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Isothiourea-catalyzed formal enantioselective conjugate addition of benzophenone imines to β -fluorinated α , β -unsaturated esters[†]

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The isothiourea-catalyzed formal enantioselective conjugate addition of 2-hydroxybenzophenone imine derivatives to α , β -unsaturated *para*nitrophenyl esters has been developed. Investigations of the scope and limitations of this procedure showed that β -electron withdrawing substituents within the α , β -unsaturated ester component are required for good product yield, giving rise to a range of β -imino ester and amide derivatives in moderate to good isolated yields with excellent enantioselectivity (20 examples, up to 81% yield and 97:3 er).

The development of methods for the enantioselective synthesis of β -amino acid derivatives^{1a} is of widespread importance due to the prevalence of this structural motif in natural products and medicinally relevant compounds.¹ Among the synthetic methods that have been developed for the preparation of β -amino acid derivatives, arguably the most simple and elegant involves the asymmetric conjugate addition of an ammonia equivalent to an α , β -unsaturated carbonyl motif. As an example of this approach, the conjugate addition of enantiomerically pure lithium amide derivatives to α , β -unsaturated esters has been developed and exploited extensively by Davies and coworkers. Conjugate addition of lithium N-benzyl-N-amethylbenzylamide to an α , β -unsaturated ester gives the corresponding β -amino ester with high diastereoselectivity (>95:5 dr), with N-deprotection through hydrogenolysis giving the corresponding β -amino ester derivatives (Scheme 1a).²

Over the last two decades, several enantioselective organocatalytic approaches to amine conjugate addition have been introduced. To date, these successful approaches rely upon enals,³ enones,⁴ *N*-acyl pyrazoles,⁵ and nitro-olefins⁶ as Michael acceptors, with the use of bifunctional thiourea^{4a,5b,7,8a-c,e} or squaramide^{4,5c,8a,b,e} organocatalysts, or Lewis basic pyrrolidines^{3,8} commonplace. Catalytic enantioselective amine conjugate additions to α , β -unsaturated esters are rare, reflecting the recognized



(a) Asymmetric synthesis of β -amino esters by lithium amide addition

recalcitrance of α,β -unsaturated esters as Michael acceptors (Scheme 1b). To date, the current state-of-the-art organocatalytic approach is represented by Seidel and co-workers'⁹ demonstration of the conjugate addition of cyclic secondary amines to β -alkyl- α,β -unsaturated benzyl esters using a selenourea-thiourea catalyst **1** (Scheme 1c). Although limited to β -alkyl substituted Michael acceptors, this impressive methodology was applicable to a range of cyclic amines and the kinetic resolution of (\pm)-cyclic 2-arylamines.

Our approach to enantioselective amine conjugate addition focused upon the use of imines as nucleophiles. The conjugate addition of (diphenylmethylene)amine to α , β -unsaturated esters, nitriles and ketones in racemic form has been demonstrated by de Meijere *et al.* MeOH was optimal as a solvent and

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Scheme 2 Previous imine conjugate additions and this work.

a basic additive (such as NEt₃) led to effective product formation (Scheme 2a).¹⁰ In 2018, Alemán and co-workers successfully demonstrated an enantioselective aza-Michael addition of nucleophilic imines to enals using secondary amine catalyst 2 (Scheme 2b).¹¹ Trapping of the resultant β -imino aldehydes with a phosphorane gave the corresponding δ -imino esters in good yield and enantioselectivity. Notably, 2-hydroxybenzophenone imines showed increased reactivity and enantioselectivity compared with the parent benzophenone imine, attributed to an increase in acidity of the imine proton caused by intramolecular hydrogen bonding.^{12,13} In previous work, we and others have demonstrated a range of enantioselective Michael-addition processes of *in situ* generated α,β -unsaturated acyl ammonium species.^{14,15} Building on these precedents, we report herein the formal isothiourea-catalyzed enantioselective addition of 2-hydroxybenzophenone imines to β -fluorinated α,β -unsaturated paranitrophenyl esters via an α , β -unsaturated acyl ammonium intermediate, giving products in up to 98:2 er (Scheme 2c).

