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Aryl-CF₃ Bond-Forming Reductive Elimination from Palladium(IV)

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Abstract



This communication describes oxidatively-induced Ar–CF₃ bond-forming reductive elimination from new Pd^{II} complexes of general structure (L~L)Pd^{II}(Ar)(CF₃). The electrophilic fluorinating reagent *N*-fluoro-2,4,6-trimethylpyridinium triflate promotes these reactions in good to excellent yields. The palladium(IV) intermediate (^{*I*}Bu-bpy)Pd^{IV}(CF₃)(F)(OTf)(C₆H₄F) has been isolated, characterized, and demonstrated to undergo high yielding Ar–CF₃ coupling upon thermolysis. This work provides an attractive conceptual framework for the development of Pd^{II/IV}-catalyzed arene trifluoromethylation reactions.

Trifluoromethyl groups feature prominently in a wide variety of medicinal compounds.¹ The substitution of CH_3 for CF_3 can dramatically change the physical properties and biological activity of organic molecules. As a result, tremendous effort has been directed at the introduction of CF_3 groups into organic structures.² This has led to numerous methods for the efficient construction of sp^3 carbon– CF_3 bonds using nucleophilic (CF_3^-), electrophilic (CF_3^+) and radical (CF_3^-)-based trifluoromethylating reagents.²

In contrast, the formation of aryl carbon– CF_3 bonds remains significantly more challenging. ³ This functional group is commonly prepared using the Swarts reaction (which requires high temperatures and reactive SbF_5)⁴ or through the use of stoichiometric quantities of ill-defined and sensitive "Cu– CF_3 " reagents.⁵ Transition metal-catalyzed cross-coupling would provide a highly attractive alternative route to aryl– CF_3 linkages. While sporadic reports have suggested the viability of this approach,⁶ general, efficient, and robust versions of such transformations have been slow to develop. The major challenge in this area is that CF_3 ligands are typically inert towards C–C bond-forming reductive elimination.⁷

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Supporting Information Available: Experimental and crystallographic details and spectroscopic data for new compounds. This material is available free of charge via the internet at http://pubs.acs.org.

Ball et al.

Several groups have made exciting recent progress towards addressing this challenge. For example, Vicic and coworkers have shown that the isolable *N*-heterocyclic carbene (NHC) copper complexes (NHC)Cu–CF₃ react stoichiometrically with aryl iodides to afford Ar–CF₃ products.⁸ Vicic's group has also established that (dippe)Ni^{II}(Ph)(CF₃) undergoes H_sO-promoted Ph–CF₃ coupling in modest (22%) yield.⁹ Finally, Grushin has demonstrated stoichiometric Ph–CF₃ coupling from (Xantphos)Pd^{II} (Ph)(CF₃).¹⁰ This latter work is the only reported example of selective Ar–CF₃ bond-forming reductive elimination from a well-defined transition metal aryl/CF₃ complex. However, this reaction was limited to Ar = Ph and was also extremely sensitive to the nature of the ancillary ligand at Pd^{II}. A variety of P- and N-donor ligands including dppe,¹¹ dppb,¹¹ tmeda,¹¹ and PPh₃ were examined,10^{,12} but only Xantphos was effective at promoting this transformation below 150 °C. Thus, the development of complementary and more general strategies for arene trifluoromethylation from M(Ar)(CF₃) species remains a topic of great current interest.

Our group¹³ and others¹⁴ have shown that Pd^{IV} complexes can participate in reductive elimination reactions that are challenging at other metal centers. Since high oxidation state palladium can be accessed following arene C–H activation¹⁵ or transmetallation¹⁶ processes, Ar–CF₃ coupling through this manifold would provide opportunities for the development of diverse new trifluoromethylation reactions. We report herein the first demonstration of Ar–CF₃ bond-forming reductive elimination from a $Pd^{IV}(Ar)(CF_3)$ complex.

