Thermal lens spectrum of organic dyes using optical parametric oscillator

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Abstract

The wavelength dependence of thermal lens signal from organic dyes are studied using dual beam thermal technique. It is found that the profile of thermal lens spectrum widely differ from the conventional absorption spectra in the case of rhodamine B unlike in the case of crystal violet. This is explained on the basis of varying contribution of nonradiative relaxations from the excited vibronic levels.

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1. Introduction

In the last three decades, we have witnessed the development of a number of techniques for non-destructive characterization of the thermal, optical and structural properties of materials based upon the photothermal techniques such as photoacoustic spectroscopy, photothermal lensing spectroscopy and photothermal deflection spectroscopy. In these methods, the signal magnitude is directly related to temperature rise in a light-illuminated material and to the amount of heat generated via optical absorption and subsequent nonradiative relaxation. Among these photothermal techniques, thermal lensing TL is a versatile and viable technique for exploring nonlinear processes taking place in organic materials and dyes. Eversince the discovery of photothermal lensing effect by Godan et al. [1] this technique of monitoring nonradiative relaxation in excited molecule has been refined by various researchers to suite for the study of various phenomena related to light-matter interactions [2–10]. Most important modification of the thermal lens technique is the dual beam method developed by Long et al. [11] so that we can record the thermal lens spectrum of sample. Detection of thermal lens signal has been proved later on by incorporating optical fibres to introduce flexibility in the experimental configuration [12]. One of the important features of the
technique is that it is a highly sensitive method for
measuring the absorbance of light in a multitude
of samples based on physical changes that occur in
the sample during irradiation.

When a medium is illuminated with Gaussian
elements, some of the incident energy is
absorbed by the molecules in the ground state, and
excited to higher energy states. The excess
energy acquired by the molecule can be dissipated
in many ways. The non-radiative decay produces
a localized temperature rise, creating a refractive
index gradient in the medium so that the medium
acts as a lens-like optical element, called thermal
lens. The focal length of the induced thermal lens
produced by laser irradiation is given by [13,14]:

\[
\frac{1}{f} = \left( 1 + \frac{2nt_e}{\lambda_c} \right)^{-2}
\]

where \( f \) is the time-dependent focal length of the
thermal lens, \( f_0 \) is the focal length just after heating,
\( t_e \) is the characteristic time constant of the
molecule, \( l \) is the sample length, \( D \) is the thermal
diffusivity, \( N \) is the number of molecules, \( \sigma \) is the
absorption cross-section of the molecule, \( h \) is the Planck's
constant, \( n \) is the number of photons, \( H \) is the total output
energy of the laser, \( J \) is the thermal conductivity, \( J \) is
the Joule's constant, \( \omega \) is the beam radius, \( \rho \) is
the density, \( C_p \) is the specific heat and \( P(t) \) is the
power of the heating laser. The probe beam
through the TL will be affected and the TL signal
usually measured as the relative change in the
probe beam center intensity given as:

\[
\Delta I = \frac{1}{2} \left( 1 + \frac{2nt_e}{\lambda_c} \right)^{-2}
\]

\[
\Delta I = \frac{5 \ln(10) \Delta E_0 \Delta n}{\omega^2 \lambda \rho C_p} \frac{dn}{dT}
\]

where \( \Delta E_0 \) is the energy of the pump laser.

In the present experiment, changes in the probe
beam intensity results from the wavelength-dependent
heating of the sample by absorption of the
pump beam. A plot of these intensity changes,
therefore, provides a characteristic signature of the
absorption mechanism of the samples. Most of the
studies reported earlier used tunable dye lasers as
the pump beam [13]. One of the drawbacks of dye
lasers is the limited range of tunability. The
present paper describes the use of TL effect to
record absorption spectrum of organic dyes in
methanol using optical parametric oscillator
(OPO) as the excitation source since optical
parametric devices provide wide and continuous
wavelength coverage, easy and rapid wavelength
tunability, high-energy output and the advantage of
being all solid state.

2. Experimental

The experimental setup of dual beam thermal
lens technique employed in the present investigation
is shown in Fig. 1. In dual beam configuration,
separate lasers are used for pump and probe
beams. This technique is more advantageous since
only a single wavelength (probe) is always detected
and there needed no correction for the spectral
response of the optical elements and detector.
Moreover one can record TL spectra only by
dual beam setup. The pump beam used for the
present study is the radiation from an OPO
(Quantaray mop-700) with tunable output in the
range 460–620 nm. Radiation of wavelength

![Fig. 1. Schematic diagram of the experimental set up. L1, L2—
Lens; DM—dichoric mirror; S—Sample; OF—optic fibre; MPA—
monochromator-PMT assembly; DSO—digital storage oscilloscope.](image-url)
632.8 nm from a low power (1.5 mw) intensity stabilized He–Ne laser source is used as the probe beam. Samples in a quartz cuvette (1 mm) are kept one confocal length past the beam waist. Hu et al. demonstrated that this configuration enhances the thermal lens signal [2]. The probe beam is made to pass collinearly through the sample using a dichroic mirror. An optical fibre mounted on XYZ translator placed at the beam center in the far field serves simultaneously as the finite aperture as well as the detector. The other end of the fibre is coupled to a monochromator-PMT assembly which is set at 632.8 nm. The signal output from PMT is processed using a digital storage oscilloscope (Tektronix, TDS 220). The present work is done at a temperature of 26 °C. The thermal lens spectra are normalized to account for the spectral profile of the OPO output.

