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Catalytic Asymmetric Synthesis of Tertiary Alkyl Fluorides: Negishi Cross-Couplings of Racemic $\alpha_i \alpha$ -Dihaloketones

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ABSTRACT: The development of new approaches to the construction of fluorine-containing target molecules is important for a variety of scientific disciplines, including medicinal chemistry. In this Article, we describe a method for the catalytic enantioselective synthesis of tertiary alkyl fluorides through Negishi reactions of racemic α -halo- α -fluoroketones, which represents the first catalytic asymmetric crosscoupling that employs geminal dihalides as electrophiles. Thus, selective reaction of a C−Br (or C−Cl) bond in the presence of a C−F bond can be achieved with the aid of a nickel/bis(oxazoline) catalyst.

The products of the stereoconvergent cross-couplings, enantioenriched tertiary α-fluoroketones, can be converted into an array of interesting organofluorine compounds.

ENTRODUCTION

Motivated by potential applications in biomedical research and other disciplines,^{[1](#page-4-0)} substantial effort has been dedicated to the development of methods for the preparation of organofluorine compounds.[2](#page-4-0) In the case of alkyl fluorides, advances have been described in the catalytic enantioselective synthesis of stereo-genic centers that bear a fluorine substituent,^{[3](#page-4-0)} particularly α to a carbonyl group. Although most studies have addressed the generation of secondary stereocenters, 4 a few reports have examined the establishment of tertiary centers; currently, the latter methods are largely limited to either cyclic or doubly activated acyclic α -fluorocarbonyl compounds.^{[5](#page-4-0),[6](#page-4-0)} To the best of our knowledge, no general catalytic enantioselective process has been discovered for the synthesis of simple tertiary α -fluorinated acyclic ketones[.7](#page-4-0)[−][9](#page-4-0)

During the past decade, we have developed an array of nickel-catalyzed asymmetric cross-coupling methods, employing racemic alkyl electrophiles as reaction partners; to date, all of the electrophiles have been secondary, with $Z = H$ (eq 1).^{10,[11](#page-4-0)}

We have now begun to explore enantioselective couplings of other families of electrophiles, beginning with geminal dihalides (eq 1; Z = F, X = halide);^{[12](#page-4-0)} such H \rightarrow F substitutions can have a dramatic impact on reactivity and/or ee.^{[13](#page-4-0)} If a suitable catalyst could achieve selective cleavage of the C−X bond, along with efficient and highly enantioselective C−C bond formation, then this would enable the catalytic asymmetric synthesis of tertiary alkyl fluorides.

In view of the high interest in the enantioselective synthesis of α -fluorocarbonyl compounds,^{[4](#page-4-0),[5](#page-4-0),[8](#page-4-0),[9](#page-4-0)} as well as the excellent functional-group compatibility of Negishi reactions, 14 we chose to examine the coupling of α -halo- α -fluoroketones with organozinc reagents. In this Article, we describe the first catalytic asymmetric cross-coupling that employs geminal dihalides as electrophiles, specifically, a nickel/bis(oxazoline)-catalyzed stereoconvergent Negishi arylation of racemic $α$ -bromo- $α$ -fluoroketones to generate tertiary α -fluorinated acyclic ketones (eq 2).

■ RESULTS AND DISCUSSION

At the time that we began our investigation in 2011, we were not aware of any precedent for selective nickel-catalyzed crosscouplings of α -halo- α -fluoro compounds. However, during the course of our studies, Ando reported diastereoselective Kumada reactions of $α$ -bromo- $α$ -fluoro- $β$ -lactams with aryl Grignard reagents,^{[15](#page-4-0)} employing a NiCl₂·glyme/bis(oxazoline) catalyst

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that we had described for enantioselective couplings of aryl Grignard reagents with α -bromoketones.^{[10c](#page-4-0)}

Unfortunately, our attempts to apply our previous methods for Kumada^{[10c](#page-4-0)} and Negishi^{[10d](#page-4-0)} arylations to the cross-coupling of the racemic α-bromo-α-fluoroketone illustrated in Table 1

Table 1. Catalytic Asymmetric Synthesis of Tertiary Alkyl Fluorides: Effect of Reaction Parameters^a

entry	variation from the "standard" conditions	ee $(\%)^b$	yield $(\%)^c$
1	none	97	68
2	no NiCl ₂ •glyme		2
3	no L*		2
4	1, instead of L*	-76	11
5	2, instead of L*	-70	8
6	3. instead of L*		2
$\overline{7}$	4, instead of L*	52	40
8	5, instead of L*		2
9	10% NiCl ₂ · glyme, 11% L*	97	49
10	1.2 equiv PhZnCl	97	51
11	PhMgCl, instead of PhZnCl		2
12	Ph ₂ Zn, instead of PhZnCl	98	56
13	r.t.	83	17
14	THF only	90	38
15	diglyme only	91	19
16	0.1 equiv H_2O	97	65
17	in air in a closed vial	94	27
	Me Me		Me Me
Ph	Ph 5 Ph N N N Рh Ph 2 Ph	Ń Ph	N Рh 3
	∩ Ph Ph … N Me(ОМе	Ph Me –NH 5	Ph $HN-Me$

 a All data are the average of two experiments. b A negative ee value signifies that the major product of the reaction is the opposite (R) enantiomer. ^cThe yields were determined through analysis by ¹⁹F NMR spectroscopy, with the aid of an internal standard.

were unsuccessful (<2% yield). However, through the appropriate choice of reaction parameters, we were able to achieve the desired α -arylation and to generate the tertiary alkyl fluoride with very good enantioselectivity (97% ee; Table 1, entry 1). NiCl₂·glyme and bis(oxazoline) L^* are commercially available and air-stable.

