Facile access to nitroarenes and nitroheteroarenes using N-nitrosaccharin

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Nitroaromatics and nitroheteroaromatics serve as key building blocks and intermediates in synthesis, and form the core scaffold of a vast number of materials, dyes, explosives, agrochemicals and pharmaceuticals. However, their synthesis relies on harsh methodologies involving excess mineral acids, which present a number of critical drawbacks in terms of functional group compatibility and environmental impact. Modern, alternative strategies still suffer from significant limitations in terms of practicality, and a general protocol amenable to the direct C-H functionalization of a broad range of aromatics has remained elusive. Herein we introduce a bench-stable, inexpensive, easy to synthesize and recyclable nitrating reagent based on saccharin. This reagent acts as a controllable source of the nitronium ion, allowing mild and practical nitration of both arenes and heteroarenes displaying an exceptional functional group tolerance.

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Electrophilic aromatic nitration is arguably one of the most extensively studied transformations in organic synthesis. The reaction has played a crucial role in our understanding of fundamental concepts underlying the reactivity of aromatics and has become an integral part of any undergraduate curriculum. Furthermore, nitroaromatics have become an essential class of compounds serving as key building blocks and versatile intermediates in organic synthesis and in the preparation of various industrial products. These compounds serve as precursors to amines, hydroxylamines, aldehydes, carboxylic acids, isocyanates and various heterocycles, and are starting materials in the nitro-aldol and Michael reactions, as well as in various cycloadditions. In an industrial context, nitrobenzene is an essential intermediate in the synthesis of aniline, while nitro(hetero)aromatic compounds are important precursors in the synthesis of azo dyes, explosives, and materials.

Despite the significance of electrophilic aromatic nitration and the value of nitroaromatic compounds, a mild and practical approach towards electrophilic aromatic nitration remains a challenge for synthetic chemists. To this day, the so-called ‘fixed acid’ approach remains the fundamental process for the production of nitroaromatics on both laboratory and industrial scales. This methodology utilizes a mixture of nitric acid (HNO₃) and sulfuric acid (H₂SO₄) as a solvent and source of the active electrophilic nitronium (NO₂⁺) species, which is not surprising that this harsh methodology suffers from several critical drawbacks (Fig. 1a). Crucial issues include the poor regioselectivity and functional group tolerance of the protocol, as it is clearly unamenable to acid-sensitive functional groups, and side products resulting from oxidation or hydrolysis are often obtained. Furthermore, nitrogen oxides (NOₓ) are generated along with superstoichiometric amounts of acidic waste, which both presents an environmental hazard and complicates isolation of the desired products.

While other classical electrophilic nitrating reagents, including acyl nitrates, nitryl halides, or nitronium salts circumvent the use of excess quantities of mineral acid, these reactions typically require harsh, and in many cases strictly anhydrous conditions, while some of these reagents are further susceptible to thermal decomposition. Various nitrate salts have been investigated in recent years as alternative nitrating agents, however, they generate stoichiometric amounts of metal waste. Modern regiospecific nitration methodologies include ipso-nitration of prefunctionalized arenes and chelation-assisted nitration using transition metal catalysis, and alternatively, aryl azides can be oxidized to the corresponding nitro compounds.

A mild and practical approach towards the nitration of a diverse range of arenes that does not require prefunctionalized

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**Fig. 1 Development of nitrating reagents.**

**a** Current methods to access nitroarenes. **b** Synthesis of a series of N–NO₂ heterocyclic compounds based on pyrrolidinone (1), succinimide (2), phthalimide (3), and saccharin (4) structures. **c** Our synthesis of reagent 4 and its application in the nitration of arenes.

Reagent
- Bench-stable reagent
- Can be handled under air
- Cheap, one-step synthesis
- Recyclable starting material

Nitration of arenes
- Mild, neutral reaction conditions
- Excellent functional group tolerance
- High chemical efficiency
- No formation of unwanted side-products
- Only recoverable saccharin as by-product
- Scalable methodology
starting materials remains in high demand at both an academic and industrial level. We speculated that the development of a nitrating reagent based on a tunable organic scaffold could offer a versatile approach towards the realization of such a process.