The absorption spectrum of the sample is recorded using a UV-Vis–IR spectrophotometer (Jasco V-570). For the fluorescence study, the front surface emission is collected and focused by a lens to the entrance slit of a 1 m Spex monochromator, which is coupled to a PMT having an S20 cathode. The PMT output is fed to a lock-in amplifier. The emission wavelength is scanned in the specified region (500–640 nm). Studies on rhodamine B and crystal violet are reported in the present paper.

An accurately weighed amount of rhodamine B (Exciton) is dissolved in methanol to give a concentration of 1.87 mmol l\(^{-1}\). From this stock solution, sample solutions with different concentrations ranging from 2.68 to 460 μmol l\(^{-1}\) are prepared.

3. Results and discussion

The absorption, TL and fluorescence spectra of rhodamine B in methanol at a concentration of 2.68 μmol l\(^{-1}\) are given in Fig. 2. Absorption spectrum shows the peak absorption at 18315 cm\(^{-1}\) and a shoulder at 19380 cm\(^{-1}\) which reveals two prominent vibronic levels at 546 nm (18315 cm\(^{-1}\)) and 516 nm (19380 cm\(^{-1}\)) with intensity variation as determined by Franck Condon principle. As is clear from the figure TL spectral peak and absorption peak do not coincide. The fact that these peaks do not coincide reveals the unequal magnitudes of nonradiative and radiative transition probabilities from the excited vibronic level. Excitation to 19380 cm\(^{-1}\) level is followed by nonradiative de-excitations to a large number of low-lying vibronic levels. One can conclude that probability of nonradiative de-excitations from 19380 cm\(^{-1}\) is more than that from 18315 cm\(^{-1}\) so as to get the TL spectral peak at 19380 cm\(^{-1}\). This conclusion is also supported by the fact that fluorescence emission (and hence radiative relaxation cross-section) from this level is almost zero as one would expect from the complementary nature of nonradiative and radiative relaxation processes. The fluorescence peak longer wavelength accounts for the Stokes shift [15,16].

Fig. 3 shows the spectra of rhodamine B in methanol at a concentration of 460 μmol l\(^{-1}\). It is evidently observed from the figure, at this concentration there is a relative enhancement of the TL signal as compared with that of fluorescence spectrum. With increasing concentration the possibility of transfer of energy between molecules by collisional means makes the nonradiative part to become prominent. Hence it is obvious that fluorescence quantum yield decreases as the concentration
Spectra of rhodamine B for a concentration of 460 μmol L⁻¹ in methanol (a) absorption spectrum (b) thermal spectrum (c) fluorescence spectrum.

Enhancement in nonradiative processes with increase in concentration will also take place due to reabsorption of fluorescence emission, thereby causing the red shift in the fluorescence peak (Fig. 4). Concentration dependence of TL signal (Fig. 5) from rhodamine B at various pump wavelengths also revealed same results. In solutions of low concentration dye dissolve practically completely into monomers. It should be mentioned that basic dyes like rhodamine B may dissociate at high dilutions into cations and anions. The absorption spectra are determined by the intrinsic absorption of the dye molecules and the dye-solvent interaction. At this concentration dye-dye interaction is negligible because of the large average distance between them. The absorption spectra (Fig. 3) contain contributions from the monomers and the aggregates which makes the spectrum broader at higher concentrations. For highly soluble dye like rhodamine B the dye-dye interaction gains importance at high concentrations since the mean distance between them becomes small [22–24]. On increasing the concentration of dye solution, the aggregate formation and reabsorption of fluorescence emission will result into the enhancement in absorption and nonradiative relaxations at higher wavelengths. Fig. 6 shows the absorption and TL spectrum of crystal violet dye for a concentration of 6.5 μmol L⁻¹. The absorption and the TL spectra have the same profile. This indicates that

The concentration dependence of peak fluorescence of rhodamine B.
unlike in the case of rhodamine B dye there are no intermediate vibronic levels from which nonradiative de-excitation becomes prominent. Our studies show that the simultaneous analysis of TL, absorption and fluorescence spectra will be helpful to understand relative magnitudes of nonradiative and radiative de-excitation probabilities from the excited states.

4. Conclusion

Thermal lens, absorption and fluorescence spectra of rhodamine B and crystal violet were studied. It has been found that, unlike in the case of crystal violet, profiles of TL spectrum widely differ from the absorption spectrum. This is due to different radiative and nonradiative de-excitation from magnitudes of branching ratios of the upper vibronic levels. In other words, we are able to identify an intermediate vibronic level from which nonradiative de-excitation is predominant.

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