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**PAPER** 

# Readily available hydrogen bond catalysts for the asymmetric transfer hydrogenation of nitroolefins†

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This paper focuses on readily accessible thiourea hydrogen bond catalysts derived from amino acids, whose steric and electronic features are modulated by their degree of substitution at the carbinol carbon center. These catalysts were applied in the asymmetric transfer hydrogenation of nitroolefins furnishing the chiral products in up to 99% yield and 86% enantiomeric excess. The proposed catalyst's mode of action is supported by mechanistic investigations.

#### Introduction

The hydrogenation of nitroolefins and nitro acrylates yields chiral nitro compounds, thus offering an efficient access to chiral β-amines and β<sup>2</sup>-amino acid derivatives. Chiral amines are extensively used as chiral building blocks, as resolving agents or as chiral auxiliaries.1 Amino acid derivatives play an important role in the generation of artificial peptide structures. This multifaceted application of amines makes the synthesis of enantiopure nitrogen-containing compounds an attractive field of research. A biomimetic access to enantiopure nitro-compounds by transfer hydrogenation of nitroolefins<sup>2</sup> and nitroacrylates<sup>3</sup> with Hantzsch's esters4,5 was first described by List et al. The applied monofunctional cyclohexyl-diamine-derived thiourea catalysts<sup>6</sup> furnished the hydrogenated products in excellent yields and enantioselectivities. Apart from those Jacobsen-type catalysts, amino alcohol related thiourea derivatives<sup>7</sup> have been successfully employed in the Morita-Baylis-Hilmann reaction,8 in the conjugate addition, 9,10 in the Diels-Alder reaction 11 and in the Friedel-Crafts-alkylation of indole with nitroolefins. 12 In the latter report Ricci et al. discussed the interaction of a highly polarized indole-NH with an alcohol group as being essential for activity and enantioselectivity.12

Our report introduces readily available amino-alcohol-derived thiourea structures as new catalysts for the asymmetric hydrogen bond mediated transfer hydrogenation of nitroolefins. Our investigations show that the dihydropyridine-NH-bond (Hantzsch's ester) significantly contributes to the efficiency and enantioselectivity of the asymmetric hydrogen transfer promoted by hydroxy substituted thiourea derivatives.

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## **Results and Discussion**

The bifunctional<sup>13</sup> thiourea structures<sup>14</sup> were obtained from the reaction of commercially available and readily accessible amino alcohols with 3,5-(trifluormethylphenyl)thioisocyanate (Fig. 1, see ESI† for synthetic procedures and full characterization). The synthesized bifunctional thiourea derivatives (Fig. 1) differ in the degree of substitution at the carbinol carbon atom (1a-1p) and can thus be divided into three classes: structures bearing a tertiary, secondary or primary alcohol functionality. This feature allows studying the catalyst's performance depending on the degree of substitution at the carbinol functionality. The results of the hydrogen bond mediated asymmetric transfer hydrogenation of (E)-2-phenyl-1-nitro-propene (2a) with tert-butyl-Hantzsch's ester (3) are summarized in Table 1.

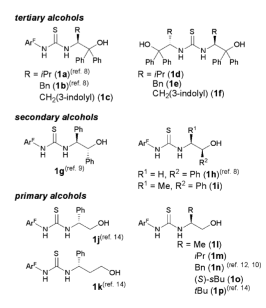


Fig. 1 Bifunctional thiourea catalysts derived from amino alcohols (Ar<sup>F</sup> =  $3,5-(CF_3)_2-C_6H_3$ ).

**Table 1** Asymmetric transfer hydrogenation of 2

Entry	Catalyst	Yield (%) <sup>a</sup>	ee (%) <sup>b</sup>
1 <sup>c</sup>	_	5	n.d.
2	1a	81	$<5(S)^{d}$
3	1b	72	13 (S)
4	1c	26	<5(R)
5	1d	46	11 (R)
6	1e	39	<5(S)
7	1f	31	16 (R)
8	1g	70	20(R)
9	1ĥ	90	40 (S)
10	1i	88	62(R)
11	1j	99	61 (S)
12	1k	78	50 (S)
13	11	99	61 (R)
14	1m	91	67 (R)
15	1n	99	60 (R)
16	1o	94	67 (R)
17	1p	99	70(R)
18	epi-1p	99	65 (S)

<sup>a</sup> Yields were determined by GC with dodecane as the internal standard. <sup>b</sup> The enantiomeric excess was determined after purification on silica gel by HPLC with chiral stationary phase. <sup>c</sup> Reaction was performed at 40 °C in toluene; n.d. not determined.  $\bar{d}$  The absolute configuration was determined by comparison with literature values (see ESI†).