Table 1 provides information on the impact of various reaction parameters on the efficiency of this catalytic asymmetric synthesis of tertiary alkyl fluorides. In the absence of $NiCl₂$. glyme or of ligand L*, essentially no carbon−carbon bond formation is observed (entries 2 and 3). The cis phenyl substituent in the 5 position of the oxazolines plays an important role in enantioselectivity and yield (entry 4), as does the substitution on the one-carbon linker that bridges the oxazolines (entries 5 and 6). A variety of pybox and 1,2-diamine ligands^{[10b,d,e](#page-4-0)} furnish inferior results (e.g., entries 7 and 8). The use of less catalyst or less nucleophile leads to a modest decrease in yield, although

no erosion in ee (entries 9 and 10). Under our optimized conditions, PhMgCl is not a useful coupling partner (entry 11), whereas good enantioselectivity but less-efficient cross-coupling is observed when $Ph₂Zn$ serves as the nucleophile (entry 12). Conducting the Negishi reaction at room temperature or with a single solvent causes a small drop in ee and a substantial loss in yield (entries 13−15). The presence of water (0.1 equiv) has essentially no impact on the course of the reaction (entry 16), whereas running the reaction under air results in decreased yield (entry 17).

Our optimized conditions can be applied to the catalytic asymmetric Negishi arylation of a variety of racemic α-bromo- α -fluoroketones, furnishing tertiary alkyl fluorides in generally good ee $(Table 2)$.^{[16](#page-4-0)} The R group of the ketone can vary in

 a All data are the average of two experiments. b Yield of purified product.

size, although a lower ee is observed with a bulky isopropyl substituent (entries 1−6). High enantioselectivity is typically obtained whether the aromatic group (Ar) is para-, meta-, or ortho-substituted, and whether it is electron-rich or electronpoor (entries 7−16); we have not previously observed high ee in related nickel-catalyzed cross-couplings with ortho-substituted Ar groups (entries 14 and 15; previously: $\leq 75\%$ ee).^{[10c](#page-4-0),[d,17](#page-4-0)} Functional groups such as an olefin, alkyl chloride,^{[18](#page-4-0)} aryl methyl ether,^{[19](#page-4-0)} and aryl fluoride²⁰ are compatible with the reaction conditions.

The scope of this method for the catalytic asymmetric synthesis of tertiary alkyl fluorides is also fairly broad with respect to the nucleophile (ArZnCl; Table 3).^{[21](#page-4-0)} Thus, para- and

^aAll data are the average of two experimens. ^bYield of purified product.
^CReaction temperature: –20 °C Reaction temperature: −20 °C.

meta- (but not ortho-) 22 substituted arylzinc reagents are suitable cross-coupling partners, furnishing the desired α -fluoroketone in good ee. Electron-rich as well as electron-poor nucleophiles can be employed; in the case of the latter, a reaction temperature of −20 °C, rather than −25 °C, is optimal. A silyl ether, aryl chloride, and aryl bromide^{[23](#page-5-0)} are compatible with the coupling conditions.

This method for the selective cross-coupling of a geminal dihalide electrophile is not limited to α -bromo- α -fluoroketones. Under similar conditions, a racemic α -chloro- α -fluoroketone also reacts with an arylzinc reagent to generate a tertiary alkyl fluoride in good ee (eq 3). In contrast, in the case of related cross-couplings, α -chlorocarbonyl compounds were not suitable reaction partners.[10c](#page-4-0)−[e](#page-4-0)

The enantioenriched organofluorine cross-coupling products can be derivatized with high stereoselectivity. For example, nucleophilic additions of allyl and aryl nucleophiles to the carbonyl group proceed with >20:1 dr to generate densely functionalized adducts (eqs 4 and 5).^{24,[25](#page-5-0)}

Moreover, regioselective Baeyer−Villiger oxidation of the enantioenriched α -fluoroketones can be achieved, thereby providing access to either an alkyl ester or an aryl ester by controlling the relative migratory aptitudes of the ketone sub-stituents (eqs 6 and 7).^{26−[28](#page-5-0)} This oxidation thereby enables, to

our knowledge, the first asymmetric synthesis of acylated fluorohydrins (eq 6),^{[29](#page-5-0)} as well as an indirect method for the catalytic enantioselective synthesis of tertiary α -fluoroesters (eq 7) and amides. 3

■ CONCLUSIONS

In summary, we have developed the first catalytic asymmetric cross-coupling method that employs geminal dihalides as electrophiles. Specifically, we have established that nickel/ bis(oxazoline)-catalyzed stereoconvergent Negishi reactions of racemic α -halo- α -fluoroketones provide access to enantioenriched tertiary alkyl fluorides, thereby complementing earlier catalytic asymmetric methods for the synthesis of organofluorine compounds, which have typically focused on the generation of secondary alkyl fluorides. The α -fluoroketones that are produced in these Negishi couplings can be transformed into a variety of interesting families of organofluorine target molecules. Additional investigations of catalytic enantioselective cross-couplings of alkyl electrophiles are underway.

■ ASSOCIATED CONTENT

6 Supporting Information

Experimental procedures and compound characterization data. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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(17) A ketone in which the aromatic group (Ar) was mesityl did not undergo cross-coupling under our standard conditions.

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(24) The stereochemistry of each product (eqs [4](#page-3-0) and [5\)](#page-3-0) has been determined by X-ray crystallography (see the [Supporting Informa](#page-4-0)[tion](#page-4-0)). Reduction of the ketone with NaBH₄ also proceeded with high diastereoselectivity (>20:1) and in good yield (>99%). However, we have not yet been able to establish the relative stereochemistry of the major diastereomer.

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