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A third generation of radical fluorinating agents based on *N*-fluoro-*N*-arylsulfonamides

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Radical fluorination has been known for a long time, but synthetic applications were severely limited by the hazardous nature of the first generation of reagents such as F_2 and the strongly electrophilic nature of the second generation of reagents such as *N*-fluorobenzenesulfonimide (NFSI) and Selecfluor[®]. Here, we report the preparation, use and properties of *N*-fluoro-*N*-arylsulfonamides (NFASs), a class of fluorinating reagents suitable for radical fluorination under mild conditions. Their N–F bond dissociation energies (BDE) are 30–45 kJ mol⁻¹ lower than the N–F BDE of the reagents of the second generation. This favors clean radical fluorination processes over undesired side reactions. The utility of NFASs is demonstrated by a metal-free radical hydrofluorination of alkenes including an efficient remote C–H fluorination via a 1,5-hydrogen atom transfer. NFASs have the potential to become the reagents of choice in many radical fluorination processes.

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he introduction of fluorine atoms into organic molecules significantly changes their physical, chemical, and biological properties, and is therefore very attractive for the preparation of innovative materials, agrochemicals, and pharmaceuticals¹⁻³. Moreover, ¹⁸F-labeled organic compounds are of high clinical interest as contrast agents for positron emission tomography (PET) $^{4-6}$. This situation has created a strong demand for efficient fluorination techniques. In the last 30 years, the introduction of fluorine atoms using nucleophilic and electrophilic reagents has led to remarkable advances. Radical fluorination has been known for a long time, but synthetic applications were severely limited by the hazardous nature of the first generation of reagents (Fig. 1a) such as F_2^7 , hypofluorites (ROF)⁸, and XeF_2^9 . Recently, a second generation of reagents, initially developed and optimized for electrophilic fluorination, changed dramatically that picture and radical fluorination is becoming an essential tool for selective fluorination under mild conditions (Fig. 1b)¹⁰⁻¹³. Sammis and co-workers¹⁴ proposed in 2012 that N-fluorobenzenesulfonimide (NFSI), Selectfluor®, and N-fluoropyridinum salts (NFPY), due to their low N-F bond dissociation energies (BDE), may be used for radical fluorination. This hypothesis was confirmed by the description of a radical fluorinative decarboxylation of tert-butyl peresters (Fig. 1b)¹⁴ and 2arvloxy carboxylic acids using NFSI15 as a source of fluorine atom. NFSI was also used by Zhang et al.¹⁶ for the coppercatalyzed aminofluorination of styrene, by Britton and coworkers¹⁷ for a tetra-*n*-butylammonium decatungstate-catalyzed $C(sp^3)$ -H bond fluorination, and by Lectka and co-workers¹⁸ for the aminofluorination of cyclopropanes. Following the work of Li Ag(I)-catalyzed fluorodecarboxylation on the with

Selectfluor^{®19,20}, this reagent became the most common reagent for radical fluorination processes¹¹. Using this reagent, the decarboxylative fluorination²¹⁻²⁴ has been thoroughly investigated and very recently the fluorination of tertiary alkyl halides was reported²⁵. Interestingly, the fluorinative deboronation of alkylpinacolboranes and alkylboronic acids catalyzed by Ag(I) with Selectfluor[®] was reported by Li (Fig. 1b)²⁶. Aggarwal and coworkers²⁷ reported that such a radical process involving Selectfluor[®] was a competing reaction during the electrophilic fluorination of boronate complexes. Boger and Barker²⁸ developed an Fe(III)/NaBH₄-mediated free radical Markovnikov hydrofluorination of unactivated alkenes with Selectfluor®. A related cobalt-catalyzed hydrofluorination reaction was reported by Hirova and co-workers²⁹ using a N-fluoropyridine source of atomic fluorine. Groves and co-workers^{30,31} developed recently an appealing manganese-catalyzed procedure for C-H fluorination process using the nucleophilic F^- as the fluorine source.

