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One-pot multistep synthesis of 1-fluoroalkylisoquinolines and fused fluoroalkylpyridines from *N*-fluoroalkyl-1,2,3-triazoles†

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An efficient one-pot microwave-assisted potassium fluoride-mediated synthesis of 1-fluoroalkyl-3-fluoroisoquinolines and fused fluoroalkylpyridines from *N*-fluoroalkylated 1,2,3-triazoles was developed. The reaction has a wide scope and allows the preparation of structurally diverse 3-fluoroisoquinolines with a fluoroalkyl group in position 1, a substituent in position 4 and a substituent on the fused benzene (or heteroaromatic) ring. *N*-Fluoroalkylated ketenimines, which undergo stereoselective formal 1,3-fluorine shift to difluoroazadienes, were identified as intermediates in the reaction sequence. The presence of fluorine in position 3 and a halogen in position 4 of the resulting isoquinolines allowed for further modification by nucleophilic aromatic substitution and cross-coupling reactions, respectively. The developed methodologies were utilized for the synthesis of derivatives of drug candidates.

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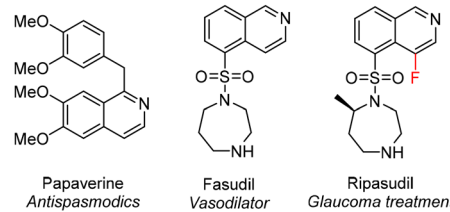
Introduction

The isoquinoline core is present in a variety of drugs and a large number of naturally occurring alkaloids, which in many cases possess compelling biological activities (Fig. 1).^{1–4} Numerous synthetic approaches exist leading to these benzopyridines, such as multistep sequences of reactions including the well-known Bischler–Napieralski,⁵ Pomeranz–Fritsch⁶ and Pictet–Spengler⁷ reactions, as well as processes involving transition metal catalysis.⁸

The introduction of fluorine atoms or fluoroalkyl groups into a lead molecule is a widely used strategy to enhance the pharmacologically relevant properties⁹ and several 1-trifluoromethylisoquinolines exploit this trend (Fig. 2), for example valiglurax¹⁰ – a positive allosteric modulator of mGlu4 receptors and a candidate for the treatment of Parkinson's disease. Yet, the procedures for their preparation remain underdeveloped, substrate-specific, low-yielding, or require expensive, non-selective and atom non-economical fluoroalkylation methods or transition metal catalysts.^{13,14}

The first multi-step approach leading to 1-perfluoroalkylisoquinolines with the Bischler–Napieralski type cyclization was demonstrated by Pastor¹⁵ in 1979. A similar approach was also used in 2019 by Lindsley (Scheme 1A) for the preparation of valiglurax, which allowed the construction of the isoquinoline core in an overall 21% yield.¹⁰ Another possible approach involves direct C–H bond perfluoroalkylation of isoquinolines¹⁶ (Scheme 1B) or isoquinoline-*N*-oxides.^{17,18} Trifluoromethylation *via* a coupling reaction of iodoisoquinolines with copper^{19–22} or palladium²³ catalysts was also reported. However, the most common strategy towards 1-fluoroalkylated isoquinolines involves the insertion of fluoroalkyl radicals into isonitriles, followed by radical cyclization (Scheme 1C).^{13,24–29}

Herein, we report a high-yielding and novel one-pot synthetic strategy to prepare substituted 1-fluoroalkyl-3-fluoroisoquinolines and fused fluoroalkylpyridines.



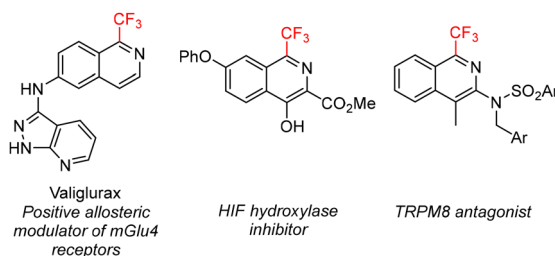
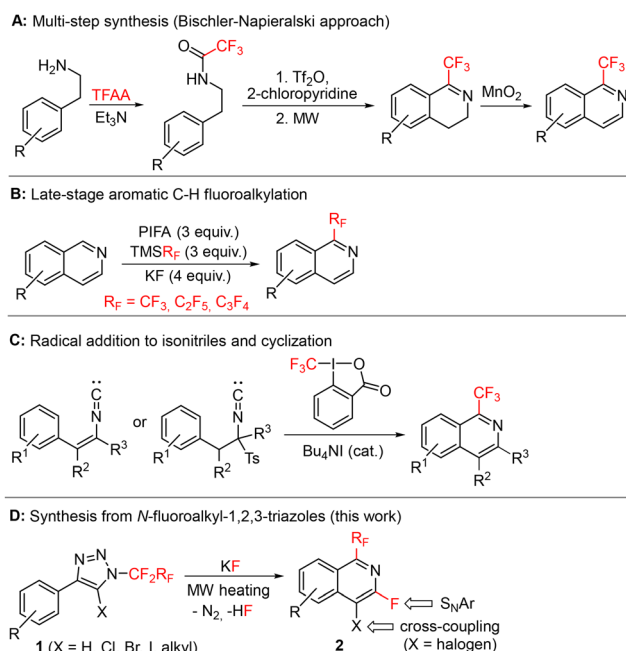
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Fig. 1 Examples of an isoquinoline containing alkaloid (papaverine) and synthetic drugs (fasudil and ripasudil).



