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Polyfluoroalkylated antipyridines in Pd-catalyzed transformations†

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In the direct C–H arylation with arylhalogenides in the presence of Pd(OAc)₂, trifluoromethyl-containing antipyridine reacts very slowly and incompletely owing to the low nucleophilicity of its C4 center. However, it was effective in modifying polyfluoroalkyl-substituted 4-bromo- and 4-iodo antipyridines by the Suzuki and Sonogashira reactions. It was established that using Pd₂(dba)₃ as catalyst and XPhos as phosphine ligand was the optimal catalytic system for the synthesis of 4-aryl- and 4-phenylethynyl-3-polyfluoroalkyl-antipyridines. Moreover, iodo-derivatives as the initial reagents were found to be more advantageous compared to bromo-containing analogs. It was found that 4-phenylethynyl-5-CF₃-antipyridine has a moderate activity against the influenza virus A/Puerto Rico/8/34 (H1N1) and 4-iodo-5-CF₃-antipyridine reveals a weak activity against the vaccine virus (strain Copenhagen) and bovine diarrhea virus (strain VC-1).

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Introduction

The pyrazole core is an acknowledged privileged scaffold in medicinal chemistry.^{1–10} In this regard, pyrazolone derivatives are of particular interest because, based on them, an entire class of analgesic-antipyretics has been developed to be successfully applied in clinical practice for a long time.^{11,12} The predecessor of this pyrazolone family is phenazone (antipyridine) (1) (Fig. 1) that is still part of some combination drugs, *e.g.*, A/B otic drops (ear drops).¹³

Research in recent years has shown that the pyrazolone core is a universal structure¹⁴ to develop substances with various types of biological action including antimicrobial,¹¹ antitubercular,^{15,16} antiviral,¹⁷ anticancer,^{18–21} analgesic,²² anti-inflammatory,²³ antioxidant,²⁴ and anti-diabetic²⁵ activities as well as action on CNS,^{26,27} *etc.*

The several drugs containing pyrazolone scaffold is presently approved for medical use (Fig. 1). These are edaravone (2) with antiradical properties for the treatment of amyotrophic lateral

sclerosis, aminophenazone (3) with antipyretic and anti-inflammatory activity, eltrombopag (4) for the treatment of low blood platelet counts, dichloralphenazone (5) for relieving tension and vascular headaches, metamizole (6) to stop severe pain and fever, sulfamazone (7) – sulfanilamide antibiotic with antipyretic properties, and analgesics propyphenazone (8) and nifenazone (9) (please see, DrugBank).

Note that fluorine-containing pyrazolone derivatives are promising because of their potential use in the pharmaceutical

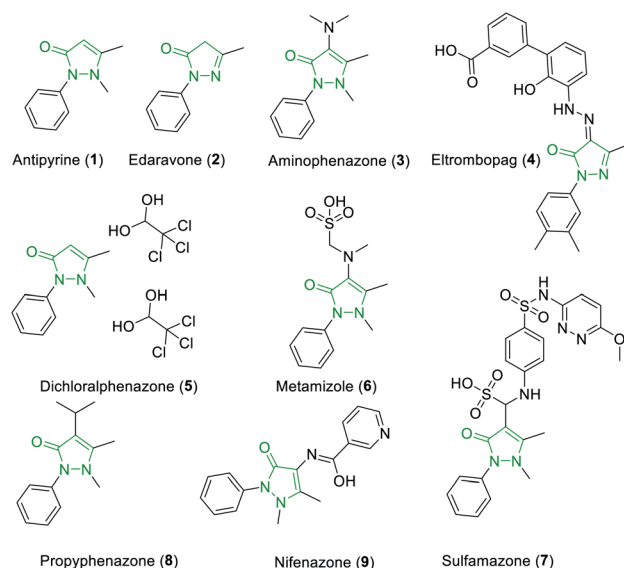


Fig. 1 Drugs based on pyrazolone core.

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industry.^{28–33} For example, the medicines containing tri-fluoromethyl–pyrazole cycle are celecoxib,³⁴ mavacoxib³⁵ (COX-2 selective inhibitors), and razaxaban³⁶ (inhibitor of coagulation blood factor Xa).

