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Metal-free di- and tri-fluoromethylation of alkenes realized by visible-light-induced perylene photoredox catalysis[†]

Regioselective amino-difluoromethylation of aromatic alkenes via $C(sp^3)-CF_2H$ and $C(sp^3)-N$ bond

formation with the C=C moiety has been achieved in a single operation by visible-light photoredox

catalysis. The combination of a shelf-stable and easy-to-handle sulfonium salt, S-difluoromethyl-S-di(p-

xylyl)sulfonium tetrafluoroborate, and perylene catalysis is the key to the successful transformation. Furthermore, this noble metal-free protocol allows for the photocatalytic trifluoromethylation of alkenes.

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Introduction

The trifluoromethyl (CF_3) and difluoromethyl (CF_2H) groups have prevailed as key structural motifs of drugs and agrochemicals.¹ In particular, the CF_2H group is regarded as a unique fluorinated group because it acts as a bioisostere to hydroxyl and thiol units as well as a lipophilic hydrogen donor. Recently, practical trifluoromethylation has been realized by the action of appropriate catalysis to a variety of CF_3 sources.² In contrast, versatile strategies for the direct difluoromethylation of various carbon skeletons are still underdeveloped.³

In the past several years, visible-light photoredox catalysis with metal catalysts such as $[Ru(bpy)_3]^{2+}$ and $fac-[Ir(ppy)_3]$ (bpy = 2,2'-bipyridine, ppy = 2-phenylpyridyl) has emerged as a useful tool for radical trifluoromethylation.⁴ In particular, shelf-stable and solid sulfonium salts such as Umemoto A5 and Yagupolskii B6 reagents readily undergo single-electron transfer (SET) from photoactivated catalysts to serve as excellent CF₃ radical precursors (Fig. 1).7,8 More recently several groups, including us, have developed novel strategies for generation of the CF₂H radical from well-designed CF₂H sources such as sulfonyl derivatives (C-E) and phosphonium salts F.9 Remarkably, the subtle change in the number of fluorine atoms in the CF_2X reagents (X = F, H) causes significant differences in their chemical properties such as redox performance and stability. For example, generation of the CF₂H radical from electrophilic CF₂H sources as presented herein demands a stronger reductant compared with the CF₃ radical. In general, the Ir

photocatalyst, *fac*-[Ir(ppy)₃], is regarded as a strong **1e**-reductant when excited by visible light irradiation. But from the viewpoint of the elements strategy initiative¹⁰ and green chemistry, development of organic photocatalytic systems has attracted great interest.¹¹ However, the design of visible-light organic photoredox catalysts with stronger reduction power still leaves room for further development. In 2014, König and co-workers developed the consecutive photoinduced electron transfer (conPET) of perylene diimide, but it requires sacrificial electron donors.¹² The groups of Hawker and Miyake reported that phenylphenothiazine and diaryl dihydrophenazine serve as strong reductants, respectively.¹³ Then, simple polycyclic aromatic hydrocarbons (PAHs) attracted our attention. It is known that some PAHs exhibit high excited state energies



Fig. 1 Reductive generation of fluoroalkyl radicals by SET photoredox catalysis. Ts = p-toluenesulfonyl.



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accompanied by relatively high HOMO levels,¹⁴ suggesting that they can serve as efficient and economical photoredox catalysts without extra reductants. Herein, we disclose that perylene can serve as an excellent visible-light organic photocatalyst for amino-difluoromethylation of aromatic alkenes in a single operation. The present noble metal-free photocatalytic system also allows trifluoromethylation of alkenes.

Results and discussion

We first tackled the synthesis of a shelf-stable and easy-tohandle electrophilic CF₂H source. In 2007, Prakash, Olah and co-workers reported on the synthesis of the S-difluoromethyl-Sphenyl-S-2,3,4,5-tetramethylphenylsulfonium G,15 reagent which reacted with various hetero-atom-nucleophiles resulting in the formation of X-CF₂H bonds (X = N, O, P), but construction of a $C(sp^3)$ -CF₂H bond has not been reported. In addition, the reagent G is semi-solid and not very stable, and so decomposes by $\sim 10\%$ after three months even when stored at -20 °C. Therefore, we designed the S-(difluoromethyl)sulfonium reagent (1), where the two methyl groups of the *p*-xylyl substituents in the proximity of the sulfur atom may hinder decomposition via ionic and carbenoid reactions due to steric and electronic effects. The reagent 1 was easily synthesized according to the procedures modified from the original ones15,16 and characterized by ¹H, ¹³C and ¹⁹F NMR spectroscopy and elemental analysis. The structure of 1 was confirmed by single crystal X-ray analysis (Scheme 1).17 Compound 1 is a stable, crystalline white solid. It is worth noting that no decomposition was observed of a solid sample on a shelf for three months at ambient temperature, while a CH₃CN solution partially decomposed ($\sim 10\%$) when left for 24 h at ambient temperature. In addition, a cyclic voltammogram of 1 exhibited a broad irreversible reduction wave at around -1.70 V vs. [Cp₂Fe].

