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Direct observation of the ultrafast electron transfer process in a polymer/fullerene blend

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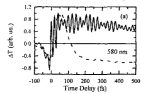
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Photoinduced electron transfer in organic molecules is a topic of fundamental interest in photophysics and for applications to artificial photosynthesis Highly efficient ultrafast electron transfer from photoexcited conjugated polymers to C₆₀ has been reported [1]; the back transfer is leavily hindered, thus providing a stabilization mechanism of the photogenerated charges. The detailed time resolution of this process is needed to shed light on the charge transfer mechanism. In this work we perform ultrafast experiments on conjugated polymer/C₆₀ blends with sub-10-fs time resolution. We are able to time resolve for the first time the charge transfer process, obtaining a time constant τ_a ≈ 45 fs. The excitation source is a visible optical parametric amplifier [2] providing ultrabroadband pulses with sub-10-fs duration and spectrum extending from 320 to 720 nm. We excite thin films of poly [2-methoxy, 5-(3.7" – dimethyl-loctyloxy)]-p-phenylene vinylene (MDMO-PPV), blended with C₆₀, with a 1:1 ratio in molecular concentration, at these high C₆₀ concentrations we measure the intrinsic charge transfer process unaffected by diffusion. For comparison we also study pure MDMO-PPV films.

Fig. 1(3) shows as solid line the differential transmission (ΔT) signal for pure MDMO-PPV at

Fig. 1(a) shows as solid line the differential transmission (ΔT) signal for pure MDMO-PPV at Fig. 1(a) shows as solid line the differential transmission (ΔT) signal for pure MDMO-PPV at the probe wavelength of 580 nm. The features at negative and near-zero delays are due to coherent artifacts. The ΔT signal has positive sign and is assigned to stimulated emission from the excited state The strong oscillations superimposed on the signal (see Fourier spectrum in Fig. 1b) reflect the therational modes coupled to the excited state. By blending with C_∞, the signal changes dramatically (dashed line in Fig. 1(a)). The initially positive ΔT rapidly changes sign, indicating the quick formation of a new photoinduced absorption (PA). Based on cw measurements, this new PA is assigned to the MDMO-PPV charged state (polaron) and thus provides a direct signature for the charge transfer process By an exponential fitting of the PA risetime, a time constant for the electron transfer process of ≈ 45 fs is calculated. Similar results are obtained at other probe wavelengths Excited state vibrational coherence is not maintained in the charge transfer process, thus indicating evolution of the system on an anharmonic charge transfer potential energy surface



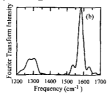


Fig. 1 (a) ΔT for pure MDMO-PPV (solid line) and PPV/C₈₀ (dashed line) at the probe wavelength of 580 nm; (b) Fourier transform of oscillatory component of the MDMO-PPV signal.

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Bright and Dark Resonances in Er:YAG: Generation, Measurement and Coherent Control

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Max-Planck-Institut fuer Feaskoerperforschung Heisenbergstr. 1, D-70569 Stuttgurt, Germany window 1 0-1.1 ps in Fig. 2. It is obvious that the two interferograms have opposite phase, making coherent control possible to chance up. Germany window 1 0-1.1 ps in Fig. 2. It is obvious that the two interferograms have opposite phase, making coherent control possible to chance up. Germany window 1 0-1.1 ps in Fig. 2. It is obvious that the two interferograms have opposite phase, making coherent control possible to chance up. Germany window 10-1.1 ps in Fig. 2. It is obvious that the two interferograms have opposite phase, making coherent form top opposite phase, making coherent form the phase opposite phase two interferograms can be explained with the excitation spectra and the coherent (In) dyramics regording the single ion and pair-Summary

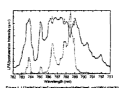
Rare earth ions (RE) with optically activated 4f electrons
present sharp spectral transitions in condensed matter due
to the shielding of the 4f electrons by the outer 5x5p
electrors. The 'quantum doi' rature of RE makes them
ideal for the study of condensed matter structures and
dynamics. In this communication, we report on the
fluorescence excitation spectra and the coherent

dynamics. In this communication, we report on the fluorescence excitation spectra and the coherent femosecond (fis) dynamics regarding the single ion and pair-transitions in EVA Oct at 77K. We further apply fa interferometric techniques to control the single ion and ionspair emissions.

The continuous wave Ti-supphire laser output with the wavelength turable near the 800 nm spectral region excites the Er from the ground state to an intermediate excited state reading in internal fluorescence (LF) Previous work demonstrated that the rare earth optical fransitions are "inhomogeneously broadened", with pair transition salelliter responsible for upconversion normally obscured by a strong linear absorption to their hand, recording of the up-converted signal can separate the pair transitions from a strong linear absorption beokeground.

Fig. 1 shows the fluorescence excitation spectra recorded for LF (upper curve) and for up-conversion (lower curve). The depth and spectral profiles of bright and drak resonances at 786.5, 781.8, 785.5, and 789.0 mm are found to be excitation intensity dependent. The bright resonances of the dark resonances in the LF excitation spectra. The excitation across sections of the two processes can be written as

durk resonances in the LF excitation spectra. The excitation cross sections of the two processes cane be written so $\frac{L(\cdot, 1) + ^{ML}(\cdot, 1) = ^{m}(\cdot, I = 0)}{\text{the Ce}^{L}, \beta^{NL}} \text{ and } \gamma^{W}$ are the absorption cross sections for the LF, upconversion and for LF recorded at very low intensity respectively. The simultaneously recorded ferniosecond interferograms for LF and for up-conversion are shown in the typical time



excitation pulse pair by the phase delay of a References:

1 W.M. Yen in "Spectroscopy of solids containing rare earth ions", Ed by A.A. Kaplyanskii and R M Macfarlane, North Holland, (1987), 185.

2. S. Mukamel, "The principles of nonlinear spectroscop Oxford University Press, New York(1995)

5 (t_d; _d)+5 (t_d) = 0 for large time oney suggether production of the phase shift of n between the two interferograms, making it possible to enhance linear or nonlinear conversion efficiency simply by delaying the

 $S(t_d; d) = \int_{-\infty}^{\infty} d \cos(t_d + d) I(t_d) dt_d + \Omega (2)$ Here t_a is the delay time between the two exciting pulses. Ω is the central frequency of the pulses and ϕ_a is the related phase delay. $I(\alpha)$ is the exciting spectrum and α' general absorption cross section for LF or for upconversion Replacing equation (1) into (2) gives rise to $S^{L}(t_{d}; d) + S^{NL}(t_{d}; d) = S^{\kappa}(t_{d}; d)$ (3) As the spectral bandwidth of the excitation spectra for LP is in general much broader than the burned spectral hole width, S^{*} goes to zero as a function of t_{c} much faster than the terms on the left hand side. As a result, $S^{L}(t_{d};\ _{d})+S^{NL}(t_{f};\ _{d})=0 \ \ \text{for large time delay suggests}$

