



12.50

San San San

A DECEMPENDAL CONTRACTORY

MICROCOPY RESOLUTION TEST CHART NATIONAL BUREAU OF STANDARDS-1963-A

1

REPORT DOCUMENTATION PAGE	READ INSTRUCTIONS BEFORE COMPLETING FOR
	I NO. 3. RECIPIENT'S CATALOG NUMBER
4. TITLE (and Sublitie)	J. TYPE OF REPORT & PENIOD COVER
Electron Donor-acceptor Quenching and Photo-	Technical, 1/1/82-10/31
induced Electron Transfer for Coumarin Dyes	6. PERFORMING ORG. REPORT NUMBE
7. AUTHOR(4)	8. CONTRACT OR GRANT NUMBER(+)
G. Jones, II, S. F. Griffin, C. Choi, and	N00014-79-C-0054
W. R. Bergmark	N00014-79-0-0094
9. PERFORMING ORGANIZATION NAME AND ADDRESS	10. PROGRAM ELEMENT, PROJECT, TA AREA & WORK UNIT NUMBERS
Department of Chemistry Boston University	NR 395-609
BOSTON UNIVERSITY BOSTON, Massachusetts 02215 11. Controlling office name and address	
11. CONTROLLING OFFICE NAME AND ADDRESS Office of Naval Research	12. REPORT DATE October 31, 1983
Eastern/Central Regional Office	13. NUMBER OF PAGES
666 Summer Street, Boston MA 02210	20 18. SECURITY CLASS. (of this report)
	unclassified
	184. DECLASSIFICATION/DOWN GRADIN
16. DISTRIBUTION STATEMENT (of this Report)	
States Government. Approved for public release	
, ,	DTIC
17. DISTRIBUTION STATEMENT (of the abstrast antered in Block 20, if differen	DTIC LECTE DEC 2 8 1983 A
17. DISTRIBUTION STATEMENT (of the obstract entered in Block 20, if differen 18. SUPPLEMENTARY NOTES	DEC 2 8 1983
17. DISTRIBUTION STATEMENT (of the electroni entered in Block 20, 11 differen 16. SUPPLEMENTARY NOTES Prepared for publication in the Journal of Physi	DEC 2 8 1983 A A A A A A A A A A A A A
<ul> <li>17. DISTRIBUTION STATEMENT (of the electroni entered in Block 20, if different</li> <li>18. SUPPLEMENTARY NOTES</li> <li>Prepared for publication in the Journal of Physic</li> <li>19. KEY WORDS (Continue on reverse elds if necessary and identify by block number of the second state of the second s</li></ul>	DEC 2 8 1983 A A A A A A A A A A A A A
<ul> <li>17. DISTRIBUTION STATEMENT (of the electron entered in Block 20, if different</li> <li>18. SUPPLEMENTARY NOTES</li> <li>Prepared for publication in the Journal of Physic</li> <li>19. KEY WORDS (Centime on reverse olds if necessary and identify by block number of the second state of the second sta</li></ul>	DEC 2 8 1983 DEC 2 8 1983 A LECTE DEC 2 8 1983 A ical Chemistry dye fluorescence quenching, is, photoredox processes enched by a variety of organi in general, donors with half- DV vs SCE and acceptors with a SCE are candidates for dif- ce. In flash photolysis experience enchers is demonstrated. Electronic dif-
<ul> <li>17. DISTRIBUTION STATEMENT (of the ebetrous entered in Block 20, 11 different 18. SUPPLEMENTARY NOTES</li> <li>Prepared for publication in the Journal of Physical Prepared for Physical Prepared for Physical Prepared for Physical P</li></ul>	DEC 2 8 1983 DEC 2 8 1983 A LECTE DEC 2 8 1983 A ical Chemistry dye fluorescence quenching, is, photoredox processes enched by a variety of organi in general, donors with half- DV vs SCE and acceptors with a SCE are candidates for dif- ce. In flash photolysis experience enchers is demonstrated. Electronic dif-

