Photodegradation of cyanine and merocyanine dyes

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Abstract

The photodegradation kinetics for some cyanine and merocyanine dyes was studied by UV-visible spectroscopy and the results show that the fading process follows quasi-first-order or zero-order kinetics in acetonitrile. In other experiments, the principal photodegradation products of the dyes were identified with the aid of GC/MS, and it was found that those cyanine dyes holding a positive charge had higher photostability than the corresponding merocyanines. The relationship between photostability and chemical structure was established using PM3 and AM1 MO calculations. Experimental results in this regard suggest that the species associated with the photodegradation of cyanine dyes may be the semioxidized dye free radical cation, \( \text{Dye}^{+} \). The transient absorption spectra of \( \text{Dye}^{+} \) were determined by nanosecond pump and probe spectroscopy. © 2001 Elsevier Science Ltd. All rights reserved.

Keywords: Photodegradation; Cyanine dyes; MO calculations; GC/MS

1. Introduction

Cyanine dyes are among the most important and widely used synthetic dyes as spectral sensitisers in conventional silver halide photography [1]. Also, it has been reported that cyanine dyes offer advantages as absorbers for optical recording disks. For instance, they are inexpensive, easy to synthesize, and exhibit excellent optical properties, giving good recording sensitivity and a favorable signal to noise ratio [2,3]. In DNA sequencing, the introduction of fluorescent labels offers a practical and safe alternative to the traditional radioactive labels. To eliminate the high fluorescence back-

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fluorescence spectroscopy, where either high sensitivity or a high signal rate is crucial [6,7]. Studies pertaining to the effects of substituents in the poly-
methine chain [8,9], end groups [10,11], and coun-
ter ions [12,13] on the photodegradation of
cyanine dyes have been undertaken.

In the present study, flash photolysis, reaction
pathways, and molecular orbital calculations were
used to investigate the dependence of cyanine dye
photodegradation on molecular charge. It was
anticipated that the positive charge on the indolo
heterocyclic moiety would play an important role in
the photostability of cyanine dyes. Merocyanine
dyes, whose indolo heterocyclic group holds no
charge, were investigated in parallel studies. Struc-
tures of the four dyes employed are given in Fig. 1.

2. Experimental

Photolytic reactions were carried out in aceto-
nitrile (HPLC grade, Fluka) at room temperature
but interaction between solvent and dye molecules
in acetonitrile were neglected. The photolysis
experiments were carried out using a photochemical
reaction apparatus (British Applied Photophys.
Limited) equipped with a 200-watt Hg lamp.
Absorption spectra were recorded on a Shimadzu
UV-260 UV-visible spectrophotometer. Dye degra-
dation was assessed by recording absorption spec-
tra at various irradiation times. After 48 h, the
photodegradation products were analysed by GC/
MS using an HP 1800B GCD GC detector. The GC
column was an HP-5 (30 m × 0.25 mm × 0.25 µm) and
the temperature was raised from 40 to 300°C at a
rate of 10°/min. The carrier gas used was high-purity
nitrogen (99.99%). MS detection was performed
under the vacuum of 52 mtorr, with a 167°C
detector temperature and a voltage of 1507 V.

The transient absorption measurements were
performed using a nanosecond time-resolved pump
and probe apparatus with an excitation wavelength
of 308 nm, pulse width of 20 ns, and output of 20
ml/pulse. The excitation laser and detection light
beam passed perpendicularly through the quartz
sample cell. The transmitted light entered a
monochromometer equipped with a R955 photo-
multiplier. The signals were recorded using a 300
MHz HP54510B digital storage oscilloscope and
then processed with a SUN-486 personal computer
[14]. Molecular geometries and electronic struc-
tures of the dyes were calculated using SCF-MO

3. Results and discussion

3.1. Photochemical reaction

All of the four dyes have strong absorptions
in the 360–450 nm range, and the actual data
are listed in Table 1. On the other hand, the

Fig. 1. Molecular structures of the dyes used in this study.
photodegradation products did not have absorptions in the visible region. Consequently, the dependence of dye concentration on irradiation time was determined by measuring visible absorbance values at various irradiation times. This information was used to determine the rate constants associated with photodegradation.

