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## Enantioselective Organocatalytic Amine Conjugate Addition

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The prevalence of amines in pharmaceutical agents and natural products places C-N stereogenicity among the most important synthon targets for enantioselective reaction development.<sup>1</sup> While the 1,4-addition of stoichiometric chiral amines to electron-deficient olefins has long been established as a principal strategy for C-N bond construction,<sup>2</sup> it is surprising that corresponding catalytic variants have only recently been realized using Lewis acids with  $\alpha,\beta$ -unsaturated ketones,<sup>3a,b</sup> imides,<sup>3b,c</sup> and amides.<sup>3d</sup> Moreover, the enantioselective conjugate amination of  $\alpha,\beta$ -unsaturated aldehydes has remained elusive, a notable deficiency given the broad utility of  $\beta$ -amino aldehydes. Over the last six years, our laboratory has demonstrated that the reversible formation of iminium ions from chiral amine catalysts and  $\alpha,\beta$ -unsaturated aldehydes is a useful platform for conjugate addition reactions involving carbon<sup>4a-c</sup> and hydrido<sup>4d</sup> nucleophiles. In this communication, we further advance this activation concept to describe the first enantioselective organocatalytic amine conjugate addition, a highly chemo- and stereoselective transform that is founded upon a rationally designed N-centered nucleophile. This operationally simple protocol allows rapid and predictable access to enantioenriched  $\beta$ -amino aldehydes and  $\beta$ -amino acids using an inexpensive amine catalyst.

## Enantioselective Organocatalytic & Amino Aldehyde Synthesis



Iminium Catalyzed Amination Requires Selective Amine Partition



## Amine A Design: Carbamate Nucleophilicity Enhanced by a-Effect



**Design Plan.** From the outset, we recognized that a number of chemoselectivity issues must be addressed if an enantioselective conjugate amination were to be realized using an iminium catalysis platform. First, an amine **A** must be identified that will selectively function as a 1,4-addition nucleophile (TS-1), yet will not participate in iminium activation (TS-2, a racemic pathway). Moreover, a second amine **B** must be found that will perform as an iminium catalyst (TS-1) while avoiding a nucleophilic role (a pathway that would lead to catalyst consumption). Third, to ensure that the reaction can proceed under asymmetric (kinetic) control, the

Me	o BnO	O M OTBS	20 mol% cat		,otbs
entry	catalyst	solvent	temp (°C)	% conversion <sup>a</sup>	% ee <sup>b</sup>
1	L-proline	CH <sub>2</sub> Cl <sub>2</sub>	0	0	
2	1•TFA	CH <sub>2</sub> Cl <sub>2</sub>	-20	25	3
3	2•TFA	CH <sub>2</sub> Cl <sub>2</sub>	-20	29	81
4	2•HC1	$CH_2Cl_2$	-20	32	69
5	2•pTSA	$CH_2Cl_2$	-20	70	87
6	2•pTSA	EtOAc	-20	12	92
7	2•pTSA	THF	-20	45	92
8	2•pTSA	CHCl <sub>3</sub>	-20	95	92

<sup>*a*</sup> Conversion determined by <sup>1</sup>H NMR analysis. <sup>*b*</sup> Enantiomeric excess determined by chiral HPLC analysis on the corresponding amino alcohol.

stereodefining heteroatom addition step must be accompanied by *irreversible* loss of the nucleophile's proton. More specifically, this deprotonation step removes the possibility of an equilibrium-controlled process involving a reversible N-addition step (a thermodynamic pathway that would demand the formation of racemic products). In consideration of these requirements, we selected *N*-silyloxycarbamates as the nucleophilic component **A** on the basis that the N–O functionality would enhance nucleophilicity at the nitrogen center via the  $\alpha$ -effect,<sup>5,6</sup> while the carbamate functionality would render the amino aldehyde product effectively nonbasic (silyloxycarbamates N–H, estimated p $K_a \sim 9.0$ ).<sup>7</sup> With respect to the catalyst component **B**, we focused upon the use of imidazolidinone amines given their established capacity<sup>4</sup> to participate in asymmetric iminium activation with enals and enones while selectively avoiding heteroconjugate addition.