Generally, the reaction is catalyzed by hydroxy-functionalized thiourea derivatives. Catalysts bearing tertiary alcohol groups furnished the nitro alkane 4a in good yields (entries 2 and 3), while the tryptophan-derived catalyst (1c, entry 4) was not sufficiently active. The C<sub>2</sub>-symmetric catalysts **1d–1f** (entries 5–7) were generally less active than the corresponding unsymmetrical thiourea derivatives. Although the catalysts 1a and 1b displayed good activity, the enantioselectivity achieved was low (<20% ee). Catalysts featuring a secondary alcohol function were more active and furnished 4a in 70–90% yield (entries 8–10). Diverging activities and enantioselectivities were observed for 1g and 1i (entries 8 and 10, 70% yield, 20% ee and 88% yield, 62% ee) resulting from the relative configuration of the 1,2-amino alcohol fragment. This may be rationalized by assuming that different conformers in 1g and 1i are present in solution. The conformations of ephedrine- and 1,2-diphenyl-2-amino ethanol derivatives have been studied in solid state<sup>15</sup> and in solution by NMR.<sup>16</sup> In most cases the gauche conformation, featuring an internal hydrogen bond, has been identified by coupling-constant analysis of the vicinal protons as the most abundant conformer. However, the catalyst systems 1g-i proved too dynamic on the NMR timescale for coupling-constant analysis. The 2-amino-2-phenyl ethanol substructure in catalyst 1j is isomeric to structure 1h. The catalyst 1j features a primary hydroxy functionality and furnished the hydrogenation product 4a in high yield (entry 11, 99% yield). The enantioselectivity is improved to 61% ee compared to catalyst 1h (entry 9). This indicates that the transfer of stereo information (from the catalyst to the product) is more efficient from the chiral vicinity of the thiourea moiety than from the asymmetric

carbinol functionality (entries 9–11). The ethylene spacer group in the catalyst's structure proved to be optimal for transfer hydrogenation. The 2-amino-2-phenyl propanol derived catalyst 1k provided the product in 78% yield and 50% ee (entry 12). For this reason we investigated the efficiency of the thiourea derivatives depending on the bulkiness of the alkyl chain at the stereogenic carbon center (entries 13-17). Surprisingly, the smallest alkyl group, such as a methyl group (11), already generated a highly efficient catalyst (entry 13, 99% yield and 61% ee). Moving along in the series of iPr (1m), Bn (1n), sBu (1o) and tBu (1p) (entry 14–17) the enantioselectivity of the catalyzed process gradually improved to 70% ee while the excellent activity of the catalyst was preserved (91-99%). When the reaction was prefromed in diethylether 4a was obtained with 77% ee but with unsatisfactory yield (55% see ESI† for further optimization studies). 17 To demonstrate that both enantiomers of the saturated product 4a are accessible by this methodology, we conducted the reaction with the catalyst's enantiomer epi-1p. The hydrogenation of 2a furnished the Sconfigured product in quantitative yield with 65% enantiomeric excess (entry 18).

We then turned our interest to the substrate scope for the thiourea-catalyzed transfer hydrogenation of unsaturated nitro compounds using our best performing catalyst 1p (Table 2). All substrates were converted to the saturated nitro compounds in good to excellent yields (84–99%). However, the enantioselectivity of these transformations differed significantly depending on the substitution pattern. Methyl-aryl-substituted nitroolefins were obtained with an enantiomeric excess of 50–70% (entries 1–6). Electron withdrawing as well as electron donatin groups on the aryl ring are tolerated and the products were furnished with 50-67% ee. Higher enantioselectivities up to 87% ee were observed for nitro alkenes bearing bulkier substituents (entry 8, tBu). Additionally, the thiourea 1p showed excellent activity in the reduction of nitro acrylates (Table 3). The products 6a-6c were formed in excellent yield. Surprisingly, the methyl (6a), ethyl (6b) and ipropyl (6c) esters were obtained with almost the same enantiomeric excess (54–60% ee) revealing an unique opportunity to synthesize a wide variety of ester derivatives.

