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Article

Opening a Pandora's Flask on a Prototype Catalytic Direct Arylation Reaction of Pentafluorobenzene: The Ag₂CO₃/Pd(OAc)₂/PPh₃ System

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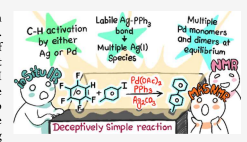
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ABSTRACT: Direct C-H functionalization reactions have opened new avenues in catalysis, removing the need for prefunctionalization of at least one of the substrates. Although C-H functionalization catalyzed by palladium complexes in the presence of a base is generally considered to proceed by the CMD/AMLA-6 mechanism, recent research has shown that silver(I) salts, frequently used as bases, can function as C-H bond activators instead of (or in addition to) palladium(II). In this study, we examine the coupling of pentafluorobenzene 1 to 4-iodotoluene 2a (and its analogues) to form 4-(pentafluorophenyl)toluene 3a catalyzed by palladium(II) acetate with the commonplace PPh3 ligand, silver carbonate as base, and DMF as solvent. By studying the reaction of 1 with Ag₂CO₃/PPh₃ and with isolated silver (triphenylphosphine)



carbonate complexes, we show the formation of C-H activation products containing the $Ag(C_6F_5)(PPh_3)_n$ unit. However, analysis is complicated by the lability of the Ag-PPh₃ bond and the presence of multiple species in the solution. The speciation of palladium(II) is investigated by high-resolution-MAS NMR (chosen for its suitability for suspensions) with a substoichiometric catalyst, demonstrating the formation of an equilibrium mixture of $Pd(Ar)(\kappa^1-OAc)(PPh_3)_2$ and $[Pd(Ar)(\mu-OAc)(PPh_3)]_2$ as resting states (Ar = Ph, 4-tolyl). These two complexes react stoichiometrically with 1 to form coupling products. The catalytic reaction kinetics is investigated by in situ IR spectroscopy revealing a two-term rate law and dependence on $[Pd_{tot}/nPPh_3]^{0.5}$ consistent with the dissociation of an off-cycle palladium dimer. The first term is independent of [1], whereas the second term is first order in [1]. The observed rates are very similar with $Pd(PPh_3)_4$, $Pd(Ph)(\kappa^1-OAc)(PPh_3)_2$, and $[Pd(Ph)(\mu-OAc)(PPh_3)]_2$ catalysts. The kinetic isotope effect varied significantly according to conditions. The multiple speciation of both AgI and PdII acts as a warning against specifying the catalytic cycles in detail. Moreover, the rapid dynamic interconversion of AgI species creates a level of complexity that has not been appreciated previously.

INTRODUCTION

Metal-mediated direct C-H bond functionalization reactions have been studied extensively as cost-effective, eco-friendly, and sustainable synthetic chemistry alternatives to conventional cross-coupling reactions, with enhanced atom economy (at least in substrate) and less metal waste. The strategies are most commonly applied to aryl-aryl bond formation as it avoids the prefunctionalization of aromatics/heteroaromatics with electropositive heteroatoms.^{2,3} A wide range of aromatic hydrocarbons have been shown to undergo C-H bond functionalization reactions in the presence of carboxylates, and this method has been successfully applied to the functionalization of electron-rich (e.g., indole), 5,6 neutral (e.g., benzene), and poor $(e.g., pyridine N-oxide)^8$ aromatic systems. The direct arylation of a fluoroarene 9-19 is an example with significant industrial interest for the potential in accessing fluorinated compounds without presynthesized organometallic species. 20,21 Furthermore, polyfluoroarenes undergo regioselective functionalizations influenced by their electronic and steric properties via the ortho-fluorine effect. 9,22-27 In this paper, we address a prototype example:

the cross-coupling of aryl iodides with pentafluorobenzene catalyzed by palladium acetate with silver carbonate as the added base.

The catalytic cycle for the direct arylation of polyfluoroarenes has been proposed to involve ambiphilic metal ligand activation (AMLA) or concerted metalation deprotonation $(CMD)^{28-31}$ between an aryl-Pd κ^1 -carboxylate intermediate and the fluoroaromatic reactant (Scheme 1a).³⁰ The AMLA(6) transition state is characterized by the agostic interaction of the arene substrate at the same time as the interaction of the arene hydrogen with the carbonyl of the coordinated carboxylate. This mechanism highlights the potential to enhance the reactivity of typically inert bonds by a combination of multiple weak interactions working in synergy; the H-bonding

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Scheme 1. (a) Commonly Proposed CMD/AMLA(6) Mechanism for Direct Arylation of Pentafluorobenzene. (b) Catalytic Cycle with Silver Performing C-H Activation Step (Adapted with Permission from ref 45, Copyright 2022, American Chemical Society)

(a)
$$F_n$$

$$Ar$$

$$Pd^0L_n$$

$$Y_2CO_3$$

$$YHCO_3$$

$$Y_1$$

$$Y_2CO_3$$

$$Y_1$$

$$Y_2$$

$$Y_2$$

$$Y_1$$

$$Y_2$$

$$Y_2$$

$$Y_3$$

$$Y_4$$

$$Y_2$$

$$Y_4$$

$$Y_2$$

$$Y_4$$

$$Y_4$$

$$Y_2$$

$$Y_4$$

interaction between the C-H bond and the carboxylate ligand increases the electron density on the C-H bond, and the resulting enhancement in the agostic interaction polarizes the C-H bond and increases the acidity of the proton.³⁰ The reactivity of isolated metal complexes in stoichiometric reactions has been used as the evidence for proposing catalytic intermediates in the AMLA(6) mechanism. This approach was highlighted by Wakioka and co-workers who reported the stoichiometric reaction of a preformed dinuclear $[Pd(Ar)(\mu OAc)(PPh_3)]_2$ (Ar = Ph, 2-MeC₆H₄, 2,6-Me₂C₆H₃) complex with 3-methylthiophene³² and benzothiazole.³³ A mononuclear complex, $Pd(Ar)(\kappa^2-OAc)(PPh_3)$, was proposed as the active catalytic species based on the equilibrium with the dinuclear $[Pd(Ar)(\mu-OAc)(PPh_3)]_2$ in solution and the isolation of stable mononuclear $[Pd(Ar)(N-BT)(\kappa^1-OAc)(PPh_3)]$ (BT = benzothiazole).

In Scheme 1a, the base is written as Y_2CO_3 with no particular role for the cation, Y, other than to remove YX, as highlighted for AgI. In the frequent situation that the base is silver carbonate or another silver salt, this mechanistic hypothesis has been unraveling. Several publications, notably from the Sanford, Hartwig, and Larrosa groups, have recently demonstrated that silver(I) complexes are active in C–H activation even in the absence of palladium. Levidence

comes from the detection of silver aryl complexes, H/D exchange reactions, and study of the kinetics of cross-coupling; a scheme summarizing the evidence may be found in ref 45. The evidence for silver participation has also been reviewed. 46-49 We recently reported 45 that the reaction of Ag₂CO₃ with XPhos in the presence of pentafluorobenzene yields Ag(C₆F₅)(XPhos) and showed that this is a mononuclear complex that is linear at silver. Its ³¹P{¹H} NMR spectra show characteristic large couplings to ¹⁰⁷Ag and ¹⁰⁹Ag (51.8 and 48.2% abundance, respectively). It undergoes ready exchange with free XPhos or its tBu analogue by an associative mechanism on a timescale of a few seconds. This complex reacts stoichiometrically with PdI(C₆H₅)(XPhos) to form the cross-coupling product C₆H₅-C₆F₅. Catalytic cross-coupling can be achieved with 5 mol % Ag(C₆F₅)(XPhos) as the sole silver source. 45 These results led us to propose a catalytic cycle (Scheme 1b) in which the C-H activation step occurs at silver and is followed by transmetalation to palladium prior to C-C bond formation. This cycle may compete with a cycle in which C-H activation occurs at palladium as in Scheme 1a. Although heterobimetallic Ag-Pd species are also plausible intermediates, no experimental evidence has been found to support their involvement. 50,51

Palladium catalyst speciation, particularly involving $Pd(0)L_n$ species, is affected by the type of phosphine, solvent, additives, and reaction conditions in general. For Xphos and PPh₃, there is experimental evidence that $Pd(0)L_n$ species can form from mixtures of $Pd(OAc)_2$ /phosphine ligand.⁵² We recognize that dinuclear Pd species (leading to other higher-order Pd clusters)⁵³ can also form, as shown by stoichiometric experiments reported by Fairlamb et al. (for PPh₃), Jutand et al. (for Xphos), and others.^{53–55} For the purposes of this paper, we refer to the active catalyst species being $Pd(0)L_n$, which is supported by our global findings (see later).

