Introducing SuFNucs: Sulfamoyl-Fluoride-Functionalized Nucleosides That Undergo Sulfur Fluoride Exchange Reaction

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ABSTRACT: The reaction between ribonucleosides and ex situ generated sulfonyl fluoride has been developed. The reaction takes place at the −NH2 groups of nucleobases, and the resulting nucleosides are equipped with a sulfamoyl fluoride moiety, dubbed SuFNucs. These species undergo a selective sulfur fluoride exchange (SuFEx) reaction with various amines, leading to sulfamide-functionalized derivatives of adenosine, guanosine, and cytidine (SulfamNucs). The scope and examples of further SuFNucs functionalization leading to nucleotides, oligonucleotides, and peptide−nucleoside conjugates are presented.

Unlike other sulfur(VI) electrophiles, the −SO2F group is an extraordinarily stable unit, even under harsh conditions. Nevertheless, it is possible to controllably activate the S(VI)−F bond, transforming fluoride into a good leaving group and enabling reactions with various nucleophiles. Different conditions have been explored to achieve this purpose (appropriate acid/base environment, presence of R3Si+ as a mediator, properly selected catalyst, and constrained environment) and used in versatile applications. S(VI)−F-containing compounds and their chemical behavior have been known and utilized for nearly a century, but their huge potential was recognized and systematized in 2014 by Sharpless and co-workers under the term sulfur(VI) fluoride exchange reaction (SuFEx, Figure 1A).1 SuFEx reactions were immediately included in the click chemistry paradigm,2 as the S(VI)−F unit in organic compounds can be used as a molecular connector in a predictable manner under mild conditions. Since this breakthrough, the expansion of SuFEx-type reactions covered general organic chemistry,3,4 material/polymer science,5 chemical biology,6 bioconjugate chemistry,7 and medicinal chemistry including drug discovery.8,9 The SuFEx approach requires installation of a −SO2F group in the molecule of interest. The emerging demand for diverse molecules containing the S(VI)−F motif can be satisfied thanks to so-called SuFEx hubs—reagents that can introduce the SO2F group into the target molecule (Figure 1B). The simplest SuFEx hub is sulfonyl fluoride (SO2F2, bp = −55 °C),5 which, in spite of its high reactivity toward phenols and secondary amines, has limited application toward the functionalization of primary amines. Moreover, SO2F2 cannot be used in laboratories lacking proper equipment to handle harmful gases, and its availability is limited by regulations. These drawbacks have been solved by the development of a solid SO2F+ donor fluorosulfuryl imidazolium triflate salt (SuFExIT, Figure 1B)10 and [4-(acetylamino)phenyl]imidodisulfuryl difluoride (AISF, Figure 1B)11 which are stable, are easy to handle, ensure precise control over the reagent ratio, and show expanded reactivity (compared to the parent gas). Alternatively, SO2F2 can be generated ex situ from the solid precursors10,12,13 [for example sulfonyl diimidazole (SDI) and KF, Figure 1C] and delivered into the reaction mixture using specially designed, commercially available laboratory glassware (COware, Figure 1D). This set of reagents and fine-tuned conditions for their reaction with various organic structures allows quick access to the vast spectrum of SuFExable compounds for further investigations including drugs9 and other biologically active molecules.14

Although SuFEx reactions already established their usefulness in bioconjugate chemistry and medicinal chemistry, there is only a limited number of nucleosides and nucleotide derivatives functionalized with SuFExable units or involved in SuFEx-type chemistry.15 This gap in the area of SuFEx chemistry inspired us to investigate the possibility of introducing the −SO2F motif into nucleosides and investigate the potential of such derivatives as novel, ready-to-functionalize building blocks.

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We aimed to add SuFEx-able groups to natural ribonucleosides bearing an \(-\text{NH}_2\) group at the nucleobase unit, namely, adenosine (A), guanosine (G), and cytidine (C). We expected that their successful reaction with an electrophilic \(\text{SO}_2\text{F}^+\) donor will result in sulfamoyl-fluoride-functionalized nucleosides, which can be excellent starting materials for the synthesis of yet unexplored sulfamide-bearing nucleosides. To the best of our knowledge, the only example of nucleobases possessing a sulfamide group was synthetized using pentachlorophenyl sulfamate and was limited to simple \(-\text{SO}_2\text{F}^-\text{NH}_2\) functionality.\(^{16}\) In preliminary experiments, we have tested SuFExIT\(^{10}\) as a \(\text{SO}_2\text{F}^+\) group donor. This reagent has been shown as a superior SuFEx hub for primary amine derivatization, and before its introduction synthetic access to the sulfamoyl fluorides was limited.\(^{17}\)

Unfortunately, by treating A, G, C, Ac₃A, Ac₃G, and Ac₃C with this reagent in various solvents, we obtained complex synthesized products that were difficult to separate into individual components.