# ELECTRON DONOR-ACCEPTOR QUENCHING AND PHOTOINDUCED ELECTRON TRANSFER FOR COUMARIN DYRS

19751

Guilford Jones, II,\* Susan F. Griffin, Chol-yoo Choi,

and William R. Bergmark<sup>1</sup>

Department of Chemistry, Boston University, Boston MA 02215

#### Abstract

The fluorescence of 7-maincocumarins is quenched by a variety of organic electron donors or acceptors in acetomitrile. In general, donors with half-wave oridation potentials less positive than 1.0 V vs SCE and acceptors with reduction potentials less negative than -1.5 V vs SCE are candidates for diffusion limited quenching of commarin singlet states. Profiles of quenching rates are consistent with calculated free energies for electron transfer between excited commarins and donors or acceptors. In flash photolysis experiments electron transfer for several dyes and quenchers (e.g., methyl viologen) is demonstrated. Relatively low yields of net electron transfer are consistently obtained due to inefficient ionic photodissociation via singlet quenching or a low yield of more photometive commarin triplets. Electrochemical properties of the coumarins have been investigated by cyclic voltanmetry with the indications of reversible oxidation and irreversible reduction as important processes.

> DTIC COPY

The 7-eminecoumarins (e.g., 1 - 5) constitute an important class of organic dyes which lase <sup>3</sup> and which in some circumstances may act as photosensitizers.<sup>3,4</sup> These structures are also related to the furocoumarins which have received much attention due to their photobiological properties.<sup>4</sup> In previous papers in this series, photophysical and photochemical properties for coumarin dyes have been reported with particular attention given to the pronounced medium dependences of spectral properties and emission yields associated with the highly polar coumarin excited states (nominally depicted by  $\underline{6}$ ).<sup>6</sup> Other recent investigations have been directed to the mechanism of coumarin photodegradation, <sup>7</sup> the behavior of coumarin dyes in water and in aqueous detergent media, <sup>6</sup> and the effects of medium and additives on photostability and emission yield under lasing conditions.<sup>9</sup> - 11

COURSE ADDRESS STATUS

THE PARTY AND A CANADA TO THE PARTY AND A PARTY AND A

The countrin dyes give rise to strong, broad absorption bands extending into the visible accompanied by solvent dependent red shifts in emission indicative of an emissive state having a large dipole moment (i.e., <u>f</u>). The interaction of such excited species with added quenchers has not been studied in detail although countrin singlet <sup>40, 12</sup> and triplet <sup>13</sup> quenching by oxygen have been reported. Our expectation was that the internal combination of donor and acceptor groups for the aninocountrins would activate the intromolecular charge transfer state toward quenching by both reducing and oxidizing agents. This feature would in turn dictate to a significant degree the bimolecular photochemistry displayed by countrin dyes and also prescribe the types of reagents that might be suitable as

account instanted have and proved to

additives to dye solutions (agents that might insure photostability). We report here the behavior of representative aminocoumarins toward electron transfer agents in acetomitrile, the detection of products resulting from photoinduced electron transfer, and related electrochemistry for the dyes.

**Experimental Section** 

North Andrews Andrews

<u>Materials</u>. Dyes <u>1</u> - <u>5</u> were laser grade materials obtained from Eastman Kodak Co. (commarins 1, 35, 102, 153, and 6, respectively). The dyes were checked for purity by tic (silica, ethyl acetate/hexane) and in most cases used as received. Several of the commercial samples were recrystallized from heptane or methanol/water. The amine quenchers and DMM were distilled prior to use; DMA was dried by addition of lithium aluminum hydride under mitrogen prior to distillation under reduced pressure (20 torr). FUN was recrystallized three times from hexane/chloroform and DCB and NV were recrystallized twice from methanol. Solvents employed were spectroquality scetomitrile used as received and triply distilled water.

