VIBRONIC RELAXATION IN THE $S_1$ STATE OF RHODAMINE DYЕ SOLUTIONS

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The relaxation of vibronic energy levels was investigated by two different techniques. (A) The transmission of picosecond pulses was measured as a function of the incident peak intensity and (B) the transmission of a second probe pulse was studied versus delay time. Population lifetimes of the excited levels between 0.5 ps and 1.0 ps were found for rhodamine 6G and rhodamine B in the solvents water, ethanol, ethylene glycol and glycerol. The lifetimes are independent of solvent viscosity.

The relaxation of excited rhodamine molecules in the $S_1$ state has been studied by numerous authors [1—7]. Several papers [1—3] were concerned with the delayed amplification of a probe pulse after excitation by an intense pump pulse. Other investigations [4—6] measured the delayed appearance of Stokes shifted spontaneous fluorescence. In all these experiments the emitting and the corresponding final states are not known. The emitting vibrational levels vary during and immediately after the excitation pulse until the molecules settle into the relatively long-lived lowest $S_1$ state.

In this letter we report on measurements which determine the population lifetime $\tau$ of vibrational energy states located several hundred wave numbers above the bottom of the first excited singlet state $S_1$. Two different experimental techniques were used to determine the same relaxation time $\tau$.

(A) The energy transmission $T_L$ of a single intense picosecond laser pulse was measured as a function of input peak intensity $I_{0L}$. The nonlinear transmission of the molecules is compared with model calculations to provide the desired time constant $\tau$. Very rapid relaxation of the excited states leads to less effective bleaching at a fixed input intensity. We note that for relaxation times $\tau$ longer than the pulse duration $t_p$, only one half of the molecules can be excited while for relaxation times $\tau$ shorter than $t_p$ all molecules can be promoted to excited energy states.

(B) After the excitation by the first strong light pulse, the energy transmission $T_L$ of a weak interrogating pulse was measured as a function of delay time $t_D$. The probe pulse had the same frequency, pulse duration, polarization and spatial distribution as the pump pulse. This probe pulse monitors the transient population of the initially excited vibrational states. The probe transmission is equal to the pump transmission when pump and probe beam coincide ($t_D = 0$). At a delay time of $t_D \approx t_p/2$ the probe transmission is largest. The transmission of the delayed probe pulse decreases while the molecules decay from the optically excited energy states to intermediate levels. If the relaxation time $\tau$ is considerably shorter than the exciting pulse duration $t_p$, the molecules relax during the excitation pulse and the transmission of the probe pulse remains constant with increasing delay.

Our experimental set-up is depicted schematically in fig. 1. A mode-locked Nd-glass laser system was used in our investigations [8]. Single pulses were selected from the beginning of the pulse trains with an electro-optical shutter and passed subsequently through a Nd-glass amplifier. The amplified single picosecond light pulses were band width limited with a duration of approximately 5 ps (fwhm), a band width of $\Delta \nu \approx 3$ cm$^{-1}$ (fwhm) and an energy of ca. 3 mJ. The light pulses were frequency doubled in an ADP-crystal with a
conversion efficiency of ≈ 20%. The fundamental light at $\lambda = 1.06 \mu m$ was eliminated by a filter F. The duration of the green light pulses ($\lambda = 5285 \text{ Å}$) was measured by a two-photon fluorescence system.

9,10-diphenyl-anthracene dissolved in cyclohexane ($10^{-3} \text{ M/£}$) was used as two-photon absorber. A colliding beam arrangement was employed and the fluorescence light was imaged onto the vidicon of an optical multi-channel analyser (OMA). Pulse durations in the range between 3 ps and 5 ps were measured with an accuracy of 0.4 ps.

In fig. 1 the experimental arrangement is shown for the simultaneous measurement of the energy transmission of the intense picosecond light pulses through rhodamine 6G dissolved in ethanol. The open circles represent averages over five laser shots. Single pulses of duration $t_p = 3.5 \pm 0.5$ ps were chosen for the data points. It is readily seen from the figure that the transmission of the sample increases strongly from a low intensity value of $T_0 = 0.08$ to $T_L = 0.57$ for input peak intensities of $I_{0L} = 6 \times 10^9 \text{ W/cm}^2$. The two curves in fig. 2 are calculated for population life-
times of $\tau = 0.5$ ps and $\tau = 2.0$ ps using an energy level model which will be discussed below. The theoretical curves account well for our experimental data suggesting a time constant of $\tau \approx 1$ ps.

