Title: Cooperative Catalysis with Coupled Chiral Induction in 1,3-Dipolar Cycloadditions of Azomethine Ylides

Authors: Jose Miguel Sansano, Alberto Cayuelas, Olatz Larrañaga, Verónica Selva, Carmen Nájera, Takahiko Akiyama, Abel de Cózar, José I. Miranda, and Fernando P. Cossío

This manuscript has been accepted after peer review and appears as an Accepted Article online prior to editing, proofing, and formal publication of the final Version of Record (VoR). This work is currently citable by using the Digital Object Identifier (DOI) given below. The VoR will be published online in Early View as soon as possible and may be different to this Accepted Article as a result of editing. Readers should obtain the VoR from the journal website shown below when it is published to ensure accuracy of information. The authors are responsible for the content of this Accepted Article.

To be cited as: Chem. Eur. J. 10.1002/chem.201801433

Link to VoR: http://dx.doi.org/10.1002/chem.201801433
Cooperative Catalysis with Coupled Chiral Induction in 1,3-Dipolar Cycloadditions of Azomethine Ylides

Alberto Cayuelas,[a][b] Olaz Larrañaga,[b][c] Verónica Selva,[a][b][c] Carmen Nájera,[a][b][c] Takahiko Akiyama,[e] José M. Sansano,[a][b][c] Abel de Cózar,[b][d][f] José I. Miranda,[g] and Fernando P. Cossio*[b][d]

Dedicated to Prof. Miguel Yus on the occasion of his retirement.

Abstract: 1,3-Dipolar cycloadditions (1,3-DC) between imine esters (as precursors of N-metalated azomethine ylides) and α-deficient alkenes are promoted by cooperative asymmetric Lewis acid-Brunsted base catalysis. The components of these catalytic pairs are silver salts derived from enantiopure commercially available BINOL-based phosphoric acids and Cinchona alkaloids. Chiral phosphoric silver(I) salts promote HOMO raising of in situ formed 1,3-dipoles, whereas protonated cinchona alkaloids generate a LUMO lowering of the dipolarophiles resulting in a global acceleration of the 1,3-DC. The best results were obtained with BINOL-derived silver phosphate and hydroquinoline. Matching between both cooperative metallo- and organocatalyst results in an enhanced enantioselective excess, superior to that reached by both separate components. NOESY experiments and DFT calculations are compatible with a non-covalent interaction (hydrogen bond) between both catalysts, which results in close contacts and mutually coupled chiral environments.

Scientists learn many key synthetic concepts from nature. The simple and sophisticated strategies employed in biocatalysis have been implemented in modern asymmetric synthesis, synergistic/cooperative catalysis 1,2 being one of the most fascinating areas (Figure 1).2 Due to the opposite nature of chiral catalysts involved in these types of processes, they are free to interact to each other and give inactive species. To overcome this drawback it is necessary to select them in such a way that a reversible binding can occur allowing the corresponding substrate activation separately. Concerning the interaction of two chiral entities,3 successful metal catalyst-metal catalyst, metal catalyst-organocatalyst, and organocatalyst-organocatalyst have been reported.3

Within a wider context, Carreira et al.5 firstly introduced the concept of stereodivergent dual catalysis for the enantioselective α-allylation of aldehydes. In that case, related to the (a) strategy of Figure 1, the selection of different combinations of catalysts provided access to all possible stereoisomers of a target compound in high enantiomeric excess. Therefore, this strategy is based on the independent chiral induction generated by both catalysts (Figure 1a). This model contrasts with the clear double chiral activation of a single substrate one (Figure 1b).

