Rotationally Resolved Electronic Spectra of trans,trans-Octatetraene and Its Derivatives

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Described herein are the rotationally resolved one-photon fluorescence excitation spectra of several vibronic bands in the S1 ← S0 electronic transitions of three linear polyenes in the gas phase, 1,3,5,7-octatetraene (OT), 1,3,5,7-nonatetraene (NT), and 2,4,6,8-decatetraene (DT). Several of the spectra are significantly perturbed by an apparent centrifugal distortion in the S1 state of OT, owing to the high frequency of rotations parallel to a and the low frequency of an in-plane bending mode (v148), by Coriolis coupling in the S1 state of NT, involving v148 and a nearby methyl torsional level, and by torsion–rotation coupling in the S1 states of NT and DT, owing to a significant reduction in the excited-state torsional barrier(s) compared to the ground state. Nonetheless, the inertial parameters of eight different S0 and S1 vibronic levels have been determined, from which it is concluded that the carriers of the spectra are in all cases the trans,trans isomers. The important role of v148 as a promoting mode for S1 ← S0 vibronic coupling, the source of the S1 ← S0 oscillator strength, is confirmed. Finally, the measured differences in the rotational constants of the S0 and S1 states (e.g., ΔA = 2532, ΔB = −11.7, and ΔC = −11.0 MHz for the vibronic origin of OT) provide new information about the changes in geometry that occur when the photon is absorbed.

Introduction

Polyenes (CnH2n−2), molecules with alternating single and double bonds between covalently attached carbon atoms, are an important class of molecules with many interesting properties.1 Among these is geometrical isomerism, by far the most significant property of polyenes in biological systems. Each 180° turn at either a single or a double bond gives a different configuration, resulting in a large number of possible structures, especially when n is large. Thus, octatetraene (n = 8) has 20 distinguishable isomers, although three or four consecutive cis configurations result in steric crowding that may prevent their formation. The net count in C8H10 is 15, excluding cis3 configurations, and 18, excluding cis4 configurations.2

Given this fact, the existence of a controversy concerning the correct assignment of the electronic spectrum of octatetraene is not surprising. Heimbrook et al.3 observed S2 ← S0 (11B0 → 11A0) emission from the gas-phase molecule and recorded its strongly allowed S2 ← S0 fluorescence excitation spectrum. However, they were not able to detect the dipole forbidden S1 ← S0 (21A1 ← 11A0) transition under isolated molecule conditions. Later, Buma et al.4 obtained the two-color REMPI spectrum of the 21A1 state in the gas phase and assigned their spectrum to a "cis, trans" isomer. They argued that only a noncentrosymmetric isomer would have a large enough oscillator strength to be detected under their experimental conditions. Then, Petek et al.5 succeeded in observing the S1 ← S0 fluorescence of octatetraene in a supersonic jet and recording its one- and two-photon S1 ← S0 fluorescence excitation spectra. A detailed comparison of these spectra led to the conclusion that the carrier is the trans,trans isomer, the most stable form. This conclusion was supported by a more recent study of the fluorescence excitation spectra of two additional “linear” tetraenes, nonatetraene and decatetraene, in which one or both terminal hydrogen atoms in octatetraene are replaced by methyl groups.6,7

Rotationally resolved electronic spectroscopy is a powerful method for establishing the identity of gas-phase chromophores. The patterns of energy levels observed in this experiment are sensitive both to the size and shape of the molecule and to how these changes when a photon is absorbed. Here, we demonstrate the utility of this approach in a study of the high-resolution one-photon fluorescence excitation spectra of octatetraene, nonatetraene, and decatetraene in the collision-free environment of a molecular beam. The results provide an unambiguous identification of the isomers responsible for the above-mentioned spectra as the trans,trans forms. The spectra also give new insight into the remarkable photochemical and photophysical properties of the polyenes and provide new experimental benchmarks for evaluating theoretical descriptions of their 21A1 states.

Experimental Section

Octatetraene (OT) was prepared by dehydrating 1,4,6-octatrien-3-ol at 80 °C using pyridinium p-toluenesulfonate (Aldrich) following the procedure developed by Yoshida and Tasumi.9 The 1,4-octatrien-3-ol was produced by mixing 2,4-hexadienal with vinylmagnesium bromide (Alfa). The dehydra-

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tion produces \textit{trans,trans}-OT of \textgreater 99\% isomeric purity as determined by an analysis of the \textit{S}_2 \rightarrow \textit{S}_0 spectrum in a free jet.\textsuperscript{5} The OT crystals were stored at \(-80^\circ\text{C}\) to preserve their isomeric purity. Nonatetraene (NT) and decatetraene (DT) were synthesized from Wittig reactions, using hexadienal and allyltriphenylphosphonium bromide (Fluka) for the former and hexadienal and crotyltriphenylphosphonium bromide (Fluka) for the latter.

Supersonic jet spectra were obtained using a standard low-resolution apparatus. Typically, the polyenes (OT at room temperature, NT and DT with mild heating) were seeded into 80 psig He and expanded through a pulsed 1 mm nozzle (General Valve, series 9). The resulting free jet was crossed with a frequency-doubled Quanta-Ray Nd\textsuperscript{3+}:YAG-pumped dye laser beam about 1 cm downstream of the nozzle. When tuned to a particular vibronic band in the absorption spectrum, the laser produced \textit{S}_1 \rightarrow \textit{S}_0 fluorescence, which was collected by a single lens system, detected by a photomultiplier tube and SRS boxcar integrator, and acquired by a MASSCOMP data acquisition system.

High-resolution spectra were obtained using a CW molecular beam spectrometer equipped with an intracavity-doubled Spectra-Physics ring dye laser, described in detail elsewhere.\textsuperscript{10} As above, OT at room temperature (or NT and DT with mild heating) was continuously expanded through a 90 \(\mu\)m nozzle in 40–60 psig He, skimmed once, and excited \textasciitilde15 cm downstream of the nozzle by the laser. Fluorescence was collected using spatially selective optics, detected by a photomultiplier tube and photon-counting system, and processed by the data acquisition system. Relative frequency calibrations of the spectra were performed using a near-confocal interferometer. Approximately 10 \times 100 mg of each sample was available for the experiments, so special care was taken to optimize the performance of the apparatus prior to each run.

\textbf{Results and Interpretation}

Figure 1 shows the low-resolution one-photon fluorescence excitation spectra of OT, NT, and DT recorded in a supersonic jet. These are similar in every respect to the published spectra.\textsuperscript{1,5} The first strong band in the excitation spectrum of OT (Figure 1a) is at 29 024.9 cm\textsuperscript{-1} (band OT-1). This band has been assigned as the \textit{0a} \textit{1} band of the \textit{S}_1 \rightarrow \textit{S}_0 transition of an OT having a cis linkage by Buma et al.\textsuperscript{4} and as the vibronic origin of \textit{trans,trans}-OT by Petek et al.\textsuperscript{5} Their assignment places the \textit{0a} \textit{1} band at 76.2 cm\textsuperscript{-1} to the red of band OT-1, at 28 948.7 cm\textsuperscript{-1}, observed only in two-photon excitation experiments.\textsuperscript{5} The frequency of 76.2 cm\textsuperscript{-1} has been assigned to \(\nu_a\), a \textit{b}_\text{a} in-plane bending mode in the \textit{S}_1 state. Also prominent in the spectrum of OT is a band at 50.9 cm\textsuperscript{-1} (band OT-2), which has been assigned to \(\nu_{16} + \nu_{16}\), where \(\nu_{16}\) is a totally symmetric (\textit{a})-in-plane bending mode.