(B) The transmission of a subsequent weak probe pulse allows an independent determination of the relaxation time $\tau$ of the initially excited energy states. For a fixed intensity of the pump pulse, the energy transmission $T_1$ of a parallel polarized interrogating pulse is determined. The ratio of the energy transmissions $R = T_1/T_L$ changes as a function of delay time $t_D$ depending upon the time constant $\tau$. For $\tau \gtrsim t_p/4$ the time dependence of the $R(t_D)$ curves allows an accurate determination of the relaxation time. For $\tau \lesssim t_p/4$ the shape of the $R(t_D)$ curves becomes similar and the relaxation time has to be determined from the absolute values of $R(t_D)$.

We have measured the energy transmission ratio $R$ as a function of delay time $t_D$ for various solutions and compared it with calculated curves as indicated below. Concentrations of $10^{-4} \text{ M/l}$ and sample lengths of 0.2 cm (rhodamine 6G) and 0.5 cm (rhodamine B) were used. In fig. 3 we present experimental data (open circles) for rhodamine 6G in ethanol. Fig. 3 shows quite vividly the rapid rise of the probe transmission around $t_D \approx 0$ and the decay of the transmission values for longer delays $t_D$ between the pump and the probe pulse. For delay times $t_D$ long compared to $\tau$ the decay of the curves is determined by the orientational relaxation time of $\tau_{or}$ which is known from the literature [11]. Our experimental points of $R(t_D)$ are well accounted for by a theoretical curve with relaxation parameter $\tau \approx 0.7$ ps.

Similar results obtained for rhodamine 6G in water, ethylene glycol and glycerol and for rhodamine B in water, ethanol and glycerol at room temperature (see table 1) and for rhodamine 6G in ethanol at 80 K.

We turn now to a quantitative discussion of our observations. A five-level system appears to be appropriate for a dynamic description of the experimental situation. Fig. 4 shows schematically the considered level system of the rhodamine molecules (right) together with the measured absorption spectrum (left) of rhodamine 6G in ethanol. The calculated model has the following features:

(i) Prior to optical excitation thermal equilibrium exists in the $S_0$ ground state position (1) with energy spread $\Delta E \approx kT \approx 200 \text{ cm}^{-1}$ at $T = 300 \text{ K}$.

(ii) A linear polarized picosecond light pulse of frequency $\tilde{v}_L = 18910 \text{ cm}^{-1}$ excites molecules to a non-equilibrium position (2) in the $S_1$ state (Franck-Condon state with energy spread equal to that of the ground state). A constant absorption cross section $a_{12}$ of the thermally populated ground state levels is assumed since transition probabilities to the respective $S_1$ states are expected to have nearly equal Franck-Condon factors.

(iii) The energy states (2) are emptied by vibrational relaxation and reorientation to a temporary equilibrium position (3) in the $S_1$ state. The position (3) is situated approximately $600 \text{ cm}^{-1}$ and $1300 \text{ cm}^{-1}$ below the position (2) in rhodamine 6G and rhodamine B solutions, respectively.

(iv) Most molecules in the $S_1$ band return to the ground state by fluorescence emission. The fluorescence lifetimes are in the nanosecond range with $\tau_F \approx 4.5 \text{ ns}$ and $\tau_F \approx 2.3 \text{ ns}$ for rhodamine 6G [12] and rhodamine B [13], respectively. Complications due to
<table>
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<th>Solution</th>
<th>$10^{16} \sigma_{12}$ (cm$^2$)</th>
<th>$10^{16} \sigma_{15}$ (cm$^2$)</th>
<th>$10^{16} \sigma_{ex}$ (cm$^2$)</th>
<th>$\eta$ (cP)</th>
<th>$\tau_{or}$ (ps)</th>
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<td>1.2 ± 1</td>
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Fig. 4. Absorption spectrum of rhodamine 6G in ethanol ($10^{-5}$ M/l, 1 cm cell length) and schematic five-level model for the transitions in rhodamine dyes.

Super-fluorescence do not occur in our experiments as checked by measurements of the fluorescence decay times and the emission spectra of our dye solutions. Repopulation of the ground state by fluorescence is neglected.

(v) Molecules in the $S_1$ state absorb pump light of $\nu_L$ by excited state absorption and thereby populate higher singlet states (position 5). A substantial accumulation of population in these higher singlet states does not occur since the excited state absorption cross section $\sigma_{ex}$ is small and the relaxation to position (3) is expected to be in the picosecond range. We assume a constant value of $\sigma_{ex}$ for all molecules which are in the electronically excited state.

