Click Chemistry

Silicon-Free SuFEx Reactions of Sulfonimidoyl Fluorides: Scope, Enantioselectivity, and Mechanism

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Abstract: SuFEx reactions, in which an S–F moiety reacts with a silyl-protected phenol, have been developed as powerful click reactions. In the current paper, we open up the potential of SuFEx reactions as enantioselective reactions, analyze the role of Si and outline the mechanism of this reaction. As a result, fast, high-yielding, “Si-free” and enantiospecific SuFEx reactions of sulfonimidoyl fluorides have been developed, and their mechanism shown, by both experimental and theoretical methods, to yield chiral products.

Introduction

Click chemistry[1] has been widely used to construct functional molecular skeletons concisely and efficiently.[2] A variety of new click reactions has been developed, such as Cu-catalyzed azide–alkyne cycloadditions (CuAAC)[3] as well as metal-free click reactions[4] such as thiol–ene,[5a] oxime ligation,[5b] (thio-)Michael,[5c] Diels–Alder,[5d] strain-promoted cycloadditions between azide–alkyne (SPAAC)[5e] cyclo-octyne-o-quinone (SPOCO),[5f,g] and cyclopropene-o-quinone[5h] reactions. More recently, the pioneering work of Gembus[6] and Sharpless[7] revived the sulfur(VI) fluoride exchange reactions. More recently, the pioneering work of Gembus[6] and Sharpless[7] revived the sulfur(VI) fluoride exchange (SuFEx) reaction from “old-school chemistry” to the latest generation of click reactions. In this reaction, the S–F bond is typically stable enough to be unreactive in a wide variety of reaction conditions, but in the presence of silyl-protected phenols, fluorosulfates readily react to form sulfates requiring only small amounts of catalyst (base or HF2−). In this process, Si-protection is useful, as Si and F form the strongest single bond in nature, allowing the rapid formation of SO2–O bonds from stable silyl ether precursors.[8] This synergy of the click reaction with the formation of the thermodynamically favorable Si–F bond (BDE = 135 kcal mol−1) results in highly reliable SuFEx reactions under metal-free conditions.

As a result, research into the SuFEx reaction has surged in the last 5 years.[9] To expand the range of useful SuFEx connectors, several SuFEx hubs and fluorosulfuryl transfer reagents have been developed, such as ethanesulfonyl fluoride,[10] SO2F2,[7] SOF4,[11] fluorosulfuryl imidazolium salt,[12] and others.[13] Hereby, a wide variety of SuFEx-able substrates with excellent functional group tolerance can be obtained, affording almost quantitative yields and/or polymers with high molecular weights and narrow polydispersity. For example, SuFEx has been used in modifications of peptides and proteins,[14] syntheses of functional polymers,[15] macrocycles,[16] MOFs[17] and functionalized surfaces,[18] preparations of ionic liquids[19] and late-stage drug functionalization.[20] In addition, SuFEx products have been used as important substrates and key intermediates in various reactions.[21]

Although the SuFEx reaction has become a very practical click reaction, three key features are as of yet unclear: its mechanism, an evaluation of the necessity and influence of Si-based protection of the phenols, and the potential of the SuFEx reaction as source of chiral molecules. To address all three facets in one study, we here report on the investigation of the SuFEx reaction of sulfonimidoyl fluorides. Sulfonimidoyl fluorides[22] bear a stable chiral sulfur-centered moiety, and form an ideal platform for mechanistic and enantioselectivity studies, whereby S1 versus S2 and addition/elimination pathways can be identified via the chirality of the substrates and products. Although some examples of reactions of unprotected phenols with SO2F2,[7,20,25] and R–SO2F[24] have been reported, formation of Si–F covalent bonds (as in Figure 1c) has generally been considered to be a, or even: 

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(wB97XD/6-311+G(d,p) density functional theory) studies (Figure 1d).

While many sulfur-fluoride reagents with different valency have been studied (Figure 1a), sulfonimidoyl fluorides have rarely been studied at all, let alone as SuFEx reagents, although they have already been labelled as useful connectors from the start of recent SuFEx literature.[25] Despite this lack of exploration, the SuFEx reactivity of sulfonimidoyl fluorides has been shown to be comparable to sulfonyl fluorides,[27] with further regulation possible by modification of substituents on the nitrogen atom, making them ideal tunable SuFEx substrates. Further motive is provided by the usefulness of the product, as the resulting sulfoximine group, and structural variations thereof, can express significant medicinal activity.[28] For example, buthionine sulfoximine is a potential adjunct with chemotherapy in the treatment of metastatic melanoma,[28] while atuveciclib is the first potent and highly selective PTEFb/CDK9 inhibitor to enter clinical trials for the treatment of cancer (Figure 1b).[29]