Fig. 2 Examples of bioactive 1-trifluoromethylisoquinolines.^{10–12}

Scheme 1 Selected literature syntheses of 1-fluoroalkylated isoquinolines (A–C) and our new approach from triazoles (D).

quinolines 2 based on thermal decomposition of *N*-fluoroalkyl-1,2,3-triazoles 1, formal 1,3-fluorine shift, and cyclization (Scheme 1D). The presence of fluorine in position 3 and a halogen in position 4 of the isoquinoline ring enabled further modifications by nucleophilic aromatic substitution and/or cross-coupling reactions, respectively. The procedure is applicable also to heteroaryl substituted *N*-fluoroalkyl-1,2,3-triazoles affording heteroarenes with fused fluoroalkylated pyridine rings. The methodology thus allows the expansion of known chemical space to new selectively substituted fluoroalkylated isoquinoline-type structures with potential applications in life sciences.

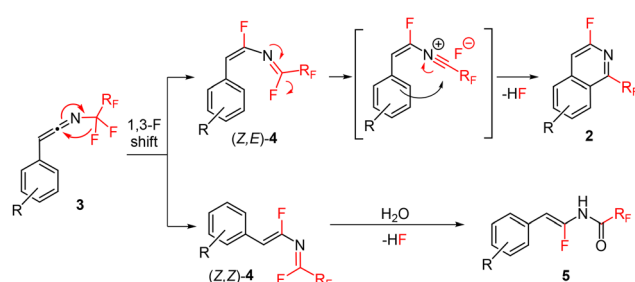
Results and discussion

Recently, we reported thermal rearrangement of *N*-fluoroalkyl-1,2,3-triazoles^{30–33} 1 leading to *N*-fluoroalkylated ketenimines 3.³⁴ We noticed that a prolonged heating of 3 led to new products identified by HRMS and NMR as isoquinolines 2 and

enamides 5. Their formation can be explained by a thermally induced 1,3-fluorine shift of ketenimines 3 to two geometric isomers of azadienes 4. Although four isomers of 4 can be theoretically formed by the fluorine shift, only the formation of two isomers was observed. The isomer (*Z,E*)-4 cyclized to isoquinolines 2 while the isomer (*Z,Z*)-4 only hydrolysed to enamides 5 (Scheme 2) (see the ESI† for three examples of isolated enamides 5). A high-temperature NMR kinetic study revealed the time course of intermediate and product formation (Fig. 3).

A related transformation was briefly reported by Lermontov in 2002.³⁵ Thermal Huisgen cycloaddition of diphenylacetylene with ethyl 3-azido-2,2,3,3-tetrafluoropropanoate afforded isoquinolines and enamides in low yields. The authors wrongly assumed antiaromatic 1*H*-azirines to be the reactive intermediates (Scheme 3A), which we disproved with *ab initio* calculations in our previous study.³⁴ In another report, Molina showed the formation of an isoquinoline by ring closure of an *N*-styryl-substituted ketenimine (Scheme 3B).^{36–38}

