ELECTRONIC SPECTRA AND STRUCTURE OF LARGE MOLECULES AND CRYSTALS

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The first detailed study of the electronic spectrum of benzophenone crystals was completed. The triplet state at 24187 cm\(^{-1}\) was characterized as \(m^*\) and shown to be the analogue of the formaldehyde lowest triplet state. We showed that at least two of the spin substates are radiatively excited.

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The difference in linewidths for the singlet and triplet states of benzophenone was used to determine a value of ca. \(10^{11}\) sec\(^{-1}\) for the intersystem crossing rate constant.
FINAL REPORT

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9. Outline of Research Findings:

The first detailed study of the electronic spectrum of benzophenone crystals was completed. The triplet state at 24187 cm⁻¹ was characterized as \( \pi^* \) and shown to be the analogue of the formaldehyde lowest triplet state. We showed that at least two of the spin substates are radiatively active and this work served as the stepping stone for later work on the lowest triplet states of ketones and aldehydes of aromatic molecules. Towards the end of the above grant period we published our studies of spin-orbit coupling and radiationless transitions in such molecules. We showed that the scheme describing the intersystem crossing in benzophenone was qualitatively applicable to other ketones. Spin-polarization experiments on anthrone in n-heptane at 4, 2 and 2K showed that the \( T_z \) spin state was populated directly by the intersystem crossing process: In the case of anthrone we concluded that \( T_z \) was predominantly populated.

Studies of the low temperature spectra of a number of ketones showed perturbations, in the form of line broadening, that could be used to expose higher excited states of many molecules. These studies along with others done in our laboratory led us to see general principles relating line broadening to vibronic interactions between electronic states and hence to radiationless transitions. An early review of these
effects was presented in my Accounts of Chemical Research paper in which it is shown how significant information about molecular structure and non-radiative processes can be gleaned from relatively casual observations of spectral bandwidths.

The difference in linewidths for the singlet and triplet states of benzophenone was used to determine a value of ca. $10^{11} \text{ sec}^{-1}$ for the intersystem crossing rate constant, and additional linebroadening of the vibrational bands was used to determine lower limits for vibrational relaxation times in molecular crystals.

We have shown by direct absorption spectroscopy at low temperature that aromatic ketones have low lying triplet state manifolds that consist of interlocking $n\pi^*$ and $\pi\pi^*$ states that strongly perturb one-another in the overlapping region. This work strongly suggests that the low-energy nearby states of such molecules will become an extremely fruitful area for learning more about the excited states and behavior of excited molecules. We present a mechanism for intersystem crossing in ketones that invoked direct spin-orbit coupling of vibronically modified $n\pi^*$ and $\pi\pi^*$ states through the $T_z$ spin substate.

In other work we developed a microspectrophotometer that was used to obtain extremely precise polarized absorption spectra of derivatives of ferrimyoglobin. We showed that a PR (polarization ratio)
spectrum magnifies spectral effects that are otherwise obscured by wide absorption bands. We confirmed general features of porphyrin spectra including polarization of the B and Q states, the vibronic nature of the $Q_v$ state, and the intensity imbalance in the $Q_o$ state. The PR spectrum of the azide uncovered a new electron-transfer excited state. These spectra are certainly the most accurate crystal absorption spectra yet to be obtained due to the dilute mixed crystal features of the heme-protein system and they exposed many new avenues for the investigation of the electronic structure of very large molecules.