As first part of our studies, the comparison between the traditional and Si-free SuFEx reaction was investigated. By mixing Si-protected (short for: tert-butyl-dimethylsilyl (TBDMS)-protected) phenol with unprotected p-cresol and performing the SuFEx reaction with N-benzoyl-4-methylbenzenesulfonimidoyl fluoride 1 (ratios: 10:10:1, to obtain pseudo first-order conditions), the relative reactivity of the respective phenol derivative (free or Si-protected) can be observed from the composition of the final product mixture. This yielded a ratio for Ph-O-Si/CH_3-Ph-OH of 5.95. The analogous experiment with Si-protected cresol and unprotected phenol yielded a ratio for CH_3-Ph-O-Si/Ph-OH of 9.91 (ratios obtained from H NMR of products upon completion of the reaction, typically < 2 min). This shows that a roughly 10:1 ratio was found for the reaction rate of unprotected phenols and Si-protected phenols (see the Supporting Information (SI) for details). During this reaction time, exchange of the TBDMS-group from the cresol to the phenol or vice versa was negligible. While the formation of the Si-F covalent bond can still be of relevance, it cannot be characterized as the driving force behind the SuFEx reaction.

Since the unprotected phenol derivative was clearly more reactive, we set out to outline the scope of Si-free SuFEx reactions of 1, prepared according to literature,[26] via a systematic variation of the nature of the phenol. The results are displayed in Table 1.

In the presence of 1 equivalent 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU), fluoride 1 reacted smoothly with a variety of phenols 2 and 3-pyrindinol to form the product sulfoximine in good to excellent yields under ambient conditions. Unlike in the normal SuFEx reaction, DBU acted both as a catalyst and an acid scavenger due to the absence of a silane group to stabilize the leaving fluoride anions in the Si-free SuFEx. Most of the reactions actually reached completion within 10 min, providing one product, the sulfoximine, in almost quantitative NMR yield without optimization of the reaction conditions. The reaction exhibited an excellent functional group tolerance, as phenol derivatives containing alkyl (2b,c), ether (2d,g), pyridinyl (2e), amino (2f), amide (2h), carboxyl (2i), halogen (2m-q,v), ester (2s), hydroxymethyl (2t), boronic acid (2u) and cyan (2y) functional groups were successfully used as substrates. Similar to the normal SuFEx reaction, alkyl alcohols, anilines and amides, which are weaker nucleophiles compared to phenolates generated from deprotonations by DBU, did not react in Si-free SuFEx either. Moreover, screening revealed the Si-free reaction was not very sensitive to steric hindrance although reactions took longer, as target products were obtained successfully for sterically hindered phenols, bearing bulky ortho substituents, such as 2-bromophenol (2p-o), 2-(tert-butyl)phenol (2r-o), mesitol (2t). Unsurprisingly, meta-substituted phenol derivatives (2b-m, 2m-m, 2p-m) were found to be excellent substrates. Di-substituted phenol derivatives resorcinol (2k) and biphenol A (BPA) (2l) also partook in the reaction, albeit with 2 equivalents of the sulfonimidoyl fluoride 1. Unfortunately, the resulting diastereomers (3k,l) could not be separated by flash column chromatography (see SI). Substrates containing strong electron-withdrawing substituents in the ortho- or para-position, such as formyl (2w), cyano (2y), trifluoromethyl (2z) or nitro (2ac, 2ad) bearing two large groups (such as 2,6-di-tert-butyl, 2aa) yielded hydrolysis by-products at a loss of the target SuFEx products (see SI) at varying degree, and only provide the SuFEx product very slowly under rigorously dry conditions.

To further demonstrate the scope of this reaction, several natural phenolic derivatives were evaluated (Table 2). Eugenol (4a), vanillic acid (4b), raspberry ketone (4c), tyrosol (4d) sesamol (4e), ferulic acid (4f), and vic-tyrhalen (4g), salicyl aldehyde (4h), and vanillin (4i) all reacted smoothly with fluoride 1 to form the corresponding sulfoximine in good to excellent yields under ambient conditions.

