Synthetic Methods

Enantiospecific Three-Component Alkylation of Furan and Indole

Mattia Silvi, Raffael Schrof, Adam Noble, and Varinder K. Aggarwal[*][a]

Abstract: Furan- and indole-derived boronate complexes react with alkyl iodides under radical (photoredox) or polar (S$_2$2) conditions to generate three-component alkylation products with high efficiency and complete stereospecificity. The methodology allows the incorporation of versatile functional groups such as nitriles, ketones, esters, sulfones, and amidcs, providing rapid access to complex chiral heteroaromatic molecules in enantioenriched form. Interestingly, while indolyl boronate complexes react directly with alkyl halides in a polar pathway, furyl boronates require photoredox catalysis. Careful mechanistic analysis revealed that the boronate complex not only serves as a substrate in the reaction but also acts as a reductive quencher for the excited state of the photocatalyst.

Heteroaromatic compounds are ubiquitous in natural products and bioactive compounds and play a central role in organic chemistry. In particular, the furan ring is a versatile moiety for organic synthesis, readily undergoing oxidations, acid-catalyzed rearrangements and cycloaddition reactions,[1] while the indole ring is one of the most common motifs in alkaloids and drug candidates.[2] Our group recently introduced an enantiospecific sp$^3$–sp$^2$ coupling between aromatic rings and chiral boronic esters 1 to access a variety of enantioenriched monofunctionalized aromatic molecules (e.g. 3, Scheme 1a).[3] Furthermore, inspired by the intriguing reactivity of unsaturated boronate complexes,[4] we showed that nucleophilic furyl-derived boronate complexes 2 react with electrophiles in an enantiospecific three-component coupling reaction (leading to 4, Scheme 1a).[5] Key to the success of this process was the use of highly activated electrophiles, such as the Umemoto trifluromethylating agent[6] or carbocationic species. Interestingly, the trifluoromethylation reaction was found to proceed through a self-initiated radical chain mechanism. Independently, Studer and our group have shown that electrophilic radicals add to vinyl boronate complexes 5 (Scheme 1b).[7] The resulting α-boronic radicals 7 undergo facile single electron oxidation to trigger 1,2-migration and form boronic ester adducts 8.

We envisioned extending this merger of photoredox catalysis[8] with boron 1,2-metallate rearrangements to aromatic boronate systems (Scheme 1c).[9,10] Photoredox radical-mediated reaction of alkyl halides 6[11] with boronate complexes derived from chiral boronic esters 9 and aryllithiums 10 should lead to dearomatized intermediates 11 and 12. As 1,2-migration is well-known to be a stereospecific process,[12] the stereochemistry within boronic ester 1 should be conserved in the process. In situ oxidation/rearomatization of 11 and 12 would then provide chiral aromatic compounds 13 and 14 in enantioenriched form. Such a transformation would constitute a novel enantiospecific three-component alkylation protocol, providing valuable routes to complex chiral heteroaromatic structures. Herein, we describe the successful development of this new methodology, which takes advantage of the versatility and mild conditions of photoredox catalysis to introduce a suite of synthetically valuable functional groups.

Attracted by the versatility of nitriles in organic synthesis,[13] we commenced our studies by investigating the reaction of furan with iodoacetonitrile (6a) (Table 1).[14] Boronate complex 2a was formed by addition of cyclohexyl boronic acid pinacol ester (1a) to 2-furyllithium, generated by lithiation of furan with n-butyllithium in THF. Solutions of iodoacetonitrile...
Photocatalyst were then added and the mixture irradiated with blue LEDs for 1 h. Gratifyingly, in a preliminary experiment carried out using acetonitrile/THF as solvent (entry 1), we found that the reaction proceeded smoothly to give the desired de aromatized intermediate 11a in 50% yield using Ru(bpy)2+ as a photocatalyst. A solvent screen led to the identification of a mixture of DMF/THF (2:1) as optimal, giving 11a in 73% yield (entry 4). The use of the more reducing photocatalyst Ir(ppy)2 gave similar results (entry 5). In contrast to the photochemical three-component alkylation of vinyl boronate complexes,2a we observed that the use of a photocatalyst was crucial for obtaining high yields in this reaction (compare entries 4 and 6). A control experiment carried out in the dark did not give any product (entry 7), showing the photochemical nature of this transformation. Intermediate 11a could be oxidized by addition of iodine and potassium acetate7a to the reaction vessel to give the desired three-component alkylation product 13a in 71% isolated yield (entry 4).