Finerescence anenching. Dye emission was recorded on a Perkin-Elmer MPF 44% fluorimeter equipped with a spectrum correction unit and using quartz cells. Solutions of  $10^{-4}$  N dye were excited at the absorption maximum and the fluorescence intensity (at  $\lambda_{max}$ ) recorded as a function of added quencher (samples were air saturated). Except for the cases noted below, no changes were recorded in the emission maximum or in the dye absorption profile as the result of addition of quencher. Intensity changes were plotted vs. quencher concentration using the Stern-Volmer equation,  $Io/I = 1 + k_q \tau$  [Q]. Linear regression analysis (r = > 0.98) provided slopes ( $k_q \tau$  values) and intercepts (typically 1.00 + 0.02).

Triplet anenching. Flash photolysis. Flash photolysis apparatus which consisted of a Xenon flash lamp with ca. 35µs duration (fwhm) (22 cm Pyrez cell) has been described previously.<sup>14</sup> Argon-purged solutions of ca. 10<sup>-6</sup>N dye were employed. Photographs of oscilloscope traces were obtained to record % transmission values which were converted to transient absorbance. For measurement of relative yield of transients, absorbance values were recorded at their maximum at the shortest practical times following lamp discharge (usually 100µs following the flash).

<u>Cyclic voltametry</u>. Current-voltage curves were obtained for 10 mM dye in reagent grade (wet) acetonitrile with 0.1 M tetraethylammonium perchlorate (TEAP) or 0.1 M LiClO<sub>4</sub> supporting electrolyte using a Bioanalytical Systems potentiostat. Other conditions included: working electrode, An or Pt; reference electrode , Ag, AgNO<sub>5</sub> (0.01 M); auxiliary electrode, Pt; operating temperature 22<sup>6</sup>.

#### Results and Discussion

CARACTER MARINE CONCERNS CONCERNS AND A CONCERNS

Connaria dve properties: evolie voltametry. Absorption and fluorescence emission data for dyes 1 - 5 are shown in Table 1 along with fluorescence lifetimes appropriate for air-saturated acetonitrile solutions at room temperature. In Table 2 are included values for oxidation and reduction potentials for the dyes obtained by cyclic voltametry for acetonitrile solutions. The electrochemical oxidations were characterized by relatively clean forward and return waves with 60 - 100 mV peak to peak separation consistent with quasi-reversible behavior.<sup>16</sup> The dyes were somewhat more difficultly reduced (a peak potential could not be measured for <u>3</u>) and reduction return waves were not observed. Due to the irreversibility of reduction of the dyes in acetonitrile, peak potentials which varied somewhat according to scan rate (100 - 500 mV/s) represent rough approximations to the thermodynamic reduction potentials.

12 L'AU

1576572

The substituent influences for dye reduction and oxidation are readily apparent in terms of the lactone moiety and other groups such as CF, providing a retarding influence on oxidation (relative to dialkylanilines for which  $E_{ox} = ca$ . 0.7 V vs SCE<sup>17</sup>). Alkyl substitution at the aniline ring provides for greater case of oxidation whereas the series is less readily reduced due to these electron domating influences (relative to unsubstituted coumarin for which  $E_{red} = -1.4$  V vs SCE<sup>16</sup>).

Sinclet sneaching by electron donors and acceptors. The selection of potential quenchers of dye fluorescence is shown in Table 3 along with the appropriate electrochemical data which reflect their relative strengths as oxidizing or reducing agents in accetonitrile. The results of Stern-Volmer analysis of steady-state emission quenching by the electron donor amines are assembled in Table 4. DMA is sufficiently potent to quench the fluorescent state of three of the dyes (1, 2, and 4) at rates which approach the diffusion controlled limit (or. 2-3 x 10<sup>10</sup> M<sup>-1</sup>s<sup>-1</sup>).<sup>18</sup> The exception involves the least readily reduced dye, 2, for which a diminished rate is observed. Consistent with this indication of dominant

donor-acceptor influence is the trend established for quenching  $\frac{2}{2}$ fluorescence in which rate constants fall in a regular fashion according to amine exidation potential. If a near-diffusion- limited value for kq is supplied for DMA quenching of  $\frac{5}{2}$  a quite reasonable value for the lifetime of emission for this dye (not presently available from photon counting measurements) is obtained (Table 1).