Our studies began with the synthesis of a series of new palladium(II) complexes of general structure (L) $_2Pd^{II}(Ar)(CF_3)$ (**1–3**).^{12b} These were prepared by treating the corresponding Pd^{II} aryl iodides with CsF followed by TMSCF₃ at 23 °C in THF (eq. 1).¹⁰,²¹ The products were obtained in 32–76% yield as yellow solids (see Supporting Information for full details).

$$\begin{array}{c} \begin{array}{c} \begin{array}{c} L \\ \end{array} \\ Pd \end{array} \begin{array}{c} Ar \\ I \end{array} \xrightarrow{1. \text{ CsF, THF, 23 °C}} \\ \hline 2. \text{ TMSCF}_3, \text{ THF, 23 °C} \end{array} \begin{array}{c} \begin{array}{c} L \\ \end{array} \begin{array}{c} Pd \end{array} \begin{array}{c} Ar \\ CF_3 \end{array} \\ \hline (1-3) \end{array} \\ \begin{bmatrix} L-L \\ I \end{array} \begin{array}{c} I \\ \end{array} \end{array}$$

Heating complexes **1–3** at 130 °C for 3 d in nitrobenzene-*d*₅ produced <5% of Ar–CF₃ coupling products (eq. 2). This is consistent with literature reports showing that most Pd^{II}(Ar)(CF₃) complexes are poorly reactive towards Ar–CF₃ bond-forming reductive elimination.^{10,12} We reasoned that $2e^-$ oxidation should yield Pd^{IV} species that might undergo more facile Ar–CF₃ coupling.^{13–15} However, treatment of **1a** with PhI(OAc)₂, NCS,¹¹ or NBS¹¹ (oxidants known to promote the oxidation of Pd^{II} to Pd^{IV})¹³ did not afford 1-fluoro-4-trifluoromethylbenzene; instead, the corresponding acetoxylated or halogenated products were obtained (eq. 2). This result suggests that Ar–X (X = OAc, Cl, Br) bond-forming reductive elimination is significantly faster than Ar–CF₃ coupling in this system.

$$\begin{array}{c} \text{Ar-CF}_{3} \xrightarrow{130 \text{ °C}} (N > \text{Pd} < Ar \\ (<5 \%) \\ \text{(1a)} \\ \text{N} \sim \text{N} = i^{2}\text{Bu-bpy} \\ \text{Ar} = \rho \text{-FC}_{6}\text{H}_{4} \end{array} \xrightarrow{\begin{array}{c} \text{PhI}(\text{OAc})_{2} \\ \text{NCS or NBS} \\ \text{NCS or NBS} \\ \text{H} \\ \text{Ar-CF}_{3} \\ \text{(<5\%)} \\ \text{H} \\ \text{H} \\ \text{Ar-CF}_{3} \\ \text{(<5\%)} \\ \text{H} \\ \text$$

JAm Chem Soc. Author manuscript; available in PMC 2011 March 10.

(2)

(1)

We next examined the use of *N*-fluoro-2,4,6-trimethylpyridinium triflate (NFTPT) to effect Ar–CF₃ coupling from **1a**. This oxidant was selected based on the hypothesis that fluoride and triflate (the X-type ligands introduced to a putative Pd^{IV} intermediate by NFTPT) might undergo slower reductive elimination than CF_3 .^{17,18} Gratifyingly, treatment of **1a** with NFTPT at 80 °C for 3 h in nitrobenzene- d_5 resulted in clean formation of 1-fluoro-4-trifluoromethylbenzene in 85% yield (as determined by ¹⁹F NMR spectroscopy) (Table 1, entry 1). As predicted, <5% of products derived from C–F or C–OTf coupling were observed. NFTPT also promoted Ar–CF₃ bond-formation from the other Pd^{II} complexes **1b–3** (Table 1). These transformations were efficient with strongly electron donating and electron withdrawing arene substituents (entries 2 and 3, respectively). In contrast, reactions of [Cu–CF₃] are often highest yielding with electron deficient aryl coupling partners.⁶ Additionally, unlike Grushin's Ar–CF₃ coupling from Pd^{II} (which is very sensitive to the nature of the ancillary ligands),¹⁰ this oxidatively induced transformation proceeded in modest to excellent yield with diverse N- and P-donor ligands, including ⁷Bu-bpy, tmeda, and dppe.¹¹

