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Continuous flow synthesis of aryl aldehydes by Pd-catalyzed formylation of phenol-derived aryl fluorosulfonates using syngas†

This communication describes the palladium-catalyzed reductive carbonylation of aryl fluorosulfonates $(ArOSO_2F)$ using syngas as an inexpensive and sustainable source of carbon monoxide and hydrogen. The conversion of phenols to aryl fluorosulfonates can be conveniently achieved by employing the inexpensive commodity chemical sulfuryl fluoride (SO_2F_2) and base. The developed continuous flow formylation protocol requires relatively low loadings for palladium acetate (1.25 mol%) and ligand (2.5 mol%). Good to excellent yields of aryl aldehydes were obtained within 45 min for substrates containing electron withdrawing substituents, and 2 h for substrates containing electron donating substituents. The optimal reaction conditions were identified as 120 °C temperature and 20 bar pressure in dimethyl sulfoxide (DMSO) as solvent. DMSO was crucial in suppressing Pd black formation and enhancing reaction rate and selectivity.

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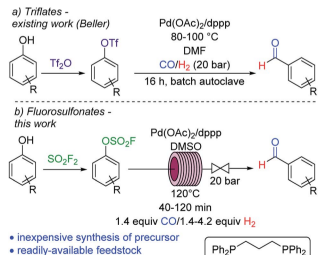
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Aryl aldehydes are ubiquitous intermediates and building blocks in the chemical and pharmaceutical industry. As such, their convenient and cost-efficient synthesis is of high interest. One of the simplest ways of forming aldehydes is through the Pd-catalyzed formylation of Ar-X (where X = I, Br, or OTf) using syngas (CO and H₂). The main limitations in the case of aryl iodides and bromides are the price and availability of the starting materials. In the case of triflates. fluoromethanesulfonic anhydride (Tf2O) is the reagent most commonly employed to prepare triflates from alcohols, but it is relatively expensive, prone to hydrolysis and atom uneconomic.2 Fluorosulfonates (-OSO₂F) have been proposed as an alternative leaving group to triflates because their reactivity is considered largely the same,^{2,3} or placed in-between bromides and chlorides.4 The fluorosulfonate leaving group is an emerging chemical motif⁵ that has been used in many types of chemical transformations including reduction,6 metal catalyzed crosscoupling,3,5,7 deoxyfluorination,8 amination,9 and methoxyearbonylation. 10 Aryl fluorosulfonates can be conveniently and inexpensively prepared by treating readily available phenols

with the commodity chemical sulfuryl fluoride (SO_2F_2) and base

Previously, Beller and co-workers successfully demonstrated the Pd-catalyzed formylation of aryl triflates with syngas as a sustainable and cost effective reagent under batch conditions (Scheme 1a).¹¹ We were interested in the synthesis of aryl aldehydes from aryl fluorosulfonates, which could be derived



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safe process intensification in flowstoichiometric amounts of syngas

Ph₂P PPh₂

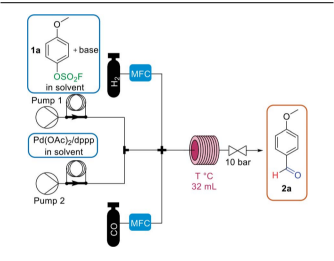
Scheme 1 Pd-catalyzed reductive carbonylation using syngas for (a) triflates (Beller)¹¹ and (b) fluorosulfonates (this work) as leaving groups.

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from their corresponding phenols. Our group previously reported the continuous flow Pd-catalyzed formylation of (hetero) aryl bromides within a continuous flow system.12 Carbonylation reactions benefit from the highly efficient mixing, enhanced mass and heat transfer characteristics, precise residence time, accessibility to high pressure and temperature regimes, and the high operational safety of continuous flow reactors. 13-16 Herein, we describe the development of a continuous flow protocol for the synthesis of aryl aldehydes from their corresponding aryl fluorosulfonates (Scheme 1b).

