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Intermediate diradical character and thermal *cistrans* isomerization of near-infrared absorbing thionated squaraine dyes[†][‡]

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We synthesized squaraine dyes incorporating 4-membered thionated oxocarbon and chalcogenopyrylium moieties, which displayed intermediate diradical character as confirmed by bond-length analysis and temperature-dependent behaviors observed in ¹H NMR and ESR spectroscopy. The ¹H NMR signals of the cyclobutenedithione dyes disappeared at lower temperatures compared to the cyclobutenedione analogues, indicating a lower thermal transition to the excited triplet state and thus greater diradical character. Moreover, we identified *cis*-*trans* isomerization in the thionated dyes. Notably, an increase in diradical character lowered the activation energy of this isomerization, facilitating the structural transition between *cis* and *trans* isomers. These dyes exhibit strong near-infrared absorption, with slight temperature-dependent shifts likely arising from the closely matched electronic transition energies of the isomers.

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Introduction

Singlet diradicaloids are molecules characterized by intermediate diradical character, which is illustrated by resonance structures that include both closed-shell and open-shell forms.¹⁻³ In these molecules, two electrons with antiparallel spins are only partially coupled in the ground state. This partial coupling causes the electrons to occupy separate spatial regions to minimize electron repulsion, thereby reducing the covalent character of the π bond and stabilizing the diradical resonance forms. As a result, singlet diradicaloids can facilitate thermal excitation from singlet to triplet states.⁴⁻⁷ Over the past two decades, various stable singlet diradicaloids have been demonstrated, including π -extended quinodimethanes like hydrocarbon,^{8–10} Chichibabin's quinoidal oligothiophenes,^{11–13} and graphene fragments.^{14,15} These compounds have attracted significant interest due to their unique physical properties stemming from diradical forms, which offer substantial potential for applications in nonlinear optics, singlet fission, and semiconductor materials.¹⁶⁻¹⁹

^bDepartment of Chemistry, Graduate School of Science, Osaka Metropolitan University, Sumiyoshi-ku, Osaka 558-8585, Japan. E-mail: hfuji@omu.ac.jp †Dedicated to Prof. Dr Frank Würthner on the occasion of his 60th birthday. ‡Electronic supplementary information (ESI) available. CCDC 2194625 and 2194626. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d4q001722f The diradical character not only influences the physical properties of these molecules but also affects their structural conversions. For instance, some singlet diradicaloids induced structural transformations through diradical configurations (Fig. 1A).²⁰ Kozaki, Okada, *et al.* reported the thermal *cis-trans* isomerization of a dithienoquinoid analogue of Chichibabin's hydrocarbon (Fig. 1B).²¹ More recently, Wu *et al.* reported that oxindolyl-based Chichibabin's hydrocarbons induced *cis-trans* isomerization due to the contribution of the diradical forms



Fig. 1 An example of a structural change in a diradical form (A). Thermal *cis*-*trans* isomerization of Chichibabin's hydrocarbon analogues (B and C).

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(Fig. 1C).²² These cases collectively highlight the pivotal role of intermediate diradical character in driving dynamic structural changes.

Until recently, the range of compounds classified as singlet diradicals was relatively narrow. However, it is now clear that a variety of compounds with extended π -electron systems exhibit sustained singlet diradical character. In fact, even molecules traditionally considered as closed-shell, such as certain diketopyrrolopyrrole derivatives and donor–acceptor conjugated polymers with quinoid repeat units, can exhibit intermediate diradical character.^{23–28} We also demonstrated that croconaine dyes, typically classified as closed-shell species, actually exhibited intermediate diradical character.²⁹ These dyes consisting of chalcogenopyrylium and five-membered oxocarbons (see Fig. 2A), displayed notable temperature-dependent magnetic properties due to their thermally accessible triplet states.

In addition to the croconaine dyes, chalcogenopyryliumbased squaraine dyes, that were four-membered oxocarbon analogues for the variable applications,^{30–33} also exhibited intermediate diradical character.³⁴ Although we previously confirmed that the chalcogenopyrylium moiety influences the contribution of the diradical structure, the impact of the central oxyallyl structure on the diradical character remains unclear. Thioxyallyl derivatives, where sulphur atom replaces the oxygen atom in the oxyallyl structure, are known to exhibit diradical character, unlike oxyallyl derivatives.³⁵ Therefore, investigating the effect of sulphur-substitutions in the central skeleton on diradical character of squaraine dyes is of particular interest.