The first strong band in the excitation spectrum of NT (Figure 1b) is a doublet (bands NT-1 and NT-2). Band NT-1 lies at 29 850.8 cm\textsuperscript{-1} and has been assigned as the true origin of the \textit{S}_1 \rightarrow \textit{S}_0 transition of \textit{trans,trans}-NT.\textsuperscript{7} According to Petek et al.,\textsuperscript{2} the doubling of the origin is produced by the hindered internal rotation of the attached methyl group. Thus, the two components are \textit{0a} \textit{1} \rightarrow \textit{0a} \textit{1} and 1e \rightarrow 1e tunneling doublets, with the latter (band NT-2) shifted to the blue by 3.34 cm\textsuperscript{-1} with respect to the former (band NT-1). At 60.1 cm\textsuperscript{-1} further to the blue of band NT-1 lies a second doublet (bands NT-3 and NT-4), also split (by 3.37 cm\textsuperscript{-1}) by the tunneling motion of the methyl group. Band NT-3 lies at 29 010.9 cm\textsuperscript{-1} and has been assigned as the vibronic origin of the \textit{S}_1 \rightarrow \textit{S}_0 transition of \textit{trans,trans}-NT. Most (but not all) of the remaining bands in the low-resolution spectrum exhibit similar splittings.

The first strong band in the excitation spectrum of DT (Figure 1c) is a triplet (bands DT-1, 2, and 3). Band DT-1 is at 29 014.0 cm\textsuperscript{-1}. This band has been assigned as the vibronic origin of the \textit{S}_1 \rightarrow \textit{S}_0 transition of \textit{trans,trans}-DT, whose true origin lies 50.9 cm\textsuperscript{-1} to the red at 28 963.1 cm\textsuperscript{-1}.\textsuperscript{6,7} The three components of the triplet are separated by 3.09 and 3.16 cm\textsuperscript{-1}. These have been assigned as the \textit{0a} \textit{1} \rightarrow \textit{0a} \textit{1}, 1g \rightarrow 1g, and (1e \textit{1} + 1e \textit{3}) \rightarrow (1e \textit{1} + 1e \textit{3}) tunneling components of two equivalent methyl groups. Further to the blue in the spectra of all three molecules are several additional vibronic bands that have not been examined at high resolution. Their assignments are discussed elsewhere.\textsuperscript{4–7}

Figure 2 shows the rotationally resolved one-photon fluorescence excitation spectrum of the band at 29 024.9 cm\textsuperscript{-1} in OT, band OT-1.\textsuperscript{11} The spectrum is that of a near-prolate symmetric top (\(k = -0.999\)) and, despite its overall appearance, is an a-type band. Band OT-2, at 29 360.6 cm\textsuperscript{-1}, is qualitatively similar. The reason for the unusual appearance of these two bands is that the pseudo-linear OT has, in its \textit{S}_0 state, a large \(\text{A}_{\text{eff}}\) constant (~20 GHz) relative to its \textit{B} and \textit{C} constants (~500 MHz). The moment of inertia about the long axis is very small relative to those perpendicular to it. Thus, OT is very nearly a symmetric top. Even a very small percentage change in the \(K_{\text{a}}\) subbands, as shown in Figure 3. These shifts are approximately proportional to \(K_{\text{a}}^2\), destroying the sharp, centrally located Q-branch structure that is normally observed in a-type bands. Similar effects have been observed in the rotationally resolved \textit{S}_1 \rightarrow \textit{S}_0 spectra of \textit{trans,trans}-diphenylbutadiene (DPB)\textsuperscript{12} and \textit{trans}-stilbene (\textit{iS}).\textsuperscript{13}
The parameters obtained from fits of these two spectra to the ground state of OT, in a molecular beam. The top trace shows the experimental spectrum, the next trace shows the computed spectrum (simulated using the constants in Tables 1 and 2), and the bottom two traces show a comparison of a portion of the experimental and computed spectra, to illustrate the quality of the fit. The line width of single rovibronic features in the experimental spectrum in 15 MHz; this line width is not included in the simulation in the bottom trace. The rotational temperature of the fit is 8 K.

The parameters obtained from fits of these two spectra to distortable rotor Hamiltonians (S-reduction) are listed in Tables 1 and 2. Bands OT-1 and OT-2 exhibit the same ground-state rotational constants within experimental error. Thus, they both originate in the same vibronic level, presumably the zero-point vibrational level (ZPL) of the S0 state. The ground-state parameters listed in Table 1 are the results of a fit of 205 combination differences obtained from the two spectra. The calculated energy differences reproduce the experimental values to within a standard deviation of 4.5 MHz. This may be compared to the observed Doppler width of about 15 MHz. Adding centrifugal distortion terms to the S0 Hamiltonian did not significantly improve the quality of the fit. Hence, the ZPL of the ground state is a rigid rotor.

Also listed in Table 1 are two sets of theoretical rotational constants for the trans,trans isomer of OT. The first set was calculated from a Hartree–Fock (HF) geometry, optimized using the 6-31G* basis set.15 It gives \( A'' = 19 \text{ MHz}, B'' = 573, \) and \( C'' = 557 \text{ MHz} \). The second set was calculated from a geometry obtained by Serrano-Andrés et al.16 using the complete active-space self-consistent-field (CASSCF) method. It gives very similar values: \( A'' = 19 \text{ MHz}, B'' = 568, \) and \( C'' = 552 \text{ MHz} \). The experimental \( A'' \) rotational constant (18 192 MHz) could not be determined accurately from the spectra owing to their near-symmetric-top nature.17 However, the experimental values of \( B'' (571 \text{ MHz}) \) and \( C'' (555 \text{ MHz}) \) are known to high precision. Moreover, there is near-perfect agreement between these values and the two sets of theoretical values. We therefore conclude that the carrier responsible for the two observed bands is trans,trans-OT, in agreement with Petek et al.5

We also have calculated the rotational constants of eight additional OT isomers, each possessing at least one cis linkage, using the 6-31G* basis set. All exhibit significantly smaller \( A'' \) values and significantly larger \( B'' \) and \( C'' \) values, as expected. For example, the tctt isomer has \( A'' = 8019, B'' = 758, \) and \( C'' = 693 \text{ MHz}. \) Thus, there can be little doubt that the carrier of these bands is the trans,trans structure.

### Table 1: Inertial Parameters of trans,trans-Octatetraene in the Zero-Point Vibrational Level of Its Ground Electronic State (1\(^1\)A\(_e\))

<table>
<thead>
<tr>
<th></th>
<th>experiment</th>
<th>theory</th>
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</thead>
<tbody>
<tr>
<td>( A'' ) (MHz)</td>
<td>18192</td>
<td>19831</td>
</tr>
<tr>
<td>( B'' ) (MHz)</td>
<td>571.04(0.1)</td>
<td>573.2</td>
</tr>
<tr>
<td>( C'' ) (MHz)</td>
<td>554.96(0.1)</td>
<td>557.1</td>
</tr>
<tr>
<td>( \kappa'' )</td>
<td>-0.999</td>
<td>-0.999</td>
</tr>
<tr>
<td>( \Delta'' ) (amu Å(^2))</td>
<td>-2.14</td>
<td>1.91</td>
</tr>
</tbody>
</table>

* Results from a fit of 205 combination—differences that appear in the rotationally resolved spectra of the two bands at 29 024.9 cm\(^{-1}\) (\( \Omega_0^0 + 76.2 \text{ cm}^{-1} \)) and 29 360.6 cm\(^{-1}\) (\( \Omega_0^0 + 411.9 \text{ cm}^{-1} \)). Numbers in parentheses represent estimates of the systematic uncertainties in the experiment. \(^{b}\) Estimated errors from the diagonal elements of the correlation matrix. \(^{c}\) Reference 16.

