Enantioselective Redox-Neutral Coupling of Aldehydes and Alkenes by an Iron-Catalyzed “Catch–Release” Tethering Approach

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ABSTRACT: The reductive coupling of aldehydes and alkenes is an emerging technology that holds the potential to reinvent carbonyl addition chemistry. However, existing enantioselective methods are limited to form “branched” products. Herein, we present a directed enantio- and diastereoselective alkylation of aldehydes with simple olefins to selectively yield linear coupling products. This is achieved by redox-neutral remote functionalization, whereby a tethering “catch–release” strategy decisively solves the key problems of reactivity and selectivity.

In recent years, coupling approaches combining aldehydes and alkenes, readily available feedstocks, have started to emerge as an appealing alternative to classical carbonyl addition reactions (Figure 1a). Krische and co-workers pioneered the development of both hydrogenative and transfer hydrogenative couplings of carbonyl compounds with a variety of unsaturated compounds, including enones, alkynes, 1,3-dienes, enynes, allenes, and allyl compounds, yielding the corresponding branched coupling products. Additionally, Buchwald and co-workers developed the asymmetric addition of alkene-derived nucleophiles to ketones and leading exclusively to the branched products. However, enantioselective methods to access the complementary, highly valuable linear products (Figure 1b) using a direct redox-neutral coupling of olefins have not yet been developed.

As part of our group’s ongoing research on chirality transfer processes, we became interested in accessing these scaffolds from readily available chiral building blocks. We therefore envisioned an enantioselective redox-neutral coupling that would take advantage of a carbocatonic mechanism to ensure complete selectivity for the linear coupling product (Figure 1b). To overcome the low intrinsic nucleophilicity of unactivated alkenes, we decided to employ a “catch-release” tethering group approach (Figure 1c). Under Lewis-acid catalysis, the alcohol moiety on the substrate would serve as anchoring point for the aldehyde reaction partner, reversibly forming an oxocarbenium ion (cf. Figure 1c). By virtue of tethering, attack of the alkyne partner onto this highly electrophilic species becomes intramolecular, thus circumventing reactivity issues and leading to a cyclic, tertiary carbocation. In order to evade the potential deleterious side reactions typical of carbocatonic intermediates, a designed 1,5-hydride shift event‡ would not only allow convergence to a stabilized oxacarbenium ion C but also ensure the formal reduction of carbocation B. Hydrolysis of C would then achieve a “release” event and yield the desired linear coupling motif.

Herein we present our reduction of this concept to practice in an asymmetric synthesis of linear aldehyde–alkene coupling motifs by redox-neutral remote functionalization under iron catalysis, featuring the possibility of forming two new challenging chiral centers starting from a single, readily chiral-pool derived, stereogenic element.

At the onset of our investigation, we tested several unsaturated alcohols, differing in the length of the spacing carbon-chain (Table 1, entries 1–3), as alkene partners. After some initial experimentation with aldehyde as model electrophile, we identified SnCl4 as a suitable Lewis Acid. To our delight, alcohol 1a (with 3 methylene units between the carbonyl and the alkene) showed promising reactivity and already excellent levels of diastereoselectivity of the final product 3a, carrying 2 stereogenic centers in a 1,3-relationship. This substrate mechanically implies the transient formation of an eight-membered ring (cf. intermediate B in Figure 1). Encouraged by these promising initial results, we continued with substrates 1a and 2a for further optimization. Changing the amount of tin(IV) chloride led to worse results (entries 4 and 5). While investigating other Lewis acids (entries 6–9), we identified iron(III) chloride as a uniquely effective catalyst for this reaction. Increasing the temperature from room temperature to 100 °C resulted in better yield and remarkably short reaction times (entry 10). Notably, product 3a was obtained as virtually a single enantio- and diastereomer. Importantly, the reaction can be run without the need for scrupulously dry solvents or the use of an inert-gas atmosphere.

With optimized conditions in hand, we turned our attention toward the scope of this transformation (Scheme 1). The reaction tolerates a variety of different aldehyde electrophiles, including sterically congested cases such as 2-methylpropanal, cyclohexylcarboxaldehyde or even pivalaldehyde (Scheme 1, 3a–e). Notably, all products are obtained in good yields after a short reaction time and essentially as single enantio- and diastereoisomers. A wide range of functional groups, including halides (3f–g), protected alcohols (3h), and alkyl thiocarbonyls (3i) as well as esters, unsaturated systems, and alkynes (3j–l) are tolerated under the present reaction conditions. On the alkene component, aromatic and other aliphatic substituents show excellent reactivity (3m–p) while retaining the same near perfect diastereo- and enantioselectivity. Additionally, products resulting from a nine-membered...
The intermediate can be accessed in equal selectivities by ortho-appending of the bridging carbon chain around an aromatic core (3q).

The reasons behind the excellent stereoselectivity observed throughout the transformations compiled in Scheme 1 intrigued us from the very first experiments. Scheme 2 depicts our stereochemical model. We assume that the aldehyde R substituent and the carbinol residue R1 adopt pseudoequatorial orientations during the cyclization event, at which point the first stereogenic center of the product is formed. During the hydride transfer step, the original chiral element of the substrate is deleted concomitantly with the setting of the second stereogenic center of the product. As hydride transfer must proceed suprafacially, this ensures perfect chirality transfer and rationalizes the excellent stereoselectivities observed even when acetaldehyde, otherwise a challenging aldehyde for enantioselective C−C bond forming reactions is employed (3r). Additionally, an α-chiral aldehyde such as 2s (i.e., (S)-2-methylbutyaldehyde) is a suitable substrate for this process, allowing the preparation of 3s without epimerization of its sensitive chiral center (3s obtained in 71% yield and >20:1 d.r.). Furthermore, we found that there is a large reactivity difference between aliphatic and aromatic aldehydes, enabling the use of a dialdehyde substrate such as 3t without the need for protecting group manipulations.

