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Lithium Amide Assisted Asymmetric Mannich-Type Reactions of Menthyl Acetate with PMP-Aldimines

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ABSTRACT

A lithium enolate of menthyl acetate added to PMP-imines, in the presence of an equimolar amount of lithium diisopropylamide, affords the Mannich-type addition products in high stereoselectivity.

There have been few reports^{1,2} on the asymmetric Mannichtype reactions of acetate lithium enolates with stable Nsubstituted imines^{3,4} of poor electrophilicity.⁵ In contrast, propionates and other α-substituted acetate analogues have been used in the asymmetric Mannich-type reactions.⁶ Success of the Mannich reaction with a chiral acetate is

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 β -lactam products in high enantioselectivities. However, an acetate itself, in place of α -substituted esters, was not a reactive partner in the reaction, and PMP-imines were recovered mostly unchanged. After many trials, we finally reached a simple solution for this problematic reaction by using a lithium amide as an assisting agent.

We systematically studied the influence of the *para* substituents of the aniline-derived aldimines $3 \, (R^1 = Ph)$ on the reaction efficiency using menthyl acetate $1 \, (Scheme 1)$.

Scheme 1. Mannich-Type Reaction of Imine 3 with Menthyl Acetate 1 via Lithium Enolate 2, Giving 4

The lithium enolate **2** was generated by treatment of a THF solution of **1** with a slight excess of lithium diisopropylamide (LDA) at -78 °C for 1 h and was then treated with **3**. Contrary to the lack of the addition product **4a** (R¹ = Ph, R² = 4-MeO) with PMP-imine **3a** (R¹ = Ph, R² = 4-MeO) (Table 1, entry 1), ¹⁰ imines **3b**-**3f** of anilines bearing

Table 1. Mannich-Type Reaction of **1** with **3** ($R^1 = Ph$) at -78 °C

entry	3 $(\mathbf{R}^1 = \mathbf{Ph})$	\mathbb{R}^2	<i>t</i> (h)	4 $(\mathbf{R}^1 = \mathbf{Ph})$	yield (%)	dr
1	a	4-MeO	5	a	0	
2	b	4-H	5	b	62	93:7
3	c	$4-CF_3$	0.3	c	87	80:20
4	d	4-CO ₂ <i>i</i> -Pr	0.3	d	80	72:28
5	e	4-CN	0.3	e	95	60:40
6	f	$4-NO_2$	0.3	f	98	56:44
7	g	2-MeO	3	g	86	84:16

electron-withdrawing substituents or no substituent at the 4-position were good partners for the Mannich-type reaction at -78 °C for 0.3 h (5 h for **3b**) to afford **4b**–**4f** (R¹ = Ph) in good to high yields [62% (93:7 diastereomer ratio (dr)) with **3b** (R² = 4-H), 87% (80:20 dr) with **3c** (R² = 4-CF₃), 80% (72:28 dr) with **3d** (R² = 4-CO₂*i*-Pr), 95% (60:40 dr) with **3e** (R² = 4-CN), and 98% (56:44 dr) with **3f** (R² = 4-NO₂) (entries 2-6)]. As expected from Yamamoto's

report,⁷ the reaction with 3g ($R^2 = 2$ -MeO) proceeded under the same conditions to give 4g in 86% yield and 84:16 dr (entry 7).

Hammett σ -constants and p K_a values of the 4-substituted anilines are likely to be the key to the success and failure of the Mannich reaction. For example, imines 3b-3f derived from anilines bearing 4-NO₂ (σ^{12} 0.81), 4-CN (0.71), 4-CO₂i-Pr (0.68 for CO₂Et), 4-CF₃ (0.53), and 4-H (0) groups are good reaction partners, and 3a bearing 4-MeO (-0.28) is not. The p K_a values¹³ of 4-NO₂ (21), 4-CN (26), and 4-H (31) anilines are smaller than or nearly equal to that (30 for MeCO₂t-Bu) of an acetate. These apparently indicate two critical points: (1) electron-withdrawing groups (R^2) at the 4-position of aniline increase the reactivity of the imine functionality, and (2) the initially formed lithium amide species of 4 plays the important role in the reaction (Figure 1). When the lithium amides of the products 4 are more stable

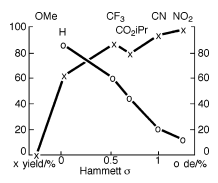


Figure 1. Hammett σ -constant, yield (\times) and dr (\bigcirc) of the reaction.

as a result of the electron-withdrawing 4-R² group or internal coordination (2-MeO), the efficiency of the reaction is higher and vice versa.

The diastereoselectivity is inversely proportional to the value of the Hammett σ -constants of R^2 , implying the important role of the reactivity of the imine functionality (Figure 1). The reaction with more reactive imines likely proceeds through a looser organization of the transition state, leading to lower diastereoselectivity down to 56:44 dr with $3f(R^2 = NO_2)$. Less reactive imines react through a tighter transition state to afford better diastereoselectivity up to 93:7 dr with $3b(R^2 = H)$.

These experiments and analyses led us to the expectation that the least activated imine **3a** would provide the highest diastereoselectivity if the reaction proceeded. As a simple solution to the problem we applied the concept of ternary reagent control⁹ that enables activation of a lithium ester enolate by forming a complex with a lithium amide and a

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⁽¹⁰⁾ The reaction at higher temperature resulted in the production of a messy mixture.

