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Observation and control of all-*trans*- β -carotene wavepacket motion using pump-degenerate four-wave mixing

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Abstract. Wavepacket dynamics on the ground and optically dark, first electronic state of all-*trans*- β -carotene are studied with 16 fs time resolution using pump-degenerate four-wave mixing spectroscopy. Moreover control over the vibrational ground state modes is shown.

Carotenoids, being part of light-harvesting complexes (LH) play an important role in photosynthesis. They absorb light in blue-green and efficiently transfer energy to chlorophylls. The electronic states of interest to this study are the electronic ground $1^1A_g^-$ (S_0), the first optically allowed $1^1B_u^+$ (S_2) and the optically dark state $2^1A_g^-$ (S_1) [1]. The long time kinetics of β -carotene have been studied extensively [1,2] and only very recent work has been successful in monitoring the coherent dynamics of the very high frequency modes of S_0 using transient absorption [3]. To observe the wavepacket dynamics in S_1 poses an extra challenge besides the short pulse duration required, since this state is not directly optically accessible from the ground state, but only via the population transfer from S_2 to S_1 . This pathway seems to be however incoherent [3], thus making simple pump probe measurements impossible. Recently it has been realized that by a pulse sequence that incorporates stimulated emission pumping (SEP) coherent motion in dark states can be observed [4]. FSRS is able to collect vibrational spectra of S_1 with high temporal and spectral resolution, but it is not sensitive to the relative phase between modes, thus a wavepacket motion cannot be reconstructed from this data.

In this work we apply degenerate four-wave mixing spectroscopy (DFWM) combined with narrowband detection (1nm spectral resolution) to probe and control the wavepacket dynamics in S_0 . In combination with a pump pulse (pump-DFWM) we are able to monitor coherent dynamics of the dark S_1 state. The vibrational spectrum of S_0 and S_1 has been observed in several picosecond resonance Raman studies and has been recently studied with <100 fs time resolution with femtosecond-stimulated Raman spectroscopy (FSRS) [5]. Compressed output pulses of 10 μ J from a NOPA are frequency tunable from 400-700nm and have a typical pulse duration of 16 fs. The output of one NOPA is split into the 3 DFWM beams. The pulses within two of the beams are passed

through an all-reflective pulse shaper to control their temporal profile. The second NOPA delivers the pump pulse for the pump – DFWM experiments used for probing S_1 dynamics. A 300 μ m sample holds all-*trans*- β -carotene dissolved in cyclohexane. The data obtained for resonant DFWM on the S_0 - S_2 transition show pronounced wavepacket oscillations for red shifted small bandwidth detection at 546 nm (Fig. 1 (a)). Fourier transformation reveals that the wavepacket is due to coherent S_0 mode dynamics (Fig. 1 (b)). The low frequency peak at $\sim 400\text{cm}^{-1}$ originates from beatings between the strongest β -carotene modes. No dynamics from S_2 can be observed, possibly due to its fast decay. The S_0 C=C stretch mode requires a more red shifted detection compared to the lower modes.

Non-resonant DFWM on the S_0 - S_2 transition (center wavelength 555 nm) at higher pulse energies excites and probes a ground state wavepacket via Raman excitation ($t_{12}=0$). Again wavepacket motion is only observed for red shifted small bandwidth detection (see Fig. 1 (a)). Shaping the time coincident pump and Stokes into a pulse train allowed the selective excitation of the C-C stretch ground state mode (see Fig. 1 (a)).

