

Selective Carbon-Bromine Bond Fission Induced by Non-Resonant Two-Photon (NRTP) Excitation of *trans*-4-Bromostilbene

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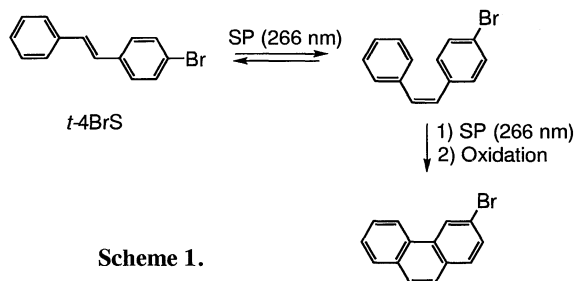
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Non-resonant two-photon (NRTP) excitation of a 1,4-dioxane solution of *trans*-4-bromostilbene induces selective homolytic carbon-bromine bond fission, in contrast to the corresponding single-photon (SP) excitation, where the *cis-trans* isomerization is predominant.

We have recently reported that non-resonant two-photon (NRTP) excitation induces dramatic change of the photochemical reaction pathways of several organic compounds in solution from the conventional single-photon (SP) excitation.¹ Typically, the NRTP excitation of hexane solutions of α,ω -diphenylpolyenes such as stilbenes and *trans*-1,4-diphenylbutadiene^{1b,c} induces selectively *cis-trans* isomerization, while the SP excitation yields several byproducts in addition to the isomerization.^{2,3} We have applied the NRTP method to *trans*-4-bromostilbene (*t*-4BrS) and found a highly selective carbon-bromine bond fission in 1,4-dioxane, in contrast to the SP excitation, which induces the *trans-cis* isomerization predominantly, while the prolonged SP irradiation yields 3-bromophenanthrene (BrPhen) via the intramolecular cyclization.^{3d} This finding constitutes the first example of the selective photoreaction by the NRTP excitation except for the *cis-trans* isomerization of α,ω -diphenylpolyenes, suggesting the biradical nature of the two-photon allowed excited state.

When a 1,4-dioxane solution of *t*-4BrS (3.5×10^{-3} M), which was purified by recrystallization from ethanol before use, was irradiated at room temperature with 266 nm pulsed light (the fourth harmonic of a Nd:YAG laser, 1.2 MW/cm², pulse width ~ 4 ns, 10 Hz) for 45 min, *cis*-4-bromostilbene (*c*-4BrS), 3-bromophenanthrene (BrPhen), and *t*-stilbene (*t*-S) were formed in the yields of 29%, and 8%, and 1.2%, respectively, together with the recovered *t*-4BrS (46%), as determined by gas chromatography. The time course of the product distribution is shown in Figure 1. These results are in good accord with a previous SP photochemistry of *t*-4BrS using a Hg arc lamp.^{3d}



Scheme 1.

On the other hand, quite different photoreactions were observed when a 1,4-dioxane solution of *t*-4BrS (3.5×10^{-3} M) was irradiated with 532 nm pulsed light (the second harmonic

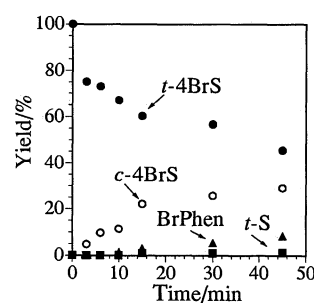


Figure 1. Time course of the product distribution during irradiation of *t*-4BrS with 266 nm laser pulses in 1,4-dioxane.

of a Nd:YAG laser, 400 MW/cm², pulse width ~ 6 ns, 10 Hz) at room temperature. Thus, irradiation of a 1,4-dioxane solution of *t*-4BrS produced selectively *unsubstituted trans*-stilbene (*t*-S), together with a small amount of *c*-4BrS as shown in Figure 2 (a). The yields of *t*-S, *c*-4BrS and the recovery of *t*-4BrS after 180 min irradiation were 37%, 1.0%, and 32%, respectively. The formation efficiency⁴ of *t*-S increased in proportion to almost the square of the light intensity as shown in Figure 2 (b).

