Rotational Isomerism and Dual Luminescence in Dipolar Dialkylamino-Compounds

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Summary The feasibility of a rotational isomerisation model in accounting for dual luminescence of p-dimethylaminobenzonitrile and related compounds is substantiated by PPP-SCF-MO-CI calculations.

The dual fluorescence of p-dimethylaminobenzonitrile (DABN) has aroused much controversy¹ concerning the emitting states, and has been ascribed to a solvent-induced inversion of electronic states, to a stepwise formation of excited complexes with solvent molecules, to protonation

of excited species, to excimers, and to isomerisation of the excited molecule via internal rotation of the NMe2 group.

The latter rotational isomerisation model advocated by Grabowski and his co-workers2 has been discounted on the basis of a CNDO/s-CI calculation by McGlynn.3

We report, qualitatively the results of PPP-SCF-MO-CI calculations on DABN and related compounds. The computer program and parametrisation were conventional. Rotation about the ring C(1)-NMe₂ bond was represented by a variation of the resonance integral, β_{cn} , in the range $0~(90^{\circ}~{
m twist})~{
m to}~-2.50~{
m eV}~({
m coplanar}).$ The two-centre repulsion integral, γ_{CN} , was taken to be 5.4 eV for all rotation angles. For the coplanar isomer of DABN we obtain results in close agreement with those obtained by the CNDO/s method. However, in the twisted isomer the $2^{1}A_{1}$ ($^{1}L_{a}$) state drops markedly in energy when $-\beta$ is less than ca. 1.5 eV, reaching 3.0 eV for $\beta = 0.00$ eV, and its character becomes increasingly dominated by excitation from the amino nitrogen lone-pair.† Whilst CI accentuates the stabilisation of the excited twisted isomer it is not the sole effect. Examination of the component energy terms reveals that the change of the repulsion term $\gamma_{\rm CN}$ $C_{\rm N}(m)^2 C_{\rm LVMO}^2$ with twist about the C(1)-NMe₂ bond dominates all others [where C_N is the amino-nitrogen coefficient in the occupied MO 'm', and CLVMO is the coefficient of the ring carbon C(1) for the lowest vacant MO corresponding to the dominant configuration, LVMO $\leftarrow m$, in the excited-state description]. This repulsion term only changes appreciably when, on twist about the C(1)-NMe₂ bond, the orbital 'm' transforms from a delocalised MO to an isolated 2p orbital on the donor NMe₂.

This analysis can be extended to other systems of the type D-Ar-A where D is an electron-donor group distinguishable by rotation and A is an electron-acceptor group.

We predict that when D is of much higher ionisation potential than NMe₂ (e.g. when it is OMe), rotation about D-Ar does not lower the energy of any electronic state significant in luminescence. For strong π electron acceptor groups, e.g. $A = NO_2$, $CH=CHNO_2$, or 3,5-dichlorotriazinyl, rotation of the donor NMe₂ group does not lead to a lowering in energy of the charge-transfer state for in these cases C_{LVMO} is too small.

It appears that, for D = NMe₂, rotational isomerisation is feasible as an explanation of dual luminescence when A is $C \equiv N$ or C(=O)X (X = H, alkyl, OH, or O-alkyl) in a para but not in a meta position with respect to the donor D. We predict that para-, but not meta-, NN-dimethylaminopyridine could exhibit dual luminescence.

These predictions are confirmed satisfactorily by further experimental data which we are accumulating. Dual luminescence is observed at room temperature for acetonitrile solutions of p-dimethylaminobenzaldehyde (380 and 530 nm), p-dimethylaminobenzoic acid (346 and 480 nm), and the methyl p-dimethylaminobenzoate (343 and 480) whilst a single fluorescence band only is observed for mdimethylaminobenzoic acid (435 nm), (p-diethylaminophenyl)dichloro-1,3,5-triazine (420 nm), and trans-1-(4-NNdimethylaminophenyl)-2-nitroethylene (520 nm).

It could be argued that both PPP and CNDO/s methods are unreliable in this area. Whilst this suggests that the results should be treated with caution, the success in correlation and usefulness as a stimulus for further experimental work justifies attention to the PPP results. The rotational isomerisation model is confirmed as feasible in a restricted area. The role of solvent polarity and other influences must, of course, be included in any detailed explanations of dual luminescence phenomena.

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[†] The π^* orbital is not the in-plane π^* orbital of the nitrile group suggested in ref. 2, but embraces the usual set of 2p AOs. In the 90°-twisted form the excitation is equivalent to $\pi^* \leftarrow n$.

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