Optical properties of cross-conjugated isopolydiacetylene oligomers as measured by ultraviolet–visible spectroscopy and the optical Kerr effect

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Abstract
A novel series of isopolydiacetylene oligomers have been characterized on the basis of their electronic absorption, emission and nonlinear optical properties. The ultraviolet–visible spectra indicate only a small shift to lower energy as a function of increasing oligomer length. To evaluate the third-order nonlinear susceptibilities, \( \chi(3) \), and second hyperpolarizabilities, \( \gamma \), of these oligomeric samples a new Kerr gate technique, differential optical Kerr effect detection, has been employed. A linear relationship between \( \gamma \) and the number of repeat units is found, suggesting that the fixed-length linearly conjugated segments dominate the electronic polarizability. The largest nonlinearity was observed in the heptamer sample, displaying a second hyperpolarizability relative to the tetrahydrofuran solvent of \( \gamma_{\text{heptamer}}/\gamma_{\text{THF}} = 181 \pm 9 \).

Keywords: Nonlinear optics, optical Kerr effect, conjugated oligomers, polydiacetylenes, alkynes

1. Introduction
The delocalized electronic structure of \( \pi \)-conjugated organic compounds offers a number of tantalizing opportunities for applications as nonlinear optical (NLO) materials. For practical applications, organic nonlinear materials will also require transparency, stability, rapid response times and processability [1–3]. The ease with which one can systematically vary the framework of organic molecules provides a remarkable opportunity to probe the relationship between structure and function, and these analyses have become the foundation of any effort toward the optimization of optical materials [4–7].

Studies of linearly conjugated polydiacetylenes (figure 1) have shown these materials to have significant third-order nonlinearities [8]. Both experimentally and theoretically it is clear that extending the \( \pi \)-conjugated path of polydiacetylenes significantly enhances the observed nonlinearities [2, 6, 9, 10]. As the conjugation length increases, however, the wavelength of the lowest-energy absorption maximum, \( \lambda_{\text{max}} \), increases as well. This ultimately yields materials with insufficient transparency for many applications [11–13]. It is clear that attempts to maximize \( \chi(3) \) by simply increasing the length of the linearly conjugated backbones are unlikely to produce practical NLO materials [4].

In an attempt to identify molecular designs that could provide both conjugated and transparent materials, we targeted cross-conjugated oligomeric compounds. Because \( \pi \)-delocalization is less efficient across a cross-conjugated bridge than in an analogous linearly conjugated arrangement, we were intrigued as to the possibility of increasing the optical nonlinearities, while at the same time preserving the electronic transparency.
2. Experimental considerations

The optical Kerr effect (OKE) is an optically induced birefringence in an isotropic medium [17]. In a standard Kerr gate [18–20], an intense, linearly polarized pulse induces a birefringence in an isotropic medium. The anisotropy in the sample’s index of refraction (\( n_\parallel - n_\perp \)), the difference in refractive index between parallel and perpendicular axes to the pump polarization) induced by the pump pulse is then probed by a second, weaker, pulse. The induced birefringence, \( n_\gamma \), manifests itself as retardation in the probe beam polarization and may be detected using a polarizer acting as an analyser. A temporal profile of the induced birefringence may be obtained by time-delays of the arrival of the probe pulse. When the pump and probe pulses arrive simultaneously at a sample dominated by an ultrafast electronic nonlinearity, the probe beam experiences the maximum rotation in polarization given by

\[
\phi = \frac{2\pi d}{\lambda} n_2 I_{\text{pump}},
\]

where \( n_2 \) is the nonlinear index of refraction and is proportional to \( \chi^{(3)} \), \( \lambda \) is the wavelength, \( d \) is the sample length and \( I_{\text{pump}} \) is the pump intensity in units of \( \text{W m}^{-2} \). The quantity \( n_2 \) is a modification to the linear index of refraction where the total index of refraction is given by \( n = n_0 + n_2 I_{\text{pump}} \), with \( n_0 \) being the linear index of refraction. \( \chi^{(3)} \) is obtained from \( n_2 \) by [21]

\[
n_2 = \frac{3.9 \times 10^{-6}}{n_0^2} \chi^{(3)} \text{(esu)},
\]

where \( n_2 \) is in units of \( \text{m}^2 \text{W}^{-1} \) and \( \chi^{(3)} \) is in esu [21]. The molecular second hyperpolarizability, \( \gamma \), may be obtained by [3]

\[
\gamma = \frac{\chi^{(3)}}{N_c L^4}.
\]

where \( N_c \) is the molecular number density in \( \text{cm}^{-3} \) and \( L^4 \) is the Lorentz field-factor which, for isotropic liquid media is approximated by \((n_0^2 + 2)/3)^4 \).

