

Metal-Free Enantioselective Electrophilic Activation of Allenamides: Stereoselective Dearomatization of Indoles**

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Dedicated to Professor Pier Giorgio Cozzi on the occasion of his 50th birthday

Abstract: The effective and unprecedented chiral BINOL phosphoric acid catalyzed (1–10 mol%) dearomatization of indoles through electrophilic activation of allenamides (*ee* up to 94%), is documented. Besides the synthesis of 3,3-disubstituted indolenine cores, a dearomatization/hydrogen transfer cascade sequence is also presented as a new synthetic shortcut toward highly enantiomerically enriched indolines.

The hydrogen bond (HB) activation mode is recognized worldwide as one of the most powerful strategies in the field of metal-free asymmetric manipulation of heteroatom-based organic functional groups and anions.^[1] On the contrary, the use of metal-free noncovalent interactions for the direct activation of carbon-based unsaturated π systems has rarely found solutions so far.

In this segment, a Brønsted acid (BA) assisted intramolecular hydroamination of isolated alkenes was reported by the group of Ackermann (*ee*_{max} = 17%)^[2a] and more recently by Toste and co-workers.^[2b] In the latter case, the efficiency of chiral dithiophosphoric acids in promoting the enantioselective intramolecular nucleophilic addition to 1,2-/1,3-dienes was demonstrated. Additionally, chiral BAs were also reported catalyzing Friedel–Crafts-type alkylations of indoles by the groups of Terada^[3a] and Zhou,^[3b] employing electron-rich alkenes (i.e., enamides and enecarbamates) as precursors of the alkylating agents. Here, classic hydrogen-bond contacts between the chiral promoter and the nitrogen atom of the aldimino tautomer, were invoked in the enantiodiscriminating event of the catalytic process. Finally, Terada and co-workers reported on the suitability of BINOL-based phosphoric acids in delivering the enantioselective addition of azlactones to 3-vinylindoles (i.e., ene-type reaction).^[3c]

To face the current growing demand for stereochemical manipulations of C–C π systems by metal-free activation, we envisaged bifunctional allenamides as a valuable organic platform to access chemical diversity.^[4] This class of “electron-rich” π systems is receiving growing attention in asymmetric synthesis through metal-assisted electrophilic activation.^[5] The coordination of chiral late-transition-metal (LTM) complexes to the allenyl framework “C=C=C” is known to deliver positively charged intermediates **A** that can smoothly undergo condensation with a variety of nucleophilic agents both at the α or γ position (Figure 1a). Amongst others, enantioselective cycloaddition reactions represent one of the most explored stereoselective transformations involving allenamides with the active participation of chiral gold complexes as promoters.^[6]

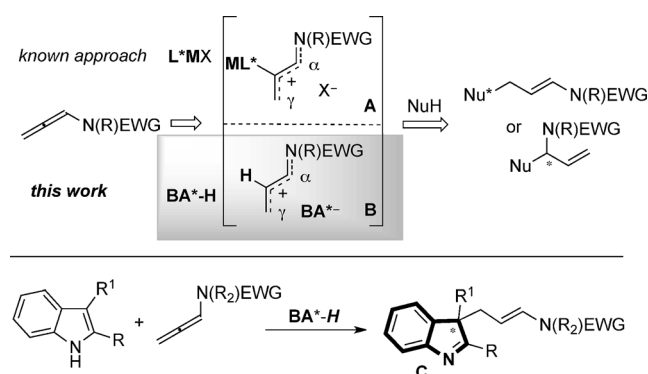


Figure 1. a) Conventional (metal) and “unconventional” (metal-free) electrophilic activation of allenamides in asymmetric synthesis. b) Enantioselective Brønsted acid catalyzed dearomatization of indoles with allenamides: the model reaction.

