

Regioselective, catalytic 1,1-difluorination of enynes

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Fluorinated small molecules are prevalent across the functional small-molecule spectrum, but the scarcity of naturally occurring sources creates an opportunity for creative endeavour in developing routes to access these important materials. Iodine(I)/iodine(III) catalysis has proven to be particularly well-suited to this task, enabling abundant alkene substrates to be readily intercepted by in situ-generated λ^3 -iodanes and processed to high-value (di)fluorinated products. These organocatalysis paradigms often emulate metal-based processes by engaging the π bond and, in the case of styrenes, facilitating fluorinative phenonium-ion rearrangements to generate difluoromethylene units. Here we demonstrate that enynes are competent proxies for styrenes, thereby mitigating the recurrent need for aryl substituents, and enabling highly versatile homopropargylic difluorides to be generated in an operationally simple manner. The scope of the method is disclosed, together with application in target synthesis (>30 examples, up to >90% yield).

The synergistic interplay of precision synthesis^{1,2} and functional small-molecule design³ continues to be a major driver of innovation in both disciplines. In the vanguard of enabling technologies, fluorination has a venerable history in tailoring the physicochemical traits of promising active pharmaceutical ingredients (APIs), and the societal impact of Fried's seminal work on fluorinated steroids is a compelling exemplar⁴. Function-driven synthesis thus continues to provide a powerful incentive to expand the current methodological arsenal under the auspices of atom and step efficiency⁵. In particular, the success of the geminal-difluoromethylene group in leading pharmaceuticals has stimulated much interest in the development of main group catalysis-based strategies to facilitate installation from readily available precursors^{6–8}.

The prominence of fluorination patterns in contemporary drug discovery^{9–14} disguises the comparative scarcity of naturally occurring organofluorines in marine and terrestrial environments^{15,16}. Although more than 5,000 halogen-containing natural products have been described so far^{17,18}, and fluorine sources are accessible, it is manifest that nature has not been compelled to evolved fluorine biochemistry to any substantial degree^{19,20}. This fluorous juxtaposition between natural and synthetic functional small molecules continues to provide

opportunities for the conception and development of new molecular entities with geometries and physiochemistries that are not encountered in biology^{21–23}, and it logically follows that this continues to expand the chemical space available for function-driven synthesis. A compelling exemplar is the bond-angle distortion that results from CH_2 to CF_2 replacement²⁴, which renders the difluoromethylene group a validated bioisostere of oxygen in phosphate mimics^{25,26}. This motif is finding increasing application in the small-molecule drug repertoire, with prominent examples including lubiprostone (Amitiza), tafluprost (Taflostan) and various 5-HT_{1D} agonists (Fig. 1a)²⁷. Motivated by the demand for new fluorinated modules for medicinal chemistry^{12,14}, and cognizant of the emergent importance of alkyne-containing APIs such as efavirenz (Estiva) and levonorgestrel (PlanB One-Step), it was envisaged that a route to homopropargylic difluorides would address a gap in the discovery portfolio: this would provide isosteric surrogates of propargylic ethers and alcohols in which the electronegativity of the fluorine atoms would emulate the non-bonding electron pairs²⁸.

Of the many enabling innovations that enable direct, geminal difluoromethylation of alkenes, hypervalent iodine (I/III) catalysis has proven particularly powerful^{29–39}. However, a precondition of this

