ABSTRACT

Transient experiments with picosecond laser pulses give valuable information on the dynamic properties of polyatomic molecules in the electronic ground state. In small molecules the decay of vibrational energy occurs via individual lower energy states; in large molecules the experimental data support a statistical model.

For a number of years we have been concerned with two elementary questions: How long does a normal mode of a polyatomic molecule vibrate after excitation and in which way and how fast is vibrational energy redistributed over the molecule? Our investigations were made in liquids, since this phase is of paramount importance in chemistry and biology. There are numerous reactions in chemistry and biology, where excess energy first goes into vibrational energy states and from there to the thermal bath. What are the relevant time constants for these processes? Another topic of current interest is the prospect of state-selected chemistry where new and efficient reaction channels are predicted after excitation of individual vibrational modes by infrared laser radiation. In this respect, the population life times of vibrational modes are of relevance. The data presented in this note shine new light on the problem raised above.

The main reason for the lack of information on vibrational life times in liquids is the strong interaction of molecules in the liquid phase. The expected time constants are very short, of the order of $10^{-10}$ to $10^{-12}$ seconds. With the advent of ultrashort laser pulses it becomes possible to start quantitative experiments in this very short time domain /1-3/. In our experiments we excite a well defined vibrational mode, e.g. a known CH-stretching mode,
by an ultrashort (~5 ps) infrared pulse of the appropriate frequency and monitor the anti-Stokes Raman spectrum with a second delayed visible pulse, again of a few picoseconds duration. In this way we are able to investigate several vibrational energy states of the same molecule. We may follow the redistribution of vibrational energy from the excited mode to lower lying states. In particular, we study for different vibrational modes (i) the rise and the delayed maximum of the excess vibrational population, (ii) the decay time of the mode, and - most recently - (iii) the absolute change in population.

In a number of previous papers we investigated the population life time of individual modes in small molecules /4-7/. CH-stretching modes were resonantly excited around 3000 cm\(^{-1}\) by tunable IR-pulses. Of major interest are the following observations:

(i) There exists rapid vibrational energy exchange between similar modes of nearby energy states. For instance, energy is rapidly transferred from the primary excited CH-stretching modes to a neighboring Raman active CH-stretching mode during the excitation time of a few picoseconds.

(ii) The decay of vibrational energy proceeds via anharmonic coupling with neighboring overtones and combination modes. As an example we point to the favorable energy transfer in certain molecules from the CH-stretching modes at ~3000 cm\(^{-1}\) to the overtones of the corresponding bending modes at approximately 1500 cm\(^{-1}\).

(iii) The degree of interaction between the fundamentals and the overtones and combination modes may be inferred from an inspection of the types of modes involved and from the investigation of the Fermi resonance between the modes of interest. The quantitative study of the infrared Raman spectra has allowed us to estimate transfer rates for vibrational energy in fair agreement with experimental observations.

As an example we present data on CH\(_2\)CCl\(_2\) and trans CHCl=CHCl, where only two atoms exchange their positions while the structure and the number of atoms are preserved /6/. In Fig. 1a experimental data of CH\(_2\)CCl\(_2\) are depicted. The \(v_1\) mode at 3036 cm\(^{-1}\) is resonantly excited by an infrared pulse and the anti-Stokes signal of the same mode is observed. The Raman signal, i.e. the population of
Figure 1: Anti-Stokes signal versus delay time of the probing pulse. (a) CH<sub>2</sub>CCl<sub>2</sub> in CCl<sub>4</sub> (c = 0.35 m.f.). The decay of the CH<sub>2</sub>-stretching mode at 3036 cm<sup>-1</sup> is shown. (b) trans CHCl=CHCl in CCl<sub>4</sub> (c = 0.35 m.f.). The CH-stretching mode at 3084 cm<sup>-1</sup> is excited and the mode at 3073 cm<sup>-1</sup> is monitored. The broken curves are the cross-correlation functions of the IR exciting and green probing pulse.

The ν<sub>1</sub> mode, decays with a life time of T<sub>1</sub> = 3±1 ps. Quite different is the time dependence of the CH-stretching modes in trans CHCl=CHCl (see Fig. 1b). The molecule is excited via the ν<sub>9</sub> mode at 3084 cm<sup>-1</sup> and the population of the ν<sub>1</sub> mode at 3073 cm<sup>-1</sup> is monitored. Now we find a longer time constant of T<sub>1</sub> = 10±2 ps. In Fig. 2 we show the IR absorption and Raman spectra between 2950 and 3250 cm<sup>-1</sup> of both molecules. We point to the strong Fermi resonances in the IR spectrum of CH<sub>2</sub>CCl<sub>2</sub> with combination modes and overtones. This strong anharmonic coupling is responsible for the short life time of the molecule. Vibrational energy is rapidly transferred to combination modes and overtones and thus to lower energy states. The smaller Fermi resonances of trans CHCl=CHCl give rise to the longer life time discussed above.

