A Chiral Pentfluorinated Isopropyl Group via Iodine(I)/(III) Catalysis
Stephanie Meyer, Joel Häfliger, Michael Schäfer, John J. Molloy, Constantin G. Daniliuc, and Ryan Gilmour*

Abstract: An I(I)/(III) catalysis strategy to construct an enantioenriched fluorinated isostere of the iPr group is reported. The difluorination of readily accessible α-CF3-styrenes is enabled by the in situ generation of a chiral ArIF2 species to forge a stereocentre with the substituents F, CH2F and CF3 (up to 95%, >20:1 vicinal:geminal difluorination). The replacement of the metabolically labile benzylic proton results in a highly preorganised scaffold as was determined by X-ray crystallography (π–α* and stereoelectronic gauche σ–α* interactions). A process of catalyst editing is disclosed in which preliminary validation of enantioselectivity is placed on a structural foundation.

Short, unfunctionalised aliphatic groups (C1–C4) are ubiquitous structural features in the natural product repertoire, and are particularly conspicuous in polyketides and terpenes.[1] This is a logical consequence of iterative biosynthesis algorithms that process low molecular weight fragments into higher homologues.[2] Introduced under the auspices of acetyl- or propionyl-CoA,[3] complemented by electrophilic para- digms involving methyltransferases (SAM),[4] these motifs appear to be vestigial in nature. However, they often encode for a highly specific function and thus delineating their biosynthetic origins has been intensively pursued. Indeed the value of harnessing small aliphatic groups to enhance the physicochemical profiles of drug candidates is exemplified by the "magic methyl effect".[5] Interrogating the stereochemical course of enzymatic methylation has a venerable history, due to the achiral nature of this motif and the pre-conditions associated with designing a chiral bioisostere to track the possible translation of stereochemical information.[6,7] Arigoni’s celebrated synthesis of chiral acetic acid remains a tour de force in stereocatalystic synthesis, and a master class in orbital symmetry to craft an isotopically orthogonal motif (1H, 2H and 3H, Figure 1, top).[8] Whilst this isoisostrateg strategy remains expansive in the field of mechanistic enzymology, small fragment-based bioisosterism in drug design relies on stable isotopes to enhance the pharmaco-inertics and -dynamics of drug candidate performance.[9] Molecular editing with fluorine (H=F) has proven to be particularly effective,[10] and is reflected in the increasing number of fluorinated small molecules reaching the market.[11] This is a consequence of fluorine’s low steric demand, low polarization and the stability of the C-F bond. Given the success of achiral perfluoroalkyl groups in drug discovery, catalysis and agrochemistry,[12] routes to small, chiral, 3D fluorooalkane motifs would be advantageous to expand the available chemical space. This includes the C3 (BITE group)[13] which is a bioisosteric hybrid of the ethyl and trifluoromethyl groups.[14] Cognisant of the prevalence of (C3) isopropyl units in active natural product leads and small molecule synthesis:
pharmaceuticals (Figure 1 centre), a catalysis-based strategy to access a differentially fluorinated analogue of the Pr group was initiated. Harnessing I(II)/(III) catalysis, it was envisaged that a formal 1,2-addition of fluorine across the alkene moieties of simple α-trifluoromethyl styrenes would generate a stereogenic centre bearing F, CH₂F and CF₃ groups (Figure 1, bottom).

The success of this catalysis-based approach would be contingent on the in situ oxidation of a chiral aryl iodide organocatalyst to generate an ArIF₂ species that would be sufficiently active to engage a sterically-congested, electron-deficient alkene. If successful, the resulting pentafluoroisopropyl surrogate would constitute a chiral C₃ building block in which the lability of the methine proton is mitigated.

Moreover, the constituent hyperconjugative interactions intrinsic to the internal vicinal-difluoro motif would manifest themselves in conformation. To identify conditions that would enable the target fluorinated isopropyl motif to be generated from simple α-trifluoromethyl styrenes, a process of reaction optimisation was conducted (Figure 2, 1a→2a).

To that end, simple aryl iodides were investigated as inexpensive catalysts in conjunction with Selectfluor as the terminal oxidant to generate the key ArI(III)F₂ species. Initial studies were performed in chloroform at ambient temperature using an amine:HF ratio of 1:7.5 and the reactions were examined by ¹⁹F NMR spectroscopy using an internal standard. Iodobenzene proved to be a perfectly effective catalyst for this transformation to generate (±)-2a and 3a in a 2.5:1 ratio (86% combined yield). The latter product arises from phenonium ion rearrangement and has been exploited in a range of catalysis-based geminal difluorination processes.