While the application of various N-nitramine-based reagents in electrophilic nitration of arenes has been investigated in the last century, these protocols suffer from high-moisture sensitivity, and the use of air-sensitive Lewis acid catalysts. Nevertheless, these studies demonstrate the potential of such R₂N–NO₂ reagents to act as viable nitrating agents. Herein we present a mild and acid-free methodology for the electrophilic nitration of a diverse range of arenes and heteroarenes displaying an exceptional functional group tolerance. The reaction is enabled by 4, an easily accessible, inexpensive and recyclable reagent that is a bench-stable and convenient source of the NO₂ group (Fig. 1c).

Results
Reaction development. We commenced by evaluating the reactivity of a series of R₂N–NO₂ heterocyclic compounds based on pyrrolidinone (1)¹⁹, succinimide (2)²⁰, phthalimide (3)²⁰, and saccharin (4) core structures. While the application of 2 in aromatic nitrations under photolysis has been investigated, the reagent demonstrated poor reactivity²¹ and likewise, we did not observe nitration of benzene using 2 during our initial reaction screening (Supplementary Table 2). However, we were pleased to observe that the nitrination of benzene proceeded in 75% yield using reagent 4a in slight excess in acetonitrile at 85 °C (Supplementary Table 3). Although the synthesis of 4a from saccharin has been reported using either unstable NO₂BF₄ or gaseous N₂O₅, with rudimentary characterization, its application as a nitrating reagent has to the best of our knowledge never been explored²²,²³.

We pursued a practical approach towards 4a, and achieved its synthesis on a 50 g scale in 95% yield from saccharin using a combination of fuming nitric acid and acetic anhydride (Fig. 1c). Structures of both reagents 4a and 4b have been confirmed by single crystal X-ray diffraction (Supplementary Tables 5 and 6). Characterization by differential scanning calorimetry and thermal gravimetric analysis revealed 4a to be stable until 173 °C whereby a rapid exothermic decomposition was observed, accompanied by a mass loss of 87% (Supplementary Fig. 2).

Further screening on the nitrination of benzene using 4a revealed the beneficial effects of fluorinated alcoholic solvents on the reaction, and employing 1,1,1,3,3,3-hexafluoroisopropanol (HFIP) as a solvent at 55 °C afforded nitrobenzene in 99% in 3 h. Additionally, a careful screening of various Bronsted or Lewis acids (Supplementary Table 3) revealed that the reaction proceeded smoothly using 10 mol% of magnesium perchlorate [Mg(ClO₄)₂] in acetonitrile at 85 °C. With two methodologies in hand, we proceeded to explore the scope and functional group compatibility in the nitrination of various small and medium-sized molecules (Fig. 2). HFIP proved to be suitably activating in almost all cases, and only for exceptionally challenging deactivated substrates was the [Mg(ClO₄)₂] protocol employed.

Reaction scope. Monosubstituted arenes containing both electron-donating and electron-withdrawing groups (5–20) were successfully nitrated in excellent yields. Halides (6–9), alkyl (10–11), and alkoxo-groups (12–13) were well tolerated, with the expected o- and nitrated products obtained in yields above 90% in all cases (Fig. 2). Remarkably, the nitrination of toluene resulted in a slight excess of the para-regioisomeric product, which is in stark contrast to the generally observed preference of o-nitrobenzene formation under mixed acid conditions or when using nitronium salts²⁴,²⁵. Likewise, the ratio of p- to o-tert-butyl-nitrobenzene is substantially higher than the expected ~ 4:1 ratio using mixed acid (11)²⁴.