### Table 2: Inertial Parameters of trans,trans-Octatetraene in Two Vibrational Levels of Its First Excited Electronic State (2\(^1\)A\(_e\))

<table>
<thead>
<tr>
<th></th>
<th>experiment</th>
<th>theoretical</th>
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<tbody>
<tr>
<td>( \Omega_0^0 + 76.2 \text{ cm}^{-1} )</td>
<td>20724</td>
<td>20435</td>
</tr>
<tr>
<td>( \Omega_0^0 + 411.9 \text{ cm}^{-1} )</td>
<td>2000</td>
<td>2000</td>
</tr>
</tbody>
</table>

* Numbers in parentheses represent estimates of the systematic uncertainties in the experiment. Band origins were not determined owing to the lack of a calibrated I\(_2\) spectrum in this region. \(^{d}\) Estimated errors from the diagonal elements of the correlation matrix. \(^{e}\) Reference 16. \(^{f}\) Neglecting distortion parameters.

Figure 2. High-resolution one-photon fluorescence excitation spectrum of band OT-1 in the S\(_1\) ← S\(_0\) transition of octatetraene, at 29 024.9 cm\(^{-1}\), in a molecular beam. The top trace shows the experimental spectrum, the next trace shows the computed spectrum (simulated using the constants in Tables 1 and 2), and the bottom two traces show a comparison of a portion of the experimental and computed spectra, to illustrate the quality of the fit. The line width of single rovibronic features in the experimental spectrum in 15 MHz; this line width is not included in the simulation in the bottom trace. The rotational temperature of the fit is 8 K.

Figure 3. Deconvolution of the spectrum in Figure 2, showing the contributions from individual \( K_a \) subbands. Note the large shift to the blue of successive subbands, determined by \( \Delta(A-B)K_a^2 \) in the rigid-rotor limit. These shifts are increased further by the addition of the \( D_aK_a^4 \) distortion term to the \( S_1 \) Hamiltonian.

\[ A'' (\text{MHz}) \]
\[ B'' (\text{MHz}) \]
\[ C'' (\text{MHz}) \]
\[ \kappa'' \]
\[ \Delta'' (\text{amu Å}^2) \]
\[ \text{assignt} \]
\[ \text{rms (OMC)} \]
The excited-state parameters of the two bands listed in Table 2 were determined separately by fixing the $S_0$ rotational constants at the experimental values shown in Table 1 and varying the $S_1$ rotational constants to obtain least-squares fits. These calculations showed that both excited-state vibrational levels are nonrigid, requiring the full Watson Hamiltonian\textsuperscript{14} for their analysis. For the band at $0_0^+ 76.2 \text{ cm}^{-1}$ (band OT-1), we fit 353 transitions with a standard deviation of 11.0 MHz. For the band at $0_0^+ 411.9 \text{ cm}^{-1}$ (band OT-2), we fit 192 transitions with a standard deviation of 7.2 MHz. Both deviations are less than the observed widths of single rovibronic lines in the two spectra. The complete set of quartic centrifugal distortion terms\textsuperscript{14} is listed in Table 2. However, only $D_{K'}$ significantly improves the fit. The effect of this term is to shift the parallel-type $K_a$ subbands even further to the blue, by $D_{K'} K_a^4$ (cf., Figure 3). No other perturbations were observed in these two rotationally resolved spectra of OT.

Four bands in the $S_1 \rightarrow S_0$ spectrum of NT have been recorded at high resolution. These include both components of the doublet located at 28 950.8 and 28 954.1 cm\textsuperscript{-1} (bands NT-1 and NT-2) and both components of the doublet located at 29 010.9 and 29 014.3 cm\textsuperscript{-1} (bands NT-3 and NT-4). These are shown in Figures 4–6. The red and blue members of these doublets have been assigned to torsional levels having a and e symmetry, respectively.\textsuperscript{7} Consistent with this interpretation, bands NT-1 and NT-3 were found to exhibit a-type contours, like the two recorded bands of OT. Thus, similar strategies could be used for their analysis. First, we fit the two bands independently, using rigid rotor Hamiltonians for all four states. Again, we found near-prolate symmetric top behavior, with large $A$ and small and nearly equal $B$ and $C$ values. We also found that the two sets of ground-state constants are identical within experimental error, showing that the two bands originate in the same vibronic level, presumably the ZPL of the $S_0$ state. Then we used the combination–difference method to determine more accurate values of the rotational constants of this level. The results are listed in Table 3. Examination of these data shows that, in its ground state, NT is very nearly a planar symmetric top ($\kappa = -0.999$) to which a single methyl group is attached ($\Delta f = -3.3$ amu Å\textsuperscript{2}). The data also show that the $S_0$ ZPL of NT is rigid; adding centrifugal distortion terms did not significantly improve the quality of the fit.

As a substituted polyene, NT also can exhibit the property of geometrical isomerism, leading to several possible structures. So again, it is important to compare our results with theory to ascertain which isomer is responsible for the observed spectra. Two such comparisons are shown in Table 3, one using the rotational constants calculated by the HF (6-31G\textsuperscript{*}) method and a second using the constants calculated by the CASSCF method, both sets referring to the $\text{trans,trans}$ structure. (The second set of constants was determined by adding a methyl group to the optimized geometry of $\text{trans,trans}$-OT obtained by Serrano-Andrés et al.\textsuperscript{16}) Again, we find excellent agreement between experiment and theory, showing that the carrier responsible for these two bands is $\text{trans,trans}$-NT.

The two $S_1$ vibrational levels reached in bands NT-1 and NT-3 have significantly different properties. As can be seen from Figure 4, the high-resolution spectrum of band NT-1

![Figure 4](image-url)

Figure 4. High-resolution one-photon fluorescence excitation spectrum of band NT-1 in the $S_1 \rightarrow S_0$ transition of nonatetraene, at 28 950.8 cm\textsuperscript{-1}, in a molecular beam. See the captions to Figures 2 and 3 for more details. The rotational temperature of the fit is 4 K.

The excited-state parameters of the two bands listed in Table 2 were determined separately by fixing the $S_0$ rotational constants at the experimental values shown in Table 1 and varying the $S_1$ rotational constants to obtain least-squares fits. These calculations showed that both excited-state vibrational levels are nonrigid, requiring the full Watson Hamiltonian\textsuperscript{14} for their analysis. For the band at $0_0^+ 76.2 \text{ cm}^{-1}$ (band OT-1), we fit 353 transitions with a standard deviation of 11.0 MHz. For the band at $0_0^+ 411.9 \text{ cm}^{-1}$ (band OT-2), we fit 192 transitions with a standard deviation of 7.2 MHz. Both deviations are less than the observed widths of single rovibronic lines in the two spectra. The complete set of quartic centrifugal distortion terms\textsuperscript{14} is listed in Table 2. However, only $D_{K'}$ significantly improves the fit. The effect of this term is to shift the parallel-type $K_a$ subbands even further to the blue, by $D_{K'} K_a^4$ (cf., Figure 3). No other perturbations were observed in these two rotationally resolved spectra of OT.