We commenced our investigation by evaluating the catalytic system palladium(II) acetate (Pd(OAc)2) and 1,3-bis(diphenylphosphino)propane (dppp), which was utilized by Beller and coworkers for the batch formylation of aryl triflates. 11,17 The reaction was optimized within a continuous flow reactor for 4methoxyphenyl sulfurofluoridate (1a) as model substrate. We used a Uniqsis FlowSyn flow system consisting of two HPLC pumps and a heated reactor coil (32 mL). Two sample loops

Table 1 Solvent, base and temperature screening for the continuous flow formylation of aryl fluorosulfonates^a



Entry	Solvent	Temp. [°C]	Conv. 1a ^b [%]	Sel. ^b [%]	Yield ^b 2a [%
1	PhMe	100	7.7	0	0
_				-	
2	PhMe	120	17.1	2.2	0.4
3	THF	100	9.6	6.9	0.7
4	THF	120	17.0	4.9	0.8
5	MeCN	100	20.7	3.0	5.9
6	MeCN	120	42.7	5.9	3.4
7	DMF	100	67.1	19.1	12.8
8	DMF	120	79.1	22.1	17.5
9	DMF^c	120	50.3	22.6	11.4
10	DMF^d	120	36.7	59.0	21.6
11	$\mathrm{DMF}^{d,e}$	120	32.8	50.0	16.4

^a General conditions: feed 1: 0.2 M 4-methoxy sulfurofluoridate (1a), 1.0 equiv. Et₃N and 0.15 equiv. Ph₂O in solvent; feed 2: 5 mol% Pd(OAc)₂ and 10 mol% dppp in solvent. Feed $1/\text{feed } 2/\text{H}_2/\text{CO} = 0.1:0.1:5:5$ mL min $^{-1}$ resulting in a 27 min residence time $(t_{\rm res})$. 10 bar system pressure. b Analyzed by GC-FID. c 13.5 min $t_{\rm res}$. d 1 equiv. of pyridine as base. e Flow rates for feed 1/feed 2/H₂/CO = 0.3 : 0.3 : 1.87 : 1.87 $\mathrm{mL}\ \mathrm{min}^{-1}$, t_{res} 35 min.

were used to deliver the substrate solution (2 mL) and catalyst feed solution (3 mL). The gases, H₂ and CO, were introduced into a four-way mixer by using two mass flow controllers (MFCs). The flow system was pressurized to 10 bar by using a back pressure regulator (BPR). Substrate 1a was mixed with triethylamine (Et₃N) as base and diphenyl ether (Ph₂O) as internal standard for preparation of the substrate feed. Pd(OAc)₂ and dppp were used as the catalyst feed. Initially, we evaluated toluene (PhMe), tetrahydrofuran (THF), acetonitrile (MeCN) and dimethylformamide (DMF) as solvents for the carbonylation reaction (Table 1).

Higher conversion was typically observed when using 120 °C instead of 100 °C. Using PhMe and THF as solvent gave only minimal conversion and trace amount of product (Table 1, entries 1-4). MeCN as solvent gave higher conversions but still very low selectivity towards product (Table 1, entry 5 & 6). Using DMF gave the best preliminary results (entry 7-9). By using pyridine as base, selectivity could be improved, from approximately 20% to 60%, albeit with a drop in conversion (entry 10). We selected to optimize conditions using pyridine since higher selectivity is more desirable. The throughput of the reaction could be almost tripled by using equimolar amounts of gas (1.4) equiv.) and higher liquid flow rates without high loss in yield (entry 11).

