

Exploiting a "Beast" in Carbenoid Chemistry: Development of a Straightforward Direct Nucleophilic Fluoromethylation Strategy

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S Supporting Information

ABSTRACT: The first direct and straightforward nucleophilic fluoromethylation of organic compounds is reported. The tactic employs a "fleeting" lithium fluorocarbenoid (LiCH₂F) generated from commercially available fluoriodomethane. Precise reaction conditions were developed for the generation and synthetic exploitation of such a labile species. The versatility of the strategy is showcased in ca. 50 examples involving a plethora of electrophiles. Highly valuable chemicals such as fluoroalcohols, fluoroamines, and fluoromethylated oxygenated heterocycles could be prepared in very good yields through a single synthetic operation. The scalability of the reaction and its application to complex molecular architectures (e.g., steroids) are documented.

The presence of fluorine in an organic framework profoundly influences the physicochemical properties, thus making the resulting compounds unique and highly valuable scaffolds across the chemical sciences. Such behavior is advantageously exploited in drug discovery not only to modulate critical parameters, including pharmacokinetics and pharmacodynamics, but also to design radiopharmaceuticals for positron emission tomography.^{1–3} Recent achievements in fluoroalkylation chemistry have culminated nowadays in established and robust methodologies for installing trifluoromethyl (CF₃) or difluoromethyl (CF₂) units, mainly via the generation of the corresponding radicals or carbenes or, alternatively, by means of other electrophilic reagents.^{4,5} Moreover, compared with trifluoro- or difluoromethylation, monofluoromethylation strategies still remain a formidable challenge. The direct introduction of a fluoromethyl unit holds great importance because of the isosteric correspondence of the CH₂F group to a CH₃ group,⁶ as showcased in some fluoromethylated drugs reported in Figure 1. As for nucleophilic fluoroalkylations, that is, the transfer of fluoroalkyl groups to an electrophile by a fluorinated carbanion equivalent, important aspects concerning the thermal and chemical stability of the intermediates were recently disclosed.⁷ Hu reported the so-called "negative fluorine effect" (NFE) to highlight the influence of fluorine on the thermal stability and nucleophilic fluoroalkylation reactivity of fluorinated carbanions.^{8,9} Conceptually, a selective nucleophilic monofluoromethylation could be accomplished through two main strategies: (a) direct transfer of a

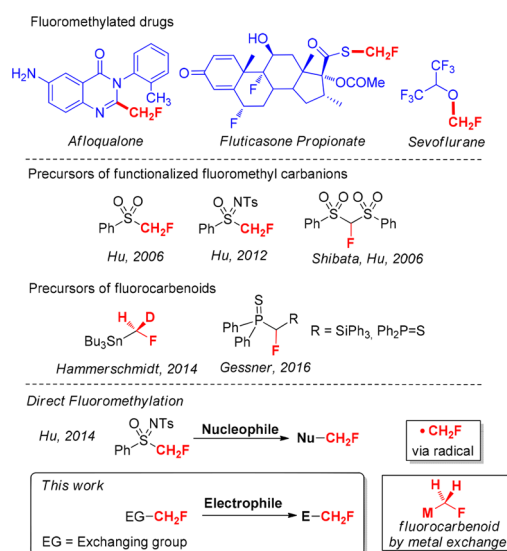


Figure 1. State of the art in monofluoromethylation.

"CH₂F" moiety and (b) transfer of the fluorinated group linked to a suitable auxiliary that must be removed at the end of the sequence.⁹ To date, the limited chemical stability of fluoromethyl carbanions has been efficiently overcome through the stabilizing effect displayed by strong electron-withdrawing functionalities (Figure 1). Accordingly, fluoromethylated sulfones, sulfoximines, and bis(phenylsulfonyl) could be advantageously employed as effective agents (Olah, Hu, and Shibata).^{10–13}