Based on an established mechanism for the oxidative photodegradation of a cyanine dye, the rate of photodegradation can be defined by Eq. (1):

$$-\frac{d[Dye]}{dt} = k_1[Dye][O_2]$$

(1)

Since the concentration of O\textsubscript{2} in the air-saturated solution is constant at \(\sim 3 \times 10^{-4} \text{ M}\) [7], Eq. (2) can be used to define the rate process:

$$\ln[Dye]_0/[Dye] = kt$$

(2)

A plot of \(\ln[Dye]_0/[Dye]\) versus irradiation time \(t\) gives a straight line, the slope of which is the rate constant for oxidative photodegradation. In the present study, we observed a linear relationship between dye concentration and absorbance at \(\lambda_{\text{max}}\). Therefore, the absorbance values were used directly as a function of changing irradiation times \(t\). The experimental results arising from dyes \textbf{D-1} and \textbf{D-2} are shown in Fig. 2. The data show a linear relationship between \(\ln A_0/A\) and irradiation time, which means that the photodegradation of \textbf{D-1} and \textbf{D-2} follows quasi-first order kinetics.

The photodegradation of dyes \textbf{D-3} and \textbf{D-4} was followed at 443 and 430 nm, respectively. In both cases, there was a linear relationship between \(A_t\) and irradiation time \(t\), as shown in Fig. 3, which means that photodegradation followed zero-order kinetics. Since oxygen is a good electron acceptor and abundant in the reaction medium, the reaction of both dyes with dissolved oxygen in the excited state would normally characterize the primary process associated with photodegradation.

$$\text{Dye} + hv \rightarrow \text{Dye}^*$$

(3)

$$\text{Dye}^* + O_2 \rightarrow \text{Dye}^+ + O_2^-$$

(4)

$$O_2^- + \text{Dye} \rightarrow \text{degradation}$$

(5)

If dye excitation [Eq. (3)] is the rate-determining step, then photodegradation follows first order kinetics. This was the case for dyes \textbf{D-1} and \textbf{D-2}. For dyes \textbf{D-3} and \textbf{D-4}, Eqs. (3) and (5) proceeded quickly, and in this case the interaction of excited
dye with oxygen [Eq. (4)] became the rate determining step and photodegradation followed zero order kinetics.

The rate constants for the photodegradation of D-1 and D-3 were smaller than those for D-2 and D-4, as shown in Table 2. This means that the positive charge present in D-1 and D-3 can greatly influence photostability. Specifically, the presence of a positive charge in the conjugated system stabilized these dyes to oxidation.

3.2. GC/MS of dye D-4

The photodegradation products of dye D-4 were characterised in this aspect of our study. It was not possible to isolate the principal products, even in small quantities using conventional techniques. Instead, we used GC/MS to directly examine the product mixtures. Following the irradiation of dye D-4 in acetonitrile at room temperature with a 200-W Hg lamp, eight major decomposition products were detected (Fig. 4). The major fragmentation ions observed in the mass spectrum of the product mixture are listed in Table 3. Based on these results, a degradation pathway was developed (Fig. 5). We believe that dye degradation begins with the rupture of the N–C(Ar) bond of the N=N–Ar moiety. Then the C=N bond of the CH=N=N moiety and the C=C bond between the indole ring and CH=N=N moiety are attacked to produce the remaining products shown in Fig. 5. It should be mentioned that the products detected by GC/MS in these experiments arise from both the photodegradation and thermal decomposition of D-4. While the mechanisms associated with these two processes may be quite different, the bonds most easily broken were clearly revealed by the MS data. This information will be very useful in the design of photostable dyes.

3.3. Pump and probe spectroscopy experiments

In order to define how cyanine dyes undergo photodegradation, the triplet state of dyes should normally be considered as the species that reacts with oxygen [7]. In our studies, nanosecond pump and probe spectroscopy was used to record the transient absorption of the cyanine dyes following excitation at 308 nm. We used dyes D-3 and D-4 in this aspect of the study. The transient absorption spectra recorded on a solution of D-3 at various times after laser excitation are shown in Fig. 6.

<table>
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<tr>
<th>Dye</th>
<th>D-1</th>
<th>D-2</th>
<th>D-3</th>
<th>D-4</th>
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<tr>
<td>$k$</td>
<td>$8.9 \times 10^{-3}$ min$^{-1}$</td>
<td>$4.45 \times 10^{-2}$ min$^{-1}$</td>
<td>$2.7 \times 10^{-3}$ mol$^{-1}$ min$^{-1}$</td>
<td>$1.2 \times 10^{-1}$ mol$^{-1}$ min$^{-1}$</td>
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Fig. 4. Total ion chromatogram from GC/MS analysis of the decomposition products of D-4.
Each spectrum shows a transient absorption centered at 390 nm and recovery of the absorption centered at 460 nm, over a 30-μs period. The curves for the 390 nm transient species and the recovery of the ground state absorption at 460 nm are shown in Fig. 7. The two curves follow quasi-first-order kinetics, the half-lives of which are shown in Table 4. Laser excitation of dye D-3 in O₂-saturated acetonitrile was performed for comparison. The resultant transient absorption and decay curves were very similar to those in Figs. 6 and 7, which indicates that O₂ did not influence the pump and probe spectroscopy of D-3 following excitation at 308 nm. As seen in Table 4, the decay of the transient absorption at 390 nm did not vary significantly in the presence and absence of O₂. Also, the addition of ferrocene (a triplet state quencher, 2×10⁻⁵ M) did not give an obvious