Silver carbonate is highly insoluble in solvents used for cross-coupling reactions but reacts with phosphines to form soluble silver phosphine carbonate or bicarbonate complexes. Tlahuext-Aca and Hartwig reported two {Ag(phosphine)} $_2(\mu-\kappa,^2\kappa^2$ -CO $_3$) complexes, one with a PtBuXPhos ligand that was characterized crystallographically and one with a Johnphos ligand (Scheme 2). We reported a direct analogue with

Scheme 2. Carbonate and Bicarbonate Complexes of Silver with Phosphine Ligands

 ${Ag(PR_3)}_2(\mu-\kappa^2,\kappa^2-CO_3)$ PR₃ = Johnphos, XPhos, tBuXPhos

 ${Ag(PPh_3)_2(\kappa^2-HCO_3)}_2$

 $\{Ag(PPh_3)_2\}_2(\mu\text{-}\kappa^2,\kappa^1\text{-}CO_3)$

XPhos, also characterized crystallographically. ⁴⁵ Importantly, $\{Ag(Johnphos)\}_2(\mu-\kappa,^2\kappa^2-CO_3)$ proved to be an active H/D exchange catalyst for thiophene. A related dinuclear complex of silver carbonate with triphenylphosphine $\{Ag(PPh_3)_2\}_2(\mu-\kappa,^2\kappa^1-CO_3)$ and a bicarbonate complex $\{Ag(PPh_3)_2(\kappa^2-HCO_3)\}_2$ have been described (Scheme 2), ⁵⁶ but their reactivity toward fluoroarenes is unknown.

The study of phosphines coordinated to silver is greatly aided by the presence of two I = $^{1}/_{2}$ isotopes of silver (see above). Muetterties and Alegranti showed that Ag(Pp-tol₃)_n species are extremely labile (activation energy for Ag-P rupture in [Ag(Pp-tol₃)₄]NO₃ is 9 ± 1 kcal/mol). The lability prevents observation of coupling at room temperature, but the coupling constants can be measured from low-temperature spectra, revealing a trend according to the number of phosphine ligands: [Ag(Pp-tol₃)_n]PF₆ $J(^{107}$ Ag-P) n = 4, 224 Hz; n = 3, 321 Hz; n = 2, 496 Hz. Subsequent work has confirmed that the coupling constants become larger as the coordination number of silver decreases. [S8-61]

detailed level, the value of J(Ag-P) is associated with the scharacter of the hybrid orbitals of Ag and P via the Fermi contact term. The s character of the phosphorus orbitals barely varies for different PPh₃ complexes, but the s character at silver is sensitive to the geometry and increases as the P-Ag-P angle increases within a set of complexes with related geometries. In keeping with Muetterties and Alegranti's early work, later authors have found that PPh₃ is very labile at silver, so measurements of J(Ag-P) typically require low temperatures.

In our previous paper, 45 we showed that silver carbonate reacts with pentafluorobenzene and PPh3 leading to a new ortho-fluorine resonance in the 19F NMR and a broad singlet in the ³¹P{¹H} NMR spectrum with no apparent coupling to ¹⁰⁷Ag or ¹⁰⁹Ag at room temperature. However, the ³¹P{¹H} signal splits into several resonances with conspicuous coupling to Ag nuclei on cooling to -100 °C, indicating the presence of multiple species at equilibrium (Table S8). At the same time, several 19F resonances were observed at low temperature around δ -100, characteristic of the ortho-F of Ag(C₆F₅) species. The complex speciation contrasted with the corresponding reaction with XPhos that led to a single species, $Ag(C_6F_5)(XPhos)$. Our earlier work as well as the related studies relied on the use of more specialist phosphines, leaving it unclear to what extent the results could be generalized. In this paper, we return to the use of the everyday phosphine, PPh₃, and investigate the effect of different parameters on the palladium-catalyzed cross-coupling reaction of 4-iodotoluene with pentafluorobenzene leading to improved understanding of the optimum reaction conditions. We also investigate the reaction of the carbonate and bicarbonate complexes of Scheme 2 with pentafluorobenzene in the absence of palladium and show that they are capable of C-H bond activation, albeit with complex speciation.

Instead of following the reaction by conventional solution NMR spectroscopy that is hampered by the suspensions formed with insoluble salts, we adopted two different techniques: high-resolution magic angle spinning (HR-MAS) NMR spectroscopy and in situ IR spectroscopy. HR-MAS NMR has been applied for analysis of biological samples and structural study on heterogeneous catalysts in suspension, 64-68 including real-time reaction monitoring.⁶⁹⁻⁷¹ We observed several Pd intermediates involved in the catalytic direct arylation reaction of pentafluorobenzene by HR-MAS NMR spectroscopy and confirmed their identity by ex situ MS analysis. These species were also studied in stoichiometric reactions to discriminate between the key catalytic intermediates. In situ IR spectroscopy allows the catalytic reaction to be monitored in a stirred, temperature-controlled flask attached to a Schlenk line, revealing that the kinetics requires a two-term rate law and that the kinetic isotope effect varies with conditions. Our study shows that (a) the reaction with PPh₃ is complicated by multiple speciation with both silver and palladium coordination; (b) Ag(I) plays a direct role in C-H bond activation; and (c) there are two very different pathways for catalytic reaction, resulting in unusual kinetic behavior.

RESULTS

Effect of Reaction Conditions on Catalytic Conversion to Biaryls. The direct arylation of 4-iodotoluene 2a with pentafluorobenzene 1 was selected as the model reaction system (Reactions R1).⁷² Quantitative conversion of substrate 2a was observed, and the product 4-(pentafluorophenyl)-

toluene 3a was isolated in 84% yield after purification. Conversion of 2a after 24 h was determined from the integration of the methyl $^1\mathrm{H}$ signals of reagent 2a and product 3a at δ 2.31 and 2.44, respectively. The homocoupling side-product 4,4′-dimethyl-1,1′-biphenyl was formed in trace quantities for the reaction of 2a with over 10 mol % of Pd catalyst loading or the reaction in the absence of the AgI additive but was otherwise absent. The methyl protons of this biaryl side-product were observed at δ 2.40 and did not interfere with the analysis.

The role of the acetate ligands on the precatalyst $Pd(OAc)_2$ (we abbreviate high-purity nitrite-free $Pd_3(OAc)_6$ to $Pd(OAc)_2$ throughout the paper) was studied by comparing the yields of biaryl product 3a for reactions using $Pd(PPh_3)_4$, $Pd(PPh_3)_2Cl_2$, and $PdCl_2$ precatalysts, in each case with 0.75 equiv Ag_2CO_3 (Table 1). Results comparable with $Pd(OAc)_2 + 10$ mol %

Table 1. Conversions of 2a and Yields of Isolated Product 3a

entry	catalyst ^a	Ag_2CO_3 (equiv)	PPh ₃ (mol %)	% conversion ^a (% yield) ^b
1	5 mol % Pd(OAc) ₂	0.75	10	quant. (84)
2	5 mol % Pd(PPh ₃) ₄	0.75	0	quant. (85)
3	5 mol % Pd(PPh ₃) ₂ Cl ₂	0.75	10	quant. (87)
4	5 mol % PdCl ₂	0.75	10	72 (61)
5	5 mol % Pd(PPh ₃) ₂ Cl ₂ ,	0.75	0	60 (57)
7	1.5 mol % Pd(OAc) ₂	0	3	0
8	1.5 mol % PdCl ₂	0	3	0
9	20 mol % Pd(OAc) ₂	0	40	(14)
10	20 mol % PdCl ₂	0	40	0
11	20 mol % Pd(PPh ₃) ₄	0	0	0

^aStandard reaction time 24 h. Determined from integration of methyl ¹H NMR peaks of the reagent 2a and the product 3a. ^bAfter purification by flash chromatography.

PPh₃ (100% NMR yield, 84% isolated yield) were obtained for reactions catalyzed by Pd(PPh₃)₄ and Pd(PPh₃)₂Cl₂. The ligand PPh₃, particularly in excess greater than 3, is understood to play a role in both the reduction of Pd^{II} to Pd⁰ and the stabilization of the oxidative addition complex in stoichiometric studies. 73

We also investigated the effect of changing the base on the catalytic reaction (Table S1). Of the metal carboxylates tested, Ag₂CO₃ resulted in the highest conversion of substrate **2a** and yield of the isolated product **3a**. Similar yields were obtained with Pd(OAc)₂ + 10 mol % PPh₃ + 1.5 equiv [Me₄N][OAc] or with Pd(OAc)₂ + 10 mol % PPh₃ + 0.75 equiv Ag₂O. Changing the base to any of K₂CO₃, Cs₂CO₃, AgOAc, or [Bu₄N][OAc] resulted in marked reductions in yields. A recent paper reports H/D exchange of pentafluorobenzene with 10 mol % Cs₂CO₃ in d_6 -acetone or d_6 -DMSO.⁷⁴ Some of the action of bases that

we observe may be caused by deprotonation of C_6F_5H . Nevertheless, only $[Me_4N]OAc$ and Ag_2CO_3 give full conversion in the catalytic reaction.

Alternative solvents utilized in the literature for C–H bond functionalization reactions include dialkyl carbonates, PEG, and water. The polar aprotic solvents such as dimethylacetamide (DMAc) and N-methyl 2-pyrrolidone (NMP) resulted in comparable yields to DMF (Table S2). Ethylene carbonate and propylene carbonate afforded 3a in reasonable yields.

A variety of monophosphines were tested for comparison to PPh₃ (Table S3). Of these, P(4-FC₆H₄)₃, P(3,5-(CF₃)₂C₆H₃)₃, and P(2-furyl)₃ were as effective as PPh₃. Substitution with methoxy groups proved deleterious, whereas PCy₃ was very poor indeed.