After having established that the Si-free SuFEx reaction proceeds smoothly and efficiently for a wide variety of phenol substrates, we aimed for further insight, by studying other catalysts and the reaction kinetics. Regarding the catalysts, HF2 anion has been introduced as a superior rate-enhancing agent.[29] Therefore we investigated the effects of this anion on...
a range of reactions. When 1 was reacted with Si-protected 2b-p, then this quantitatively yielded product 3b-p at rt in 1 h with one equivalent of catalyst. When using catalytic amounts of HF2 anion (0.2 equiv), the same reaction reached completion, albeit at a much slower pace (30 h). When the unprotected phenol was used with this catalyst (even with 1 equiv), then only trace amounts of product were observed. The likely role the HF2 anion plays in these reactions is clarified if it is added to Si-protected 2b-p in the absence of any 1. Within 10 min, full deprotection of the TBDMS group is then achieved. Therefore, when used in a standard SuFEx reaction, HF2 anion might act as follows: it effects the rapid deprotection of Si-protected phenols. In aprotic solvents, this then yields a highly reactive phenolate, which—upon reaction—frees up a fluoride anion, allowing the reformation of HF2 anion, or fluoride by itself might affect the deprotection of a subsequent Si-protected phenol. The high rate of both the deprotection and of the SuFEx reaction of the phenolate would then allow the use of only small amounts of catalysts, as has been observed experimentally. In line with this, for unprotected phenols HF2 is a poor catalyst: since fluoride is only a poor base, it will—like DBU—not induce the formation of the reactive phenolates; as a result, nothing happens.

As a start of our kinetics study, we compared the reaction rate of the Si-free SuFEx reaction of the sterically hindered 2-(tert-butyl)phenol (2opo) with a traditional SuFEx reaction with the Si-protected 2-(tert-butyl)phenol analogue 5. At room temperature, both these reactions were finished in under two minutes; therefore the kinetics were studied in CD3CN using low-temperature NMR. Under pseudo first-order conditions, the second-order rate constant found for 2opo at 30°C was (3.0 ± 0.1) × 10^-3 M^-1 s^-1. For 5 the kinetics were clearly slower, but interestingly showed an activation phase, with a clear S-shape appearance of the product over time (see SI). To test whether this could be due to the appearance of unprotected phenol 2opo upon a rate-determining desilylation of 5, we studied this desilylation under these reaction conditions (−30°C) with equimolar concentration of 5 and (CH3)4NF (both 0.12 M). This yielded a full desilylation within 2 min, suggesting that for silylated phenols the reaction is initiated by freeing up some F anion, which can deprotect a Si-protected phenol, which can subsequently react and free up another F, etc. Here DBU can play a double role, as it has been reported to catalyze the desilylation,[30] while as a base, of course, it also increases the nucleophilicity of the phenol.

Table 1: Scope of Si-free SuFEx reactions of sulfonimidoyl fluoride 1 with phenolic derivatives.^[a]

| Compound | Reaction Conditions | Yield (%) |
|----------|---------------------|-----------|
| BzN | 30°C, 30 min | >95% |
| BzN | 20°C, 30 min | >95% |
| BzN | 30°C, 30 min | >95% |
| BzN | 30°C, 30 min | >95% |
| BzN | 30°C, 30 min | >95% |
| BzN | 30°C, 30 min | >95% |
| BzN | 30°C, 30 min | >95% |

[a] Conversion was determined by 1H NMR measurements. Isolated yield in brackets. Reaction conditions for 1H NMR conversion: 1 (0.05 mmol), phenolic derivative 2 (1.05 equiv) and DBU (1.0 equiv) in CD3CN (0.55 mL), rt. For isolated yield: Fluoride 1 (0.5 mmol), phenolic derivatives 2 (0.5 mmol), DBU (1.5 mmol) in anhydrous CH3CN (1 mL), rt. [b] Hydrolysis by-product was formed (see SI). [c] 2 equiv of DBU were used. [d] 2 equiv of 1 were used.
Table 2: Scope of Si-free SuFEx reactions of sulfonimidoyl fluoride 1 with naturally occurring phenols.\(^{[a]}\)

| Reaction Conditions | Products |
|---------------------|----------|
| 1 mL, 3.6 mmol in CH₃CN, phenolic derivative 2 (1 equiv), DBU (1 equiv), rt. | BzN₂SO₂R, BzN₂SO₂OH, BzN₂SO₂COOH |

[a] Conversion was determined by \(^1\)H NMR measurements. Reaction conditions for \(^1\)H NMR yield: 1 (0.03 mmol) phenolic derivative 2 (1.05 equiv) and DBU (1.0 equiv) in CH₃CN (0.55 mL), rt. [b] Hydrolysis by-product was formed (see SI). [c] 2 equiv of DBU were used.

Further mechanism investigations were made possible by utilizing the chirality of sulfur atom in the sulfonimidoyl fluoride 1 and its SuFEx products. Previous mechanism-oriented reports have typically focused on substitution reactions on disulfides, yielding addition–elimination and Sn₂ as the most widely accepted mechanisms, with the detailed mechanism being sensitive to the strain, counter cations, leaving groups, substituents on sulfur atom, nucleophiles, solvents, and so on (see for an overview). Analogously, we considered as the three most likely mechanisms those summarized in Figure 2.