Recently, we have developed the methods for synthesis of polyfluoroalkyl-containing antipyrines.³⁷ CF₃-antipyrine (1-methyl-2-phenyl-5-trifluoromethyl-1,2-dihydro-3H-pyrazolon-3-one) was found to show an anti-inflammatory and analgesic activity at the level of diclofenac and metamizole or significantly higher, and its antipyretic effect was higher than that of paracetamol.³⁸ Therefore, the promising challenge is the development of methods for functionalization of polyfluoroalkyl-antipyrines.

The modification of organic compounds *via* the generation of new carbon–carbon bond is known to be one of the most used approaches to generate new molecules. In the mid-seventies of the 20th century, a wide range of C–C bond formation reactions catalyzed by metal complexes was discovered, which made it possible to put these transformations in a row of the most effective and successful tools of organic synthesis. The great research interest in these transformations is caused primarily by their synthetic capabilities, which allow a wide range of organic substrates to be involved in the reactions, as well as by high selectivity in the formation of target products and by the comparatively low requirements to purity of the reagents.^{39,40}

The cross-coupling reactions allow C–C bonds not only to be generated between aryl components but also to involve various hetaryl fragments in the process, which significantly extends their synthetic potential to modify the natural substances and pharmacologically active compounds.^{41,42} Recently, the methods have been enthusiastically investigated for modifying pyrazoles, including polyfluoroalkyl-containing ones, at the position 4. For example, the Suzuki cross-coupling reaction of 3-chloro-4-iodo-1-methyl-5-(trifluoromethyl)-1H-pyrazole with phenylboronic acid in the presence of Pd(PPh₃)₄ led to 3-chloro-1-methyl-4-phenyl-5-(trifluoromethyl)-1H-pyrazole.⁴³ We used an analogous catalyst to modify 4-bromo-1,5-diphenyl-3-(polyfluoroalkyl)-1H-pyrazoles in the conditions of microwave irradiation to obtain 4-(het)aryl-substituted pyrazoles.⁴⁴ The 4-iodo-derivatives of a celecoxib and its analogs were used in the cross-coupling reactions with copper(I) cyanide to give 4-cyano-3-CF₃-pyrazoles, with phenylboronic acid in the presence of Pd(PPh₃)₄ and K₂CO₃ – triaryl-substituted 3-CF₃-pyrazoles, with 4-methoxythiophenol in the presence of the catalytic system of Pd(dba)₂-Xantphos – 4-arylthio-3-CF₃-pyrazoles, and in the conditions of Sonogashira reaction – 4-phenylethynyl-3-CF₃-pyrazoles in.⁴⁵

A great research attention has also been paid to the modification of antipyrine and its analogs in the cross-coupling reactions. A convenient and ligand-free method has been described for the synthesis of a series of 4-aryl-antipyrines based on 4-unsubstituted derivatives and arylhalogenides *via* an activation of C–H bond by palladium(II) acetate in the presence of Ag₂CO₃.⁴⁶ Reactivity of *para*-, *meta*- and *ortho*-substituted aryl-bromides has been investigated in the reaction with antipyrine by Pd(OAc)₂ and K₂CO₃.⁴⁷ A method has been suggested for the

direct introduction of arylthiol and arylselenium residues into antipyrines by di-*p*-tolyldisulfide or diphenyldiselenide using catalyst system AgOTf/AgOAc.⁴⁸ An effective way of carrying out the cross-coupling reaction of 4-iodo-antipyrine with arylboronic acids has been found on Pd/nanoglobular carbon support.⁴⁹ The reactions have been described for Sonogashira cross-coupling of antipyrine with electrophile alkynylation reagent – 1-[(triisopropylsilyl)ethynyl]-1,2-benziodoxol-3(1H)-one (TIPS-EBX).⁵⁰

In this report, we investigated and compared the capabilities of polyfluoroalkyl-containing antipyrines in Pd-catalyzed C–H arylations and Suzuki and Sonogashira cross-coupling reactions, and studied antiviral activity of the synthesized compounds.