In this direction, the well-known isolobal analogy often interconnecting $[\text{Au}^I]$ species and the proton,^[7] introduces also chiral σ acids (i.e., Brønsted acids; BAs) as effective promoting agents for the stereochemical manipulation of allenamides. Here, the realization of tight contact ion pairs between the chiral anion and the obtained α,β -unsaturated iminium species **B** (Figure 1a) could provide the ideal stereochemical environment to control the stereochemical pathway of the entire process. However, it should be mentioned that no examples of metal-free electrophilic activation of allenamides in enantioselective transformations have been reported so far.^[8]

To verify the feasibility of this working hypothesis, we considered the challenging C3-site-selective intermolecular

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[**] We acknowledge Progetto FIRB “Futuro in Ricerca” Innovative sustainable synthetic methodologies for C–H activation processes (MIUR, Rome) and the University of Bologna. M.B. thanks Prof. Pier Giorgio Cozzi and Dr. Gianpiero Cera for stimulating scientific discussions.

Supporting information for this article is available on the WWW under <http://dx.doi.org/10.1002/anie.201407518>.

enantioselective dearomatization of *N*(H)-free indoles as the benchmark process.^[9] The realization of this transformation would result in the direct synthesis of densely functionalized enantiomerically enriched 3,3-disubstituted indolenine cores featuring an all-carbon quaternary stereogenic center at the C3 position (**C**, Figure 1b).^[10] This transformation has been a recent focus of our research program.^[11] In particular, we recently reported on the key role of the gold counterion in controlling both reactivity and regioselectivity in the condensation of indoles with allenamides.^[12] These findings inspired and supported the working hypothesis of the present enantioselective metal-free variant.

The condensation of 2,3-dimethylindole (**1a**) and readily available, moisture-tolerant *N*-phenyl-*N*-sulfonylallenamide (**2a**) was selected as the model transformation. A range of chiral BINOL-based phosphate catalysts (**I–VII**, 10 mol %) was screened at room temperature in anhydrous benzene as the reaction media (Table 1).^[13]

Table 1: Optimization of the catalytic system.^[a]

I: X = 3,5-(CF ₃) ₂ C ₆ H ₃ , R = H II: X = SiPh ₃ , R = H III: X = 9-anthracenyl, R = H IV: X = 2,4,6-(iPr) ₃ C ₆ H ₂ , R = H V: X = 2-naphthyl, R = H VI: X = 2,4,6-(Cy) ₃ C ₆ H ₂ , R = H VII: X = 2,4,6-(Cy) ₃ C ₆ H ₂ , R = nC ₈ H ₁₇				
Entry	BA	T [°C]/t [h]	Yield [%] 3aa ^[b]	ee [%] 3aa ^[c]
1	(<i>R</i>)- I	25/16	53	28 (<i>S</i>)
2	(<i>R</i>)- II	25/16	61	8 (<i>S</i>)
3	(<i>R</i>)- III	25/16	92	72 (<i>S</i>)
4	(<i>R</i>)- IV	25/16	75	82 (<i>S</i>)
5	(<i>R</i>)- V	25/16	53	25 (<i>S</i>)
6	(<i>S</i>)- VI	25/16	88	84 (<i>R</i>)
7	(<i>R</i>)- VII	25/16	98	92 (<i>S</i>)
8 ^[d]	(<i>R</i>)- VII	10/16	72	94 (<i>S</i>)
9 ^[e]	(<i>R</i>)- VII	25/72	57	87 (<i>S</i>)
10 ^[f]	(<i>R</i>)- VII	rt/16	95	85 (<i>S</i>)

[a] All reactions were performed in dry benzene under nitrogen conditions (**1a**/**2a**/**BA** = 2:1:0.1, unless otherwise specified). [b] Determined after flash chromatography. [c] Determined with chiral HPLC. Absolute configuration in brackets. [d] At 10 °C. [e] 1 mol % of catalyst was used. [f] With activated 5 Å molecular sieves.