In order to develop a general synthesis of 1-fluoroalkylated-2-fluoroisoquinolines 2 from triazoles 1 or ketenimines 3 we studied the effect of additives on the formation of 4. Ideally, the formation of 4 should proceed stereoselectively to the *Z,E*-isomer. Therefore, the influence of additives on the stereoselectivity of the formal 1,3-fluorine shift of 3a at room temperature was studied. While the addition of Et₃N, DBU, or BF₃·OEt₂ did not lead to efficient formation of 4a, the addition of other basic additives or fluoride salts proved beneficial (Table 1). Carbonates induced the stereoselective transformation to the required (*Z,E*)-isomer of 4a with Cs₂CO₃ reacting much faster than K₂CO₃ (entries 2 and 3) and Na₂CO₃ being unreactive (presumably due to its low solubility). However, decomposition of 4a was observed in the basic conditions over time. A similar trend was observed in the case of inorganic fluorides with NaF being unreactive and CsF inducing the formation much faster than KF, but product decomposition and isomerization precluded its use in preparative experiments (entries 4–6; see the ESI† for the isomerization study of 4a with CsF). Therefore, mildly basic KF was used as the additive of choice, accelerating the formal 1,3-fluorine shift of ketenimines 3 and providing a high selectivity to the required isomer of 4 for further cyclization. The origin of the stereoselectivity



Scheme 2 Proposed reaction mechanism of thermal additive-free decomposition of ketenimines 3 to isoquinolines 2 and side-products enamides 5.



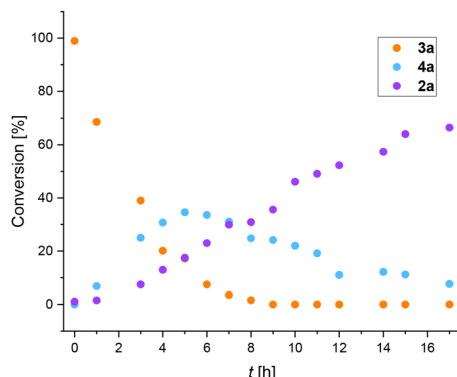
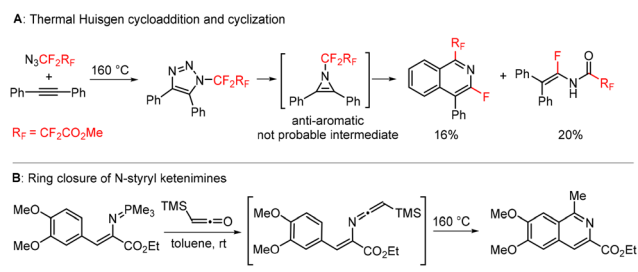
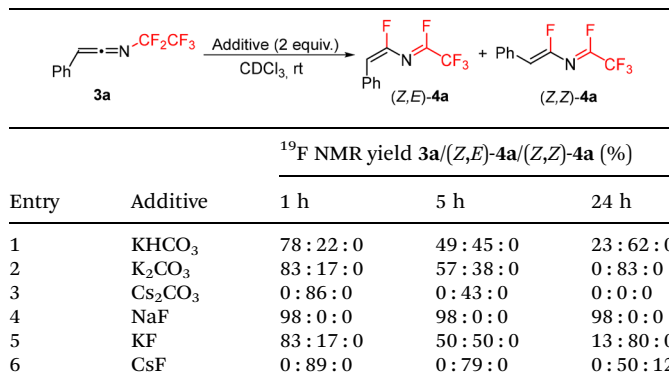


Fig. 3 Conversion of **3a** ($R = H$, $R_F = CF_3$) vs. reaction time for the formation of intermediates **4a** and product **2a** determined by 1H NMR ($C_2D_4Cl_2$; $140^\circ C$).



Scheme 3 Published preparations of isoquinolines by thermal Huisgen cyclization (A) and from *N*-styryl-substituted ketenimines (B).

Table 1 Influence of additives on the formation of isomers of **4a**

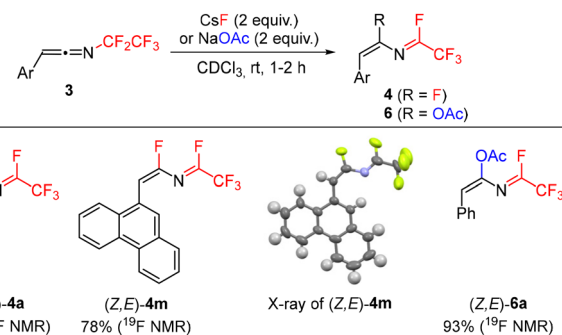


of the thermal or heterogeneous additive-mediated 1,3-fluorine shift is unknown; however, we propose the steric factor to be dominant with fluoride addition to the central sp carbon atom of the ketenimine proceeding *trans* to the large aryl group followed by fluoride elimination from the CF₂ group (Scheme 4).