A similar series of quenching data involving various electron acceptors is reproduced in Table 5. Again, one common quencher, FUM, is successful in sequestering coumarin singlets at a high rate. The more potent acceptor, NV, 1,1'-dimethy1-4,4'-bipyridinium dication, requiring for solubility considerations a mixed aqueous medium, is somewhat more successful in quenching 1 fluorescence (reaching the diffusion limited rate). When less effective oxidizing quenchers are inspected for 5, fall-off of the quenching constants is again observed, consistent with a diminished quencher reduction potential.

Quenching data may be analyzed using the Weller equation, 19

 $\Delta G_{et} = E_{ox} - E_{red} - E_{oo} - C$ 

with which the free energy change for an encounter pair undergoing electron transfer is computed using redox potentials (converted to energies), the excitation energy of the excited state participating in quenching ( $E_{00}$ , for commarin singlets, computed from absorption and emission curves), and a

coulombic term, C, usually assumed to value ca. 0.1 eV for acetomitrile solvent.

ANALY STATES ARACHINE STATES

AND A CONTRACT OF A CONTRACT OF

AND THE A

Computed free energies are provided in Tables 4 and 5. The "rule of thumb"<sup>19</sup> which suggests that near diffusion limited quenching behavior is observed in bimolecular quenching via electron transfer when  $\Delta G_{et} \leq -5.0$  kcal/mol is again valid for the coumarin quenching data. The pattern of free energy dependence shown by the data for 2 and 5 reflects a diminution in quenching constant of about one order of magnitude for each reduction in electron transfer exothermicity of 5 - 10 kcal/mol. A similar dependence for organic donors and acceptors in excited state quenching has been found in a variety of other systems.<sup>20,21</sup>

Triplet guesching. Flash photolysis. The coumarins in combination with several quenchers were subjected to flash irradiation using conventional equipment (Xenon flash lamp, 35µs fwhm). Under conditions where quencher concentrations dictated that coumarin singlets were substantially quenched (from Stern-Volmer analyses, vide supra), transients were not generally observed in the 450 - 700 nm range (100 µs to 100 ms regime). Such combinations included 1 and 2 with DNA and FUN. Spectra for coumarin radical-ions have not yet been reported, but transients such as DMA<sup>+</sup> ( $\lambda_{max}^{2}$  500<sub>mm</sub>)<sup>22</sup> are readily observed. The failure of singlet quenching in providing even moderate yields of radical- ions in bulk solution is consistent with a number of recent findings which show that the fate of singlet radical-ion pairs resulting from exothermic electron transfer quenching is predominantly in-cage recombination.<sup>33,34</sup>

The search for electron transfer products from quenching was combined with the detection of triplet-triplet absorption for two of the dyes. For 1 alone in acetomitrile or in 85% acetomitrile/water, a transient with  $\lambda_{max}$ at 600 - 625 mm was observed and assigned to the coumarin triplet which has been detected previously.<sup>7,13,35</sup> The triplet of 1, which showed a first order decay and a lifetime of 120-180 us (several runs) in acetonitrile could be completely quenched by 1.0 mM FUM (where singlet quenching is not important). In this experiment the 600 nm transient is not replaced by another absorbing species in the 400 - 700 region (the FUN radical-anion absorbs at 350 mm,<sup>26</sup> an area obscured by dye bleaching and recovery). DNA (1.0 mM), on the other hand, is not successful in intercepting triplet 1. Both of these results are consistent with electron (or energy) transfer quenching by FUN but not by DNA if the energetics of Tables 4 and 5 are modified by reducing the exothermicity of electron transfer by ca. 10 keal/mol, the estimated difference in singlet and triplet energies for similar dyes as shown by recent spectroscopic measurements. 3,37

