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Palladium-catalysed difluoroolefination of benzyl tosylates toward the synthesis of gem-difluoro-2 trifluromethyl styrene derivatives†

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We have presented an efficient method to access gem-difluoro-2-trifluromethyl styrene derivatives via palladium catalysis. This method features mild reaction conditions, broad substrate scope and good product yields. Moreover, gram–scale reactions demonstrated the robustness and potential of this method. Control experiments revealed that the $-CF_3$ group was essential to the success of this transformation. Finally, the practicality of this method was successfully proven by three synthetic applications.

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Introduction

Fluorinated compounds have found wide applications in various fields due to their unique properties.¹ Among them, gem-difluorostyrenes have been frequently used in the design of potential enzyme inhibitors.² Introducing α –CF₃ group into gem-difluoroolefins could not only retain its high electrophilicity towards many nucleophiles at the terminal carbon, but also increase the biological activity of the molecules (Fig. 1a).³

Compared with considerable efforts devoted to the development of gem-difluorostyrenes, $4,5$ the synthesis of gemdifluoro-2-trifluromethyl styrenes is less investigated. The most common method is Wittig gem-difluoroolefination of trifluoroacetophenone (Fig. 1b(1)).⁶ Another pathway is a multistep strategy involving nucleophilic addition of an aryl metallic reagent to chloropentafluoroacetone, S_N2 type substitution of chloride anions and dechlorination with Zn $(Fig. 1b(2))$. However, stochiometric phosphine oxide as a by-product, utilization of organometallic reagents and multistep operation have greatly restricted substrate scope and applications of the methods above. Therefore, it is of great significance to develop a complementary method for the synthesis of gem-difluoro-2trifluromethyl styrenes.

On the other hand, transition-metal catalysis plays an irreplaceable role in modern organic synthesis.⁸ We hypothesize that a method including two elementary reactions to access

gem-difluoro-2-trifluromethyl styrenes from trifluoromethylsubstituted benzyl tosylate by transition metal catalysis could be developed (Fig. 1c). From the perspective of elementary reactions, the oxidative addition of palladium catalyst into Csp³-O bond⁹ and β -F elimination of palladium complex¹⁰ have been realized in different transformations in the reported work respectively. Therefore, the key to success of this strategy is to find a suitable catalyst system which is compatible with the two elementary reactions above. PAPER
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b) Previous pathways to access gem-difluoro-2-trifluromethyl olefins

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Ar \begin{matrix} CF_3 & \text{Wittig reagent} \\ 0 & \text{Ar} \end{matrix} \qquad Ar \begin{matrix} CF_3 & & & (1) \\ CF_2 & & & \end{matrix}
$$

$$
CF_3COCF_2Cl \xrightarrow{1) \text{ArLi} \atop{2)} SOCl_2} Ar-CCF_2Cl \xrightarrow{Ch \atop{C}} Ar-CCF_2Cl \xrightarrow{Zn} Ar-CCF_3
$$
 (2)

c) Our strategy (this work)

Fig. 1 Importance of gem-difluoro-2-trifluoromethyl olefins and synthetic strategy.

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Results and discussion

To demonstrate our hypothesis, we began the study by evaluating gem-difluoroolefination of trifluoromethyl-substituted benzyl tosylate (1a) via palladium catalysis (Table 1). Compound 1a could be easily synthesized from corresponding aryl carboxylic acid.¹¹ After evaluation of all reaction parameters, reaction conditions which could provide a high yield of 2a was identified. The optimum reaction conditions consisted of $PdI₂$ (5 mol%) with bidentate ligand DPPP (5 mol%) as catalyst, zinc (2.0 equiv.) as reductant, and DMA as solvent at 80 $^{\circ}$ C (entry 1). Using other palladium sources as catalyst resulted in lower yields (entries 2–5). Variation of monodentate and other bidentate phosphine ligands from DPPP led to moderate yields of 2a (entries 6–11), while nitrogen ligands would inhibit the reaction with the majority of 1a unconverted (entries 12–13). Solvents screening revealed that DMA was the best choice for this transformation (entries 14–18). Lastly, reaction temperature investigation suggested that the desired product 2a could be formed in the highest yield at 80 \degree C, although the yield was acceptable at 40 \degree C (entries 19–21).