⁽¹¹⁾ The diastereomer ratio was determined by ¹H NMR (doublet Me protons appearing at around 1 ppm) of the crude product.

⁽¹²⁾ March's Advanced Organic Chemistry: Reactions, Mechanisms, and Structure, 5th ed.; Smith, M. B., March, J., Eds.; John Wiley & Sons Inc.: NY, 2001.

⁽¹³⁾ The pK_a values in DMSO quoted from the Bordwell pK_a table.

chiral ligand. For the reaction in THF, a binary complex of a lithium enolate with a lithium amide would be the reagent of choice.¹⁴

To our delight, the reaction at -30 °C for 4 h of a binary complex of menthyl acetate enolate (2-LDA), generated by treatment of 1 with 2.25 equiv of LDA in THF, converted PMP imine 3a to the desired adduct 4a ($R^1 = Ph$, $R^2 = 4-MeO$) in 76% yield and 96:4 dr (Table 2, entry 1). The

Table 2. Mannich-Type Reaction of Imine 3 with 1 via Binary Reagent of 2-LDA

$$1\xrightarrow[\text{THF, }-78\text{ °C, }1\text{ h}]{\text{LDA}}}2\text{-LDA}+3\xrightarrow[-40\text{ to }-20\text{ °C, }3-4\text{ h}]{\text{LDA}}}4$$

entry	3	\mathbb{R}^1	\mathbb{R}^2	4	yield (%)	dr
1	a	Ph	4-MeO	a	76	96:4
2	b	Ph	Н	b	58	96:4
3	h	$4-CF_3C_6H_4$	4-MeO	h	74	97:3
4	i	4-MeOC_6H_4	4-MeO	i	56	96:4
5	j	1-Naph	4-MeO	j	65	82:18
6	k	2-Naph	4-MeO	k	83	94:6

use of other lithium amides, derived from dicyclohexylamine, isopropylcyclohexylamine, or tetramethylpiperidine, had little influence on the reaction to afford **4a** with the same high selectivity of 94:6–93:7 dr and 56–80% yields. The binary reagent (**2**-LDA) also converted **3b** in 3.5 h to **4b** with 96:4 improved dr (entry 2). Under these conditions, no improvement in the diastereoselectivity was observed in the reaction with activated imines **3c**–**3f**.

The 4-substituents of benzaldehyde derivatives were not so influential on the reaction. The 4-trifluoromethyl- and 4-methoxybenzaldehyde imines $\bf 3h$ and $\bf 3i$ ($\bf R^1=4\text{-}CF_3C_6H_4$, 4-MeOC₆H₄, $\bf R^2=4\text{-}MeO$) were converted at -30 °C in the presence of an excess of LDA to $\bf 4h$ in 74% yield and 97:3 dr and $\bf 4i$ in 56% yield and 96:4 dr (entries 3 and 4). In the absence of an excess of LDA, the starting imines were recovered without formation of detectable amount of the adducts. Other than benzaldehyde imines, PMP imines of 1- and 2-naphthaldehydes $\bf 3j$ and $\bf 3k$ ($\bf R^1=1$ - and 2-Naph) were converted at -40 °C for 3 h to $\bf 4j$ in 65% and 82:18 dr and $\bf 4k$ in 83% and 94:6 dr (entries 5 and 6).

The deuterium oxide quench of the reaction with **3a** revealed that the α-position of ester **4a** was deuterized at 80%, indicating the conversion of the initially formed lithium amide of the adduct to its dianion. The stability of the dianion of the adduct was confirmed by treatment of **4a** with 3 equiv of LDA to recover **4a** in 90% yield without any loss of diastereo-integrity (Scheme 2). On the other hand, treatment of **4a** with 1.2 equiv of LDA gave the recovery of **4a** in a trace amount, along with 4-methoxyaniline in 13% yield, *N*-PMP cinnamoylamide **5** in 28% yield, and its PMP-amine

Scheme 2. Stability of 4a upon Treatment with LDA

1,4-adduct **6** in 12% yield. These products seemed to arise through elimination of 4-methoxyaniline from **4a** via its lithium enolate and subsequent 1,2-addition to a carbonyl group of cinnamate followed by 1,4-addition.¹⁵ Thus the stability of the lithium amide of **4a** is also one of the controlling factors of the reaction. It is important to note that LDA plays a dual role in the activation of the lithium enolate and the lithiation of the unstable initial adduct into the stable dianion.

The PMP group of (-)-4a, diastereomerically enriched to over 99:1 dr in 70% yield by recrystallization from methanol, was easily removed by a CAN oxidation to afford optically pure (R)- 7^{15} (Scheme 3).

Scheme 3. Conversion of Enantiomerically Pure 4a to 7

In conclusion, we have developed a new and simple methodology for the Mannich-type reaction of a chiral acetate with arylaldimines, in which the use of a binary reagent composed of a lithium acetate enolate and a lithium amide is key to the success. It is also important to note that a simple chiral auxiliary, menthol, is operative as an efficient stoichiometric stereocontrolling group.

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Supporting Information Available: Experimental procedures and characterization data. This material is available free of charge via the Internet at http://pubs.acs.org.

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