The second generation of radical fluorinating agents has transformed the field. However, they are often penalized by the necessity to use a transition metal catalyst and by their strong electrophilic/oxidative character. A careful look at the reaction mechanisms shows that they are frequently involved in electron transfer processes and that carbocation intermediates are generated by overoxidation processes. This was clearly demonstrated by Li and co-workers¹⁹ for the non-catalyzed fluorinative decarboxylation of peresters with Selectfluor[®] in the absence of a Ag (I) catalyst. A third generation of reagents designed to work efficiently under mild radical reaction conditions without being involved in electrophilic or electron transfer processes is clearly needed^{32,33}. We report here that *N*-fluoro-*N*-arylsulfonamides



Fig. 1 The three generations of reagents for radical fluorination. **a** Fluorination reagents of first generation. **b** Example of fluorinative decarboxylation and deboronation reactions using reagents of the second generation. **c** General structure of the *N*-fluoro-*N*-arylsulfonamides (NFASs) described in this work

(NFASs) belong to this third generation of radical fluorinating reagents (Fig. 1c). NFASs have been optimized for the catecholborane-mediated hydrofluorination of alkenes and tested in the fluorinative decarboxylation of peresters.

Results

Design of radical fluorinating agents. Initial investigations of the hydrofluorination of alkenes started with the hydroboration of 1-phenyl-1-cyclohexene **1a** with catecholborane followed by reaction with Selectfluor[®] and NFSI as fluorinating agents (Fig. 2a). Reaction with Selectfluor[®] was highly exothermic and led to decomposition of the intermediate *B*-alkylcatecholborane. No trace of the fluoride **2a** was detected by GC analysis. The reaction with NFSI afforded **2a** in 15% yield. In order to suppress undesired side reactions caused by the electrophilicity of the fluorinating agents, less electrophilic N–F reagents were tested. Benzenesulfonamides **3a–3b** and benzamide **3c** were prepared by fluorination of the corresponding amides³⁴ and tested, but all three *N*-fluoroamides proved to be inefficient (yields \leq 4%).

The disappointing results obtained with the N-fluoro-Nalkylamides 3a-3c were interpreted as a consequence of a too high BDE of the N-F bonds. In order to put this hypothesis on a quantitative basis, N-F BDEs were calculated for Selectfluor®, NFSI, and **3a-3c** in the gas phase and in DMF solution (Fig. 2b). As in previous studies on radical stabilities of N-centered radicals, geometry optimizations have been performed at the (U)B3LYP/ 6-31G(d) level of theory³⁵. Thermochemical corrections to 298.15 K have been calculated at the same level of theory using the rigid rotor/harmonic oscillator model. Improved relative energies were obtained using the (RO)B2PLYP/G3MP2Large and G3(MP2)-RAD scheme proposed by Radom and co-workers^{36,37}. The stabilities for N-centered radicals obtained from fluoramides R₂N-F have been determined with reference to fluoroamine (H₂N-F) using the isodesmic fluorine exchange reaction shown in Eq. (1).

$$R_2 N - F + \cdot NH_2 \rightarrow R_2 N \cdot + F - NH_2 \quad \Delta H_{298} = RSE(R_2 N \cdot)$$
(1)

$$BDE(R_2N - F) = RSE(R_2N \cdot) + BDE(H_2N - F)$$
(2)