![Figure 5](image-url)

Figure 5. High-resolution one-photon fluorescence excitation spectrum of band NT-2 in the $S_1 \rightarrow S_0$ transition of nonatetraene, at 28 954.1 cm\textsuperscript{-1}, in a molecular beam and an approximate deconvolution of its $K_a$ subband structure (cf. Figure 3). Note that the $K_a > 0$ transitions are split into two subbands. This splitting is approximately proportional to $D_{K'} K_a$, where $D_{K_a}$ is the coefficient of the linear angular momentum operator parallel to the $a$ axis.
exhibits the same large, monotonic shifts of its $K_a$ subbands that were observed in bands OT-1 and OT-2 of trans,trans-Octatetraene, signaling a large change in the $A$ rotational constant on $S_1$ excitation. However, a fit of band NT-1 to the distortable rotor Hamiltonian (using the ground-state constants listed in Table 3) revealed no significant additional shifts due to the $D_kK_a$ distortion term, similar to those observed in OT. The $S_1$ vibrational level reached in band NT-1 is rigid. Its inertial parameters are listed in Table 4. These parameters include the quartic distortion terms, but these are relatively small and give only a slight improvement to the fit. We conclude from this observation that this level is the ZPL of the $S_1$ state, as suggested by Petek et al.\(^7\).

The situation is different in band NT-3. Figure 6 shows its rotationally resolved spectrum (as well as that of band NT-4, vide infra). Transitions observed in this spectrum have been successfully assigned, as demonstrated by their use in the combination–difference calculations. However, these assignments showed that the observed $K_a$ structure in band NT-3 does not conform to expectations for either a rigid rotor or a distortable rotor. In particular, as shown in Figure 6, the shift of the $K_a$ subbands is not monotonic, proportional to either $K_a^2$ or $K_a^4$ or some combination thereof. Instead, the shifts of particular subbands appear to be larger for odd values of $K_a$.

This effect is so dramatic that the $K_a = 1$ and 2 subbands exhibit nearly identical shifts from the origin. The most likely explanation of this effect is Coriolis coupling.\(^9\) The $S_1$ vibrational level reached in band NT-3, at $63.2 \text{ cm}^{-1}$, is believed to be the analogue of the $S_1$ level in band OT-1 of octatetraene, a $b_h$ bending mode. According to Petek et al.,\(^7\) it is nearly degenerate with the $3a_1$ methyl torsional level built upon the electronic origin of NT, at a (calculated) frequency of $63.2 \text{ cm}^{-1}$. Therefore, it is reasonable to suppose that the two zero-order levels are coupled by a Coriolis interaction, leading to the observed perturbations in the spectrum, especially since these perturbations appear to be larger for odd $K_a$. (A first-order treatment of the Coriolis interaction in symmetric top molecules leads to terms in the rotational Hamiltonian that are linear in $K_a$.)\(^11\) A more complete analysis of this behavior is in progress.

We turn next to the results for bands 2 and 4 of trans,trans-Octatetraene. These bands also are perturbed but for a different reason. The rotationally resolved spectrum of band NT-2, the blue member of the doublet located at $28\,954.1 \text{ cm}^{-1}$, is shown in Figure 5. Our analysis of this spectrum reveals that each $K_a$...
subband with $K_a \geq 1$ is split into two components, with the center of gravity of the two components shifting to the red with increasing $|K_a|$. Qualitatively similar behavior is exhibited by the $K_a$ subbands in the rotationally resolved spectrum of the blue member of the doublet located at 29 014.3 cm$^{-1}$, band NT-4, as shown in Figure 6.

The origin of these perturbations is the torsion–rotation interaction.\textsuperscript{17} Recall that bands NT-2 and NT-4 have been assigned to transitions that connect the 1e methyl torsional component of the ZPL of the $S_0$ state with the corresponding components of two different vibrational levels of the $S_1$ state. As discussed in detail elsewhere,\textsuperscript{20} the torsion–rotation interaction affects a and e torsional levels differently. The a torsional levels are affected only in second order, leading to modified values of their rotational constants, whereas the e torsional levels are affected both in first and in second order, leading to splittings that depend on $K_a$ as well as modified rotational constants. The observation of such splittings in the high-resolution spectra of bands NT-2 and NT-4 thus confirms their assignment as 1e–1e bands.

We now use the observed splittings to obtain information about the barriers to internal rotation of the methyl group in the $S_0$ and $S_1$ states of trans,trans-NT. The effective Hamiltonian is

$$\hat{H} = \hat{H}_T + \hat{H}_R + \hat{H}_{TR}$$

where

$$\hat{H}_T = \frac{F_p^2 + V_3(1 - \cos 3\phi)}{2}$$

$$\hat{H}_R = A P_a^2 + B P_b^2 + C P_c^2$$

$$\hat{H}_{TR} = D_a P_a + D_b P_b$$

and the rest of the terms have their usual meaning.\textsuperscript{20} We first fit the high-resolution spectra of bands NT-2 and NT-4 using eq 1 to model the rotational energy level patterns in the e torsional components of the connected vibrational levels of the two electronic states. This led to two sets of ground-state constants that were found to be identical within experimental error. Then we used the combination–difference method to extract more accurate values of the inertial parameters of the e subtorsional level of the ZPL of the $S_1$ state from the spectrum of band NT-2. A similar refinement of the fit of band NT-4 was not attempted owing to the presence of additional Coriolis perturbations in this spectrum.

Table 5 lists the values of the parameters that were derived using this procedure. Considering first the rotational constants of the a and e torsional components of the ZPL of the $S_0$ state, we see from a comparison of the values in Tables 3 and 5 that they are nearly the same within experimental error, showing that the second-order contribution of the torsional–rotation interaction is small in the ground state. Making the same comparison for the $S_1$ state (Tables 4 and 5), we see again that the $C^\prime$ rotational constants are the same but that the $A^\prime$ and $B^\prime$ rotational constants are significantly different, showing that the second-order contributions of the torsion–rotation interaction are large in this case, especially for motions about $a$ and $b$. In this connection, it is interesting to note that the errors in the determinations of the $A$ rotational constants of the e components of the ZPL’s of the $S_0$ and $S_1$ states are very small, despite the near-symmetric-top nature of the spectra. This significant improvement in the accuracy of the experiment can be traced to a mixing of $K_a$, $K_e$ states in the affected levels by the torsion–rotation interaction.\textsuperscript{20} These perturbations cause a splitting of the nearly degenerate rotational levels differing only in $K_a$, which are exactly degenerate in the symmetric top limit. Increases in these splittings relative to the resolution of the experiment provide increased accuracies of the determinations of the $A$ constants of the affected vibrational levels.

