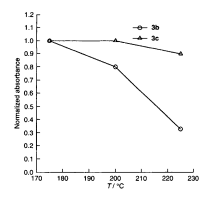
| Table 1                                        | Electronic | absorption a | and the | first m | nolecular | hyperpolarizability |
|------------------------------------------------|------------|--------------|---------|---------|-----------|---------------------|
| obtained for nonlinear optical compounds 1a-3c |            |              |         |         |           |                     |



a 2,2-dimethylpropane-1,3-diyl moiety attached to the triene to give a 6-membered ring system that provides a configurationlocked geometry of *trans*-triene to prevent the thermally induced *cis-trans* isomerization. This allows us to synthesize a series of NLO chromophores with broad variation of electron acceptors to fine tune the linear absorption property ( $\lambda_{max}$ ) and  $\beta\mu$ . We also report the preliminary results of the E-O coefficients and thermal stability.

The synthesis of a series of configuration-locked triene chromophores 1c-3c is shown in Scheme 1. Compound 4 (trans-trans) was prepared by using the Knoevenagel condensation of 4-(N,N-diethylamino) benzaldehyde with isophorone (3,5,5-trimethylcyclohex-2-en-1-one) in the presence of Bu'OK-Bu'OH. Compound 5 was derived from the Wittig-Horner condensation of diethyl 2-thienylmethylphosphonate with 4 in Bu<sup>t</sup>OK-THF. Compound 5, which does not have an electron-withdrawing group at the end of the conjugating moieties, was obtained as a mixture of trans-trans-trans and trans-trans-cis isomers in a ratio of 2:1, as determined by <sup>1</sup>H NMR spectroscopy. Lithiation of 5 (mixture of trans- and cisisomers) with BuLi and subsequent quenching with dimethylformamide (DMF) and tetracyanoethylene (TCNE) gives 6<sup>†</sup> and tricyanovinylsubstituted chromophore 3c. Treatment of 6 (all-trans) with malononitrile and diethylthiobarbituric acid gave the all-trans 1c and 2c in 80-90% yields. All the double-bond linkages in 1c-3c were confirmed to be *trans* by careful NMR analysis.

Electronic absorption spectra of these compounds were obtained to compare their intramolecular charge-transfer properties (Table 1). The trienes 1c-3c have a substantially red shifted charge-transfer band when compared to the stilbenes 1a-3a and dienes 2a-2c, indicating the more efficient electron delocalization nature of the triene bridge. Second-order hyperpolarizabilities of these compounds were determined by using the electric field-induced second harmonic generation (EFISH) technique described previously.14 In order to minimize possible resonance enhancement which would exaggerate the measured NLO responses, measurements were performed at a fundamental wavelength of 1.9 µm in dioxane solvent. Comparison of the  $\beta\mu$  values of the compounds in Table 1 reveals that the combination of the conformation-locked trans-triene bridge with the tricyanovinyl acceptor provides a very efficient mechanism to enhance molecular nonlinearity. The absorption maximum ( $\lambda_{max}$ ) for **3c** was at 684 nm and thus correcting this value for dispersive enhancement using the two-level correction<sup>15</sup> gave a zero frequency value,  $\mu\beta(0)$ , of 5480  $\times$  10<sup>-48</sup> esu, which is 15 times greater than that for the commonly employed 4-(N,N-diethylamino)nitrostilbene NLO chromophore [DANS,  $\mu\beta(0)$  of  $370 \times 10^{-48}$  esu]. Thermal stability studies were performed by dissolving 3b and 3c in polyquinoline PQ-100 (Maxdem) and heating the thin film samples isothermally at 175, 200 and 225 °C for 20 min, respectively. The  $\pi$ - $\pi$ \* chargetransfer absorption band was used to monitor the decomposition



**Fig. 1** Thermal stability of **3b** and **3c** heated isothermally at 175, 200 and 225 °C for 20 min each, measured by following the change in absorbance of a 20 wt% dye-doped polyquinoline thin film vs. T

temperature. Compound 3c with a configuration-locked transtriene bridge possesses much better thermal stability at 200 °C (100%) than **3b**, which has a *trans*-diene bridge (<70%) (Fig. 1). It is well known that thermal stability of the polyene NLO chromophores decreases with an increase in the number of double bond linkages. However, by using this configurationlocked triene approach, the thermal stability of compounds with even longer chain lengths is significantly enhanced. Furthermore, when 3c was doped (15 wt%) in a polyquinoline PQ-100 host and poled at 210 °C with a poling field of 1.4 MV  $cm^{-1}$ , the resulting polymer had an electro-optic coefficient,  $r_{33}$ , of 27 pm V<sup>-1</sup> measured at 1.3  $\mu$ m,<sup>16</sup> consistent with the large molecular nonlinearity. The thermal stability of the poled polymer was demonstrated by heating the poled sample in an oven at 100 °C for over 1000 h. The E-O activity of the sample shows an initial drop to 75% of its original value within 80 h, then remains unchanged for a similar period of time.

We thank the National Science Council (ROC) (NSC84-2113-M-009-003) and the Office of Naval Research (ONR) (N 00014-95-1-1319) for partial financial support for this research. Dr Yongming Cai is acknowledged for help with EFISH measurements.

## Footnote

† Both **6** and **3c** were obtained as a mixture of *trans-trans-trans* and *trans-trans-cis* isomers. However, the *trans* isomer can be separated from the *cis* isomer by column chromatography.  $\delta_{\rm H}$  (CDCl<sub>3</sub>, 300 MHz) for **3c** (*trans*) 1.08 (s, 6 H), 1.19 (t, 6 H), 2.35 (s, 2 H), 2.54 (s, 2 H), 3.40 (q, 4 H), 6.33 (s, 1 H), 6.63 (s, 1 H), 6.64 (d, J 9.0, 2 H), 6.72 (d, J 16.8, 1 H), 6.78 (d, J 17.0, 1 H), 7.12 (d, J 4.8, 1 H), 7.35 (d, J 9.0, 2 H), 7.95 (d, J 4.8, 1 H); for **3c** (*cis*): 1.03 (s, 6 H), 1.22 (t, 6 H), 2.32 (s, 2 H), 2.38 (s, 2 H), 3.40 (q, 4 H), 6.37 (s, 1 H), 6.64 (d, J 9.0, 2 H), 6.65 (s, 1 H), 6.96 (d, J 15.6, 1 H), 6.80 (d, J 15.3, 1 H), 7.10 (d, J 4.8, 1 H), 7.37 (d, J 9.0, 2 H), 7.95 (d, J 4.8, 1 H). H).

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Received, 3rd June 1996; Com. 6/03796H