In order to comprehend the varied performance of the synthesized catalysts a hydrogen-bonded structure can be conceived. The formation of a ternary complex incorporating the thiourea derivative (1p), the nitroolefin (4a) and the reducing agent 3 can serve as a rationalization for the observed activities and enantioselectivities. Literature precedent supports the assumption that hydroxyl groups are beneficial for organocatalytic transformations where polarized N-H bonds are involved. 2c,11,12 The formation of a N-H · · · O hydrogen bond between a heterocycle and the catalyst has already been proposed and investigated for the Friedel-Crafts alkylation of indole.12 We decided to study the role of the hydroxy group of our catalyst in the asymmetric transfer hydrogenation by selective suppression of important substrate-catalyst interactions. For this purpose we identified two significant interactions apart from the nitro-thiourea-interaction: 1) hydrogen-bonding between the nitro- and the hydroxy group (N=O··· H-O); 2) hydrogenbonding between the Hantzsch's ester and the hydroxy-group (N- $H \cdots O$ ) (Scheme 1).

1) To suppress the simultaneous binding of the nitro-compound to the thiourea and the hydroxy group the silylated thiourea 7 was synthesized. In this structure, the postulated coordination of the

**Table 2** Substrate scope for the catalyzed transfer hydrogenation of nitroolefins

Entry	Nitroolefin	2	Product 4	Yield (%) <sup>a</sup>	ee (%) <sup>b</sup>
	Me NO <sub>2</sub>		Me NO <sub>2</sub>		
1 2 3 4 5 6	R = H $R = Me$ $R = OMe$ $R = Cl$ $R = F$ $R = CN$	(2a) (2b) (2c) (2d) (2e) (2f)	4a 4b 4c 4d 4e 4f	99 99 99 97 95 88	70 50 62 67 63 56
	NO <sub>2</sub>		$R$ $NO_2$		
7 8	R = Et $R = tBu$	(2g) (2h)	4g 4h	95 76	68 87
9	Me NO <sub>2</sub>		$\bigvee^{Me}NO_2$	84°	40
		(2i)	<b>(4i)</b>		

<sup>&</sup>lt;sup>a</sup> After purification by column chromatography. <sup>b</sup> The enantiomeric excess was determined by HPLC with chiral stationary phase. <sup>c</sup> The yield was determined by GC with dodecane as the internal standard.

 Table 3
 Asymmetric reduction of nitroacrylates

	CO <sub>2</sub> R Ph NO <sub>2</sub>	3 1p (20 mol%), DCE, 0 °C, 3d, (1M)	CO <sub>2</sub> R NO <sub>2</sub>	
Entry	Nitroacrylate	Product	Yield (%) <sup>a</sup>	ee (%) <sup>b</sup>
1 2 3	R = Me (5a) $R = Et (5b)$ $R = iPr (5c)$	6a 6b 6c	95 99 93	60 58 54

<sup>&</sup>lt;sup>a</sup> after purification by column chromatograpy; <sup>b</sup> the enantiomeric excess was determined by HPLC with chiral stationary phase.

nitro-compound by both the thiourea- and the hydroxy-group is not possible, while the interaction of the Hantzsch's ester (3) to the Lewis-basic oxygen atom is still feasible. The application of 7 in the transfer hydrogenation of 2a furnished the product 4a in lower yield and in slightly diminished enantioselectivity (52% yield, 51% ee; Scheme 1a). This may be explained by the lack additional hydrogen bonds between the hydroxy- and nitro-functionality. Additionally, the free hydroxy group in 1p might facilitate the proton transfer to the formed nitronate species from the conjugate hydride attack to the nitroolefin. To probe the interaction of the nitro group with the unmasked hydroxy group in 1p we conducted <sup>1</sup>H NMR experiments. However, the addition of 0.5, 1.0, 5.0 and 10 equiv. of nitroolefin 2a to a 0.2 M solution of catalyst 1p did not result in significant changes in the <sup>1</sup>H NMR-spectrum (see ESI†).