The reaction enthalpies (ΔH_{298}) obtained from Eq. (1) (commonly referred to as radical stabilization energies of the substrate radicals R₂N•) can be combined with the reference value for the H₂N-F parent system $(+286.6 \text{ kJ mol}^{-1})^{38}$ to obtain N-F BDE values of the fluoroamines R_2N-F as expressed in Eq. (2). The trends in N-F BDE values are very similar at all levels of theory and also in the gas phase and in DMF solution (see Supplementary Figs. 219-221 and Supplementary Tables 4-7). For the sake of brevity we will only discuss the results obtained at the G3(MP2)-RAD level. In DMF solution the N-F BDEs of 3a, **3b**, and **3c** are calculated to be 263.0, 263.6, and 274.6 kJ mol⁻¹ (62.9, 63.0, and 65.6 kcal mol⁻¹), which is close to the N-F BDE in Selectfluor[®] (265.7 kJ mol⁻¹, 63.5 kcal mol⁻¹), but slightly higher than in NFSI (259.3 kJ mol⁻¹, 62.0 kcal mol⁻¹) (Fig. 2b). These results are in line with the fact that such N-alkylamidyl radicals are only weakly stabilized³⁵ and have been used recently for C-H chlorination, bromination, and xanthylation reactions³⁹⁻ 41.

In order to decrease the N-F BDE while maintaining enough polar effects to favor the fluorination of (nucleophilic) alkyl radicals, N-fluoro-N-arylsulfonamides (NFASs) 4 were investigated (Fig. 3). A solution phase N-F BDE of 222.3 kJ mol⁻¹ (53.1 kcal mol⁻¹) was calculated for N-Fluoro-N-(4-(trifluoromethyl)) phenyl)benzenesulfonamide 4a, supporting our assumption that N-aryl substituents should lead to lower N-F BDEs due to stabilization of the corresponding amidyl radical by delocalization onto the aromatic ring. Analyzing the impact of electronwithdrawing substituents in the anilide moiety and of electrondonating substituents in the arylsulfonyl moiety of 4a, we find neither of these to lead to large alterations in the N-F BDE values. In fact, all N-F BDE values calculated for NFASs 4a-4i cluster in the range from $220.0-226.1 \text{ kJ mol}^{-1}$ (52.6-54.0 kcal mol⁻¹), which is well below that for NFSI (62.0 kcal mol^{-1} , this value is in good accordance with the one of $63.4 \text{ kcal mol}^{-1}$ calculated recently by Xue, Cheng and co-workers)³³.

Attempts to prepare the simple *N*-fluoro-*N*-phenylbenzenesulfonamide were not successful, presumably due to side reactions involving reaction of NFSI with the electron-rich aromatic anilide moiety. After deactivation of the anilide moiety with electronwithdrawing groups (CF₃, F), the NFASs **4a**–**4i** were readily prepared by fluorination of the amides upon treatment with Cs_2CO_3 and NFSI and they could be purified by flash chromatography followed by recrystallization from heptane



Fig. 2 Initial attempts of hydrofluorination via formation of *B*-alkylcatecholboranes. **a** Hydrofluorination of 1-phenyl-1-cyclohexene (**1a**) with Selectfluor[®], NFSI, and *N*-fluoro-*N*-alkylamides **3a-3c**. **b** Solution phase (DMF) N-F bond BDEs (ΔH_{sol}) calculated at the G3(MP2)-RAD level of theory



Fig. 3 Preparation and characterization of NFASs 4a-4i. X-ray single crystal structure of 4a and 4f and solution phase (DMF) N-F BDEs calculated at the G3(MP2)-RAD level of theory

(Fig. 3). The structures of **4a** and **4f** have also been determined by X-ray crystallography and are depicted in Fig. 3. The N–F bond lengths in **4a** and **4f** (1.43 and 1.44 Å, respectively) were found to be marginally longer than the N–F bond length in NFSI (1.42 Å). The structures obtained by X-ray crystallography match well with those calculated at (U)B3LYP/6–31G(d) level (see Supplementary Figs. 1, 2 and 218).