Preliminary investigations used β -CF₃-substituted α , β unsaturated *para*-nitrophenyl ester 4 (1.0 equiv.) in toluene as standard. Given the moderate reactivity of α , β -unsaturated acyl ammonium ions, imine 3 (2.0 equiv.) bearing an electron donor 4-OMe-substituent was postulated to enhance nucleophilicity (Table 1). Attempted isolation of the *para*-nitrophenyl ester product led to low and irreproducible product yields, so addition of pyrrolidine to give the isolable amide 5 was adopted. Screening of isothiourea catalysts 6–8 (10 mol%) at 1:2 substrate ratio of ester 4: imine 3 (entries 1–3) showed that tetramisole 6 and BTM 7 gave promising product yield (~50%) whereas HyperBTM 8 showed poor catalytic activity (<10% yield). Excellent enantioselectivity (96:4 er) was observed using BTM 7. Altering the
 Table 1
 Reaction optimisation



	$\begin{array}{c ccccc} OMe & & OMe \\ 3 & & O & (i). \ 6-8 \ (2.5-20 \ mol\%) \\ Solvent \ (0.1 \ M) \\ 6-48 \ h, \ rt \ to \ 60 \ ^{\circ}C \\ + & H & (ii). \ Pyrrolidine \\ F_{3}C & 4 & OAr & Ar = 4-NO_{2}C_{6}H_{4} \\ \end{array} \begin{array}{c} OMe \\ OMe \\ H \\ OH \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ OH \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ OH \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ OH \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ OH \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ OH \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\ H \\ Solvent \ (0.1 \ M) \\ F_{3}C \\$					>
	Ph= $\langle N \\ N \\ S \rangle$ -Tetramisole 6	Ph		ⁱ Pr ^{,,,} , N Ph ^{,,,,} N S (2 <i>S</i> ,3 <i>R</i>)-HyperBTM 8		
Ent	ry Catalyst (mol%)	Temp. (°C)	Solvent	3:4	Yield ^a (%)	er ^b
$\frac{1^{c}}{2^{c}}$	6 (10) 7 (10)	rt rt	Toluene Toluene	1:2 1:2	50 54	12:88 96:4
3^{c}	8 (10)	rt	Toluene	1:2 1:2	54 <10	68:32
4^c	7(10)	rt	Toluene	1:1.5	42	95:5
5^c	7 (10)	rt	Toluene	1.5:1	38	97:3
6 ^{<i>c</i>}	7 (10)	40	Toluene	1:2	52	94:6
7 ^c	7 (10)	60	Toluene	1:2	47	91:9
8 ^c	7 (2.5)	rt	Toluene	1:2	<10	91:9
9 ^c	7 (5.0)	rt	Toluene	1:2	18	96:4
10^{c}	7 (20)	rt	Toluene	1:2	71^d	96:4
11^c	7 (20)	rt	THF	1:2	31	96:4
12^c	7 (20)	rt	Et_2O	1:2	30	96:4
13 ^c	7 (20)	rt	CH_2Cl_2	1:2	37	96:4
14^e	7 (20)	rt	Toluene	1:2	31	98:2
15^{f}	7 (20)	rt	Toluene	1:2	42	96:4
16^g	7 (20)	rt	Toluene	1:2	36	96:4

^{*a*} Using ¹H NMR spectroscopic analysis and 1,3,5-trimethoxybenzene as internal standard. ^{*b*} Ratio of (*R*): (*S*) enantiomers determined by HPLC analysis on a chiral stationary phase. ^{*c*} Ar = 4-NO₂C₆H₄. ^{*d*} Isolated yield. ^{*e*} Ar = 2,4,6-Cl₃C₆H₂. ^{*f*} Ar = C₆F₅. ^{*g*} Ar = 3,5-(CF₃)₂C₆H₃.

reaction stoichiometry (entries 4 and 5) led to reduced product yield. A detrimental effect on product enantioselectivity (91:9 er) was observed when the reaction temperature was increased to 40 °C or 60 °C (entries 6 and 7). Lowering the catalyst loading showed a significant decrease in product yield and enantioselectivity (entries 8 and 9), while using 20 mol% BTM 7 gave increased yield (71% yield, 96:4 er, entry 10). Screening of a alternative solvents gave high product enantioselectivity but reduced yields (entries 11–13). Further optimisation probed the effectiveness of alternative electron-deficient aryl esters. Comparison of *para*-nitrophenyl with 2,4,6-trichlorophenyl, pentafluorophenyl, and 3,5-bis(trifluoromethyl)phenyl esters (entries 14–16) showed that excellent enantioselectivities were observed in each case (up to 98:2 er), with the *para*-nitrophenyl ester leading to the best product yield (71%).