During the initial trials with DMF as solvent, we noticed that the obtained results were irreproducible without pre-washing the reactor coil with 20% aqueous nitric acid (HNO₃) at 60 °C in-between experimental runs. Subsequent reaction runs without a wash run in-between resulted in a drop of 4methoxybenzaldehyde (2a) yield and ultimately resulted in a black output solution (Fig. 1). The drop in yield and black output solution indicates a well-known phenomenon of Pd⁰ agglomerating and forming clusters.18 These clusters then irreversibly precipitate as Pd black particles. These Pd black particles coat the reactor wall and can catalyze further Pd black formation.12,19 In order to tackle this issue, we looked for an additive or a method to prevent or slow the rate of Pd black agglomeration.

In 2005, Zierkiewicz and Privalov studied the Pd(OAc)₂/ dimethyl sulfoxide (DMSO) system and identified that DMSO can coordinate to free Pdo, which can help the re-oxidation of Pd⁰ to Pd^{II}.²⁰ When applied to our reaction system using 4-

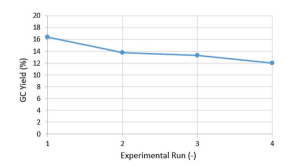


Fig. 1 Drop in 4-methoxybenzaldehyde (2a) GC yield after repeated reaction runs without wash runs in-between. Conditions used are provided in Table 1, entry 11.

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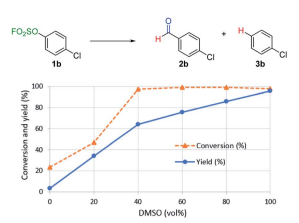
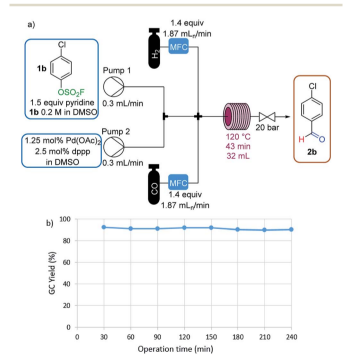


Fig. 2 Influence of DMSO on conversion and yield for Pd-catalyzed formylation of 4-chloro sulfurofluoridate (1b) with syngas measured by GC-FID (Ph₂O as internal standard). For 0–80 vol% DMSO, 0.2 M 4-chloro sulfurofluoridate (1b), 1.0 equiv. pyridine and 0.15 equiv. Ph₂O in DMF/DMSO; feed 2: 5 mol% Pd(OAc)₂ and 10 mol% dppp in DMF/DMSO. 120 °C temperature and 10 bar system pressure. Flow rates for feed 1/feed 2/H₂/CO = 0.3 : 0.3 : 1.87 : 1.87 mL min⁻¹, corresponding to $t_{\rm res}$ 35 min. 100 vol% DMSO experiment was performed using 1.25 mol% Pd(OAc)₂ and 2.5 mol% dppp.

chloro sulfurofluoridate (1b) as substrate, the use of DMSO as co-solvent alleviated the issues associated with Pd black formation (Fig. 2). Furthermore, we could drastically increase the yield of desired product 2b, due to reduced catalyst decomposition. The only side product in all cases was the hydrogenated product 3b, with the loss of the fluorosulfonate



Scheme 2 (a) Continuous flow setup for the Pd-catalyzed formylation of 4-chloro sulfurofluoridate (1b) long run. (b) 4-Chlorobenzaldehyde (2b) GC yield at 30 min intervals over 4 h operation time. 0.15 equiv. Ph_2O was used as an internal standard.

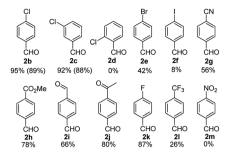


Fig. 3 Substrate scope for aryl fluorosulfonates containing electron withdrawing groups. Yields were determined by GC-FID using Ph_2O as internal standard. Yields in parentheses were isolated yield. Conditions are the same as given in Scheme 2a.

group (-OSO₂F). Increased pressure (20 bar) and base (1.5 equiv.) resulted in higher selectivity to the aldehyde product **2b** (Fig. S1†). The fraction of DMSO solvent could be increased to no more than 80% owing to catalyst solubility.