Under the standard conditions, the reaction solution remains clear until the reaction is ca. 70% complete, but subsequently, the solution darkens, suggesting the formation of Pd metal (or possibly Ag). The use of preformed stabilized Pd nanoparticles (Pd-NPs) as an alternative to Pd(OAc)₂ precatalyst was therefore examined to establish the type of catalyst (homogeneous and/or heterogeneous) involved in the reaction, recognizing that stabilized PdNPs are usually less active than naked, polar aprotic solvent stabilized PdNPs. The catalytic activities of Pd-NPs supported on polyvinylpyrrolidone (PVP) of different polymer weights and particle sizes were tested (Table S4) while maintaining the molar quantity of the Pd at 5 mol %. The reaction at 70 °C required PPh₃, and the isolated yields were significantly lower than when using Pd(OAc)₂ precatalyst.

The structure—reactivity relationship between the electronic properties of the fluoroarenes and the reaction rates was studied to characterize the transition state based on a modified Hammett equation. The electronic properties of fluoroarenes were tuned by varying the substituents on the C1 position of 2,3,5,6-tetrafluorobenzene. Functional groups with electronic properties ranging from electron-donating dimethylamino to electron-withdrawing trifluoromethyl were selected. The reactions of these fluoroarenes achieved quantitative conversion of the starting material to the desired products after 20 h at 70 °C. The substituent effect was determined from relative yields obtained by competition reaction between 4-iodotoluene with 10 equiv each of the 1-X-2,3,5,6-tetrafluorobenzene (X = NMe₂, OMe, F, Cl, and CF₃) and pentafluorobenzene analyzed by 19 F NMR (eq 2).

The Hammett equation was modified by substitution of the relative reaction rates represented by the ratio of the two product concentrations (P^X/P^F) with the standard set as X=F instead of X=H (eq 1). Modest increases in yields were achieved with more electron-withdrawing functional group in the C1 position (Table S5). A linear free-energy relationship (LFER) was observed with the logarithms of the relative yields directly proportional to σ^+ (Figure 1) yielding R^2 of 0.954. The reaction constant (ρ) determined from the slope of the LFER was +0.28 \pm 0.02, consistent with the AMLA(6) pathway that is expected to favor electron-withdrawing substituents stabilizing the negative charge at the TS.

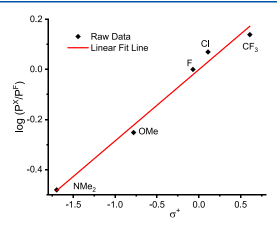


Figure 1. Hammett plot of the competition reaction shown in Reaction R2.

$$\log\left(\frac{P^{X}}{P^{F}}\right) = \rho \cdot \sigma^{+} \tag{1}$$

where $P^{\rm X}$ = integration of 1-X-2,3,5,6-tetrafluorobiaryl, $P^{\rm F}$ = integration of pentafluorobiaryl, ρ = reaction constant, and σ^+ = substituent constant.

Ag Speciation and Possible Intermediates. Following the reports of C–H activation by silver salts, we investigated H/D exchange between pentafluorobenzene and D_2O catalyzed by Ag(I) salts, monitoring the reaction by ^{19}F NMR spectroscopy. Replacement of H by D causes an isotopic shift of the ortho-fluorine resonance of +0.3 ppm and loss of F–H coupling. The exchange with 1 equiv D_2O in DMF at 50 $^{\circ}C$ catalyzed by Ag₂CO₃ (10 mol %) and PPh₃ (20 mol %) resulted in 99% deuteration (Reaction R3). When the triphenylphosphine was omitted, the reaction resulted in 51% deuteration, whereas the corresponding reaction using Ag-

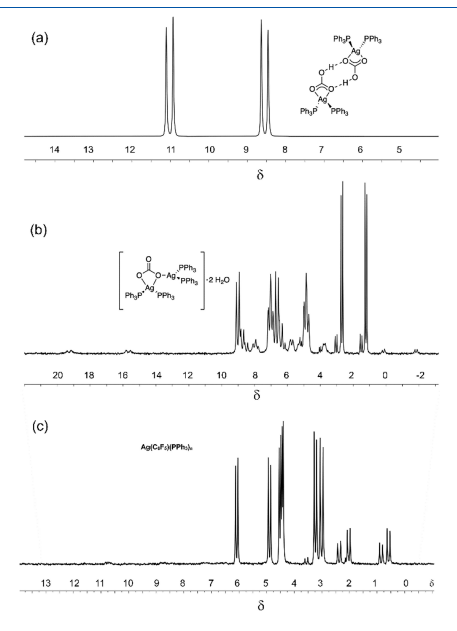


Figure 2. Low-temperature $^{31}P\{^{1}H\}$ NMR spectrum in toluene/dichloromethane (20:80) of (a) $[Ag(PPh_3)_2(\kappa^2-HCO_3)]_2$ at -80 °C, (b) $[Ag(PPh_3)_2]_2CO_3\cdot 2H_2O$ at -100 °C, and (c) $Ag(C_6F_5)(PPh_3)_n$ at -100 °C.

 $(PPh_3)_2(\kappa^2\text{-OAc})$ (see below) in place of $Ag_2CO_3 + PPh_3$ yielded 37% deuteration.

When AgOAc (1 equiv) and PPh₃ (2 equiv) were reacted with pentafluorobenzene (10 equiv), no Ag(C_6F_5)-containing product was detected by $^{19}F\{^1H\}$ NMR spectroscopy. Replacement of AgOAc by Ag₂CO₃ (1 equiv) resulted in the detection of an Ag(C_6F_5)-containing product with ortho-F resonances centered at δ –106.5 (in CH₃CN). The reaction of AgOAc (1 equiv) or Ag₂CO₃ (1 equiv) with C_6F_5H (10 equiv) in the absence of phosphine did not generate any detectable Ag(C_6F_5) complex, indicating that the PPh₃ ligand is critical for C–H activation (Table S6). To investigate this reactivity further, the PPh₃-coordinated Ag carbonate complexes in Scheme 2 were synthesized following literature methodologies, ⁵⁶ together with Ag(PPh₃)₂(κ ²-OAc). ⁸¹

The Ag-PPh₃ bond was found to be highly labile at room temperature, resulting in a singlet at room temperature in the ³¹P{¹H} NMR spectrum of these complexes. The complexes $\{Ag(PPh_3)_2(\kappa^2-HCO_3)\}_2$ and $\{Ag(PPh_3)_2\}_2(\mu-\kappa, \kappa^2\kappa^1-CO_3)$ were characterized using low-temperature \$\frac{31}{2}P\{^1H\} NMR spectroscopy. The $\{Ag(PPh_3)_2(\kappa^2-HCO_3)\}_2$ complex existed as a well-defined species in solution, and the ³¹P{¹H} NMR spectrum was fully resolved at -80 °C to reveal a pair of overlapping doublets centered at δ 9.78 with coupling constants ${}^{1}J_{^{31}P} - {}^{107}Ag = 465 \text{ Hz} \text{ and } {}^{1}J_{^{31}P} - {}^{109}Ag = 537 \text{ Hz (Figure})$ 2a). These values may be compared with those for [Ag- $(XPhos)_{2}(CO_{3})$ of 634 and 731 Hz; the reduction in the PPh3 complex relative to the XPhos complex reflects the increased coordination number of Ag. 57-60 In contrast, the low-temperature ${}^{31}P\{{}^{1}H\}$ NMR showed that $\{Ag(PPh_3)_2\}_2(\mu$ - $\kappa_1^2 \kappa^1$ -CO₃) exists in equilibrium with multiple Ag-PPh₃ containing species. Most of the resonances appeared as overlapping doublets with characteristic Ag-P coupling; the chemical shifts and coupling constants of the major species could be identified clearly, but other peaks were difficult to distinguish due to peak overlap (Figure 2b, Table S9). The distinguishable coupling constants varied from 317 to 778 Hz for ¹⁰⁹Ag, indicating the presence of species with a wide range of coordination numbers. Thus, the published crystal structure⁵⁶ represents one of many different species that are formed in solution. Because carbonate can act as a bridging ligand in several different ways, 82 we can envisage numerous different oligomeric structures.

The phosphine-coordinated Ag complexes were reacted with C_6F_5H (10 equiv) to examine the C–H activation capabilities of these complexes. The presence of a C–H activation product, $Ag(C_6F_5)(PPh_3)_n$, was detected by the ortho-F resonance at δ –106 using $^{19}F\{^1H\}$ NMR spectroscopy with $[Ag(PPh_3)_2]_2CO_3\cdot 2H_2O$. Because the structure of the resulting $Ag(C_6F_5)(PPh_3)_n$ complex is poorly defined, percent conversions could not be measured accurately. Only when C_5CO_3 is added is a reaction between $\{Ag(PPh_3)_2(\kappa^2-HCO_3)\}_2$ and C_6F_5H observed, either at 60 °C or at room temperature (Table 2). We also tested $Ag(PPh_3)_2(\kappa^2-OAc)$ for reaction with C_6F_5H but found no evidence for C–H activation. These experiments show that the carbonate complex $[Ag^2-C_5H]_3$

Table 2. Stoichiometric Reaction of PPh₃-Coordinated Ag Complexes and C₆F₅H^a

entry	conditions	temperature (°C)	detection of $Ag(C_6F_5)$ product
1	$[Ag(PPh3)2(\kappa^2-HCO3)]2 (0.5 equiv)$	60	no
2	$[Ag(PPh3)2(\kappa^2-HCO3)]2 (0.5 equiv) + Cs2CO3 (1 equiv)$	60	yes
3	$[Ag(PPh_3)_2]_2CO_3\cdot 2H_2O$	60	yes
4	$[Ag(PPh_3)_2]_2CO_3\cdot 2H_2O$	RT	yes
5	$Ag(PPh_3)_2(\kappa^2-OAc)$ (1 equiv)	60	no

^a10 equiv C_6F_5H . ^bDetected by the ortho-F resonances at δ –106.0.