Considering next the first-order torsion–rotation constants in Table 5, we find $D_{\alpha''} = 0$ and $D_{\alpha''} = 0.19$ MHz for the e component of the $S_0$ state and $D_{\beta'} = 200$ and $D_{\beta'} = 15430$ MHz for the e component of the $S_1$ state. The first-order term $D_{\beta'}$ is very large, as large as the $A^\prime$ rotational constant itself, requiring the inclusion in eq 4 of a third-order correction term (which is neglected in the subsequent analysis). These data show that there is a large decrease in the methyl internal rotation barrier in the $S_1$ state. Combining the data in the usual way\textsuperscript{20} and using as well the observed tunneling splittings in the low-resolution spectrum,\textsuperscript{7} we find $V_3(S_0) \approx 900$ cm$^{-1}$ and $V_3(S_1) = 42.8$ and $V_3(S_1) = -29.4$ cm$^{-1}$. (The $V_3$ term was added to eq 2 to improve the fit.) These values are in good agreement with those determined by Petek et al.\textsuperscript{7} Thus, the observed tunneling splittings are almost entirely attributable to a low torsional barrier in the $S_1$ state.

Finally, we turn to DT. Figure 7 shows the high-resolution spectra of two members (DT-1 and DT-2) of the triplet observed in the DT $S_1 \rightarrow S_0$ spectrum located at 29 014.0 and 29 017.1 cm$^{-1}$ (DT-3 was not recorded owing to its low intensity.) Given the results for OT and NT, it is reasonable to expect (as did Petek et al.\textsuperscript{6,7}) that bands DT-1, 2, and 3 are the tunneling “triplet” of the vibronic origin of DT, 50.9 cm$^{-1}$ above the true origin of the $S_1 \rightarrow S_0$ transition. Examination of Figure 7 shows that these two observed bands confirm qualitatively to this expectation. Both are $a$-type bands. Band DT-1 appears to be a $0a_0 \rightarrow 0a_0$ transition, unperturbed by torsion–rotation coupling, and band DT-2 appears to be a significantly perturbed $1g \rightarrow 1g$ transition. Apparently, the species responsible for these bands is trans,trans-DT. Unfortunately, both bands are too weak to permit detailed analysis at this time. This is probably a result of several factors. Addition of the second methyl group has restored the center of inversion that makes the $S_1 \rightarrow S_0$ transition forbidden in the $C_{2h}$ point group. In addition, its intensity is

### Table 5: Inertial Components of trans,trans-Nonatetraene in the 1e Torsional Components of the Zero-Point Levels of Its Ground and First Excited Electronic States

<table>
<thead>
<tr>
<th></th>
<th>$S_0$ (&quot;1(A_g^2))</th>
<th>$S_1$ (&quot;2(A_g^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>exp\footnote{a}</td>
<td>std error\footnote{a}</td>
<td>exp\footnote{a}</td>
</tr>
<tr>
<td>$A$ (MHz)</td>
<td>16102</td>
<td>2.3</td>
</tr>
<tr>
<td>$B$ (MHz)</td>
<td>390.23(0.1)</td>
<td>0.02</td>
</tr>
<tr>
<td>$C$ (MHz)</td>
<td>382.07(0.1)</td>
<td>0.02</td>
</tr>
<tr>
<td>$D_a$ (MHz)</td>
<td>0.19</td>
<td>0.62</td>
</tr>
<tr>
<td>$D_b$ (MHz)</td>
<td>-99.42</td>
<td>0.06</td>
</tr>
<tr>
<td>$\kappa$</td>
<td>-0.999</td>
<td>-0.999</td>
</tr>
<tr>
<td>$\Delta^f$ (amu $\AA^2$)</td>
<td>-3.56</td>
<td>0.08</td>
</tr>
</tbody>
</table>

\footnote{a} Results from a fit of 423 combination–difference appearances that appear in the rotationally resolved spectra of the two blue components of the doublets that appear in the 0\(_0^e\) and 0\(_0^e + 60.1\) cm$^{-1}$ bands, located at 29 954.1 and 29 014.3 cm$^{-1}$, respectively, using the torsion–rotation Hamiltonian (eq 1). Numbers in parentheses represent estimates of the systematic uncertainties in the experiment. Band origin was not determined owing to the lack of a calibrated $I_2$ spectrum in this region.

\footnote{b} Estimated errors from the diagonal elements of the correlation matrix.

\footnote{c} Results from a fit of 159 assigned transitions in the spectrum of band NT-2, the blue component of the doublet of the 0\(_0^e\) band, located at 29 954.1 cm$^{-1}$. The ground-state parameters were fixed to the values shown in this table. The standard deviation of this fit is 7.03 MHz.

\footnote{d} Neglecting internal rotation parameters.
now spread over three torsional levels having different nuclear spin symmetries. And perhaps most importantly, the vapor pressure decreases as one goes from OT to DT, resulting in progressively lower concentrations in the molecular beam. Owing to the instability of these compounds, this cannot be compensated for by heating.

The inability to observed the blue member of the triplet, DT-3, may indicate the existence of a coupling between the two methyl rotors, despite their large separation. In the absence of such coupling, consideration of nuclear spin statistics suggests that the relative intensities of bands DT-1 and DT-3 should be the same. However, in the presence of such coupling, the normally degenerate members of an ε torsional level, associated with clockwise and counterclockwise rotations of the first methyl group, would have different energies, depending on the sense of rotation of the second methyl group. A splitting would result, leading to a further diminution of the intensities of the affected band, DT-3 in this case. As shown elsewhere, the bands of two coupled rotors also would be expected to exhibit significant torsion—rotation interactions, especially for rotors that are hindered by only small barriers.

Discussion

Eight vibronic bands in the $2^1A_g - 1^1A_g$ electronic transitions of OT, NT, and DT have been recorded at high resolution. Given the thermal instability of the compounds, their near-symmetric-top nature, and the existence of several perturbations, the analyses of these spectra posed significant challenges. Nevertheless, five of the bands have been fully assigned and three have been partially analyzed, providing valuable new information about these polyenes in their ground and first electronically excited states.

Paramount among our findings are the rotational constants of OT and NT in their ground electronic ($1^1A_g$) states (Tables 1, 3, and 5). Alone among all known spectroscopic parameters of an isolated molecule in the gas phase, its rotational constants are the most sensitive to its three-dimensional structure. This information is contained in its principal moments of inertia $I_a$, $I_b$, and $I_c$, which are related to its rotational constants $A$, $B$, and $C$ by

$$I_a = \frac{h}{8\pi^2 A}, \text{ etc.}$$

and to the displacements of each of its component atomic masses from the relevant axes by

$$I_a = \sum_i m_i(b_i^2 + c_i^2), \text{ etc.}$$

Thus, the fact that OT has a large $A$ constant is a consequence of its small moment of inertia $I_a$; the displacements of each of the C and H atoms from the $a$ axis ($b_i$ and $c_i$) are all relatively small (I). In contrast, OT has relatively small $B$ and $C$ constants because the displacements of many of the atoms from the $b$ and $c$ axes are relatively large.