This necessary reversible binding, associated with the high stereocore control exhibited, for example, by cycloadditions in their corresponding transition states allows to expand the scope and synthetic utility of them.6,7,8 Particularly, catalytic enantioselective 1,3-dipolar cycloaddition (1,3-DC)9,10 involving stabilized azomethine ylides and electrophiic alkenes have been promoted using different strategies: a) monofunctional chiral metal complexes10a,c,b) bifunctional chiral complexes10c) double chiral activation of metal catalysts;11c) double chiral activation of metal catalysts;11d) monofunctional chiral organocatalysts;11a,c,g) and e) polyfunctional chiral organocatalysts.11 However, no cooperative catalysis generated by two chiral entities, bound by non-covalent interactions, activating both the dipole and the dipolarophile has been reported to date (Figure 1c).14

In this work, the effects of introducing chirality in the Lewis acid catalyst, together with the chirality of the Brunsted base, in the diastereoe- and enantioselective 1,3-DCs between imino esters 1 and electrophilic alkenes 2 to give enantiomerically enriched prolines 3, will be surveyed (Scheme 1). For this purpose, several Cinchona alkaloids 4-9 as chiral Brunsted bases and a series of chiral BINOL-derived silver phosphates,15,16 generated by mixing the corresponding chiral phosphoric acids with silver carbonate for 1 h, at room temperature, will be evaluated.

Initially, the solution containing the chiral silver phosphate was allowed to react with imino ester 1a and N-maleylmaleimide (NMM, 2a), in the presence (or absence) of the chiral base (reagents added in this order), in toluene as solvent, at rt avoiding

---

[a] Dr. A. Cayuelas, V. Selva, Prof. Dr. C. Nájera, Prof. Dr. J. M. Sansano
Departmento de Química Orgánica, Universidad de Alicante.
Ctra. Alicante-San Vicente s/n, 03080-Alicante, Spain.
[b] Centro de Innovación en Química Avanzada (ORFE-CINQA).
[c] Institute of Organic Synthesis, Universidad de Alicante.
[d] Dr. O. Larrañaga, Dr. A. de Cózar, Prof. Dr. F. Cossio
Departamento de Química Orgánica I. Facultad de Química, Universidad del País Vasco/ Euskal Herriko Unibertsitatea UPV/EHU, P. K. 1072, E-20018 San Sebastián, Spain.
[e] Takahiko Akiyama
Department of Chemistry, Faculty of Science, Gakushuin University, 1-5-1 Meijo, Toshima-ku, Tokyo, 171-8588 Japan.
[g] Dr. José I. Miranda
SGiker NMR Facility, Universidad del País Vasco.

*Corresponding Author for experimental: cnajera@ua.es, jmsansano@ua.es.
*Corresponding Author for calculations: fcossio@ehu.es.
COMMUNICATION

light exposure (Scheme 1 and Table 1). 

Preliminary results, obtained in the absence of a base, revealed that the best chiral silver phosphate was formed by silver carbonate and acid (R)-10 giving ent-endo-3a in excellent both conversion and diastereoselectivity (>96:2 determined by 1H NMR) although with moderate enantioselectivity (Table 1, entry 1). The mixture (S)-10/Ag2CO3 afforded the same results yielding cycloadduct endo-3a instead. The employment of phosphoric acids 11-13 gave lower enantioselectivity (Table 1, entries 2-4) whilst reactions performed with hindered silver phosphates, derived from acids 14-16, furnished almost racemic endo-3a (data not included in Table 1). The bifunctional character of these chiral silver phosphates in this 1,3-DC was disrupted by the introduction of an external base, such as trimethylamine, lowering the enantioselectivity of the process (Table 1, entry 5). In the presence of Cinchona alkaloids as chiral bases (4a, 6-9), the results were satisfactory, especially when cinchonine 6, cinchonidine 7 (in this example ent-endo-3a was obtained), and hydrocinchonine 4a were used (Table 1, entries 6-10). Hydrocinchonine 4a was selected for the next experiments because a higher purity of the crude endo-3a product was observed by 1H NMR. Under these conditions, other solvents such as THF, CH3Cl2, tert-butyl methyl ether, acetonitrile and methanol were tested and the ee was not improved (data not included in Table 1). The lowering of the temperature to 0°C was not efficient in terms of conversion and enantioselectivity. The matched combination between phosphoric acid (R)-10/Ag2CO3/4a was demonstrated after analysis of the enantioselectivity obtained in the reaction mediated by the catalytic system formed by mismatched (S)-10/Ag2CO3/4a triad (Table 1, entry 11). According to this set of results, it was considered that the order of the addition of reagents (Cinchona alkaloid was added at the end) facilitated the reaction of the (R)-10/Ag2CO3 by itself with low enantioselectivity such as it was demonstrated in entry 1 of Table 1. So, a different addition order was attempted (Figure 2) where a better interaction chiral base-imino ester was ensured. This new procedure allowed to obtain cycloadduct endo-3a in 99% ee (Table 1, entry 12). Identical result of 3a was achieved employing the chiral phosphate (R)-13/Ag2CO3 and 4a (Table 1, entry 14), but phosphoric acids 11, 12, and 14-16 furnished poor enantioselections of 3a in low ee. The most important control in the enantioselection was provided by the chiral base (Table 1, entry 13) meanwhile the chiral silver phosphate cooperated to get the maximum level of chiral induction (Table 1, entry 12). The opposite configuration was easily generated using cinchonidine 7 together with (R)-10/Ag2CO3 giving ent-endo-3a in good conversion and 93% ee (data not included in Table 1). The absolute configuration of endo-3a was assigned by comparison of HPLC data and the specific optical rotations of purified samples with the reported ones in previous articles. 

