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The ubiquitous cross-coupling catalyst system 'Pd(OAc)₂'/2PPh₃ forms a unique dinuclear Pd^I complex: an important entry point into catalytically competent cyclic Pd₃ clusters†

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Palladium(II) acetate 'Pd(OAc) $_2$ '/nPPh $_3$ is a ubiquitous precatalyst system for cross-coupling reactions. It is widely accepted that reduction of in situ generated trans-[Pd(OAc)₂(PPh₃)₂] affords [Pd⁰(PPh₃)_n] and/or $[Pd^{0}(PPh_{3})_{2}(OAc)]^{-}$ species which undergo oxidative addition reactions with organohalides – the first committed step in cross-coupling catalytic cycles. In this paper we report for the first time that reaction of $Pd_3(OAc)_6$ with 6 equivalents of PPh_3 (i.e. a Pd/PPh_3 ratio of 1:2) affords a novel dinuclear Pd^1 complex [Pd₂(µ-PPh₂)(µ₂-OAc)(PPh₃)₂] as the major product, the elusive species resisting characterization until now. While unstable, the dinuclear Pd^I complex reacts with CH₂Cl₂, p-fluoroiodobenzene or 2bromopyridine to afford Pd3 cluster complexes containing bridging halide ligands, i.e. [Pd₃(X)(PPh₂)₂(PPh₃)₃]X, carrying an overall 4/3 oxidation state (at Pd). Use of 2-bromopyridine was critical in understanding that a putative 14-electron mononuclear 'Pd^{II}(R)(X)(PPh₃)' is released on forming $[Pd_3(X)(PPh_2)_2(PPh_3)_3]X$ clusters from $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$. Altering the Pd/PPh₃ ratio to 1:4 forms Pd⁰(PPh₃)₃ quantitatively. In an exemplar Suzuki-Miyaura cross-coupling reaction, the importance of the 'Pd(OAc)₂'/nPPh₃ ratio is demonstrated; catalytic efficacy is significantly enhanced when n=2. Employing 'Pd(OAc)₂'/PPh₃ in a 1:2 ratio leads to the generation of $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$ which upon reaction with organohalides (i.e. substrate) forms a reactive Pd3 cluster species. These higher nuclearity species are the cross-coupling catalyst species, when employing a 'Pd(OAc)2'/PPh3 of 1:2, for which there are profound implications for understanding downstream product selectivities and chemo-, regio- and stereoselectivities, particularly when employing PPh₃ as the ligand.

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Introduction

Palladium(II) acetate 'Pd(OAc)₂' is commonly used in combination with tertiary phosphine ligands, *e.g.* PPh₃, to generate active catalyst species for an eclectic array of cross-coupling reactions, where it is universally accepted that 'Pd⁰(PPh₃)_n' species are formed.² Such species enter into oxidative reactions with organohalides, *e.g.* iodobenzene to generate *trans*-[Pd(I)(Ph)(PPh₃)₂].³ Considerable and notable efforts have been made by Amatore and Jutand⁴ to understand how varying the 'Pd(OAc)₂'/nPPh₃

ratio affects the generation of reduced palladium species in both THF and DMF. Following extensive NMR spectroscopic and electrochemical measurements, conclusions were drawn implicating phosphine-induced reduction of 'Pd(OAc)2'/nPPh3 mixtures, via trans-[Pd(OAc)2(PPh3)2], by an intramolecular process (independent of phosphine concentration, once the latter complex is formed).4 The global findings from Amatore and Jutand are detailed in Scheme 1, showing the key intermediate species observed by ³¹P NMR spectroscopic studies. Comparisons of these data were made with complexes generated from [Pd(PPh₃)₄] in the presence of n-Bu₄NOAc, under electrochemical conditions. The conclusions were that 'Pd⁰(PPh₃)_n' species are generated in situ from the reaction of Pd(OAc)₂'/2PPh₃ mixtures.4a Later studies showed that increasing the Pd/PPh3 ratio to 1:3 and above led to the clean generation of $[Pd^{0}(PPh_{3})_{n}(OAc)]^{-}$ species (n = 2 or 3), with $O = PPh_{3}$ being a key side product, *i.e.* formed during the formal $Pd^{II} \rightarrow Pd^{0}$ reduction process. 4b,c Both 'Pd⁰(PPh₃)₂' and [Pd⁰(PPh₃)₂(OAc)] species react by oxidative addition with organohalides.

