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lation of the corresponding hydroxytellurides in a one-pot operation.

The soft nucleophilicity of organotellurolates driving the S_N2 -type lactone ring-opening reaction

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article info

ABSTRACT

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One of the most expressive characteristics of chalcogenolate anions is their soft nucleophilicity and low basicity.¹ The lithium, sodium, and magnesium chalcogenolates are able to undergo S_N 2-type reactions in the presence of acidic functionalities, for example, alcohols, $²$ allowing the incorporation of the chalcogen</sup> moiety into organic substrates. This property of the metal chalcogenolates associated with their peculiar reactivity has been explored in the preparation of multi-functionalized chalcogen compounds, which can be submitted to further transformations based on the rich chalcogen chemistry.^{[3,4](#page-3-0)}

For many years we have been devoting our attention to the chemistry of organotellurides, 5 and more recently we became especially interested in applying these compounds as starting materials for natural product synthesis, exploring their inherent chemical properties.^{[6](#page-3-0)}

Organotellurium entities present a dual reactivity, that is, in metal organotellurolates the tellurium atom behaves as a soft nucleophile and in diorganotellurides it is attacked by nucleophiles, especially organolithiums, generating a new anionic center at the carbon atom where the tellurium atom was originally attached. This concept has been applied by us in the preparation of functionalized organolithium reagents by Te/Li exchange, and other organometallics by further transmetallation of the originally obtained organolithiums.[5,6](#page-3-0) These organometallics have been used in the synthesis of bioactive compounds. $6,7$

In this work, taking advantage of the high soft character of the metal organotellurolates, we performed S_N 2-type lactone ringopening reactions by lithium organotellurolates, δ aiming to obtain multifunctional building blocks with potential application in organic synthesis.

Lithium and magnesium organotellurolates were reacted with lactones producing the corresponding tellurocarboxylic acids. Treatment of the reaction mixture with lithium aluminum hydride allowed the iso-

> The most practical method to prepare metal tellurolates, which has been extensively employed by us in recent years, is the direct reaction of an organolithium or Grignard reagent with elemental tellurium in THF at room temperature, so avoiding the manipulation of bad-smelling alkaneditellurides.⁸

> Reaction of metal organotellurolates with lactones in THF under reflux gave the corresponding tellurocarboxylic acids.⁹ Treatment of the reaction mixture with lithium aluminum hydride led to the corresponding hydroxytellurides in a one-pot operation^{[10](#page-3-0)} ([Scheme 1](#page-1-0)).

> Initially, screening was carried out looking for the best nucleophilic tellurium species for the lactone ring opening of the γ -butyrolactone 1a (Table 1).

> As can be seen in Table 1, n-butyltellurol, the supposed nucleophilic tellurium species resulting of the reaction of lithium n-butyltellurolate with a proton source, failed to react with lactone 1a even at long reaction times under reflux (Table 1, entries 1–3). Sodium n-butyltellurolate generated in situ by reacting the badsmelling n-dibutylditelluride with sodium borohydride in THF gave the desired product in 42% isolated yield (Table 1, entry 4). Two equivalents of sodium n-butyltellurolate increased the yield, but not satisfactorily (Table 1, entry 5). On the other hand, when lithium n-butyltellurolate, prepared from elemental tellurium and n -butyllithium in THF, was reacted with $1a$, compound $4a$ was isolated in 80% yield after 9 h under reflux (Table 1, entry 6). Use of 1.5 and 2.0 equiv of the nucleophile did not improve significantly the product yield (Table 1, entries 7 and 8). The magnesium

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Scheme 1. Lactone ring opening by metal organotellurolates and reduction of the obtained carboxylate to the corresponding alcohol.

Table 1

Influence of the metal butyltellurolate in the ring-opening reaction of the γ -butyrolactone 1a

 a Generated by reacting elemental tellurium with *n*-butyllithium.

b Generated by reacting elemental tellurium with *n*-butylmagnesium bromide.

^c Reactions performed at room temperature gave 4a in low yields.

n-butyltellurolate presented reactivity similar to that of the lithium analog (Table 1, entry 9).