With this new reagent in hand, we explored the photoredoxcatalyzed amino-difluoromethylation of styrene **2a**, which would lead to potentially useful β -CF₂H substituted amines.¹⁸ To design an organic photocatalytic system under visible light irradiation, absorption bands in the visible light region are vital. Pale-colored PAHs with large π -conjugated systems may work as visible-light catalysts. We commenced the reaction of **2a**



Scheme 1 Synthesis and an ORTEP drawing (BF₄ anion: omitted) of 1.

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in the presence of 10 mol% perylene (absorption maxima: 434, 407 nm) in CD₃CN containing an equimolar amount of D_2O_1 , under visible light irradiation with 425 nm blue LEDs. To our delight, deuterated N-(3,3-difluoro-1-phenylpropyl)acetamide $3a-d_4$ was obtained in a 96% yield (entry 1 in Table 1). In contrast, analogous PAHs such as anthracene and pyrene were totally ineffective, presumably because of a lack of a visible absorption band, while 9,10-dimethylanthracene worked to some extent (entries 2-4). It is worth noting that the Ir photocatalyst, fac-[Ir(ppy)₃], was sluggish (entry 5). The estimated reduction potential of the photoexcited perylene was remarkably high $(-2.23 \text{ V vs. } [Cp_2Fe] \text{ in CH}_3CN$, see the ESI[†]) and even higher than that of fac-[Ir(ppy)₃] (-2.14 V (ref. 19)), which has been regarded as the most strongly reducing visible-light photoredox catalyst. The quantum yield of the emission of *perylene (94% (ref. 14)) is far superior to that of *[fac-[Ir(ppy)₃] (38% (ref. 19)), but the emissive excited state of pervlene has a very short lifetime (8.2 ns). Thus, perylene has been studied extensively as a fluorescent molecule, but less attention has been paid to it as a photoredox catalyst.²⁰ Using 1.1 equivalents of 1 decreased the yield (entry 6). The reaction did not proceed at all either in the dark or in the absence of perylene (entries 7 and 8).



Furthermore, the reactions of neutral halogen-free sulfonyl derivatives (**D** and **E**) and sulfonium reagent **G** were conducted under optimized conditions (entries 9–11). It should be noted that the sulfonium reagents (**1** and **G**) are superior to **D** and **E** in this photocatalytic reaction. The sulfonium reagent **G** also served as an effective CF_2H source (entry 11), but the stability and handling of reagent **1** were significantly improved.

We further investigated the scope of this reaction (Table 2) and found that the catalyst loading could be reduced to 5 mol%. The reaction of styrene derivatives with a variety of functional groups such as Me (2b), F (2c),¹⁷ Cl (2d), Br (2e), AcO (2f), Bpin (2g) and aldehyde (2h) groups afforded the corresponding β -CF₂H substituted amino compounds (3b-g) in 30-76% yields in a regioselective manner. To demonstrate the scalability of this organic photocatalytic system, the amino-difluoromethylation of 2e was carried out on a gram scale, and the product 3e was isolated in a 64% yield (1.1 g) (eqn (1)). It is worth noting that this reaction could be applied to a structurally more complex estrone derivative (2i) (3i: 38%). An alkene with a bulky mesityl substituent (2j) was also a substrate suitable for this transformation (3j: 52%), but 1,1-diphenylethylene (2k) afforded the substituted CF_2H -alkene (4) in a 55% yield via deprotonation from the carbocationic intermediate (vide infra). Furthermore, the system was amenable to the regioselective reaction of the internal alkenes. The reactions of *trans*- β -methylstyrene (21),

Table 1 Optimization of the photocatalytic amino-difluoromethylation of 2a^a

Entrv

	CF ₂ H reagent + Ph 2:1 2a CD ₃ CN/D ₂ O rt, 3 h 425 nm blue LED photocatalyst (PC)	Ph CF ₂ H S 3a-d ₄	
	$\begin{array}{c} \qquad \ \ \ \ \ \ \ \ \ \ \ \ \ $	fac-[lr(ppy) ₃]	
CF_2H			
reagent	PC	λ_{\max} , nm	Yield of $3\mathbf{a} \cdot d_4^{\ b}$, %
1	Pervlene	434, 407	96
1	Anthracene	376, 357	0
1	9,10-Dimethyl-anthracene	398, 377	34
1	Pyrene	334, 319	0
1	fac - $[Ir(ppy)_3]$	375 (ref. 19)	29
1	Perylene		68
1	Domilono		0

