Catalysis

Enantioselective Nickel-Catalyzed *anti*-Arylmetallative Cyclizations onto Acyclic Ketones

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Abstract: Domino reactions involving nickel-catalyzed additions of (hetero)arylboronic acids to alkynes, followed by cyclization of the alkynenickel intermediates onto tethered acyclic ketones to give chiral tertiary-alcohol-containing products in high enantioselectivities, are described. The reversible E/Z isomerization of the alkynenickel intermediates enables overall *anti*-arylmetallative cyclization to occur. The ring system of the products are substructures of certain diarylindolizidine alkaloids.

Stereoegenic cyclic tertiary alcohols are important structural units that feature prominently in biologically active natural products and therapeutic compounds. Accordingly, new methods for the enantioselective construction of these units provide valuable tools for target synthesis. Of the strategies available, the catalytic asymmetric addition of carbon nucleophiles to ketones ranks highly in directness, versatility, and overall synthetic efficiency.[1] One subset of these reactions are metal-catalyzed domino sequences initiated by the addition of an arylboron reagent to an alkyne, followed by enantioselective cyclization of the resulting alkenenickel species onto a tethered ketone, which are applicable to the synthesis of diverse carbocyclic and heterocycles.[2] Recently, nickel-catalyzed variants of these reactions have been developed in which reversible E/Z isomerization of the alkenenickel intermediates is essential for cyclization.[3] Application of this general method to achiral products has also been reported,[4] and other related nickel-catalyzed processes have also appeared.[5–7] In addition to nickel being much less expensive than comparable rhodium- or palladium-catalyzed reactions,[3a] the diverse reactivity of nickel catalysis[5] often enables unique transformations not available to other metal catalysts.

Our first contribution to this field included a study of enantioselective nickel-catalyzed desymmetrizations of cyclic 1,3-diketones, which give fused bicycles in high diastereo- and enantioselectivities (Scheme 1A).[3a] Although effective, the ability to use acyclic ketones in non-desymmetrizing cyclizations would also be valuable to significantly broaden the substrate scope and provide simpler, non-fused products. However, acyclic ketones are potentially less reactive than cyclic 1,3-diketones because of their greater conformational flexibility, and because they lack the activation from the second ketone through its electron-withdrawing effect as well as the electronic repulsion caused by having aligned dipoles. Two individual examples of non-asymmetric nickel-catalyzed arylation cyclizations onto acyclic ketones have been reported recently,[6d] but to our knowledge, corresponding enantioselective processes have yet to be described.

Herein, we describe the successful use of acyclic dialkyl and alkyld-aryl ketones in these reactions in the enantioselective preparation of aza- and carbocyclic tertiary alcohols

Scheme 1. Enantioselective nickel-catalyzed arylation cyclizations onto ketones.
Most of the products contain the 4,5-diaryl-1,2,3,6-tetrahydropyridine ring system, which appears in indolizidine natural products such as (−)-phyllosteminine, (−)-septicine, and (−)-fistulopsine A (Scheme 1C). This investigation began with the reaction of \( \text{PhB(OH)}_2 \) with acyclic substrates 1, which contain an alkyne tethered to a ketone through a sulfonamide (Table 1). Chiral phosphine-oxazoline (PHOX) ligands have proven to be excellent ligands in related studies and we found (S)-Bu-PHOX \( (L_1) \) to be highly effective in these reactions. Heating a mixture of the substrate 1 and PhB(OH)\(_2\) (2.0 equiv) in the presence of 10 mol% each of Ni(OAc)\(_2\)·4H\(_2\)O and \( L_1 \) in TFE (2,2,2-trifluoroethanol) at 60 or 80 °C for 24 h provided azacycles 2a–2k in 44–90% yield and up to 99% ee. Small quantities of minor arylative cyclization products 3, resulting from initial phenylnickelation of the alkyne with the regioselectivity opposite to that required for the formation of the major products 2, were also observed by \(^1\)H NMR spectroscopy, but with the exception of the reaction producing 2k and 3k, these were not isolated. A range of aromatic ketones are tolerated in these reactions, with substrates containing phenyl (2a), 4-chlorophenyl (2b), (3-trifluoromethyl)phenyl (2c), or 2-methylphenyl ketones (2d) readily undergoing arylative cyclization. Simple dialkyl ketones are also competent electrophiles, with ketones containing methyl (2e, 2f, 2i, and 2j), ethyl (2k), isopropyl (2g), or methyl propanoate (2h) groups reacting successfully. Regarding the alkyln sub- stituent, the process is tolerant of phenyl (2a–2h), (4-carbomethoxy)phenyl (2i), and 3-methoxyphenyl groups (2j).

Next, different boronic acids were investigated in reactions with substrate 1f, and we were pleased to observe that arylative cyclization products 2l–2p were obtained in up to 79% yield and uniformly high enantioselectivities (98% to >99% ee, Table 2). Various substituted phenylboronic acids are compatible with this process including, notably, 3-hydroxyphenylboronic acid (2m). 2-Naphthylboronic acid (2o) and 3-thienylboronic acid (2p) also readily underwent the reaction.