Use of HMPA or 18-crow-6 ether did not improve the yields. With these results in hands the best reaction conditions were applied to other lactones and metal organotellurolates. Table 2 summarizes the results obtained.

Lithium and magnesium organotellurolates reacted with five, six, and seven-member ring lactones in moderate to good yields (Table 2, entries 1–8). Some 4-substituted five-member ring lactones (Table 2, entries 7–10) were also reacted with metal organotellurolates. Albeit lactone 1d underwent reaction with both lithium and magnesium tellurolates in good yields (Table 2, entries 7 and 8), the more sterically hindered homologous 1e and 1f failed to react in the same conditions (Table 2, entries 9 and 10).

As mentioned before, we are interested in developing functionalized tellurides aiming their use in the synthesis of biologically active compounds. In this context, we developed extensive studies on the preparation of C,O-dianions from hydroxytellurides.^{[6,7](#page-3-0)} As an exten-

^a Isolated yields.

sion of these studies it would be interesting to transform the organyl tellurocarboxylic acids 2 into hydroxytellurides 5 (Table 3).

To this end, initially several reducing agents were tested to transform 3 into 5c, as shown in Table 3.

Stoichiometric or excess amounts of sodium borohydride under reflux gave the hydroxyl telluride 5c in only modest yields (Table 3, entries 1 and 2). Under similar conditions the use of DIBAL-H led to the desired product in better yields (Table 3, entries 3 and 4). However lithium aluminum hydride gave the best results (Table 3, entries 5–7) allowing the preparation of 5c in a one-pot process in 75% yield when 0.8 equiv of LiAlH $_4$ was used. When higher amounts of $LiAlH₄$ were used lower yields were obtained (Table 3, entries 5 and 6).

The conditions employed for entry 7 (Table 3) were applied in the preparation of hydroxytellurides 5a–5e as presented in Table 4.

Only moderated yields were achieved in the reaction of lactones 1c and 1d with lithium n-butyltellurolate (Table 4, entries 4 and 5). By using phenyltelluro magnesium bromide as the nucleophile in the reaction with lactone 1a compound 5a was obtained in 65% isolated yield, after treating the reaction medium with $LiAlH₄$ (Table 4, entry 1). The carboxylate, produced by opening γ -valerolac-

Table 3

One-pot reduction of carboxylates to the corresponding alcohols

^a Isolated yields.

tone^{1b} with 2a, was reduced in situ to the corresponding alcohol 5c in 75% isolated yield (Table 4, entry 3).

As mentioned before, the Te/Li exchange reaction is one of the most synthetically useful reactions of the organotellurium compounds. In view of this fact, we submitted the prepared hydroxytellurides to the reaction conditions for this transformation. The hydroxytellurides **5b** and **5c** were treated with *n*-butyllithium in THF and then octanal (6) was added to the reaction mixture. In the case of compound 5b, the diol 7a was obtained in 75% yield. Under the same reaction conditions 5c failed to react, being recovered unchanged on the work-up. When 5c was transformed into the corresponding THP ether, the Te/Li exchange was successful, leading to compound 7b in 69% yield on reaction with 6 ([Scheme 2\)](#page-3-0).

Scheme 2. Te/Li exchange reaction and capture of the corresponding organolithium with 6

Finally, it must be pointed out that all the organotellurides prepared in this work, except dibuylditelluride and dibutyltelluride, are not bad smelling and are stable to air and light.

In conclusion, a practical method to prepare organotellurocarboxylic acids in a single operation has been developed by using in situ-generated metal organotellurolates in a S_N2 -type reaction with lactones. In addition, the intermediate organyl telluride carboxylates can be directly reduced to the corresponding hydroxyl tellurides, which are synthetic equivalents of the corresponding organolithiums by Te/Li exchange reaction.

