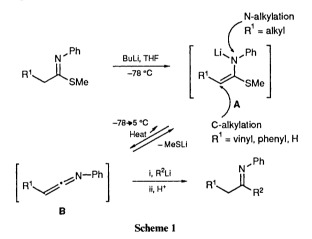
Synthesis of Silylated Ketenimines from a Metallated N-Phenyl α -Trimethylsilyl Imido Thioester

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Lithiated methyl *N*-phenyl-2-trimethylsilylethanimidothioate is regioselectively *C*-alkylated at -78 °C but, at a higher temperature in the presence of 1 equiv. of butyllithium, it is converted into a lithiated ketenimine which can be protonated or alkylated to give new and stable silylated ketenimines

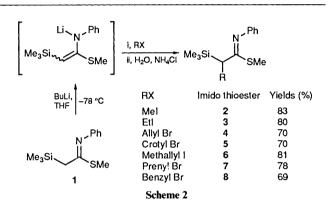
Our previous work concerning the reactivity of metallated imido thioesters¹ has shown that the regioselectivity of their alkylation, at low temperatures, is sensitive to the nature of the R^1 group (Scheme 1). The alkylation occurs at the nitrogen atom when R^1 is an alkyl substituent and at the α -carbon atom when R^1 is a phenyl or a vinyl group involving a greater delocalization of the negative charge. Moreover, at higher temperatures, the metallated imido thioester (lithium or magnesium methylthio enaminate), in the presence of organometallics (R^2Li or R^2MgX), leads to imines resulting formally from a substitution of the methylthio group by R^2 . This was interpreted (Scheme 1) as the result of the elimination of MeSLi



leading to the formation of an equilibrium between the enaminate A and the ketenimine $B^{2,3}$. This heterocumulene then undergoes a nucleophilic attack of the carbanionic part of the organometallic (R^2) at its central carbon atom. However, in these reactions such an intermediate ketenimine was not isolated.

A study of the reactivity of metallated methyl N-phenyl-2trimethylsilylethanimidothioate 1 presented herein firstly shows the influence of the trimethylsilyl group towards the regioselectivity of the alkylation at -78 °C and secondly confirms the existence, at higher temperatures of the assumed equilibrium by the obtention of new stable silylated ketenimines.

The silylated imido thioester 1 was prepared from trimethylsilylmethylmagnesium chloride and phenyl isothiocyanate.⁴ The metallation of 1 by butyllithium and the addition of the alkylating agent was performed in THF at -78 °C. The temperature was then allowed to warm slowly to 20 °C and the new α -silylated imido thioesters 2–8 were isolated in 70–83% yield (Scheme 2). No product resulting from an *N*-alkylation was detected. This demonstrates that the alkylation is selec-



tively orientated towards the C-alkylation by the trimethylsilyl group.

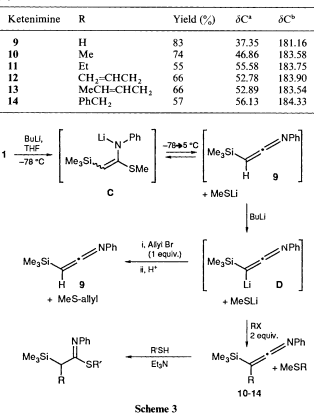
Formation of these new silvlated heterocumulenes can be rationalized by the initial formation of the lithiated enaminate C at -78 °C and with warming, the silvlated ketenimine 9 is formed. Then, due to the presence of the second equivalent of BuLi and to the carbanion stabilizing effect of the trimethylsilyl group, the deprotonation of the ketenimine 9 occurs (instead of the addition of BuLi as observed previously when R¹ = alkyl). This shifts the equilibrium towards the formation of the lithiated and silvlated ketenimine D which is then regioselectively C-alkylated.[†] The second equivalent of halide used in these reactions, alkylated MeSLi and this avoids a partial addition of MeSH to the ketenimines at the hydrolysis step.[‡]