(PPh₃)₂]₂CO₃·2H₂O is capable of C-H bond activation but not the bicarbonate complex or the acetate complex.

To study $Ag(C_6F_5)(PPh_3)_n$ further, an authentic sample of $Ag(C_6F_5)$ was prepared by literature methods⁸³ and then reacted with PPh_3 (2 equiv) in acetonitrile (Reaction R4). The resulting product was isolated from solution as brown crystals and characterized by low-temperature NMR spectroscopy (Figure 2c). The ³¹P{¹H} NMR spectrum showed five major pairs of Ag-coupled doublets and one minor pair (Table S10). Considering the tetrameric structure of several silver aryl complexes ^{84–86} and the infinite chain structure of $Ag(C_6F_5)$ -(EtCN)₂, ⁸³ the presence of multiple species at low temperature may indicate both variable numbers of coordinated phosphines and formation of different oligomers.

The complex and multiple speciation demonstrated first by the low-temperature spectra of $\{Ag(PPh_3)_2\}_2(\mu-\kappa, \kappa^2\kappa^1-CO_3)$ and second by the reaction of Ag(C₆F₅) with PPh₃ is reminiscent of the spectrum obtained by the reaction of Ag₂CO₃ with PPh₃ and C₆F₅H,⁴⁵ suggesting that there may be species in common. Comparison of the spectra (Tables S8-S10) shows three sets of resonances that are sufficiently close in coupling constant and chemical shift to belong to the same species for all three spectra (Table S11). Quoting the values for $\{Ag(PPh_3)_2\}_2(\mu-\kappa,^2\kappa^1-CO_3)$, the first two are at δ 2.0 (*J* 275, 317 Hz) and δ 2.3 (J 285, 329 Hz). Literature data for $[Ag(PPh_3)_3X]$ (X = BF₄, Cl, I with $J(^{107}Ag-P)$ 318, 227, and 262 Hz, respectively)⁶¹ and for $[Ag(PPh_3)_4][BF_4] J(av) = 238$ Hz^{63} (hence, $J(^{107}Ag-P) = 221 Hz$) suggest that the species we observe close to δ 2.0 are most likely of the type [Ag(PPh₃)₃X]. There is one further species common to $\{Ag(PPh_3)_2\}_2(\mu-\kappa_1^2\kappa^1-CO_3)$ and $Ag_2CO_3/PPh_3/C_6F_5H$ at δ 17.5 (J 677, 778 Hz) that may be due to $[Ag(PPh_3)]_2(CO_3)$ or similar species.

It seemed likely that the amount of silver salt could be reduced to catalytic quantities if an alternative stoichiometric base was employed. The catalytic abilities of silver carbonate and the Ag-carbonate triphenylphosphine complexes were

tested employing Cs_2CO_3 (0.75 equiv) as the base in each reaction (Table 3). When Cs_2CO_3 is used alone for the

Table 3. Influence of Catalytic Amounts of Ag Salts and Complexes on Direct Arylation

			conv	NMR ersio- (%)
entry	PPh ₃ (mol %)	base Cs_2CO_3 (0.75 equiv) + Ag salt	3 h	23 h
1	10		15	69
2	5	$[Ag(PPh_3)_2(\kappa^2\text{-HCO}_3)]_2 \text{ (5 mol \%)}$	51	98
3	5	$[Ag(PPh_3)_2]_2CO_3$ (5 mol %)	62	99
4	10	Ag ₂ CO ₃ (5 mol %)	73	94
5	5	$[Ag(PPh_3)_2(\kappa^2-HCO_3)]_{2,3}$ (2.5 mol %)	54	93
6	5	[Ag(PPh ₃) ₂] ₂ CO ₃ ·2H ₂ O (2.5 mol %)	59	96
7	10	Ag ₂ CO ₃ (2.5 mol %)	57	89
8	10	Ag ₂ CO ₃ (1 mol %)	44	80
9	10	$Ag(C_6F_5)$ (10 mol %)	70	96

^aThe conversion was calculated by integrating the CH_3 resonance of **2a** (δ 2.30) with respect to 1,3,5-trimethoxybenzene (δ 3.78) in the ¹H NMR spectrum of the mixture with D1 = 1 s.

standard catalytic direct arylation reaction, the conversion after heating for 23 h is 69%. Trace amounts of side-product were also detected in the 1H NMR spectrum. When 5 mol % of either $[Ag(PPh_3)_2(\kappa^2\text{-HCO}_3)]_2$ or $\{Ag(PPh_3)_2\}_2(\mu\text{-}\kappa,^2\kappa^1\text{-CO}_3)$ was used with Cs_2CO_3 (0.75 equiv) as the base, the conversion improved to 98 and 99%, respectively (entries 2 and 3), and the side-product was no longer detected. With 5 mol % of $Ag_2\text{CO}_3$, the greatest conversion is 73% at 3 h, and the reaction reaches near-completion by 23 h (94% conversion) (entry 4). Trace amounts of the homocoupling side-product were also detected. The amount of Ag used can be decreased to 2.5 mol

% without an appreciable decrease in the conversion, but when 1 mol % of Ag_2CO_3 was used, the 1H NMR yield dropped to 80% (entries 4 and 8). The use of 10 mol % $Ag(C_6F_5)$ with 10 mol % PPh₃ and Cs_2CO_3 (0.75 equiv) resulted in almost identical conversion (entry 9) to the experiments with other Ag complexes. These reactions demonstrate that the conversion can be increased to well above 90% by combining sub-stoichiometric Cs_2CO_3 with catalytic quantities of any of Ag_2CO_3 , $[Ag(PPh_3)_2(\kappa^2\text{-HCO}_3)]_2$, $[Ag(PPh_3)_2]_2CO_3$, or $Ag(C_6F_5)$, thus eliminating the need for large quantities of silver salts.

Pd Speciation: Studies by HR-MAS and Conventional NMR. *Identification of Catalytic Resting States*. The direct arylation of 4-iodobenzene 2b with pentafluorobenzene 1 was selected as the model reaction system to be studied by HR-MAS NMR spectroscopy due to the ease of synthesis of phenyl-Pd species compared to their 4-tolyl analogues (Reaction R5). The line-broadening observed by standard NMR spectroscopy for the inhomogeneous reaction mixture was successfully resolved by the application of HR-MAS NMR spectroscopic analysis as shown in a test sample of 1, 2a, and Ag₂CO₃ (Figure 3).

The catalyst loading was increased from catalytic (i.e., 5 mol %) to sub-stoichiometric (i.e., 0.5 equiv) to observe possible species involved in the catalytic cycle. After heating at 56 °C in the rotor, broad phenyl signals were observed that disappeared on completion of the reaction. The reaction was repeated, but this time, the probe was cooled after 20% conversion to products. The broad signals observed at δ 6.83 and 6.69 during the reaction at 56 °C in the ¹H NMR spectrum were resolved into two triplets at δ 6.91 and 6.75 at -24 °C. The carboxylic acid proton of the AcOH at δ 12.63 was also observed upon cooling. 2D ¹H-¹H COSY spectra revealed the presence of two species in the aromatic region (Figure S12). The triplet

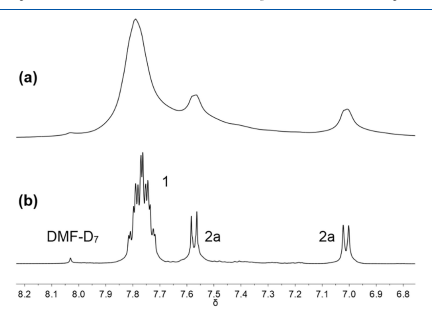


Figure 3. ¹H NMR spectra of a mixture of 1, 2a, and silver carbonate in DMF-d₇ at 13 °C (a) without spinning and (b) spinning at 3 kHz.

signal at δ 6.91 (J=7.1 Hz) was correlated with the other triplet at δ 6.75 (J=7.1 Hz) and to a signal at δ 7.29 overlapping with other aromatic signals. A second species was observed with correlation between weak signals at δ 6.29 and 6.53. Five well-known Pd complexes were synthesized for comparison and were heated to 56 °C for 90 min in DMF- d_7 to simulate the catalytic conditions before measurement of NMR spectra at -24 °C (Scheme 3, Table S12).

Scheme 3. Potential Pd(II)-Resting States

Comparison of the phenyl resonances and the ³¹P NMR signals of the Pd complexes with those of the intermediate observed during the reaction revealed that the reaction mixture contained Pd2-OAc with trace quantities of Pd1-OAc (Figure 4). Additionally, of the five Pd species analyzed, Pd2-OAc was the only complex retaining complete structural integrity after heating to 56 °C. Decomposition products such as OPPh₃ were observed for other complexes; dinuclear Pd2-OAc and Pd2-I species were observed in the ¹H and ³¹P NMR spectra of mononuclear Pd1-OAc and Pd1-I complexes, respectively.

The well-resolved phenyl signals of Pd2-OAc in DMF- d_7 at 56 °C were broadened when the dinuclear complex was used as the catalyst for the reaction of 1 with 2b (Reaction R6). It is likely that Pd2-OAc is in equilibrium with Pd1-OAc. Furthermore, the use of Pd2-OAc as the catalyst for the direct arylation reaction of 2a with 1 yielded a mixture of two biaryl products in 3.5:1 ratio (3a:3b). Thus, Pd2-OAc reacted with 1 to form pentafluorophenylbenzene 3b and then

catalytically turned over **2a** to form the 4-(pentafluorophenyl)-toluene **3a**.