Results and discussion

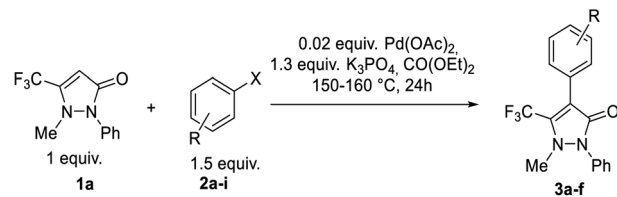
The literature data analysis has shown that antipyrine and its non-fluorinated analogs readily undergo the Pd-catalyzed direct arylation with the different arylhalogenides.^{46,47} However, our numerous efforts to involve 5-trifluoromethyl-antipyrine **1a** into the direct arylation with arylhalogenides were ineffective, despite the use of various palladium catalysts {Pd(OAc)₂, Pd₂(dba)₃}, bases (AcOK, K₂CO₃, Cs₂CO₃, K₃PO₄), ligands (ligand-free, XPhos), solvents (CO(OEt)₂, EtOH–H₂O, DMA, toluene, 1,4-dioxane), temperature modes (100–160 °C) and ratios of reagents in these reactions. The extended experiments are presented in Table S1 (please see, ESI†).

The attempts to introduce phenyl iodide **2a** and 3-cyanophenylbromide **2b** in the reaction with CF₃-antipyrine **1a** (entry 1–4, Table S1†) may be recognized as practically unsuccessful, since the content of the expected 4-aryl-antipyrines **3a**, **b** in accordance with GLC-mass spectrometry was only 5 and 12%, respectively. The conversion of CF₃-antipyrine **1a** in the reaction with 4-cyanophenyl halides **2b**, **c** and with 4-nitrophenyl iodide **2e** into the corresponding products **3c–d** was no more than 21% (entries 5–23, Table S1†). However, the use of aryl bromides **2f**, **h**, **i** in the reaction with antipyrine **1a** led to the higher content of the target products **3d–f** in a reaction mixture in the range of 32–42% (entries 24, 26 and 27, Table S1†). We were successful in isolating the substances **3d–f** in a pure form and the compound **3c** in a mixture with the initial antipyrine **1a** (the **3c** : **1a** ratio was 83 : 17) (Scheme 1).

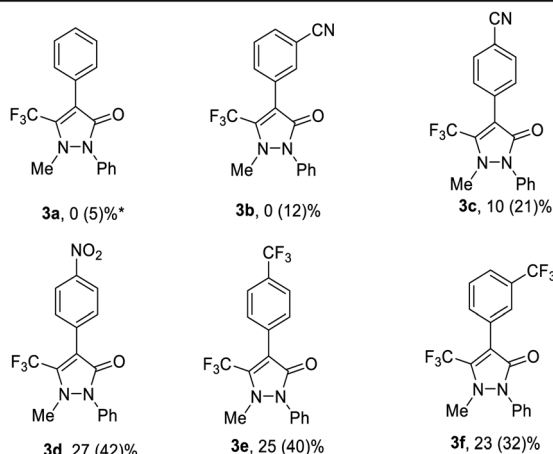
The analysis of all the reaction conditions showed that the catalyst Pd(OAc)₂ was more effective than Pd₂(dba)₃ in the direct arylation, and using the base K₃PO₄ was more successful. Among all the solvents, the definitely best one was eco-friendly CO(OEt)₂ (ref. 51–54) at 150–160 °C. In addition, all transformations did not require a phosphonium ligand to be used. Using aryl bromides was found to be more effective than aryl iodide.

Apparently, in the reactions with electrophile reagents compared to non-fluorinated analogs, the reduced reactivity of 5-trifluoromethyl-antipyrine **1a** was assumed to be caused by the deactivation of carbon C4 due to the influence of the neighboring electron-withdrawing fluorinated substituent. To explain this assumption, we evaluated the Fukui dual





2: R = H, X = I (a); R = 3-CN, X = Br (b); R = 4-CN, X = I (c), Br (d);
R = 4-NO₂, X = I (e), Br (f); R = 4-CF₃, X = I (g), Br (h); R = 3-CF₃, X = Br (i).



*preparative yield (conversion according to GC/MS)

Scheme 1 Direct C–H arylation of antipyrine 1a.

descriptors for centers C4 of antipyrine (**Ant**) and CF₃-antipyrine **1a** (Fig. 2) using quantum-chemical calculations.