(iv) Very small molecules have only a few normal modes and thus a limited number of decay routes. The individual population life times may vary drastically within the same molecule. Acetylene re-
Figure 2: Infrared absorption (a) and Raman (b) spectrum of CH₂CCl₂ between 2950 and 3250 cm⁻¹. Two combination tones are in strong Fermi resonance with the two CH-stretching modes at \( \nu_1 \) and \( \nu_7 \). Infrared absorption (c) and Raman (d) spectrum of trans CHClCHCl. There is less Fermi resonance mixing than in CH₂CCl₂.

presents a particularly interesting example /7/. The CH-stretching modes couple quite effectively to combination modes in the frequency range of 3200 cm⁻¹ making the life time of the CH-stretching modes short, approximately 2 ps. We note that all combination modes contain the C=O-stretching mode at \( \nu_2 = 1968 \) cm⁻¹. What is the life time of this lower vibrational state? Inspection of the energy levels in acetylene suggests that the transfer of energy to neighboring combination modes is very unfavorable on account of symmetry selection rules. One expects a long life time of the \( \nu_2 \) mode. Indeed, we found an exceptionally long life time. In Fig. 3 we show

Figure 3: Time dependence of the population of the C=O mode of acetylene in liquid CC₁₄ following excitation of the \( \nu_3 \) CH-stretching mode at 3265 cm⁻¹. After a rapid rise the population decays slowly with a decay time of 240 ps.
the experimental data. The $v_2$ mode is very rapidly populated after excitation of the infrared active CH-stretching mode and it decays very slowly with a life time of $T_1 = 240$ ps. This life time is the longest found so far for a vibrational mode of a polyatomic molecule at room temperature.

We now turn to the discussion of medium size molecules, where the mode density is quite large around the primarily excited mode. As an example we present data on naphthalene ($C_{10}H_{8}$), where infrared active CH-stretching modes are excited by IR pulses of $\nu = 3055$ cm$^{-1}$ /8/. The transient change of the population of the three strong Raman transitions at $\nu_1 = 3050$ cm$^{-1}$ (CH-stretching mode), $\nu_3 = 1380$ cm$^{-1}$ (ring deformation mode), and $\nu_8 = 765$ cm$^{-1}$ (skeleton mode) is depicted in Fig. 4 a to c. The three figures show quite vividly the build-up and decay of the different vibrational levels. The $\nu_1$ mode is rapidly populated, i.e., energy is quickly transferred from the excited modes, $\nu_1$ decays quite fast with a time constant of 2 ps. The $\nu_5$ mode rises more slowly and decays with a time constant of 9 ps. Model calculations suggest that the $\nu_5$ mode was populated by the decay of the CH-stretching modes. Also the $\nu_8$ mode shows a delayed maximum of excess population and a decay with a similar time constant of 7 ps.

Of special interest for the elucidation of the energy transfer processes are the transient peak occupation numbers of the various monitored modes. For the $\nu_5$ and $\nu_8$ modes we find $n = 0.01$ and $n = 0.1$, respectively. Since the density of vibrational states in the energy range of the CH-stretching modes is quite large, namely 500/cm$^{-1}$, we considered an intramolecular energy relaxation process in terms of a statistical energy randomisation. When $E(300 \text{ K})$ represents the ensemble average of the vibrational energy at room temperature, we may formally evaluate a vibrational temperature $T^*$ for those molecules which have absorbed one quantum of 3050 cm$^{-1}$. From $E(T^*) = E(300 \text{ K}) + hc 3050$ we estimate $T^* = 537$ K for naphthalene. The occupation numbers calculated for $T^*$ are in good agreement with the values found experimentally giving support to the statistical model.
Figure 4: Redistribution of vibrational energy in naphthalene. (a) Anti-Stokes Raman signal of the mode $v_1 (3050 \text{ cm}^{-1})$. Rapid transfer of vibrational energy from the initially pumped IR-active CH-stretching mode to the monitored Raman active CH-stretching mode. The correlation curve between IR and green pulse is shown for comparison (broken line). (b) Anti-Stokes Raman signal of $v_5 (1380 \text{ cm}^{-1})$. This ring deformation mode is populated by the relaxation of the CH-stretching modes. (c) Anti-Stokes Raman signal of $v_8 (765 \text{ cm}^{-1})$.

It appears from our data that the vibrational excess energy is rapidly transferred from the set of CH-stretching modes to the many neighboring combination modes. The coupling between the latter modes tends to randomize the vibrational energy within the very short time of 5 to 10 ps. More recent data of even larger molecules confirm the picture of rapid intramolecular energy randomisation. Our investigations suggest that state-selected chemistry in liquids will be highly restricted on account of the extremely short vibra-
tional life times and the rapid randomisation of vibrational energy in large polyatomic molecules.

REFERENCES

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