In this study, we synthesized novel chalcogenopyryliumbased squaraine dyes featuring a cyclobutenedithione central skeleton (**SQ2a**: X = O, **SQ2b**: X = S) to evaluate their intermediate diradical character (Fig. 2B). These dyes exist as *cis-trans* isomers, where their intermediate diradical character is likely to influence the isomerization behavior. To investigate this further, we examined the *cis-trans* isomerization of the obtained dyes and compared their behaviour with existing squaraine dyes containing cyclobutenediones (**SQ1a-b**).

Results and discussion

SQ2a and **SQ2b** were synthesized by thionation of precursor dye **SQ1a** and **SQ1b** with cyclobutenedione, respectively, using Lawesson's reagent in moderate yields (77% for **SQ2a** and 74% for **SQ2b**, see ESI[‡]).³⁶ They are stable in the solid state even at high temperature and in solution under dark conditions (Fig. S1[‡]), allowing for reliable measurements of ¹H NMR, ESR, and electronic absorption spectra.

Fig. 3 displays the X-ray structures of SQ2a and SQ2b, with detailed information available in the ESI.[‡] Both dyes were analysed as planar *trans* isomers. In the crystal structure of SQ2a, two crystallographically independent molecules are present. We investigated the degree of diradical character by examining four major resonance forms (α to δ) chosen from 16 possible resonance forms (Fig. 3C, and Fig. S3[‡]). To evaluate their contribution, we analysed the difference in average bond lengths along the thio-oxyallyl part, denoted as $\Delta D = \operatorname{ave}(g, p) - \operatorname{ave}(h, p)$ q), where ave(a, b) represents the average of bond lengths a and b. For closed-shell mesoionic structures (α and β), $\Delta D = 0$ due to identical bond lengths in the double and single bond pairs (g, p) and (h, q). In contrast, for the diradical δ form, ΔD > 0 because the outer parts (g and p) are single bonds while the inner parts (*h* and *q*) are double bonds. The γ form shows the reverse trend with $\Delta D < 0$. Importantly, the ΔD values from the X-ray structures of **SQ2a** and **SQ2b** are positive ($\Delta D > 0$), suggesting that the large contribution of diradical δ form, with unpaired electrons on the chalcogenopyrylium components (Table 1).



Fig. 2 (A) A resonance structure contributed by a diradical form of croconaine dyes consisting of a 5-membered oxocarbon moiety and chalcogenopyryliums. (B) Chalcogenopyrylium-based squaraine dyes with cyclobutenedithione moiety (SQ2a–b) synthesized from corresponding cyclobutenedione analouges (SQ1a–b). (C) Thermal *cis–trans* isomerization of the series of squaraine dyes.



Fig. 3 Single crystal X-ray structures of **SQ2a** (A) and **SQ2b** (B) at 100 K. Bond lengths of the methine part (g and p) are shown in the Table 1. Resonance contributors of mesoinonic structures (α and β), thio-oxyallyl diradical (γ), and chalcogenopyranyl radical (δ) (C). ΔD is defined as ΔD = ave(g, p) – ave(h, q).

Table 1 Bond length difference ΔD for singlet and triplet states of present squaraine dyes

$\begin{array}{c} {}^{\prime Bu}_{Pu} & \stackrel{\gamma}{\underset{Bu}{}^{\prime }} \stackrel{q}{\underset{Bu}{}^{\prime }} \stackrel{p}{\underset{C}{}^{\prime }} \stackrel{m}{\underset{Bu}{}^{\prime }} \stackrel{n}{\underset{C}{}^{\prime }} \stackrel{g}{\underset{H}{}^{\prime }} \stackrel{m}{\underset{K}{}^{\prime }} \stackrel{n}{\underset{K}{}^{\prime }} \stackrel{Bu}{\underset{Bu}{}^{\prime }} $						
$\Delta D = \operatorname{ave}(g, p) - \operatorname{ave}(h, q)$						
Dye	Singlet ^a	XRD	Triplet ^a			
SQ2a $(X = O, Y = S)$ SQ2b $(X = Y = S)$	$\begin{array}{c} 0.004\\ 0.011\end{array}$	0.002/0.007 0.024	0.030 0.039			

^a Obtained from optimized structures at the UB3PW91/cc-pVDZ level.

The singlet diradical nature of **SQ2a-b** was also confirmed by DFT calculations (see ESI, Table S3[‡]).³⁷⁻⁴⁰ The ΔD value obtained from the crystal structure was intermediate between the values for the singlet and triplet states from DFT-optimized structures at the UB3PW91/cc-pVDZ level (Table 1). This suggests that **SQ2a-b** is in a state that lies between a pure singlet and triplet state.