The hydrofluorination of 1-phenyl-1-cyclohexene (1a) with NFASs 4a-4i was examined. Results are summarized in Table 1. The N-fluorosulfonamide 4a was tested first using 0.1 equivalent of DTBHN as the initiator in DMF. The fluorinated product 2a was obtained in 30% yield together with 8% of phenylcyclohexane and 10% of 1a. Since DMF is a good hydrogen atom donor, the reaction was tested in benzene and acetonitrile³⁵. However, the desired fluoroalkane 2a was not formed in these less Lewis-basic solvents (Table 1, entries 2-3). Other solvents such as Nmethylformamide, N-methyl-2-pyrrolidone, and hexamethylphosphoramide were also tested, but they provided no improvement over DMF. Using a larger amount of the radical initiator DTBHN led to a slight but reproducible increase of the yield (Table 1, entries 4-5, 45%). The other NFASs 4b-4i were tested under the optimized reaction conditions of entry 4 (0.5 equivalent DTBHN, DMF at 80 °C). NFASs bearing a second electron-withdrawing group such as 4b-4d gave lower yields (Table 1, entries 6-8). The other fluorinating agents 4e-4i provided the desired fluoride 2a in similar yields (Table 1, entries 9-13, 40-47%). For practical reasons, ease of preparation, and stability, the NFASs 4a and 4f were selected for further studies. All the reactions reported in

1) CatBH, DMA (cat.) 2) 4a-4i (3 equiv) initiator solvent 1 2 4a-4i (3 equiv) 1 a 2 a a a b b b b b b b b					
ntry	F-reagent yield [%] ^a	Initiator (equiv)	Solvent	T (°C)	2a
	4a	DTBHN (0.1)	DMF	80	30
	4a	DTBHN (0.1)	Benzene	80	-
	4a	DTBHN (0.1)	CH₃CN	80	-
	4a	DTBHN (0.5)	DMF	80	45
	4a	DTBHN (1)	DMF	80	45
	4b	DTBHN (0.5)	DMF	80	9
	4c	DTBHN (0.5)	DMF	80	23
3	4d	DTBHN (0.5)	DMF	80	30
9	4e	DTBHN (0.5)	DMF	80	41
0	4f	DTBHN (0.5)	DMF	80	47
1	4g	DTBHN (0.5)	DMF	80	43
2	4h	DTBHN (0.5)	DMF	80	41
3	4i	DTBHN (0.5)	DMF	80	40
4	4a	DTBPO (0.5)	DMF	60	47
5	Дf	DTBPO(0.5)	DMF	60	51

Table 1, except for the bulky **4g** (entry 11), were finished in less than 10 min. Therefore, running the reaction at lower temperature was attempted. At 60 °C, the use of di-*tert*-butyl peroxyoxalate (DTBPO, easily prepared by reacting oxalyl chloride with *tert*-butyl hydroperoxide in the presence of pyridine in DMF) as an initiator^{42,43} provided highly reproducible and slightly improved yields of 47% (**4a**) and 51% (**4f**) (Table 1, entries 14 and 15). The reaction is believed to be a chain process involving the reaction of the *N*-arylsulfonamidyl radical with the alkylcatecholborane to provide the desired alkyl radical. By comparison, the yield obtained with NFSI under these optimized conditions was significantly lower (29%). Beside the fluoride **2a**, small amounts of phenylcyclohexane were detected by gas chromatography in similar quantities with all three fluorinating agents. Interestingly, the presence of the starting alkene 1a was also observed but in significantly larger proportion with NFSI than with 4a and 4f (see Supplementary Table 1 and Supplementary Figs. 3–5). Since the hydroboration process takes place with complete conversion, the formation of the alkene 1a results from undesired side reactions (see Discussion).