Whereas the reaction catalyzed by 25 mol % Pd2-OAc achieved 12% product 3b formation after 60 min, the addition of 25 mol % PPh $_3$ improved the yield of 3b to 25% after the same time. For the reaction with additional 75 mol % PPh $_3$, 57% yield was achieved after 20 min. The addition of extra PPh $_3$ resulted in increased formation of species with $^1\mathrm{H}$ and $^{31}\mathrm{P}\{^1\mathrm{H}\}$ NMR spectra very similar to those of Pd1-OAc.

The involvement of Pd2-OAc and Pd1-OAc as catalytic resting states was further supported by ex situ LIFDI mass spectrometric studies. ^{87–89} Reference spectra of Pd(tol)(κ^1 -OAc)(PPh₃)₂ and [Pd(tol)(μ -OAc)(PPh₃)]₂, the tolyl analogues of Pd2-OAc and Pd-1-OAc, are shown in Figure 5 (the LIFDI spectra of Pd2-OAc and Pd-1-OAc are in Figures S14 and S15). LIFDI spectra from the direct arylation reaction of 2a with 1 revealed the [Pd(tol)(+ μ -OAc)(PPh₃)]₂ and Pd(tol)(κ^1 -OAc)(PPh₃)₂ complexes at m/z=518.05 and 1038.16, respectively, during the reaction (Table S13 and Figures S18–S21).

Stoichiometric Reaction of Palladium Species with Pentafluorobenzene. The dinuclear species $[Pd(Ph)(\mu-X)-(PPh_3)]_2$ (where $X = OAc\ Pd2-OAc$, OH Pd2-OH, and I Pd2-I) were heated at 70 °C in the presence of 20-fold excess of pentafluorobenzene 1 in DMF (Scheme 4, reaction A, compare ref 32 and 33). Formation of the product 3b was observed for the reaction of Pd2-OAc and Pd2-OH in quantitative and 60% yields, respectively, but not for Pd2-I (Table 4, entries 1–3). However, partial formation of the product 3b was observed for Pd2-I in the presence of Ag_2CO_3 or AgOAc (entries 4 and 5). The mononuclear species Pd1-OAc and Pd1-I were heated similarly (Scheme 4, reaction B). The Pd1-OAc complex formed the product 3b in 45% yield (Table 4, entry 6). However, excellent conversion of the complex was achieved in

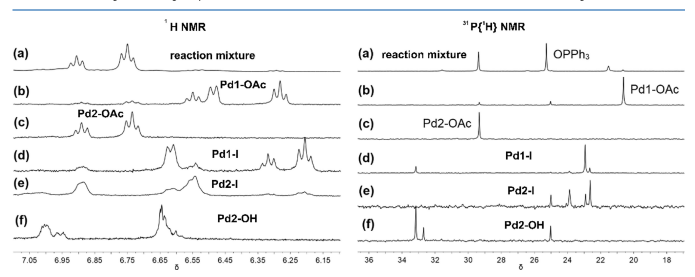
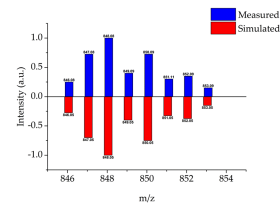


Figure 4. ¹H NMR spectra in the aromatic region (left) and ³¹P{¹H} NMR spectra (right) of (a) the reaction shown in Reaction R5, (b) $Pd(Ph)(\kappa^{1}-OAc)(PPh_{3})_{2}$ Pd1-OAc, (c) $[Pd(Ph)(\mu-OAc)(PPh_{3})]_{2}$ Pd2-OAc, (d) $Pd(Ph)(1)(PPh_{3})_{2}$ Pd1-I, (e) $[Pd(Ph)(\mu-I)(PPh_{3})]_{2}$ Pd2-I, and (f) $[Pd(Ph)(\mu-OH)(PPh_{3})]_{2}$ Pd2-OH. Spectra collected after heating at 56 °C and measured at -24 °C in DMF- d_{7} .



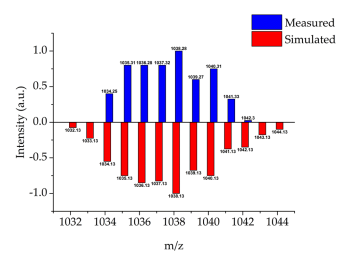


Figure 5. LIFDI-MS spectrum of (above) $Pd(4-tolyl)(I)(PPh_3)_2$ ($PdC_{43}H_{37}IP_2$ requires 848.05 for ^{106}Pd) and (below) [$Pd(4-tolyl)(\mu-OAc)(PPh_3)$]₂ ($Pd_2C_{54}H_{50}O_4P_2$ requires 1038.13 for $^{106}Pd^{108}Pd$).

Scheme 4. Stoichiometric Reaction of 1 in DMF [Reaction A: with Dinuclear Pd Species (X = OAc Pd2-OAc, OH Pd2-OH, and I Pd2-I); Reaction B: with Mononuclear Pd Species Pd1-OAc and Pd(Ph)(I)(PPh₃)₂ Pd1-I]

the presence of Ag_2CO_3 (99%) and AgOAc (98%) (entries 7 and 8). As with the dinuclear iodide complex, the mononuclear $Pd(Ph)(I)(PPh_3)_2$ **Pd1-I** complex was unreactive toward pentafluorobenzene 1 unless in the presence of Ag_2CO_3 (49%) and AgOAc (67%) (entries 9–11). No product was formed in the presence of $AgBF_4$ (entry 12). In interpreting these results, it should be recalled that only **Pd2-OAc** is stable on heating to 70 °C.

Speciation in the Absence of Substrate. Recent studies have demonstrated the formation of Pd^I clusters with bridging phosphide ligands in the reactions of Pd(OAc)₂ and PPh₃ in

Table 4. NMR Yields of 3b from Stoichiometric Reactions of Pd Species and 1 in DMF^a

entry	complex	base	NMR conversion (%) ^b
1	Pd2-OAc	none	99
2	Pd2-OH	none	60
3	Pd2-I	none	0
4	Pd2-I	3 equiv Ag ₂ CO ₃	36
5	Pd2-I	7 equiv AgOAc	22
6	Pd1-OAc	none	45
7	Pd1-OAc	2 equiv Ag ₂ CO ₃	99
8	Pd1-OAc	7 equiv AgOAc	98
9	Pd1-I	none	0
10	Pd1-I	2 equiv Ag ₂ CO ₃	49
11	Pd1-I	4 equiv AgOAc	67
12	Pd1-I	7 equiv AgBF ₄	0

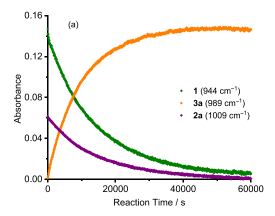
"Conditions as shown in Scheme 4 other than base. ^bBased on integration of C_6F_6 (internal standard) and 3,5-fluorines of 3a and 3b.

THF. ^{53,90} We were concerned to establish whether these species are formed under the conditions of our cross-coupling reactions. We therefore examined the nature of the Pd species formed in the absence of substrates by conventional $^{31}P\{^1H\}$ NMR spectroscopy, searching for characteristic phosphide resonances in the δ 200 region. Stoichiometric reactions of Pd(OAc)₂ with PPh₃ (2 equiv) in DMF for 20 min at 60 °C with and without Ag₂CO₃ yielded no evidence for phosphide species. There were traces of phosphide species after prolonged reaction (15 h) in the presence of Ag₂CO₃. Corresponding reactions in the presence of AgOAc yielded no evidence of phosphide species. The dominant product in all these reactions is OPPh₃.

REACTION KINETICS

Monitoring the Reaction by IR Spectroscopy. The course of the catalytic cross-coupling reaction could be monitored by in situ IR spectroscopy. Distinctive absorption bands were identified for pentafluorobenzene, iodotoluene, and their cross-coupling product in a region where the DMF solvent does not absorb significantly (Reaction R7). The in situ FTIR spectroscopic measurements were recorded by the ReactIR with a silicon probe dipping into a stirred flask with a thermocouple measuring the temperature in the solution. An example of the reaction profiles is shown in Figure 6a for a 1:1 substrate ratio and 5 mol % Pd(OAc)2. To validate the method, the conversion of 4-iodotoluene 2a to biaryl product 3a was determined by integrating the methyl peaks observed by ¹H NMR analysis of aliquots sampled at regular intervals in a reaction with a 10:1 ratio of 1:2a (Figure 6b). The kinetic profiles observed by FTIR and NMR were in good agreement. The reaction exhibited no induction period and reached 50% completion in ca. 6600 s and 90% completion after 15,000 s at 56 ± 1 °C under the conditions illustrated, making it clear that the temperature could be lowered and the reaction time reduced compared to the standard of Reaction R1.