10 mal/ DC

NDCOCD₃

6^d	1	Perylene	68
7 ^e	1	Perylene	0
8	1	_	0
9	D	Perylene	0
10	Ε	Perylene	Trace
11	G	Perylene	88
^a The reaction	was carried out under N	atmosphere and irradiation of 425 nm blue LEDs at ro	om temperature using the photocatalyst (2.5 umol)

arried out under N_2 atmosphere and irradiation of 425 nm blue LEDs at room temperature using the photocatalyst CF₂H reagent (50 µmol), 2a (25 µmol), and CD₃CN (0.50 mL: containing 25 µmol of D₂O) in an NMR tube. ^b Yields were determined by ¹H NMR spectroscopy using SiEt₄ as an internal standard. ^c A 71% NMR yield of 3a- d_4 was obtained in 24 h. ^d The ratio of 1 : 2a is 1.1 : 1. ^e In the dark. LED = light-emitting diode, ppy = 2-phenylpyridyl.

trans-stilbene (2m), and 1,2-dihydronaphthalene (2n) provided the CF₂H-substituted amino products (3l-n) in 44-64% yields but as mixtures of diastereomers. Remarkably, cinnamic acid ester (20) could also be used for this transformation, resulting in the production of a CF₂H-substituted β-amino acid derivative (30: 60%). These results showed that this metal-free photocatalytic system with the CF₂H reagent 1 is useful for regioselective and simultaneous construction of $C(sp^3)$ -CF₂H and $C(sp^3)$ -N bonds onto a C=C moiety regardless of the functionalities.

Next, to examine the scope with respect to fluoroalkylation, perylene-catalyzed system was applied to trithe fluoromethylation (Scheme 2). The reaction of styrene 2a with the reagent B afforded the amino-trifluoromethylated product 5 in a 66% yield.7b Perylene also promoted chlorotrifluoromethylation of the aliphatic alkene 2p with CF₃SO₂Cl to give the product 6 in a 57% yield.²¹

To gain insight into the reaction mechanism we conducted some experiments. The reaction of 1-phenyl-2-(1phenylethenyl)cyclopropane (2q) afforded the difluoromethylated, ring-opened product 7 (23% yield), indicating the involvement of radical processes in the photocatalytic reaction

(Scheme 3). Moreover, the reaction of 2a with 1 required continuous visible light irradiation for steady conversion (see the ESI[†]), suggesting that a radical chain mechanism was not the main reaction pathway.

On the basis of the above-mentioned observations, a possible reaction mechanism for pervlene-catalyzed difluoromethylation is depicted in Scheme 4. Perylene excited by visible light irradiation (*perylene) undergoes SET to the electrophilic CF_2H reagent 1 to form the difluoromethyl radical 'CF2H via C-S bond cleavage, and the radical cation of perylene ([perylene]⁺). The very short lifetime of *perylene may be compensated by its highly emissive quantum yield to promote the SET process. Fluorescence quenching experiments support the SET process (see the ESI^{\dagger}). The generated 'CF₂H radical reacts with alkene 2 to form the adduct 8, which is oxidized by [perylene]^{\cdot +} to provide the carbocationic intermediate 8⁺. Subsequent Ritter amination²² of 8⁺ with CH₃CN/H₂O affords the aminodifluoromethylated product 3. When an α -substituted styrene $2\mathbf{k}$ is used as a substrate, deprotonation of $\mathbf{8}^+$ gives the CF₂H-alkene 4.





^{*a*} For detailed reaction conditions, see the ESI.† ^{*b*} Yields of the isolated products are lower than those before purification. The purification processes decreased the isolated yields. The diastereomer ratios (dr) were determined using ¹H NMR spectra of crude reaction mixtures. ^{*c*} 12 h. Ac = acetyl, Bpin = boronic acid pinacol ester.



Scheme 2 Perylene-catalyzed trifluoromethylation. $^{\rm a} Anhydrous$ CH_3CN was used as a solvent.









Conclusions

In conclusion, we have developed noble metal-free photocatalytic di- and tri-fluoromethylation of alkenes using a perylene catalyst. The combination of the new *S*-(difluoromethyl) sulfonium salt (1) and perylene catalysis allows for facile aminodifluoromethylation of aromatic alkenes through radical processes, for which the Ir photocatalyst works much less efficiently. Thus, the unprecedented simple synthesis of β -CF₂Hsubstituted amines from alkenes has now become feasible. Further development of perylene-catalyzed reactions is currently under way in our laboratory.

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