Hu recently succeeded in directly fluoromethylating O, S, N, and P nucleophiles through CH₂F radical species generated from fluoromethylated sulfoximines (Figure 1).¹⁴ Unfortunately, such methodology was not suitable for C nucleophiles, thus leaving undisclosed the development of a direct C–CH₂F bond formation strategy.¹⁵ In this context, the availability of a reagent that can introduce the CH₂F group in one direct synthetic operation would be highly desirable. Conceptually, the ideal generation of a putative M–CH₂–F reagent—i.e., a carbenoid (M = metal)—would represent de facto a straightforward synthetic tactic toward the immediate one-pot functionalization of a given electrophile (Figure 1). In this context, very recently

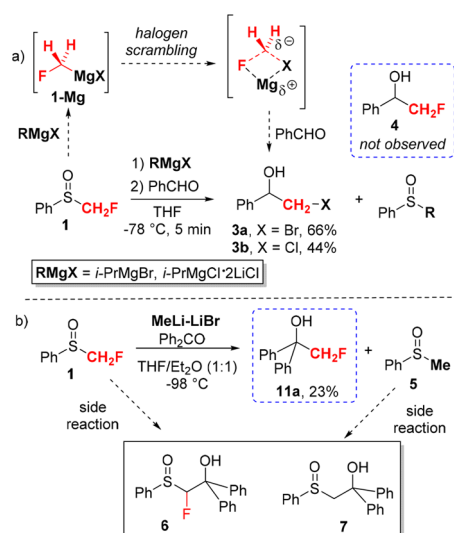
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70 Gessner succeeded in isolating and characterizing Li, Na, and K
71 fluorocarbenoids stabilized by electron-withdrawing groups
72 (Figure 1). In this interesting report,¹⁶ the authors stated that
73 “Li/F systems are still regarded as the ‘beast’ in carbenoid
74 chemistry. This is due to their extreme sensitivity and reactivity
75 connected with the facile LiF elimination typically at temper-
76 atures as low as $-78\text{ }^{\circ}\text{C}$. Hence, applications are extremely
77 limited.” Additionally, a seminal contribution by Hammersch-
78 midt¹⁷ demonstrated the high configurational stability of a chiral
79 lithiated, fluorinated deuterocarbenoid (LiCHDF) as well as the
80 dramatic chemical instability of this species even at very low
81 temperature ($-95\text{ }^{\circ}\text{C}$), thus limiting its synthetic potential.

82 Moved by this challenge and inspired by Hammerschmidt’s
83 report, we embarked on a research endeavor aimed at exploiting
84 the reactivity of Mg and Li fluoromethyl carbenoids. The study
85 commenced by considering fluoromethyl sulfoxide **1** and
86 fluoriodomethane (**2**) as simple potential precursors of
87 fluoromethylating reagents via metalation chemistry. Upon
88 treatment of **1** with a Grignard reagent (*i*-PrMgBr or *i*-
89 PrMgCl·2LiCl) followed by external electrophilic trapping with
90 benzaldehyde, bromohydrin **3a** or chlorohydrin **3b** was
91 obtained as the main reaction product (Scheme 1). Surprisingly,

Scheme 1. Metalation of Fluoromethyl Sulfoxide 1



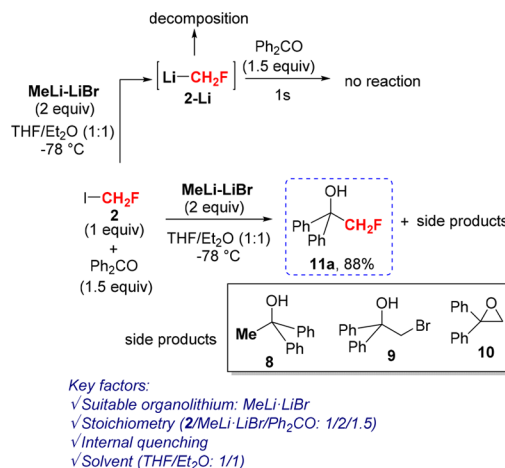
92 after extensive optimization, the expected fluoromethylated
93 adduct **4** could not be formed.¹⁸ Presumably, adducts **3** were
94 formed as a consequence of halogen scrambling at the level of
95 magnesium fluorocarbenoid **1-Mg** (Scheme 1a).