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- 9. Typical procedure for the lactone ring opening by lithium n-butyltellurolate. 4-(nbutyltellanyl)butanoic acid (4a). To a dry two-necked 50 mL round-bottomed flask equipped with magnetic stirring, reflux condenser, and a rubber septum under nitrogen, was added elemental tellurium (0.51 g, 4.0 mmol) previously dried overnight in an oven at 100 °C. Then dry tetrahydrofuran (15 mL) was added. To the stirred suspension was added n-butyllithium (2.9 mL, 1.4 M in hexane). The mixture was then heated to 80 °C and the lactone 1a (0.22 g, 2.6 mmol) was added all at once. The reaction was monitored by gas chromatography. After 9 h at 80 $^{\circ}$ C the reaction medium was allowed to reach the room temperature and then it was diluted with AcOEt (20 mL) and washed with saturated solution of NH4Cl (15 mL). The phases were separated and the aqueous phase was extracted with AcOEt $(2 \times 20 \text{ mL})$. The organic phase was dried over MgSO₄ and the solvents were evaporated under reduced pressure. The residue was purified by flash column chromatography eluting first with hexane to remove dibutylditelluride and then with hexane/AcOEt $(8:2)$ to remove the product. Yield: 0.61 g $(86%)$. ¹H NMR: $(300 \text{ MHz}, \text{CDCl}_3)$. ppm) δ 10.79 (br s, 1H); 2.66 (t, J = 7.5 Hz, 2H); 2.65 (t, J = 7.5 Hz, 2H); 2.48 (t, $J = 7.2$ Hz, 2H); 2.06 (quint., $J = 7.5$ Hz, 2H); 1.73 (quint., $J = 7.2$ Hz, 2H); 1.38 (sext., J = 7.5 Hz, 2H); 0.92 (t, J = 7.2 Hz, 3H). ¹³C NMR (75 MHz, CDCl₃, ppm) δ 179.43; 36.10; 34.28; 27.12; 25.07; 13.41; 2.83; 1.10. LRMS m/z (rel. int.) 274 (M⁺, 15); 87 (100); 57 (54); 41 (70). IR v_{max} (neat) 3053, 2957, 1646, 1370.
- 10. Typical procedure for the reduction of the organotelluro carboxylic acids. 4- (Phenyltellanyl)but-1-ol (5a) (one-pot procedure). To a dry two-necked 50 mL round-bottomed flask equipped with magnetic stirring, reflux condenser, and a rubber septum under nitrogen, was added elemental tellurium (0.51 g, 4.0 mmol) previously dried overnight in an oven at 100° C. Then dry tetrahydrofuran (15 mL) was added. To the stirred suspension was added phenylmagnesium bromide (2.9 mL, 1.4 M in hexane). The mixture was then heated to 80 °C and the lactone 1a (0.22 g, 2.6 mmol) was dissolved in THF (5 mL) and added all at once. The reaction was monitored by gas chromatography. After heating at 80 °C for 10 h, LiAlH₄ (0.079 g, 2.1 mmol) was added in small portions. After 5 h at 80 $^{\circ}$ C the reaction mixture was allowed to reach the room temperature and then it was diluted with AcOEt (30 mL), treated with NaOH (1 mL, 1 M), and filtered under vacuum. The organic phase was washed with saturated solution of NH4Cl (15 mL). The phases were separated and the organic phase was washed with AcOEt $(2 \times 20 \text{ mL})$. The organic phase was dried with MgSO₄, filtered, and the solvent was evaporated. The residue was purified by column flash chromatography eluting first with hexane to remove diphenylditelluride and then with hexane/AcOEt (8:2) to remove the product. Yield: 0.47 g (65%). ¹H NMR: (300 MHz, CDCl₃, ppm) δ 7.71 (dd, J = 6.9 Hz and 1.2 Hz, 2H); 7.16–7.29 (m, 3H); 3.61 (t, J = 6.3 Hz, 2H); 2.91 (t, J = 7.5 Hz, 2H); 2.31 (br s, 1H); 1.87
(quint., J = 7.2 Hz, 2H); 1.63 (quint., J = 6.6 Hz, 2H). ¹³C NMR (75 MHz, CDCl₃ ppm) d 138.31; 129.13; 127.53; 111.63; 62.03; 34.77; 28.08; 8.23. LRMS m/z (rel. int.) 280 (M⁺, 18); 73 (96); 55 (100); 43 (48). IR v_{max} (neat) 3372, 2930 1059, 732, 691.