According to previous report,⁵⁵ the dual descriptor is able to unambiguously expose truly nucleophilic and electrophilic regions on a molecule. The reaction center C4 of CF₃-antipyrine **1a** was found to be characterized with more positive values of the dual descriptor of Fukui function Δf , which points to its higher electrophilicity in comparison with the same center of **Ant**.⁵⁶ It explains the weak reactivity of compound **1a** in the reactions of electrophile arylation comparing to the non-fluorinated antipyrine in analogous transformations.^{46,47}

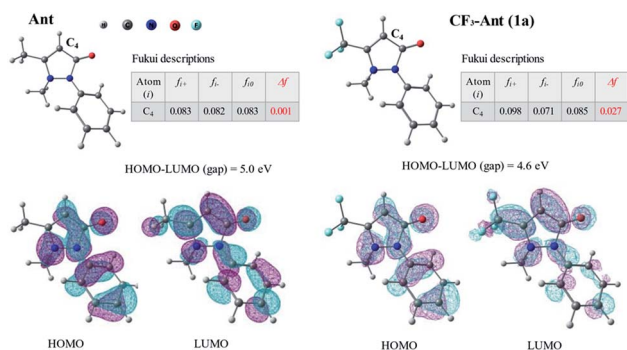


Fig. 2 The dual descriptor of Fukui functions for antipyrine (**Ant**) and CF₃-antipyrine **1a**, HOMO–LUMO gap and visualization.

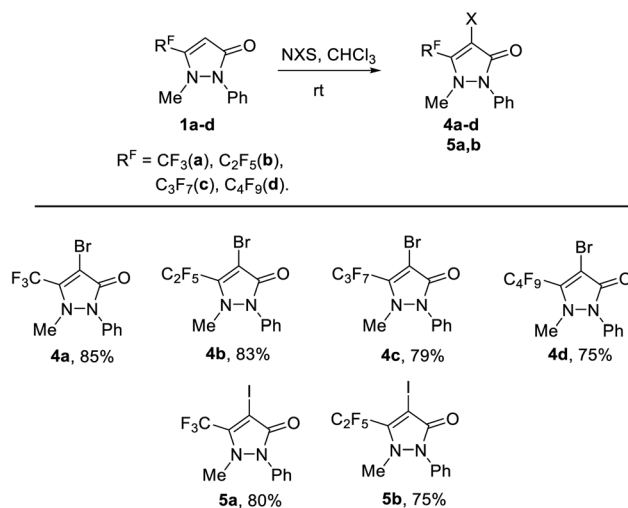
Then, we investigated the possibility of modifying polyfluoroalkyl-antipyrines **1** at the center C4 in the Pd-catalyzed Suzuki and Sonogashira cross-coupling reactions. At first, 4-bromo-5-polyfluoroalkyl-antipyrines **4a–d** and 4-iodo-5-polyfluoroalkyl-antipyrines **5a, b** were synthesized by the treatment of the initial heterocycles **1a–d** with *N*-bromo- or *N*-iodo-succinimide (NXS) (Scheme 2).

The reactions were running readily at room temperature with the good yields. The structure of the obtained halogen-containing antipyrines **4a–d** and **5a, b** was confirmed by IR and NMR spectroscopy and the elemental analysis. The halogenation reaction is confirmed by the absence of the singlet signal of =CH proton at $\delta \sim 5$ –6 ppm in ¹H NMR spectra of the compounds **4a–d** and **5a, b**.

Further, we studied the Suzuki reaction of halogen-substituted antipyrines **4a–d** and **5a, b**. To search for optimal conditions, we chose the synthesis of 4-phenyl-5-trifluoromethyl-antipyrine **3a** via cross-coupling reaction of 1 equiv. of 4-bromo-5-trifluoromethyl-antipyrine **4a** with 1.2 equiv. of phenylboronic acid in the presence of 2.5 equiv. of base K₂CO₃, varying the solvent, palladium catalyst, and phosphine ligand. The reactions were performed at 100 °C in the closed vials under inert gas. A ratio of the reaction products was determined by GLC-mass spectrometry. The data on selecting the reaction conditions are given in Table S2.†

Initially, a mixture of THF–H₂O (3 : 4) was applied as solvent, and Pd(PPh₃)₄ as catalyst/ligand system. There was a partial conversion of the initial bromo-antipyrine **4a** under these conditions (entry 1, Table S2†). The replacement of THF by aqueous ethanol led to full conversion of the initial heterocycle **4a**, but the reaction was accompanied by debromination of bromo-antipyrine **4a** into the parent antipyrine **1a** and the by-products formation (entry 2, Table S2†).