Having identified the optimum conditions, we explored the scope of our photochemical transformation (Scheme 2). A wide range of synthetically versatile functional groups could be efficiently introduced by varying the alkyl iodide radical precursor. Iodoacetophenones proved to be good substrates, leading to the corresponding ketones in good yields (13b and 13c). Ethyl iodoacetate and iodoacetamide gave ester 13d and unprotected amide 13e. Iodomethyl phenylsulfone led to sulfone 13g in 64% yield.16,17 In addition, perfluoroalkyl chains could be connected to the furan scaffold (13f) starting from readily available perfluoroalkyl iodides. The employment of tertiary and primary boronic esters led to products 13h and 13i in good yields, demonstrating that the process tolerates a wide spectrum of steric demand. Furthermore, starting from enantioenriched chiral boronic esters, compounds 13j and 13k were obtained with complete stereospecificity.

Attracted by the importance of the indole ring in synthetic and medicinal chemistry, we sought to extend our enantiospecific three-component alkylation reaction to N-methyl indole (Scheme 3). Pleasingly, the reaction of boronate 15 with iodoacetone (6a) gave the desired product 14a in 70% yield. However, in control experiments we found that neither light nor the photocatalyst were required for the transformation. Furthermore, addition of the radical inhibitor 1,1-diphenylethylamine7a had no effect on the reaction outcome (see Supporting Information-3.5 for details), ruling out a possible electron-transfer initiated radical chain process7b and supporting a polar

**Table 1. Reaction optimization.**

<table>
<thead>
<tr>
<th>Entry</th>
<th>Solvent</th>
<th>Photocatalyst (PC)</th>
<th>Yield of 11a [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CH3CN/THF</td>
<td>Ru(bpy)2Cl2·6H2O</td>
<td>50</td>
</tr>
<tr>
<td>2</td>
<td>DMSO/THF</td>
<td>Ru(bpy)2Cl2·6H2O</td>
<td>66</td>
</tr>
<tr>
<td>3</td>
<td>DMF/THF</td>
<td>Ru(bpy)2Cl2·6H2O</td>
<td>66</td>
</tr>
<tr>
<td>4</td>
<td>DMF/THF</td>
<td>Ir(ppy)2</td>
<td>73 (71)</td>
</tr>
<tr>
<td>5</td>
<td>DMF/THF</td>
<td>–</td>
<td>26</td>
</tr>
<tr>
<td>6</td>
<td>DMF/THF</td>
<td>Ru(bpy)2Cl2·6H2O</td>
<td>0</td>
</tr>
</tbody>
</table>

[a] All the reactions were carried out using 1.2 equiv of furan, 1.15 equiv of iodoacetone, 6a on a 0.2 mmol scale. [b] Mixture of solvents are intended solvent/THF 2:1. [c] DMF: 1,3-dimethyl-2-imidazolidinone. [d] Yield measured through 1H NMR analysis of the crude mixture using dibromomethane as an internal standard. Intermediate 11a was obtained as a 1:1 mixture of diastereomers. Number in parenthesis is the isolated yield of compound 13a after oxidation and chromatographic purification. [e] Photochemical step time: 2 hours. [f] Reaction carried out in the dark.

**Scheme 2. Scope of the enantiospecific three-component alkylation of furan.**

All the yields refer to isolated product after chromatographic purification. [a] Intermediate oxidation conditions: NaClO (aq.), DMF, –20 °C. [b] 4 equiv of alkyl iodide were used. [c] Volatile product, number in parenthesis is the NMR yield using CH3Br2 as internal standard.