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change, indicating that the formation of triplet states or free radicals do not characterise dye degradation. It is more likely that the transient absorption at 390 results from the semioxidized dye free radical cation (Dye$^{+}$) \[17\] and that this species is the key intermediate in the photodegradation process. These results are also consistent with a photofading mechanism that is not dependent upon oxygen. The lifetime of the semioxidized dye free radical cation for D-3 was about 14–15 $\mu$s, but the corresponding triplet state and free radical species were not observed.

Laser excitation (308 nm) of dye D-4 in acetonitrile led to two absorption bands and bleaching of ground state absorptions centered at 360, 500 and 420 nm, respectively (Fig. 8). The decay curves for the 360 and 500 nm transient absorptions and the recovery of the ground state absorption at 420 nm are depicted in Fig. 9. All three decays follow first-order kinetics, the half-lives of which are shown in Table 5. When laser excitation of D-4 in O$_2$-saturated acetonitrile was performed, the resultant transient absorption spectrum was very similar to that obtained in deaerated solution. These results indicated that the transient absorption at 360 and 500 nm were not due to the triplet state or free radicals of D-4. It is also likely that both transient absorptions are due to the semioxidized dye free radical cation (D-4$^{+}$). The lifetime of D-4$^{+}$ (4–5 $\mu$s) was much short than that of D-3$^{+}$, and the photostability of D-4 was lower than that of D-3, which may be associated with the short lifetime of D-4$^{+}$. Our experimental results were similar to those from a recent study involving the irradiation of related thiacarbocyanine dyes in methylene chloride,

![Fig. 6. Transient absorption spectra of a de-aerated D-3 solution at various times after irradiation.](image)

![Fig. 7. Transient behavior of irradiated D-3 at 390 and 460 nm.](image)

| Table 4 The decay half-life ($\tau$) of D-3 solutions at the transient absorption wavelengths |
|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| In N$_2$-rich solution | In O$_2$-rich solution | With ferrocene |
| $\tau$ (390 nm) | 14.7 $\mu$s | 15.6 $\mu$s | 14.1 $\mu$s |
| $\tau$ (460 nm) | 13.2 $\mu$s | 13.5 $\mu$s | 12.2 $\mu$s |

![Fig. 8. Transient absorption spectra of a de-aerated D-4 solution at various times after irradiation.](image)
in which long-lived reduction in the intensity of the ground state absorptions was observed [18]. De Schryver has attributed this kind of behavior to conformation changes in the ground state of the cyanine dye [19]. It was also suggested that thiacarbocyanine dyes have at least two spectroscopic minima at the excited surface.

3.4. MO calculations

With the aid of the SCF-MO methods PM3 and AM1, fully optimized molecular geometries for the cyanine dyes examined in this study were calculated. The PM3 optimized geometries for all four dyes are shown in Fig. 10. Selected bond distances, bond angles, dihedral angles, and partial charges associated with D-1 and D-2 are given in Tables 6 and Table 7. The frontier MO energy levels ($E_{HO}$ and $E_{LU}$) and the constituent of HOMO and LUMO (i.e. the coefficients of atomic orbital) are shown in Tables 8 and 9, respectively.

It is evident from the data in Table 6 that there were significant differences in the molecular structures of cationic dye D-1 and neutral dye D-2. For dye D-1, N(15), C(8), C(9), C(10) and N(11) form an extended conjugated system and angle C(5)–N(15)–C(8) (110.2°) departs far from 120°, as would be expected. N(15) has approximately sp² hybridization. As for dye D-2, the C(9)–C(10) bond length (1.459 Å) is larger than that of C(8)–C(9) (1.344 Å), and the N(11)–C(12) bond length (1.433 Å) is larger than that of N(11)–C(10) (1.294 Å). This indicates that D-2 would have inferior conjugation.