1. No.

1.10 miles

Exceptions to the general pattern of negligible reactivity (net electron transfer) were encountered for the acceptor quencher, MV.<sup>34</sup> Flash photolysis of 1 and 5 was examined in some detail under conditions appropriate for quenching by MV of dye singlets or triplets. A transient similar to triplet 1 was observed on flash irradiation in the presence of MV (Table 6). The absorption maximum was again at about 600 nm, consistent with formation of the reduced species  $MV^+$ .<sup>34</sup> (The other well characterized absorption of MV+ at 395 nm is obscured by strong absorption by the dye in that region.) That this transient observed with added MV is clearly a species other than the dye triplet is indicated by the extended decay time

page 8

(millisecond range) which is more consistent with the appearance of a radical-ion. Decay data for the presumed NV<sup>+</sup> intermediate did not uniformly obey first or second order kinetics and decay times (reported as half-lives, Table 6) varied somewhat from run to run and were sensitive to sample preparation (Ar purging).

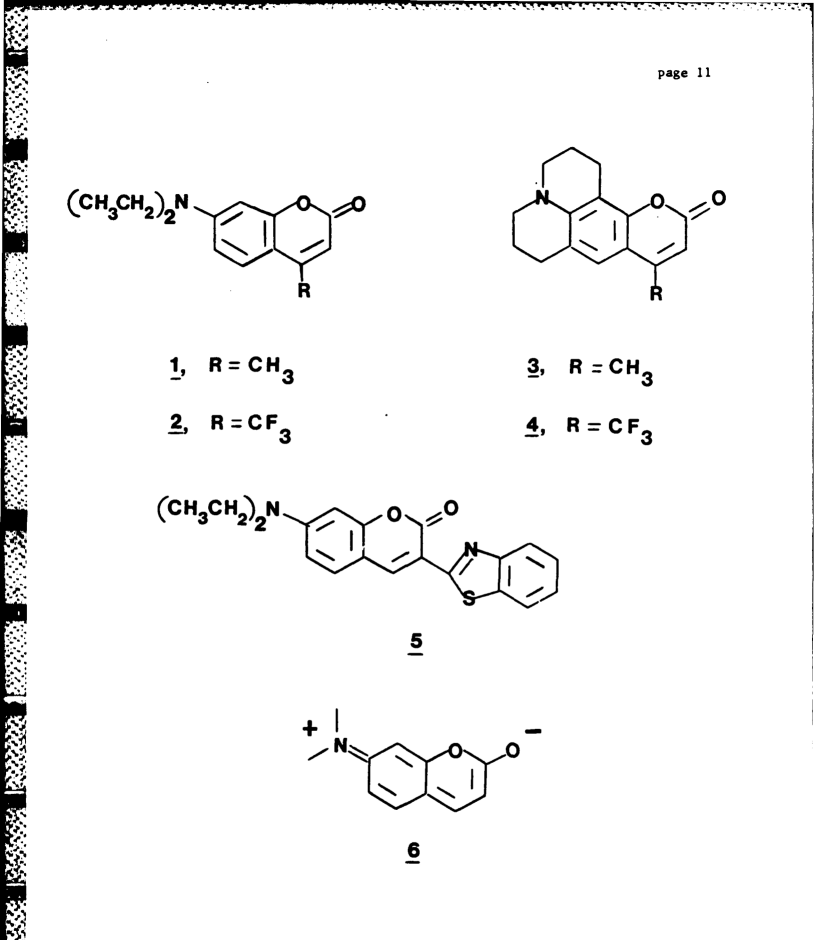
The dependence of yield of electron transfer was inspected for  $\underline{1}$  as a function of MV concentration in acetonitrile/water. A moderate increase in maximum absorbances (Table 6) recorded at about 100 $\mu$ s following the lamp flash ( prior to the onset of significant decay) is noted for increased concentrations of MV. Absolute values for electron transfer yield were not determined but were clearly quite low, as suggested by data for viologen quenching in other well characterized systems.<sup>24,39</sup> The relative yields reported here are consistent with an inefficient photoinduced electron transfer for coumarin singlet quenching (vide supra) and a more robust reaction of the coumarin triplet whose yield in the absence of quencher (via intersystem crossing) is known to be very low (< 1% for  $\underline{1}$  in acetonitrile;<sup>4</sup> (Concentrations of MV = 4.0 and 10.0 mM correspond to 20% and 40% singlet quenching , respectively.)