Determination of the Rate Law. Detailed kinetic analysis was performed using this *in situ* IR approach by traditional



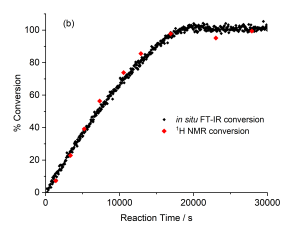


Figure 6. (a) Changes in the IR absorbance observed during the reaction shown in Reaction R7. Concentrations: pentafluorobenzene 1 (201 mM), 4-iodotoluene 2a (200 mM), Pd(OAc)₂ (10.1 mM), PPh₃ (20.2 mM), with Ag₂CO₃ (0.75 equiv) in DMF at 56 ± 1 °C. (b) Progress of the reaction with 10-fold excess of 1 (*i.e.*, 0.18 M) generated by (black) *in situ* FTIR spectroscopy and (red) conversion determined by NMR spectroscopic analysis of sampled aliquots.

kinetic methods and by variable time normalization analysis (VTNA) methods. Full details of the concentrations used together with the results from the VTNA methods are given in the Supporting Information (Figures S24-S26). Here we report the results from traditional isolation methods. Initial experiments were performed with a 10-fold excess of either 1 or 2a (initial concentrations 0.018 and 0.18 M). With excess 1, the decay of 2a was linear to 50% conversion ($R^2 = 0.997$); with excess 2a, the decay of 1 was exponential with a linear fit to $\ln [1]$ to 50% conversion ($R^2 = 0.998$). These results are consistent with a reaction that is zero order in 2a and first order in 1. A more detailed analysis of the dependence of pseudo-zero-order rate constant on [1] was carried out with [1]/[2a] ranging from 10 to 50, revealing a two-term rate law with slope $(2.83 \pm 0.26) \times 10^{-6} \text{ s}^{-1}$ and intercept $(1.12 \pm 0.26) \times 10^{-6} \text{ s}^{-1}$ $0.10) \times 10^{-6} \text{ mol dm}^{-3} \text{ s}^{-1} \text{ (Figure 7)}.$

The rate dependence on $[\mathrm{Pd}_{\mathrm{tot}}/2\mathrm{PPh_3}]$ at 56 ± 1 °C was measured between 0.19 and 3.7 mM catalyst concentration under pseudo-zeroth-order reaction conditions with a 10-fold excess of pentafluorobenzene 1 (0.18 M) to 4-iodotoluene 2a (0.018 M) (Table S14). Because the $\mathrm{Pd}(\mathrm{OAc})_2/\mathrm{PPh_3}$ ratio was maintained at 1:2 for each kinetic measurements, we refer to the concentration $[\mathrm{Pd}_{\mathrm{tot}}/2\mathrm{PPh_3}]$. The plot of k_{obs} vs $[\mathrm{Pd}_{\mathrm{tot}}/2\mathrm{Ph_3}]$

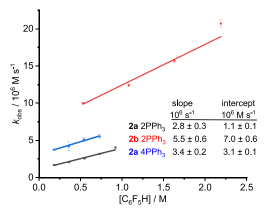


Figure 7. Dependence of pseudo-zero-order rate constant on [1]: black, reaction with 2a with Pd/PPh₃ 1:2; red, reaction with 2b with Pd/PPh₃ 1:2; blue, reaction with 2a with Pd/PPh₃ 1:4. Higher concentrations were used for 2b than for 2a because of lower IR absorption coefficients. [Catalyst] 5 mol % wrt 2a/2b: 0.92 and 2.8 mM for reaction with 2a and 2b, respectively. [2a] 18 mM, [2b] 55 mM. Ag_2CO_3 0.75 equiv, temp 56 ± 1 °C.

 $^{2PPh_3}]^{0.5}$ gave better correlation coefficients (Figure 8) than the plot of $k_{\rm obs}$ vs $[{\rm Pd_{tot}}/{\rm 2PPh_3}]^{0.75}$ whose correlation

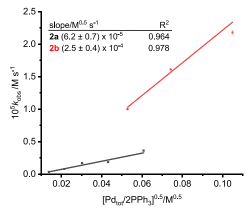


Figure 8. Plot of $k_{\rm obs}$ against $[{\rm Pd}_{\rm tot}/2{\rm PPh}_3]^{1/2}$ for the direct arylation reaction of 4-iodotoluene ${\bf 2a}$ (black) and iodobenzene ${\bf 2b}$ (red) at 56 \pm 1 °C recorded under pseudo-zeroth-order conditions with 10-fold excess of 1. Higher concentrations were used for ${\bf 2b}$ than for ${\bf 2a}$ because of lower IR absorption coefficients. [1] 180 mM for reaction with [2a] 18 mM, [1] 560 mM for reaction with [2b] 55 mM; Ag₂CO₃ 0.75 equiv, temp ${\bf 56} \pm 1$ °C.

coefficients, in turn, were much better than those for the plot of $k_{\rm obs}$ vs $[{\rm Pd_{tot}/2PPh_3}]$. The kinetics was measured similarly for iodobenzene **2b** as substrate. The concentration of the limiting reagent **2b** was increased to 55 mM (from 18 mM used for **2a**) to allow for the lower peak intensities observed for **2b** (at $1016~{\rm cm^{-1}}$) and the product **3b** (at $989~{\rm cm^{-1}}$). The kinetics follows a similar pattern to that for **2a** (Figures 7 and 8).

Thus, the overall rate law is given by eq 2:

Rate =
$$k_1[Pd_{tot}/nPPh_3]^{0.5} + k_2[Pd_{tot}/nPPh_3]^{0.5}[C_6F_5H]$$
(2)

The values of k_1 and k_2 are estimated by dividing the slopes of Figure 7 by $[Pd_{tot}/2PPh_3]^{0.5}$ (Table 5). The value of k_1 is about 3 times larger for **2b** than for **2a**, but the values of k_2 are very similar.

Table 5. Overall Rate Constants for Reaction of 1 with 2a and 2b^a

entry	$k_1/10^{-4} \text{ M}^{1/2} \text{ s}^{-1}$	$k_2/10^{-4} \text{ M}^{-1/2} \text{ s}^{-1}$		
2a Pd 2PPh ₃	0.336 ± 0.045	0.999 ± 0.075		
2b Pd 2PPh ₃	1.14 ± 0.21	1.21 ± 0.14		
2a Pd 4PPh ₃	1.02 ± 0.03	1.29 ± 0.081		
a 56 \pm 1 °C, DMF, Ag ₂ CO ₃ 0.75 equiv.				

Triphenylphosphine can play several roles including acting as a reducing agent for Pd(OAc)2, ligand for silver, and ligand for palladium. Moreover, several species may form in each role. For these reasons, we investigated the effect of the Pd/PPh₃ ratio on the reaction kinetics. At a constant 5 mol % Pd(OAc)₂, the reaction kinetics was studied with [PPh₃] ranging from 5 mol % (1:1 Pd/PPh₃) to 40 mol % (1:8 Pd/ PPh₃). The rate increased linearly up to 1:4 Pd/PPh₃ but decreased slightly at higher concentrations than 1:4 and then leveled off (Figure S31). The dependence of the rate on [C₆F₅H] (at 0.93 mM catalyst) exhibited a very similar slope to that for 1:2 Pd/PPh3 but a markedly larger intercept (Figure 7). The variation of the rate with [Pd_{tot}/4PPh₃] was also determined (Figure S32); whereas the slope of $k_{\rm obs}$ vs [Pd_{tot}/ $(6.2 \pm 0.7) \times 10^{-5} \, \text{M}^{0.5} \, \text{s}^{-1}$, that of $[\text{Pd}_{\text{tot}}/2\text{PPh}_3]^{0.5}$ was $(6.2 \pm 0.7) \times 10^{-5} \, \text{M}^{0.5} \, \text{s}^{-1}$, that of $[\text{Pd}_{\text{tot}}/2\text{PPh}_3]^{0.5}$ was $(15.6 \pm 2.0) \times 10^{-5} \, \text{M}^{0.5} \, \text{s}^{-1}$, an increase of a factor of 2.5. The overall rate constants are given in Table 5; the most marked change is the increase in the k_1 term by a factor of 3. This change can be understood if the oxidative addition of aryl iodide precedes the rate determining step. Thus, the rate is determined by the reactivity of Pd- $(PPh_3)_p(Ar)I$ or related species (see below).

The reaction kinetics was also determined for a range of 4-substituted iodoarenes in place of iodotoluene. Additionally, the conversions were determined in competition between 4-iodotoluene and alternative 4-substituted iodoarenes. The effects of substitution were very minor (Table S22).

Alternative Catalysts. The rate of coupling of penta-fluorobenzene 1 with 4-iodotoluene 2a using Ag_2CO_3 (0.75 equiv) and $Pd(OAc)_2/2PPh_3$ catalyst was compared to that with $Pd(PPh_3)_4$ at 50 ± 1 °C. The rates were $(3.05 \pm 0.03) \times 10^{-5}$ and $(9.12 \pm 0.04) \times 10^{-5}$ mol dm⁻³ s⁻¹, respectively (Table S23, Figure S33). Considering that we know that the rate for $Pd(OAc)_2$ is enhanced by a factor of 2.5 (at 56 °C) by increasing the PPh_3/Pd ratio to 4, we conclude that the difference in performance of $Pd(OAc)_2/4PPh_3$ and $Pd(PPh_3)_4$ is very slight.