With the optimized conditions in hand, the substrate scope of this transformation was investigated and the results are summarized in Scheme 1. Initially, substrates with electron-

^a Standard reaction conditions: 1a (0.2 mmol), PdI₂ (5 mol%), DPPP (5 mol%), Zn (2 equiv.), DMA (1.0 mL), 80 \degree C, 12 h. \degree Yields were determined by GC analysis using dodecane as an internal standard. ϵ Isolated yield in the parenthesis.

Scheme 1 Substrate scope. a The temperature is 100 °C and PdI₂ (10 mol%), DPPP (10 mol%) were used.

neutral aryl groups, such as naphthalene ring and benzene ring, were examined. The reactions proceeded smoothly and produced the corresponding gem-difluoroolefins in excellent

yields (2b–2c). Benzene ring bearing a phenyl substituent at the para and ortho position also afforded the corresponding products in good yields (2d–2e). In addition, vinyl group were also well tolerated (2f). Next, compounds with electron-rich substituents were evaluated. The usage of isopropyl-, methyl-, methoxy-substituted substrates led the formation of corresponding products in moderated to good yields (2g–2k). Various electron-poor substrates were also investigated. Compounds containing halogen and trifluoromethyl groups in para position were adapted to the reactions and gave desired products in moderate yields (2l–2o). The configuration of compound 2o was confirmed by X-ray crystallography.¹² Notably, aryl ring bearing bromide at the ortho position had positive hindrance effect on the reaction, resulting in good yields (2p–2q). In addition, both Boc- and sulfonamide-substituted tosylates were tolerated, giving the corresponding products 2r and 2s in 70% and 87% yields respectively. Aryl ring containing methoxy at the meta position (2t) obtained 85% yield, while substrate with an ester group (2u–2v) led to a lower yield. Pleasingly, the reaction was compatible with a range of heterocycles, as demonstrated by the excellent yields obtained for a series of substrates containing pyridine, furan and quinoline ring (2w–2z). Lastly, diaryltrifluoromethyl tosylate could also be converted to corresponding gem-difluorostyrene (2aa) in moderate yield. Unfortunately, nitro (2ab), amino (2ac) and cyano (2ad) groups were found to unsuitable for the reaction. Publis (28-24) Downloade in a strong or published on 10/16/2024 11:34:00 PM. This are not common and the common and

Scheme 2 Gram-scale reactions and control experiments. There are no conflicts to declare.

Scheme 3 Synthetic applications.

The robustness and potential of this method have also been successfully demonstrated by 2a (88% yield) and 2c (72% yield) in gram–scale reactions (Scheme 2a). Next, the effect of $-CF_3$ group was investigated (Scheme 2b). Mono- CF_3 -substituted benzyl tosylate 3 was subjected to the standard conditions, resulting in β -F elimination product 3a (10%), β -H elimination product 3b (36%) and protonated product 3c (28%). To gain more insight into the mechanism, a control experiment was carried out (Scheme 2c). The reaction was carried out in the presence of D_2O (2.0 equiv.) or CD_3OD (2.0 equiv.) under the standard conditions, leading to the formation of protonated product D-4 or D-4'. This result indicated that Pd(0) was oxidatively added into C–OTs bond rather than C–F bond.

To illustrate synthetic utility of this methodology, previously synthesized 2a was subjected to subsequent transformations (Scheme 3). Firstly, the reaction of compound 2a with imidazole in the presence of K_3PO_4 could provide the N-(α -fluorovinyl) azole product 5. 13,15 Likewise, treatment of 2a with sodium phenyl thiolate in THF at room temperature for 12 h resulted in the formation of vinyl sulfide 6 (Z : $E = 7 : 1$) in 75% yield.^{14,15} Lastly, in the presence of palladium catalyst, allylic alkylation between 2a and allyl tert-butyl carbonate 7 could take place, in which the nucleophilic addition of external fluoride onto gemdifluoroalkenes was the initial step.¹⁶

Conclusions

In conclusion, we have developed an efficient pathway to access gem-difluoro-2-trifluromethyl styrene derivatives via palladium catalysis. This transformation features mild reaction conditions, broad functional group tolerance and good yields. Gramscale reactions have demonstrated the robustness and potential of this method, and various synthetic applications have proved the practicality of this strategy.

Conflicts of interest

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