If one enantiomer of 1 is used as the substrate in SuFEx reactions, the Sn₁ reaction pathway will provide the racemized products. Conversely, stereospecific products will be obtained through the Sn₂ or addition-elimination pathway. Firstly, enantiomers of sulfonimidoyl fluoride 1 were shown to be very stable in solution without DBU, but readily racemize when DBU was present. The rate of this racemization reaction, with excess DBU under pseudo first-order conditions, is \((1.9 \pm 0.1) \times 10^{-3} \text{ M}^{-1} \text{s}^{-1}\) at \(-30^\circ\text{C}\), showing its facile occurrence and competitive character (see SI). Since this would reduce the usefulness of the reaction, further studies were performed.

To this aim, a good and a poor nucleophile—p-cresol and 4-methyl-2-nitrophenol, respectively—were adopted in SuFEx reactions. With the good nucleophile a good stereoselectivity (ca. 70% ee) was achieved, whereas with the poor nucleophile only <2% ee was obtained (Table 3). This provided important mechanistic information, as it shows that SuFEx proceeds through Sn₂ or addition-elimination channels rather than via an Sn₁ mechanism when a good nucleophile was used. The racemization or Sn₁ mechanism was a possible reason of low stereoselectivity when a poor nucleophile (4-methyl-2-nitrophenol) was used. If the Sn₂ or addition-elimination is a competitive reaction with the racemization, we reasoned that the racemization could be inhibited when a large excess of a good nucleophile (10 equiv p-cresol) was used. In fact, the experimental results showed that under such conditions the %ee was significantly improved (95% ee). However, the key finding was avoiding DBU altogether, and using sodium phenolate and sodium 4-methyl-2-nitrophenolate as nucleophiles. In this case stereospecific products were obtained, and no racemization of the reactants was observed, even for 7, which takes overnight to get to 80% conversion. This further demonstrated that the silicon-free SuFEx reaction mechanism proceeds via an Sn₂ or addition-elimination mechanism.

To further distinguish concerted Sn₂ and addition-elimination reaction paths, quantum chemical calculations were performed at the wB97X-d/6-311+G(d,p) level of theory, using the SMD solvent model to represent CH₃CN as implemented in Gaussian. We computed the potential energy surfaces (PES) for the SuFEx reaction between fluoride 1 and

Table 3: Enantioselective silicon-free SuFEx reactions.\(^{[a]}\)

| Enantiomer \(^{[b]}\) | Phenol | ee\(^{[c]}\) |
|-------------------|--------|----------|
| enantiomer-1      | 2b-p   | 73% (95\(^{[d]}\)) |
|                   | 2ac    | <2%      |
|                   | 6      | >99%     |
|                   | 7      | >99%     |
| enantiomer-2      | 2b-p   | 71% (95\(^{[d]}\)) |
|                   | 2ac    | <2%      |
|                   | 6      | >99%     |
|                   | 7      | >99%     |

[a] Reaction conditions: enantiomer-1/enantiomer-2 (1 mL, 3.6 mmol in CH₃CN), phenolic derivative (1 equiv), DBU (1 equiv), rt. [b] enantiomer-1 and enantiomer-2 refer to two enantiomers with retention times of 16.8 and 18.8 min with chiral HPLC, respectively. [c] ee determined using chiral HPLC and calculated by ee = \(\frac{\text{minority} - \text{majority}}{\text{majority} + \text{minority}}\) \times 100\%. [d] 10 equiv 2b-p used.

Figure 2. Possible mechanisms of SuFEx reactions.
phenolate 6 (Scheme 1; to slightly simplify the calculation, we replaced the 4-methylbenzenesulfonylimidoyl with a benzene-
sulfonylimidoyl group). Because of the strong S-F covalent
bond (\(\approx 90 \text{kcal mol}^{-1}\)), fluoride 1 is very stable, and the
leaving of fluoride is generally considered to be one of the
rate-limiting steps. However, for the reaction under study, this
does not seem to be the case. Compound 1 undergoes the
nucleophilic attack by 6 via TS1, with a rate-limiting enthalpic
barrier of only 2.74 kcal mol\(^{-1}\). This confirms the experimen-
tally observed high reaction rate. However, in TS1 the S-F
bond was still largely unchanged (length increase from 1.64 to
1.72 Å; Wiberg bond order decrease from 0.520 to 0.496),
likely due to a relatively early TS (\(r(\text{O} \cdots \text{S}) = 2.33 \text{ Å}; \text{B.O.} = 0.151\)).