**Scheme 3. Scope of the enantiospecific three-component alkylation of indole.**

All the yields refer to the isolated product after chromatographic purification. [a] Intermediate oxidation conditions: H2O2(aq.)/NaOH(aq.), 0 °C, DMF/THF 2:1.
S$_n$2-like pathway. The nucleophilic reactivity of indole-derived borates generated from difficult-to-handle trialkyl boranes has been described,$^{[18]}$ however the difficulties in accessing enantioenriched chiral boranes has prevented the use of this methodology for the synthesis of chiral compounds in enantioenriched form. Surprisingly, reactions of boronates derived from stable boronic esters are rare with only a single report very recently disclosed by Ready et al. (with $\pi$-allyl palladium complexes).$^{[19]}$

Through our simple protocol, indole-derived boronates could be alkylated to introduce a diverse range of functional groups, including nitriles, ketones, esters and unprotected amides, providing functionalized indoles 14a–d in good yields. Various other alkyl boronic esters were also applied to the coupling reaction, including primary (14e) and enantioenriched chiral secondary (14f and 14g) examples, all proceeding in high yields and with complete stereospecificity. However, using the bulky menthyl boronic ester provided the desired product 14h in only 18% yield (albeit with excellent stereospecificity) showing that steric hindrance has an impact in this reaction. Finally, a simple control experiment showed that N-methylindole does not undergo Friedel–Crafts alkylation with iodoacetonitrile 6a under our reaction conditions (see Supporting Information–3.5 for details), thus highlighting the importance of the boronate π-donation to the indolyl π-system for the nucleophilicity of 15. Indeed, Mayr has shown that a BF$_2$K moiety (which is not an electron-donating as RBpinLi) at the 2 position increases the nucleophilicity of N-Boc indole by $> 10^2$.$^{[20]}$

To glean insights into the mechanism of the furan three-component coupling reaction, selected spectroscopic and electrochemical studies were carried out. Quantum yield measurements gave a value of $\Phi = 27.8$ (see Supporting Information–3.4 for details), suggesting a radical chain pathway to be operative.$^{[21]}$ Fluorescence quenching analysis revealed that boronate complex 2a was an effective quencher of the excited state photocatalyst, whereas iodoacetonitrile (6a) was ineffective (see Supporting Information–3.2 for details). Based on these results, we propose the mechanism depicted in Scheme 4. The highly reducing Ru$^0$ is generated by single electron transfer (SET) from a sacrificial amount of boronate complex 2a ($\epsilon_{\text{Ru}^{II} \rightarrow \text{Ru}^{I}} = +0.26$ V vs. SCE, Figure S5) to the excited Ru$^0$ catalyst ($\epsilon_{\text{Ru}^{I}} = +0.77$ V vs. SCE).$^{[22,23]}$ Since one electron oxidation of analogous boronate complexes has been shown to lead to the generation of alkyl radicals through C–B bond fragmentation,$^{[24]}$ this reductive quenching phenomenon is expected to lead to by-product 16, which was indeed observed in the crude reaction mixtures. Once formed, the electron-rich Ru$^0$ species ($\epsilon_{\text{Ru}^{I}} = −1.33$ V vs. SCE) undergoes single electron transfer with iodoacetonitrile (6a, $\epsilon_{\text{I}} = −1.24$ V vs. SCE, Figure S6) leading to the formation of the reactive electrophilic radical 17 and the regeneration of the Ru$^0$ catalyst. Radical 17 then adds to the furyl system of 2a generating radical anion 18. The electron-rich radical anion is expected to undergo SET with another molecule of iodoacetonitrile (6a), forming radical 17 and a zwitterionic species (not shown), which undergoes 1,2-migration to release intermedia-

**Scheme 4. Proposed reaction mechanism for the three-component alkylation of furan.**

In conclusion, we have developed a novel stereospecific three-component alkylation reaction of furans and indoles with boronic esters and electron-deficient alkyl iodides. Mechanistically, the more electron-rich indole boronates are sufficiently nucleophilic to react directly with alkyl iodides through a polar pathway. Conversely, alkylation of the less reactive furyl boronates proceeded through a radical pathway induced by photoredox catalysis. Careful mechanistic analysis showed that the furyl boronate complex 2 plays a dual role, acting both as sacrificial reductive quencher for Ru$^0$ (giving the reductant RuI) and as reactant for the three-component alkylation reaction.

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**Conflict of interest**

The authors declare no conflict of interest.
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[15] An extensive screen of different bases revealed that potassium acetate (instead of potassium carbonate [see Ref. 5]) led to a signiﬁcant improvement in oxidation yield. With other substrates we found that NaClO (aq.), in DMF was required. See the Supporting Information for details.


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