To separate the real and imaginary components of \( \chi^{(3)} \) and to obtain larger signals, optical heterodyne detection–optical Kerr effect (OHD–OKE) is used by many groups [22, 23]. In OHD–OKE, a quarter-wave plate (\( \lambda/4 \)) is placed between the probe polarizer and sample such that the fast axis of the quarter-wave plate is parallel to the polarizer. By slightly rotating the polarizer, \( \pi/2 \) out-of-phase light is added to the probe beam, and this technique can provide the real or imaginary components of the nonlinear response directly from the Kerr signal. In this arrangement, for small responses, transmitted signals are linear with pump intensities.

The DOKE setup shown in figure 2 uses pump and probe pulses generated by an amplified Ti:sapphire laser, producing 800 nm, < 100 fs pulses at a 1 kHz repetition rate. A computer-controlled retroreflector is used to vary the delay between the pump and probe pulses. The pump pulse is polarized 45° to the horizontal before being focused onto a 1 mm sample-path-length quartz cuvette filled with a sample solution. The weaker probe beam is initially polarized vertically and then passes through a quarter-wave plate oriented to produce circular polarized light. Typical pump and probe powers at the sample location are between 0.5 and 6 mW for the pump beam and 50 and 80 \( \mu \text{W} \) for the probe beam. The probe light is focused to overlap in the sample with the pump beam at a near coincident angle, \( \theta < 5^\circ \). Typical spot sizes for the pump and probe beams are \( \sim 500 \text{ and } 240 \mu \text{m} \) respectively. After passing through the sample, the pump beam is blocked while the probe beam is allowed to travel to a Glan-laser polarizer acting as the analyser. Here, the transmitted and rejected beams are directed to a pair of balanced photodiodes where photodiode ‘A’ receives the rejected beam and photodiode ‘B’ receives the transmitted beam. The pump beam is chopped at 125 Hz, and lock-in detection is used to monitor the A – B signal. The A + B signal is detected at 1 kHz using a separate lock-in amplifier. Finally, these signals are sent to a computer used for data acquisition and delay-stage control.

The key feature of the DOKE setup is that A – B acts as the Kerr effect signal detection and, if \( \text{Im}(\chi^{(3)}) \) is small, A + B acts as a probe beam reference. In the presence of nonlinear absorption, such as two-photon absorption, the A + B signal provides independent detection of \( \text{Im}(\chi^{(3)}) \) processes. The photodiodes are calibrated such that in the absence of any birefringence, A and B signals are equal and A – B is zero.

Consider a vertically polarized probe beam and a sample with an induced birefringence of \( \phi = \phi' + i\phi'' \), where \( \phi' \) and \( \phi'' \) are the real and imaginary components of the phase retardation, respectively. If after passing through a quarter-wave plate set with fast axis at 45° the beam is incident on the sample, the detected signal ratio is given by

\[
\frac{A - B}{A + B} = \frac{\sin \phi' \cosh \phi''}{\sinh \phi'}.
\]

This relationship is valid when the pump and probe beams are initially polarized at 45° with respect to each other. In the absence of nonlinear absorption, \( \phi'' = 0 \), and equation (4) is reduced to equation (5), a purely real response:

\[
\frac{A - B}{A + B} = \sin \phi'.
\]
signal than the typical heterodyne scheme. We have also observed that the DOKE signals are larger than those obtained with OHD–OKE, making detection easier. A full description of the DOKE technique will be presented elsewhere.

The iso-PDA oligomers used in this study are shown in figure 3. (Details of their synthesis will be published elsewhere [24].) NLO samples were prepared by dissolving the oligomer sample in high-purity tetrahydrofuran (THF) to give solutions of 0.1–0.23 M. All samples were measured within 24 h of preparation.

3. Results and discussion

The electronic absorption data for the iso-PDA dimer through the heptamer are shown in figure 4, and the lowest-energy absorption maxima ($\lambda_{\text{max}}$) are listed in table 1. There is a slight shift in $\lambda_{\text{max}}$ values as the chain length is extended from the dimer through the heptamer. By the stage of the heptamer, this effect appears to have reached saturation at 378 nm, and longer oligomers are expected to remain transparent throughout the remainder of the visible region. Furthermore, the molar absorptivity increases with the number of fixed-length linearly conjugated paths. This is in significant contrast to the behaviour of linearly conjugated polydiacetylenes that show large increases of $\lambda_{\text{max}}$ with similar chain length extension, while maintaining a relatively constant molar absorptivity at $\lambda_{\text{max}}$ [25, 26].