With aqueous ethanol chosen as a solvent, we investigated the possibility to apply Pd₂(dba)₃ as a catalyst in combination with various phosphine ligands: P(*o*-Tol)₃, P(*p*-Tol)₃, XPhos – 2-



Scheme 2 Synthesis of 4-bromo-antipyrines **4a–d** and 4-iodo-antipyrines **5a, b**.

dicyclohexylphosphino-2',4',6'-triisopropylbiphenyl, Xantphos – 4,5-bis(diphenylphosphino)-9,9-dimethylxanthene and BINAP – (±)-2,2'-bis(diphenylphosphino)-1,1'-binaphthalene, [1,1'-Binaphthalene]-2,2'-diylbis[diphenylphosphine]. It was found that in all the cases the reaction proceeded with almost complete conversion of the starting bromo-antipyryne **4a** (entries 3–7, Table S2†).

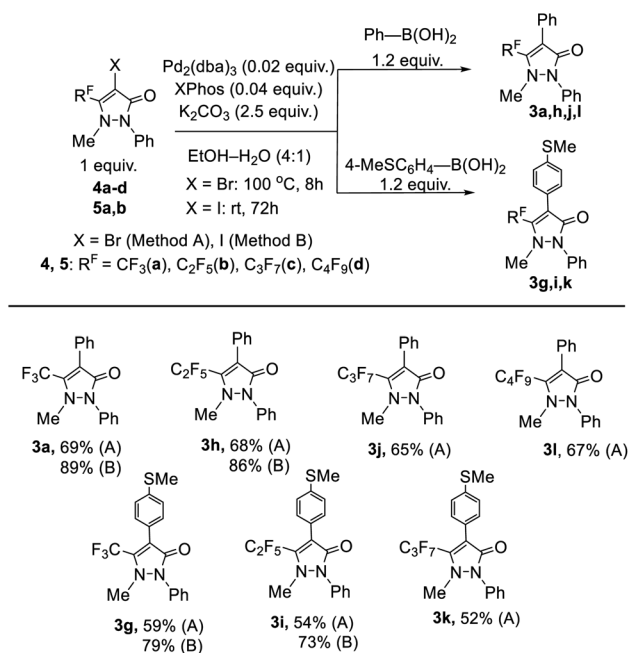
However, the catalyst system $\text{Pd}_2(\text{dba})_3/\text{XPhos}$ (entry 5, Table S2†) was found to be the most effective. In these conditions the content of targeted compound **4a** reached 86%, and the contents of debromination product and impurities were 6% and 8%, respectively. Note that the reaction in the same catalyst system under the microwave irradiation (entry 8, Table S2†) led to the formation of a large number of by-products, and as a result the product **3a** yield decreased to 62%.

It is known that (het)aryl-containing iodides should often enter into cross-coupling reactions easier.⁵⁷ Thus, we carried out a series of reactions of 4-iodo-5-trifluoromethyl-antipyrynes **5a** with phenylboronic acid to give 4-phenyl-5-trifluoromethyl-antipyryne **3a** using similar catalytic systems as those used in the reactions of bromo-substituted analog **4a**. The performed experiments showed, regardless of the use of various phosphonium ligands and Pd catalysts, the main process in these reactions at high temperature was the deiodation of 4-iodo-antipyryne **5a** into antipyryne **1a**. The yield of the desired product **3a** was reduced significantly (entries 1–4, Table S3†). The replacement of base K_2CO_3 by Cs_2CO_3 did not have influence on the reaction (entry 5, Table S3†). However, the reaction under the optimal conditions for transformations of bromo-antipyryne **4a** (1.2 equiv. of phenylboronic acid in the presence of 2.5 equiv. of base K_2CO_3 , 0.02 equiv. of $\text{Pd}_2(\text{dba})_3$ and 0.04 equiv. of XPhos), carried out at room temperature, allowed us to realize 100% conversion of the initial iodo-antipyryne **5a** into 4-phenyl-5-trifluoromethyl-antipyryne **3a** (entry 6, Table S3†). The yield of compound **3a** was 89% after purification. Therefore, these conditions may be considered as the most effective, although the reaction at room temperature proceeded for longer time but with almost quantitative yield. The comparative reaction with 0.05 equiv. of $\text{Pd}(\text{PPh}_3)_4$ was found to be ineffective (entry 7, Table S3†).