Repeating the reaction with p-iodotoluene (5) led to a notable improvement in yield (>95%) with comparable regioselectivity in favour of the desired vicinal product 2a (2.2:1). Electronic modulation was not well tolerated with the ester derivative 6 proving to be a less active catalyst under comparable conditions (26%). Control experiments in the absence of catalyst led to <5% yield and demonstrate the strongly deactivating nature of the trifluoromethyl group that inhibits background reactions such as those reported by Lal and co-workers using HF sources and Selectfluor.

To explore the scope and limitations of this catalysis-based difluorination of α-CF₃-styrenes, the effect of Brønsted acidity was probed as a function of the amine:HF ratio. The success of this catalysis-based approach would be contingent on the in situ oxidation of a chiral aryl iodide organocatalyst to generate an ArIF₂ species that would be sufficiently active to engage a sterically-congested, electron-deficient alkene. If successful, the resulting pentafluoroisopropyl surrogate would constitute a chiral C₃ building block in which the lability of the methine proton is mitigated. Moreover, the constituent hyperconjugative interactions intrinsic to the internal vicinal-difluoro motif would manifest themselves in conformation. To identify conditions that would enable the target fluorinated isopropyl motif to be generated from simple α-trifluoromethyl styrenes, a process of reaction optimisation was conducted (Figure 2, 1a→2a).

To that end, simple aryl iodides were investigated as inexpensive catalysts in conjunction with Selectfluor as the terminal oxidant to generate the key ArI(III)F₂ species. Initial studies were performed in chloroform at ambient temperature using an amine:HF ratio of 1:7.5 and the reactions were examined by ¹⁹F NMR spectroscopy using an internal standard. Iodobenzene proved to be a perfectly effective catalyst for this transformation to generate (±)-2a and 3a in a 2.5:1 ratio (86% combined yield). The latter product arises from phenonium ion rearrangement and has been exploited in a range of catalysis-based geminal difluorination processes.

Repeating the reaction with p-iodotoluene (5) led to a notable improvement in yield (>95%) with comparable regioselectivity in favour of the desired vicinal product 2a (2.2:1). Electronic modulation was not well tolerated with the ester derivative 6 proving to be a less active catalyst under comparable conditions (26%). Control experiments in the absence of catalyst led to <5% yield and demonstrate the strongly deactivating nature of the trifluoromethyl group that inhibits background reactions such as those reported by Lal and co-workers using HF sources and Selectfluor.

To explore the scope and limitations of this catalysis-based difluorination of α-CF₃-styrenes, the effect of Brønsted acidity was probed as a function of the amine:HF ratio. The success of this catalysis-based approach would be contingent on the in situ oxidation of a chiral aryl iodide organocatalyst to generate an ArIF₂ species that would be sufficiently active to engage a sterically-congested, electron-deficient alkene. If successful, the resulting pentafluoroisopropyl surrogate would constitute a chiral C₃ building block in which the lability of the methine proton is mitigated. Moreover, the constituent hyperconjugative interactions intrinsic to the internal vicinal-difluoro motif would manifest themselves in conformation. To identify conditions that would enable the target fluorinated isopropyl motif to be generated from simple α-trifluoromethyl styrenes, a process of reaction optimisation was conducted (Figure 2, 1a→2a).

To that end, simple aryl iodides were investigated as inexpensive catalysts in conjunction with Selectfluor as the terminal oxidant to generate the key ArI(III)F₂ species. Initial studies were performed in chloroform at ambient temperature using an amine:HF ratio of 1:7.5 and the reactions were examined by ¹⁹F NMR spectroscopy using an internal standard. Iodobenzene proved to be a perfectly effective catalyst for this transformation to generate (±)-2a and 3a in a 2.5:1 ratio (86% combined yield). The latter product arises from phenonium ion rearrangement and has been exploited in a range of catalysis-based geminal difluorination processes.

Repeating the reaction with p-iodotoluene (5) led to a notable improvement in yield (>95%) with comparable regioselectivity in favour of the desired vicinal product 2a (2.2:1). Electronic modulation was not well tolerated with the ester derivative 6 proving to be a less active catalyst under comparable conditions (26%). Control experiments in the absence of catalyst led to <5% yield and demonstrate the strongly deactivating nature of the trifluoromethyl group that inhibits background reactions such as those reported by Lal and co-workers using HF sources and Selectfluor.

