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SUBPICOSECOND SPECTROSCOPIC STUDIES OF SINGLET EXCITON FUSION IN MOLECULAR SOLIDS

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Abstract – The first kinetic data displaying a time-dependent \( t^{-1/2} \) singlet exciton annihilation rate has been observed. Subpicosecond optical excitation of polycrystalline thin-film samples of \( \beta \)-hydrogen phthalocyanine resulted in excitonic saturation of the crystal lattice. The initial exciton decay yields the only unambiguous measurement of a characteristic annihilation interaction rate for molecular excitons on adjacent lattice sites. Spectroscopic evidence suggests that fusion events produce vibrationally athermal ground state molecules.

Energy transfer together with exciton motion and decay in molecular solids has been persistently studied over the past twenty years.\(^{1-8}\) Several microscopic theories and pictures have been presented to explain frequency as well as time domain measurements and observations. To date however, no time resolved measurements on any molecular crystal system have shown any evidence of the the microscopically detailed and discrete nature of these dynamical processes. In this paper we present and discuss the first such data.

Previous time-resolved studies on singlet exciton annihilation have monitored wavelength integrated fluorescence intensity following pulsed optical excitation to probe excited-state population densities. In the present study, we employ the technique of sub-picosecond time-resolved absorption spectroscopy which has the advantages of combining laser pulsewidth-limited temporal resolution with valuable spectroscopic information obtainable as broad-banded optical absorption spectra.

Samples consisted of polycrystalline thin-films of \( \beta \)-hydrogen phthalocyanine (H\(_2\)Pc). While not the classic subject of investigation that for instance anthracene has been, H\(_2\)Pc has several important properties critical to the success of the present measurements. The material is highly absorbing at our laser fundamental (16,000 cm\(^{-1}\)) with \( \alpha \approx 2 \times 10^5 \) cm\(^{-1}\). High excitation densities (ca. \( 10^{21} \) cm\(^{-3}\)) are therefore obtainable with available excitation energies and intensities. H\(_2\)Pc is readily sublimed to form good optical quality amorphous films, which can be subsequently annealed to yield polycrystalline samples. These films have been well characterized.\(^{9-12}\) Spectroscopic and crystallographic evidence supports the notion that the individual microcrystallites (which have a flake-like morphology and in-plane dimensions of 10-100\( \mu \)) are isomorphous with the \( \beta \)-form of the macroscopic single crystal. The films showed a remarkable resistance to optical damage at all reported excitation intensities. Finally, H\(_2\)Pc exhibits an intense excited state absorption spectrum, with transitions in the visible at least five times stronger than those observed in anthracene.
Measurements were performed utilizing amplified laser pulses obtained from a colliding pulse ring dye laser. These pulses were 0.15 ps FWHM, centered at 625 nm (16,000 cm$^{-1}$) and occurred at a repetition rate of 10 Hz. Pump and probe measurements were performed utilizing pulses at the fundamental wavelength and white light continuum pulses respectively. Details of the experimental technique have been presented previously.

Figure 1 displays the ground state absorbance spectrum of a typical H$_2$Pc sample together with transient absorbance difference spectra taken at three different delay times. The sample was held at 4K. The difference spectra reveal regions of bleaching (due to ground state depopulation) in addition to regions of excited state absorption.

As the delay time was increased, a dramatic evolution of the entire difference spectrum was observed. Over the first 10 ps subsequent to excitation, there was a rapid decay of the initially induced transient spectrum. This is due to a nonradiative process which efficiently depopulates the initially prepared exciton population. We choose to probe kinetics at a spectral position where the observed signal can be unambiguously related to the exciton number density. Such a spectral region occurs in the area of excited state absorption between 19,000 cm$^{-1}$ and 22,000 cm$^{-1}$, where due to a minimal ground state absorption, there can be little contribution from ground state recovery processes manifest in the signal. We observe the spectral shape of the excited state absorption in this region to remain constant throughout the decay. In contrast, the spectral evolution occurring at energies below 16,000 cm$^{-1}$ is quite complex. In particular, a dramatic increase in sample absorption occurs at roughly 13,500 cm$^{-1}$, where an initial net bleaching rapidly evolves into a persistent net increase in sample absorbance lasting longer than 200 ps.

Figure 2 displays single wavelength pump and probe data obtained with pump pulses at 16,000 cm$^{-1}$ and probe pulses at 19,500 cm$^{-1}$ for two different pump intensities. The decay of these signals was observed to be highly non-exponential. These kinetic traces were observed to be independent of sample temperature over the range 4K to 300K.
We note that the initial rapid decay of the induced absorbance signal is due to singlet exciton-exciton annihilation. This non-radiative phenomenon has been observed in many kinetic studies of molecular crystals where high densities of excitons are created. This certainly corresponds to our experimental conditions, where with the highest intensity excitation pulses (55 μJ/pulse, $7 \times 10^{10}$ W/cm²), we estimate exciton densities of approximately $1.0 \pm 0.5 \times 10^{21}$ cm⁻³. Single crystal specific gravity measurements indicates a value for total saturation to be $1.68 \times 10^{21}$ cm⁻³. (15)

A simple phenomenological rate equation describing exciton decay is often written:

$$\frac{dN}{dt} = -kN - \gamma N^2$$

(1)

where $k$ is the time-independent "unimolecular" decay rate describing a reciprocal exciton lifetime in the low density limit, and $\gamma$ is the time-independent "bimolecular" annihilation rate constant. For the present discussion we disregard the unimolecular term in Eq. (1). At the exciton densities at which our experiments were performed this term describes a rate process much slower than the bimolecular term.