Electron transfer involving  $\underline{5}$  and methyl viologen appears also to be important (Table 6). Tields of electron transfer for  $\underline{1}$  and  $\underline{5}$  appear similar but the comparison is complicated since triplet yields (or triplet extinction coefficients) for both dyes are not known.<sup>30,31</sup>

In summary, the quenching of the fluorescence of countrin laser dyes by a var'ty of  $e^1$  stron donors and acceptors has been observed. Net electron tr' sfor is not a dominant path for donor-acceptor quenching,

although the photoreduction of methyl viologen on quenching of dye singlets or triplets is confirmed by flash photolysis results. The ranges of oxidation and reduction potential for additives which will diminish the fluorescence of several representative coumarin dyes have been established.

Acknowledgements. This work was supported by the Office of Naval Research. We thank also Drs. W. R. Jackson and S. Kanoktanaporn for technical assistance, and Professors N. Z. Hoffman and M. F. Delaney for the loan of equipment.



#### References

- 1. On subbatical leave from Ithaca College, Ithaca, New York
- Drexhage, K. D. "Dye Lasers," Topics in Applied Physics, vol. 1, Schafer, F. P., ed.; Springer-Verlag, New York, 1977.
- 3. Specht, D. P.; Martic, P. A.; Farid, S. <u>Tetrahedron</u> 1982, <u>38</u>, 1203.
- Jones, G., II; Xuan, P. T.; Schwarz, W. <u>Tetrahedron Lett.</u> 1982, <u>23</u>, 5505.
- Melo, M. T.; Averbeck, D.; Bensasson, R. V., Land, E. J.; Salet, C.
   <u>Photochem. and Photobiol</u>. 1979, <u>30</u>, 645, and ref cited therein.
- 6. a. Jones, G., II; Jackson, W. R.; Halpern, A. M. <u>Chem. Phys. Lett</u>.
  1980, <u>72</u>, 391; b. Jones, G., II; Jackson, W. R.; Kanoktanaporn, S.;
  Halpern, A. M. <u>Optics Commun.</u> 1980, <u>33</u>, 315; c. Jones, G. II; Jackson,
  W. R.; Choi, C.; Bergmark, W. R. , submitted.
- 7. Jones, G., II; Bergmark, W. R. J. Am. Chem. Soc., submitted.
- Jones, G., II; Jackson, W. R.; Kanoktanaporn, S.; Bergmark, W. R. submitted.
- 9. von Trebra, R.J.; Koch, T.H. Appl. Phys. Lett. 1983, 42.
- 10. von Trebra, R. J.; Koch, T. H. Chem. Phy. Lett., 1982, 93, 315.
- 11. Fletcher, A. N.; Knipe, R. H.; Pietrak, M. E. Appl. Phys. 1982, B27,

93, and ref cited therein.

