Recent Advances in C-H Fluorination

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Recent Advances in C-H Fluorination

Aijun Lin,3 C. Bryan Huehls3 and Jiong Yang*

Incorporation of fluorine atom(s) in organic compounds is often desirable in the discovery of new pharmaceuticals, agrochemicals, and materials. However, development of transformations to incorporate fluorine is challenging because of its highly electronegative nature. Recent advances in transition metal catalysis have allowed new approaches to C-F bonds. Herein we highlight progresses in the synthesis of C-F bonds in the context of C-H functionalization, which is arguably the most efficient approach to incorporate fluorine since it obviates the need of pre-functionalization of organic compounds.

Organofluorine compounds occupy a unique place in organic chemistry: despite fluorine being the 13th most abundant element in the Earth's crust, only a handful of organofluorine natural products are known.1 However, organofluorine compounds have been often used as pharmaceutical,2 agrochemical,3 material,4 and positron-emission tomography (PET) imaging agents.5 Indeed, 20% of modern pharmaceuticals and 30% of agrochemicals contain fluorine. This popularity of organofluorine compounds has been attributed to the improved pharmacological properties that fluorine may provide, such as enhanced thermal and metabolic stability, bioavailability, lipophilicity, and overall biological activity. These effects can be traced to the small size and the strong electron-withdrawing nature of fluorine, the most electronegative element of the periodic table. However, the very electronegative nature of fluorine also makes it challenging for chemical synthesis of C-F bonds due to the high hydration energy of fluoride, strong metal-fluorine interaction, and the highly polarized nature of bonds to fluorine.6 Despite the challenge, recent development in organo- and transition metal-catalysis has allowed new methods to fluorinate organic compounds.7 Herein we highlight progresses in the synthesis of C-F bonds in the context of C-H functionalization, which is arguably the most efficient approach to C-F bonds since it obviates the need of pre-functionalization of organic compounds.

The past decade witnessed the rapid development of transition metal-catalyzed C-H functionalization as a powerful tool for organic synthesis, with heteroatom directed C-H activation as a commonly used strategy for regioselection. In 2006, Sanford and co-workers described the first Pd-catalyzed aryl and benzyl C-H fluorination of 2-phenylpyridine and 8-methylquinoline derivatives with N-fluoro-trimethylpyridinium tetrafluoroborate or its trimethyl congener as the electrophilic fluorine source under microwave conditions (Scheme 1, eq. 1).8

For C(sp2)-H fluorination, 2,6-difluorinated products were formed except when ortho- or meta-substituted 2-phenylpyridines were used, which led to monofluorinated products. While quinoline and pyridine were used as the directing group in this pioneering study, other N-heterocycles, such as quinoxaline, pyrazole, benzo[d]oxazole, and pyrazine, were found to be equally effective for directed ortho-fluorination under similar conditions.9 Building upon their success on electrophilic C-H fluorination, Sanford and co-workers also described the first Pd-catalyzed nucleophilic C-H fluorination in which 8-methylquinoline derivatives were fluorinated at the benzyl position by Pd(OAc)2 and PhI(OPiv)2, with AgF as the source of the nucleophilic fluoride (Scheme 1, eq. 2).10

Yu and co-workers described ortho- C-H fluorination of triflamide-protected benzylamines with N-fluoro-2,4,6-trimethylpyridinium triflate using Pd(Ott)2 as the pre-catalyst and N-methylpyrrolidinone (NMP) as an essential additive.11 2,6-Difluorinated products were formed unless the ortho- or meta- position of the substrates had been substituted (Scheme 2, eq. 3). Hypothesizing that the slow displacement of the monofluorinated products from the PdIII center caused
difluorination of the substrates, Yu and co-workers successfully
developed a method for C-H monofluorination using the amide
derivatives of benzoic acid and 2,3,5,6-tetrafluoro-4-
(trifluoromethyl) aniline as the substrates (Scheme 2, eq. 4).

The monoselectivity of the reaction was attributed to the
weaker coordination of the L-type acidic amide directing group,
which allowed rapid dissociation of the monofluorinated
products.

Scheme 2 Pd-Catalyzed C(sp^2)-H fluorination by Yu

While a mechanistic pathway involving reductive
elimination of Pd^{IV}(R)(F) intermediates appears to be operative
in the Pd-catalyzed C-H fluorination reactions,
Daugulis and co-workers hypothesized that the amide derivative of 8-
aminoquinoline, which they introduced as a directing group for
Pd-catalyzed C-H arylation,
might promote Cu-catalyzed C-H
fluorination through Cu^{III} intermediates.
Indeed, ortho-fluorination of 8-aminoquinoline benzamide was effected with
AgF and NMO using CuI as the catalyst (Scheme 3, eq. 5).
Selective mono- or difluorination could be achieved by
adjusting the loading of the reagents and the reaction time.

Scheme 3 Cu-Catalyzed C(sp^2)-H fluorination by Daugulis

The hetero atom-directed C-H functionalization provided a
powerful approach for fluorination of organic compounds, but
the limitations of this approach are also obvious. For example,
in addition to requiring the presence of a directing group, the
Pd-catalyzed C-H fluorination of 2-phenylpyridine incorporated
fluorine into the phenyl only \textit{vide supra} and fluorination of
pyridine could not be achieved. Because of the important role
of fluorinated heterocycles in pharmaceuticals and
agrochemicals, there is an urgent need of new approaches to
fluorinate heterocycles. Inspired by the classic Chichibabin
reaction of pyridine and NaNH\textsubscript{2} to form 2-aminopyridine, Fier
and Hartwig developed a practical approach for C-H
monofluorination of pyridine by AgF\textsubscript{2} with exclusive site
selectivity at the C-H bonds adjacent to nitrogen under mild
conditions (Scheme 4, eq. 6). For example, reaction of 2-
phenylpyridine with AgF\textsubscript{2} in MeCN gave 2-fluoro-6-
phenylpyridine as the only fluorinated product in 88% yield.
This excellent site-selectivity was rationalized by a mechanism
of initial N-coordination of AgF\textsubscript{2} with pyridine followed by
addition of the Ag-F bond across the \pi system to give an
amido-silver(II)-fluoride intermediate, and abstraction of
hydrogen by a second equivalent of AgF\textsubscript{2} to give the product.
The reaction also was found to be effective for C-H
monofluorination of other 6-membered N-heterocycles, such as
quinolines, pyrazines, pyrimidines, and pyridazines.