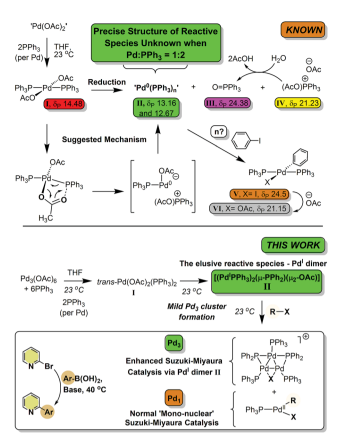
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[†] Electronic supplementary information (ESI) available: Full experimental details and characterization data for all compounds is provided, including NMR spectra, X-ray diffraction data and computational data (as a PDF file). CCDC 1894927–1894931 and 1901195. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9sc01847f

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Scheme 1 Reactions of 'Pd(OAc)2' with PPh3 (1:2 ratio). 31P NMR spectral data are taken from ref. 4a

Later, Kollár et al. examined the reaction of 'Pd(OAc)2'/nPPh3 in DMF,5 amongst other phosphines, concluding that 'Pd⁰(PPh₃)_n' species are formed under ambient reaction conditions. Taken together these studies suggest that a Pd/PPh3 ratio of 1:3 is necessary for satisfactory catalytic cross-coupling performance.

Over the last 20 years we have regularly debated the differences in cross-coupling catalyst system performance on changing the Pd/PPh₃ ratio from 1:2 to 1:3.6 When papers are reported employing a Pd/PPh3 ratio of 1:2 we have asked the why, as 1:3 would be ideal based on the outcomes of previous studies;4 in other words, optimal conditions for forming catalytically active $[Pd^{0}(PPh_{3})_{n}(OAc)]^{-}$ species requires ≥ 3 equivalents of PPh3 per Pd, "not 2 equivalents", when 'Pd(OAc)2' is used as the initial Pd^{II} precatalyst.

A superb recent example is found in the high-throughput automated reaction screening study conducted by a team from Pfizer, where a Pd/PPh₃ ratio of 1:2 was used for 480 Suzuki-Miyaura cross-coupling (SMCC) reactions, involving changes in solvent and base, against relatively minor changes in substrate structure, correlated alongside many other phosphine ligands (over 5760 reactions in total). Cronin et al. further applied a machine learning algorithm based on the product percentage yields.8 With such important developments being made in automation, reaction optimization and machine learning,9 knowing precisely the reactive Pd species,

formed under working reaction conditions, has never been more important. Thus, herein we report that the reaction of Pd₃(OAc)₆ with 6 equivalents of PPh₃ (Pd: PPh₃, 1:2), in both THF and DMF, generates a well-defined [Pd₂(μ-PPh₂)(μ₂- $OAc)(PPh_3)_2$ complex II. The formation of this unusual species adds to the mechanistic debate concerning the activation pathways for Pd(II) precatalysts, particularly papers reported by: (a) Balcells and Hazari^{10a} showing formation of Pd^I dimers with NHC ligands and bridging allyl and chloride ligands (eqn (1)); (b) Colacot and Schoenebeck showing formation of Pd^I dimers with phosphines and bridging bromide ligands (eqn (2)); (c) Bedford10c showing SPhos activation on reaction with Pd(OAc)2 (eqn (3)); (d) Jutand and Grimaud 10d showing XPhos reactions with Pd(OAc)2 leading to a proposed Pd^I dimer (egn (4)).

The stability and reactivity of these Pd^I dimers appear to be critical in understanding the delivery of active 'L-Pd⁰, species, a process dependent on L/Pd ratios and additives. From our study we find that $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$ II, exhibits unique reactivity toward organohalides, e.g. CH₂Cl₂ 1a, p-fluoroiodobenzene 1b and 2-bromopyridine 1c, which affords Pd3 cluster species, namely [Pd₃(X)(µ-PPh₂)₂(PPh₃)₃]X VII (later referred to as $Pd_3X \cdot X$, where X = Cl, Br, I or OAc). Our results naturally connect to a recent report showing that Pd₃Cl·Cl is a highly active catalyst for SMCC reactions, including the activation of substrates containing harder to activate C-Cl bonds.11 Pd₃Cl·Cl invokes an unusual switch in cross-coupling steps from oxidative addition then transmetallation to transmetallation and then oxidative addition.11

Balcells, Hazari et al.

Pd
0
—IPr

IPr = IPr = 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene)

Colacot, Schoenebeck et al.