<table>
<thead>
<tr>
<th>Entry</th>
<th>Chiral acid</th>
<th>Base</th>
<th>Conversion (%)(^a)</th>
<th>ee (%)(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(R)-10</td>
<td>2a</td>
<td>100</td>
<td>60 ent-endo-3a</td>
</tr>
<tr>
<td>2</td>
<td>(R)-11</td>
<td></td>
<td>96</td>
<td>34 ent-endo-3a</td>
</tr>
<tr>
<td>3</td>
<td>(R)-12</td>
<td></td>
<td>100</td>
<td>20 ent-endo-3a</td>
</tr>
<tr>
<td>4</td>
<td>(R)-13</td>
<td></td>
<td>100</td>
<td>26 ent-endo-3a</td>
</tr>
<tr>
<td>5</td>
<td>(R)-10</td>
<td>(\text{EtN})</td>
<td>100</td>
<td>52 ent-endo-3a</td>
</tr>
<tr>
<td>6</td>
<td>(R)-10</td>
<td>4a</td>
<td>100</td>
<td>83</td>
</tr>
<tr>
<td>7</td>
<td>(R)-10</td>
<td>6</td>
<td>91</td>
<td>83</td>
</tr>
<tr>
<td>8</td>
<td>(R)-10</td>
<td>7</td>
<td>97</td>
<td>83 ent-endo-3a</td>
</tr>
<tr>
<td>9</td>
<td>(R)-10</td>
<td>8</td>
<td>97</td>
<td>63 ent-endo-3a</td>
</tr>
<tr>
<td>10</td>
<td>(R)-10</td>
<td>9</td>
<td>100</td>
<td>65</td>
</tr>
<tr>
<td>11</td>
<td>(S)-10</td>
<td>4a</td>
<td>97</td>
<td>20</td>
</tr>
<tr>
<td>12(^c)</td>
<td>(R)-10</td>
<td>4a</td>
<td>100</td>
<td>99</td>
</tr>
<tr>
<td>13(^c)</td>
<td></td>
<td>4a</td>
<td>100</td>
<td>83</td>
</tr>
<tr>
<td>14(^c)</td>
<td>(R)-13</td>
<td>4a</td>
<td>100</td>
<td>99</td>
</tr>
</tbody>
</table>

\(^a\) Determined by analysis of crude 1H NMR spectra. \(^b\) Determined by standard or SCF HPLC using columns with chiral stationary phases. \(^c\) Reaction performed using the addition protocol described in Figure 2.

![Scheme 1. Cinchona alkaloids 4-9 and chiral BINOL-derived phosphoric acids 10-16 employed in 1,3-DC between imino ester 1a and N-methylmaleimide 2a.](image-url)
COMMUNICATION

Figure 2. Optimized addition sequence of cooperative catalysts 1-2+3.