To our great delight, exposure of crotonaldehyde to benzyl *tert*butyldimethylsilyloxycarbamate in the presence of imidazolidinone catalyst **2**·TFA did indeed provide the desired  $\beta$ -amino aldehyde product with good levels of enantiocontrol (Table 1, entry 3, 81% ee). Further evaluation of a variety of catalyst salts (entries 3–5) revealed that imidazolidinone **2**·*p*TSA exhibited superior conversion and selectivity. Last, a survey of reaction media (entries 5–8) demonstrated that CHCl<sub>3</sub> was the optimal solvent (entry 8, 92% ee). The superior levels of induction and reaction efficiency exhibited by catalyst **2**·*p*TSA in CHCl<sub>3</sub> to provide the amino aldehyde **3** in 92% ee and 92% yield prompted us to select these conditions for further reaction exploration (Table 2, entry 1).

The scope of the  $\alpha,\beta$ -unsaturated aldehyde component in this enantioselective heteroconjugate addition has been examined. As

Table 2.	Scope	of	Enantioselective	Organocatalytic	Conjugate
Aminatior	า่				

	⇒ → PG、	OTBS	20 mol% <b>2•</b> <i>p</i> TSA	PG OTBS	
R´	$\checkmark$ $\checkmark$ <sub>0</sub>	N H	–20 °C, CHCl <sub>3</sub>	R	$\sim$
entry	R <sup>a</sup>	PG	product	% yield	% $ee^{b,c}$
1	Me	Cbz	Cbz OTBS	92	92
2	<i>n</i> -Pr	Cbz	Cbz NOTBS	77	95
3	<i>n</i> -Pr	Boc	Boc Nr OTBS	85 <sup>d</sup>	92
4	<i>n</i> -Pr	Fmoc	Fmoc OTBS	78	89
5	My y	Cbz	Cbz NOTBS	87	96
6	PhCH <sub>2</sub> CH <sub>2</sub> -	Cbz	$Ph$ $V_2$ OTBS $0$	69	90
7	BnOCH <sub>2</sub> -	Cbz	Cbz NOTBS	70	96
8	N N N	Boc	Boc OTBS	85 <sup>d</sup>	87
9	O CO <sub>2</sub> Me	Boc		78 <sup>d</sup>	97

<sup>a</sup> Performed with 3 equiv of enal. <sup>b</sup> Enantioselectivity determined by HPLC or SFC analysis. <sup>c</sup> Stereochemistry assigned by chemical correlation or by analogy. d Performed with catalyst 2. TFA.

highlighted in Table 2, enal substituents, including alkyl, alkenyl, aryl, ether, amine, and ester groups, are readily tolerated (Table 2, entries 2-9, 87-97% ee).8 Moreover, variation of the carbamate protecting group from Cbz to Boc to Fmoc can be realized without loss in enantiocontrol (entries 2-4, 95, 92, and 89% ee). Variation of the silvloxy protecting group is also possible as illustrated by the TBDPS derivative (entry 9, 97% ee).9 In accord with our design plan, it is important to note that  $\beta$ -amino aldehyde products arising from 1,4-catalyst incorporation were not detected in this study.

A demonstration of the utility of this organocatalytic amine addition and the accompanying products is presented in the onepot (two-step) conversion of simple aldehydes to enantioenriched  $\beta$ -amino acids. As revealed in eq 1, exposure of 2-hexenal to our asymmetric amination conditions followed by in situ Pinnick oxidation provided the corresponding  $\beta$ -amino acid 4 with excellent enantioselectivity (92% ee). Notably, N-O bond removal can be accomplished under mildly reducing conditions (Zn/AcOH).<sup>10</sup>

Alternatively, the amino-oxy moiety can be strategically exploited to generate 1,3-amino alcohols with valuable levels of absolute and relative stereocontrol (Scheme 1). In this case, hexenal amination

Enantioselective Two-Step Synthesis of B-Amino Acids



was followed by in situ Wittig homologation to afford the unsaturated ester 5 in a single operation (71% yield, 92% ee). Subsequent exposure of amino enoate 5 to fluoride ion then enabled both silyl group removal and intramolecular oxy-Michael addition to afford isoxazolidine 6 with excellent diastereocontrol (99% yield, 10:1 syn/anti). Reduction of the N-O bond was accomplished with SmI<sub>2</sub> to afford the 1,3-amino alcohol 7 in excellent yield (99% yield, 70% yield over three steps).<sup>11</sup>

Scheme 1. Enantioselective Synthesis of 1,3-Amino Alcoh
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Supporting Information Available: Experimental procedures, structural proofs, and spectral data for all new compounds are provided (9 pages, print/PDF). This material is available free of charge via the Internet at http://pubs.acs.org.

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