Nitrination of cyclopropylbenzene (14) proceeded smoothly, with no ring opening observed. The method proved amenable to sulfides (15), whereby none of the oxidized side products generally observed when using mixed acid were formed, as well as tertiary amines (16). In the case of 15 the use of [Mg(ClO₄)₂] catalysis led to a noticeable increase in yield. Similarly [Mg(ClO₄)₂] was employed in the nitrination of deactivated trifluoromethylbenzene (17) and benzonitrile (18), furnishing the anticipated o- and n-substituted products in good yields. The smooth nitrination of benzonitrile under our conditions is noteworthy, as aryl nitriles are susceptible to hydrolysis under standard mixed acid conditions²⁶. The nitrination of methyl benzoxide (19) and benzophenone (20) also proceeded smoothly without any hydrolysis of the ester or ketone moieties. Biphenyl underwent nitrination selectively on one aromatic ring (21), while nitrination of napthalene (22) and anthracene (23) delivered only the expected isomers. The reaction proceeded well using various other d- and trisubstituted arenes containing both electron-donating and strongly withdrawing groups (24–31). Notably, the nitrination proved to be compatible with aldehydes (25), carboxylic acids (26), phenols (26–27), and amides (32–33), as well as sulfoxides (34) and boronate esters (35). To our surprise, nitrination of methyl 4-tert-butyl benzoate resulted in introduction of the nitro group ortho to the ester substituent (24) as the major regioisomeric product. While the nitrination of boronate esters remains poorly explored in the literature²⁷, nitrination of boronic acids using nitric acid results in formation of the para-regioisomeric product in 70% yield²⁸. In contrast, we observed an almost 1:1 ratio of para to ortho products in the nitrination of phenyl boronic acid pinacol ester 35. This could be rationalized on the grounds of a stabilizing interaction between the reagent’s oxygen atom and boron’s empty p-orbital, directing nitration to the ortho position (vide infra, Fig. 4e).

Our protocol also proved applicable in the nitrination of heteroarenes. While the nitrination of 2,6-dichloro-and 2,6-dimethylpyridine typically involves mixed acid at temperatures above 100 °C²⁹,³⁰, we were pleased to observe the formation of 36 and 37 in acceptable yield using 4a under our Lewis acid catalyzed conditions. This protocol was also suitable in the nitrination of substituted pyrimidines (38–39), while substituted furan (40) and thiophenes (41–42) were nitrated in good yields in HFIP.

In all cases, we were unable to observe polynitrination under established conditions. To further demonstrate the applicability and recyclability of 4a, the nitrination of benzene in HFIP was performed on a 7.5 g scale with no decrease in yield, and both saccharin and HFIP were recovered in 97% and 90%, respectively.

We subsequently investigated the application of our nitrination protocol in late-stage functionalization. Nitroaromatics have in recent decades proven to be an important class of molecules for the preparation of biologically active substances, finding application as pesticides³¹, and as anticancer, antiparasitic and antitubercular agents, as well as antibiotics and tranquilizers³² and as such, we foresaw that our methodology could be of particular value in the context of medicinal and agrochemistry. To this end, a series of nitrated derivatives of known drugs, including ibuprofen (43), lidocaine (44), naproxen (45), clofibrate (46), phenytoin (47), and nimesulide (48) were synthesized in good to excellent yields, the latter furthermore demonstrating the protocol’s tolerance to sulphonamides (Fig. 3). In addition, our protocol enabled the direct preparation of antibiotic Secnidazol (49)³³. A precursor to the Alzheimer’s drug Aricept (50), the commonly used fragrant and intermediate in the synthesis of various pharmaceuticals ventraldehyde (51), and drug
intermediate 52 all smoothly underwent nitration. Furthermore, a nitrated derivative of pesticide procymidone (53) was prepared, and the herbicide fluorodifen (54) could be synthesized directly from the corresponding diphenyl ether in 88% yield and good regioselectivity. Our protocol was also successfully applied in the nitration of naturally occurring estrone (55), arbutin (56), vitamin E (57), nodihydrocapsaicin (58), and L-phenylalanine (59). To show the applicability of our protocol beyond pharmaceuticals and agrochemicals, nitrated derivatives of chiral building blocks binol (60) and [2,2]-paracyclophane (61) were synthesized, thereby providing a convenient strategy for the development of chiral ligands for asymmetric catalysis. The reaction also proved suitable for the modification of organic materials, demonstrated through the nitration of a liquid crystalline material (62).