(cod)PdBr₂ + P(t-Bu)₃
$$\longrightarrow$$
 (t-Bu)₃P-Pd $\stackrel{\text{Br}}{\underset{\text{Dr}}{\triangleright}}$ Pd-P(t-Bu)₃ \longrightarrow Pd⁰-P(t-Bu)₃ (2)

Jutand, Grimaud et al.

$$Pd^{II}(OAc)_{2}(XPhos) + Pd^{0}(XPhos) \longrightarrow [Pd^{I}(OAc)(XPhos)]_{2}$$

$$Structure \ not \ defined$$

$$XPhos \ iPr$$

$$XPhos \ iPr$$

$$(4)$$

Results and discussion

The reaction of ultra-pure $Pd_3(OAc)_6$ (>99% purity) with PPh₃, in varying ratios, in THF-d₈ were conducted at room temperature (25 °C) and monitored by ¹H and ³¹P NMR spectroscopic analysis (where [Pd] = 20 mM; T = 298 K, external reference = 85% H₃PO₄ in H₂O). A wide spectral window (–50 to +250 ppm) was required to allow full characterization of the array of phosphorus signals and associated species formed under these reaction conditions (Fig. 1 and 2).

Where the ratio of $Pd: PPh_3$ was 1:1, degradation of $Pd_3(OAc)_6$, leading to the formation of large perfectly spherical Pd particles (sized $\sim 0.1-0.4~\mu m$, by TEM) and many P-containing species (by ^{31}P NMR) was observed (Fig. 1(a)). Alteration of the Pd/PPh₃ ratio to 1:2 (Fig. 1(b)) led to the

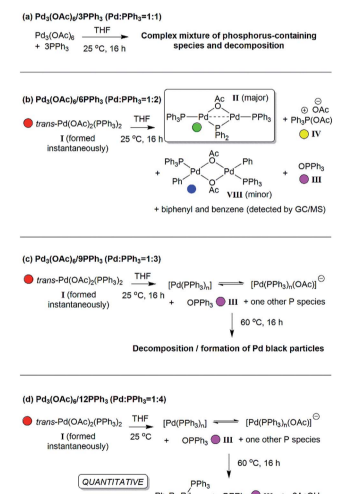


Fig. 1 The room temperature formation of dinuclear Pd^{II} complexes from trans- $Pd(OAc)_2(PPh_3)_2$ in THF. (a) Ratio of $Pd: PPh_3 = 1:1$; (b) ratio of $Pd: PPh_3 = 1:2$; (c) ratio of $Pd: PPh_3 = 1:3$; (d) ratio of $Pd: PPh_3 = 1:4$. The Pd^0 species $Pd^0(PPh_3)_3$ IX and $[Pd^0(PPh_3)_3OAc]^-$ IX' are indicated by cyan circles (appearing as coincident signals by ^{31}P NMR spectroscopic analysis when present together – compare top two ^{31}P NMR spectra in Fig. 2 with the authentic sample of $Pd^0(PPh_3)_3$ IX, bottom spectrum, Fig. 2).

formation of a major new phosphorus-containing species at δ 199.01 (t, 1P) and δ 13.41 (d, 2P), with a $^2J_{PP}$ coupling constant of 83.5 Hz (i.e. an AX₂ type spin system). The high ³¹P chemical shift of δ 199.01 indicates that the PPh₃ ligand has been activated by P-C bond-cleavage to give a bridging phosphido-group at Pd, with concomitant loss of 'C₆H₅'. The ¹H NMR spectrum shows a methyl resonance at δ 2.08 (s, 3H), due to a bridging acetoxy ligand, which balances with aromatic proton integrals (40H). Running the reaction at lower Pd₃(OAc)₆ concentration (between 3 and 20 mM) allowed this species to be isolated in a form that could be crystallized. X-ray diffraction analysis of dark red single crystals of this species confirmed its structure as $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$ II, possessing both bridging μ_2 acetoxy and µ-phosphido ligands and terminal-capping PPh3 ligands. Complex II is a diamagnetic species. The Pd-Pd bond distance was found to be 2.5958(3) Å, which is in-keeping with other dinuclear Pd^I complexes with bridging μ -acetoxy ligands known in the literature (typical Pd-Pd distances 2.532 to 2.711 Å), and shorter than a related structure, $[Pd_2(\eta^3-allyl)(\mu-OC(O)i-$ Bu)(PPh₃)₂] where the Pd···Pd bond distance equals 2.6267(3).¹²