The kinetics of reactions in the presence of catalytic quantities of the isolated stable PdII species studied earlier by HR-MAS NMR was monitored by in situ FTIR spectroscopic analysis (Table 6). The mononuclear and the dinuclear complexes were added in 5 and 2.5 mol % loading, respectively, to give 5 mol % Pd-atom loading per reaction. It should be noted that some thermal decomposition at 56 ± 1 °C was observed by HR-MAS NMR study for every complex except for $[Pd(Ph)(\mu-OAc)(PPh_3)]_2$. The kinetic profiles and the observed rate constants for the reactions catalyzed by the dinuclear Pd complex were compared with those for the catalyst mixture of 5 mol % Pd(OAc)₂ and 10 mol % PPh₂. Of the dinuclear Pd complexes tested, Pd-2-OAc and Pd1-OAc gave rates very close to Pd(OAc)₂/2PPh₃. Pd1-I was significantly faster, comparable to Pd(OAc)₂/3PPh₃. Pd2-OH and Pd2-I were ineffective when used alone but reached comparable rates with added AcOH.

Table 6. Observed Rate Constants for the Direct Arylation Reaction of 1 with 2a Catalyzed by Isolated Pd Species

entry	catalyst (5 mol % Pd-atom)	$k_{\rm obs}/10^{-6}$. mol dm ⁻³ s ⁻¹
1	$Pd(OAc)_2 + 2PPh_3$	9.90 ± 0.04
2	Pd2-OAc	7.90 ± 0.05
3	$Pd2-OAc + 2PPh_3$	7.56 ± 0.06
4	Pd2-OH	
5	Pd2-I	
6	Pd2-OH + 2AcOH	9.71 ± 0.10
7	Pd2-I + 2AcOH	8.32 ± 0.05
8	Pd1-OAc	8.52 ± 0.09
9	Pd1-I	14.83 ± 0.01
10	$Pd(OAc)_2 + 3PPh_3$	14.58 ± 0.01
11	Pd1-I + AcOH	7.34 ± 0.04
12	Pd1-I + 2AcOH	11.10 ± 0.14
13	Pd1-I + 4AcOH	14.00 ± 0.01

The role of Pd-NPs in the model reaction was considered by monitoring the reaction kinetics using preformed DMF-stabilized Pd-NP solution (0.9 mM). The reaction achieved 54% conversion of 1 after 40 h. In comparison, the reaction catalyzed by $Pd(OAc)_2$ achieved quantitative conversion of substrate in 3 h (Figure S36). The result suggested that the two reactions are catalyzed by different active species and that the Pd-NPs were much less active.

Kinetic Isotope Effect. The kinetic isotope effect was investigated by monitoring the reaction under pseudo-zerothorder kinetics with 10-fold excess (i.e., 0.18 M) of deuteropentafluorobenzene (C_6F_5D) 1-d at 56 \pm 1 °C with Pd(OAc)₂/2PPh₃. The IR spectrum of 1-d was significantly different from that of C₆F₅H 1. The characteristic peak of 2a at 1009 cm⁻¹ overlapped with the band of 1-d at 1007 cm⁻¹. However, it was possible to follow the reaction progress by observing the peak of the product 3a at 989 cm $^{-1}$. The $k_{\rm obs}$ values of separate reactions of 1 and 1-d were obtained from the gradient between 20 and 80% conversion yielding $k_{\rm H}/k_{\rm D}$ 4.36 ± 0.06 at 56 ± 1 °C (Figure 9). These measurements were repeated with a higher concentration of 1/1-d at both 56 and 40 °C, revealing a substantial reduction in KIE at higher concentrations of 1/1-d and an increase in KIE on reduction of the temperature (Table 7). The KIE was also determined with Pd/PPh₃ 1:4 as 3.86 ± 0.12 at 56 ± 1 °C with a 10-fold excess

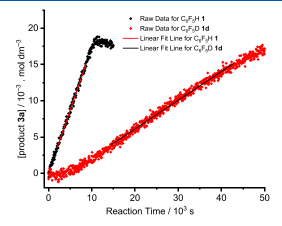


Figure 9. Change in the concentration of product **3a** over time for the reaction of **2a** with 10-fold excess of (black) C_6F_5H **1** and (red) C_6F_5D **1-d.** [Catalyst] 5 mol % wrt **2a**: 0.92 mM, [PPh₃] 1.8 mM, [1] 180 mM, [2a] 18 mM. Ag₂CO₃ 0.75 equiv, temp 56 ± 1 °C.

Table 7. KIE for the Direct Arylation Reaction of 2a by 1 or 1-d at Different Reaction Temperatures and Different Concentrations of 1/1-d^a

T/°C	catalyst	[1] or [1-d]/M	isotope	$k_{\rm obs}/10^{-7}~{ m M~s^{-1}}$	$\mathrm{KIE}^{a} \left(k_{\mathrm{obs(H)}} / k_{\mathrm{obs(D)}} \right)$
40 ± 1	$Pd(OAc)_2/2PPh_3$	0.18	Н	4.19 ± 0.02	5.47 ± 0.05
			D	0.765 ± 0.002	
		0.73	Н	9.50 ± 0.12	2.97 ± 0.05
			D	3.20 ± 0.02	
56 ± 1	$Pd(OAc)_2/2PPh_3$	0.18	Н	17.3 ± 0.2	4.36 ± 0.06
			D	3.96 ± 0.02	
		0.73	Н	29.4 ± 0.3	2.30 ± 0.04
			D	12.8 ± 0.1	
56 ± 1	$Pd(OAc)_2/4PPh_3$	0.18	Н	27.4 ± 0.4	3.86 ± 0.12
			D	7.09 ± 0.11	

 $^{{}^{}a}k_{\text{obs}}$ obtained under pseudo-zeroth-order rate law with [2a] = 18 mM.

of 1/1-d. A control reaction showed that no formation of 1 occurred according to 1H and ^{19}F NMR spectroscopy under the standard reaction conditions when the direct arylation was carried out with 1-d, showing that the C–H bond activation step is irreversible. Exchange does occur, however, with 1 in the presence of D_2O and Ag_2CO_3 and PPh_3 as described above.

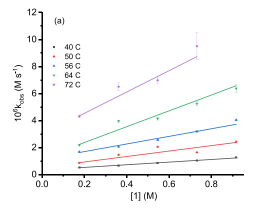
The values of the KIE reported above are derived directly from $k_{\rm obs}$ and therefore represent a composite of the k_1 and k_2 terms. Because the k_1 term is independent of [1], it is expected to have a KIE of 1.0. However, close inspection of the data shows that the k_1 term is much less important for 1-d. The k_2 term is dominant at higher [1], suggesting that the true value of the KIE for this term is ~2.30. The origin of the paradoxical behavior is probed further in the discussion.

Temperature Dependence of Rates. The temperature dependence of the rate of the catalytic reaction between 1 and 2a was studied at varying concentrations of 1 under the same conditions as in Figure 7 over the temperature range 323–345 K (Figure 10a). After conversion of the resulting slopes and intercepts to k_1 and k_2 assuming half-order in [Pd_{tot}], Eyring plots yielded values of ΔH^{\ddagger} and ΔS^{\ddagger} (Figure 10b). The values of ΔH^{\ddagger} for the k_1 and k_2 terms were 57.4 \pm 4.8 and 57.5 \pm 2.6 kJ/mol, and those for ΔS^{\ddagger} were -166 ± 15 and -157 ± 8 J/K mol. These values are the same within error, suggesting that there is a link between them; but at present, we have not identified the cause.

DISCUSSION

The results in this paper show that direct arylation of 4-iodotoluene 2a with pentafluorobenzene 1 is effective using readily available Pd^{II} and Pd^{0} precatalysts $Pd(OAc)_{2}/PPh_{3}$, $Pd(PPh_{3})_{2}Cl_{2}$, and $Pd(PPh_{3})_{4}$. The most effective bases were found to be $Ag_{2}CO_{3}$ and $[Me_{4}N]OAc$ when used in conjunction with polar aprotic solvents such as DMF, DMAc, and NMP. The system with $Pd(OAc)_{2}/PPh_{3}$ and $Ag_{2}CO_{3}$ in DMF was studied in detail.

The first indications of the reactivity of Ag_2CO_3 toward 1 came from H/D exchange experiments. Investigation of the reactions of PPh₃ and pentafluorobenzene with Ag_2CO_3 showed the formation of $Ag(C_6F_5)(PPh_3)_n$ species, but low-temperature $^{31}P\{^1H\}$ NMR revealed the presence of numerous species at equilibrium. Likewise, the previously described silver carbonate $[Ag(PPh_3)_2]_2CO_3\cdot 2H_2O$ proved highly labile and reactive toward 1 even at room temperature. $[Ag(PPh_3)_2(\kappa^2-HCO_3)]_2$ was also reactive if Cs_2CO_3 was added. Related species could be obtained by reaction of $Ag(C_6F_5)$ with PPh₃.



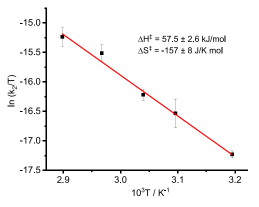


Figure 10. (a). Plot of $k_{\rm obs}$ against [1] for the catalytic reaction of 1 with ${\bf 2a}$ catalyzed by ${\rm Pd}({\rm OAc})_2$ with 2 equiv PPh₃ at ${\bf 40\pm 1}$, ${\bf 50\pm 1}$, ${\bf 56\pm 1}$, ${\bf 64\pm 1}$, and ${\bf 72\pm 1}$ °C with lines of best fit. (b) Eyring plot for values of k_2 derived assuming the reaction is half order in [Pd_{tot}/2PPh₃].