The replacement of alkylation by protonation allowed us to obtain the ketenimine 9 together with *ca.* 20% of the starting imido thioester 1. The formation of 1, which very probably results from the addition of methanethiol to the ketenimine 9 at the hydrolysis step, was minimized by the addition, at low temperature, of only 1 equiv. of alkylating agent (allyl bromide gave the best result) before hydrolysis. This can be rationalized by a selective allylation of MeSLi at -20 °C without any significant allylation of the lithiated ketenimine **D**. Under these

[†] Generally the IR spectrum of the crude ketenimine obtained by this pathway, presents, beside the strong band at 2000 cm⁻¹, a smaller band at 2140 cm⁻¹. This can be attributed to the formation of a silylated ynamine resulting from the protonation or alkylation on the nitrogen of the *N*-lithiated ynamine form of the intermediate **D**. In some cases, the silylated ynamines was also detected by ¹H NMR and their formation did not exceed 12%. However, after flash distillation, the ketenimines only were isolated.

[‡] We indeed observed that thiols readily add to the silylated ketenimines to give imido thioesters, but some desilylation occurs. This desilylation can be minimized by using the trimethylammonium salts of the thiols.

Table 1 Yields and $^{13}\mathrm{C}$ NMR characteristics of (Me_3Si)-(R)C^a=C^b=NPh



conditions, the ketenimine 9 was isolated in a 80% yield after flash distillation.

These silvlated heterocumulenes are new potential intermediates in organic synthesis and their reactivity will be examined. In particular, the lithiated silvlated ketenimine **D** could possibly be compared to the phosphaketenimine ylide [*N*-phenyl(triphenylphosphoranylidene)ethenimine] which is a useful synthetic tool described by Bestmann.⁵ We are also at present studying the generality of this new route to ketenimines starting from imido thioesters substituted at the α -position by other carbanion stabilizing groups.* Typical Experimental Procedure (for 4).—BuLi (~1.5 mol dm⁻³ solution in hexane; 1 mmol) was added to a stirred solution of the imido thioester 1 (0.2 g, 0.85 mmol) in tetrahydrofuran (10 cm³) at -78 °C under nitrogen. Stirring was continued at this temperature for 2 h after which allyl bromide (0.086 g, 1 mmol) was added and the reaction mixture was allowed to warm slowly (~15 h) to room temperature. The reaction mixture was poured into saturated aqueous ammonium chloride and extracted with light petroleum. The combined organic phases were dried (Na₂SO₄) and evaporated under reduced pressure to give an oil which was purified by bulb-to-bulb distillation (~160 °C; 0.1 Torr) to afford **4** as a pale yellow oil.

In contrast, when 2 equiv. of butyllithium were added for the metallation at -78 °C, and the reaction mixture allowed to warm to *ca*. 5 °C before the addition of 2 equiv. of alkyl or allylic halides, the silylated ketenimines **10–14** were isolated in 55–74% yield (Scheme 3 and Table 1). These were sufficiently stable to be purified by flash distillation.

Typical Experimental Procedure (for 9 and 12): BuLi (ca. 1.5 mol dm⁻³ solution in hexane; 1.7 mmol) was added to a stirred solution of the imido thioester 1 (0.2 g, 0.85 mmol) in tetrahydrofuran (10 cm³) at -78 °C, under nitrogen. Stirring was continued at this temperature for 2 h after which the reaction mixture was allowed to warm to ~ 5 °C (out of the cooling bath) and then cooled again to <-20 °C. Allyl bromide (respectively: 0.85 mmol and 1.7 mmol for the preparation of 9 and 12) was added and the temperature was allowed to warm slowly to room temperature. After work-up (identical with that for 4) and bulb-to-bulb distillation, 9 (~ 100 °C; 0.03 Torr) and 12 (~ 150 °C; 0.05 Torr) respectively were isolated as pale yellow oils.

All the silvlated ketenimines (9–14) present a strong band at 2000 cm⁻¹ in their IR spectra and their ¹³C NMR characteristics are given in Table 1.

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^{*} From the methyl *N*-phenyl-2-phenylethanimidothioate, we have already been able to obtain, *via* the corresponding lithiated ketenimine, the corresponding *N*-phenyl(methyl)phenylketenimine [N-(2-phenyl-propenylidene) aniline].