

# DIRECTED 2-METALATION OF *N,N*-DIETHYL *O*-3-(4-TRIMETHYLSILYL)PYRIDYL CARBAMATE. A SILICON PROTECTION ROUTE FOR HIGHLY SUBSTITUTED PYRIDINES†

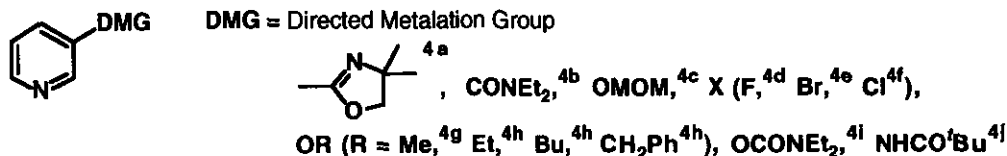
Masao Tsukazaki and Victor Snieckus\*

Guelph-Waterloo Centre for Graduate Work in Chemistry, University of Waterloo,  
Waterloo, Ontario, Canada N2L 3G1

**Abstract**— LiTMP metalation of *N,N*-diethyl *O*-3-(4-trimethylsilyl)pyridyl carbamate (**4**) followed by quenching with a number of carbon and heteroatom electrophiles affords a variety of 2-substituted derivatives (**5**) (Table 2) thus providing a versatile new methodology for the synthesis of polysubstituted pyridines.

The Directed *ortho* Metalation (DoM) reaction is currently a fertile playground for the regiospecific construction of polysubstituted aromatics.<sup>1</sup> In the context of heteroaromatics, the DoM reaction for  $\pi$ -excessive (furan, thiophene) rings is also a long recognized and widely explored process;<sup>2</sup> however, its application to  $\pi$ -deficient (pyridine, diazine) systems<sup>3</sup> was detained by the prior evidence that such low-LUMO MO substrates undergo facile nucleophilic attack by RLi and RMgX reagents. Despite this precognition, the past decade has witnessed the development of a number of DMGs, already of demonstrated utility in aromatic DoM chemistry, which promote pyridine *ortho* deprotonation thus allowing the emergence of new pyridine synthetic methodologies.<sup>3</sup> The body of knowledge for 3-DMG-substituted pyridines (Scheme 1) indicates primarily a 4-metalation regiochemistry for a variety of carbon- and heteroatom-based DMGs, 2-deprotonation has been achieved only for DMG = X and OR under carefully chosen conditions.<sup>4</sup> In continuation of our DoM studies on isomeric *O*-pyridyl carbamates,<sup>4i</sup> we have found that 2-deprotonation of *O*-3-pyridyl carbamate can be achieved by adapting the silicon protection tactic<sup>5</sup> for the more reactive C-4 metalation site (**4**). These results, disclosed herein, provide new synthetic regimens for the regiospecific construction of substituted pyridines which are inaccessible by available routes.

Scheme 1

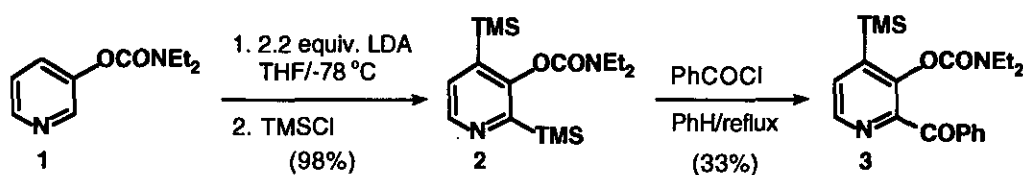


† Dedicated to Professor Ted Taylor in celebration of his 70th birthday and in recognition of his inspiring adventures in heterocyclic chemistry.



Powerful DMGs (e.g.  $\text{CONEt}_2$ , oxazoline, OMOM) promote kinetic 4-deprotonation presumably due to a strong base-DMG complexation effect while weak DMGs (e.g. X, OR) may favor 2-metalation owing to a preferred base-*N*-lone pair coordination. For the *O*-3-pyridyl carbamate, we found that the earlier observations of 4-deprotonation using *s*-BuLi/TMEDA<sup>4i</sup> or LDA<sup>6</sup> were duplicated also by metalation with *t*-BuLi or mesityl-Li in THF or Et<sub>2</sub>O solution as evidenced by CD<sub>3</sub>OD quench. However, when **1** (Scheme 2) was subjected to 2.2 equiv. of LDA followed by treatment with excess TMSCl, the 2,4-disilylated derivative (**2**) was obtained in 98% yields.<sup>7</sup> Following original observations by Pinkerton and Thames<sup>8</sup> of *ipso* carbo-desilylation of 2-TMS-pyridine, **2** was treated with excess benzoyl chloride to give **3**, an indication of a potential mild alternative to the metalation protocol for the introduction of carbon substituents into the 2-position (Table 2).

### Scheme 2



The observed formation of **2** suggested that 2-deprotonation could be achieved on the 4-TMS-carbamate (**4**).<sup>7</sup> In the first set of experiments using several RLi reagents and  $(\text{PhS})_2$  as the electrophile (Table 1), low to modest yields of products (**5f**) (E = SPh) were obtained. Thus both *s*-BuLi/TMEDA and *t*-BuLi gave unsatisfactory results in THF (Entries 1 and 2). Somewhat higher yields were observed using *t*-BuLi in Et<sub>2</sub>O at -78 °C (Entry