In order to better understand our experimental results, we analyzed in detail the parent 1,3-DC between imino ester 1a and NMM 2a in the presence of Ag2CO3 and catalysts (R)-10 and hydroquinonone 4a. On the basis of previous studies on the mechanism of asymmetric catalyzed 1,3-DC between azomethine ylides and s-deficient alkenes, the catalytic cycle gathered in Figure 3 was considered. In addition, we optimized several stationary points associated with this mechanism by means of DFT calculations, which were carried out at the M06(PCM,solvent=PhMe)6-31G(d) & LANL2DZ/6-31G(d)&LANL2DZ level of theory.

According to this catalytic cycle, the protocol described in Figure 2 ensures that when the substrates 1a and 2a are added, the Ag+ cation formed a chiral salt with the corresponding phosphoric acid. In the presence of both catalysts, imino ester 1a generated an N-metalated azomethine ylide INT1 that can react with dipolarophile 2a or with its activated complex INT2. However, when the structure of INT1 was optimized, and its reaction with 2a was analyzed, we found that the chiral information of the phosphoric acid is not optimal to reach an effective blockage of one prochiral face of the dipole and low theoretical enantioselectivity towards ent-endos-3a was obtained (ee=77%, see the SI for further details), in qualitative agreement with the experimental results (Table 1, entries 1-5). Similarly, DFT analysis of the reaction of INT2 with the N-metalated azomethine ylide of 1a in the absence of (R)-10 resulted in a modest ee of 77%, also in qualitative agreement with the experimental results (see also the SI for additional details).

In parallel to activation of 1a, phosphoric acid 10 or its silver salt 10Ag+ (Figure 3) can interact with Brensted base 4a to form non-covalent complex INT3 or, after forming the azomethine ylide moiety, INT4. In both intermediates, hydrogen bonding between the hydroxy group of 4a and the phosphate group was shown by our DFT calculations. In addition, when the O-aryl derivative 4b, was tested as chiral base in the 1,3-DC between 1a and 2a in the presence of Ag2CO3 and (R)-10, the formation of racemic endo-3a was observed. Besides, 1H-NMR experiments with the 4a/(R)-10 pair showed that the quinucleide moiety of 4a was protonated and the surrounding proton resonances changed accordingly. Finally, NOESY correlations between 4a-H+ and the aromatic protons of (R)-10 were observed, thus confirming the presence of complexes INT3 and INT4 gathered in Figure 3. In contrast, no NOEY contacts were observed in solution experiments with the 4b/(R)-10 pair. From intermediate INT4, cycloadducts INT5 were formed via saddle points TS. This step of the catalytic cycle leads to endo-3a with concomitant release of the catalysts at 10-Ag+/4a pair or INT3 intermediate.

We also calculated the possible transition structures leading to the INT5 precursors connected with endo-3a and ent-endo-3a, denoted as TSa-d in Figure 4. We considered both OH⋯O=P and NH⋯O=P contacts between 10-Ag+ and 4a-H+. Our calculations showed concerted although asynchronous [x=1+n2] topologies, in which the C⋯C σ-bonds being formed are more advanced at the distal position with respect to the 4a-H+⋯O=P(NMM) interaction. These saddle points also show double FMO activation between both reactants. The phosphoric moiety promotes a HOMO raising of the azomethine ylide and the protonated alkaloid unit generates a LUMO lowering via the above-mentioned hydrogen bond between 4a-H+ and 2a. Among the four saddle points shown in Figure 4, TSa is the one of lowest energy because of optimal interaction between the phosphoric and alkaloid units. The solvent accessible surface (SAS) shows close contacts between both chirality sources, in agreement with the NOESY experiments. Consequently, a quite rigid structure is formed, which generates and enhances chiral induction with respect to the separate catalysts.

This induction results in a preferential [a(Si)⋯d(Si)b(Re)⋯c(Re)] supra-supra interaction (Figure 4) that ultimately leads to cycloadduct, endo-3a, in nice agreement with the experimental results. The other transition structures TS-c-d exhibit less than optimal interactions. In particular, the N⋯H⋯O=P hydrogen bonding between both catalytic units were calculated to be less efficient that the OH⋯O=P interactions, a result compatible with the lack of enantioselectivity observed in the presence of O-aryl derivative 4b (vide supra). All these calculations result in a computed enantiomeric excess of 92%, in good agreement with our experimental result of 99% (see the SI for further information).