(vi) Transfer of molecules from the $S_1$ state to a triplet state is negligibly small [14].

(vii) Strong electric dipole transitions are responsible for the excitation of the molecules to higher lying electronic states. The dipole transition moments have fixed directions within the molecule. The complex dye molecules can be described classically as linear oscillators [15]. The strength of the light absorption depends upon the angle $\theta$ between dipole transition moment and polarization of the light beam. The absorption cross section is given by

$$\sigma(\theta) = \sigma_{12} \cos^2 \theta + \sigma_{ex} \sin^2 \theta$$

for the molecules investigated here. The transition moments for ground state absorption $\sigma_{12}$ and excited state absorption $\sigma_{ex}$ have the same direction as shown by fluorescence polarization measurements [16]. At the beginning of the absorption process the transition moments are oriented isotropically and the average

* An experimental check of this formula will be given in a following paper. Small deviations from this ideal linear oscillator model are found. An extended model with the ansatz $\sigma = \sigma_{12} \cos^2 \theta + \sigma_{ex} \sin^2 \theta$ leads to an increase of the $\tau$ values by approximately 0.2 ps and reduces the excited state absorption cross sections by approximately 5% compared to the values listed in table 1.
The intense picosecond pump pulse excites preferentially those molecules with their dipole moments oriented under a small angle to the linearly polarized light field.

(viii) Rotational brownian motion requires times exceeding $10^{-10}$ s depending upon the solvents used in our experiments. As a result the reorientation of the dye molecules has no influence on the bleaching process by rotational redistribution.

Rate equations are used for the interaction of the light pulses with the dye molecules. This classical approach is justified since the dephasing time $T_2$ is expecting to be considerably shorter than the laser pulse duration ($T_2 < 0.5$ ps; $t_p \approx 4$ ps). The following equations describe the bleaching of a strong pump pulse and the transmission behavior of a weak delayed probe beam of parallel polarization.

\[
\frac{\partial I_L}{\partial t} + \frac{n}{c} \frac{\partial I_L}{\partial t} = -I_L \int_0^{\pi/2} \sin \theta \cos^2 \theta \left\{ \sigma_{12} [N_1(\theta) - N_2(\theta)] + \sigma_{\text{ex}} [N - N_1(\theta)] \right\} d\theta ,
\]

\[
\frac{\partial I_p}{\partial t} + \frac{n}{c} \frac{\partial I_p}{\partial t} = -I_p \int_0^{\pi/2} \sin \theta \cos^2 \theta \left\{ \sigma_{12} [N_1(\theta) - N_2(\theta)] + \sigma_{\text{ex}} [N - N_1(\theta)] \right\} d\theta,
\]

\[
\frac{\partial N_1(\theta)}{\partial t} = -\frac{1}{h\nu_L} \sigma_{12} \cos^2 \theta (I_L + I_p) [N_1(\theta) - N_2(\theta)]
- \frac{N_1(\theta) - \int_0^{\pi/2} \sin \rho N_1(\rho) d\rho}{\tau_{\text{or}}},
\]

\[
\frac{\partial N_2(\theta)}{\partial t} = -\frac{1}{h\nu_L} \cos^2 \theta \left( \sigma_{12} (I_L + I_p) [N_1(\theta) - N_2(\theta)] - \sigma_{\text{ex}} (I_L + I_p) [N_2(\theta)] - N_2(\theta) \right) / \tau
- \frac{N_2(\theta) - \int_0^{\pi/2} \sin \rho N_2(\rho) d\rho}{\tau_{\text{or}}}
\]

The initial conditions are

\[
I_L(t, r, z = 0) = I_{0L} \exp \{ - (t/t_0)^2 - (r/r_0)^2 \},
\]

\[
N_1(t - nz/c = -\infty, r, z, \theta) = N_1(\theta),
\]

\[
N_2(t - nz/c = -\infty, r, z, \theta) = 0.
\]