From the data collected in Table 1 several conclusions can be drawn. Firstly, the reaction outcome generally shows a regiochemical preference for the nucleophilic C3 attack by indole on the γ position of the allenamide, with respect to the N1 analogues, even in the presence of nitrogen-free substrate **1a**. Secondly, the screening of differently decorated chiral BINOL phosphoric acids led to important conclusions about the impact of the BA structure on the course of the stereochemical reaction. In particular, (*S*)-C₈-TCyP (**VII**) performed exceptionally in the model reaction, providing the indolenine **3aa** in quantitative yield and 92% *ee* (entry 7).^[14] The enantiomeric excess could also be slightly increased up to

94% by reducing the temperature to 10 °C (entry 8), but no significant changes were recorded in the presence of molecular sieves (entry 10). Last but not least, the efficiency of the present metal-free stereoselective electrophilic activation of allenamides was further emphasized by the synthetically acceptable performance of **VII** when utilized at a loading of 1 mol % (*ee* 87%, entry 9).

The scope of the catalytic methodology was investigated by subjecting a range of 2,3-disubstituted indoles (**1b–p**) to dearomatization under the optimized catalytic conditions (**VII** = 1–10 mol %, benzene, 16/72 h, rt). The chemical and stereochemical outcomes are summarized in Table 2.

Table 2: Scope of the enantioselective protocol.

 3ba ^[a] (Y 75%, <i>ee</i> 91%)	 3ca ^[a] (Y 51%, <i>ee</i> 91%)	 3da ^[a] (Y 44%, <i>ee</i> 88%)
 3ea ^[a] (Y 87%, <i>ee</i> 93%)	 3fa (Y 72%, <i>ee</i> 91%)	 3ga ^[b] (Y 71%, <i>ee</i> 92%)
 3ha (Y 71%, <i>ee</i> 92%)	 3ia ^[c] (Y 64%, <i>ee</i> 91%)	 3ja ^[c] (Y 91%, <i>ee</i> 89%)
 3ka ^[c] (Y 60%, <i>ee</i> 87%)	 3la ^[b] (Y 98%, <i>ee</i> 72%)	 3ma ^[c] (Y 60%, <i>ee</i> 92%)
 3na ^[b] (Y 77%, <i>ee</i> 92%)	 3pa ^[b] (Y 69%, <i>ee</i> 87%)	 3ab ^[c] (Y 82%, <i>ee</i> 94%)

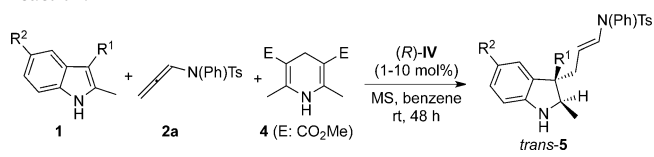
[a] (*R*)-**VII** was used. [b] Catalyst loading 5 mol %, time 72 h. [c] Catalyst loading 1 mol %, time 72 h. Ts = *p*-toluenesulfonyl, Bs = benzenesulfonyl, Y = yield.

Remarkably, all the substrates underwent the dearomatization process with high degrees of stereoselectivity. In particular, indoles carrying cyclic C(2,3)-fused substituents (**1b–d**) performed efficiently, leading to the desired indolenines **3ba–3da** in moderate to good yields and excellent enantiomeric excess up to 93%. Saturated aliphatic as well as benzylic organic residues at the C3-position of the indole (**1e–h**) also provided high stereodifferentiation (*ee* 87–93%). Similar conclusions can be drawn for C2-Et- and C2-*n*Pr-substituted indoles **1n–p**, which provided the corresponding dearomat-

ized compounds in acceptable yields (69–77%) and *ee* (87–92%). Subsequently, the tolerance of the organocatalytic approach toward substituents at the indolyl periphery was verified. A range of differently C5-substituted indoles (**1i–m**) was treated with allenamide **2a** in the presence of **VII**. To our delight, synthetically acceptable levels of stereoselection were achieved with *ee* up to 92% in the case of 5-F and 5-MeO indole derivatives. Finally, variations on the allenamide units were also investigated (see the Supporting Information, SI), highlighting arylsulfonyl groups as the best electron-withdrawing group (EWG; yield 82%, *ee* 94% with 1 mol% of **VII**).^[15]

