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Deprotonative Cadmation of Functionalized Aromatics.

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This communication describes the deproto-metalation of a large range of aromatics including heterocycles using a newly developed lithium-cadmium base. The reaction proceeds at room temperature with an excellent chemoselectivity and efficiency, 10 and proved to be regioselective in most cases.

The deprotonative metalation of aromatic rings has been widely used as a powerful method for regioselective functionalization. Various strong bases such as alkyllithiums and lithium dialkylamides have been largely employed for this 15 purpose because of their solubility in ethers and alkanes, and also because many of them are commercially available. However, the use of alkyllithiums on their own as bases has been limited to substrates with C-H acidity enhanced by directing groups. In addition, recourse to simple lithium 20 dialkylamides for aromatics bearing reactive functions (e.g. ester or cyano groups) or sensitive π -deficient heterocycles required strictly controlled conditions (extremely low reaction temperatures, in situ trapping...) due to the high reactivity of the corresponding aryllithiums. The use of additives for 25 lithium compounds in order to get more efficient or more chemoselective bases, or else to modify the deprotonation site, is a challenging area. Representative activation ways are the formation of chelates between alkyllithiums and N,N,N',N'tetramethylethylenediamine (TMEDA), as well as the 30 formation of complexes between alkyllithiums and potassium tert-butoxide (e.g. LIC-KOR superbase).

By combining soft organometallic compounds with alkali additives (e.g. LiTMP, TMP = 2,2,6,6-tetramethylpiperidino, or LiCl), bases such as ${}^{t}Bu_{2}Zn(TMP)Li_{1}^{2} {}^{i}Bu_{3}Al(TMP)Li_{2}^{3}$ 35 (Me₃SiCH₂)₂Mn(TMP)Li·TMEDA,⁴ MeCu(TMP)(CN)Li₂⁵ and $(TMP)_2Zn\cdot 2\ MgCl_2\cdot 2\ LiCl^6$ have been prepared and used to generate functionalized aromatic compounds.

Mulvey introduced the term alkali metal-mediated metalation to depict the reactions of ate bases because the 40 reactivity ("synergy") they exhibit cannot be attained by the homometallic compounds on their own. When performed in tetrahydrofuran (THF), the reactions proved to be chemoselective, but require 1 or 2 equiv of base.

Herein we report an efficient regio- and chemoselective 45 direct cadmation on functionalized aromatics including very sensitive heterocycles using a newly designed lithium cadmate base. Among organometallics, organocadmium reagents have been mainly prepared by reaction of organic halides with cadmium metal or by transmetalation, and used as soft 50 nucleophilic reagents in organic synthesis. 8 Wittig and coworkers documented in 1951 the synthesis of Ph₃ZnLi and

Ph₃CdLi, and their efficiency to deprotonate fluorene in diethyl ether. Quenching with CO₂ and subsequent acidic work-up afforded diphenyleneacetic acid in a low yield of 55 16% after 10 days reaction time using Ph₃ZnLi whereas a satisfying 64% yield was obtained after 3 days using Ph₃CdLi, a result attributed to the size of the central metal. This prompted us to study the use of lithium cadmates for the deproto-metalation of sensitive aromatic substrates.

A recent study showed LiTMP and (TMP)₂Zn, even if not associated in the form of a zincate, could behave synergically, combining both the efficiency of LiTMP and the chemoselectivity of (TMP)₂Zn. ¹⁰ In order to seek more efficient and direct methods for introducing functionalities 65 into heteroaromatic rings, we focused the deprotonative metalation using the corresponding mixture with cadmium instead of zinc on the difference of metal size. First attempts using anisole (1a) as substrate indicated that an in situ prepared mixture of CdCl₂·TMEDA¹¹ (0.5 equiv) and LiTMP 70 (1.5 equiv) was suitable for an efficient reaction, when used in THF at room temperature. Indeed, subsequent trapping with iodine after 2 hours afforded the expected derivated 2a in 74% yield, against 30% yield using ZnCl₂·TMEDA (0.5 equiv) and LiTMP (1.5 equiv). Since (TMP)₂Cd (1 equiv) and LiTMP (1 75 equiv) give much lower conversions when used separately under the same reaction conditions, both of them play a role in the reaction mechanism. In order to obtain additional information about the active species of a basic mixture obtained from a THF solution of LiTMP and CdCl₂·TMEDA 80 (1/3 equiv), NMR and DFT studies were carried out. The analysis of the ¹³C NMR spectra revealed that LiTMP was not present in solution, suggesting the formation of a lithium cadmate. This was confirmed by the B3LYP-calculated equilibrium between LiTMP and (TMP)2Cd on one side and 85 (TMP)3CdLi on the other side, which is in sharp contrast to the corresponding zinc-lithium mixture obtained from LiTMP and ZnCl₂·TMEDA (Scheme 1).¹⁰

LiTMP +
$$(TMP)_2Zn$$
 $\Delta G = +3.8 \text{ kcal/mol}$ $N = 1.95 \text{ kcal/mol}$ $N = 1.95 \text{ kcal/mol}$ $N = 1.95 \text{ kcal/mol}$ $N = 1.96 \text{ kcal/mol}$ $N = 1.$

Scheme 1 Bond lengths at the B3LYP/6-31G*&SVP(Zn) level in Å

Representative results from the metalation-trapping sequences of benzenes bearing various directing metalation tris(2,2,6,6groups (DMG) using lithium tetramethylpiperidino)cadmate (TMP-cadmate) 5 summarized in Table 1. Veratrole (1b) was similarly regioselectively deprotonated. Polar functional groups including amide, ester, nitrile and even ketone (substrates 1c, 1d, 1e and 1f, respectively) are tolerated in the reaction. Aromatic halides 1g and 1h are chemoselectively converted to 10 the metalated derivatives, with a complete regioselectivity for the position far from the heavy halogen atom. Bromo ester 1i behaved similarly to give after trapping the iodide 2i.