- 12. Kubin, R. F.; Fletcher, A. N., Chem. Phys. Lett., in press.
- 13. Pavlopoulos, T. G. IERE J. Quant. Elect. 1973, QB-9, 510.
- 14. Caspari, G.; Hughes, R. G.; Endicott, J. F.; Hoffman, M. Z. J. Am. Chem. Soc., 1970, 92, 6801.
- 15. Nicholson, R. S.; Shain, I. Anal. Chem. 1964, 36, 706.
- 16. Orlov, Yu. E. Russ. Chen. Rev. 1977, 46, 671.
- Weinberg, N. ed., Techniques of Electroorganic Synthesis," part 2, Wiley Interscience, New York, 1975; b. Zuman, M., "Handbook Series in Organic Electrochemistry," vol 1, CRC Press, Boca Ratam, Florida, 1976.
- Murov, S. L. "Handbook of Photochemistry" Marcel Dekker, Inc. New York, 1973, p 55.
- 19. Rohm, D.; Weller, A. Israel J. Chem. 1970, 259.
- See, for example, Kawenoki, I.; Keita, B.; Kossanyi, J.; Nadjo, L. <u>Nouveau</u> <u>J. Chim. 1982, 6</u>, 387; Takamuku, S.; Kigawa, H.; Susumu, T.; Tsumori, K.; Sakurai, H. <u>Bull. Chem. Soc</u>. <u>Jpn. 1981, 54</u>, 3688; Jones, G., II; Santhanam , M.; Chiang, S.-H. <u>J. Photochem.</u>, 1980, <u>12</u>, 267.
- For discussions of factors governing quenching rates as a function of free energy change, see Sutin, N. <u>Acct. Chem. Res. 1982, 15</u>, 275; and Balzani, V.; Bolletta, F.; Scandola, R. J. <u>Am. Chem. Soc</u>. <u>1980</u>, <u>102</u>, 2152.

22. Holcman, J.; Schested, K. J. Phys. Chem., 1977, 81, 1963.

- 23. a. Masuhara, H.; Mataga, N. <u>Acc. Chem. Res. 1981</u>, <u>14</u>, 312; b. Jones,
  G., II; Chiang, S.-H.; Becker, W. G.; Welch, J. A. <u>J. Phys. Chem.</u>
  <u>1982</u>, <u>86</u>, 2805.
- 24. We cannot exclude the appearence of transients which are short-lived (<100µs), have small extinction coefficients, or have overlapping absorption with dative dye.
- Dempster, D. N.; Morrow, T.; Quinn, M. F. J. Photochem. <u>1973/74</u>, <u>2</u>,
   29.
- 26. Hayashi, H.; Nagakura, S. <u>Chem. Phys. Lett.</u>, <u>1978</u>, <u>53</u>,201.
- 27. The FUN triplet energy has been placed at 59 kcal/mole, a value similar to that expected<sup>3</sup> for <u>1</u>: Weng, P.G.; <u>Can. J. Chem., 1982</u>, <u>60</u>, 339.
- 28. For a review of data concerning electron transfer to MV, see Kalyanasundaram, <u>Coord.</u> <u>Chem. Rev.</u>, <u>1982</u>, 159.
- 29. In a crude comparison (where absorbanced cannot be exactly matched) the well characterized system, ruthenium(II)-trisbipyridyl and NV in  $H_00$ ,  $p_{et} = 0.2-0.3$ ,<sup>38</sup> produces on flash photolysis the viologen radical in a yeild at least tem-fold greater than the system,  $\frac{1}{NV}$  in acetonitrile/water.
- 30. Dempster, et al.,<sup>35</sup> report & = 19,000±2,000M<sup>-1</sup>cm<sup>-1</sup> for triplet <u>1</u> in ethanol, also to be compared with a value of 14,000M<sup>-1</sup>cm<sup>-1</sup> for MV in water.

A weak tranient ( $\lambda_{max}$ -500nm) can be observed on quenching 5 with is DMA in acetonitrile. This absorption, much like that of the DMA radical-cation.<sup>22</sup>

			***************************************
	λ	٨f	τ <sub>f</sub> <sup>b</sup> , ns
1	367	434	2.8
2	396	501	0.60
<u>3</u>	3 80	451	2.8
4	418	521	4.6
5	454	501	(3.1) <sup>e</sup>

#### Table 1 Absorption and emission properties of coumarin dyes in acetomitrile

<sup>8</sup>Absorption  $(\lambda_{g})$  and emission  $(\lambda_{g})$  maxima in m.

<sup>b</sup>Fluorescence lifetimes reported for N<sub>3</sub> or Ar purged solutions (ref. 6) corrected for flu orescence quenching by  $O_3$  in air saturated solutions (ref. 6c)

. . . . . . .

<sup>6</sup>Obtained from dimethylamiline quenching results (see text).