Using the selected optimal conditions, a series of 4-phenyl-5-polyfluoroalkyl-antipyrynes **3a, h, j, l** was synthesized (Scheme 3). The yields of heterocycles **3a, h** obtained from iodo-antipyrynes **5a, b** were slightly higher than in the case of bromo-substituted analogs **4a, b**, because the reaction at room temperature allowed dehalogenation of the initial compounds **5a, b** to be avoided. Note that bromo-antipyryne **4a** practically did not react at room temperature.

In addition, the optimal conditions were successfully applied to the reactions of halogen antipyrynes **4a–d** and **5a, b** with [4-(methylthio)phenyl]boronic acid to generate new 4-[4-(methylthio)phenyl]-5-polyfluoroalkyl-antipyrynes **3g, i, k** (Scheme 3).

The structure of the obtained 4-aryl-5-polyfluoroalkyl-antipyrynes **3a, d–f, g–l** was confirmed by the elemental analysis, IR and NMR spectroscopy. The IR spectra of compounds **3a, d–f, g–l** had a similar character. For example, the absorption



Scheme 3 Synthesis of 4-aryl-5-polyfluoroalkyl-antipyrynes **3a, g–l**.

bands of the carbonyl groups of compounds **3a, d–f, g–l** were observed at ν_{CO} 1660–1680 cm^{-1} . The ^1H NMR spectra of antipyrynes **3a, d–f, g–l** were characterized with the presence of methyl and additional aryl protons.

We also performed the X-ray diffraction analysis (XRD) for the compound **3e** to determine its exact structure (Fig. 3).

Then, we studied the Sonogashira cross-coupling reactions of 4-halogen-antipyrynes **4a–d, 5a, b** with phenylacetylene. We also began the investigation of these reactions with the search for the optimal conditions, varying the phosphonium ligands and palladium catalyst.

In contrast to the Suzuki reactions, the Sonogashira cross-coupling reactions are commonly carried out in acetonitrile in the presence of copper(i) iodide as co-catalyst and DIPEA as a base. Therefore, we performed a series of the reactions of 1 equiv. of 4-bromo-5-trifluoromethyl-antipyryne **4a** with 1.5 equiv. of phenylacetylene in the presence of 2.0 equiv. of DIPEA and 0.1 equiv. of co-catalyst CuI , using various palladium catalyst and phosphine ligands in acetonitrile at 80 °C in the closed vials

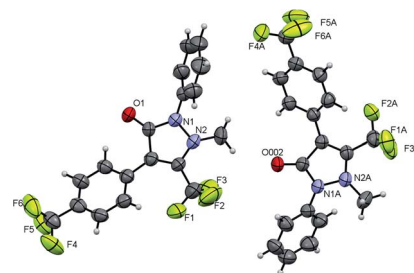


Fig. 3 The structure of 1-methyl-2-phenyl-5-(trifluoromethyl)-4-[4-(trifluoromethyl)phenyl]-1,2-dihydro-3H-pyrazol-3-one **3e** according to XRD (CCDC 2110385†).

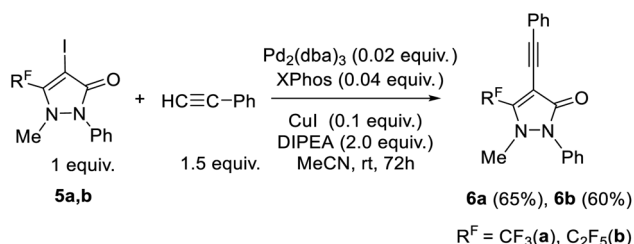


under argon for 12 h (entries 1, 3–5, Table S4†). However, these conditions did not lead to satisfactory results owing to incomplete conversion of the starting bromo-antipyridine **4a** and the formation of a large number of by-products (mainly phenylacetylene crosslinking product – 1,4-diphenyl-1,3-butadiyne). According to GLC-mass spectrometry data, the content of the targeted 1-methyl-2-phenyl-4-(phenylethynyl)-5-(trifluoromethyl)-1,2-dihydro-3H-pyrazol-3-one **6a** in the reactions mixtures was recorded in the range of 8 to 30%. The reaction with 0.05 equiv. of $\text{Pd}(\text{PPh}_3)_4$ (entry 2) or with 0.02 equiv. of $\text{Pd}_2(\text{dba})_3$ and 0.04 equiv. of XPhos (entry 6) at 100 °C for 6 h resulted in a slight increase of product **6a** to 35 and 36%, respectively. The yield of the targeted heterocycle **6a** was 26% after purification (entry 6, Table S4†).