The scope of the metal-free hydrofluorination process was examined with non-terminal alkenes **1a-1i** and NFASs **4a** and **4f** (Fig. 4a). The corresponding secondary and tertiary fluorides **2a-2i** were isolated in 48–68% yields. In many cases, the mesitylenesulfonamide **4f** gave higher yields than the benzenesulfonamide **4a**. Cyclopentene **1b** was obtained with a good *trans*-



Fig. 4 Hydrofluorination of non-terminal alkenes. **a** The reaction works efficiently with secondary and tertiary radicals derived from di- and trisubstituted alkenes, respectively. **b** The radical nature of the process is demonstrated by the ring-opening process observed with (+)-2-carene **1i**. **c** Preparation of the enantioenriched fluoride (-)-*trans*-**2b** from alkene **1b** is possible using (+)-isopinocampheylborane in the hydroboration step. Isolated yields are reported

selectivity (trans/cis 88:12). The hydrofluorination of the apinene- and nopol-derivatives 1c-1e afforded 2c-2e with high diastereoselectivities (dr \geq 95:5). The β -citronellyl benzoate 1f and the cholesteryl benzoate 1g were successfully hydrofluorinated in 53% and 48% vield, respectively. Preparation of the tertiary fluoride **2h** from 1,1'-bi(cyclohexylidene) (**1h**) worked as expected (68% yield). The presence of a free radical intermediate was demonstrated with (+)-2-carene 1i that produced the ringopening product 2i in 67% vield (Fig. 4b). Finally, based on our recent work on the enantioselective hydroazidation⁴⁴, a one-pot enantioselective hydrofluorination of 1b was performed (Fig. 4c). This one-pot procedure includes a hydroboration of the alkene with (+)-IpcBH₂, conversion to the diethyl boronate, transesterification to the B-alkylcatecholborane and a final radical fluorination. The fluoride 2b was isolated in 52% yield and 91:9 enantiomeric ratio.

Kinetic data. The rate constants for the fluorine atom transfer process between a secondary alkyl radical and NFSI, 4a and 4f were estimated using the cyclooct-1-en-5-yl radical clock⁴⁵⁻⁴⁷. The Bcyclooct-1-en-5-ylcatecholborane 5 was prepared by hydroboration of 1,5-cyclooctadiene and treated with the three fluorinating agents ([N-F] reagent = 1.2 M, three-fold excess) (Fig. 5a). The reaction with NFSI afforded a 75:25 mixture of the 5-fluorocyclooct-1-ene 6 and 2-fluorobicyclo[3.3.0]octane 7. Both 4a and 4f afforded a nearly equimolar mixture of 6 and 7. Based on this single concentration experiment and the published rate constant for the cyclization reaction $(k_c = 3.3 \times 10^4 \text{ s}^{-1} \text{ at } 80 \text{ °C})^{47}$, a rough estimation of the rate constants for fluorine transfer can be made, which for NFSI amounts to $k_{\rm F} \approx 10^5 \,{\rm M}^{-1} \,{\rm s}^{-1}$ and for the two *N*-fluoro-*N*-aryl (arenesulfonamides) 4a and 4f to $k_{\rm F} \approx 3 \times 10^4 \,{\rm M}^{-1} \,{\rm s}^{-1}$ at 80 °C (Fig. 5b). A preparative reaction was performed with 4f on 4 mmol scale. It afforded the pure fluorides 6 (31% yield) and 7 (22%) (Fig. 5a).

Remote fluorination. The hydrofluorination of terminal alkenes **8a**, **8b**, and **11** was examined next (Fig. 6). The alkene **8a** gave the fluorinated product **9a** in only 7% yield together with 7% of its isomer **9a'** resulting from a radical mediated 1,5-hydrogen shift and 50% of the corresponding alkane **10a**. Running this reaction in DMF-d₇ gave **9a** (12%) and **9a'** (11%) together with 29% of the alkane **10a** with less than 5% D-incorporation. The improved hydrofluorination/reduction ratio demonstrates that the nondeuterated DMF is probably acting as a hydrogen atom donor. However, the absence of deuterium incorporation demonstrates that other sources of hydrogen atoms are also present in the reaction mixture (including the intermediate organoborane and the fluorinating reagent itself). The methylated alkene 8b was also investigated. The presence of the methyl group was expected to favor the hydrogen atom transfer step. Indeed, product 9b' (30%) vield) became the major fluorinated product. However, a significant amount of alkane 10b (34%) was still produced. Based on these observations, it became clear that with suitable substrates, the radical hydrofluorination process can be used for efficient remote fluorination via 1,5-hydrogen atom transfer. A related remote fluorination process involving photoredox generated iminyl radicals has been recently reported²⁴. This point is demonstrated by the hydrofluorination of the terminal alkene 11 that afforded the fluoride 12 in 68% yield with an excellent trans diastereoselectivity.