96 Effectively, the attempted nucleophilic displacement of **1** with
97 MgX₂ and LiX (X = Br, Cl) in a THF solution resulted in full
98 recovery of starting material **1**, thus making such a possibility
99 unlikely. Analogous F/I halogen scrambling was noticed by
100 Charette with electrophilic zinc fluorocarbenoids.¹⁹

101 Switching to a lithium reagent (MeLi·LiBr) was beneficial:
102 pleasingly, with a 1:1 THF/Et₂O mixture at $-98\text{ }^{\circ}\text{C}$, the desired
103 fluoromethylated adduct **11a** could be isolated in 23% yield
104 (Scheme 1b). Further attempts to improve the reaction
105 performance were elusive since collateral products (**6** and **7**)
106 resulting from the reaction between **1** and **5** were detected
107 (Scheme 1b). Taking into consideration the well-established
108 applicability of dihalomethanes as carbenoid precursors,²⁰ we
109 deemed commercially available **2** to be a convenient source for
110 the MCH₂F reagent.²¹ In striking contrast to sulfoxide **1**, both *i*-

PrMgCl·2LiCl and *i*-PrMgBr were ineffective in promoting the
111 metalation of **2**, and only attack of the Grignard to the
112 electrophile was observed.¹⁸ After extensive reaction tuning, **2**
113 was identified as the optimal substrate for lithiation,¹⁸ and the
114 desired fluorohydrin **11a** was obtained in an excellent 88% yield
115 (Scheme 2). Crucial factors enabling the success of the reaction
116 s2

Scheme 2. Reactivity of Fluoriodomethane (2)



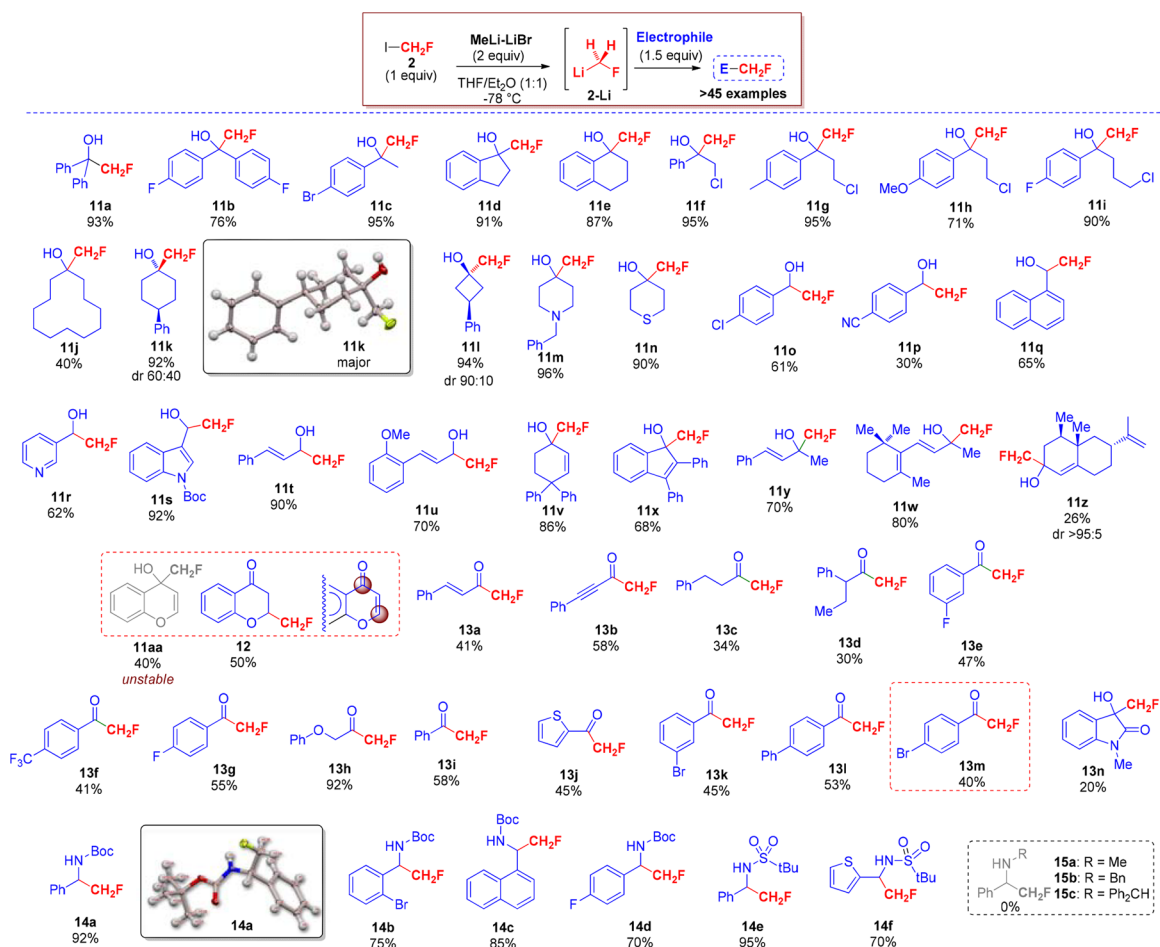
117 under Barbier-type conditions (i.e., internal quenching) were (1)
118 the use of MeLi·LiBr as the lithiating agent at $-78\text{ }^{\circ}\text{C}$, (2) the use
119 of 1:1 (v/v) THF/Et₂O as the medium, and (3) a precise 2/
120 MeLi·LiBr/electrophile stoichiometry of 1/2/1.5.¹⁸