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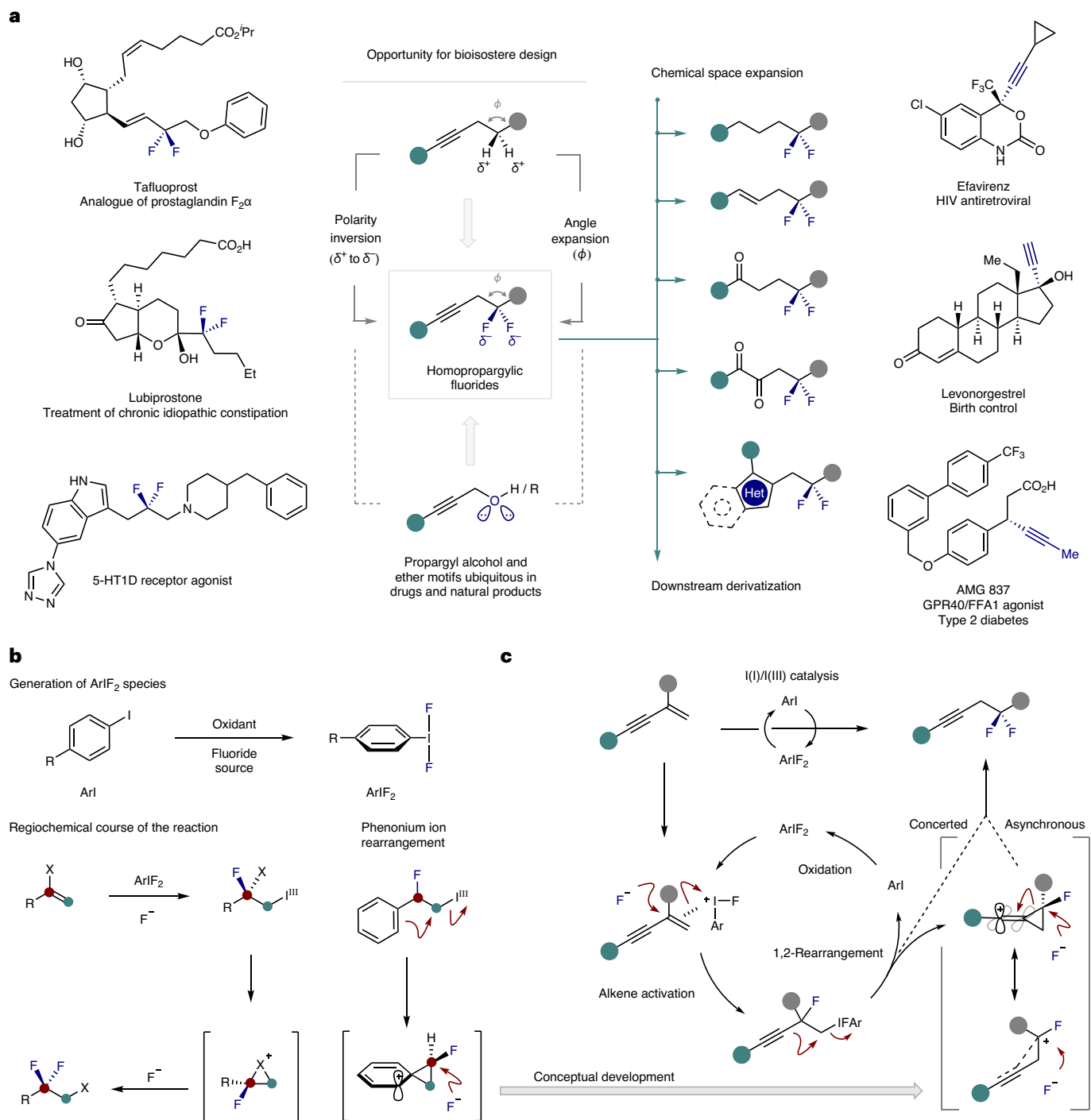
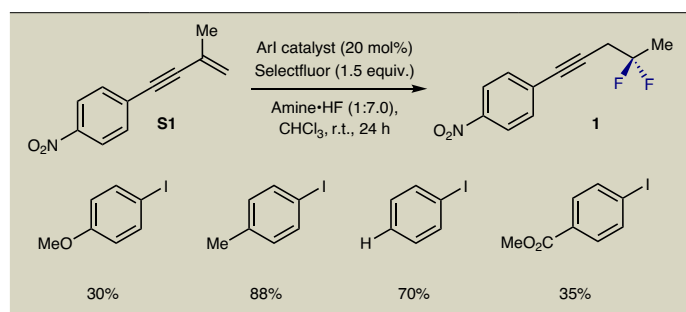


Fig. 1 | Development of a catalytic *gem*-difluorination of enynes. **a, Bioisostere design and examples of bioactive molecules containing a CF_2 or alkyne moiety. **b**, Hypervalent iodine-catalysed *gem*-difluorination of alkenes and the phenonium-ion rearrangement. **c**, Reaction blueprint to enable catalysis-based**

fluorinative alkyne-migration. The electron-rich alkyne is envisaged to be a competent proxy for phenyl, enabling the phenonium-ion rearrangement to be replaced by a formal 1,2-alkynyl shift via a stabilized vinyl cation.

strategy is the requirement for substrates that undergo skeletal rearrangement to ensure that the desired 1,1-regioselectivity of the fluorination is reached (Fig. 1b)⁴⁰. This restraint continues to limit the scope of the transformation to styrene derivatives in which a phenonium-ion rearrangement occurs^{41–45}. Although the introduction of heteroatom substituents partially circumvents this limitation^{35,36,46–48}, geminal difluorination in the presence of carbon-based groups, in the absence of aryl substituents⁴³, remain conspicuously challenging. To address

this, it was envisaged that enynes would be attractive substrates in which the electron-rich alkyne would serve as a phenyl proxy. This would enable the regiochemical paradigm predicated on the phenonium-ion rearrangement to be replaced by a formal 1,2-alkynyl shift via a stabilized vinyl cation (Fig. 1c). Homopropargylic fluorides would also enable direct access to homoallylic and alkyl difluorides, thereby expanding the impact of catalytic difluorinations enabled by I(I)/I(III) catalysis.