In principle, time-dependent energy transfer rates are expected in measurements of exciton trapping and annihilation. This arises from the fact that subsequent to the initial creation of a spatially homogeneous exciton population, proximate pairs of excitons, or excitons closest to traps, will interact first. Progressively greater and greater remaining mean interparticular distances result in decreasing characteristic interaction rates. Both motion-limited diffusion theory, and the Förster long range dipole-dipole interaction picture (in the absence of diffusion) lead to a $t^{-1/2}$ dependence for either $k$ or $\gamma$. (4,16)

For annihilation dominated kinetics, equation 1 is written to explicitly include the time dependence of the rate:

$$\frac{dN}{dt} = -\gamma t^{1/2} N^2$$

(2)

Integration of equation (2) yields an expression for the exciton number density as a function of time:

$$N(t) = \left(2\gamma t^2 + \frac{1}{N_0}\right)^{-1}.$$
interaction of excitons on adjacent lattice sites. The extrapolation of the straight line though the
data in Figure 3 is therefore expected to indicate excessive values of $N_0$ at the origin of time.

Excitonic saturation of the $\text{H}_2\text{Pc}$ lattice facilitates a clear and simple interpretation of the initial
rapid signal decay. Since the majority of excitons will have excitons as nearest neighbors, we
propose that a single exponential fit to the data at the earliest delay times corresponds to the
lifetime of proximate excitons. Over the first picosecond, such a fit is quite good yielding a
temperature-independent lifetime of 5.7 ps (Fig. 4). This observed lifetime was independent of
pump intensity for intensities above ca. $3 \times 10^{10}$ W/cm$^2$. We expect that as the sample becomes
saturated, the early time decay rate will be dominated by the single exponential rate associated
with proximate exciton pairs. As such, this study presents the only measurement of an exciton
annihilation rate free from the ambiguity or possible influence of exciton motion.

![Figure 3](image3.png)

![Figure 4](image4.png)

Data shown in Figure 2 is the first to manifest a clear and unmistakable time-dependent
annihilation rate. The rate law (Eq. 2) is followed down to delay times of roughly 0.7 ps. We
suggest that at least for the case of $\text{H}_2\text{Pc}$, the rate associated with pure annihilistic interactions is
quite small compared with approximations sometimes made for exciton hopping rates. As
Kenkre and Schmid suggest, the "motion limited" nature of previous kinetic observations of
energy transfer in molecular crystals should not be taken for granted.\(^{(17)}\)

Finally we discuss the spectral dynamics appearing in the low energy region of Figure 1. As
excitons annihilate, ground state molecules reappear. If these molecules return to the ground
state in exactly the same vibrational states as they left, the spectral features in Figure 1 should
monotonically relax toward zero. This is clearly not happening. We attribute the increase in
spectral density at ca. 13,500 cm$^{-1}$ with increasing delay time to the reappearance of
vibrationally hot ground state molecules.

With the aid of Figure 5, we argue that annihilation events produce electronic ground state
molecules with athermal distributions of vibrational energy. Early delay time difference spectra
(1 ps) taken at three different ambient temperatures (Figure 5, top) reveal spectral broadening
with increasing temperature. This is most evident in the region of bleaching between 13,000 and
14,000 cm$^{-1}$.

Molecules leaving the ground state at 4K and returning at 300K would result in the spectrum
modeled in the bottom of Figure 5. By subtracting a percentage of the 300K spectrum from the
4K spectrum, the dashed curve in the bottom of Fig. 5 is derived. The weighting factor is determined by the observed degree of recovery of the ground state (monitored at 20,000 cm\(^{-1}\)). While qualitatively delivering spectral density in the region of 13,500 cm\(^{-1}\), this construct is insufficient to quantitatively explain the observed results.

![Figure 5](image)

Furthermore, a 300 degree increase in lattice temperature is unreasonable. We calculate a maximal 80 degree temperature rise upon final thermalization of all excitation photons. Rather, we suggest a vibrationally selective conversion of electronic energy into vibrational bottlenecks as a plausible explanation for these results.

Although there are certainly other explanations for this spectral evolution, we expect that transition probabilities describing exciton annihilation must contain vibrational overlap integrals. Franck-Condon considerations should therefore be in evidence. Recent observations of nanosecond \(T_1\) times in molecular crystals add further plausibility for this argument\(^{(18)}\). One might easily envision far-IR solid state dye lasers pumped by visible light based on these phenomena.

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**References**