The scope and limitations of the developed process was explored through variation of the nucleophilic imine reaction component (Fig. 1). Variation of the electronic bias of the 4-aryl substituent within the imine component showed that decreased product yield was observed upon changing from an electron-donating 4-MeO- (5, 70% yield) to 4-Me (9, 49% yield), 4-H (11, 36% yield) and electron-withdrawing 4-Br substituent (10, 24% yield) all with >96:4 er. This is consistent with increasing



Fig. 1 0.10 mmol scale. Isolated product yield; er determined by HPLC analysis on a chiral stationary phase; [a] 40 °C for step i; [b] DMAP 20 mol% in step ii.

electron density within the imine component leading to increased product yield. Interestingly, comparing the yield and er of products **11** and **12** indicates that the 2-hydroxy-substituent within the imine is essential for high product er, but does not affect product yield. The incorporation of an additional electron-donating 4-MeO substituent led to product **13** in reduced yield but maintained high product er. Variation of the β -substituent within the α , β -unsaturated ester indicated that the incorporation of polyhalogenated or ester electron-withdrawing groups was necessary for reactivity as alkyl, aryl, ketone and amide substituted acceptors gave no significant product formation. For example, the introduction of halogenated (CF₂H) and polyhalogenated



Scheme 3 Gram scale synthesis of product 5.

substituents (CF₂Cl, CF₂Br, and C₂F₅) led to products **14–17** in up to excellent yields with high enantioselectivity (40% to 81%; >96:4 er), while the incorporation of ester substituents gave **18–19** in poor 20% product yield in up to 96:4 er. Variation of the post catalysis nucleophilic component (Nuc-H) to incorporate alcohols as well as cyclic secondary and acyclic primary amines gave a range of ester and amide products **20–24** in good yield (42% to 64%) and excellent enantioselectivity (\geq 96:4 er).

To further demonstrate the synthetic utility of this transformation, it was applied to the gram-scale synthesis of product 5 with consistent yield and enantioselectivity (67%, 96:4 er, Scheme 3). Hydrolysis gave the free β -amino amide product **26** in high yield and enantioselectivity (95%, 96:4 er).¹⁶

A proposed mechanism of this transformation is shown in Scheme 4. Reversible acylation of the isothiourea with the α , β -unsaturated ester **1a** generates the key α , β -unsaturated acyl isothiouronium ion pair **26**.



Scheme 4 Proposed reaction mechanism.

An intramolecular chalcogen 1,5-S···O interaction $(n_O \rightarrow \sigma^*_{S-C})^{17}$ provides a plausible stabilising effect and conformational lock. Hydrogen bonding between the 2-hydroxy-substituent and the imine N serves to conformationally restrict this functionality and facilitate deprotonation.^{11–13} Subsequent conjugate addition to the s-*cis* conformation of the α,β -unsaturated acyl isothiouronium **26** *anti*- to the stereodirecting phenyl substituent of the isothiourea catalyst generates the ammonium enolate intermediate **27**. Proton transfer generates the β -imino acyl isothiouronium intermediate **28**, with catalyst turnover facilitated by the aryloxide counterion to form the product and release the isothiourea catalyst BTM 7.¹⁸

In summary, enantioselective organocatalytic conjugate addition of 2-hydroxybenzophenone imines to α , β -unsaturated esters using the isothiourea BTM as an organocatalyst gives enantioenriched β -imino amides in modest to good yield (20–81%) and excellent enantioselectivity (typically >95:5 er).¹⁹

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Conflicts of interest

There are no conflicts of interests to declare.

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