The results in Table 7 indicate that the positive charge on dye D-1 resides primarily on the N(11) and N(15) atoms and that C(12) and C(5) have

![Graph of transient behavior of irradiated D-4 at 360, 420 and 500 nm.](image)

![Table 5](image)

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<tr>
<th>Parameter</th>
<th>D-1</th>
<th>D-2</th>
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<tr>
<td>τ (360 nm)</td>
<td>5.92 µs</td>
<td>5.78 µs</td>
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<td>τ (420 nm)</td>
<td>5.04 µs</td>
<td>5.82 µs</td>
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<tr>
<td>τ (500 nm)</td>
<td>4.22 µs</td>
<td>4.04 µs</td>
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</table>

![Table 6](image)

| Bond lengths (Å) and bond angles (°) for PM3 optimized D-1 and D-2 structures |
|-----------------|-----------|-----------|-----------|-----------|-----------|
| Parameter       | D-1       | D-2       | Parameter       | D-1       | D-2       |
| 4–5             | 1.387     | 1.389     | 9–10            | 1.394     | 1.459     |
| 5–6             | 1.405     | 1.406     | 10–11           | 1.347     | 1.294     |
| 5–15            | 1.451     | 1.449     | 11–12           | 1.443     | 1.433     |
| 7–8             | 1.542     | 1.532     | 12–13           | 1.405     | 1.402     |
| 8–9             | 1.394     | 1.344     | 12–14           | 1.399     | 1.397     |
| 8–15            | 1.372     | 1.454     | 15–16           | 1.464     | 1.477     |
| 8–9–10          | 126.4     | 125.1     | 5–15–8          | 110.2     | 106.8     |
| 9–10–11         | 121.8     | 119.0     | 5–15–16         | 121.6     | 116.2     |
| 10–11–12        | 123.4     | 121.7     | 8–15–16         | 128.1     | 119.5     |
| 1–6–5–15        | 178.6     | 177.1     | 8–9–10–11       | 177.6     | 154.2     |
| 5–6–7–8         | 0.838     | 0.895     | 9–10–11–12      | 175.6     | 180.0     |
| 7–8–15–5        | 0.316     | 2.883     | Ph–Ph           | 26.76     | 22.47     |
more electron rich character. Therefore, it is not surprising that the positive and negative charges are delocalised across the N(15), C(8), C(9), C(10), N(11) and C(12) conjugation system.

We used frontier MO calculations to examine the interactions between the HOMO of superoxide radical anion (O$_2^-$) and the LUMO of D-1/D-2 and D-3/D-4 combinations.

Using PM3, we found that $E_{HO} = -0.817$ eV for O$_2^-$, and $E_{LU} = -4.82$ eV and $-0.227$ eV for D-1 and D-2, respectively, and that $E_{LU} = -4.931$ and $-0.612$ eV for D-3 and D-4. The symmetry of 2p-AO in the HOMO of O(1)–O(2) matched that of 2p-AO in the LUMO (C(10)–N(11) or N(10)–N(11)) of D-1 (or D-3), and the gap (4.065 or 4.114 eV) between their energy levels was larger. However, the gap between the $E_{LU}$ of D-2 (or D-4) and $E_{HO}$ of O$_2^-$ was 0.590 (or 0.205) eV and smaller than that for D-1 (or D-3). It was also determined that the symmetry of the LUMO for C(8)–C(9) and C(10)–N(11) or N(10)–N(11) matched that of the HOMO of O$_2^-$ and the number of active positions on D-2 (or D-4) was greater than the number on D-1 (or D-3). This suggests that electron transfer from the HOMO of O$_2^-$ to the LUMO of D-2 (or D-4) is more facile than that to the LUMO of D-1 (or D-3). This could explain why the photostability of D-2 (or D-4) is lower than that of D-1 (or D-3). Similarly results were

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<th>Atom</th>
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<td>N(15)</td>
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Fig. 10. PM3 optimized geometries of D-1 – D-4.
obtained from AM1 calculations and were consistent with experimental data.

4. Conclusion

It has been found that the photostability of cyanine and merocyanine dyes that hold a positive charge in their system was higher than the corresponding neutral forms. These results were determined experimentally and from MO calculations utilising PM3 and AM1 methods. The photodegradation of these dyes took place after excitation at 308 nm and it was shown by nanosecond pump and probe spectroscopy that the triplet states or free radicals of the dyes were not involved in this process. Instead, it is believed that photodegradation occurs via formation of the semioxidized dye free radical cation.

Acknowledgements

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References