A scaled-up synthesis of II was found possible from Pd₃(OAc)₆/6PPh₃, formed in 31% yield (note: some Pd is lost as large particles during its preparation), which was fully characterized. Interestingly, the LIFDI-MS data showed that the dinuclear Pd^I complex was present in solution ($M^{+*} = m/z$ 982, with the correct isotopic distribution). The reference ³¹P NMR spectrum for purified II is given in Fig. 2 (externally-referenced to H₃PO₄). GC-MS analysis of the crude reaction mixture containing II indicated that benzene and biphenyl were present, the former most likely derived from protonation of 'Pd-Ph' species by adventitious water/AcOH and the latter by reductive elimination. Acetoxybenzene, a possible reductive elimination product, was not detected by GC-MS analysis. These species are accompanied by O=PPh3 III, [Ph3P(OAc)]X IV and another dinuclear PdII species VIII, the latter only in minor amounts. Complex VIII was previously reported as a major product of a reaction of 'Pd(OAc)₂' with 2 equivalents of PPh₃ on heating in methanol (41% yield).13 It is worth noting that complex II is stable in dry THF solutions over 12 hours, which allows for its spectroscopic characterization, but decomposition is seen after ca. 5 days at 22-25 $^{\circ}$ C.

On changing the Pd/PPh3 ratio to 1:3 complex II was not formed, simply a broad resonance at δ 5.71 (FWHM *ca.* 550 Hz) characterized as $Pd^{0}(PPh_{3})_{n}/[Pd^{0}(PPh_{3})_{n}(OAc)]P(OAc)Ph_{3}$ (n = 1,2 or 3), see Fig. 1(c). The chemical shift alters with time, with concomitant formation of O=PPh3, by hydrolysis of [Ph3-P(OAc) X IV, yielding AcOH also. Heating this mixture to 60 °C, over 16 h, eventually ended in decomposition to form large Pd black particles. Indeed, similar 31P NMR spectra were seen on changing the Pd/PPh3 ratio to 1:4, see Fig. 1(d), leading to a mixture of $Pd^{0}(PPh_{3})_{3}$ **IX** and $[Pd^{0}(OAc)(PPh_{3})_{3}]^{-}$ **IX**'. At the same Pd/PPh3 ratio, subsequent heating to 60 °C resulted in clean conversion of II into Pd⁰(PPh₃)₃ (IX), O=PPh₃ (III) and 2AcOH, quantitatively, as shown by both ³¹P and ¹H NMR spectra. Layering this solution with hexane, after t = 16 h, led to the formation of yellow-orange crystals, which were found suitable for X-ray diffraction, establishing the compound as

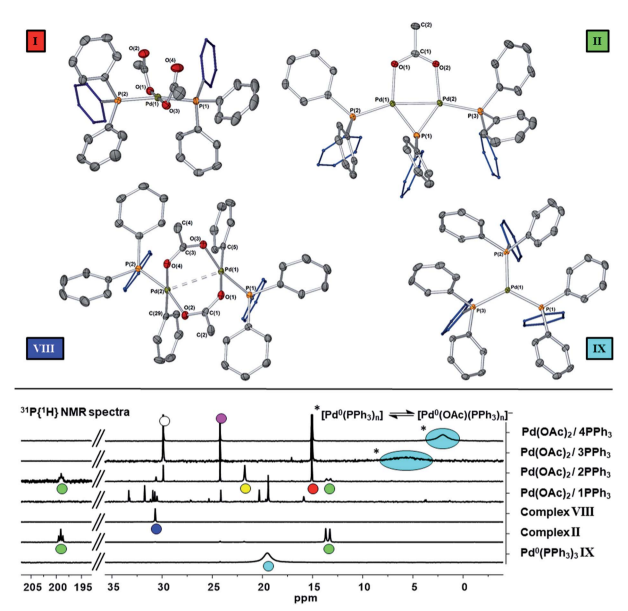


Fig. 2 Top: Single crystal X-ray diffraction structures of II, VIII and IX are shown (thermal ellipsoids shown at 50%, H-atoms and solvent of crystallization not shown, for clarity). Bottom: ${}^{31}P\{^{1}H\}$ NMR (202 MHz) spectra of mixtures of 'Pd(OAc)₂' with nPPh₃ (n=1 to 4) in THF at 23 °C for 16 h, showing differences in phosphorus speciation. Reference spectra are given for II (green), VIII (blue) and IX (cyan); other species are OPPh₃ (pink), [AcOPPh₃]X (yellow, where X is likely the OAc anion) and trans-Pd(OAc)₂(PPh₃)₂ I (red). Several phosphorus species are uncharacterized for the Pd(OAc)₂/1PPh₃ experiment (also resulting in PdNP formation). For the equilibrium shown against the ratio of Pd(OAc)₂/4PPh₃ spectral data, acetate anion and free PPh₃ are involved, explaining the substantially lower chemical shift (compare also the reference spectrum of pure Pd⁰(PPh₃)₃ IX – bottom).