The next stationary point, however, was not the set of
products, but a five-coordinated sulfur intermediate, INT1,
which was found to be lower in energy than the reactant pair.
This intermediate is characterized by an \(r(\text{O} \cdots \text{S}) = 1.78 \text{ Å},\) and a \(r(\text{S} \cdots \text{F}) = 1.93 \text{ Å},\) indicating bonding of the O to S
(B.O. = 0.544), but also no loss of F yet (B.O. = 0.315).
Interestingly, the S–O bond does not elongate significantly
during the reaction (1.44–1.45 Å, with the bond length of
a typical S–O bond = 1.64 Å\[^{[34]}\]), nor does the charge on this O
atom vary a lot during the reaction.

This intermediate readily loses fluoride via TS2 (\(\Delta H^\ddagger\) of
this step is only 0.53 kcal mol\(^{-1}\)), giving rise to the products.
Activation of S–F loss by the base has been one of the most
widely accepted mechanism models to date.\[^{[6,11,35]}\] This might
be the case for much poorer nucleophiles, including the Si-
protected phenols, but such activation is apparently not
needed for phenolate SuFEx reactions. In summary, these
DFT calculations thus suggest that SuFEx with a phenolate
nucleophile attacking a sulfonimidoyl fluoride takes places
with bimolecular kinetics via an addition–elimination mech-
anism, although a concerted S_N2 reaction under experimental
conditions cannot be excluded given the very low barrier
observed for loss of F\(^{-}\).

This SuFEx reaction has a reaction enthalpy of about
\(-5 \text{kcal mol}^{-1}\), which is nice to drive a reaction quietly to
completion, and better than many other click reactions that
are simply too exothermic.\[^{[6]}\]

We also aimed to characterize the nucleophilic attack by
DBU. However, no TS could be found, and the energy simply
increases upon decreasing the \(r(\text{N} \cdots \text{S})\) distance (to \(>19 \text{kcal}
\text{mol}^{-1}\) for \(r(\text{N} \cdots \text{S}) = 1.7 \text{ Å}\)). Apparently, the steric bulk of DBU
makes this a really non-nucleophilic base in this reaction.

**Conclusion**

In conclusion, we have demonstrated that Si-free SuFEx
reactions with phenolate anions can be readily performed in
a stereoselective manner with excellent functional group
tolerance and good to quantitative yields in minutes at room
temperature. The Si-free SuFEx is faster than the traditional
one, and detailed kinetics of the latter might point to a more
complex reaction scheme there. A combination of these
experiments with theoretical calculations show that the Si-
free SuFEx reaction is a fast, bimolecular, enantiospecific and
slightly exothermic click reaction, making it amenable for
a wide range of substrates. These studies contribute to
a clearer understanding and continuing development of
SuFEx reactions, as well as providing a practical approach
for the construction of optically pure sulfoximines.

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**Conflict of interest**

The authors declare no conflict of interest.

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T. J. Kim, I. Lee, J. Phys. Chem. A 2009, 113, 7073 – 7079; d) R. F. Höckendorf, Q. Hao, Z. Sun, B. S. Fox-Beyr, Y. Cao, O. P. Balaj, V. E. Bondybay, C.-K. Siu, M. K. Beyer, J. Phys. Chem. A 2012, 116, 3824 – 3835; e) M. Paranjothy, M. R. Siebert, W. L. Hase, S. M. Bachrach, J. Phys. Chem. A 2012, 116, 11492 – 11499; f) R. P. P. Neves, P. A. Fernandes, A. J. C. Varandas, M. J. Ramos, J. Chem. Theory Comput. 2014, 10, 4842 – 4856; g) M. Bortoli, L. P. Wolters, L. Orian, F. M. Bickelhaupt, J. Chem. Theory Comput. 2016, 12, 2752 – 2761.

[32] T. A. Hamlin, M. Swart, F. M. Bickelhaupt, ChemPhysChem 2018, 19, 1315 – 1330.

[33] R. B. Gaussian 16, Revision B.01, M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmanì, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marecich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman, and D. J. Fox, Gaussian, Inc., Wallingford CT, 2016.

[34] T. Chivers, R. S. Laitinen, Chem. Soc. Rev. 2017, 46, 5182 – 5192.
[35] a) H. Vorbrüggen, Synthesis 2008, 1165 – 1174; b) J. Yatvin, K. Brooks, J. Locklin, Chem. Eur. J. 2016, 22, 16348 – 16354.

[36] G. O. Jones, K. N. Houk, J. Org. Chem. 2008, 73, 1333 – 1342.

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