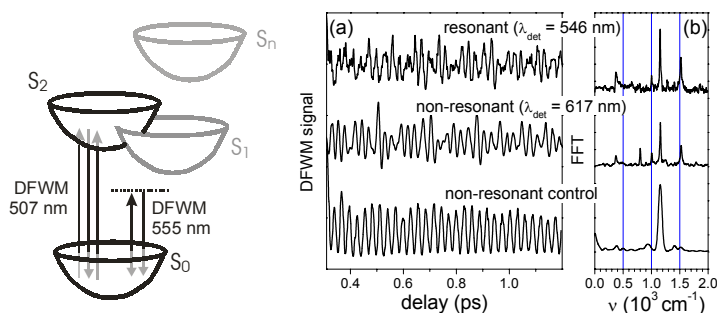
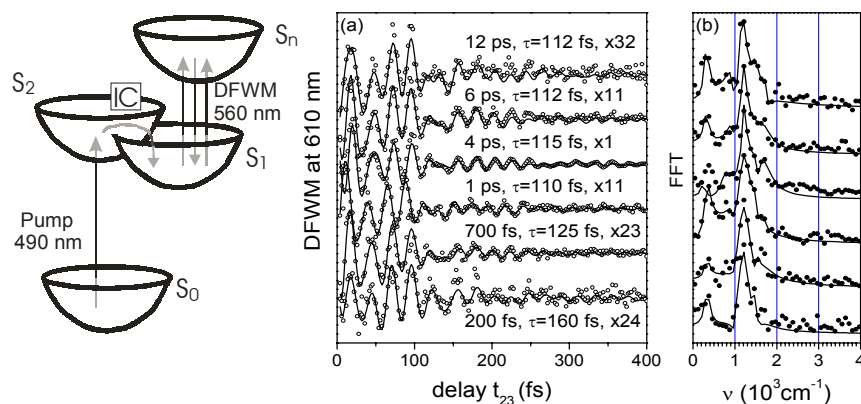


Fig. 1. Resonant and non-resonant DFWM data of the ground state S_0 . Also shown is the control of the S_0 wavepacket using pulse train excitation.

We proceed with the first time observation of coherent motion in the S_1 / hot S_1 state of all-*trans*- β -carotene. The S_1 state can not be directly excited from S_0 , and thus can only be populated via transfer through the conical intersection between S_2 and S_1 . Since strong evidence exists that this population transfer occurs incoherently, it is necessary to create coherent motion in S_1 by a second optical excitation process in order to be able to probe the coherent time dynamics of S_1 modes. Therefore in our experiments we use a pump pulse at 490 nm previous to the DFWM sequence at 560 nm (the center wavelength of the S_1 - S_n absorption band) to transfer population from S_0 to S_2 (see schematic in Fig. 3). This technique is known in literature as pump-DFWM [6]. The DFWM sequence at 560 nm is used to probe and excite the incoherent population originating from the population pumped into S_2 and transferred via conical intersection to S_1 . In order to avoid non-resonant excitation of the ground state a much reduced pump and DFWM pulse energy were adjusted. A phase-locked chopper was used to periodically block the S_0 - S_2 resonant pump. In all transients the DFWM pump and

Stokes were time coincident ($t_{12}=0$). No signal is observed when the pump is in its off state, while for “pump on” S_1 wavepacket dynamics could be observed when detecting at 610 nm (see Fig. 2 (a)). The oscillatory motion is located on top of an exponential decaying signal, with a decay constant of $\tau \sim 112$ -160 fs depending on the pump delay. Note that the wavepacket motion is strongest if the pump



precedes the DFWM sequence by 4 ps.

Fig. 2. Wavepacket dynamics of the optically dark S_1 state of all-trans- β -carotene probed with pump-DFWM. (a) DFWM signal with “pump on” detected at 610nm, for different pump delays. Without pump no signal is observed. The oscillations are on top of a signal that decays exponentially with rate τ . Scaling factor x of the data, with respect to the transient recorded at a pump delay of 4 ps. (b) Fourier transform of the transients in (a).

Evidence for the S_1 nature of the observed modes is provided by a multimode fit model that fits all transients and spectra with good accuracy.

In summary we have shown that pump-DFWM in combination with spectral detection of 1nm resolution can be successfully applied to the study of complex molecular systems, where optically dark electronic states are common.

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