The reaction conditions in terms of residence time and catalyst loading were then re-optimized for using DMSO as solvent. The catalyst loading was lowered to 1.25 mol% Pd(OAc)₂ and 2.5 mol% dppp to ensure full dissolution of the catalyst system. A further decrease in loading did not provide satisfactory results, with a drop in yield observed (Fig. S2†). Under these conditions 40 min of residence time was sufficient to achieve a high yield of 4-chlorobenzaldehyde (2b) (Fig. S3†). With these new conditions (Scheme 2a), a long run was successfully operated which was stable for 4 hours with no apparent drop in yield, thus demonstrating no or minimal catalyst decomposition (Scheme 2b).

The applicability of the optimized conditions, shown in Scheme 2a, was demonstrated on a range of substrates. In most cases, substrates bearing an electron withdrawing group (EWG) in the *para* position underwent full conversion and the corresponding aryl aldehyde was formed in moderate to excellent yields (Fig. 3). The hydrogenated product was the sole side product observed. However, the reaction of substrates **1e** and **1f** resulted in the formation of a mixture of double formylation and hydrogenated products. The formylation of 4-nitro sulfurofluoridate (1 m) was problematic because of deactivation of the catalyst.

Less reactive substrates, **1k** and **1l**, were only fully converted when modified reaction conditions were applied. In these cases, the liquid flow rates were lowered, which prolonged the residence time to approximately 2 h (Fig. 4a). Moreover, the H₂ flow rate was modified to deliver 4.2 equiv. The same modified conditions were also used for substrates containing electron donating groups (EDG). The formation of the hydrogenated side product was not a significant problem for substrates containing EDG substituents with the aldehyde products obtained in good to excellent yields (Fig. 4b). Gratifyingly, 6-methoxy-2-naphthalaldehyde (**2p**) could be isolated in 82% yield after purification by column chromatography. Substrate **2b** is a potential precursor to naproxen, an important non-steroidal anti-inflammatory active pharmaceutical ingredient.

Fig. 4 (a) Unreactive substrates containing electron withdrawing groups; (b) substrates containing electron donating groups. For 43 min conditions, see Scheme 2a. Conditions: feed 1: 0.2 M fluorosulfonate, 1.5 equiv. pyridine and 0.15 equiv. Ph₂O in DMSO; feed 2: 1.25 mol% Pd(OAc)₂ and 2.5 mol% dppp in DMSO. 120 °C temperature and 10 bar system pressure. Feed 1/feed 2/H₂/CO = 0.08: 0.08: 1.5: 0.5 mL min $^{-1}$, corresponding to a $t_{\rm res}=123$ min.

Substrates bearing a substituent in *ortho* position to the fluorosulfonate group afforded the corresponding aldehyde either in low yield, for **1r** and **1u**, or no yield, for **1d** and **1o**. The drop in yield could be caused by steric hindrance from the *ortho*-substituent. The steric block on the catalytic center will favor the addition of the much smaller hydrogen molecule instead of a larger CO molecule, resulting in the hydrogenated product.

In conclusion, we have described a continuous flow method for the Pd-catalyzed synthesis of valuable aryl aldehyde building blocks from aryl fluorosulfonates and syngas as an inexpensive, atom-economic, and environmentally friendly source of CO and H₂. The continuous flow approach enabled the precise addition of gas by using mass flow controllers. Meta and para substituted aryl fluorosulfonates could be converted to their corresponding aldehydes in good to excellent yields. Catalyst decomposition was successfully avoided by using DMSO as solvent. DMSO coordinates to Pd⁰ and facilitates the re-oxidation to Pd^{II}. Additionally, reaction rates and selectivity could be enhanced by the use of DMSO. Starting materials for the reductive carbonylation could be conveniently derived in excellent yields from readily-available phenols and the commodity chemical sulfuryl fluoride. The developed process is especially appealing for the chemical industry, where reagent cost and availability are important factors in process feasibility.

Conflicts of interest

The authors declare no competing financial interest.

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