$I_{0L}$ and $I_{0P}$ are the peak intensities of the incident picosecond pump and probe pulse, respectively. Gaussian pulse shapes are assumed. $2t_0$ and $r_0$ are the 1/e values of the pulse duration and the beam radius of both pulses. $\nu_L$ is the frequency of the laser pulses (both pulses have the same frequency), $n$ denotes the refractive index. $N = \int_0^{\pi/2} N(\theta) \sin \theta d\theta$ is the total number density of dye molecules in the solution. $N_1(\theta)$ and $N_2(\theta)$ are the number densities of molecules in the ground state (1) and the Franck–Condon state (2). The rotational redistribution of the molecules is taken into account phenomenologically by the last terms in eqs. (3) and (4). The equations are simplified by the transformation $z' = z$ and $t' = t - nz/c$. The resulting system is solved numerically and the transmitted intensities $I_1(t', r, z' = l)$ and $I_p(t', r, z' = l)$ are calculated. The energy transmissions for the incident pump pulse, $T_L$, and for the probe pulse, $T_p$, are

\[
T_L = \frac{\int_0^{\infty} 2\pi r \int_{-\infty}^{\infty} I_1(t', r, l) dt'dr}{\int_0^{\infty} 2\pi r \int_{-\infty}^{\infty} I_1(t', r, 0) dt'dr}.
\]

The energy transmissions $T_L$ and $T_p$ are determined by the material parameters $N$, $\sigma_{12}$, $\sigma_{\text{ex}}$, $\tau_{\text{or}}$, $\tau$, the pulse parameters $I_{0L}$, $I_{0P}$, $t_0$ and the delay time $t_D$. $N$ is determined by the chosen dye concentration. $\sigma_{12} = 3\sigma_m$ is obtained by absorption measurements with a spectral photometer. $\tau_{\text{or}}$ is known from the literature [11]. For each laser shot we measured the peak intensities $I_{0L}$, $I_{0P}$ and the pulse duration $t_p = 2(\ln 2)^{1/2} t_0$ (fwhm) together with the energy transmission $T_L$ and $T_p$.

To (A). The nonlinear transmission of an intense picosecond laser pulse is calculated from eqs. (1)–(8) with the simplification of $I_{0P} = 0$ (no probe pulse). Two calculated curves for the population lifetimes $\tau = 0.5$ and 2 ps are depicted in fig. 2. The excited state absorption cross section $\sigma_{\text{ex}}$ was chosen in such a way that the curves fit the experimental data at the highest intensity value of $I_{0L} = 6 \times 10^9 \text{ W/cm}^2$. In
table 1, the experimentally determined values of \( \sigma_{\text{ex}} \) are compared with the absorption cross sections \( \sigma_{15} = 3\sigma_{15m} \) which start from the ground state \((\nu = 37800 \text{ cm}^{-1})\). In rhodamine 6G the two cross sections are practically the same, in rhodamine B, the values of \( \sigma_{15} \) appear to be somewhat larger.

The comparison of the experimental points with the calculated curves allows to estimate the time constant \( \tau \) to lie between 0.5 ps and 2 ps. The numbers obtained for various solutions are listed in table 1 together with estimated uncertainties.

To (B). The energy transmission of a probe pulse was calculated as a function of delay time \( t_D \) for different population lifetimes \( \tau \). The five curves in fig. 3 show strongly varying shapes for \( \tau \)-values between 0 ps and 4 ps. The comparison between the calculated curves and the experimental points allows us to determine the relaxation time \( \tau \) with an uncertainty of \( \pm 0.2 \text{ ps} \). This accuracy is considerably better than found in the simple pulse transmission of point A. The obtained times are summarized in table 1 for \( T = 300 \text{ K} \). Values of \( \tau \) between 0.5 ps and 1 ps are found for all investigated solutions of rhodamine 6G and rhodamine B, practically independent of the viscosity of the solvent. We note that even in the solid solution at \( T = 80 \text{ K} \) the short lifetime of \( \tau = 0.9 \pm 0.2 \text{ ps} \) is observed.

The measured \( \tau \)-values determine the lifetime of the molecules in the initially excited states \((2)\). The decay of the molecules from the resonant Franck–Condon states by reorientation of the solvent molecules is ruled out on account of the viscosity data. Reorientation is too slow in highly viscous solutions and in particular in the glassy sample at 80 K. Two physical processes contribute to the observed rapid changes of optical transmission. (1) Energy transfer to neighboring vibrational states with smaller absorption cross sections and (2) vibrational decay to lower lying vibrational levels within the \( S_1 \) state. It appears at the present time that both processes \((1)\) and \((2)\) contribute to our observations.

Our numbers of the vibrational relaxation times differ somewhat from previous reports \([1-7]\). This result is expected since the relaxation values are deduced from different experiments. In this letter the transient population of certain vibrational energy levels is studied while previous investigations determined the delay between excitation and the appearance of gain or measured the delayed appearance of fluorescence spectra.

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References