121 During the optimization study, side products **8** and **9** were
122 found in the crude reaction mixture, likely as a consequence of
123 halogen scrambling induced by LiBr or direct insertion of a
124 carbene into the C=O bond of the electrophile. The amounts of
125 **8** and **9** were strictly dependent on the reaction conditions and
126 possibly on the chemical stability of **2-Li**.²² As expected, lithium
127 fluorocarbenoid **2-Li** was found to be extremely reactive, fully
128 decomposing under external trapping conditions even when the
129 electrophile was added after only 1 s (Scheme 2).¹⁸ Similarly,
130 polar solvents such as THF and higher temperatures enhanced
131 the decomposition of **2-Li**. In fact, when the temperature was
132 increased to $-40\text{ }^{\circ}\text{C}$ under internal quenching conditions,
133 epoxide **10** was formed in 10% yield compared with **11a** in 72%
134 yield. The use of toluene or Et₂O as the solvent was ineffective.¹⁸

135 Remarkably, our protocol could be conveniently applied to a
136 wide range of electrophiles, including carbonyls, imines, and
137 Weinreb amides (Scheme 3). Useful β -fluoroalcohols **11a–z**
138 were obtained in good to excellent yields with high chemo-
139 control, as showcased by adducts **11c**, **11o**, and **11f–i** featuring
140 additional potentially exchangeable halogens.²³ Carbocyclic and
141 heterocyclic enolizable ketones furnished the corresponding
142 fluoromethylated products **11j–n** in very good yields.²⁶ The
143 reaction proceeded with stereocontrol in the case of a small-sized
144 cyclic ketone, providing fluoroalcohol **11l** in 94% yield with
145 90:10 dr. Aromatic and heteroaromatic aldehydes provided
146 fluorohydrins **11o–u** in good yields in almost all cases, with the
147 exception of **11p** (because of its volatility), where chemocontrol
148 in the presence of a nitrile electrophilic functionality was fully
149 preserved. α,β -Unsaturated carbonyls reacted chemoselectively
150 in 1,2-fashion to give fluorohydrins **11t–z** without affecting the
151 chemical integrity of the double bond.