Difluoroazadiene (*Z,E*)-**4a** was prepared using CsF (Scheme 5). The structure of its derivative **4m** was confirmed by X-ray crystallography. Furthermore, addition of sodium acetate to **3a** efficiently afforded acetate **6a**, confirming that



Scheme 4 Fluoride-mediated formation of azadienes **4a** from ketenimine **3a**.



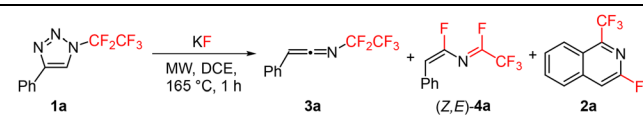
Scheme 5 Characterized azadienes **4** and **6**

indeed a suitable nucleophile can add to the sp carbon of ketenimine **3a**, followed by fluoride elimination. Attempts to use chloride or iodide salts were unsuccessful.

An optimization study revealed that under microwave heating conditions a slight excess of KF afforded the formation of isoquinoline **2a** directly from triazole **1a** without the need to isolate the intermediates ketenimine **3a** or difluoroazadiene **4a** (Table 2). As for the solvent effect, we previously reported that the formation of ketenimines **3** from triazoles **1** works best in DCE but other solvents (chloroform, THF, toluene, cyclohexane, acetone) can also be used.³⁴ In this study we chose DCE as the optimal solvent.

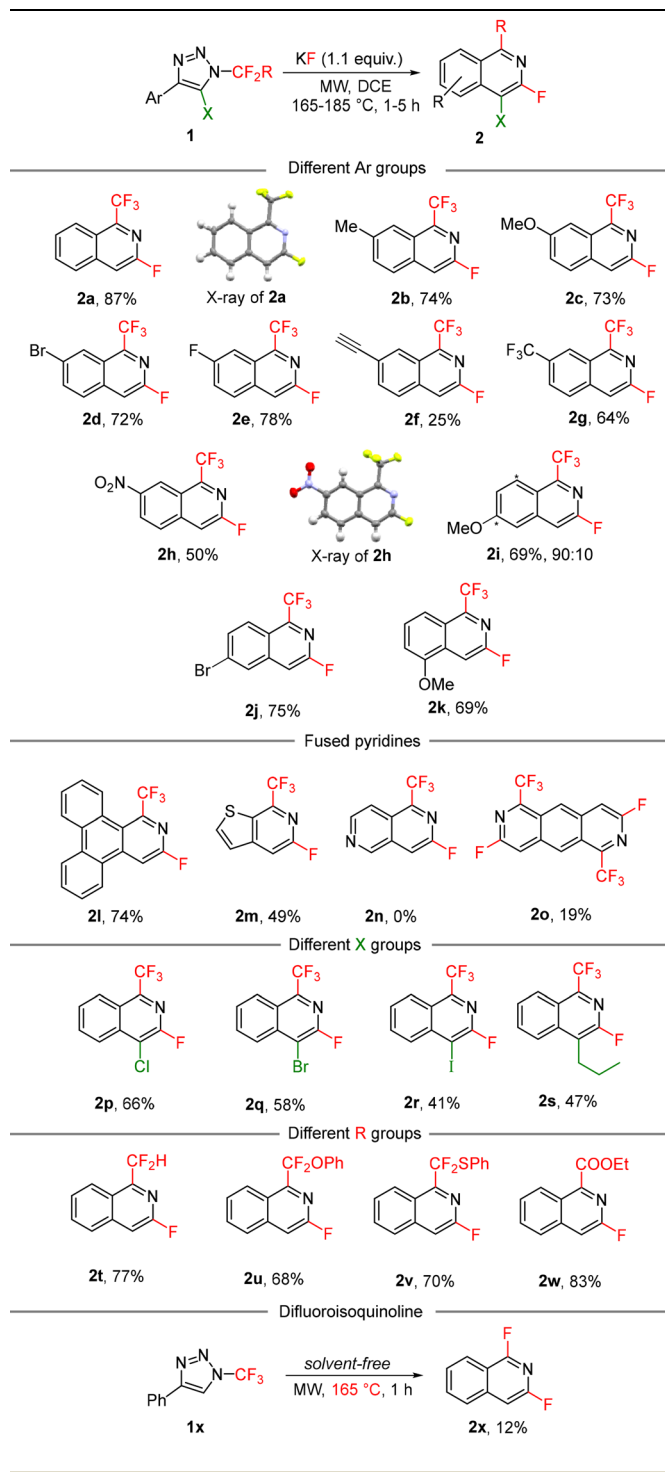
With the optimized set of conditions, we expanded the multistep one-pot process to diversely substituted *N*-fluoroalkylated 1,2,3-triazoles **1** (Table 3). The method tolerated various functional groups on the aryl moiety, including electron-neutral, electron-rich and electron-poor substituents on the phenyl group with slightly decreased yields in the last