Further, we used 4-iodo-5-trifluoromethyl-antipyridine **5a** as the initial reagent in the reaction with phenylacetylene (Table S5†). To perform a series of experiments, the already proven catalytic system $\text{Pd}_2(\text{dba})_3$ -XPhos was chosen. The reaction of antipyridine **5a** with 1.5 equiv. of phenylacetylene was carried out in the closed vials under argon in the presence of 0.02 equiv. of $\text{Pd}_2(\text{dba})_3$, 0.04 equiv. of XPhos, 2.0 equiv. of DIPEA and 0.1 equiv. of CuI in different solvents at heating or at room temperature (Table S5†).

The reaction in toluene at 100 °C was found to result in a mixture of by-products, and the content of the targeted 4-(phenylethynyl)-5-(trifluoromethyl)-antipyridine **6a** was only 2% (entry 1, Table S5†). Using 1,4-dioxane (entry 2) or acetonitrile (entry 3) at the same temperature led to significant deiodation of the initial iodo-antipyridine **5a**, but the content of the desired heterocycle **6a** was slightly higher (25%) in acetonitrile than in 1,4-dioxane (13%). At a lower reaction temperature (50 °C) in acetonitrile the content of targeted compound **6a** increased up to 35% (entry 4, Table S5†).

The most effective conditions for the Sonogashira reaction of iodo-antipyridine **5a** with phenylacetylene was found to hold the reaction mass in acetonitrile at room temperature for 72 h (entry 6, Table S5†). Here, the full conversion of the initial iodo-antipyridine **5a** occurred but this transformation was also accompanied by a side process: self-condensation of phenylacetylene to form 1,4-diphenyl-1,3-butadiyne. The content of the desired product **6a** in the reaction mixture was 75% and the yield after purification was 60%. The found optimal conditions were used to generate pentafluoroethyl substituted analog **6b** (Scheme 4).



Scheme 4 Synthesis of 4-(phenylethynyl)-5-(polyfluoroalkyl)-antipyridines **6a**, **b**.

The structure of 4-(phenylethynyl)-5-(polyfluoroalkyl)-antipyridines **6a**, **b** was confirmed by the elemental analysis, IR and NMR spectroscopy. The existence of triple bond was affirmed by the presence in the ^{13}C NMR spectra of the chemical shifts at δ 76.4–76.6 and 97.8–98.0 ppm, which are typical of two sp -hybridized carbon atoms.^{45,58}

Biological testing

Iodoantipyridine has been known to be applied in the medicinal practice to treat and prevent tick-borne encephalitis,^{59,60} influenza and infections caused by nonpoliomyelitis enteroviruses of Cocksackie and ECHO groups.⁶⁰ We evaluated an antiviral activity of the synthesized polyfluoroalkyl-containing iodo-antipyridines **5a**, **b**, and aryl-**3a**, **g** and phenylethynyl-substituted **6a** antipyridines against the influenza virus A/Puerto Rico/8/34 (H1N1) on MDCK cell line (Table 1) using ribavirin as a reference drug according the published procedure.⁶¹

Studying the cytotoxicity of compounds on the MDCK cell line, it was found that compounds **5a**, **6a** have lower cytotoxicity ($\text{CC}_{50} > 815 \mu\text{M}$) compared to the derivatives **5b**, **3a** and **3g**. The elongation of polyfluoroalkyl chain in iodo-antipyridines **5a**, **b** led to the increase of cytotoxicity (**5a** $\text{CC}_{50} > 815 \mu\text{M}$ vs. **5b** CC_{50} 378 μM). The introduction of phenyl substituent resulted in the increased cytotoxic properties of compound **3a** (CC_{50} 789 μM) in comparison with the methylthiophenyl analog **3g** (CC_{50} 263 μM). Phenylethyl derivative **6a** ($\text{CC}_{50} > 877 \mu\text{M}$) showed the lowest cytotoxicity.

The virus-inhibiting activity of iodo-antipyridines **5a**, **b** and phenyl-antipyridines **3a** was low ($\text{IC}_{50} > 314$ –177 μM), therefore their selectivity index did not exceed 5. The activity of methylthiophenyl-antipyridine **3g** increased up to IC_{50} 91 μM , but it has a low SI = 3 owing to the raised cytotoxicity. The best ratio of cytotoxic and inhibitory properties was shown by phenylethynyl antipyridine **6a** with SI = 15.