When no base was present in the reaction mixture or weak bases were used, the starting material was fully recovered, but in the presence of DBU (4 equiv) the overnight reaction resulted in full conversion of the starting material and highly selective formation of the expected product. The reaction proceeded equally well in any solvent tested (see Table S1), and the product 1 was isolated in nearly quantitative yields and excellent purity only by extraction techniques (to remove DBU and its salts). In the case of Ac₃G and Ac₃C, the same reaction conditions and purification method worked alike, giving 2 and 3 in 95% and 75% yield, respectively (Scheme 1, see SI for details). The structures of the products were confirmed by a series of NMR analyses, which indicated the presence of a fluorine nucleus with chemical shifts in the range characteristic for sulfamoyl fluorides: \(\delta_F = 53.93\) ppm for 1 (recorded in CDCl₃), \(\delta_F = 50.89\) ppm for 2 (recorded in DMSO-\(d_6\)), and \(\delta_F = 52.49\) ppm for 3 (recorded in CDCl₃). The \(^{13}\)C NMR analysis of 2 revealed a spin–spin coupling between C2 carbon and fluoride \((J_{C-F} = 3\) Hz), confirming that the reaction took place at the N2 position.

The reaction scale up did not generate any losses in selectivity and maintained high yields and purity, allowing for the preparation of larger quantities (up to 3 mmol/>1 g in one batch when 100 mL of reactor was applied, see Figure S1) of SuFNucs 1–3 without compromising the reaction performance. After establishing a practical method for the preparation of SuFNucs 1–3, we investigated their reactivity toward amines. Heating 1 with selected amines (benzyl amine, morpholine, and toluidine) in the presence of TEA in various solvents\(^ {18}\) did not
lead to the formation of the desired product, while slight degradation of the starting material was observed. Hence, we focused on the approach toward reacting "SuFExable" units with amines developed by am Ende and Ball.20,21 In this concept, the activation of the −SO2F group is achieved by the addition of a stoichiometric amount of Lewis acid in combination with a proper amine mediator. Application of Ca(NTf2)2 and DABCO has allowed us to transform sulfamoyl fluorides, fluorosulfates, and sulfonyl fluorides into respective sulfamides, sulfamates, and sulfonamides by applying a single set of optimized conditions.21 To our delight, the model reaction between 1 and benzyl amine in the presence of the Ca(NTf2)2/DABCO system resulted in the formation of the expected sulfamide along with some unidentified polar byproducts, which could be easily removed during workup (see Table S2 and Figure S2 for details).

The yield and selectivity were affected by the nature of the solvent used (Table S2 and Figure S2), with DCM and MeCN performing the best. This finding opened an entry to a wide range of different sulfamide-modified nucleosides (SulfamNucs, Figure 2). The reaction worked well for ammonia (4a, 30%) and simple primary amines like methyl amine (4b, 41%), t-Bu amine (4c, 55%), cyclopropyl amine (4d, 44%), benzyl amine (4f, 50%), and adamantane (4e, 45%). The utility of amines possessing an additional functional group, which can be used for further transformations, gave high yields of valuable compounds 4h (69%) and 4i (70%) functionalized with terminal alkyne and azide, respectively. The yields were not affected upon scaleup to 2 mmol of the starting material. A conjugate with an amino acid derivative, methyl ester of glycine, was also obtained in good yield (4j, 46%). Secondary amines reacted equally well, giving pyrrolidine (4k), morpholine (4l), and proline methyl ester (4m) SulfamNucs derivatives in 70%, 64%, and 43% yield, respectively. It is worth mentioning that some of the amines (volatile ones and amino acids) were used as hydrochlorides, which did not affect the yields if a proper excess of DABCO was applied. The functionalization of guanosine derivative 2 also gave satisfying results, after a slight amendment of the reaction conditions was applied to overcome solubility issues (use of the DCM/THF mixture as a solvent and lowering the concentration). Yields of the respective SulfamNucs were slightly lower compared to adenine derivatives; nevertheless, the simplicity of workup and purification allowed us to obtain a collection of SulfamNucs 5a−g (Figure 2) with simple and more complex substituents (for additional information, see Figure S3). In the case of cytidine, the reactivity under the established conditions was similar to adenine. Compound 3 reacted smoothly with different primary and secondary amines (Scheme 2) in moderate to good yields, keeping the workup/isolation of SulfamNucs 6a−e simple and not particularly laborious.