## Table 2 Redox potentials for coumarin dyes from cyclic voltammetry

÷	E <sub>ox</sub> (V vs SCE) <sup>a</sup>	E <sub>red</sub> (V vs SCE) <sup>b</sup>
1	1.09	-2.2
2	1.20	-1.8
1	0,72	<-2.3
£	0.89	-1.8
٤	1.02	-1.5

<sup>a</sup>Oxidation potentials from reversible waves calculated as  $[B_p(ox) +$ 

E\_(red)]/2 (CE,CN)

West Contracts

<sup>b</sup>Reduction (peak) potentials from irreversible waves (CH<sub>1</sub>CN)

.

and the state of the second state of the sta

#### Table 3. Electron donor or acceptor quenchers<sup>2</sup>

ANALY MANYARY ANALY ANALY ANALY ANALY ANALYSIN

Ĵ,

1. <u>6. 7. 7</u>. 1

Electron donor	E <sub>ox</sub> (V vs SCE)	Electron acceptor	E <sub>red</sub> (V vs SCE)
diethylamine (DEA)	2.2	dimethyl maleate (DNM)	-1.6
triethylamine (TEA)	1.1	p-dicyanobenzene (DCB)	-1.7
dimethylamiline (DM	A) 0.73	fumaronitrile (FUN)	-1.3
		methyl viologen (MV)	-0.69

\*Redox potentials = half-wave potentials from polarography or cyclic voltammetry of quenchers in acetomitrile (ref 17).

Table 4.	. Fluorescend	e quenching d	ata for countries and	electro
Dy∙	Quonchor	k <sub>q</sub> τ,∦ <sup>−1</sup>	k <sub>q</sub> (x 10 <sup>-9</sup> M <sup>-1</sup> s- <sup>1</sup> )	۵G
1	DNA	27.	9.7	-4.6
2	DBA	<0.10	<0.17	28.
	TEA	1.5	2.5	4.0
	DKA	11.	18.	-5.7
3	DNA	6.7	2.4	>-1.2
4	DHA	50.	11.	-4.6
٤	DMA	31.	(10.)	-8.3

## Table 4. Fluorescence quenching data for coumarins and electron donors

Dy •	Quescher	لا <sub>و</sub> د , ا <sup>-1</sup>	k <sub>q</sub> (x10 <sup>-9</sup> M <sup>-1</sup> s <sup>-1</sup> )	۵G <sup>4</sup> et
1	fun	29,	8.7	-17
	MAp	81.	29.	-31
2	fun	3.5	5.8	-6.8
3	fun	66,	24.	-23 .
4	FUX	25,	5.4	-11.
5	DCB	<0,2	<0.06	2.2
	Dome	0,6	0.2	0.8
	FUN	31,	9.7	-6.8

### Table 5. Fluorescence quenching data for countries and electron acceptors

<sup>a</sup>Free energy change for electron transfer (kcal/mol) calculated using the Weller equation (see text).

<sup>b</sup>Quenching results for 85% CH<sub>2</sub>CN/H<sub>2</sub>O, assuming  $\tau_f(\underline{1}) = 2.8$  ms, the value for CH<sub>2</sub>CN (Table 1).

	(NV) , =M	A	۲ <sub>1/3</sub> ,85	Assignment
1	-	0.28	0.16	triplet <u>1</u>
	0.02	0.25	1.3	NV radical-ion
	4.0	0.57	4.8	MV radical-ion
	10.0	0.67	1.3	NV radical-ion
٤	-	0.02	0.3	tripløt <u>5</u>
	0.04	0.65	1.4	NV radical-ion

Table 6. Transient absorption on flash photolysis of 1 and 5 with MV<sup>4</sup>

THE REAL PROPERTY OF THE PARTY OF THE PARTY

Provide the state of the second of the second

<sup>a</sup>Absorbances recorded at 600 nm, 100 us after flash (argon-purged 85% v/v CH<sub>2</sub>CN/H<sub>2</sub>O).