Decarboxylative fluorination. The radical fluorination ability of the NFASs 4a and 4f was further tested in the decarboxylative fluorination of *tert*-butyl peresters and compared with NFSI and Selectfluor[®] (Fig. 7). This reaction, due to its non-chain nature, is not expected to be particularly efficient and recent methods have clearly surpassed this procedure^{19,23}. However, this simple reaction is very suitable to compare reagents involved in a radical mediated metal-free fluorination process. The decarboxylative fluorination of 13a using 5 equivalents of NFSI at 110 °C (sealed tube) according to the condition of Sammis, except for the use of benzene instead of benzene-d₆, gave 3-fluoropentadecane 14a in 5% yield together with a complex mixture of alkenes. This outcome is in line with the result of Li who ran the same reaction in benzene at 110 °C and did not observe the formation of the fluoride 14a. All subsequent reactions were run in chlorobenzene instead of benzene to avoid the use of a sealed reaction vessel and only 2 equivalents of the fluorinating agent were used. Under these conditions, the reaction was run with NFSI, Selectfluor[®], and NFASs 4a and 4f. NFSI provided the fluoride 14a in 7% yield, while Selectfluor[®] gave only traces of 14a (<2%, due to the high polarity of Selectfluor[®], the reaction was performed in a 1:1 mixture of chlorobenzene and DMPU). Interestingly, both 4a and 4f gave the fluoride 14a in moderate 48% and 47% yield. Similar



Fig. 5 Rate constant determination using the (Z)-cyclooct-1-en-5-yl radical clock. a Fluorination of boronate 5 with NFSI, 4a and 4f affords mixtures of mono and bicyclic fluorides 6 and 7. b Estimated rate constants for the radical fluorination



Fig. 6 Hydrofluorination of terminal alkenes. The lower nucleophilicity of primary alkyl radical slows the direct fluorination and favors hydrogen atom abstraction processes leading to remote fluorination of unactivated C-H bonds (*p*-ClBz = *para*-chlorobenzoyl)



Fig. 7 Decarboxylative fluorination of *tert*-butyl peresters. The fluorinating reagents of the second generation provide the desired fluorides in significantly lower yield than the one of the third generation due mainly to the formation of alkene side products

results were obtained with the tertiary radical derived from 13b. Reactions with 4a and 4f gave 14b in 46% and 47% yield accompanied by 35% of the alkenes. NFSI afforded only traces of the product 14b (3%) together with larger amounts of 2methyltetradec-2-ene and 2-methyltetradec-1-ene (64%) (see Supplementary Figs. 10 and 11). The cholic acid derivative 13c was examined next. In that case too, NFSI (22% yield) was inferior to 4a and 4f (39% and 33%, respectively). Sammis and co-workers¹⁴ reported a yield of 50% for this reaction when it was performed in deuterated acetonitrile on a 0.05 mmol scale.

Transition states and discussion. The higher fluorination rate observed with NFSI relative to the NFASs results is best rationalized by polar effects. The paramount importance of polar effects on the rate of radical reactions is well-established and has