The diradical index y_0 , where $y_0 = 0$ and $y_0 = 1$ correspond to a perfect closed-shell and an open-shell electronic structure, respectively, is commonly used to express the degree of contribution of the diradical form in singlet diradicaloids (see ESI[‡]).⁴¹ We estimated y_0 values for *trans* isomers of **SQ2a** and **SQ2b** using the spin-projected unrestricted Hartree–Fock (PUHF) theory using the 6-31G(d,p) basis set, obtaining values of 0.24 and 0.37, respectively (Table 2). These results suggest that the thiopyrylium skeleton enhances the contribution of the diradical form.

In addition to the bond length analyses, ESR spectra of these dyes displayed a signature of singlet diradicaloids. While the signal intensity of SQ2a was very low, both SQ2a and SQ2b exhibited signals with g values of 2.006, indicating the presence of carbon-centred radicals (Fig. 4). While SQ2a displayed negligible temperature-dependent behaviour, SQ2b showed increased signal intensity with rising temperature, indicating a thermal transition from singlet to triplet states that reflects its singlet diradical nature. This observation suggests that the thiopyrylium skeleton likely enhances the diradical contribution.



Fig. 4 ESR spectra of SQ2a (A) and SQ2b (B) in the solid state.

Unlike metal-free organic dyes with closed-shell structures, **SQ2a–b** exhibit characteristic temperature-dependent behavior in their ¹H NMR spectra in DMSO- d_6 due to their diradical character (Fig. 5A for **SQ2a**, Fig. 5B for **SQ2b**). At 303 K, **SQ2a** showed signals attributed to the proton on the pyrylium component. As the temperature increased, the signal broadened and completely disappeared at 333 K (Table 2). In **SQ2b**, no signal corresponding to the proton on the thiopyrylium components was observed, even at 303 K, where DMSO- d_6 barely freezes. These results indicate an increase in thermally excited triplet species with increasing temperature, confirming the intermediate diradical nature of **SQ2a–b**. Thus, ¹H NMR studies, along with ESR data, clearly show that the intermediate diradical character of **SQ2b** is higher than that of **SQ2a**.

We previously reported that **SQ1a-b**, with cyclobutenedione skeletons (precursors to **SQ2a-b**), also exhibits intermediate diradical character. For **SQ1a-b**, the proton signals disappear

Table 2	Indices associated with	diradical character a	nd cis-trans isome	erization for SQ2a,	SQ2b, SQ1a, and SQ1b
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Dye	Signal-disappearance temperature in ¹ H NMR ^{<i>a</i>}	Vibrational frequency $(cm^{-1})^{b}$	$\lambda_{\max} \left(\operatorname{nm} \right) \left(\log \varepsilon \right)^{c}$	$\Delta G^{\ddagger d}$ (kcal mol ⁻¹)	y_0^{e}
SQ2a	333	1480	754 (5.22)	13.7	0.24
SQ2b	>303	1470	849 (5.28)	13.4	0.37
SQ1a	<423	1487	721 (5.37)		0.37
SQ1b	403	1478	822 (5.44)	14.5	0.47

^{*a*} The temperatures at which the signals assigned to the protons of the chalcogenopyrylium skeleton disappear in DMSO- $d_{6,}^{b}$ Frequencies of the stretching bands around methine groups in CHCl₃. Details are shown in ESI.[‡] ^{*c*} Electronic absorption maxima in DMSO. The values in parentheses indicate logarithm of molar absorptivity. ^{*d*} The Gibbs free energies of activation (ΔG^{\ddagger}) for the *cis–trans* isomerization calculated by dynamic NMR method. Details are shown in ESI.[‡] ^{*e*} The diradical index calculated by the spin-projected unrestricted Hartree–Fock (PUHF) theory using the 6-31G(d,p) basis set. Details are shown in ESI.[‡]



Fig. 5 (A) VT-¹H NMR spectra of **SQ2a** in DMSO-*d*₆ (333–303 K) and CDCl₃ (278–213 K). (B) VT-¹H NMR spectra of **SQ2b** in DMSO-*d*₆ (303 K) and CDCl₃ (293–213 K). (C) Schematic diagram of the thermal *cis*–*trans* isomerization and the Gibbs free energies of activation (ΔG^{\ddagger}) for the isomerization.

above 400 K, whereas for **SQ2a–b**, the proton signals vanish at a lower temperature, around 300–330 K (Table 2). This is because **SQ2a–b** reach their thermally excited triplet state at a lower temperature, indicating a smaller energy gap between the singlet and triplet states and a significant contribution from the diradical forms. Thus, thionation of the central skeleton enhances the contribution of the diradical structure, even though the diradical index y_0 values for **SQ1a–b** are estimated to be larger than those for **SQ2a–b** due to limitations in computational accuracy.