Mechanistic considerations. Despite arene nitration via electrophilic aromatic substitution being one of the oldest studied reactions, mechanistic aspects of the transformation are still being elucidated today. In light of recent studies considering the effects of implicit solvent molecules on electrophilic nitration of arenes under acidic conditions, we were motivated to gain mechanistic insight into the transfer of the NO2 group and the role of HFIP in facilitating this reaction. We commenced by performing the nitration of benzene in the presence of radical scavengers, and did not observe significant inhibition (Fig. 4a).

**Fig. 2** Electrophilic nitration of arenes and five- and six-membered heteroarenes using 4a. Standard procedures: I (Hetero)arene (0.5 mmol), 4a (0.65 mmol), HFIP (0.5 M), 55 °C, 3 h ([a], 19 h); II (Hetero)arene (0.5 mmol), 4a (0.65 mmol), [Mg(ClO4)2] (10 mol%), CH3CN (0.5 M), 85 °C, 5 h ([a], 19 h). Yields refer to the isolated product of a single regioisomer or a sum of separately isolated regioisomers. [*] The minor regioisomeric position is labeled. Scale up synthesis of 5 under conditions I (reaction run on 84.0 mmol of benzene; yield of 5 is 97%). a Nitration of small and medium-sized building blocks. b Nitration of heteroaromatic compounds.
An intermolecular competition experiment with a secondary isotope effect of 0.89 is consistent with the known fact that C–N bond formation is the rate-determining step in electrophilic aromatic substitution (Fig. 4b)38. Subsequently, a Hammett study revealed a small negative ρ-value of −0.66 (Fig. 4c). In the presence of reagent 4a, the half-life (t1/2) of the first order reaction on the substrate was ~92 min (Supplementary Fig. 9), while using reagent 4b the reaction rate drastically increased, with a t1/2 of ~8 min (Supplementary Fig. 10).

Owing to its unique properties, HFIP has in recent years proven to be a versatile solvent and additive with broad applications39–41. While HFIP may promote nitration due to its cation stabilization ability, we postulated that its strong hydrogen bonding capacity may also activate 4a.

To further distinguish between a polar electrophilic process or radical mechanism, and to understand the role of HFIP, we turned to density functional theory (DFT) studies. Prior to nitronium transfer we did not observe formation of a π complex, but the NO2 group of 4a was found to interact with HFIP via a hydrogen bond (Supplementary Fig. 15). The reaction then proceeds with a concerted and strongly asynchronous mechanism (Fig. 4e) whereby the C–N bond formation was found to be the rate-determining step (Fig. 4d), in agreement with the first-order reaction in benzene (Supplementary Fig. 9) and the observed KIE (Fig. 4b). In this transformation, the σ complex was not located as a discreet intermediate, although it is observed in the reaction profile and is rapidly followed by H+ elimination assisted by saccharin’s sulfoxide group (Fig. 4d, e)36,42.

Fig. 3 Direct electrophilic nitration of complex molecules and biorelevant compounds using 4a. Standard procedures: I (Hetero)arene (0.5 mmol), 4a (0.65 mmol), HFIP (0.5 M), 55 °C, 3 h ([a], 19 h); II (Hetero)arene (0.5 mmol), 4a (0.65 mmol), [Mg(ClO4)2] (10 mol%), CH3CN (0.5 M), 85 °C, 5 h ([a], 19 h). Yields refer to the isolated product of a single regioisomer or a sum of separately isolated regioisomers. [*] The minor regioisomeric position is labeled.
The reagent’s activity as a controllable source of the nitronium ion is elucidated when considering the high level of organization in the transition state, where HFIP (O atom) interacts with the NO2 group (N atom, 2.79 Å), assisting the cleavage of the N−N bond and the concomitant addition of the nitronium species to the aromatic ring (Fig. 4e). It is this highly ordered transition state that enables delivery of the nitronium ion from 4a in a controlled fashion.