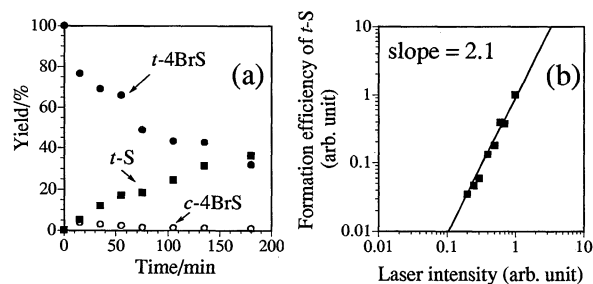
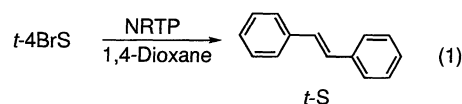


Figure 2. (a) Time course of the product distribution during irradiation of *t*-4BrS with 532 nm laser pulses in 1,4-dioxane. (b) Square dependence of the relative formation efficiency of *trans*-stilbene (*t*-S) on the incident laser intensity (532 nm) for NRTP excitation of *t*-4BrS.

In order to confirm the two-photon nature of the *t*-S formation *via* the excitation of *t*-4BrS using the 532-nm light, we have examined (1) a photoreaction of *t*-4BrS by direct singlet-triplet excitation with a CW Ar⁺ laser and (2) the wavelength dependence of the efficiency of *t*-S formation from *t*-4BrS, which

is compared with the two-photon absorption spectrum of *t*-4BrS obtained by the two-photon fluorescence (TPF) method.^{1c,5}

Irradiation of a 1,4-dioxane solution of *t*-4BrS with CW 514 nm light (1 W) from Ar⁺ laser did not yield *t*-S at all, but gave a very clean mixture of *t*-4BrS and *c*-4BrS; after 900 min irradiation, *c*-4BrS and *t*-4BrS were found to be 65% and 34%, respectively. The triplet states of *trans*- and *cis*-4-bromostilbenes would be related to the selective *cis-trans* isomerization, since the direct singlet-triplet absorption of stilbenes is observed at around 500 nm.⁶ The results also indicate that the triplet state of *t*-4BrS does not produce *t*-S via the photodissociation of the C-Br bond.

Irradiation of *t*-4BrS using pulsed light from 460 to 630 nm (80 MW/cm², pulse width 3 ns, 10 Hz)⁷ gave *t*-S as the major product, in similar to the irradiation with the pulsed 532-nm light. Figure 3(a) shows a plot of the formation efficiency of *t*-S⁴ vs a half of the wavelength of the incident laser light ($\lambda_t/2$). The action spectrum showed a maximum at around $\lambda_t = 490$ nm. The feature is in good agreement with the two-photon absorption spectrum in 1,4-dioxane obtained by monitoring dependence of the S₁→S₀ fluorescence intensity at 350 nm on $\lambda_t/2$ as shown in Figure 3.⁸ The above results evidence that the selective carbon-bromine bond fission occurs from the two-photon excited state.⁹

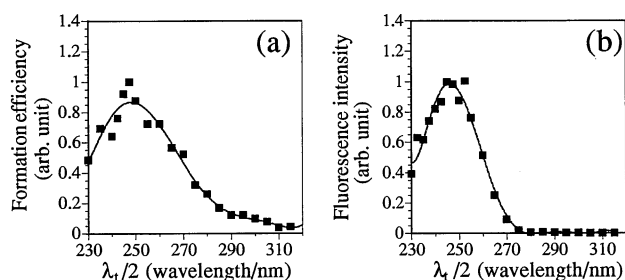


Figure 3. The wavelength dependence of (a) formation efficiency of *t*-S from *t*-4BrS and (b) relative fluorescence intensity of *t*-4BrS monitored at 350 nm.

A plausible mechanism for formation of *t*-S during the NRTP irradiation of *t*-4BrS is the spontaneous C-Br bond fission from the excited state to give *t*-stilben-4-yl radical, which abstracts hydrogen from a solvent molecule. In order to verify this mechanism, a similar NRTP photoreaction of *t*-4BrS was investigated in 1,4-dioxane-d₈. The mass spectrometry showed 100% incorporation of deuterium in *t*-S produced. Whereas formation of *t*-S was also observed for the NRTP reaction of *t*-4BrS in hexane, the pathway was less significant than in 1,4-dioxane, probably due to the lower efficiency of hexane as a hydrogen source. A possible intervention of a polar reactive intermediate like *t*-stilben-4-yl cation would be excluded, since the NRTP photoreactions of *t*-4BrS in methanol and ethanol does not yield the corresponding alkoxy substituted stilbene at all.⁹

Although the electronic nature of the two-photon allowed excited state (A_g symmetry) of *t*-S has not yet been depicted in

detail,^{2e} the observed homolytic C-Br bond fission via NRTP excitation of *t*-4BrS may suggest the nonionic biradicaloid nature of the excited state. Further works will be needed to obtain a reasonable picture of the two-photon allowed excited states of stilbenes.

References and Note

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