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Photochemical Myers—Saito and C²—C⁶ Cyclizations of Enyne—Allenes: Direct Detection of Intermediates in Solution

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Natural enediyne antitumor antibiotics utilize two thermally triggered biradical cyclizations to ultimately damage DNA by hydrogen abstraction: the Bergman¹ and Myers—Saito² reaction. In recent years, photochemical analogues³ of the Bergman cyclization⁴ and alternative photochemically triggered enediyne⁵ processes have attracted numerous research activities due to their potential applicability for photodynamic therapy (PDT).⁶ In contrast, photochemical analogues of the Myers—Saito or the C²—C⁶ cyclization of enyne—allenes have remained a *terra incognita*.

To design photochemically ignitable enyne—allenes, we resorted to the recently described photochemical reactions of enyne—heteroallenes, 7 that is, enyne—carbodiimides and enyne—ketenimines, for some meaningful clues. These cyclizations occur very effectively along the C^2-C^6 , but interestingly not along the C^2-C^7 (Myers—Saito) path. Theoretical studies by Engels explained the preference for the C^2-C^6 route and suggested, as did our experimental results, 7 that the process is initiated by triplet sensitization. Moreover, for a photochemical enyne—allene cyclization to be successful, the study recommended to avoid benzannulated derivatives due to their high excitation energy. Herein, we wish to illustrate that photochemical C^2-C^7 and C^2-C^6 cyclizations of enyne—allenes can be ignited when specially designed systems are used.

Scheme 1. Thermal and Photochemical Myers–Saito (C^2-C^7) and C^2-C^6 Cyclizations of Enyne–Allenes

Scheme 2. Model Compounds 1a-e

Enyne—allenes 1a-e (Scheme 2) were composed along the input requirements: (i) avoid benzannulation, (ii) use cycloalkenes to prevent cis—trans isomerization, (iii) attach internal triplet sensitizer units as in enyne—heteroallenes⁷ at position R' (e.g., naphthalene,

triphenylamine, and carbonyl groups), and (iv) use a TIPS group at the alkyne to raise the cyclization barrier and thus to avoid thermal reactions during photolysis.^{9a}

As programmed, enyne—allenes **1** underwent thermal cyclization only at elevated temperatures (DSC^{9b} results: **1a**, $T_{\text{onset}} = 152$ °C; **1b**, $T_{\text{onset}} = 135$ °C; **1c**, $T_{\text{onset}} = 143$ °C; **1d**, $T_{\text{onset}} = 151$ °C; **1e**, $T_{\text{onset}} \approx 190$ °C [affected by polymerization]). In line with other enyne—allenes^{10a–c} carrying bulky groups^{10d} at the alkyne terminus, the thermal cyclization of **1a–e** furnished only C²–C⁶ products (i.e., **2a–e**) by a stepwise ene process. Formal Diels—Alder products^{10c} were not formed except for trace amounts of **3** from **1b**. Loss of the TIPS group in **3** may be due to desilylation of the highly strained **3**′ in the chromatographic process.¹¹

All efforts to photocyclize ${\bf 1a}$ in hexane or toluene were met with failure, most likely due to the low triplet energy $(E_{\rm T})$ of bromonaphthalene $(E_{\rm T}=59~{\rm kcal~mol^{-1}})$, 12 suggesting to go to stronger internal triplet sensitizers. Indeed, irradiation of ${\bf 1b}$ (R' = biphenyl; $E_{\rm T}=65.0~{\rm kcal~mol^{-1}})^{12}$ in toluene at 300 nm in the presence of 1,4-cyclohexadiene led to photocyclization (Table 1) revealing both the ${\rm C^2-C^6}$ product ${\bf 3}$ (12%) and the Myers–Saito product ${\bf 4}$ (24%). Their structures were confirmed using NMR techniques and also by spectral comparison of ${\bf 4}$ with independently synthesized ${\bf 6}$. ${\bf 4}$ and ${\bf 6}$ showed a characteristic triplet for the benzhydrylic hydrogen at $\sim 3.8~{\rm ppm}$ ($^3J\sim 7.5~{\rm Hz}$). Notably, calculations⁸ predicted parallel formation of both Myers–Saito and ${\rm C^2-C^6}$ products via triplet excitation of simple enyne–allenes, although with a preference for ${\rm C^2-C^6}$ products.