Three rotational constants are clearly insufficient to determine the complete structure of OT and its derivatives, making necessary a comparison with theory in order to establish the identity of the species responsible for the spectra. Fortunately, this comparison leads to an unambiguous result. OT has 20 distinguishable isomers, 15 including $cis^3$ configurations.
TABLE 6: Modeling the Geometry of trans,trans-Octatetraene in Its Ground (1A_g) and Electronically Excited (2A_g) States

<table>
<thead>
<tr>
<th>parameter*</th>
<th>S_0 (1A_g)</th>
<th>S_1 (2A_g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( r(C\sim C) )</td>
<td>model 1 CASSCF</td>
<td>model 2 CASSCF</td>
</tr>
<tr>
<td>( r(C\sim C) )</td>
<td>1.340</td>
<td>1.343</td>
</tr>
<tr>
<td>( r(C\sim C) )</td>
<td>1.460</td>
<td>1.456</td>
</tr>
<tr>
<td>( \angle(C\sim C\sim C) )</td>
<td>1.340</td>
<td>1.340</td>
</tr>
<tr>
<td>( \angle(C\sim C\sim C) )</td>
<td>1.460</td>
<td>1.452</td>
</tr>
<tr>
<td>( \angle(C\sim C\sim C) )</td>
<td>124.0</td>
<td>124.3</td>
</tr>
<tr>
<td>( \angle(C\sim C\sim C) )</td>
<td>12.4</td>
<td>124.0</td>
</tr>
<tr>
<td>( \angle(C\sim C\sim C) )</td>
<td>124.0</td>
<td>124.1</td>
</tr>
<tr>
<td>( A ) (MHz)</td>
<td>19,649</td>
<td>19,711</td>
</tr>
<tr>
<td>( B ) (MHz)</td>
<td>570.5</td>
<td>568.0</td>
</tr>
<tr>
<td>( C ) (MHz)</td>
<td>554.4</td>
<td>552.2</td>
</tr>
</tbody>
</table>

* Bond lengths in angstroms and bond angles in degrees. A C−H bond length of 1.078 Å and H−C−C angles of 1/360 − \( \angle \)C−C−C were used in all calculations.

However, only the trans,trans form (I) has calculated \( B' \) and \( C' \) rotational constants that agree with those measured in this work. For NT, the best agreement between experiment and theory also is obtained for the trans,trans isomer. Here, we find from our fit of band NT-2 the values \( A' = 16.102 \pm 2.3 \), \( B' = 390.2 \pm 0.1 \), and \( C' = 382.1 \pm 0.1 \) MHz, whereas theory (CASSCF)\(^{16} \) gives the values \( A' = 16.090 \), \( B' = 390.0 \), and \( C' = 381.9 \) MHz. Thus, we confirm the conclusions of Petek et al.\(^{2-7} \) The carriers observed in their spectra of OT, NT, and DT are primarily the trans,trans forms. These data also show that modern ab initio theory gives an accurate description of the structures of the isolated molecules in their ground electronic states.

Now, the patterns of energy levels observed in the high-resolution spectra also are sensitive to the size and shape of the molecules in their electronically excited states and therefore to the changes in structure that occur when the photon is absorbed. We find (Tables 2, 4, and 5) that excitation of all three molecules to their S_1 (2A_g) states results in large increases in \( A \) and small decreases in \( B \) and \( C \). This shows that the molecular dimensions perpendicular to \( a \) decrease significantly while those perpendicular to \( b \) and \( c \) increase slightly, compared to the ground state. Despite uncertainties in the \( A' \) constants themselves, the changes in the rotational constants, \( \Delta A (= A' - A) \), \( \Delta B \), and \( \Delta C \), are all known to high precision (±0.1 MHz). These are 2532, -11.7, and -11.0 MHz for the 0\( ^0 \) + 76.2 cm\(^{-1} \) band of OT; 2243, -11.7, and -11.1 MHz for the 0\( ^0 \) + 411.9 cm\(^{-1} \) band of OT; and 457, -6.3, and -6.5 MHz for the 0\( ^0 \) band of NT, respectively. The magnitudes of the changes are smaller in NT. Ab initio theory\(^{16} \) predicts the values \( \Delta A = -88.8 \), \( \Delta B = -12.3 \), and \( \Delta C = -11.7 \) MHz for trans,trans-OT. The agreement between the observed and calculated values of \( \Delta B \) and \( \Delta C \) is very good. However, the difference between the observed and calculated \( \Delta A \) value is very large, much too large to be attributed to the vibrational contributions of an in-plane bending motion\(^{22} \) and/or possible inadequacies of the dihedral angle Hamiltonian. In all cases, we find \( \Delta A > 0 \), whereas theory predicts \( \Delta A < 0 \).\(^{16} \) This shows that modern ab initio descriptions of the structures of the excited electronic states of isolated polyenes are still inadequate.

Intrigued by this result, we have performed some simple calculations to explore the connections between different model OT geometries and their rotational constants. The properties of the model structures are listed in Table 6. For the ground state (model 1), we chose \( r(C\sim C) = r_1 = 1.340 \) Å, \( r_2 = r_4 = 1.460 \) Å, and \( \angle_1 = \angle_2 = \angle_3 \) (C−C−C) = 124.0°, thereby capturing the essence of its structure, alternating C−C bond lengths and C−C−C angles close to 120°. The specific values of these parameters were chosen to most closely match those of the CASSCF geometry of the ground state,\(^{16} \) also listed in Table 6. We ignore the small calculated decrease in the difference between “single” and “double” bond lengths as one moves toward the center of the structure, and the small differences in the bond angles. Despite these facts, model 1 has the rotational constants \( \Delta A' = 19.574 \), \( \Delta B' = 571.1 \), and \( \Delta C' = 554 \) MHz, in reasonable agreement with the experimental results.

Next, we distorted the molecule along one or more of the chosen coordinates to model its possible behavior on excitation to the S_1 state. According to theory,\(^{1,6,24,25} \) the primary characteristic of the S_1 → S_0 transition in the polyenes is bond-order “reversal”. Thus, as shown in Table 6, the CASSCF calculation\(^{16} \) predicts that single bonds shorten and double bonds lengthen on S_1 excitation, whereas the C−C−C angles remain essentially unchanged. Model 2 was chosen to mimic this behavior. Here, we find that decreasing \( r_1 \) and \( r_2 \) from 1.460 to 1.340 Å increases \( A \), \( B \), and \( C \), owing to decreases in the displacements of all atoms from all three inertial axes. Increasing \( r_3 \) and \( r_5 \) from 1.340 to 1.460 Å has the opposite effect. But the magnitudes of the observed changes in \( \Delta A \)−\( \Delta C \) are different in the two cases, since OT (in its ground state) has fewer C−C single bonds than double ones. Thus, when we simultaneously increased \( r_1 \) and \( r_3 \) while decreasing \( r_2 \) and \( r_5 \), we found that \( A \) increases while \( B \) and \( C \) decrease. The primary reason for the increase in \( A \) is that a decrease in \( r_1 \) moves all atoms closer to the \( a \) axis.

The model 2 structure, with \( r_1 = r_5 = 1.460 \) Å, \( r_2 = r_4 = 1.340 \) Å, and \( r_3 = 1.340 \) Å, and \( \theta = 124^\circ \), has \( \Delta A = 137 \), \( \Delta B = -7.9 \), and \( \Delta C = -7.4 \) MHz. To improve the agreement with experiment, we also explored the effect of changes in the C−C−C bond angles. Increasing these angles increases \( A \), since the displacements of the atoms with respect to \( a \) decrease, and decreases \( B \) and \( C \), since the displacements with respect to \( b \) and \( c \) increase. Our model 3 structure, which has all C−C−C angles set to 125.2°, has \( \Delta A = 572 \), \( \Delta B = -14.6 \), and \( \Delta C = -13.5 \) MHz. The calculated \( \Delta A \) value is still in rather poor agreement with the observed one. Nonetheless, it is large and positive, as observed. Thus, it appears that S_1 excitation of the typical polyene results in significant changes in the bond angles as well as the bond lengths. Studies of isotopically labeled species and/or further refinements of theory will be necessary to confirm this finding.

