

Am Chem Soc. Author manuscript; available in PMC 2013 January 25.

Published in final edited form as:

J Am Chem Soc. 2012 January 25; 134(3): 1494–1497. doi:10.1021/ja211422g.

A New Reagent for Direct Difluoromethylation

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Abstract

Molecular scaffolds containing alkylfluorine substituents are desired in many areas of chemical research from materials to pharmaceuticals. Herein, we report the invention of a new reagent $(Zn(SO_2CF_2H)_2, DFMS)$ for the innate difluoromethylation of organic substrates via a radical process. This mild, operationally simple, chemoselective, and scalable difluoromethylation method is compatible with a range of nitrogen-containing heteroarene substrates of varying complexity as well as select classes of conjugated π -systems and thiols. Regiochemical comparisons suggest that the CF_2H radical generated from the new reagent possesses nucleophilic character.

The difluoromethyl group (CF₂H) is an intriguing structural motif that has great potential in the areas of pharmaceuticals, agrochemicals, and materials. ^{1–5} In the area of medicinal chemistry, the CF₂H unit is of special interest for its use in isostere-based drug design. ⁶ As a lipophilic hydrogen bond donor, CF₂H substitution offers an alternative to more traditional hydrogen bond donors while improving membrane permeability. For example, the difluoromethyl group has been utilized as a thiol mimic in the context of HCV NS3 protease inhibitors, successfully mimicking the cysteine CH₂SH P1 element in the parent compounds. ⁷ It has also been used as a hydroxamic acid hydroxyl isostere in a series of COX-2 and 5-LOX dual inhibitors, likely diminishing the metabolic toxicity expressed by some hydroxamic acids. ⁸ While several methods exist for the fluorination and alkylfluorination of organic substrates, strategies for direct difluoromethylation are less common, particularly in the context of heteroarene substrates. In this communication, the invention of a reagent to accomplish the mild, direct, scalable, and predictably selective difluoromethylation of heteroarenes and related structures is reported.

Currently, all known methods for heteroarene difluoromethylation rely on prefunctionalization through a programmed approach (Figure 1A). Perhaps the most widely known strategy involves difluorination of aldehydes using (diethylamino)sulfur trifluoride (DAST). Heteroarene-CF $_2$ H compounds may also be accessed through a coppercatalyzed cross-coupling/decarboxylation sequence, or via radical debromination from a heteroarene-CF $_2$ Br precursor. Although these methods boast impressive levels of

ASSOCIATED CONTENT

Supporting Information.

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reactivity and site selectivity, the need for methods achieving the direct transfer of the CF₂H unit has been explicitly voiced in a recent review. ¹³

The pioneering work reported by Minisci on the decarboxylative generation of alkyl radicals and subsequent addition to heteroarenes (a form of innate 14 C–H functionalization) inspired initial forays into this area. 15 Thus, difluoroacetic acid was the first CF $_2$ H source that was tested on caffeine (1) under a variety of oxidative conditions (Figure 1B). However, this strategy was unsuccessful, necessitating the exploration of alternative methods for generating CF $_2$ H-radicals. The only previously reported example of a direct radical transfer of a CF $_2$ H unit was reported by Chen and co-workers in the context of additions to olefins and alkynes (no other substrate classes reported) using non-commercial and operationally problematic HCF $_2$ I gas as a precursor. 16

In contrast, we have previously demonstrated the ease and practicality associated with the use of stable fluoroalkyl metal sulfinate complexes for the generation of trifluoromethyl radicals and subsequent addition to heteroarenes. The seemingly simple extension of this approach to difluoromethylation was without precedent and required extensive experimentation. Ultimately, after several strategies were evaluated, difluoromethanesulfonyl chloride was recognized as a convenient, commercially available starting point for the synthesis of a variety of different difluoromethylsulfinate reagents (Figure 1C). Using caffeine (1) as a model substrate, systematic variation of the metal counterion led to the identification of Zn(SO₂CF₂H)₂ (3, DFMS) as the optimum precursor for the CF₂H-radical. DFMS is an easily prepared, air stable, free flowing white powder whose structure was confirmed by X-ray crystallographic analysis and shown to exist as polymer in the solid state (see SI for more details).