Scheme 4 Monofluorination of pyridine by AgF\textsubscript{2}

The challenge of selective C(sp^3)-H fluorination without a
directing group lies in identifying a catalytic system that is both
sufficiently reactive and predictably selective. In an elegant
study, Groves and co-workers discovered that the manganese
porphyrin system Mn(TMP)Cl could be used for regioselective
C(sp^3)-H fluorination by silver fluoride/tetrabutylammonium
fluoride trihydrate with iodosylbenzene as the oxo-transfer
agent. The unique reactivity of the system was demonstrated
in the fluoridation of a number of polycyclic natural/unnatural
compounds. For example, among the 26 unactivated C(sp^3)-H
bonds of sclareolide, only those at C2 and C3 reacted to give
methylene-fluorinated products (Scheme 5, eq. 7). The
proposed reaction mechanism involved abstraction of hydrogen
by the \textit{in situ} formed oxomanganese species O=Mn^{V}(TMP) and
capture of the resulting C-centered radical with fluorine by a
\textit{trans}-Mn^{IV}(TMP)F\textsubscript{2} species. A related manganese-salen system
was also reported by the same research group for selective
benzyl C-H fluorination with triethylamine trihydrofluoride
(TREAT·HF) as the source of nucleophilic fluoride (Scheme 5,
eq. 8). Even though less efficient, potassium fluoride in the
presence of 18-crown-6 could also be used as the source of
fluorine.
Scheme 5 Mn-catalyzed C(sp³)-H fluorination

The electrophilic Pd(II)-sulfoxide system developed by White is a powerful tool for C(sp³)-H functionalization. Recently, Braun and Doyle demonstrated this catalytic system to be effective for allylic C-H fluorination as well. Their optimized reaction conditions include Pd(OTf)₂·1,2-bis(phenylsulfinyl)ethane (L₁), Lewis acidic co-catalyst (salen)CrCl, and benzoquinone as the oxidant, with the inexpensive Et₃N·3HF as the nucleophilic fluoride source. (Scheme 6) The allyl fluoride product was formed with high branch:linear regioselectivity, and in good yields.

Scheme 6 Pd-catalyzed allylic C(sp³)-H fluorination

A multicomponent system of CuI-bis(imine) (L₂) complex, KB(C₆F₅)₃, KI, and N-hydroxyphthalimide (NHPI) was reported by Lectka and co-workers for aliphatic, allylic, and benzylic monofluorination with Selectfluor (Scheme 7, eq. 10). A radical mechanism was proposed with NHPI, known to form the phthalimide N-oxygen (PINO) radical in situ in the presence of redox active metals, serving as a radical initiator. Evidence to support this hypothesis included reaction inhibition by the radical trapping agent 2,2,6,6-tetramethylpiperidin-1-yloxy (TEMPO). Interestingly, in a related study, Inoue and co-workers found that NHPI alone was capable of initiating benzylic C-H fluorination with Selectfluor, but synthetically useful yields for the benzylic and aliphatic C-H fluorination required N,N-dihydroxypyrromellitimide (NDHPI) to be used (Scheme 7, eq. 11). A recent report by Lectka and co-workers showed that benzylic C(sp³)-H monofluorination could also be effected with Selectfluor using Fe(acac)₃ as the catalyst (Scheme 7, eq. 12).

Scheme 7 Some radical C(sp³)-H fluorination

A transition metal-free photochemical method also was described recently for benzylic C(sp³)-H fluorination. In this elegant study, Chen and co-workers reported that a wide range of substrates underwent benzylic C(sp³)-H fluorination with Selectfluor using 9-fluorenone as the catalyst (Scheme 8, eq. 13). No special equipment was necessary as the reaction could be carried out in regular glassware with household CFL as the source of visible light. Under similar conditions, the challenging C(sp³)-H benzylic difluorination could be effected with Selectfluor II using xanthone as the catalyst (eq. 14).

Scheme 8 Photolytic benzylic C(sp³)-H fluorination

Chemists have gone a long way developing new methods for synthesis of C-F bonds. Particularly, recent advances in transition metal-catalysis and radical reactions have allowed innovative approaches for C-H fluorination. However, significant challenges remain. Some of these challenges are inherent in C-H activation, such as developing new directing groups for mild and regioselective C-H activation, new catalytic systems with unique selectivity, and new reaction conditions with broader substrate scope. Further challenges have to be met for effective C-H fluorination, such as...
developing economical electrophilic fluorine sources, fluorination processes based on nucleophilic fluoride, and identifying operationally simple reaction conditions to address the special needs in synthesizing $^{18}$F PET imaging agents, which would benefit from a rapid reaction (due to the short, 110 min half-life of $^{18}$F) with dilute aqueous [$^{19}$F]fluoride (produced by proton bombardment of $^{18}$O-enriched water).$^{14,26}$ These challenges are expected to stimulate further efforts to invent innovative approaches to incorporate fluorine into organic compounds.

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Highlighted in this article are recent advances in C(sp²)-H and C(sp³)-H fluorination reactions.