Table 1 | Reaction optimization


Entry	Modified conditions	Yield (%)
1	Amine·HF 1:4.5	55
2	Amine·HF 1:9.2	41
3	Oxone as oxidant	51
4	<i>m</i> -CPBA as oxidant ^a	74
5	10 mol% <i>p</i> -Toll	78
6	Reaction at 0 °C	24
7 ^b	No catalyst	<5
8	No oxidant	<5

Standard reaction conditions: enyne **1** (0.1 mmol), catalyst (20 mol%), amine·HF 1:7.0 (0.25 ml), CHCl₃ (0.25 ml) and Selectfluor (0.15 mmol). Yields were determined by ¹⁹F NMR using ethyl fluoroacetate as an internal standard. ^ameta-Chloroperoxybenzoic. ^bAmine·HF ratio changed to 1:7.5.

A catalytic cycle was conceived based on the in situ generation of an ArIF₂ species, via a process of ligand exchange, that would promote an alkene-activation and fluorination sequence. Should the key rearrangement be successful, then the product cation would benefit from fluorine as a stabilizing auxiliary⁴⁹. This would provide a facile route to homopropargylic difluorides, in which the alkyne handle would facilitate downstream functionalization.

Results and discussion

To validate the working hypothesis delineated in Fig. 1c, enyne **1** was prepared and exposed to catalysis conditions using various inexpensive aryl iodides, oxidants and HF sources (Table 1). Initially, *p*-Toll was combined with Selectfluor and amine·HF (1:7 ratio) in chloroform at ambient temperature. This enabled the desired homopropargylic fluoride **1** to be generated in 88% yield. Importantly, the *vicinal* regioisomer was not formed under these conditions, as determined by ¹⁹F NMR (<5%). However, in the absence of the catalyst, the *vicinal* difluoride was formed in 13% yield. A screen of electronically modulated catalysts confirmed the superiority of *p*-Toll, and revealed the following trend: *p*-Me > *p*-H > *p*-CO₂Me > *p*-OMe. Modifying the amine:HF ratio or the oxidant were found to have a detrimental effect on the reaction outcome (Table 1).

Having identified optimized conditions for the title reaction, the scope and limitations of the transformation were investigated. In the course of this process, reactivity differences were noted in response to subtle changes in the amine:HF ratio. This is in line with early observations related to the impact of trifluoroacetic acid on the reactivity of iodobenzene dichloride⁵⁰. For that purpose, a gradient of amine:HF ratios was considered starting from 1:4.5 and increasing to 1:7.0 (denoted A–F). For simplicity, only the most effective conditions are indicated in Table 2. Initially, the impact of modifying the capping aryl group was investigated while keeping the alkene substituent constant (R = Me). This enabled a series of *gem*-difluorides to be generated, and demonstrated functional-group compatibility with electron-withdrawing groups, halogens and small alkyl fragments (**1–10**, up to 83%). In the case of product **2**, it was possible to rigorously