These results show that electronic characteristics of the iso-PDAs are dominated primarily by contributions from the longest linearly conjugated segment present in the molecules (shown in bold in for the dimer and the heptamer in figure 3),

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**Table 1.** Linear and NLO data for iso-PDAs.

<table>
<thead>
<tr>
<th>n</th>
<th>$\lambda_{\text{max}}$ (nm)</th>
<th>$\lambda_{\text{emission}}$ (nm)</th>
<th>$\gamma_S$ ($\times 10^{-36}$ esu)</th>
<th>$\gamma_S/\gamma_{\text{THF}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>373 (18 500)</td>
<td>475</td>
<td>30.3 ± 0.3</td>
<td>58 ± 0.5</td>
</tr>
<tr>
<td>3</td>
<td>367 (31 600)</td>
<td>489</td>
<td>49 ± 4</td>
<td>94 ± 7</td>
</tr>
<tr>
<td>5</td>
<td>377 (49 300)</td>
<td>498</td>
<td>69 ± 5</td>
<td>134 ± 9</td>
</tr>
<tr>
<td>7</td>
<td>378 (76 800)</td>
<td>501</td>
<td>95 ± 4</td>
<td>181 ± 9</td>
</tr>
</tbody>
</table>
while contributions from cross-PDAs are minimal or absent.

Unlike the absorption energies for these four compounds, the photoluminescence emission energy is more dependent on oligomer length. Figure 5 shows the emission spectra for the iso-PDAs. The emission wavelength of the heptamer is redshifted by about 25 nm in comparison to that of the dimer. The similar emission wavelength for the pentamer and heptamer, however, suggests that saturation has already been reached.

Table 1 lists the molecular hyperpolarizabilities for each of the oligomers, as well as the nonlinearity relative to THF. Figure 6 shows the concentration-normalized DOKE signals for the oligomer series, with each time-resolved plot the result of five averaged scans. The molecular second hyperpolarizabilities were obtained from the peak signals in figure 6, using

$$\chi^{(3)}_S = \frac{\gamma_S (\phi_S / \phi_R - 1)}{L^2 N_C}.$$  \hspace{1cm} (6)

The requisite $\chi^{(3)}$ values were obtained from the following relationship:

$$\chi^{(3)}_S = \chi^{(3)}_R \left( \frac{\phi_S}{\phi_R} \right),$$  \hspace{1cm} (7)

where $\chi^{(3)}_S$ and $\chi^{(3)}_R$ are, respectively, the sample and reference susceptibilities and $\phi_S$ and $\phi_R$ are, respectively, the measured rotation in polarization of the sample and reference.

THF was used as a reference for all samples. Using the treatment described above, we obtain a reference value of $\chi^{(3)}_R = 2.1 \times 10^{-14}$ esu. It should be noted that $\chi^{(3)}_R$ is not $\chi^{(3)}_{THF}$, as it includes contributions of the quartz cuvette. Separate the effects of the sample holder yields $\chi^{(3)}_{R,THF} = (1.2 \pm 0.3) \times 10^{-14}$ esu and $\gamma_{THF} = (5.2 \pm 1.3) \times 10^{-15}$ esu, which compare reasonably well with reported values of $\chi^{(3)}_{THF} = 3.7 \times 10^{-14}$ esu ($\lambda = 620$ nm) [27] and $\gamma_{THF} = 4.8 \times 10^{-15}$ esu [28]. Maximum signals are observed at zero probe delay times, suggesting that the ultrafast electronic response is the dominant contributor to the observed nonlinearities.

4 A full discussion of signal analysis and blank separation is beyond the scope of this paper. Briefly, $\chi^{(3)}_R$ and $\gamma_R$ refer to signals that include the sample holders, whereas $\chi^{(3)}_{R,THF}$ and $\gamma_{THF}$ refer to pure THF values.

5 Uncertainty in these values comes from uncertainties in pulse-shape/power diagnostics as well as signal analysis. The error in $\gamma$ reported is due to uncertainties in signal analysis alone, as a precise $\chi^{(3)}$ is assumed.

4. Conclusions

The linear and third-order NLO properties of cross-conjugated, iso-PDAs have been explored as a function of oligomer length. These compounds show a linear increase in molecular second hyperpolarizability, $\gamma$, as the chain length is increased from the dimer to heptamer. This increase in $\gamma$ can be attributed to an increase in the number of fixed-length linearly conjugated paths. No added contribution to the nonlinear response attributable to the cross-conjugated backbone has been observed. Extension of this study to include longer oligomers is under way, and should provide more concrete trends in sample nonlinearities.
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References

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