In an effort to detect intermediates in the Ar–CF₃ coupling process, we examined the reaction of **1a** with NFTPT at room temperature. In both nitrobenzene and DCE, a single major palladium-containing product (**4**) was observed. This species was isolated from DCE in 53% yield, and the analytically pure yellow solid exhibits four ¹⁹F NMR signals in a 3 : 3 : 1 : 1 ratio. X-ray quality crystals were obtained by vapor diffusion of pentanes into a DCE solution of **4**. The X-ray crystal structure is shown in Figure 1 and confirms that **4** is the octahedral Pd^{IV} species (^{*t*}Bu-bpy)Pd^{IV}(CF₃)(F)(OTf)(C₆H₄F). To our knowledge this is the first isolated example of a Pd^{IV} complex containing a perfluoroalkyl ligand.¹⁹

$$(N_{N} = {}^{CF_{3}}_{(1a)} = {}^{DCE_{3}}_{23 \circ C}, (4) = {}^{N-F_{3}}_{OTf} = {}^{N-CF_{3}}_{Ar-CF_{3}} = {}^{Ar-CF_{3}}_{Ar-CF_{3}} = {}^{Ar-CF_{3}}_{OTf} = {}^{NO_{2}Ph}_{Ar-CF_{3}} = {}^{NO_{$$

(3)

The reactivity of **4** towards Ar–CF₃ bond formation was next investigated. Heating a nitrobenzene- d_5 solution of **4** at 80 °C for 3 h resulted in smooth reductive elimination to form 1-fluoro-4-trifluoromethylbenzene in 77% yield (eq. 3). Similar to the reactions in Table 1, none of the products derived from Ar–F or Ar–OTf coupling were observed by ¹⁹F NMR spectroscopy. This result demonstrates for the first time that Ar–CF₃ coupling can be kinetically accessible from mon- σ -aryl Pd^{IV} complexes; as such, it provides an attractive conceptual framework for the development of Pd^{II/IV}-catalyzed arene trifluoromethylation reactions.

In summary, we describe herein a new Ar–CF₃ bond-forming reaction mediated by Pd^{IV} centers. These transformations proceed under mild conditions with diverse nitrogen and phosphorus-based ancillary ligands. Efforts to gain further insights into the mechanism as well as to develop related catalytic transformations are currently underway in our laboratory and will be reported in due course.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References