### Scheme 3

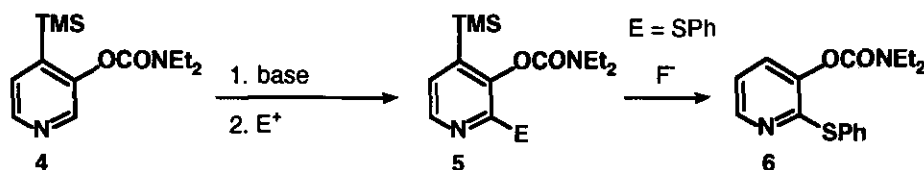


Table 1. Reactions of **4** with RLi Bases Using  $(\text{PhS})_2$  as the Electrophile

| Entry | Base                 | Solvent           | Temp, °C  | Time, min | Yield of <b>5f</b> , % |
|-------|----------------------|-------------------|-----------|-----------|------------------------|
| 1     | <i>s</i> -BuLi/TMEDA | THF               | -78       | 10        | 31                     |
| 2     | <i>t</i> -BuLi       | THF               | -78       | 15        | 29                     |
| 3     | <i>t</i> -BuLi       | Et <sub>2</sub> O | -78       | 15        | 43                     |
| 4     | <i>t</i> -BuLi       | Et <sub>2</sub> O | -110      | 10        | 49                     |
| 5     | <i>t</i> -BuLi/HMPA  | Et <sub>2</sub> O | -110      | 10        | 58                     |
| 6     | <i>n</i> -BuLi       | THF               | -78       | 10        | 0 <sup>a</sup>         |
| 7     | Mesityl-Li           | THF               | -78 → -15 | 60        | 0 <sup>b</sup>         |

<sup>a</sup> Addition of *n*-BuLi to the pyridine ring of **4** was observed. <sup>b</sup> Starting material was recovered.

3) and -110 °C (Entry 4) while the best result was realized using *t*-BuLi/HMPA (Entry 5) at the latter temperature. *n*-BuLi and mesityl-Li proved completely unsatisfactory (Entries 6 and 7).

**Table 2. Synthesis of Pyridine Derivatives Using LiTMP as Base**

| Entry | Method <sup>a</sup> | E <sup>+</sup>                    | Product   | E                     | Yield, %        |
|-------|---------------------|-----------------------------------|-----------|-----------------------|-----------------|
| 1     | A                   | CD <sub>3</sub> OD                | <b>5a</b> | D                     | 87 <sup>b</sup> |
| 2     | A                   | <i>i</i> PrCHO                    | <b>5b</b> | CH(OH)Pr <sup>i</sup> | 48              |
| 3     | A                   | PhCHO                             | <b>5c</b> | CH(OH)Ph              | 82              |
| 4     | A                   | Ph <sub>2</sub> CO                | <b>5d</b> | C(OH)Ph <sub>2</sub>  | 56              |
| 5     | A                   | PhCOCl                            | <b>3</b>  | COPh                  | 41              |
| 6     | B                   | CCl <sub>3</sub> CCl <sub>3</sub> | <b>5e</b> | Cl                    | 64              |
| 7     | B                   | (PhS) <sub>2</sub>                | <b>5f</b> | SPh                   | 79              |
| 8     | B                   | Et <sub>3</sub> SiCl              | <b>5g</b> | SiEt <sub>3</sub>     | 89              |
| 9     | B                   | PhSeCl                            | <b>5h</b> | SePh                  | 72              |

<sup>a</sup> Method A: 1.5 equiv. LiTMP. Method B: 1.2 equiv. of LiTMP. <sup>b</sup> 58% d<sub>1</sub>-incorporation by ms.

Conditions for satisfactory and convenient 2-deprotonation of **4** were achieved using the hindered strong base, LiTMP (Table 2). Thus, in appreciation of the previous experience of Iwao on pyridine amides,<sup>9</sup> a short metalation time (5 min) at -78 °C followed by rapid electrophile quench led to 2-substituted products.<sup>10</sup> Preliminary results show that a variety of carbon and heteroatom electrophiles can be introduced in modest to good yields.<sup>11</sup> Furthermore, desilylation in a selected case using TBAF<sup>12</sup> led to the formation of **6** thus opening the door for further DoM chemistry.

In summary, using optimized conditions (LiTMP/THF/-78 °C/5 min), 2-deprotonation of *O*-3-pyridyl carbamates has been achieved, **4**→**5** by taking advantage of the silicon protection protocol<sup>5</sup> at C-4. In view of the facile unmasking of the reactive C-4 metalation site in **5** and the regioselective *ipso* carbo-desilylation reactivity of **2**, broader use of the developed methodology for the construction of substituted pyridines may be anticipated.<sup>13</sup>

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  7. Use of 1.1 equiv. of LDA led to a separable mixture of **4** (88%) and **2** (9%). See also ref 6.
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  10. Metalation for 45 min followed by benzaldehyde quench gave **5c** (51%) together with 2-(*N,N*-diethylcarbamoyl)-4-TMS-3-hydroxypyridine (20%), the result of an anionic Fries rearrangement. See also ref 6.
  11. The following procedure (Method A) is representative. To a solution of LiTMP (1.50 mmol, prepared *in situ* from *n*-BuLi and tetramethylpiperidine at 0 °C) in THF at -78 °C was added a solution of **4** (0.226 g, 1.00 mmol) in THF (0.5 ml). After 5 min, PhCHO (0.209 ml, 2.00 mmol) was added and the mixture was stirred for 15 min at -78 °C. The solution was then warmed to -20 °C and quenched with sat. NH<sub>4</sub>Cl solution. Extraction with Et<sub>2</sub>O followed by drying (MgSO<sub>4</sub>) and column chromatography afforded pure **5c** (0.306 g, 82%) as colorless crystals, mp 71-72 °C (Et<sub>2</sub>O-hexane).
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  13. All new compounds show spectroscopic (ir, <sup>1</sup>H and <sup>13</sup>C nmr, ms) and combustion analysis data fully consistent with their proposed structures.

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