Figure 3. Proposed catalytic cycle for the successful cooperative non-covalent catalysis involving catalysts 10 and 4a via double FMO activation.

Figure 4. Transition structures TSa-d associated with 1,3-DCs between 1a and 2a to give endo-3a and ent-endo-3a in the presence of catalysts Ag2CO3, 4a and (R)-10. All structures were fully optimized at the M06(PCM,solvent=Ph Me).
With this information, the survey of the versatility of this cooperative enantioselective 1,3-DC was carried out using BINOL-derived silver phosphate and hydrocinchonine (Figure 5). NMM dipolarophile 2a was a suitable dipolarophile for these transformations and reacted satisfactorily with methyl, isopropyl and tert-butyl benzylideneglycinates in good yields and very high to excellent diastereo (>98:2 dr) and enantioselectivities of endo-3a-c, methyl ester being the most appropriate starter. Alanine derivative 1 was also allowed to react with the catalytic system and 2a affording endo-3d in 93% ee. Prolinate endo-3e was diastereoselectively obtained in 61% yield and 95% ee (Figure 5). More hindered substrates and different dipolarophiles as ortho-substituted arenes in the imino group of 1 and tert-butyl acrylate were tested together in the same reaction affording endo-3f in good yield (78%) and excellent ee (>99%).

β-Nitrostyrene is an excellent dipolarophile that, generally, promotes an exo-approach towards the dipole, but in this reaction, unexpected endo-cycloadduct 3g was the major diastereoisomer isolated with a 95% ee. This result also supports the proposed interaction dipolarophile (nitro group)-protonated quinuclidine described in INT4 (Figure 3). Chalcone and 4-methoxychalcone were also essayed with methyl iminoglycinate 1a giving excellent enantioselectivity and high yields of endo-3h-j.

In conclusion, in this work an efficient endo-diastereo and enantioselective 1,3-DC of azomethine ylides based on a cooperative asymmetric Lewis acid-Bronsted base catalysis has been presented. Silver(I) salts of chiral phosphoric acids are poor asymmetric catalysts for this reaction. Cinchona alkaloids, in turn, are efficient in terms of chiral induction. In contrast, matched non-covalent coupling between both cooperative catalysts results in a significant improvement of the observed ee’s. We think that the concepts presented in this communication can be applied to other reactions involving doubly FMO-activated nucleophilic and electrophilic substrates.

Acknowledgements

We gratefully acknowledge financial support from the Spanish Ministerio de Economía y Competitividad (MINECO) (projects CTQ2013-43446-P and CTQ2014-51912-RED), the Spanish Ministerio de Economía, Industria y Competitividad, Agencia Estatal de Investigación (AEI) and Fondo Europeo de Desarrollo Regional (FEDER, EU) (projects CTQ2016-76782-P, CTQ2016-80375-P and CTQ2016-81797-RED), the Generalitat Valenciana (PROMETEOII/2014/017), the Gobierno Vasco/ Eusko Jaurlaritza (GV/EJ, Grant IT673-13) and the University of Alicante. This work was partially supported by a Grant-in-Aid for Scientific Research from JSPS (17H03060).

Keywords: asymmetric catalysis • cooperative catalysis • cycloadditions • DFT calculations • enantioselective synthesis
Better together. Enhanced non-additive enantiocontrol is observed in asymmetric 1,3-DC between imino esters and alkenes in the presence of commercially available Lewis acid-Bronsted base catalytic pairs. This cooperative catalysis is based on FMO-mediated activation of both reactants. The reinforced combined chiral induction relies on contacts between both catalysts mediated by a non-covalent interaction under mild conditions.
In nature, for example, reductases or oxidases are combining with NADP+/NADPH cofactor allowing hydrgenly efficient specific cooperative asymmetric transformations.


[1] When two or more substances that produce an effect greater than the sum of their individual effects can be considered as synergism. In addition, two or more entities working or acting together willingly for a common purpose or benefit involving mutual assistance in working towards a common goal is considered cooperation. According to these simple definitions we establish that a synergy can be produced by two catalysts in a double activation catalysis or in a single activation catalysis. In both cases it is convenient to point that a dual activation catalysis.