With this new reagent in-hand, the scope of heteroarene difluoromethylation was evaluated on a broad cross-section of heterocyclic space. Many substrates showed good reactivity towards DFMS under the standard reaction conditions using tBuOOH in a CH₂Cl₂/H₂O solvent system, although select substrates suffered from poor conversion. In these cases, a second addition of DFMS and tBuOOH was added after 12-24 hours to drive the reaction towards completion. It was also found that TFA showed improved rate and conversion for selected nitrogen heteroarene substrates, but was not essential to achieve the desired reactivity for most cases. Pyridines (5–10, 12, 22), pyrroles (14, 15), pyrimidines (18), quinoxalines (16), pyrazines (17), xanthines (2, 19, 20), purines (21), quinoline (23), thiadiazoles (13) and pyridinones (11) are all competent participants in the described method, which is tolerant of several potentially sensitive functional groups. In most cases, substrates with multiple potential reaction sites exhibit high levels of regioselectivity, commonly producing only one observable regioisomer with C-H functionalization occurring at electron-deficient positions. ¹⁷ It should be noted that all of the compounds illustrated in Table 1 are chemical entities with heretofore unreported synthetic procedures. In addition to the heteroarene substrates listed in Table 1, other organic substrates are also reactive towards difluoromethylation by the described process (Table 2). Aromatic thiols exhibit unexpected reactivity towards the CF₂H-radical, leading to the generation of difluoromethyl thioethers. Radical difluoromethylation was also successful in the context of other electrondeficient π -systems such as α , β -unsaturated enones.

In addition to the predictable site-selectivity observed in the difluoromethylation of simple small-molecule substrates, innate C–H difluoromethylation of complex reacting partners with many potential reactive sites is also predictable. As shown in Figure 2, the site-selectivity of difluoromethylation depends on the combined electronic properties of the reacting π -system and incoming radical species. This point is illustrated through a comparison between CF₃- and CF₂H-radical additions to dihydroquinine and Chantix. In

both cases, high levels of selectivity are observed for CF_3 - and CF_2H -radical addition, in spite of the multiple, potentially reactive sites. For both dihydroquinine and Chantix, innate radical C–H trifluoromethylation takes place at the most electron-rich position within the arene rings (C7 and C5, **32** and **33**, respectively). Toonversely, difluoromethylation occurs exclusively at electron-poor sites adjacent to heteroatoms within the heteroarene rings (both at C2, **30** and **31**). The orthogonality of these two approaches is a powerful example of the ability to fine-tune an approach to alkylfluorination when considering the innate reactivity and electronic nature of the two reacting components.

Although many substrates listed above in Table 1 produce a single observable regioisomer, it is possible to access alternative substitution in some cases by judicious choice of organic solvent media. As shown in Figure 3, 4-acetylpyridine shows exceptional regioselectivity in CH₂Cl₂/H₂O, leading to exclusive formation of the C2-CF₂H product (**6-C2**). However, substituting DMSO for CH₂Cl₂ leads to significant reversal of regiochemistry, favoring the C3-CF₂H isomer (**6-C3**) in 1.5:1.0 ratio. Although C3-selectivity is modest under these conditions, the change in selectivity is significant considering that the C3-CF₂H product is inaccessible under standard reaction conditions.

In summary, a new reagent (DFMS) for direct difluoromethylation has been invented to access compounds of high value that, in almost all cases, were hitherto unknown. DFMS (3) is effective for the direct transfer of a CF $_2$ H unit to various organic substrates including heteroarenes, α , β -unsaturated enones, and aromatic thiols via user-friendly, scalable, openflask conditions. ^{19,20} Changes in regioselectivity may be promoted by varying the organic co-solvent in the context of 4-substituted pyridines. Further examination of this and other mechanistic features are ongoing and will be reported in due course. DFMS has recently been commercialized by Sigma-Aldrich (product number L510084).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We are grateful to Professor Arnold Rheingold and Dr. Curtis E. Moore for X-ray crystallographic analysis. Financial support for this work was provided by the NIH/NIGMS (GM-073949), the Uehara Memorial Foundation (post-doctoral fellowship for Y.F.) the NSF (predoctoral fellowship for R.A.R) and Pfizer Inc. (postdoctoral fellowship for R.D.B.).