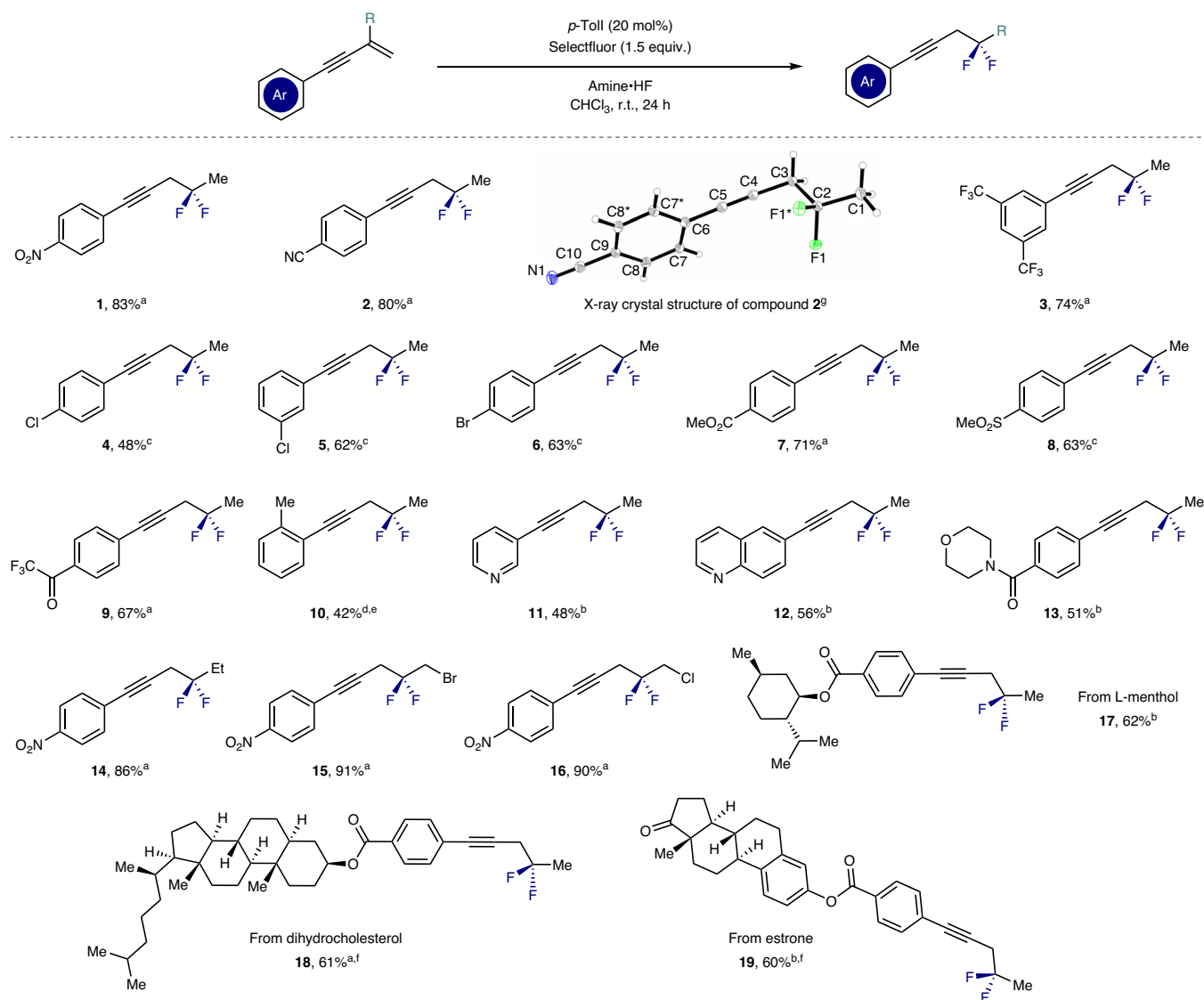
establish the molecular connectivity by single-crystal X-ray diffraction (Table 2; CCDC 2256836). Gratifyingly, the method also proved to be compatible with medicinally relevant heterocycles such as pyridines, quinolines and morpholines (**11–13**, up to 56%). Furthermore, it was possible to replace R = Me with R = CH₂X (X = Br and Cl) to create linchpins that could be functionalized at the proximal C(*sp*³) position (**15** and **16**, up to 91%). Finally, the compatibility of the method with more complex natural-product-derived scaffolds was validated (**17–19**).

To advance the scope of the transformation beyond aryl-substituted enynes, aliphatic derivatives were then explored with a view to applying the method to target synthesis (Table 3). Simple alkyl and cycloalkyl derivatives were tolerated (**20** and **21**, up to 63%) and it was possible to introduce functionality in the form of phthalimides (**22**, 50%) and ethers (**23**, 71%). Substrates with potentially challenging benzylic/propargylic positions such as **24** were smoothly converted to the desired product. The transformation was found to be chemoselective for the enyne versus cinnamoyl motifs (**25**, 64%), and alkyneic esters (**26**, 63%), tosylates (**27**, 56%) and alcohols (**28**, 41%) were compatible. Modifying the alkyl substituent was possible (**29**, **30**) and enabled the 1,2,2-trifluoro motif to be generated in a facile manner. The introduction of more complex heterocycles, such as in febusostat (Adenuric) derivative **32**, is an encouraging validation of the method in a drug-discovery setting. With a view to accessing the parent motif derived from the unsubstituted enyne, the triisopropylsilyl (TIPS)-acetylene **33** was prepared in 67% yield.