 $Pd^{0}(PPh_{3})_{3}$ (**IX**) (Fig. 2). It is worthy of note that $Pd^{0}(PPh_{3})_{3}$ **IX** is a relatively stable Pd^{0} complex in the solid-state (note: discoloration is noted in air after ~ 1 day).

Computational studies using DFT calculations with $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$ II at the B3LYP/DEF2SVP-D3 level of theory. The calculations reveal a short Pd-Pd bond (2.58 Å), supporting its diamagnetic properties. The HOMO resides primarily on the Pd-Pd centers, whereas the LUMO can be found over the phosphide and Pd-Pd centers (Fig. 3). The HOMO/LUMO provide potential clues about the underlying reactivity of $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$ toward other species such as electrophiles and nucleophiles.

We believe that the mechanism for formation of ${\bf II}$ is different to the Pd^I dimer stabilised by a bridging arene, as reported by Bedford. In the latter case a sequential reaction in methanol was used, followed by treatment with a non-coordinating anion leaves a suitably-disposed arene to stabilise the cationic Pd^I dimer species, though $Pd-\pi$ -arene interactions. In ${\bf II}$ acetate takes on that role.

Reactivity of II towards organohalides

The reaction of $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$ II with CH_2Cl_2 1a (~10-fold excess) occurred at room temperature to afford a new

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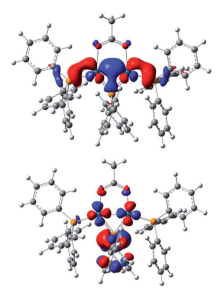


Fig. 3 The HOMO (left) and LUMO (right) for $[Pd_2(\mu-PPh_2)(\mu_2-\mu_2)]$ OAc)(PPh₃)₂] II, computed by density functional theory (b3lyp/def2svp functional/basis set used for optimization, single point orbital and total energies; CPCM implicit solvent (tetrahydrofuran) and Gimme's D3 empirical correction used).

Pd species, which was identified as [Pd₃(Cl)(PPh₂)₂(PPh₃)₃]OAc $Pd_3Cl \cdot OAc$ by ESI studies (+ve mode, detected by the $[M - OAc]^+$ ion) (Fig. 4).14 Real-time reaction monitoring by 31P NMR spectroscopic analysis showed that Pd3Cl·OAc formed over several hours at the expense of II. The data for Pd3Cl·OAc closely matches the data obtained from the independent synthesis of [Pd₃(Cl)(μ-PPh₂)₂(PPh₃)₃]Cl **Pd₃Cl·Cl**, starting from PdCl₂(PPh₃)₂ in aniline under H_{2(g)} at 90 °C, for which an X-ray structure of a single crystal was determined (Fig. 5).15 We were unable to trace the 'CH2Cl' fragment derived from CH2Cl2, 'Pd(PPh₃)₁' and acetate anion required to balance the overall chemical reaction. However, balance of overall charge and mass allows one to postulate a putative 14-electronic Pd^{II} species Xa. Similarly, reaction of II with p-fluoro-iodobenzene 1b afforded $[Pd_3(I)(\mu-PPh_2)_2(PPh_3)_4]OAc$ $Pd_3I\cdot OAc$, as shown by ³¹P NMR and ESI data (as the $[M - OAc]^+$ ion), which degraded rapidly to form Pd black. As with the reaction of II with CH₂Cl₂ 1a, the 'p-F-C₆H₄-' fragment derived from **1b**, 'Pd(PPh₃)₁' and OAc anion could not be fully traced (PdII species Xb is postulated).

To reveal whether putative 14-electron PdII species were formed in the reactions of II with organohalides (R-X) we hypothesized that a reaction with 2-bromopyridine 1c would enable characterization by the stabilization conferred by Ncoordination from the pyridine ring, leading to formation of a stable 16-electron dinuclear Pd^{II} species (i.e. 4a-c, Fig. 4).