Table 1 Deprotonative cadmation of functionalized benzenes

Entry	Substrate (1)		Product (2)		Yield
1	OMe	1a	OMe	2a	74%
2	OMe MeO	1b	OMe MeO I	2b	79%
3	CONEt ₂	1c	CONEt ₂	2c	91%
4	CO ₂ Me	1d	CO ₂ Me	2d	62%
5	CN	1e	CN	2e	68%
6	COPh	1f	COPh	2f	66%
7	OMe OMe	1g	Br OMe	2g	97%
8		1h		2h	83%
9	Br CO ₂ Me	1i	Br CO ₂ Me	2i	60%

^a Using ZnCl₂·TMEDA (0.5 eq) and LiTMP (1.5 eq).

We next demonstrated that TMP-cadmate was suitable for the chemoselective metalation of a large range of aromatic heterocycles, giving in general higher yields than using Zn^{10,12} (Table 2). The reaction with both π-excessive (substrates 1j, 1k and 1l) and π-deficient (substrates 1m, 1n, 1o, 1p and 1q) heterocycles was found to proceed smoothly at RT. The expected iodides formed regioselectively, except 3-iodopyridazine (2o), which was accompanied by the 4-iodo derivative 2'o (about 60/40 ratio for 2o/2'o).

Table 2 Deprotonative cadmation of aromatic heterocycles^a

	•				
Entry	Substrate (1)		Product (2)		Yield (%)
1	S	1j	S	2j	97 (73) ^b
2		1k		2k	84 (69) ^b
3	N Boc	11	N Boc	21	68 (68) ^b
4	N S	1m	N S	2m	97 (52) ^b
5	N	1n	N	2n	63 (57) ^b
6	N-N	10	N-N	20	55°
			N N	2'0	41 ^c
7	N N	1p	N I	2 p	71 (57) ^b
8		1q	$\binom{N}{N}$	2 q	63 ^d (59) ^b

^a Reactions carried out using CdCl₂·TMEDA (0.5 eq) and LiTMP (1.5 eq). ^b Using ZnCl₂·TMEDA (0.5 eq) and LiTMP (1.5 eq). ^{10,12 c} Using CdCl₂·TMEDA (1 eq) and LiTMP (3 eq). ^d Using CdCl₂·TMEDA (0.33 eq) and LiTMP (1 eq).

Starting from pyrazine (1q), the 2,5-diodo derivative 3q was isolated concomitantly in 20% yield using 0.5 equiv of TMP-cadmate, probably through dideprotonation, whereas it was avoided using 1/3 equiv. The formation of dimetalated derivatives being described using zincate¹³ or manganate^{4,14} type bases, the use of a larger amount of TMP-cadmate (1 sequiv) was attempted to deprotonate 1q. Under the same reaction conditions, the diiodide 3q was isolated in 58% yield. The method was successfully extended to five-membered substrates 1r, 1s, 1t and 11 (Table 3).

Table 3 Deprotonative dicadmation of aromatic heterocycles^a

Entry	Substrate (1)		Product (3)		Yield (%)
1		1q	I N	3q	58
2	SN SN	1r	I S	3r	50
3	s	1s	I S I	3s	74
4	o o	1t	o o	3t	81
5	N Boc	11	I N I Boc	31	60

⁴⁰ Reactions carried out using CdCl₂·TMEDA (1 eq) and LiTMP (3 eq).

In summary, highly chemoand regioselective deprotonative cadmation of functionalized aromatics including heterocycles was realized using a newly developed TMP-Cd-ate base. The latter is compatible with very sensitive 5 substrates such as diazines for which classical lithium bases can hardly be used, even at very low temperatures. 15 The aromatic lithium cadmates were evidenced using iodine as electrophile. Trapping of the furylcadmate was attempted using other electrophiles: benzoyl chloride^{8b} to afford the 10 ketone 4u, and 4-bromoanisole to give the expected coupling 16 product 5u under palladium catalysis, as depicted in Scheme 2.

Scheme 2 Other trapping reactions of heterocyclic cadmates

- Comparisons with previously described combinations of LiTMP on the one hand, and Zn,² Al,³ Mn⁴ and Cu⁵ organometallic compounds on the other hand, showed the basic mixture we here described is both more efficient and/or more chemoselective.
- Due to the toxicity of cadmium compounds, we actually try hard to develop basic mixtures containing cadmium salts as catalysts. In addition, works in order to develop new mixed lithium-metal bases of ate type still efficient and chemoselective but less toxic are in parallel under 25 investigation, profitting from the present study.

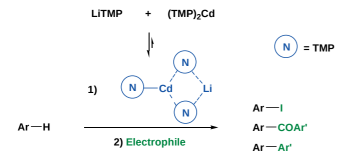
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Notes and references

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 - † Electronic Supplementary Information (ESI) available: Experimental procedures and characterizations (1H and 13C NMR spectra for all compounds), and theoretical data (details of computational methods,
- 45 cartesian coordinates and Gibbs free energies). See DOI: 10.1039/b000000x/

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Graphical and textual abstract for the contents pages.



The deproto-metalation of a large range of aromatics including heterocycles is described using a newly developed lithium-cadmium base. The reaction proceeds at room temperature with excellent chemoselectivity and efficiency, and proved to be 10 regioselective in most cases.