been thoroughly discussed by Giese⁴⁸, Fischer and Radom⁴⁹ in their leading review articles. Polar effects have been reported to override thermodynamic effects for radical addition to alkenes⁵⁰. Recently, Xue, Cheng and co-workers have reported that NFSI has a fluorine plus detachment (FPD) value lower than that of Nmethyl-*N*-fluoro-*p*-toluenesulfonamides by 145.6 kJ mol⁻¹ (34.8) kcal mol⁻¹) in acetonitrile solution. FPD values correlated well with the reactivity of electrophilic fluorinating N-F reagents³². The free energy surfaces for the fluorination of the isopropyl radical in DMF solution have therefore been calculated at the (RO)B2PLYP/G3MP2large level for NFSI, 4a, and 4f. The calculations show slightly lower barriers for NFSI than for 4a $(\Delta G^{\neq}_{298} = +46.1 \text{ vs.} +51.3 \text{ kJ mol}^{-1})$ and a somewhat higher barrier for 4f ($\Delta G_{298}^{\neq} = +56.7 \text{ kJ mol}^{-1}$). The transition states (TSs) for NFSI and 4f are depicted in Fig. 8a, b, respectively. They are characterized by long C-F (2.32-2.33 Å) and short N-F



Fig. 8 Calculated transition states for the fluorination of the isopropyl radical. **a** Free energy surfaces ($\Delta G_{sol-opt}$, in kJ mol⁻¹) in DMF solution for the reaction of isopropyl radical (*i*-Pr•) with NFSI and **b 4f** calculated at the (RO)B2PLYP/G3MP2Large level of theory. Distances (in Å), NPA charges and NPA spin distributions have been calculated at the SMD(DMF)/(U)B3LYP/6-31G(d) level of theory. Free energies in solution $\Delta G_{sol-opt}$ have been obtained by adding $\Delta G_{solv}[(U)B3LYP/6-31G(d)/SMD(DMF)]$ to $\Delta G_{298}[(RO)B2PLYP/G3MP2Large//SMD(DMF)/(U)B3LYP/6-31G(d)]$

(1.56–1.60 Å) distances typical for very early transition states. The transition state charge distribution is very similar for all three fluorination reagents and indicates a charge transfer component of ca. 0.15–0.19e from the radical to the reagent. This charge transfer component is quite important for such an early transition state, where only 20–24% of the spin density has left the substrate isopropyl radical. Interestingly, the most significant difference between the NFSI and NFAS transition states concerns the length of the N–F bond (1.56 Å for NFSI against 1.58–1.60 Å for NFASs). In other words, the more electrophilic NFSI is able to accommodate the extra electron density caused by the charge transfer with less cleavage of the N–F bond relative to the NFASs leading to an extra stabilization of the transition state in full accordance with the polar effects aforementioned.

Both in the hydrofluorination and the decarboxylation processes, NFSI provided the desired fluorinated products in significantly lower yields than NFASs despite the observed higher rate constant for the fluorine atom transfer. For both reactions, the analysis of the crude reaction mixture showed the formation of larger quantities of alkenes for reactions involving NFSI relative to NFASs. The alkenes may result from at least three competitive processes: a single electron transfer (SET) between the fluorinating agent and the secondary alkyl radical leading to a cation followed by loss of a proton; a post fluorination acid catalyzed HF elimination; a radical cross-disproportionation process involving the alkyl radical and the imidyl radical (NFSI) or the amidyl radicals (NFASs). All these three processes are expected to be more prominent when reactions are run with NFSI relative to NFASs. Indeed, the electrophilic nature of NFSI should favor the SET process (pathway a). The HF elimination (pathway b) was experimentally found to be trigger by HF itself. The presence of HF may result from electrophilic reactions between the fluorinating agents and DMF or tert-butanol (hydrofluorination reaction) or traces of water (decarboxylation reaction)^{51,52}. Finally, the radical cross disproportionation process (pathway c) is expected to be favored by the more reactive NFSI-derived imidyl radical over the amidyl radicals derived from NFASs. The difference of reactivity of these radicals is well-illustrated by the calculated N-H BDE for the corresponding amides (H-NFSI: BDE 454.2 kJ mol⁻¹; H-**4a** BDE 393.0 kJ mol⁻¹; H-**4f** BDE 390.6 kJ mol⁻¹ (see Supplementary Fig. 222 and Supplementary Table 8).