Another dramatic consequence of the change in electronic structure on S_1 → S_0 excitation is the large decrease in the methyl group internal rotation barriers in NT and DT. We find \( V_{\theta}(S_0) \geq 900 \) cm\(^{-1} \), \( V_{\theta}(S_1) = 42.8 \), and \( V_{\theta}(S_1) = -29.4 \) cm\(^{-1} \) in NT. Petek et al.\(^{7} \) find comparable values for NT and DT from their analyses of the low-resolution spectra. Now it is known that the ground states of propene and other molecules having CH\(_3\)−CH=\( X \) functionalities prefer methyl group conformations that have a C−H bond syn or eclipsed with the double bond, as shown below (II). The barrier to rotation is on the order of 700 cm\(^{-1} \). The origin of this barrier can be traced to repulsive interactions between the occupied methyl group orbitals of \( \pi \) symmetry and the \( \pi \) orbitals of C=\( X \).\(^{36} \) However, the situation is different in \( \pi \pi^* \) excited states. Here, the staggered conformation (III) is preferred because the dominant interaction is an attractive one, owing to overlap of the singly
occupied $\pi^*$ orbital and the vacant $\pi^*$ orbital.26 Thus, the magnitude (and sign) of the methyl group internal rotation barrier is a sensitive probe of the $\pi$-orbital bond order in the adjacent $\text{C}=$C bond(s).20,27

Theory (HF/6-31G*)18 predicts that the stable conformation of NT in its ground electronic state is the eclipsed form II, consistent with the existence of a double bond between $C_7$ and $C_8$ and with its measured torsional barrier in the $S_0$ state. However, this barrier drops precipitously on excitation to the $S_1$ state. This shows that there is a large decrease in the $\pi$ bond order in the adjacent $C=$C bond in the $S_1$ state. Thus, the small torsional barrier in the $S_1$ state of NT also provides compelling evidence for bond order reversal in the $\text{S}_1 \rightarrow \text{S}_0$ transitions of these linear polyenes. The $C_2=$C$_8$ bond in trans,trans-NT must be nearly a single bond in the $S_1$ state.

It is interesting to note that the methyl group internal rotation barrier in the $S_1$ state of trans,trans-NT ($V_J = 42.8, V_K = -29.4$ cm$^{-1}$) is comparable to that of the ethyl radical, in its ground state ($V_J \lesssim 50$ cm$^{-1}$).28 As is well-known, Hudson and Kohler29 were the first to show that the $S_1$ state of most linear polyenes is a mixture of singly and doubly excited $\text{A}_g$ configurations rather than the singly excited $B_g$ (HOMO–LUMO) configuration predicted by simple MO theory. Theoretical calculations by Schulten et al.24 confirmed this conclusion. One chemical consequence of this finding is bond-order reversal. A second consequence is two “separated but paired” electrons, mostly localized on the terminal carbon atoms in the $S_1$ state. The ethyl radical also has a largely localized (unpaired) electron, in a $\pi$-type orbital adjacent to the CH$_3$ group. Thus, the fact that the internal rotation barriers in these two species are similar suggests that, in accord with theory, the $S_1$ state of trans,trans-NT also has “single” $\pi$ electrons localized on $C_7$ and $C_8$. The measured barrier is a very sensitive probe of the interaction of these electrons with attached methyl groups.

The rotationally resolved spectra presented in this work also provide compelling evidence for the model proposed by Petek et al.5–7 to explain the source of $S_1 \rightarrow S_0$ oscillator strength in trans,trans-OT, NT, and DT. This information comes from our analysis of the two bands of OT, in which it was found that the two excited-state vibrational levels have unusually large values of the centrifugal distortion term $D_K$ (10.4 MHz in band OT-1 and 13.7 MHz in band OT-2; Table 2). Both levels exhibit small, positive inertial defects (1.08 and 0.80 amu Å$^2$, respectively), whereas the ZPL of the $S_0$ state exhibits a small, negative $\Delta I$ value (−2.14 amu Å$^2$). Both in-plane and out-of-plane vibrations contribute to the inertial defects of planar molecules.17 Usually, in-plane vibrations make a positive contribution and out-of-plane vibrations make a negative contribution. We believe, therefore, that out-of-plane vibrations dominate in the ZPL of the $S_0$ state, given its observed negative inertial defect. Further, since the sign of $\Delta I$ changes on excitation of trans,trans-OT to both of the two examined $S_1$ vibrational levels, we also believe that these two levels involve significant in-plane motions.

C$_8$H$_{10}$ has 48 (nondegenerate) vibrational modes, 33 in-plane ($a_g$ and $b_u$) and 15 out-of-plane ($a_u$ and $b_g$).5 Six in-plane modes have frequencies less than 600 cm$^{-1}$ in the ground state. These are all $C$-$C$-$C$ bending modes: $v_{15}$ ($a_g$) at 543, $v_{16}$ ($a_g$) at 332, $v_{17}$ ($a_g$) at 222, $v_{46}$ ($b_u$) at 566, $v_{47}$ ($b_u$) at 390, and $v_{48}$ ($b_u$) at 86 cm$^{-1}$. (The quoted frequencies are scaled MP2/6-31G* values.)30 Given the frequency displacements in the spectra (76 and 412 cm$^{-1}$), the observed levels are likely the excited-state analogues of two of these modes or linear combinations of them (owing to possible Duschinsky rotations). Petek et al.5 have assigned them as $v_{48}$ and $v_{48} + v_{16}$, placing $v_{48}$ at 76.2 and $v_{16}$ at 335.7 cm$^{-1}$ in the $S_1$ state, close to the ground-state values. However, significant changes in both the nature and the frequencies of these modes could occur on $S_1$ excitation.

The values of the quartic distortion terms derived from the fits of bands OT-1 and OT-2 provide new information about the identity of these modes. The largest distortion term by far is $D_K$, which is at least 3 orders of magnitude larger than $D_J$ and $D_K'$, and much larger than in other “typical” large molecules. Thus, if the source of this nonrigid behavior were a genuine centrifugal effect, we would conclude that the largest force produced by rotation is for rotation about the $a$ axis, when $K_a$ is large. That rotation of OT about its (near) symmetric top axis could lead to distortions of the molecule in directions perpendicular to the axis of rotation is reasonable. But the $D_K'$ value of NT in the ZPL of its $S_1$ state, where similar effects might be expected, is relatively small, $D_K' = 0.73$ MHz (Table 4). Further, only $g$ vibrational modes can be involved in centrifugal distortion.31 Therefore, it is likely that the large $D_K'$ values for bands OT-1 and OT-2 of octatetraene are not truly centrifugal in origin but instead are due to Coriolis coupling involving $v_{48}$ and an out-of-plane vibration. This vibration must be of $a_u$ symmetry to couple about the $a$ axis. As suggested by Watson,31 this mechanism also would explain why $\Delta A$ for OT is unexpectedly large.