2) The significance of the Lewis-basic oxygen atom in the proposed N-H···O-interaction was demonstrated by two experiments: by the complete removal of the hydroxy functionality and by masking the NH-group of the Hantzsch's ester  $(3 \rightarrow 9)$ with a methyl group (Scheme 1b and 1c). The defunctionalized thiourea-derivative 8 afforded the saturated nitro-compound 4a as racemic material in DCE and Et<sub>2</sub>O (Scheme 1b). Consequently, the complete removal of the Lewis-basic oxygen atom from the catalysts structure generated an unselective catalyst. This underlines the importance of the hydroxy-functionality tethered to the catalyst for an enantioselective reaction. Use of the protected Hantzsch's ester 919 together with our best catalyst 1p and one equivalent of ethanol as a proton source<sup>20</sup> also furnished the product 4a as racemic material (Scheme 1c, 30% yield). The control experiment using 3 and one equiv. of ethanol as proton source supplied 4a in 90% yield with significant enantioenrichment (Scheme 1d, 50% ee). Hence, the constructive interaction between the catalyst and the Hantzsch's ester is essential for the enantioselective hydrogen transfer. This conclusion is supported by the fact, that racemic 4a was obtained when DMSO was applied as cosolvent (DMSO/DCE, 1:1; see ESI†) under best reaction conditions. Concluding this experimental data set, a ternary complex consisting of the catalyst (1p), the nitroolefin (2a) and the Hantzsch's ester (3) as transient species according to Fig. 2 seems instructive.

The catalyst is able to coordinate the nitroolefin in two different ways: one in which the hydroxy group is located on the *Re*-face of the nitroolefin (Fig. 2 left) and one in which the hydroxy group is

Scheme 1 Mechanistic investigations for the enantioselective transfer hydrogenation catalyzed by 1p.

Fig. 2 Postulated ternary substrate-catalyst complex leading to the stereoselective hydride transfer.

located on the *Si*-face (Fig. 2 right). This results in the selective delivery of the hydride to the enantiotopic faces, hence determining the stereoselectivity of the reduction.<sup>21</sup> The coordination of the nitroolefin for the *Re*-face attack is less stabilized by encountering steric clashes between the methyl group and the catalyst (Fig. 1 left). On the contrary, the coordination of the nitroolefin for the *Si*-face attack is favored because now the vinylic hydrogen atom in **2a** is oriented towards the catalyst's backbone resulting in lower steric interactions. This model supports our experimental observations of high selectivity with bulky substrates, together with the observed stereochemical outcome of the product. Nitroolefin **2h** (*t*Bu-Ph) should coordinate with its *Si*-face pointing towards the Hantzsch's

ester (3) to avoid steric interactions of the *t*Bu-group with the catalyst's side chain, resulting in the hydride delivery to the *Si*-face with 76% yield and 87% enantiomeric excess.

### **Conclusions**

We have disclosed an efficient synthesis and derivatization of chiral bifunctional thiourea catalysts for the asymmetric transfer hydrogenation of nitroolefins. The configurational features of **1i** and **1g** led us to the development of highly active and enantioselective catalyst structures, *e.g.* **1p**. This catalyst displayed a rather wide substrate scope and furnished the products in excellent yields (up to 99%) and moderate to good enantioselectivities (up to 87% ee). The mode of action of the catalyst was probed by selective suppression of three distinct substrate-catalyst interactions. These mechanistic investigations suggest a transient, ternary substrate-catalyst-structure for the described enantioselective transfer hydrogenation of nitroolefins.