The deprotection of SulfamNucs was tested with a range of standard reagents (ammonia, MeNH2 in MeOH, and MeNH2 in EtOH). Methylamine in absolute ethanol (33%) was the most effective, leading to unprotected nucleosides in high yields and only minimal workup necessary (Scheme 2; see SI for details). To demonstrate the reactivity of SuFNucs toward more complex molecules, we reacted 1 with a short peptide-containing

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**Figure 2.** Scope of Ca(NTf2)2/DABCO-mediated SuFEx reaction between SuFNucs and amines.
lysine (Scheme 3). Due to the solubility issues, the reaction was performed in DMSO, under high dilution and with an excess of 1 relative to the peptide. After 4 h of incubation, the whole peptide was consumed, and the desired conjugate was easily isolated using RP-HPLC. Deprotection of OH groups was performed as described above and lead selectively to the peptide−nucleotide conjugate 10 connected via sulfamide linkage in 24% overall yield (the conditions and workup/isolation procedure were not optimized).

To investigate further transformation possibilities, adenosine SulfamNuc 7d was transformed into the corresponding 5′-phosphate under standard Yoshikawa phosphorylation conditions (Scheme 4). Treatment of 7d with POCl₃ in (MeO)₃PO under cooling resulted in monophosphate 11 in good yield and purity.

Finally, we introduced sulfamide-functionalized adenosine into short RNA oligonucleotide using standard phosphoramidite-based solid phase synthesis (SPS). The synthesis of the proper phosphoramidite building block required for SPS started from 2′-O-Me adenosine (2′-O-MeA; Scheme 5). This excluded the necessity of selective 2′-OH protection and simplified the synthesis. The synthesis of proper SulfamNuc 14 was accomplished in 3 steps with 72% yield. Introduction of DMT at the 5′ position and phosphoramidite at the 3′ position was achieved using standard nucleoside chemistry and led to 16 in 36% yield (26% overall yield for 5 steps). 16 was used in manual SPS (see SI for details) of trinucleotide 17 composed of uridine, SulfamNuc 2′-O-Me adenosine 14, and guanosine. Upon deprotection and cleavage from the support, the trinucleotide 17 was purified using standard chromatographic techniques in 57% yield.

In summary, by applying ex situ generation of SO₂F₂, we were able to prepare adenosine, guanosine, and cytidine derivatives bearing −NHSO₂F motif as “SuFEx-able” nucleosides (SuF-Nuc) in high yields. Such a reaction between primary amine and SO₂F₂ is very rare and is in contrast to previous observations. The −NHSO₂F functional group was then used to functionalize these nucleosides with diverse amines leading to previously unknown sulfamide-functionalized nucleosides (SulfamNuc) in good yields. All these transformations are operationally simple, are high yielding, and can be easily upscaled to the gram scale, and obtained compounds are stable upon storage. Further prospects of SulfamNuc chemistry were demonstrated by conjugation with peptide and incorporation into nucleotides including oligonucleotides. These findings potentially open several new avenues in nucleoside and nucleotide chemistry, including more robust preparation of highly functionalized structures, bioconjugates, and nucleotide/nucleoside libraries. Further exploration of SuFNuc- and SulfamNuc-type compounds is under investigation in our group and will be presented in due course.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.orglett.2c02034. Supporting tables and figures, detailed synthetic procedures, and compound characterization (PDF)

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. M.C., J.J., and J.K. designed the study. M.C. conducted experiments. K.Z. performed additional experiments. M.C., J.J., and J.K. prepared the first draft of the manuscript. J.J. acquired funding. All authors reviewed, edited, and approved the final version of the manuscript.

Notes

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