To demonstrate the synthetic utility of this geminal difluorination of enynes, two representative experiments were validated on a 4.0 mmol scale (Fig. 2a), and a series of product derivatization reactions were conducted (Fig. 2b). Initially, alkyne **20** was fully and partially reduced⁵¹ to the alkane and alkene products **34** and **35**, respectively. To demonstrate the value of the homopropargylic fluorides in heterocycle formation, compound **4** was converted to the quinoxaline **36** through Ru-catalysed oxidation of the alkyne and concomitant condensation with 1,2-phenyldiamine^{52,53}. Desilylation of compound **33** with tetrabutylammonium fluoride (TBAF) furnished the terminal alkyne **37** in 85% yield: this could then be processed further to triazole **38** via a copper-catalysed click reaction⁵⁴. In situ deprotection of **33** and subsequent Sonogashira cross-coupling proved facile, enabling the electron-rich aryl alkyne **39** to be generated in 64% yield. Because electron-rich enynes undergo uncatalysed side reactions with the Selectfluor⁵⁵, this approach enables the geminal difluorination products to be generated by an alternative route. Finally, the compatibility of the motif under Suzuki–Miyaura conditions was demonstrated through the generation of compound **40** (93% yield)⁵⁶.

A concise route to the CF₂-modified palmitic acid **44** was conceived to validate the method in target synthesis (Fig. 2c). Initially reported by O'Hagan and co-workers in the context of a wider study of the conformational preferences of palmitic acids and nonadecane containing CF₂ groups⁵⁷, this molecule remains a benchmark in difluorination method development⁵⁸. With the aim of complementing the existing reagent-based approaches, enyne **41** was exposed to the catalytic geminal-difluorination conditions: this furnished the key intermediate **42** in 51% yield. Chemoselective reduction of the alkyne and saponification of the methyl ester enabled the desired compound **44** to be generated in 98% yield over two steps.

Finally, control reactions were performed to establish that deletion of the alkyl substituent was tolerated (Fig. 3). Interest in the difluoromethyl group as a surrogate of primary alcohols²⁸ renders such products appealing in the wider context of molecular design on account of their hydrogen-donor character⁵⁹. Pleasingly, both the aryl- and alkyl-substituted enynes **45** and **47** could be processed to their

Table 2 | Establishing the scope of aryl alkynes

Reaction conditions: enyne (0.2 mmol), catalyst (20 mol%), amine-HF (0.5 ml), CHCl₃ (0.5 ml) and Selectfluor (0.3 mmol). Isolated yields are given. ^aAmine:HF=1:7.0. ^bAmine:HF=1:6.0. ^cAmine:HF=1:5.0. ^dAmine:HF=1:4.5. ^e¹⁹F NMR yield using ethyl fluoroacetate as an internal standard. 23% of the vicinal difluorination product was formed due to an uncatalysed background reaction (details are provided in Supplementary section 1.3). ^fReaction performed on a 0.10 mmol scale. ^gThermal ellipsoids are shown at 50% probability. Care should be exercised during isolation due to the volatility of many of the products.

respective homopropargyl difluoride products **46** and **48**, respectively. Replacing the substituent with an aryl group (**49** Ar = *p*-CF₃) was then explored to identify which regioisomer was predominantly formed. The isolation of compound **50** as the sole product of the reaction (40% yield) is consistent with the 1,2-shift out-competing phenonium-ion rearrangement. The skeletal rearrangement that is central to the working hypothesis was supported by deuterium labelling to generate **29-d** (56%, 76% D incorporation; Fig. 3c).

