Synthesis and First Hyperpolarizabilities of Acceptor-substituted β-apo-8′-Carotenal Derived Compounds

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The synthesis and second-order nonlinear optical properties of acceptor-substituted biologically derived β-apo-8′-carotenal compounds are reported; electric field-induced second harmonic generation (EFISH) measurements give values of β(0) which are 2–6 times greater than for 4,N,N-dimethylamino-4′-nitrostilbene (DANS).

In an attempt to understand the structural basis of molecular polarizabilities and hyperpolarizabilities in conjugated organic chromophores we have been investigating the parameters of bond length alternation (BLA) and aromaticity.1–3 Computational studies using MOPAC,4,5 together with experimental electric field-induced second harmonic generation (EFISH)6,7 and third harmonic generation (THG) results on asymmetric cyanines and polarized polyenes have lead us to believe that there is a specific amount of BLA (the difference in length between the carbon double and single bonds in a conjugated chain) which can maximize the molecular polarizability (α), the first hyperpolarizability (β)1–3–8 or the second hyperpolarizability (γ)9 for a given molecular length. Experimental and theoretical evidence1,3,9 points to the fact that α is maximized when BLA is zero, and that β is peaked at a value of around 0.04 Å. Calculations for relatively short systems, also predict that positive γ is peaked at greater BLA than for β, and that negative γ is maximized at zero BLA.

At present most donor–acceptor nonlinear optical chromophores in their ground-state contain aromatic groups, which from simple resonance structure considerations become quinoidal when the molecule is in its charge-separated state. Investigations on a typical neutral donor–acceptor substituted compound1 indicate that for a given chain length β is larger when the molecule does not have to lose aromaticity upon charge-transfer polarization. As aromaticity is stabilizing, its loss upon charge-separation could hinder efficient charge-transfer polarization.

We wished to probe the effects of aromaticity and the introduction of ground- and excited-state dipole moments on both the second- and third-order hyperpolarizabilities of a series of compounds very closely related to the well-studied molecule, β-carotene.10 Blanchard-Desce and coworkers11–13 have investigated the nonlinear optical properties of some

![Scheme 1 Synthesis of acceptor-substituted β-apo-8′-carotenoid compounds](image-url)
related compounds with chain lengths shorter than and equal to β-apo-8'-carotenal, 1 (Scheme 1), substituted with a series of donors and an aldehyde acceptor, and Iked and co-workers have examined a series of retinal-derived compounds. In this contribution we report on the synthesis and second-order nonlinear optical properties of β-apo-8'-carotenal derivatives with terminal acceptor substituents, whose structures are more closely related to β-carotene than those in the previous studies.

The new compounds were readily synthesized (Scheme 1) using standard Knoevenagel conditions for the synthesis of 2 and Wadsworth–Emmons conditions for 3. The compounds were purified by chromatography on silica, recrystallized and then characterized by UV–VIS, 1H and 13C NMR and mass spectroscopy, as well as elemental analysis. β Values were measured in chloroform solution by EFISH using 1.907 μm fundamental radiation.6,7

Table 1 shows values of the optical absorption maximum (λmax), the extinction coefficient (ε), the dipole moment (μ), first hyperpolarizability (β), the zero frequency of β (β(0)) corrected for dispersion using a two-state model and the dot product of β and μ (β·μ), which is of interest in polarized polymer applications. Although these chromophores have no heteroatom donor, their hyperpolarizabilities are reasonably large, with each having a β(0) significantly larger than that of the prototypical stilbene compound 4-N,N-dimethylamino-4'-nitrostilbene (DANS) [β(0) = 55 × 10⁻³⁰ esu]. These rather large nonlinearities can be explained by the large intrinsic polarizability of the extended π-electron system. For polymers, the HOMO–LUMO gap decreases with increasing conjugation length resulting from the energy of the LUMO being lowered and that of the HOMO being raised. In molecular orbital terms, a donor is a high lying occupied orbital, and thus, in long polyenes, the HOMO itself can act as a donor. As the length of the conjugated chain is increased, the HOMO becomes a more effective donor.

Dicyanovinyl groups are stronger acceptors than aldehydes, and this is illustrated in compound 2 having a β(0) value of over three times greater than for compound 1. Nitro groups are also stronger acceptors than aldehydes, but on looking at the β(0) value for 3 it is not substantially greater than for 1, even although it is 6 atoms longer. This low value can be explained by the nitrophenyl acceptor containing a stabilizing aromatic group which hinders the formation of the charge-separated form of the molecule and hence reduces the polarizability and therefore β(0).8

In summary, β(0) values of aldehyde 1, dicyanovinyl 2 and nitrophenyl 3 acceptor-substituted β-apo-8'-carotenal chromophores measured by EFISH, are all large in view of the fact that no conventional donor is present, with 2 giving the largest value. The aromatic ring in compound 3 may hinder the formation of the charge-separated state, causing β(0) to be greatly reduced.

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Table 1 Selected linear and nonlinear optical properties in chloroform of substituted β-apo-8'-carotenoids in Scheme 1

<table>
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<tr>
<th>Compound</th>
<th>λmax/νm</th>
<th>ε/mol⁻¹ · cm⁻¹</th>
<th>μ/10⁻¹⁸ esu</th>
<th>β(0)/10⁻³⁰ esu</th>
<th>β(0)/10⁻³⁰ esu</th>
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<td>1</td>
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References