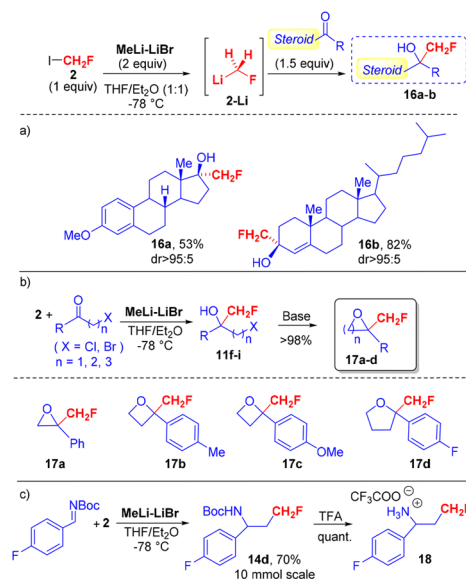
152 Surprisingly, the use of chromanone led to both 1,2- and 1,4-
153 addition products (**11aa** and **12**, respectively). However, **11aa**

Scheme 3. Scope of the Direct Nucleophilic Fluoromethylation Strategy



154 was found to be highly unstable, and only adduct **12** was isolated
 155 in 50% yield (Scheme 3).²⁴ Less electrophilic Weinreb amides
 156 were excellent acylating agents for LiCH₂F, thus enabling direct
 157 access to α -fluorinated ketones **13a–l**. Unsaturated motifs
 158 (alkenes and alkynes) were perfectly tolerated in terms of
 159 chemocontrol (i.e., **13a** and **13b**), as were heterocycles and
 160 halogenated aromatics (i.e., **13j** and **13k**). The special ketone
 161 isatin was fluoromethylated to give adduct **13n** in a lower yield of
 162 20% due to its low solubility in the reaction medium. To further
 163 benchmark the methodology, aromatic and heteroaromatic
 164 imines were employed as electrophiles to obtain highly valuable
 165 β -fluoroamines **14a–f**.^{25,26} The process requires imines bearing
 166 electron-withdrawing N substituents (Boc, *t*-BuSO, *t*-BuSO₂),
 167 whereas the use of *N*-alkyl- or *N*-benzylimines was unsuccessful
 168 (i.e., **15a–c**; Scheme 3). Biologically relevant and complex
 169 scaffolds such as 3-*O*-methylsterone and 4-cholesten-3-one
 170 efficiently underwent the transformation (Scheme 4a). Remark-
 171 ably, the reaction of fluorocarbenoid **2-Li** occurred with superb
 172 stereoselectivity, furnishing **16a** and **16b** as single stereoisomers.
 173 The functional group compatibility was pivotal for the design of
 174 an unprecedented two-step access to α -fluoromethylated
 175 oxygenated heterocycles such as the challenging epifluoroalcohol
 176 **17a**, fluoromethylated oxetanes **17b** and **17c**, and tetrahy-
 177 **drofuran** **17d** by the chemoselective intramolecular cyclization
 178 (Scheme 4b). Next, given the importance of β -fluoroamines, a 10
 179 mmol preparation of **18** was achieved using a two-step sequence
 180 involving direct fluoromethylation of *N*-Boc-imine followed by
 181 acidic removal of the Boc group (Scheme 4c).

Scheme 4. Further Applications of the Direct Fluoromethylation Strategy



In conclusion, a novel one-pot strategy for direct nucleophilic
 fluoromethylation has been developed. This method overcomes
 the drawbacks associated with the use of auxiliary groups
 requiring proper removal after introduction of the fluorinated
 fragment. This work demonstrates that the fleeting carbenoid

187 fluoromethyl lithium can be efficiently exploited for synthetic
188 purposes. Through fine-tuning of the reaction conditions it is
189 possible to avoid decomposition of the intermediates, allowing
190 the reaction to be carried out with various electrophiles,
191 including structurally complex molecules. We believe that this
192 work paves the way for further progress in fluoromethylation
193 strategies and fluorinated organometallics.

194 ■ ASSOCIATED CONTENT

195 ● Supporting Information

196 The Supporting Information is available free of charge on the
197 ACS Publications website at DOI: 10.1021/jacs.7b07891.

198 Procedures, optimization tables, and spectral data (PDF)
199 Crystallographic data for 14a (CIF)
200 Crystallographic data for 11k (CIF)

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209 Notes

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