Considering the reactivity of the silver compounds toward 1, we tested whether the standard 0.75 equiv Ag_2CO_3 used in the catalytic reaction of 1 with 2a could be reduced to a catalytic amount. Indeed, catalytic quantities of any of Ag_2CO_3 , $[Ag(PPh_3)_2]_2CO_3\cdot 2H_2O$, or $[Ag(PPh_3)_2(\kappa^2-HCO_3)]_2$ were effective if used in conjunction with Cs_2CO_3 as substoichiometric base. These experiments indicate that the Ag_2CO_3/PPh_3 system is competent to activate the C-H bond of 1 in a similar way to $Ag_2CO_3/XPhos^{45}$ but the PPh_3 system is complicated by the extreme lability of the $Ag-PPh_3$

bonds. We therefore have to consider the possibility that C–H activation by Ag^I competes with C–H activation by Pd^{II}.

Studies of the reactivity of 1 with iodobenzene 2b in the presence of a variety of Pd complexes and Ag_2CO_3 by HR-MAS NMR led to the identification of $[Pd(C_6H_5)(\mu-OAc)(PPh_3)]_2$ Pd2-OAc and $Pd(C_6H_5)(\kappa^1-OAc)(PPh_3)_2$ Pd1-OAc under high $Pd(OAc)_2$ concentrations. The stoichiometric reaction of the isolated Pd species Pd2-OAc with 1 at 70 °C yielded the expected coupling product 3b quantitatively, whereas the reaction of Pd1-OAc with 1 achieved 45% yield without Ag_2CO_3 and 100% with Ag_2CO_3 . These investigations lead to the postulate of Pd1-OAc and Pd2-OAc as likely resting states in the catalytic reaction. Other species such as Pd1-I and Pd2-I are also active but less effective than the acetate complexes. Evidently, multiple Pd species are present, and the speciation will vary with PPh3 concentration and temperature.

We now consider the interpretation of the kinetic results. The k_2 term of the rate law (eq 2) shows that the rate depends on $[C_6F_5H]$ and $[Pd_{tot}/2PPh_3]^{0.5}$, a similar rate expression to that described by Rosner et al. This expression is consistent with a catalytically active mononuclear Pd⁰ species that reacts rapidly with ArI to form Pd¹¹ species present in monomeric and dimeric forms. The monomer is active in the catalytic cycle, but the dimeric species with which it is at equilibrium is inactive. The concentration of the monomer is given by $K_{\rm eq}[{
m dimer}]^{0.5}$, leading to the half-order dependence. The monomer proceeds to react with C₆F₅H and base before reductive elimination of product ensues. The small value of ΔH^{\ddagger} and the large negative ΔS^{\ddagger} are consistent with a bimolecular reaction in the rate determining transition state. 92 This catalytic cycle is similar to that shown in Scheme 1a with the addition of the monomer-dimer equilibrium. It is tempting to identify the monomer and dimer as Pd1-OAc and Pd2-OAc (or their tolyl equivalents) that were identified as resting states through the HR-MAS NMR experiments. The value of k_2 is little changed when using a 4:1 ratio of [PPh₃]/ [Pd(OAc)₂] in place of a 2:1 ratio, indicating that the Pd^{II} speciation is little affected by this change. The significant KIE (~2.30 taken from the higher [1] at 56 °C) and substantial entropy of activation are consistent with this model. There are several indications that this model of the k_2 reaction may be oversimplified and that multiple species contribute. First, the experiments with carbonate complexes of silver show that multiple complexes of AgI with PPh3 and carbonate are present and could be active as bases; second, the HR-MAS NMR experiments show the presence and activity of several PdII species. The variation of the KIE with reaction conditions is a likely consequence of the complex speciation.

The k_1 term of the rate law shows dependence on $[Pd_{tot}/2PPh_3]^{0.5}$ and is increased by a factor of ~2.5 with a 4:1 ratio of PPh₃ to Pd_{tot}. The absence of a dependence on $[C_6F_5H]$ is consistent with C–H activation of 1 by Ag^I(PPh₃)_n followed by transmetalation to any of Pd1-OAc, Pd2-OAc, Pd1-I, or Pd2-I. The dependence on $[Pd_{tot}/2PPh_3]^{0.5}$ indicates that the monomer–dimer equilibrium remains important and that the rate-determining transition state is likely to be associated with the transmetalation and/or reductive elimination. Considering the negative value of ΔS^{\ddagger} and small value of ΔH^{\ddagger} , transmetalation is more likely. Direct participation of the solvent could also be possible. The dependence on $[PPh_3]$ is consistent with the requirement for sufficient ligand to solubilize the Ag₂CO₃ and activate 1; the concentrations of

PPh3 are suboptimal at the lower ratio because PPh3 is consumed both by reduction of Pd^{II} to Pd⁰ with formation of OPPh₃ and by coordination to Ag^I. To understand the variation of KIE with conditions, we recall that the KIE for the stoichiometric reaction of $[Ag(Xphos)]_2(\mu-\kappa,^2\kappa^2-CO_3)$ with pentafluorobenzene is 3.7 ± 0.3 , a value that we may assume is close to that for PPh3 analogues. This value would mean that the k_1 term is slowed more than the k_2 term by deuteration, and we postulate that the AgI route to C-D activation is uncompetitive with the PdII route. Unfortunately, we cannot monitor the rate of C-H/C-D activation by Ag(PPh₃)_n species because of their lability and the consequent multiple speciation. The similarity of the rate constants for different catalysts Pd(OAc)₂/nPPh₃, Pd(PPh₃)₄, Pd-2OAc, Pd1-OAc, and Pd1-I points to common mechanisms. We anticipate that the catalytic mechanism involving bond activation by Ag¹ is similar to that shown in Scheme 1b with the proviso of multiple speciation of both silver and palladium complexes that vary with conditions.

CONCLUSIONS

The direct arylation of iodoarenes by pentafluorobenzene 1 is a prototype reaction that is catalyzed by palladium complexes in the presence of triphenylphosphine and silver carbonate. The reaction is typically performed with 0.75 equiv Ag₂CO₃, but the use of Ag₂CO₃ may be reduced to catalytic quantities in the presence of Cs₂CO₃. The simplicity of the components does not mean, however, that the reaction mechanism is straightforward. Our study has demonstrated a Pandora's flask of multiple species that can only be interpreted through previous research on the reactivity of analogous systems using phosphines such as XPhos. Both in situ IR spectroscopy and high-resolution magic angle spinning NMR spectroscopy proved useful in our analysis, in addition to solution NMR methods. We have shown that the reaction of Ag₂CO₃ with PPh₃ and 1 leads to C-H bond activation with multiple Ag complexes present in solution. The same applies when starting with isolated complexes such as [Ag(PPh₃)₂]₂CO₃·2H₂O. The potential for C-H activation by AgI aligns with our own experiments using XPhos in place of PPh3 and the reactions investigated notably by Larrosa, Sanford, and Hartwig with their co-workers.^{34–43} The problem of multiple speciation also applies to the palladium complexes, although it is not so acute. Our HR-MAS experiments point to $[Pd(Ar)(\mu-OAc)(PPh_3)]_2$ and $Pd(Ar)(\kappa^{1}-OAc)(PPh_{3})_{2}$ (Ar = Ph, 4-tolyl) as likely resting states that are at equilibrium, with the dimer as the major species. The reaction kinetics studied by in situ IR spectroscopy demonstrates that synthetic experiments may be performed with a 1:1 ratio of [ArI]/[C₆F₅H] under milder conditions (56 °C, 5 h) than are typically used (70 °C, 24 h). Rates are improved with Pd(OAc)₂ + 4PPh₃ or Pd(PPh₃)₄ compared with the standard Pd(OAc)₂ + 2PPh₃ in keeping with the multiple roles of PPh3 for coordination of both Ag and Pd, as well as reduction of Pd(OAc)₂. Higher [PPh₃] also favors monomeric Pd complexes over dimeric complexes. Similar rate constants are found with $[Pd(Ph)(\mu-OAc) (PPh_3)$ ₂ and $Pd(Ph)(\kappa^1-OAc)(PPh_3)_2$ as catalysts. A twoterm rate law is found with both terms dependent on Pdtot/ 2PPh₃]^{0.5}, consistent with an off-cycle Pd dimer dissociating to form an on-cycle Pd monomer. The k_1 term is independent of [1], whereas the k_2 term is first order in [1]. Neither term shows dependence on [ArI]. We interpret the k_1 term as arising from C-H bond activation by Ag¹, whereas the k_2 term

involves C–H activation by Pd^{II}. Both processes require base-assistance and may proceed by AMLA/CMD mechanisms. The multiple speciation of both Ag^I(PPh₃) and Pd^{II}(PPh₃) complexes leads to a situation where it is no longer appropriate to define a mechanism in full. In particular, the implications of the extraordinary lability of Ag^I(PPh₃) species have not been recognized previously. Thus, a reaction with apparently well-behaved kinetics disguises extraordinary complexity. It may be possible to suppress C–H bond activation by one of Pd^{II} or Ag^I by adding appropriate inhibitors.⁴⁹

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.organomet.3c00309.

General information on chemicals and instruments, general procedures, synthetic procedures and compound data, studies on conversion, speciation of silver complexes, analysis of catalytic intermediates by HR-MAS NMR spectroscopy, and reaction kinetics by *in situ* IR spectroscopy (PDF)

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Notes

The authors declare no competing financial interest.

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