Table 2 Optimization of potassium fluoride-accelerated synthesis of isoquinoline 2a from triazole 1a



Entry	KF (equiv.)	^{19}F NMR ratio 3a / $(\text{Z},\text{E})\text{-4a}$ / 2a
1	0.05	70 : 20 : 10
2	0.2	48 : 28 : 24
4	0.7	7 : 5 : 88
5	1.0	3 : 3 : 94
6	1.1	0 : 0 : 99
7	2.0	0 : 0 : 99

Table 3 Substrate scope of KF-mediated synthesis of isoquinolines 2



case. Different substitution positions on the aryl group were also well tolerated with differently substituted isoquinolines being produced from *o*-, *m*- or *p*-substituted aryls. In the case of *m*-substituted aryls, two isomers of the products were formed (**2i**) with good regioselectivity. In another case, the reaction was regiospecific (**2j**). Substrates with large (**1l**) or heteroaromatic (**1m**) groups also underwent the reaction to afford

unique isoquinolines or fused pyridines; however, the pyridyl-substituted triazole (**1n**) was found to be unreactive and only decomposition to a complex mixture of products was observed.

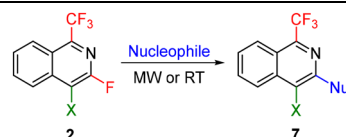
Position 5 of the starting triazole ring can be substituted with a halogen or alkyl group, which introduced these functions into position 4 of the final isoquinoline with various degrees of efficiency. The observed trend can be explained by steric factors where bulky substituents on the ketenimine sp^2 carbon atom hindered the attack of the fluoride ion to form the productive isomer of azadienes **4**.

The methodology was found to display an excellent robustness with regards to the fluoroalkyl substituent in position 1 of the products. Not only the trifluoromethyl group, but also difluoromethyl, substituted difluoromethylene and ethoxycarbonyl substituents can be introduced efficiently. Under solvent free conditions difluoroisoquinoline **2x** was prepared in low yield due to its high volatility and some side reactions.

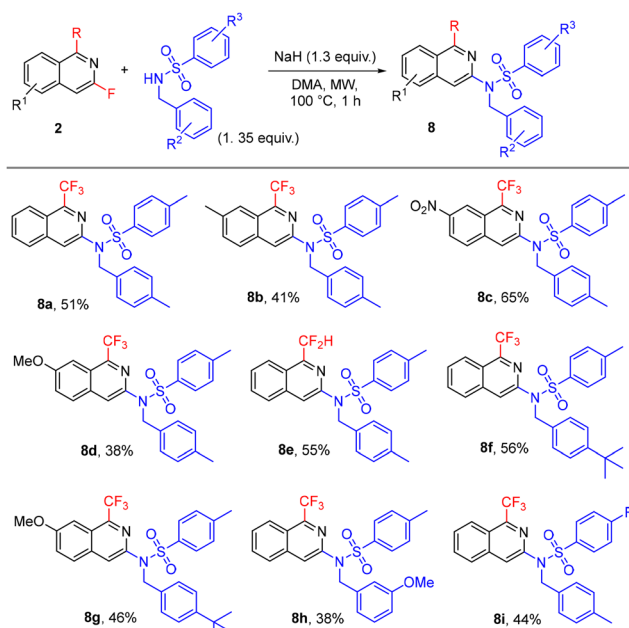
The presence of a fluorine substituent in isoquinolines **2** in the activated position called for the investigation of substitution with various nucleophiles by S_NAr which expanded the diversity of accessible 1-fluoroalkylated isoquinolines. Thus, the fluorine atom of isoquinolines **2** was readily substituted with various oxygen, sulfur, and nitrogen nucleophiles in polar solvents to obtain heteroatom-substituted 1-trifluoromethyl isoquinolines **7** (Table 4).

Furthermore, isoquinolines **2** were used for the preparation of a small library of nine analogues of the TRPM8 antagonist shown in Fig. 2. Compounds **8** were easily accessed by nucleophilic sulfonamidation of **2** (Scheme 6), demonstrating the value of our approach in the synthesis of fluorinated and fluorooxylated isoquinolines and their structurally diverse derivatives in drug development.