To explore the scope and limitations of this catalysis-based difluorination of α-CF₃-styrenes, the effect of Brønsted acidity was probed as a function of the amine:HF ratio. The success of this catalysis-based approach would be contingent on the in situ oxidation of a chiral aryl iodide organocatalyst to generate an ArIF₂ species that would be sufficiently active to engage a sterically-congested, electron-deficient alkene. If successful, the resulting pentafluoroisopropyl surrogate would constitute a chiral C₃ building block in which the lability of the methine proton is mitigated. Moreover, the constituent hyperconjugative interactions intrinsic to the internal vicinal-difluoro motif would manifest themselves in conformation. To identify conditions that would enable the target fluorinated isopropyl motif to be generated from simple α-trifluoromethyl styrenes, a process of reaction optimisation was conducted (Figure 2, 1a→2a).

To that end, simple aryl iodides were investigated as inexpensive catalysts in conjunction with Selectfluor as the terminal oxidant to generate the key ArI(III)F₂ species. Initial studies were performed in chloroform at ambient temperature using an amine:HF ratio of 1:7.5 and the reactions were examined by ¹⁹F NMR spectroscopy using an internal standard. Iodobenzene proved to be a perfectly effective catalyst for this transformation to generate (±)-2a and 3a in a 2.5:1 ratio (86% combined yield). The latter product arises from phenonium ion rearrangement and has been exploited in a range of catalysis-based geminal difluorination processes.

Repeating the reaction with p-iodotoluene (5) led to a notable improvement in yield (>95%) with comparable regioselectivity in favour of the desired vicinal product 2a (2.2:1). Electronic modulation was not well tolerated with the ester derivative 6 proving to be a less active catalyst under comparable conditions (26%). Control experiments in the absence of catalyst led to <5% yield and demonstrate the strongly deactivating nature of the trifluoromethyl group that inhibits background reactions such as those reported by Lal and co-workers using HF sources and Selectfluor.

To explore the scope and limitations of this catalysis-based difluorination of α-CF₃-styrenes, the effect of Brønsted acidity was probed as a function of the amine:HF ratio.
This led to the identification of methods A, B and C, reflecting amine:HF ratios of 1:4.5, 1:6 and 1:7.5 respectively (Figures 3 and 4). Method A proved to be highly effective in generating the electron rich products 2b–2d with high levels of regioselectivity favouring formation of the desired vicinal product (> 20:1, up to 86 %). The presence of the CF₃ group clearly distinguished this substrate class from the parent styrenes, which rearrange to generate the geminal product.[20] Control reactions again confirmed the necessity for the catalyst. The seemingly subtle change to Method B proved to be optimal for substrates 2e–2k, enabling the generation of alkyl derivatives (2e–2h, up to >20:1, vic:gem) as well as the electron deficient aniline derivative 2i (74 %, 15.5:1).