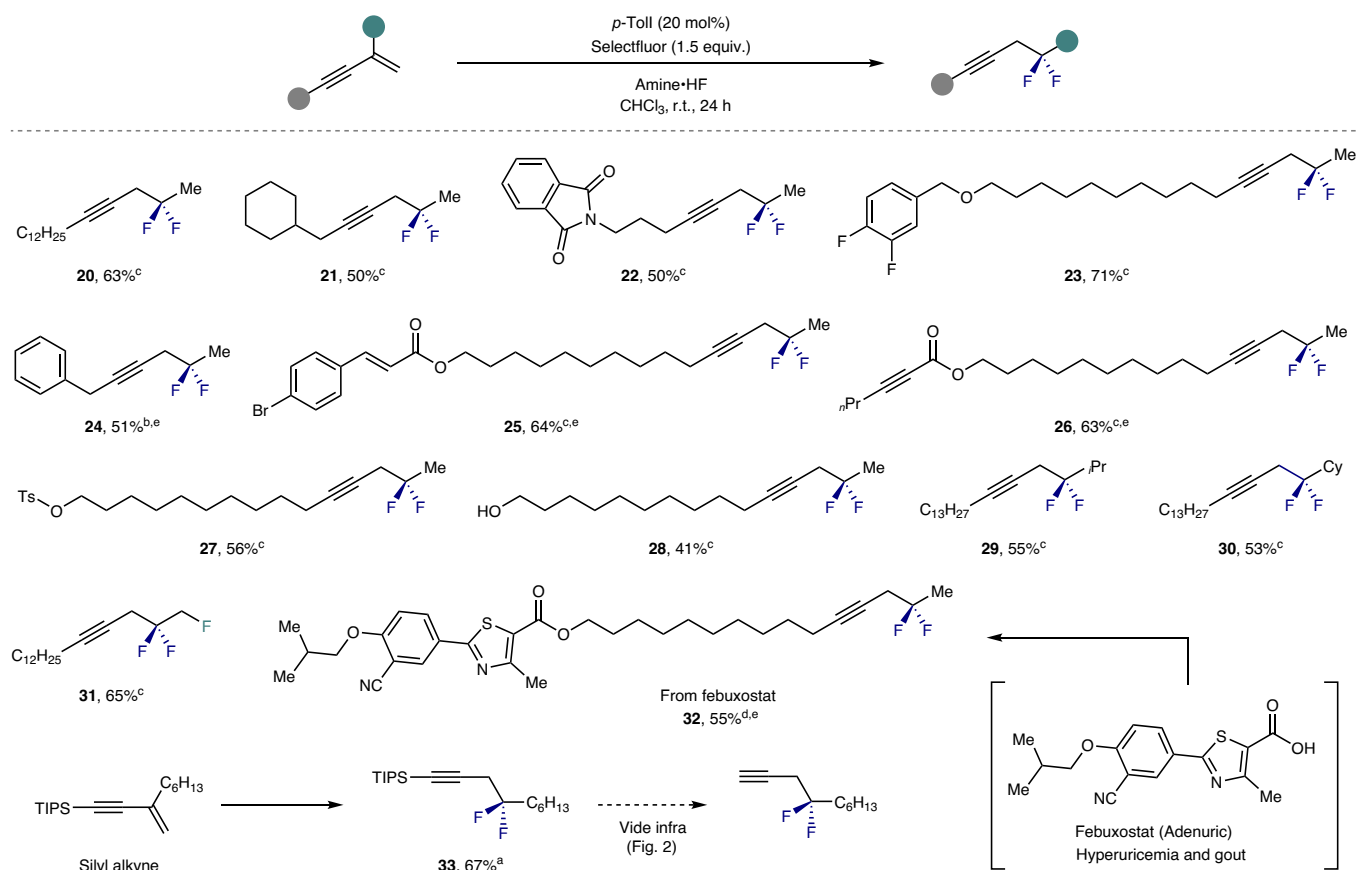
Conclusions

The direct, geminal difluorination of alkenes under the auspices of hypervalent iodine catalysis remains a powerful paradigm to expand organofluorine chemical space for contemporary drug discovery. In situ-generated λ^3 -iodanes regulate regiocontrol by inducing C(*sp*³)-F bond-forming/rearrangement sequences with exquisite efficiency: the latter step is conditional on substrates that are predisposed to undergo a phenonium-ion rearrangement. To circumvent this limitation, enynes have been validated as competent substrates that

deliver the desired 1,1-selectivity, where the phenonium-ion rearrangement can be replaced by a formal 1,2-shift of the alkyne. Computational support for the tentative mechanism outlined in Fig. 1 is available in Supplementary section 1.7. Utilizing the alkyne as a phenyl proxy, it has been possible to achieve the title reaction and deliver homopropargylic difluorides that are highly amenable to downstream functionalization. A broad substrate scope is demonstrated (>30 examples) together with selected derivatization protocols, as well as a short, catalysis-based synthesis of CF₂-modified palmitic acid. It is envisaged that this enabling method will find application in the conception of new drug-discovery modules.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41557-023-01344-5>.

