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Highly regioselective and chemoselective titanocene mediated Barbier-type allylation reactions[†]

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Titanocene carboxylate 1 is an excellent chemoselective reagent for unprecedented α -regioselective Barbier-type reactions. It constitutes the first titanocene(III) able to tolerate epoxides and readily reduced carbonyl compounds, such as aromatic and α , β -unsaturated aldehydes.

Titanocene(\mathfrak{m}) complexes are useful tools for mediating and catalyzing a number of useful transformations,¹ such as homolytic epoxide opening,² pinacol coupling reactions,³ and Barbier-type additions of allylic and propargylic halides or carbonates to carbonyl compounds.⁴ A drawback of these procedures is the lack of chemoselectivity in the electron transfer step. For this reason, aromatic and α , β -unsaturated carbonyl compounds are usually unsuitable substrates in Barbier-type reactions.⁴ Here, we demonstrate that this shortcoming can be resolved in the case of complex **1** (Scheme 1) in the presence of Mn dust *via* complex **2a** (Fig. 1).^{5,6}

Gratifyingly, **3** and **4**, model compounds for aromatic and α , β -unsaturated aldehydes, reacted with **5** in the presence of complex **1** and Mn dust to exclusively yield α -prenylated compounds **6** and **7**. Thus, the highly useful α -regioselectivity of prenylation noted earlier



Scheme 1 Ti-mediated regioselective α -prenylation of benzaldehyde (3) and citral (4).

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Fig. 1 CV of 2 mM solution of 1 (red) and Mn–1 (black) recorded at ν = 100 mV s⁻¹ in 0.2 M TBAPF₆–THF.

could be maintained.^{4*a*,6} Remarkably, in the absence of **5**, no pinacol products are formed.⁷ Further exploring this new reactivity concluded that epoxides were also tolerated by this reagent. This suggests that the aldehydes or epoxides cannot bind to Mn–**1**.

To rationalize these findings, cyclic voltammograms (CVs) of 1 and Mn-1 in THF were recorded (Fig. 1). The oxidation of electrochemically reduced 1 reveals the presence of two species. 1^{-} (E_{pc} = -1.38 V vs. Fc⁺/Fc) is formed via electron transfer at the electrode. The process is irreversible due to the formation of $2a (E_{pa1} = -1.00 \text{ V})$ through loss of chloride. For Mn-1, 2a ($E_{pa1} = -0.95$ V) is formed as expected, due to the more efficient abstraction of the chloride and formation of $MnCl_2$. The more negative value of E_{pa1} compared to $Cp_2TiCl(E_{pa1} = -0.83 \text{ V})$ is in line with the carboxylate that is a better donor ligand than chloride. The second peak ($E_{pa2} = -0.65$ V) is also observed at higher sweep rates ($\nu = 1-50 \text{ V s}^{-1}$). Thus, the species being oxidized is an initial component⁸ of Mn-1 in THF and is not formed during the sweep. We suggest that 2b and not cationic 2c is the second component of Mn-1 for a number of reasons. 2c should have a potential similar to that of $[Cp(C_5H_4^{t}Bu)Ti]^+$ ($E_{pa} = -0.47$ V). It has been recently shown that cationic titanocene(III) complexes open epoxides.9 Both findings are in contradiction to the behavior of Mn-1. In agreement with the observations, 2b is more difficult to oxidize than 2a because the electron density is withdrawn from Ti through carboxylate coordination. The steric shielding of Ti in 2b will

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Communication

be similar to that in **2a** and thus epoxide and aldehyde binding is as difficult for **2b** as for **2a**.



To understand the unprecedented chemoselectivity, we carried out the model reactions with Mn-reduced Cp₂TiCl(OMe),¹⁰ Cp₂TiCl(OAc),¹¹ and Cp₂TiCl(OBz).¹² Unselective electron transfer reactions with Cp₂TiCl₂ were observed. This suggests that the tethering of the carboxylate to the ligand is essential.

To highlight this point, the coordination of 2a to THF, benzaldehyde, and trimethyloxirane was studied by DFT calculations (Table 1).⁷

The generation of these coordination complexes is characterized by: (a) a only slightly negative enthaply of formation and (b) a highly unfavorable entropy of formation. This results in ΔG values of about +8 to +12 kcal mol⁻¹.¹³ Therefore, aldehyde and epoxide complexation seems to be precluded as observed experimentally.

Next, we explored the chemoselectivity of electron transfer further (Scheme 2 and Table 2). Gratifyingly, our Barbier-type reactions using activated halides as pronucleophiles are general and take place with moderate to excellent yields.⁷ A variety of aromatic and α , β -unsaturated aldehydes are suitable. Even acetophenone, as an example of aromatic ketones, is an excellent substrate (Scheme 2). All products are valuable building blocks for terpene synthesis.^{1d} Even more interesting and



+12.5

+12.0



Scheme 2 Ti-mediated regioselective α -prenylation of aromatic and α , β -unsaturated carbonyl compounds. Conditions: for aromatic aldehydes: **1** (1 mmol), Mn dust (8 mmol), aldehyde (1 mmol), and activated halide (2 mmol) in THF. Otherwise: **1** (1.5 mmol), Mn dust (8 mmol), carbonyl compound (1 mmol), and activated halide (3 mmol) in THF. ^aPrenyl chloride can be used with similar yield (77%).

demanding are Barbier-type reactions with electrophiles (19-20) or nucleophiles (23), incorporating epoxides (Table 2).

In agreement with our DFT calculations, epoxides are readily tolerated by Mn–1.⁷ Thus, functionalized epoxypolyprenes that are highly attractive intermediates for synthesis of natural products^{1d} can be prepared from either epoxide containing carbonyl compounds or activated halides. As an attractive additional feature, our novel method allows the elimination of the often difficult regioselective epoxidation step of the polyprenic starting materials.

An important aspect of this study is the regioselectivity of the addition of the allylic nucleophile. In contrast to the numerous γ -selective additions, α -regioselective additions are considerably less common.¹⁴ Here, we obtained the α -regioisomers in very high selectivity (>92: <8) when prenyl derivatives **5**, **8** and **23** were used as pronucleophiles. Moreover, commercially available Mn dust could be used at room temperature.

Finally, we also investigated Wurtz-type coupling reactions mediated by Mn–1. Such reactions can be carried out in the presence of $Cp_2TiCl.^{15}$ However, the chemoselectivity of these reactions is low. This is not the case in our system and, hence, epoxide containing substrates can be readily coupled (Table 2, entry 8). The reaction took place with excellent yield and also with a high α, α -regioselectivity (86:14).

In summary, we have demonstrated that Mn–1 is a reagent for Barbier-type allylation reactions delivering the α -addition products with almost complete regioselectivity. The reactions proceed with an unprecedented chemoselectivity because epoxides and readily reduced carbonyl compounds, such as aromatic and α , β -unsaturated aldehydes, are tolerated. The compounds obtained are difficult to prepare using other methodologies and are valuable substrates in terpene synthesis. Moreover, the reactions can be performed at room temperature and no preformed organometallic reagents (B or Li) or especially activated metals (Ba) have to be employed.

-0.5

2a-Ep

Table 2 Regioselective Barbier-type reactions of electrophiles and nucleophiles presenting an oxirane ring mediated by $Mn-1^a$



^{*a*} Complex **1** (1.5 mmol), Mn dust (8 mmol), carbonyl compound (1 mmol), and activated halide (3 mmol) in THF. ^{*b*} This compound was obtained as a 6:1 mixture of α, α and α, γ -regioisomers.

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