The single crystal of **SQ1b** contains both *cis* and *trans* isomers, confirming that *cis–trans* isomerization occurs in solution.⁴² Similar to **SQ1b**, **SQ2a–b** also exhibited *cis* and *trans* isomers at lower temperatures in CDCl₃ (Fig. 5A for **SQ2a**, Fig. 5B for **SQ2b** and Fig. S9–10‡ for **SQ1a–b**). Signals corresponding to the protons on the chalcogenopyrylium component coalesced at 278 K for **SQ2a** and at 273 K for **SQ2b**, suggesting that the isomerization is promoted at higher temperature condition and the signals for isomers is no longer observed in the NMR timescale. The Gibbs free energies of activation (ΔG^{\ddagger}) for the *cis–trans* isomerization of **SQ2a** and **SQ2b** were estimated using the dynamic NMR method, result-

ing in values of 13.7 and 13.4 kcal mol⁻¹, respectively (Fig. 5C, and Table 2).⁴³ Although some error was introduced by the inability to observe peak shifts at temperatures below 213 K due to the freezing of CDCl₃ used in the measurements, **SQ2b**, with a cyclobutenedithione skeleton, exhibits a lower ΔG^{\ddagger} than **SQ1b**, which has a cyclobutenedione skeleton ($\Delta G^{\ddagger} = 14.5$ kcal mol⁻¹).⁴⁴ The higher diradical contribution of **SQ2b** compared to **SQ1b** may lower the activation barrier and promote isomerization.

To measure the strength of the C–C double bond involved in isomerization, FT-IR spectra of these dyes in CHCl₃ were recorded. The IR absorption corresponding to the stretching vibration around the methine group in **SQ2b** was observed at a lower wavenumber compared to **SQ2a**, indicating that the bond in the methine group of **SQ2b** is relatively weaker (Table 2, Fig. S11 and 12‡). Thus, increased diradical character appears to diminish the double bond character of the methine group in the ground state, thereby promoting *cis–trans* isomerization.

The electronic absorption spectra for the dves SO2a and SQ2b, compared with those of SQ1a and SQ1b, are illustrated in Fig. 6 and summarized in Table 2. SO2a and SO2b each exhibit strong and sharp absorption peaks at 754 nm and 849 nm, respectively. The dyes with the cyclobutenedithione skeleton (SO2a, SO2b) exhibited absorption bands in a longer wavelength region compared to those with corresponding cyclobutenedione analogues (SQ1a, SQ1b). Notably, SQ2b, which has a high contribution from the diradical structure, demonstrated a lower transition energy. This is consistent with our previous results that a greater diradical character results in reduced transition energy according to the perturbation theory proposed by Fabian et al.^{34,45} We also examined the temperature-dependence in absorption spectra of these dyes (Fig. 6B, and Fig. S15[‡]). With increasing temperature, the absorption bands of SQ2a and SQ2b were red-shifted by 2 nm and 5 nm, respectively. This change in absorption band with temperature might be attributed to cis-trans isomerization. DFT calculations estimate that the transition energy of the cis isomer is slightly lower than that of the *trans* isomer, suggesting that the equilibrium shifts towards the trans isomer at low temperatures due to cis-trans isomerization (Table S3 and S4[‡]).



Fig. 6 (A) The electronic absorption spectra of SQ2a (green), SQ2b (purple), SQ1a (red) and SQ1b (blue) in DMSO (5 μ M). (B) Normalized electronic absorption spectra of SQ2a and SQ2b in the temperature range from 373 to 303 K.

Conclusions

In summary, we synthesized cyclobutenedithione-based dyes SQ2a-b and analysed their intermediate diradical character using ¹H NMR, ESR, and theoretical calculations. Our results reveal that dyes incorporating a thiopyrylium component exhibit large degree of diradical character compared to those with a pyrylium component. Furthermore, SQ2a-b demonstrate a higher degree of diradical character than their cyclobutenedione counterparts (SQ1a-b). This enhanced diradical character promotes cis-trans isomerization, which is attributed to the weakening of the C-C double bonds involved. Additionally, SQ2a-b show subtle temperature-dependent changes in their electronic absorption spectra within the nearinfrared region. Our study exemplifies the impact of diradical character on the physical and chemical properties of squaraine dyes, offering valuable insights for the design of advanced near-infrared absorbing dyes.

Data availability

The data supporting this article have been included as part of the ESI.[‡] Crystallographic data for **SQ2a**, **SQ2b**, **SQ1a**, and **SQ1b** are available in the CCDC under deposit numbers 2382744, 2382745, 2194625 and 2194626,[‡] respectively.

Conflicts of interest

There are no conflicts to declare.

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