### **Experimental section**

# N-((S)-(2-amino-3,3-dimethylbutanol))-N'-(3,5-bis-(trifluoromethyl)phenyl)thiourea (1p)

To a solution of (S)-tert-leucinol (800 mg, 6.83 mmol, 1.0 equiv.) in dry CH<sub>2</sub>Cl<sub>2</sub> (7.0 mL) was added 3,5bis(trifluoromethyl)phenylisothiocyanate (1.37 mL, 7.51 mmol, 1.10 equiv.). After stirring at 40 °C for 3 h, the solvent was removed under reduced pressure. The crude product was purified by column chromatography (SiO<sub>2</sub>, cyclohexane/ethyl acetate 80/20 v/v). The compound was obtained as a white foamy solid (2.49 g, 6.40 mmol, 94%). m.p. 56 °C (capillary);  $R_{\rm f}$  (cyclohexane/ethyl acetate, 80:20) = 0.34;  $[\alpha]_D^{20}$  -75.6 (c 0.5 in chloroform);  $\delta_H$ (400 MHz, acetone-d<sub>6</sub>) 9.50 (br s, 1H, OH), 8.38 (s, 2H, HAr), 7.67 (s, 1H, HAr), 7.58 (d, J = 7.4 Hz, 1H, NH), 4.64–4.55 (m, 1H, CH), 3.94–3.83 (m, 2H, CH<sub>2</sub>, NH), 3.81–3.73 (m, 1H, CH<sub>2</sub>), 1.04 (s, 9H, CH<sub>3</sub>);  $\delta_{\rm C}$  (100 MHz, acetone-d<sub>6</sub>) 184.0 (C=S), 144.1 (CAr), 132.9 (q, J = 33.0 Hz, 2C, CCF<sub>3</sub>), 125.4 (q, J = 271.9 Hz, 2C, CF<sub>3</sub>), 123.9 (2C, CHAr), 118.1 (CHAr), 64.8 (CH), 62.9 (CH<sub>2</sub>), 36.4 (C), 28.6 (3C, CH<sub>3</sub>) ppm;  $\delta_F$  (376 MHz, CDCl<sub>3</sub>) -63.10 (m, 6F, CF<sub>3</sub>); IR (Platinum ATR)  $v_{\text{max}}/\text{cm}^{-1}$  3265 (vw), 2965 (vw), 1532 (w), 1471 (vw), 1379 (w), 1342 (vw), 1273 (m), 1169 (w), 1124 (m), 1042 (w), 996 (vw), 972 (vw), 885 (w), 847 (vw), 700 (w), 680 (w), 571 (vw), 401 (vw); MS (FAB), m/z 389.1 ([M + H]<sup>+</sup>, 100%), 370.1 (20%), 355.1 (10%), 289.1  $([C_9H_7F_6N_2S]^+$ , 15%); HRFABMS calcd for  $C_{15}H_{19}F_6N_2OS$ : 389.1119, found 389.1122 [M + H]<sup>+</sup>; elemental analysis: Found: N 6.97, C 45.97, H 4.51. C<sub>15</sub>H<sub>18</sub>F<sub>6</sub>N<sub>2</sub>OS requires N 7.21, C 46.39, H 4.67.

# Procedure for the asymmetric transfer hydrogenation of nitroolefins and nitroacrylates

A solution of the respective nitroolefin or nitro acrylate (0.3 mmol) in dichloroethane (0.3 mL) was cooled to 0 °C. Subsequently catalyst 1p (0.06 mmol) and tBu-Hantzsch's ester 3 (0.36 mmol) were added successively as a solid. The reaction was stirred at 0 °C for 3 d. The mixture was diluted with pentane/Et<sub>2</sub>O (99:1 v/v, 0.7 mL) and subjected to column chromatography (SiO<sub>2</sub>, pentane/Et<sub>2</sub>O 99:1–98:2 v/v for nitroalkanes and 95/5 v/v for esters).

### (R)-2-(4-Methylphenyl)-1-nitropropane (4a)

 $\delta_{\rm H}$  (250 MHz, CDCl<sub>3</sub>) 7.45–7.23 (m, 5H, HAr), 4.63–4.46 (m, 2H,  $CH_2NO_2$ ), 3.75–3.59 (m, 1H, CH), 1.42 (d, J = 7.0 Hz, 3H,  $CH_3$ );  $\delta_{\rm C}$  (125 MHz, CDCl<sub>3</sub>) 140.8 (C), 129.0 (2C, CHAr), 127.6 (CHAr), 126.9 (2C, CHAr), 81.9 (CH<sub>2</sub>), 38.6 (CH), 18.7 (CH<sub>3</sub>).

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