TIME-RESOLVED RESONANCE RAMAN SPECTROSCOPY OF EXCITED-STATE PORPHYRINS

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Time-resolved resonance Raman (TR\textsuperscript{3}) and absorbance difference studies of the excited states of Cu(TPP) (TPP = 5,10,15,20-tetraphenylporphyrin) have been carried out with < 10 ps times resolution in THF and pyridine solvents. In THF the distinctive transient Raman bands in the $\nu_2$ and $\nu_4$ regions, previously observed with ns laser pulses, grow in the first 55 ps before decaying in 100's of ps. The \AA\ spectra also show biphasic decay. This behaviour is associated with attack by solvent on the 4-coordinate excited state to form the longer lived species observed in TR\textsuperscript{3} experiments.

In pyridine two component decay is also observed but it is the shorter-lived species which gives the transient Raman bands seen previously with ns laser excitation. This state is different from that seen in THF. At 5 ps delay $\nu_4$ is broader than in the ground state and, more importantly, there is a significant shift in the two pyridine bands at ca. 1000 cm\textsuperscript{-1}. This implies a significant involvement of the pyridine-based orbitals in the excited state.

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Time-resolved resonance Raman studies of the excited states of porphyrins have now become established as an excellent means of characterising the electronic structure of such species [1]. Metalloporph-
phyrins with closed-shell metals have a relatively simple photochemistry with well-defined singlet and triplet excited states but in Cu(II) porphyrins additional $d-d$ and charge-transfer (LMCT and MLCT) excited states [3–5] now lie close in energy to the porphyrin $\pi - \pi^*$ states [2]. The relative energies of all these states depend strongly on the solvent, which can coordinate the Cu centre in either the ground or excited states. Cu(TPP) in THF gives distinctive transient Raman spectra, when excited with ns laser pulses at wavelengths 415–450 nm. These spectra typically show double peaks in the $\nu_2$ and $\nu_4$ regions, one set of which are in positions close to those of the ground state species and a second major pair which are shifted ca. 20 cm$^{-1}$ lower and are due to an excited 5-coordinate Cu(TPP)-THF complex [3–5].

The $\Delta A$ spectra of Cu(TPP) in THF recorded at 5 and 100 ps (Fig. 1, insert) show two components, one of which maximises near the Soret band and one which is farther to the red. The latter band (which decays by 100 ps) is very similar to that of simple triplet Cu(TPP). The former absorption band is associated with the species giving rise to the
distinctive transient Raman bands. TR^3 measurements also show evidence for biphasic decay. Loss of the transient bands occurs on the timescale of hundreds of ps but spectra taken at short (10–55 ps) delays (Fig. 1), show that, although both the transient and residual ground state bands are present at 10 ps, both these features grow in intensity up to 55 ps, after which the transient bands decay. At 10 ps some of the excited Cu(TPP) has not yet undergone solvent attack and so is still in the excited triplet state which can be observed in the ∆A spectra but is Raman silent at this probe wavelength. In the next 45 ps solvent attack proceeds faster than decay of either the triplet state or the 5-coordinate excited exciplex so that the population of the 5-coordinate exciplex rises before ultimately decaying in 100’s of ps.

With pyridine solvent the excited state lifetime is significantly shorter than in THF but again the ∆A spectra at 5 ps show a double feature. However, in this case the rate of decay of the 5-coordinate species is faster than that of solvent attack on the 4-coordinate porphyrin triplet present, so that it is the shorter wavelength feature which decays first in the ∆A spectra. The nature of this short lived 5-coordinate state is clearly different from that observed in THF. The data in Figure 2 show

![Figure 2](image_url)

**FIGURE 2** Time-resolved resonance Raman spectra of Cu(TPP) in pyridine. Two spectral regions are shown. λ_pump = 540 nm, λ_probe = 410 nm. Pump-probe time delay: (a) and (e) 5 ps; (b) and (f) 15 ps; (c) and (g) 25 ps; (d) and (h) 35 ps.
that at the shortest time delay $\nu_4$ is broader than in the ground state (which is consistent with literature data [4, 5]) but the most important observation for structural assignment of the state is that in the spectrum recorded at 5 ps there is a significant shift in the two pyridine bands at ca. 1000 cm$^{-1}$ compared to their positions in the spectra recorded at longer time delays and those of the ground state. This implies a significant involvement of the pyridine-based orbitals in the excited state.

References