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Stefano Menichettia; Mattia Moria; Cristina Nativia

^a Dipartimento di Chimica Organica, Polo Scientifico Università di Firenze, Sesto Fiorentino, Firenze, Italy

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Inverse Electron Demand Hetero Diels–Alder Reactions of Solid Supported α -Acilthiones

Stefano Menichetti Mattia Mori Cristina Nativi

Dipartimento di Chimica Organica, Polo Scientifico Università di Firenze, Sesto Fiorentino, Firenze, Italy

A solid supported dienic α,α' -dioxothione, obtained from the corresponding resinlinked β -chetoester, is able to react with vinyl ethers to give chemo- and regiospecifically the expected oxathiin cycloadducts. Trans-esterification allowed a quantitative and very clean cleavage of the products from the solid support.

Keywords Diels-alder reactions; dioxothione; oxathiins; solid phase synthesis

INTRODUCTION

In the past few years we demonstrated the utility of the phthalimidesulfenyl chloride (PhthNSCl, Phth = Phthaloyl)-mediated generation of α -oxothiones and several related thiocarbonyl species. This methodology is based on the electrophilic introduction of the *N*-thiophthalimide group on a suitable α -functionalized carbon nucleophile followed by base-catalyzed formation of the carbon-sulfur double bond via phthalimide anion elimination (Scheme 1).

By applying this procedure to β -diketons, β -ketoesters, and N-sulfonyl- β -iminoesters, it is possible to generate the corresponding α,α' -difunctionalized tioketones, which behave as efficient electron-poor dienes in chemo- and regiospecific and highly stereoselective inverse electron demand Diels–Alder reactions with a plethora of electron-rich alkenes, to give oxathiin cycloadducts with interesting synthetic applications (Scheme 1).^{1,2}

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Address correspondence to Stefano Menichetti, Dipartimento di Chimica Organica, Polo Scientifico Universitá di Firenze, Via della Lastruccia 13, 50019 Sesto Fiorentino, Firenze, Italy. E-mail: stefano.menichetti@unifi.it

Phth = Phthaloyl; XH = OH, NHSO₂Ph; Y = COR, COOR; EDG = OR, SR, NCOR, Ar

SCHEME 1

Among the features required in a successful synthetic protocol, the possibility of its exploitation on solid phase is nowadays particularly appreciated.

In this communication we report our preliminary results on the generation of a solid supported α,α' -dioxothione from a β -ketoester functionalized resin.

RESULTS AND DISCUSSION

 β -Ketoesters are among the more versatile reagents in organic synthesis, and consequently several methods have been reported for their loading on solid support.³ For this project we decided to use the Wang resin on which the β -ketoester moiety can be introduced by either simple trans-esterification with t-butylacetoacetate or using an acyl Meldrum's acid derivative.³ Both procedures gave the expected modified resin **1** as demonstrated by FTIR and ¹H NMR analysis even though, in our hands, the trans-esterification protocol was more convenient (Scheme 2).

The next step was the sulfenylation with PhthNSCl, which was the crucial point of the procedure because the co-formation of HCl could degrade Wang resin. The reaction was performed using 1.5 equiv of PhthNSCl, in dry CH_2Cl_2 at rt, and FTIR analysis showed the formation of the linked α,α' -dioxothiophthalimide 2 without evidence of cleavage of the intermediates from the resin either under this condition or using an excess (3 equiv) of the sulfenyl chloride (Scheme 2).

The final step was the generation of the solid-supported dioxothione **3** by reaction with a base. In a first attempt, functionalized resin **2** was

SCHEME 2

reacted with pyridine (1 equiv) in the presence of an excess of ethyl vinyl ether 4 in CHCl₃ at 60°C for 22 h. The formation of the supported oxathiin cycloadduct 5 was confirmed by spectroscopic analysis and eventually by cleavage from the resin with MeONa in THF,⁴ which allowed us to isolate cycloadduct 6 in 23% yield as a pure compound (Scheme 2).

Several example of Diels–Alder reactions on solid phase have been reported⁵ but, to the best of our knowledge, this is the first example of a hetero cycloaddition involving a solid-supported thicketone. Moreover, it must be considered that in this reaction the diene is a reactive intermediate that has to be generated on the resin and trapped by the electron-rich dienophile.

Crucial improvements to the methodology were obtained using a one-step procedure. Thus the resin-linked β -ketoester 1 was reacted in CH_2Cl_2 with PhthNSCl and, in sequence, with Et_3N and 4, and the mixture heated at $60^{\circ}C$ for 19 h. The same cleavage with an excess of MeONa in THF (5 h at rt) of the oxathiin modified resin 5 afforded the pure cycloadduct 6 in 81% yield (Scheme 2). Because the Wang resin requires five reactions (trans-esterification, sulfenylation, dioxothione generation, cycloaddition, and cleavage) to the final product 6, the overall yield indicates that every step occurs with an almost quantitative conversion.

Using resin 1 as a suitable source of the supported dienic dioxothione 3, the reaction was carried out with vinyl ethers 7–9 under the aforementioned conditions and, in any case, the expected oxathiin cycloadducts 10–12 (Table I) were isolated as single isomers in satisfactory yield and as pure compounds directly from the washing of the resin after cleavage.

Reactions of a α, α -Dioxothione on Sond Phase		
Dienophile	Product	Yield^a
	S 6	81%
4	0 0 0 0 10	$50\%^b$
, O 8	0 0 0	$53\%^b$
9	0 12	$40\%^b$

TABLE I Oxathiins 6, 10–12 Prepared by Hetero Diels–Alder Reactions of a α,α' -Dioxothione on Solid Phase

Together with their spectroscopic data, the definitive attribution of the structural identity of derivatives **6**, **10–12** was easily achieved by comparison with authentic samples obtained in solution using the 3-oxo-2-thioxo-butyric acid methyl ester¹ as diene.

CONCLUSION

In conclusion the phthalimidesulfenyl chloride-mediated synthesis of α , α' -dioxothiones can be successfully applied on solid phase. Using a β -ketoester functionalized Wang resin, it is possible to generate in situ an α -oxothione that participates as electron-poor diene to hetero Diels–Alder reactions with vinyl ethers. The solid-supported cycloadducts can be quantitatively cleaved from the resin by trans-esterification, which afforded the required oxathiins, which do not deserve further purification.

The application of this methodology to other dienophiles as well as different sulfur-containing dienes is under investigation.

^aIsolated yield of the pure command (>95% by ¹H NMR) after cleavage.

^bIsolated yield on a single run without optimization.

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