Curious about the high para:ortho ratio in the nitration of tert-butyl-benzene, we calculated the selectivity according to the transition state distribution. A good match was found between the experimental and calculated ratios, and the high para selectivity can be explained by steric hindrance in the transition state, whereby 4a shields one ortho position, and the HFIP molecule the other (Supplementary Fig. 22). This steric hindrance likewise also accounts for the introduction of the nitro group ortho to the ester substituent in 24.

Finally, a trend was found between the activation Gibbs free energy ($\Delta G^{\ddagger}$) and the dissociation Gibbs free energy for the heterolytic cleavage of the N−N bond ($\Delta G_{\text{DISS}}^\ddagger$) for the series of nitrating reagents 1–4. The findings are in agreement with the drastically increased rate of reaction for 4b compared to 4a.

**Fig. 4** Mechanistic studies into the nitration of arenes using 4 and procedure I. a Radical trapping experiments. b Kinetic isotope effect by intermolecular competition experiment. c Hammett study. d Gibbs-free energy profile for nitration of benzene using 4a and 4b. e Transition state of nitronium addition to benzene and intrinsic reaction coordinate for nitration of benzene after rate-determining step.
Discussion

Using the bench-stable and recyclable electrophilic nitrating reagent 4a, we have developed a general protocol for the direct synthesis of a broad range of nitroarenes and nitroheteroarenes demonstrating an exceptional functional group tolerance. Furthermore, our method was applicable to the late-stage C–H functionalization of a range of complex molecules. Mechanistic studies strongly support a classical electrophilic aromatic nitrination, which was found to proceed through a unique and highly sterically demanding transition state. As such, our reagent behaves as a controllable source of the nitronium ion, enabling such a mild and functional group tolerant reaction.

Methods

General procedure I for the nitration of (hetero)arenes. An oven-dried 25 ml micro-vial was charged on the benchtop with a magnetic pTFE-coated stirbar and 4a (148 mg, 0.65 mmol, 1.3 equiv.) and [Mg(ClO₄)₂] (11.2 mg, 0.05 mmol, 10 mol%). The vial was sealed and the atmosphere was cycled 3x with Ar/vac. (Hetero)arene substrate (0.5 mmol, 1.0 equiv.) in HFIP (1 ml) was added by plastic syringe and the reaction mixture was heated at 55 °C with vigorous stirring for 3 h. After cooling to room temperature, the solvent was removed under reduced pressure, and the product was purified by flash column chromatography (SiO₂, ethyl acetate/n-hexane gradient).

General procedure II for the nitration of (hetero)arenes. An oven-dried 25 ml micro-vial was charged on the benchtop with a magnetic pTFE-coated stirbar, 4a (148 mg, 0.65 mmol, 1.3 equiv.) and [Mg(ClO₄)₂] (112 mg, 0.55 mmol, 10 mol%). The vial was sealed and the atmosphere was cycled 3x with Ar/vac. (Hetero)arene substrate (0.5 mmol, 1.0 equiv.) in MeCN (1 ml) was added with a plastic syringe, and the reaction mixture was heated at 85 °C with vigorous stirring for 5 h. After cooling to room temperature, the solvent was removed under reduced pressure, and the product was purified by flash column chromatography (SiO₂, ethyl acetate/n-hexane gradient).

Data availability

The authors declare that the data supporting the findings of this study, including synthetic procedures, NMR spectra, characterization for all new compounds and further details of computational studies, are available in the article and its Supplementary Information, or from the corresponding author upon reasonable request. The X-ray crystallographic coordinates for structures of 4a, 4b, 54, and 61 have been deposited at the Cambridge Crystallographic Data Center (CCDC), under deposition numbers CCDC 1903468, CCDC 1903467, CCDC 1903470, and CCDC 1456828, respectively. These data can be obtained free of charge from The Cambridge Crystallographic Data Center via https://www.ccdc.cam.ac.uk/.

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**Author contributions**

D.K. conceived the project and performed initial reaction optimization; D.K. and R.C. carried out the experiments; K.Z. designed and prepared nitrating reagents; A.P. performed computational studies; D.K. and R.C. wrote the manuscript.

**Additional information**

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