The extraordinary efficiency of the BINOL-based phosphoric acid **C₈-TCYP (VII)** in the regio- and stereoselective dearomatization of indoles offered the unique opportunity to extend the methodology to the preparation of enantiomerically enriched 3,3-disubstituted indolines^[16] by transfer hydrogenation of the newly formed imine moiety of **3**.^[17] In this direction, we envisioned a Brønsted acid catalyzed one-pot dearomatization/hydrogenation transfer sequence through a three-component reaction of the indole, the allenamide, and a Hantzsch ester (HE) **4**.^[18] Remarkably, commercially available (*R*)-TRIP (**IV**) demonstrated exceptional efficiency in the proposed approach, leading to densely functionalized *trans*-C2/C3 indolines (**5**) in moderate yield (44–72%) and with excellent stereoselection (Table 3). In particular, both

Table 3: Enantioselective dearomatization/hydrogen transfer cascade reaction.^[a]



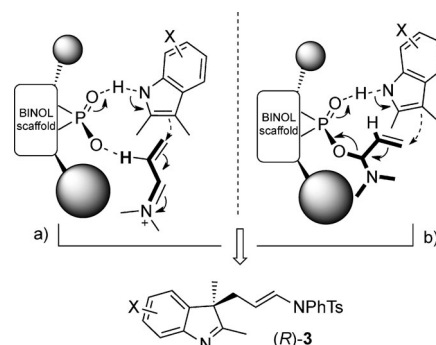
Entry	R ¹ /R ² (1)	Yield [%] ^[b] <i>trans</i> + <i>cis</i>	d.r. ^[c] <i>trans</i> / <i>cis</i>	<i>ee</i> [%] ^[d] <i>trans</i> / <i>cis</i>
1	Me/H (1a)	51	19:1	98/82
2	Et/H (1f)	49	15:1	99/33
3	Me/Cl (1j)	72	11:1	95/70
4	Me/Br (1k)	60	10:1	95/82
5	Me/Me (1l)	44	12:1	94/n.d.

[a] All the reactions were performed under inert conditions using nitrogen gas (**1**:**2a**:HE:BA = 2:1:2:0.1, unless otherwise specified).

[b] Determined after flash chromatography. [c] Determined on the reaction crude. [d] Determined with chiral HPLC. n.d. not determined.

EWGs and electron-donating groups (EDGs) located at the benzenoid ring of the indole were adequately supported by the protocol, leading to very high diastereomeric ratios (d.r. up to 19:1) as well as enantiomeric excesses (*ee* up to 99%).

Mechanistically, the selective protonation of the allenamide **2** at the β position by the **BA-H** would provide the necessary electrophilic activation of the electron-rich π system, delivering a transient α,β-unsaturated iminium intermediate **B** (Figure 1a). Subsequent Michael-type addition with the indole would result in the final product **3** (Scheme 1a). The basic site located at the phosphoryl oxygen atom



Scheme 1. Possible activation modes: noncovalent (a), and covalent (b) BA–allenamide interactions.

could also “regulate” the enantiodiscriminating step of the process by controlling the approaching trajectory of the heterocycle through hydrogen bond interaction (acid/base dual function catalysis).^[19] The bifunctional catalysis could also account for the observed regiospecific C3 versus N1 functionalization of the indole. Differently, a covalent interaction between the catalyst and the π system, resulting from the hydrophosphorylation of the activated allenamide (i.e., an S_N2'-type mechanism, Scheme 1b) cannot be ruled out at the present.^[2b,20]

Both hypothetical approaching trajectories depicted in Scheme 1 justify the overall stereochemistry recorded in the final product that was determined to be *R* by single-crystal X-ray analysis of indolenine **3ka** (see SI).^[21]

In conclusion, an unprecedented electrophilic metal-free activation of allenamides is documented in the enantioselective intermolecular dearomatization of indoles. Chiral BINOL-based Brønsted acids (*S*)-**C₈-TCYP** and commercially available (*R*)-TRIP provide access to a library of 3,3-disubstituted indolines as well as indolenines in enantiomerically enriched form. Attempts to extend the present protocol to different organocatalyzed enantioselective manipulations of allenamides are currently under way in our laboratories.

Received: July 23, 2014

Revised: September 18, 2014

Published online: October 24, 2014

Keywords: Brønsted acids · dearomatization · enantioselectivity · homogeneous catalysis · indoles

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