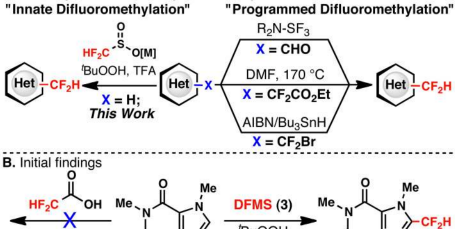
References

- 1. Smart BE. Chem Rev. 1996; 96:1555-1556. [PubMed: 11848803]
- Filler, R.; Kobayashi, Y.; Yagupolskii, LM. Organofluorine Compounds in Medicinal Chemistry and Biomedical Applications. Elsevier; Amsterdam: 1993.
- 3. Welch JT. Tetrahedron. 1987; 43:3123-3197.
- Rewcastle GW, Gamage SA, Flanagan JU, Frederick R, Denny WA, Baguley BC, Kestell P, Singh R, Kendall JD, Marshall ES, Lill CL, Lee W-J, Kolekar S, Buchanan CM, Jamieson SMF, Sheperd PR. J Med Chem. 2011; 54:7105–7126. [PubMed: 21882832]
- 5. Furuya T, Kuttruff C, Ritter T. Curr Opin Drug Disc Dev. 2008; 11:803–819.
- 6. Meanwell NA. J Med Chem. 2011; 54:2529-2591. and references therin. [PubMed: 21413808]
- 7. Narjes F, Koehler KF, Koch U, Gerlach B, Colarusso S, Steinkuhler C, Brunetti M, Altamura S, De Francesco R, Matassa VG. Bioorg Med Chem Lett. 2002; 12:701–704. [PubMed: 11844705]
- 8. Chowdhury MA, Abdellatif KRA, Dong Y, Das D, Suresh MR, Knaus EE. J Med Chem. 2009; 52:1525–1529. [PubMed: 19296694]
- 9. Markovski LN, Pahinnik VE, Kirsanov AV. Synthesis. 1973:787–789.

10. (a) Kuduk, SD.; Schlegel, K-A.; Yang, Z-Q. Quinoline Amide M1 Receptor Positive Allosteric Modulators. International Patent. PCT/US2010/060007. (b) Rodgers, JD.; Shepard, S.; Arvanitis, AG.; Wang, H.; Storace, L.; Folmer, B.; Shao, L.; Zhu, W.; Glenn, JP. N-(Hetero)aryl-pyrrolidine Derivatives of Pyrazol-4-yl-pyrrolo[2, 3-d]pyrimidines and Pyrrol-3-yl-pyrrolo[2, 3-d]pyrimidines as Janus Kinase Inhibitors. International Patent. PCT/US2010/035783.

- 11. Fujikawa K, Fujioka Y, Kobayashi A, Amii H. Org Lett. 2011; 13:5560–5563. [PubMed: 21955064]
- 12. (a) Dolbier WR, Medebielle MA–M. Tetrahedron Lett. 2001; 42:4811–4814.(b) Burkholder CR, Dolbier WR, Medebielle M. J Fluorine Chem. 2001; 109:39–48.
- 13. Hu J, Zhang W, Wang F. Chem Commun. 2009:7465-7478.
- 14. Bruckel T, Baxter RD, Ishihara Y, Baran PS. Acc Chem Res. 201110.1021/ar200194b
- (a) Minisci F, Fontana F, Vismara E. J Heterocycl Chem. 1990; 27:79–96.(b) Minisci F, Vismara E, Fontana F. Heterocycles. 1989; 28:489–519.(c) Duncton AJ. Med Chem Commun. 2011; 2:1135–1161.
- 16. Cao P, Duan JX, Chen QY. J Chem Soc, Chem Commun. 1994:737-738.
- 17. Ji Y, Brueckl T, Baxter RD, Fujiwara Y, Seiple IB, Su S, Blackmond DG, Baran PS. Proc Natl Acad Sci. 2011; 108:14411–14415. [PubMed: 21844378]
- 18. Attempts at synthesis of NaSO₂CF₂H based on Chen Q, Long Z, Faming F. Shenquing Gongkai Shuomingshu. 1999CN 1221735 A 19990707. were fruitless. Synthesis of various [M]SO₂CF₂H reagents from fluoroalkanesulfinic acid hydrazinium salts based on Nanmyo T, Kashiwaba T, Morinaka T, Kawamoto H, Inoue S, Kume T. 2010PCT Int Appl. 2010013687. was ineffective. Transferring CF₂H via reaction of fluorocarboxylic acids of the general structure [E.W.G.]–CF₂–CO₂H analogous to Forat G, Mas J-M, Saint-James L. 2002US Pat Appl. 20020042542. was unsuccessful.
- 19. A sample of DFMS left open to air on the bench-top retained its original purity and reactivity after two months.
- 20. A provisional patent on this work has been filed, application number 61565756.