Discussion

We have developed NFASs, a class of fluorinating reagents suitable for radical fluorination under mild conditions. The bond dissociation energies of the NFASs are 30-45 kJ mol⁻¹ lower than the one of NFSI and Selectfluor". This favors smooth radical processes over side reactions caused by the electrophilic and oxidant properties of the previous generations of radical fluorinating agents. NFASs were successfully used in a metal-free hydrofluorination method involving hydroboration with catecholborane followed by a radical deborylative fluorination. By using monoisopinocampheylborane (IpcBH₂) in the hydroboration step, the asymmetric hydrofluorination of trisubstituted alkenes can easily be performed. Remarkably, NFASs also proved to be superior to NFSI in decarboxylative fluorination of tertbutyl peresters demonstrating that they are attractive reagents for a broad range of radical mediated fluorination processes. They have the potential to deeply transform the field of radical fluorination by enabling powerful transformations under milder conditions than the former generations of fluorinating agents.

Methods

N-Fluoro-N-(4-(trifluoromethyl)phenyl)benzenesulfonamide (4a). To a solution of *N*-(4-(trifluoromethyl)phenyl)benzenesulfonamide (12.05 g, 40.0 mmol) in DCM (400 mL) was added Cs_2CO_3 (16.90 g, 52.0 mmol) and stirred at room temperature for 60 min. Then, NFSI (16.40 g, 52.0 mmol) was added and the mixture was allowed to stir at room temperature for 5 h. The mixture was diluted with pentane (400 mL), filtered, and concentrated. The product was purified by rapid column chromatography (heptane/TBME 85:15). Concentration of the collected chromatography fractions to a volume of 100–150 mL promoted the crystallization. The solution was stored for one night at 4 °C to yield **4a** (10.15 g, 80%) as a slightly yellow solid. R_f 0.40 (heptane/TBME 9:1); m.p. 74–75 °C.

N-Fluoro-2,4,6-trimethyl-N-(4-(trifluoromethyl)phenyl)-benzenesulfonamide

(4f). According to the procedure for 4a, starting from *N*-(4-(trifluoromethyl) phenyl)-2,4,6-trimethylbenzenesulfonamide (17.17 g, 50.0 mmol). Crystallization at 4 °C yielded 4f (16.16 g, 89%) as a slightly yellow solid. R_f 0.55 (heptane/TBME 9:1); m.p. 116–117 °C.

General procedure for the hydrofluorination of alkenes. To a solution of the alkene (1.0 mmol), *N*,*N*-dimethylacetamide (14 μ L, 0.15 mmol) in dry DCM (1 mL) was added dropwise catecholborane (0.23 mL, 2.2 mmol) at 0 °C. The reaction was allowed to stir at 30 °C for 16 h. The mixture was cooled to 0 °C and *t*-BuOH (0.124 mL, 1.3 mmol) was added. The reaction mixture was stirred at room temperature for 15 min, concentrated under vacuum, and the residue was dissolved in dry DMF (2 mL). DTBPO (117 mg, 0.5 mmol) and **4a** or **4f** (3.0 mmol) were added. The mixture was heated to 60 °C (preheated oil bath was used) and stirred at this temperature for 30–45 min. The crude product was purified by column chromatography.

Data availability

Data supporting the findings of this work are available within the paper and its Supplementary Information files and from the corresponding authors on request. Source data for Supplementary Tables 9, 16 and 17 are provided as supplementary data. CCDCs 1828679 and 1828684 contain the supplementary crystallographic data for compound **4a** and **4f**, respectively. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via http://www.ccdc. cam.ac.uk/data_request/cif

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Author contributions

P.R. proposed the research direction and guided the project. D.M. designed and run the experimental work with the assistance of F.W. Calculations were designed by H.Z. and H. J., and performed by H.J. The manuscript was jointly written by D.M., P.R., and H.Z.

Additional information

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