Table 3 | Expanding the scope to alkyl and silyl alkynes

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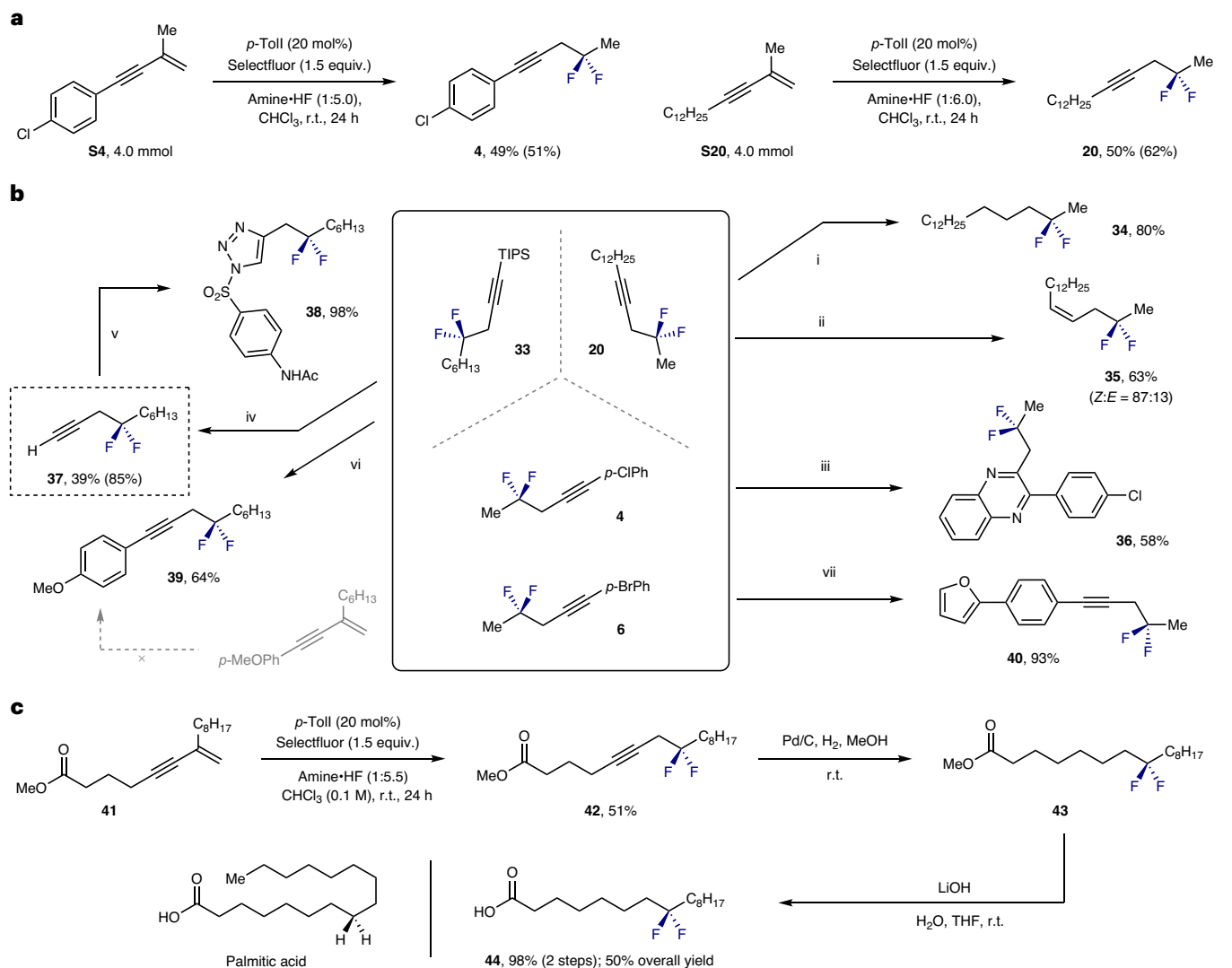


Fig. 2 | Synthetic applications. **a**, Scale-up experiments. **b**, Product derivatization. Conditions. (i) Conversion of **20** to form **34**: Pd/C (10 mol%), H₂, MeOH (0.1 M), r.t., 24 h. (ii) Semi-reduction to generate the *Z*-alkene **35**: TiCl₂Cp₂ (10 mol%), LiAlH₄ (2.0 equiv.), tetrahydrofuran (THF) (0.2 M), r.t., overnight. (iii) Generation of quinoxaline **36**: (1) RuCl₃ (1 mol%), PhI(OAc)₂ (3.0 equiv.), DCM, H₂O (4:1, 0.2 M), r.t., 3 h; (2) saccharin (5 mol%), 1,2-phenylenediamine (1.1 equiv.), MeOH (0.2 M), r.t., 12 h. (iv) TIPS deprotection to generate terminal alkyne **37**: TBAF (2.0 equiv.), THF (0.4 M), r.t., 2 h. (v) Formation of triazole **38** via a copper-catalysed click reaction: CuTc (10 mol%), 4-acetamidobenzenesulfonyl

azide (1.2 equiv.), toluene (0.2 M), r.t., 12 h. (vi) In situ deprotection of **33** and subsequent Sonogashira cross-coupling: Pd(PPh₃)₂Cl₂ (4.5 mol%), CuI (5 mol%), 4-iodoanisole (1.2 equiv.), NEt₃ (7.0 equiv.), TBAF (2.0 equiv.), THF (0.2 M), 45 °C, 12 h. (vii) Suzuki-coupling to generate compound **40**: Pd(PPh₃)₄ (10 mol%), 2-furanboronic acid (2.5 equiv.), K₂CO₃ (2.5 equiv., 2 M in H₂O), dimethoxyethane (DME) (0.2 M), 85 °C, overnight. **c**, Synthesis of CF₂-modified palmitic acid. Isolated yields are given. ¹⁹F NMR yields are given in parentheses and determined by ¹⁹F NMR using ethyl fluoroacetate as an internal standard.