Comparable efficiency and selectivity were also noted for the biphenyl system 2j and adduct 2k. Augmenting the amine:HF ratio further to Method C provided optimised conditions to access products 2a,l–2q. Whereas employing higher HF ratios/Bronsted acids under the conditions developed by this laboratory tend to favour 1,1-difluorination,[20] electron-deficient a-CF₃-styrenes proved to be notably more recalcitrant to rearrangement and the vicinal products predominated throughout. Given the importance of aryl bromides in contemporary medicinal chemistry, where the C(sp²)-Br provides a handle for subsequent cross-coupling, the synthesis of 2I was conducted on a 1 mmol scale. Despite the volatility of the product, the vicinal product could be isolated in 43 % yield. Products 2m, 2n and 2o behaved similarly and were generated in a vicinal:geminal ratio of ca. 3:1. Given the prominence of aniline fragments bearing isopropyl units in drug and agrochemical discovery (See Figure 1), the phthalimide 2p was generated cleanly in 61 % yield. Finally, access to the disubstituted aryl 2q was realised, this time with an amine:HF ratio of 1:7.5 in CHCl₃ at ambient temperature. Initially, the effect of modifying the substituent X was assessed using the methyl esters 10–13. Counterintuitively, augmenting the steric footprint at site X had a detrimental effect on selectivity. Catalyst 10 (X = Me) proved to be most effective, generating compound 2a with 86:14 e.r. (> 95 % conversion, 88 % combined yield). Structural editing at site Y was not tolerated as exemplified by catalysts 14–16. As a control series, the C₂-symmetric catalysts 17–19 were examined (Figure 5, lower). Direct comparison of 17 with the most promising scaffold 10 confirmed the importance of C₂ symmetry (72:28 versus 86:14 e.r.). Interestingly, substituting the methyl ester for benzyl (catalyst 18) did not erode selectivity, although efficiency was decreased. Moreover, the a-Bn catalyst (19) proved to be less efficient than the C₂-symmetric derivative 12. Having identified catalyst 10 as the most promising scaffold to validate an effective, balancing the electronic effects of the resorcinol with a p-CO₂Me in catalyst 9 led to notably superior catalysis (87 % yield, vicinal:geminal 3:1). As the logical next step, C₂-symmetric resorcinol derivatives were investigated as summarised in Figure 5.[23,24] Reactions were performed under standard conditions with an amine:HF ratio of 1:7.5 in CHCl₃ at ambient temperature. Initially, the effect of modifying the substituent X was assessed using the methyl esters 10–13. Counterintuitively, augmenting the steric footprint at site X had a detrimental effect on selectivity. Catalyst 10 (X = Me) proved to be most effective, generating compound 2a with 86:14 e.r. (> 95 % conversion, 88 % combined yield). Structural editing at site Y was not tolerated as exemplified by catalysts 14–16. As a control series, the C₂-symmetric catalysts 17–19 were examined (Figure 5, lower). Direct comparison of 17 with the most promising scaffold 10 confirmed the importance of C₂ symmetry (72:28 versus 86:14 e.r.). Interestingly, substituting the methyl ester for benzyl (catalyst 18) did not erode selectivity, although efficiency was decreased. Moreover, the a-Bn catalyst (19) proved to be less efficient than the C₂-symmetric derivative 12. Having identified catalyst 10 as the most promising scaffold to validate an effective, balancing the electronic effects of the resorcinol with a p-CO₂Me in catalyst 9 led to notably superior catalysis (87 % yield, vicinal:geminal 3:1). As the logical next step, C₂-symmetric resorcinol derivatives were investigated as summarised in Figure 5.[23,24] Reactions were performed under standard conditions with an amine:HF ratio of 1:7.5 in CHCl₃ at ambient temperature. Initially, the effect of modifying the substituent X was assessed using the methyl esters 10–13. Counterintuitively, augmenting the steric footprint at site X had a detrimental effect on selectivity. Catalyst 10 (X = Me) proved to be most effective, generating compound 2a with 86:14 e.r. (> 95 % conversion, 88 % combined yield). Structural editing at site Y was not tolerated as exemplified by catalysts 14–16. As a control series, the C₂-symmetric catalysts 17–19 were examined (Figure 5, lower). Direct comparison of 17 with the most promising scaffold 10 confirmed the importance of C₂ symmetry (72:28 versus 86:14 e.r.). Interestingly, substituting the methyl ester for benzyl (catalyst 18) did not erode selectivity, although efficiency was decreased. Moreover, the a-Bn catalyst (19) proved to be less efficient than the C₂-symmetric derivative 12. Having identified catalyst 10 as the most promising scaffold to validate an effective, balancing the electronic effects of the resorcinol with a p-CO₂Me in catalyst 9 led to notably superior catalysis (87 % yield, vicinal:geminal 3:1). As the logical next step, C₂-symmetric resorcinol derivatives were investigated as summarised in Figure 5.[23,24] Reactions were performed under standard conditions with an amine:HF ratio of 1:7.5 in CHCl₃ at ambient temperature. Initially, the effect of modifying the substituent X was assessed using the methyl esters 10–13. Counterintuitively, augmenting the steric footprint at site X had a detrimental effect on selectivity. Catalyst 10 (X = Me) proved to be most effective, generating compound 2a with 86:14 e.r. (> 95 % conversion, 88 % combined yield). Structural editing at site Y was not tolerated as exemplified by catalysts 14–16. As a control series, the C₂-symmetric catalysts 17–19 were examined (Figure 5, lower). Direct comparison of 17 with the most promising scaffold 10 confirmed the importance of C₂ symmetry (72:28 versus 86:14 e.r.). Interestingly, substituting the methyl ester for benzyl (catalyst 18) did not erode selectivity, although efficiency was decreased. Moreover, the a-Bn catalyst (19) proved to be less efficient than the C₂-symmetric derivative 12. Having identified catalyst 10 as the most promising scaffold to validate an effective, balancing the electronic effects of the resorcinol with a p-CO₂Me in catalyst 9 led to notably superior catalysis (87 % yield, vicinal:geminal 3:1). As the logical next step, C₂-symmetric resorcinol derivatives were investigated as summarised in Figure 5.[23,24] Reactions were performed under standard conditions with an amine:HF ratio of 1:7.5 in CHCl₃ at ambient temperature. Initially, the effect of modifying the substituent X was assessed using the methyl esters 10–13. Counterintuitively, augmenting the steric footprint at site X had a detrimental effect on selectivity. Catalyst 10 (X = Me) proved to be most effective, generating compound 2a with 86:14 e.r. (> 95 % conversion, 88 % combined yield). Structural editing at site Y was not tolerated as exemplified by catalysts 14–16. As a control series, the C₂-symmetric catalysts 17–19 were examined (Figure 5, lower). Direct comparison of 17 with the most promising scaffold 10 confirmed the importance of C₂ symmetry (72:28 versus 86:14 e.r.). Interestingly, substituting the methyl ester for benzyl (catalyst 18) did not erode selectivity, although efficiency was decreased. Moreover, the a-Bn catalyst (19) proved to be less efficient than the C₂-symmetric derivative 12. Having identified catalyst 10 as the most promising scaffold to validate an
enantioselective process (please see the ESI for additional details) a representative selection of α-CF₃-styrenes were subjected to the general catalysis conditions using 10 (Figure 6).