Table 1. Scope of alkynyl-tethered ketones.  

| 2a | 84% (11:1), 94% ee \(^{[a]}\) |
| 2b | 76% (10:1), 98% ee \(^{[a]}\) |
| 2c | 76% (10:1), 98% ee \(^{[a]}\) |
| 2d | 72% (10:1), >99% ee \(^{[a]}\) |

[a] Reactions were conducted using 0.30 mmol of \( 1 \) in TFE (3 mL). Yields are of isolated products. Values in parentheses refer to the ratio of 2:3 as determined by \(^1\)H NMR analysis of the crude reactions. Unless stated otherwise, the minor isomers 3 were not evident in the isolated products. Enantiomeric excesses were determined by HPLC analysis on a chiral stationary phase. [b] At 80 °C. [c] At 60 °C. [d] Product 2k was obtained as an inseparable 12:1 mixture together with the minor product 3k in 90% combined yield.

Table 2. Scope of boronic acids.  

| 2l | 78% (>10:1), 50% ee |
| 2m | 67% (10:1), >99% ee |
| 2n | 70% (19:1), >99% ee |
| 2o | 51% (>19:1), >99% ee |
| 2p | 94% (>19:1), 98% ee |

[a] Reactions were conducted using 0.30 mmol of \( 1f \) in TFE (3 mL). Yields are of isolated products. Values in parentheses refer to the ratio of 2:3 as determined by \(^1\)H NMR analysis of the crude reactions. Unless stated otherwise, the minor isomers 3 were not evident in the isolated products. Enantiomeric excesses were determined by HPLC analysis on a chiral stationary phase.
Further experiments to explore the scope of this process in the reactions of various other substrates with PhB(OH)₂ are shown in Schemes 2–6. Changing the alkynyl substituent to a chloride was only moderately successful; substrate 11 reacted to give chloroalkene-containing tetrahydropyridine 2q in 12 % yield and 71 % ee, with the remainder of the material being predominantly unreacted 11 (Scheme 2). Next, the preparation of carboxyclic products was attempted by changing the sulfonamide connecting the alkyne and the ketone to a malonyl group. Interestingly, the reaction of substrate 4 did give the six-membered product 5 in 25 % yield and 84 % ee, but the cyclopent-2-ene 6a resulting from cyclization of the intermediate alkenynickel species onto one of the ester groups was also obtained in 14 % yield[12] and 85 % ee (Scheme 3). The enantioselective formation of cyclopent-2-enones in this manner was described by our group previously[3a] and in this case, it appears that despite the lower electrophilicity of the methyl esters in 4 compared with the phenyl ketone, the kinetic preference to form a five-membered ring over a six-membered ring makes the formation of 6a competitive with 5. An attempt to prepare a five-membered cycloalkanol by the reaction of PhB(OH)₂ with substrate 7 was unsuccessful, and gave a complex mixture of unidentified products (Scheme 4). However, it should be noted that the formation of five-membered rings by anti-carbometallative cyclizations onto cyclic 1,3-diketones was successful in our previous work (see Scheme 1A).[3a]

Attempts to form seven-membered ring products are shown in Schemes 5 and 6. Substrate 8 reacted with PhB(OH)₂ to give cyclopent-2-ene 6b in 55 % yield and 19 % ee[13] but no product resulting from cyclization onto the ketone was observed (Scheme 5). In addition, the reaction of substrate 9 with PhB(OH)₂ led to the trisubstituted alkene (Z)-10 resulting from alkyne hydroarylation as the only isolable product in 34 % yield (Scheme 6). The identity of the remainder of the material in this reaction was not clear; although this did not appear to contain an appreciable quantity of the corresponding E-isomer of 10, we cannot rule out its presence resulting from E/Z isomerization of the alkenynickel intermediate.

In summary, we have described enantioselective nickel-catalyzed anti-arylmallative cyclizations of (hetero)arylboronic acids with substrates containing an alkyne tethered to an acyclic ketone, which proceed to give chiral tertiary alcohols with high enantioselectivities in most cases (often ≥ 99 % ee). Compared with a previous study,[3a] this work demonstrates a substantial increase in scope of ketones that can be used as electrophiles. The products are 4,5-diaryl-1,2,3,6-tetrahydropyridines, a ring system that is seen in certain indolizidine alkaloids. The formation of carboxyclic products is also possible.[14]

Scheme 2. Reaction of chloroalkyne 11.

Scheme 3. Formation of carboxyclic products 5 and 6a.

Scheme 4. Attempted reaction of substrate 7.

Scheme 5. Attempted formation of a seven-membered carbocycle.

Scheme 6. Attempted formation of a seven-membered azacycle.

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Conflict of interest

The authors declare no conflict of interest.

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was accompanied by inseparable, unidentified
and the assignment of its absolute con-
figure, which was made by analogy to the results reported in
ref. [3c], is tentative.

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