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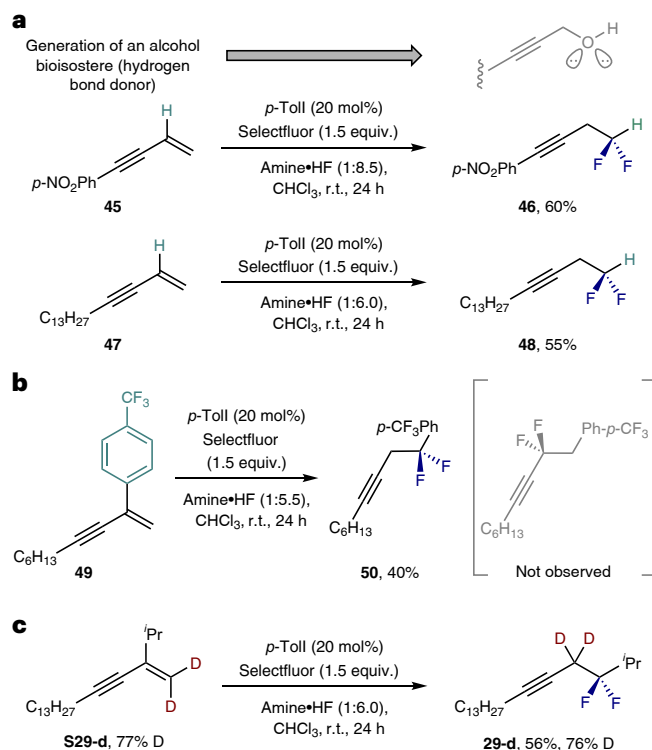


Fig. 3 | Control experiments. **a**, Removal of the alkyl group. Enynes **45** and **47** could be converted to **46** and **48**, containing the difluoromethyl group, respectively. **b**, Investigation of regioselectivity. Compound **50** was obtained as the sole product. **c**, Deuterium labelling experiment. Deuterium atom incorporated exclusively at the propargylic position, unambiguously demonstrating that 1,2-alkynyl migration took place. Isolated yields are given.

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Methods

General procedure for 1,1-difluorination of enynes

Unless otherwise stated, a Teflon vial was equipped with a 1-cm stirring bar followed by the addition of enyne (0.2 mmol, 1.0 equiv.), *p*-iodotoluene (9 mg, 0.04 mmol, 20 mol%) and CHCl₃ (0.5 ml). The stated amine:HF mixture was added (0.5 ml) via syringe. After stirring for 1 min, Selectfluor (106 mg, 0.3 mmol, 1.5 equiv.) was added in one portion. The reaction vessel was then sealed with a Teflon screw cap. After stirring (350 r.p.m.) at ambient temperature for 24 h, the reaction mixture was poured into 100 ml of a saturated solution of NaHCO₃ (caution! generation of CO₂!). The Teflon vial was rinsed with dichloromethane (DCM) and dropped into another flask of saturated aqueous solution of NaHCO₃ to guarantee the removal of excess HF. The organics were extracted with DCM (3 × 30 ml), the combined organic layers were dried over Na₂SO₄, filtered, and the solvent was carefully removed under reduced pressure. An internal standard (ethyl fluoroacetate) was added to the crude residue and the NMR yield was analysed by ¹⁹F NMR spectroscopy against the internal standard. The NMR sample was recombined with the crude residue and purification by column chromatography or preparative thin-layer chromatography yielded the desired product.

Data availability

Crystallographic data for the structure reported in this Article have been deposited at the Cambridge Crystallographic Data Centre, under deposition nos. CCDC 2256836 (2). Copies of the data can be obtained free of charge via <https://www.ccdc.cam.ac.uk/structures/>. All data are available in the main text or the Supplementary Information.

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Author contributions

Conceptualization was provided by Z.-X.W. and R.G., and methodology by Z.-X.W., K.L., C.H., C.G.D., C.M.-L. and R.G. Investigations were carried out by Z.-X.W., C.H., K.L., C.G.D., C.M.-L. and R.G. Funding acquisition was performed by Z.-X.W. and R.G. Project administration was carried out by Z.-X.W., K.L. and R.G. The manuscript was written by Z.-X.W. and R.G.

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Competing interests

The authors declare no competing interests.

Additional information

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