The approximate Hamiltonian for the $K$ rotational energies is31

$$E(K) = \begin{bmatrix} v_{48} + AK^2 & 2iA\zeta K & 2AK^2 \\ -2iA\zeta K & v_{25} + AK^2 & \end{bmatrix}$$

Expanding yields

$$E(K) = v_{48} + AK^2 + \frac{4A^2\zeta^2K^2}{v_{48} - v_{25}} - \frac{16A^4\zeta^4K^4}{(v_{48} - v_{23})^3} + \ldots$$

Here, $v_{25}$ is the frequency of an $a_u$ out-of-plane bending mode, calculated to be 60 cm$^{-1}$. If we take $\Delta A = 2$ GHz, $A = 20$ GHz, and $v_{48} - v_{25} = 16$ cm$^{-1}$, we find from eq 8 the Coriolis parameter $\zeta = 0.77$, a not unreasonable result. Therefore, we conclude that the large value of $D_K'$ in band OT-1 is most likely a consequence of Coriolis effects involving $v_{48}$ and $v_{25}$. A similar explanation would account for the large $D_K'$ value in band OT-2, in which $v_{48} + v_{16}$ interact with $v_{25} + v_{16}$. We further conclude that such couplings also could account for the large $\Delta A$ values of these two bands. And finally, we confirm the suggestion of Petek et al.,5 band OT-1 terminates in the analogue of the “overall” $b_u$ in-plane bending mode ($v_{16}$) in the $S_1$ state.

This spectroscopic finding has significant implications for the properties of the isolated molecule. As shown by Petek et al.,5 the origin in the two-photon excitation spectrum of octatetraene is $\sim 76$ cm$^{-1}$ lower in energy than the origin of the one-photon spectrum. We have shown that the latter band originates in the ZPL of the $S_0$ state of the trans,trans form, is an $a$-type spectrum,32,33 and terminates in a vibronic level of the $S_1$ state that has $b_u$ symmetry, lying $\sim 76$ cm$^{-1}$ above the ZPL. Thus, this band (OT-1 at 0$_0^+$, 76.2 cm$^{-1}$) is the “false” origin of the $S_1 \rightarrow S_0$ transition of trans,trans-OT. It gains its intensity via Herzberg–Teller coupling with the $S_1(1^1B_u)$ state, via the $b_u$ promoting mode, which is the origin of most of the remaining vibrational activity in the spectrum.5 As also shown by Petek et al.,5 the complete lack of overlap between vibronic bands in the one- and two-photon spectra proves that the one-photon ($g \rightarrow u$) and two-photon ($g \rightarrow g$) selection rules are strictly obeyed, thus excluding a $cis$ form of OT as the carrier of the spectrum.
It is interesting to note that the CASSCF geometry of the \(1^1B_u\) state of \(trans,trans\)-OT has rotational constants of \(A = 20.148, B = 559,\) and \(C = 544\) MHz. The calculated values of \(B\) and \(C\) for the \(1^1B_u\) state are nearly the same as those for the \(2^1A_g\) state. However, the calculated value of \(A\) for the \(1^1B_u\) state is significantly larger than the corresponding value for the \(2^1A_g\) state. Thus, if the degree of mixing of the two states via the \(b_g\) promoting mode is large, this may account for at least some of the discrepancy between the observed and calculated values of \(\Delta A\) for the \(2^1A_g\) state.

Making such arguments quantitative is a challenging task.\(^{40}\)

In early work, Kohler and co-workers\(^{5–7}\) invoked vibronic coupling between the ground \(1^1A_g\) and excited \(2^1A_g\) states to explain the increase in frequency of the Franck–Condon active, totally symmetric \(\text{C}=\text{C}\) stretch that was observed on excitation of \(2,10\)-dimethylundecapentaene and \(2,12\)-dimethyltetradeca-hexaene. Later, Orlandi and Zerbetto\(^{38}\) showed that, whereas vibronic couplings between states of the same parity are strongly forbidden, and those between states of opposite parity are comparably weak and involve mainly \(\text{C}=\text{C}\) bending modes. More recently, Buma and Zerbetto\(^{39}\) used CI methods to investigate vibronic intensity patterns in the spectrum of \(trans,trans\)-OT. They found that the low-frequency \(b_g\) mode \((\nu_{bg})\) is most active in coupling the two \(1^1A_g\) and \(1^1B_u\) states. They also confirmed that the intensity of the false origin in the one-photon excitation spectrum of \(trans,trans\)-OT is comparable to that of the true origin in an OT containing a \(cis\) linkage, as previously discussed by Petek et al.\(^5\) Future modeling of this effect should focus on \(trans,trans\)-NT, in which both the true and false origins are observed (cf., Figure 1). Quantitative estimates of the vibronic coupling matrix elements are needed to account for the large variations in time scales that have been observed in the excited-state relaxation behavior of several polyenes.\(^{40}\)

**Conclusions**

Rotationally resolved electronic spectra of several vibronic bands in the \(S_1 \rightarrow S_0\) transitions of octatetraene (OT), nonatetraene (NT), and decatetraene (DT) in the gas phase have been observed and analyzed. These analyses show that (1) the first strong band in the one-photon spectrum of NT originates in the zero-point vibrational level (ZPL) of the ground \((1^1A_g)\) state of the \(trans,trans\) isomer and terminates in the \(v_{bg}\) vibrational level of the excited \((2^1A_g)\) state, which lies \(\approx 76\) cm\(^{-1}\) above its ZPL, (2) the one-photon spectrum, strictly forbidden by parity selection rules, gains its intensity via Herzberg–Teller coupling with the \(S_1\) \((1^1B_u)\) state, via the \(v_{bg}\) \((b_g)\) promoting mode, (3) the \(S_1\) state of \(trans,trans\)-OT has a different equilibrium geometry from that of the \(S_0\) state, exhibiting significant changes in the \(\text{C}=\text{C}\) bond angles as well as the \(\text{C}=\text{C}\) bond lengths, as required by bond-order reversal models of the \(S_1\) state, (4) the first two strong bands in the one-photon spectrum of NT originate in the ZPL of the ground \((1^1A_g)\) state of the \(trans,trans\) isomer and terminate in the ZPL and \(v_{bg}\) vibrational levels of the excited \((2^1A_g)\) state, which lies \(\approx 60\) cm\(^{-1}\) above its ZPL (the two bands are, respectively, the true and false origins of the \(S_1 \rightarrow S_0\) transition, the former being no longer strictly forbidden), and (5) the methyl group internal rotation barrier in \(trans,trans\)-NT, large in the \(S_0\) state, decreases significantly on excitation to the \(S_1\) state, further validating the bond reversal model. The observed spectra of DT are consistent with its assignment to the \(trans,trans\) structure. All of these conclusions agree with the earlier assignments and interpretations of Petek et al.\(^{5–7}\) The excited-state rotational constants and other experimental parameters derived in this work will serve as benchmarks for future theoretical studies of the structural and dynamical properties of linear polyenes.

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**References and Notes**


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\(^{22}\) The vibrational contributions of \(v_{bg}\) to the rotational constants of the ground state are estimated to be \(\Delta A = -1402, \Delta B = +3,\) and \(\Delta C = 95\).
+3 MHz from the HF/6-31G* calculations (ref 18). Thus, $\Delta A$, although having the right order of magnitude, has the wrong sign to explain this discrepancy.

(23) Negative $\Delta A$ values have been found in DPB (ref 12) and $\ell$s (ref 13), but in these cases the observed changes appear to be associated primarily with increases in the lengths of the C–C bonds in the phenyl rings that are most perpendicular to $\alpha$.


(32) In this connection, it is interesting to note that we find no evidence for an “off-axis” orientation of the $S_1 - S_0$ transition moment in any of our spectra, such as has been observed for some linear conjugated polyenes as guests in the channels of urea crystals by Hudson and co-workers (ref 33). However, we cannot rule out the possible existence of such a tilt; it could be as large as 15°.