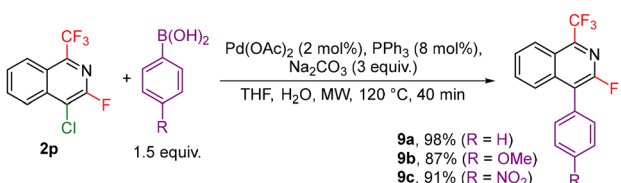
Other investigated follow-up derivatizations of compounds **2** were the cross-coupling reactions. Suzuki–Miyaura coupling of arylboronic acids with chloroisooquinoline **2p** afforded coupling products **9a–c** in high yields (Scheme 7). Heck, Sonogashira and Buchwald–Hartwig reactions of bromoisooqui-

Table 4 S_NAr of isoquinolines **2**

Entry	Nucleophile (equiv.)	X	Solvent	Temp. (°C)	7, yield (%)
1	NaOH (15)	H	H_2O	155	7a , 88
2	EtONa (12)	H	EtOH	80	7b , 96
3	EtONa (12)	Cl	EtOH	155	7c , 99
4	<i>t</i> -BuOK (1.2)	H	<i>t</i> -BuOH	80	7d , 80
5	PhONa (1.5)	H	DMA	80	7e , 89
6	MeSNa (5)	H	DMA	20	7f , 85
7	MeSNa (2)	Ph	DMF	20	7g , 91
8	<i>p</i> -Tol-SNa (1)	H	DMA	80	7h , 91
9	<i>p</i> -Tol-SO ₂ Li (2.5)	H	DMSO	155	7i , 58
10	NH ₂ NH ₂ (20)	H	i-PrOH	100	7j , 95
11	<i>p</i> -Tol-CH ₂ NH ₂ (2)	H	DMSO	155	7k , 42



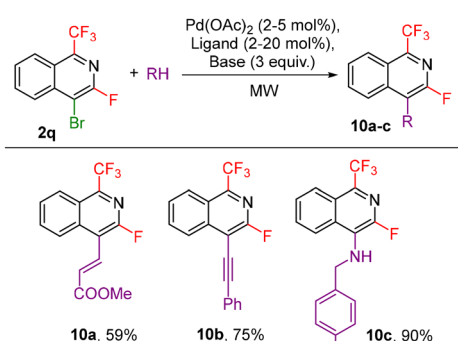
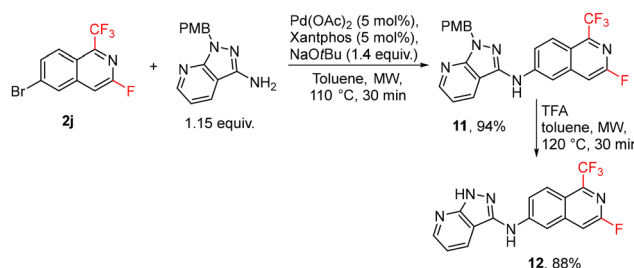
Scheme 6 Analogues of the TRPM8 antagonist.



Scheme 7 Suzuki–Miyaura coupling reactions with 2p.

noline **2q** also worked well giving the coupling products **10a–c** (Scheme 8).

The developed method for the synthesis of fluorinated isoquinolines was used for the preparation of 3-fluoro analogue **12** of the drug candidate valiglurax. The brominated isoquinoline **2j** was used for Pd-catalyzed amination, followed by protecting group removal to give analogue **12** in high yields (Scheme 9).

Scheme 8 Heck, Sonogashira and Buchwald–Hartwig coupling reactions of **2r** (see the ESI† for detailed conditions).Scheme 9 Synthesis of valiglurax analogue **12**.

Conclusions

In conclusion, microwave heating of *N*-fluoroalkyl-1,2,3-triazoles in the presence of potassium fluoride led to a series of events involving triazole ring opening, nitrogen molecule elimination, rearrangement, stereoselective formal 1,3-fluorine shift, and finally cyclization to produce diverse 1-fluoroalkylated 3-fluoroisoquinolines in good yields and with excellent substrate scope. Nucleophilic aromatic substitution of the fluorine atom in position 3 with heteroatom nucleophiles afforded 1-fluoroalkylated 3-substituted isoquinolines. Cross-coupling reactions of halogen atoms in position 4 of the isoquinolines gave derivatives with aryl, alkenyl, alkynyl or alkylamino groups. This synthetic approach to novel selectively fluorinated isoquinolines was applied in the synthesis of analogues of two families of drug candidates.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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