- 1. Kirk KL. Org. Process Res. Dev 2008;12:305.
- (a) Ma J-A, Cahard D. Chem. Rev 2004;104:6119. [PubMed: 15584697] (b) Ma J-A, Cahard D. Chem. Rev 2008;108 PR1. (c) Prakash GKS, Chacko S. Curr. Opin. Drug Disc Dev 2008;11:793. (d) Shibata N, Mizuta S, Kawai H. Tetrahedron: Asymmetry 2008;19:2633.
- 3. Grushin VV. Acc. Chem. Res 2010;43:160. [PubMed: 19788304]
- 4. Swarts F. Bull. Acad. R. Belg 1892;24:309.
- For select examples, see:(a) Kobayashi Y, Kumadaki I. Tetrahedron Lett 1969;10:4095. (b) Konderatenko NV, Vechirko EP, Yagupolskii LM. Synthesis 1980:932. (c) Matsui K, Tobita E, Ando M, Kondo K. Chem. Lett 1981:1719. (d) Suzuki H, Yoshida Y, Osuka A. Chem. Lett 1982:135. (e) Burton DJ, Wiemers DM. J. Am. Chem. Soc 1985;107:5014. (f) Urata H, Fuchikami T. Tetrahedron Lett 1991;32:91.
- Cu-catalyzed coupling between of ArI and Et₃SiCF₃:Oishi M, Kondo H, Amii H. Chem. Commun 2009:1909.Pd-catalyzed coupling between ArI and [Zn-CF₃]:Kitazume T, Ishikawa N. Chem. Lett 1982:137.
- 7. For reviews on [M]–CF₃ and [M]-R_f (R_f = perfluoroalkyl) complexes, see:(a) Hughes RP. Adv. Organomet. Chem 1990;31:183. (b) Morrison JA. Adv. Organomet. Chem 1993;35:211.
- (a) Dubinina GG, Furutachi H, Vicic DA. J. Am. Chem. Soc 2008;130:8600. [PubMed: 18543912] (b) Dubinina GG, Ogikubo J, Vicic DA. Organometallics 2008;27:6233.
- 9. Dubinina GG, Brennessel WW, Miller JL, Vicic DA. Organometallics 2008;27:3933.
- 10. Grushin VV, Marshall WJ. J. Am. Chem. Soc 2006;128:12644. [PubMed: 17002347]
- 11. Xantphos = 4.5-bis(diphenylphosphino)-9,9-dimethylxanthene; ^tBu-bpy = 4,4'-di-*tert*-butyl-2,2'-bipyridyl; tmeda = *N*,*N*'-tetramethylethylene diamine; dppe = 1,2-bis(diphenylphosphino)ethane; dippe = 1,2- bis(diisopropylphosphino)ethane; NCS = *N*-chlorosuccinimide; NBS = *N*-bromosuccinide.
- (a) Culkin DA, Hartwig JF. Organometallics 2004;23:3398. (b) Grushin VV, Marshall WJ. J. Am. Chem. Soc 2006;128:4632. [PubMed: 16594700]
- (a) Dick AR, Kampf JW, Sanford MS. J. Am. Chem. Soc 2005;127:12790. [PubMed: 16159259] (b) Whitfield SR, Sanford MS. J. Am. Chem. Soc 2007;129:15142. [PubMed: 18004863] (c) Ball ND, Sanford MS. J. Am. Chem. Soc 2009;131:3796. [PubMed: 19249867] (d) Racowski JA, Dick AR, Sanford MS. J. Am. Chem. Soc 2009;131:10974. [PubMed: 19459631] (e) Arnold PL, Sanford MS, Pearson SM. J. Am. Chem. Soc 2009;131:13912. [PubMed: 19788324]
- 14. (a) Alsters PL, Engel PF, Hogerheide MP, Copijn M, Spek AL, van Koten G. Organometallics 1993;12:1831. (b) Kaspi AW, Yahav-Levi A, Goldberg I, Vigalok A. Inorg. Chem 2008;47:5. [PubMed: 18052157] (c) Furuya T, Ritter T. J. Am. Chem. Soc 2008;130:10060. [PubMed: 18616246] (d) Canty AJ. Dalton Trans 2009:10409. [PubMed: 20023859]
- For recent reviews, see:(b) Lyons TW, Sanford MS. Chem. Rev. [Online early access]. DOI: 10.1021/ cr900184e. Published online: Jan 15, 2010, http://pubs.acs.org/doi/full/10.1021/cr900184e. (b) Muniz K. Angew. Chem., Int. Ed 2009;48:9412.
- 16. Furuya T, Kaiser HM, Ritter T. Angew. Chem., Int. Ed 2008;47:5993.
- 17. For a similar strategy to generate C-N bonds using NFTPT, see:Mei TS, Wang X, Yu JQ. J. Am. Chem. Soc 2009;131:10806. [PubMed: 19606861]
- For use of similar oxidants in Ar-F coupling, see:(a) Hull KL, Anani WQ, Sanford MS. J. Am. Chem. Soc 2006;128:7134. [PubMed: 16734446] (b) Wang X, Mei TS, Yu JQ. J. Am. Chem. Soc 2009;131:7520. [PubMed: 19435367]

 Notably, related structures have been proposed as transient intermediates in the reactions of perfluoroalkyl iodides with Pd^{II}. See:Hughes RP, Overby JS, Williamson A, Lam K-C, Concolino TE, Rheingold AL. Organometallics 2000;19:5190.





Figure 1. ORTEP Diagram of **4**.

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Table 1

Oxidatively Induced Ar–CF $_3$ Coupling from Complexes 1–3

			FF PhNO₂ 80 °C, 3 h	Ar–CF ₃
Entry	Compound	L ₂	Ar	Yield Ar–CF ₃ ^a
1	1 a	'Bu-bpy	p-FC ₆ H ₄	85%
2	1b	^t Bu-bpy	p-CF ₃ C ₆ H ₄	61%
3	1c	^t Bu-bpy	p-CH ₃ OC ₆ H ₄	86%
4	1d	'Bu-bpy	p-PhC ₆ H ₄	83%
5	1e	^t Bu-bpy	p-CH ₃ C ₆ H ₄	85%
6	2	tmeda	p-FC ₆ H ₄	89%
7	3	dppe	Ph	29%

 a Determined by 19 F NMR spectroscopy

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