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Table 1. Yields of Photochemical Myers-Saito (C2-C7) and C2-C6 Cyclization Products (1,4-CHD: 1,4-cyclohexadiene) at 17 ± 2 °C

compound (conc)	solvent (irradiation time)	$\lambda_{ ext{max}}$ /nm (in hexane)	C ² –C ⁶ products (yield)	C ² –C ⁷ products (yield)
1a (1.78 mM)	toluene + 1,4-CHD ^a (6 h)	283, 300		
1b (1.96 mM)	toluene $+$ 1,4-CHD ^a (7 h)	290, 300 (sh)	$\frac{3}{(12\%)^b}$	$\frac{4}{(24\%)^b}$
1c (1.36 mM)	toluene $+$ 1,4-CHD ^{a} (6 h)	320	2c (15%) + 7 (15%) ^c	
1d (5.28 mM)	2:1 hexane/1,4-CHD (14 h)	277, 290 (sh)	$(32\%)^d$	
1d (5.28 mM)	2:1 hexane/1,4-CHD (4 h)	277, 290 (sh)	2d (61%) ^e	
1e (7.29 mM)	toluene + 290 equiv of 1, 4-CHD (22 h)	282, 296	2e (7%) ^c	

^a 100-fold amount of 1,4-CHD compared to 1. ^b Isolated yield based on **1b.** ^c Isolated yield. ^d Isolated yield based on **1d**: 75:25 mixture of **2d** and a geometrical isomer. e Yield based on 1d after 4 h (18% conversion).

In contrast to that of 1b, the photochemical reaction of 1c-d provided only C²-C⁶ products. Irradiation of **1c** at 300 nm furnished a 1:1 mixture of 2c and 7. The latter is again a formal Diels-Alder product that lost a TIPS group.11 Irradiation of 1d under analogous conditions afforded 2d in 32% yield after isolation as a mixture of cis-trans isomers. Due to the photolability of 2d, the reaction was studied at low conversion (18%). Now, 2d was the sole isolable product in 61% yield. Due to ring strain effects, 13 we had originally expected that photolysis of cyclopentenyne-allene 1e could be directed toward the Myers-Saito pathway, but again a C²-C⁶ product (i.e., **2e**) was furnished (Table 1).

A laser flash photolysis (LFP) study of 1c (excitation at 355 nm) revealed the existence of two transients. The first one (λ_{max} = 440 and 505 nm) with a decay rate constant $k = (3 \pm 1) \times 10^7 \text{ s}^{-1}$ $(\tau \sim 30 \text{ ns})$ was quenched by ${}^3\text{O}_2$ at a diffusion-controlled rate. Its absorption maximum at 505 nm is actually typical for a triplet state of triphenylamine derivatives.¹⁴ Hence, we assigned this transient to ${}^{3}\mathbf{1c}^{*}$. The second transient ($\lambda_{max}=470$ nm) exhibited a lifetime $\tau = 33 \pm 5 \ \mu s$ with $k = (3.0 \pm 0.4) \times 10^4 \ s^{-1}$. Stern-Volmer quenching of the long-lived transient by 3O_2 indicated that ${}^3\mathbf{1c}^*$ is a precursor. Since the long-lived transient itself did not react with ³O₂, nBu₃SnH, and by 1,4-cyclohexadiene, we assigned it to the singlet biradical. Due to a high barrier (ca. 16 kcal mol⁻¹),¹⁵ it should exist as two noninterconverting rotamers (8 and 9), whose ratio is reflected in the product ratio 7/2c. After the decay of the long-lived transient, a residual broad absorption band remained that coincided largely with that of 2c.

It is interesting to see that the products of the photocyclization of cyclohexenyne-allenes 1a-d are related to the triplet energies of the substituents R'. Hence, with a triplet energy of about 59 kcal mol⁻¹, as in **1a**, no photocyclization occurred, whereas cyclization was seen with 1b containing a biphenyl ($E_T = 65.0$ kcal mol⁻¹), ¹² remarkably, mostly toward the Myers-Saito pathway. With even higher triplet energies (E_T: triphenylamine, 70 kcal mol⁻¹; ketone, 80 kcal mol⁻¹)¹² of groups R', the photocyclization exclusively furnished C²-C⁶ products.

To summarize, we have realized the first photochemical Myers-Saito and C²-C⁶ cyclizations of enyne-allenes. The presence of a

1c
$$N(\rho C_6 H_4 Br)_2$$
 TIPS H_{τ}^{H} $N(\rho C_6 H_4 Br)_2$ $N(\rho C_6 H_4 Br)_2$

triplet sensitizing unit at the allene terminus and the LFP results suggest that the cyclization proceeds along the triplet manifold as predicted by DFT calculations.⁸ An intermediate with $\tau = 33 \pm 5$ us was tentatively assigned to a singlet biradical. Further studies to elucidate the details of the photochemical initiation for enyneallenes and other Cope-type¹⁶ cyclizations are underway.

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Supporting Information Available: Experimental procedures, ¹H, ¹³C spectra for all compounds, and LFP results. This material is available free of charge via the Internet at http://pubs.acs.org.

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