The products of this methodology are chiral diads represented in a range of natural products and bioactive substances. As an example we chose 4, a key intermediate in Maio’s study on the synthesis of the neuroactive marine macrolide Palmynolide A and its analog (Scheme 3). In the course of Maio’s original work, amide 4 was synthesized from pivalaldehyde (Scheme 3b). The sequence includes a proline-assisted aldol addition, setting the first chiral center. Evans–

Table 1. Selected Optimization of Reaction Conditions

<table>
<thead>
<tr>
<th>entry</th>
<th>alcohol</th>
<th>Lewis acid (%)</th>
<th>time (temperature)</th>
<th>yield [%] (dr)</th>
<th>ee [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>rac-1a</td>
<td>SnCl4 (20%)</td>
<td>1 min (r.t.)</td>
<td>65 (&gt;20:1)</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>rac-1b</td>
<td>SnCl4 (100%)</td>
<td>15 min (r.t.)</td>
<td>&lt;5 (n.d.)</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>rac-1c</td>
<td>SnCl4 (20%)</td>
<td>1 h (r.t.)</td>
<td>25 (&gt;20:1)</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>(S)-1a</td>
<td>SnCl4 (5%)</td>
<td>3.5 h (r.t.)</td>
<td>45 (&gt;20:1)</td>
<td>&gt;99%</td>
</tr>
<tr>
<td>5</td>
<td>(S)-1a</td>
<td>SnCl4 (30%)</td>
<td>1 min (r.t.)</td>
<td>35 (&gt;20:1)</td>
<td>&gt;99%</td>
</tr>
<tr>
<td>6</td>
<td>(S)-1a</td>
<td>AlCl3 (20%)</td>
<td>10 min (r.t.)</td>
<td>trace (n.d.)</td>
<td>n.d.</td>
</tr>
<tr>
<td>7</td>
<td>(S)-1a</td>
<td>TMSOTf (20%)</td>
<td>24 h (r.t.)</td>
<td>trace (n.d.)</td>
<td>n.d.</td>
</tr>
<tr>
<td>8</td>
<td>(S)-1a</td>
<td>HCl (20%)</td>
<td>24 h (r.t.)</td>
<td>trace (n.d.)</td>
<td>n.d.</td>
</tr>
<tr>
<td>9</td>
<td>(S)-1a</td>
<td>FeCl3 (20%)</td>
<td>10 min (r.t.)</td>
<td>71 (&gt;20:1)</td>
<td>&gt;99%</td>
</tr>
<tr>
<td>10</td>
<td>(S)-1a</td>
<td>FeCl3 (5%)</td>
<td>5 min (100 °C)</td>
<td>76 (&gt;20:1)</td>
<td>&gt;99%</td>
</tr>
</tbody>
</table>

“Reaction conditions: alcohol (0.2 mmol), aldehyde (0.24 mmol), Lewis acid in dichloromethane (DCM, 0.1 M).” Determined by 1H NMR analysis of the crude reaction mixture. “Determined by chiral stationary phase HPLC.” “Isolated yield. Reaction performed in 1,2-dichloroethane (DCE, 0.1M). See SI for further screening of reaction conditions.”
Tishchenko reaction followed by installation of a sulfate ester gives compound 5, which can be converted into 6 via allylation and protection of the alcohol-moiety. Hoveyda–Grubbs metathesis and hydrogenation results in key intermediate 4 in 24% overall yield. Using the method reported herein, that sequence can be cut in half. Commercially available and cheap \((S)-\text{propylene oxide}\) was first ring-opened to 1a in gram-scale. The novel redox-neutral coupling of 1a with pivalaldehyde gives ketone 3e in good yield and perfect stereochemical outcome. Simple Lieben-haloform reaction directly followed by amide coupling using standard conditions gives key intermediate 4 in 37% overall yield over only 4 steps from cheap commercially available materials. In addition, direct comparison of product 4 to the material obtained by Maio also allowed unambiguous assignment of stereochemistry to be as shown (C5R, C7S).

In conclusion, we have developed an efficient method to access linear aldehyde/alkene coupling products as essentially single diastereo- and enantiomeric materials. An easily accessible chiral alcohol functions as a tether which, by virtue of a “catch−release” process, overcomes the reactivity and selectivity limitations typically associated with olefins as nucleophiles. This method allows the assembly of two chiral centers in near-perfect stereoselectivity, starting from one lone, chiral-pool-derived stereocenter.

Scheme 1. Scope of the Redox-Neutral Coupling of Aldehydes and Alkenes

Scheme 2. Proposed Stereochemical Model for the Redox-Neutral Coupling of Aldehydes and Alkenes

Scheme 3. Synthesis of Key Intermediate 4 and Determination of the Absolute Configuration
Iridium-Catalyzed Carbonyl Allylation from the Alcohol or Aldehyde

References

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Notes

The authors declare no competing financial interest.

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