The evaluation of antiviral activity of CF_3 -iodo-antipyridine **5a** was carried out *in vitro* against the vaccine virus (VV, strain Copenhagen), herpes simplex virus type 1 (HSV-1, strain VR-3) and bovine diarrhea virus (BDV) (strain VC-1), a surrogate of the hepatitis C virus on the cells of the calf coronary vessels (CCV). It was found that heterocycle **5a** was inactive against

Table 1 Antiviral activity of compounds **5a**, **b**, **3a**, **g**, **6a** against influenza virus A/Puerto Rico/8/34 (H1N1) on the MDCK cell line^a

No.	Compound	CC_{50} , μM	IC_{50} , μM	SI
1	5a	>815	177 ± 21	5
2	5b	378 ± 26	205 ± 19	2
3	3a	789 ± 57	>314	3
4	3g	263 ± 19	>91	3
5	6a	>877	58 ± 8	15
6	Ribavirin	>2130	36 ± 5	59

^a CC_{50} – 50% cytotoxic concentration, the concentration resulting in death of 50% of cells in culture; IC_{50} – 50% inhibiting concentration, the concentration resulting in a 50% decrease of virus production as compared to the control; SI – selectivity index ($\text{CC}_{50}/\text{IC}_{50}$).



Table 2 Cytotoxic and antiviral activity of compound **5a** on the Vero cells against VV, HSV-1, and on the CCV cells against BDV^a

CC ₅₀ , μ M	IC ₅₀ , μ M		IC ₅₀ , μ M		CC ₅₀ , μ M	IC ₅₀ , μ M	
Vero	VV	SI _{BOB}	HSV-1	SI _{HSV-1}	CCV	BDV	SI _{VDC}
298 $\pm$ 50	103 $\pm$ 10	2.9	Inactive	—	298 $\pm$ 50	46.88 $\pm$ 2	6.4

^a CC₅₀ – 50% cytotoxic concentration, the concentration resulting in death of 50% of cells in culture; IC₅₀ – 50% inhibiting concentration, the concentration resulting in a 50% decrease of the virus production as compared to control; SI – selectivity index.

HSV-1, but it showed a weak activity against strains of SVV and VDC with SI 2.9 and 6.4, respectively (Table 2).

Conclusions

In summary, we found a weak reactivity of CF₃-antipyrine in Pd-catalyzed reactions of the direct arylation owing to a decrease in the nucleophilicity of its C4 site compared to antipyrine. For the purpose of modification of polyfluoroalkyl-substituted antipyrines at the C4 center, the Suzuki and Sonogashira reactions *via* the preliminary synthesis of 4-bromo- and 4-iodo-derivatives were found to be effective. Using Pd₂(dba)₃ as catalyst and XPhos as phosphine ligand were found to be the optimal conditions for the preparation of 4-aryl- and 4-phenylethynyl-3-R^F-antipyrines. In addition, the application of iodo-R^F-antipyrines as the initial reagents was more advantageous compared to bromo-containing analogs, since they were easily and more efficiently transformed into the target products at room temperatures. The synthesis under such mild conditions allowed us to avoid dehalogenation of the starting halogen-R^F-antipyrines. Note that this process was the main side reaction in the studied transformations, which reduces significantly the yield of the target products.

The investigation of antiviral activity of the new derivatives of polyfluoroalkyl-containing antipyrines showed a moderate activity of 4-phenylethynyl-5-CF₃-antipyrine against the influenza virus A/Puerto Rico/8/34 (H1N1) and a weak activity of iodo-5-CF₃-antipyrines against vaccine virus (strain Copenhagen) and bovine diarrhea virus (strain VC-1), a surrogate of the hepatitis C virus.

Author contributions

E. V. Shchegolkov: conceptualization, methodology, validation, investigation, data curation, writing – original draft, writing – review & editing, visualization. Y. V. Burgart: conceptualization, methodology, validation, writing – original draft, writing – review & editing, visualization. D. A. Matsneva: investigation, formal analysis. S. S. Borisevich: formal analysis, visualization. R. A. Kadyrova: investigation, formal analysis. I. R. Orshanskaya: investigation, formal analysis. V. V. Zarubaev: data curation, validation, resources, writing – original draft. V. I. Saloutin: conceptualization, resources, writing – review & editing, funding acquisition.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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