Gratifyingly, the methyl derivative underwent smooth difluorination to generate 2e (> 95%, 17.1 vicinal:geminal, 83:17 e.r.). The fluorinated biaryl system 2j was compatible with the conditions and could be prepared with a regioselectivity of 7:1 vicinal:geminal and 85:15 e.r. after recrystallisation. Gratifyingly, compounds 2j and 2p were crystalline allowing the (S)-configuration of the new stereocentre to be assigned (vide infra). [25] Finally, the phthalimide derivative 2p was processed to an analogue of the TRPA1 antagonist HC-030031 in a short synthetic sequence (Figure 6).

To complement the plenum of methods available to construct short, unfunctionalised aliphatic groups for drug discovery, a catalysis-based strategy to access chiral, fluorinated surrogates of the isopropyl group has been developed. This serves to expands the current portfolio of fluorine drug modules for drug discovery (Figure 7, centre). [26] Despite the intrinsic steric and electronic challenges associated with generating highly fluorinated stereocentres, this I(I)/I(III) catalysis platform enables α-CF₃-styrenes to undergo smooth vicinal difluorination (up to > 20:1 vicinal:geminal). Importantly, the CF₃ group effectively inhibits the dominant phenonium ion rearrangement associated with electron rich styrenes, allowing products such as 2b to be generated with excellent levels of regiocontrol (> 20:1). Finally, preliminary validation of an enantioselective variant is disclosed. Whilst the sterically demanding phenyl and trifluoromethyl substituents (VvdW (CF₃) = 39.8 Å⁸/C₁₃₈₃)[10e,27] render this intermolecular process challenging, it is gratifying to observe encouraging levels of enantioselectivity. A tentative induction model is proposed in which facial discrimination in the enantiodetermining fluorination is a precondition of selectivity. Since X-ray analyses of 2j and 2p confirm that the major enantiomer is (S)-configured (Figure 7), it is conceivable that stabilising electrostatic interactions (RCF₂d/C₀F···d+H-CH₂R), [28] may bias catalyst-substrate preorganisation. [29] Simple steric discrimination (CF₃ vs. Ph) is not consistent with the selectivities observed the C₁-symmetric catalyst. The solid-state analysis also reveals a stereoelectronic gauche effect (α-α*-; θ_FCCF = 69.9° and 51° for 2j and 2p, respectively) and that the CF₃ group is orthogonal to the plane of the π system (π-π*-). [30]

Exploring the physicochemical profile of this new motif in the context of drug discovery and contemporary agrochemistry is the focus of ongoing studies and will be reported in due course.

Figure 6. The yield is the sum of vicinal and geminal difluorination products. The regioselectivity ratio (vic:gem) and yield were determined by ¹⁹F NMR spectroscopy using α,α,α-trifluorotoluene as internal standard. Isolated yield of the vicinal product is given in parentheses. Enantioselectivity determined by chiral HPLC. *After recrystallisation. N.B.: The products are often highly volatile and care must be taken in the isolation.

Figure 7. Postulated induction model and the X-ray structure of compound 2j and 2p. Thermal ellipsoids are shown at 15% probability. CCDC 2044630 (2j) and 2044631 (2p).
Acknowledgements

We acknowledge generous financial support from the Westfälische Wilhelms-Universität Münster, the European Research Council (ERC Consolidator Grant—Project Number 818949-RECON, to RG), the DFG (Cluster of Excellence “Cells in Motion—CiM” (FF-2013-10) and SFB 858) and the Alexander von Humboldt Foundation (post-doctoral research fellowship to JMJ). We thank Mr. Tomáš Neveselý (WWU Münster) for helpful discussions. Open access funding enabled and organized by Project DEAL.

Conflict of interest

The authors declare no conflict of interest.

Keywords: agrochemistry · bioisostere · conformation · fluorine · organocatalysis