A. Pathways to difluoromethylate heterocycles



C. Invention of zinc difluoromethanesulfinate (DFMS) Nal, acetone [Na]SO₂CF₂H Lil, acetone (30%) $Zn](SO_2CF_2H)_2$ Me (85%)(73%)DFMS (3) 2 [35 g scale] · air stable free-flowing powder · X-ray (polymeric crystal) 150g prepared

Figure 1. Invention of a new difluoromethylation reagent.

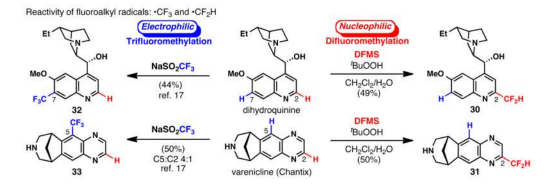
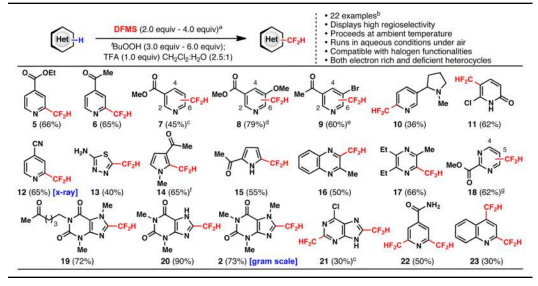


Figure 2.Regiochemical comparison of innate difluoro- and trifluoromethylations. Isolated yields after preparative HPLC purification.

Figure 3. Solvent effects on regioselectivity. ^aStandard conditions were followed. ^bSecond addition of DFMS (2.0 equiv) and *tert*-butyl hydroperoxide (3.0 equiv) was added.

Table 1Scope of C–H difluoromethylation of heteroarene substrates.



^aHeterocycle (1.0 equiv), DFMS (2.0 equiv), tert-butyl hydroperoxide (3.0 equiv), TFA (1.0 equiv), 23 °C; isolated yields of chromatographically pure products are displayed, unless otherwise noted.

 $[\]begin{tabular}{ll} b Both indole and 1-(2,4-dimethyl furan-3-yl) ethanone were unreactive, giving recovered starting material. \end{tabular}$

^cC2:C4:C6; 1.5:1.5:2.

 $^{^{}d}$ C2:C4:C6; 1:1:2.

^еС2:С4:С6; 3:1:2.

fReaction showed incomplete conversion after 12 hours, and a second addition of DFMS (2.0 equiv) and tert-butyl hydroperoxide (3.0 equiv) was added

^cHeterocycle (1.0 equiv), DFMS (3.0 equiv), *tert*-butyl hydroperoxide (4.0 equiv), 23 °C.

^gC4:C5; 4:1.

Table 2

Difluoromethylation of thiols and enones.

 $^{^{}a}$ Standard conditions were followed and yields were obtained after silica gel chromatography.

 $[^]b\mathrm{DFMS}$ (1.0 equiv) and tert-butyl hydroperoxide (1.5 equiv) was used.

 $^{^{\}it C}{\rm Second}$ addition of DFMS (2.0 equiv) and $\it tert\text{-}{\rm butyl}$ hydroperoxide (3.0 equiv) was added.

 $[\]stackrel{d}{\alpha}, \alpha, \alpha\text{-trifluorotoluene}$ was used instead of dichloromethane.

eketo:enol (1:5).