To verify findings concerning formation of [Pd₃(Br)(μ- $PPh_2_2(PPh_3)_3$ OAc ($Pd_3Br \cdot OAc$) vide supra, a closely related sample was prepared by treatment of [Pd₃(Cl)(μ-PPh₂)₂(PPh₃)₃] Cl (Pd₃Cl·Cl) with excess KBr in CH₂Cl₂, giving [Pd₃(Br)(μ-PPh₂)₂(PPh₃)₃]Br (Pd₃Br·Br).¹⁶ The latter material possessed identical ³¹P NMR and MS data to that seen for Pd₃Br·OAc from the reaction of II with 2-bromopyridine 1c. A reasonable single crystal X-ray diffraction structure for Pd3Br·Br was further determined (Fig. 5). Whilst a detailed comparison between $Pd_3Cl \cdot Cl$ and $Pd_3Br \cdot Br$ cannot be made (R_1 factors for $Pd_3Cl \cdot Cl$ = 3.58% and $Pd_3Br \cdot Br = 7.04\%$), there are striking structural differences that necessitate additional comment. The cyclic 6membered 'Pd-P-Pd-Cl-Pd-P' fragment is essentially flat in Pd₃Cl·Cl, leaving the second chloride anion as noncoordinating. However, in Pd₃Br·Br we see something quite different - the cyclic 6-membered 'Pd-P-Pd-Br-Pd-P' fragment is highly twisted, which is associated with an interacting second bromide anion.

It is tempting to draw an analogy here to a bromonium ion interacting with a bromide anion (i.e. reactions of alkenes with bromide proceeding via bromonium ion intermediates). The structural differences between Pd₃Cl·Cl and Pd₃Br·Br suggest that they could be distinctly different in how they operate in catalysis.

The relevance of our findings concerning reaction of II with organohalides requires contextualisation with the results recently reported by Schoenebeck and co-workers.17 It has been shown that $[Pd(\mu-I)P(t-Bu)_3]_2$ reacts with PHPh₂ (slight excess relative to the Pd^I dimer) in toluene at room temperature to give a Pd₃ cluster containing three bridging phosphide ligands (Fig. 6). Subsequent reaction with an aryl halide then delivers a Pd₃-type cluster containing a bridging iodide ligand, similar to the $Pd_3X \cdot X$ clusters vide supra. The pathways to these Pd_3 clusters are not the same. Complex II reacts directly with organohalides to give $Pd_3X \cdot X$ clusters (where X = Cl, Br or I), i.e. additional phosphine is not necessary at this point. Indeed, if additional PPh₃ (2 equiv.) is reacted with II (1 equiv.) in THF at room temperature we see the generation of $Pd(PPh_3)_n$ species (where n = 3, this species was detected by LIFDI-MS, see ESI†). This finding is in-keeping with what was observed when Pd₃(OAc)₆ was reacted with nine equivalents of PPh₃ (i.e. Pd/ PPh_3 ratio of 1:3, Fig. 2). We expect that Pd^0 complexes are generated from disproportionation of the Pd^I dinuclear complex II, upon addition of PR3, akin to the observations reported by Schoenebeck and Colacot. 10b

Importance of our findings in an exemplar SMCC reaction

To better understand the importance of the 'Pd(OAc)2'/nPPh3 ratio in catalysis, the cross-coupling of 2-bromopyridine 1c with p-fluorophenylboronic acid 2 to give 2-arylpyridine 3 was examined,6b using 1 M n-Bu4NOH as the base, in a THF/water mixture (1:1, v/v) at 40 °C. We carefully selected 1 M n-Bu₄NOH as the base, drawing on the recent findings concerning the importance of both the hydroxide anion and cation-type in SMCCs.¹⁸ Also pertinent to mention is that our SMCC reaction is homogeneous, i.e. not biphasic, simplifying the discussion concerning which phase the Pd catalyst and organoboron species reside in. Furthermore, operationally NMR spectroscopic analysis in operando was made feasible by use of aqueous n-Bu₄NOH in THF.

SMCC reactions of $1c + 2 \rightarrow 3$ were monitored in operando by ¹H NMR spectroscopic analysis, allowing pre-stirred mixtures of $Pd_3(OAc)_6/nPPh_3$ (n = 6 and 12, i.e. $Pd/PPh_3 = 1:2$ or 1:4

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2x Pd₃CI.OAc CIHC Exact Mass: 1509.06 1503 1506 1509 1512 1515 1518 1521 THF. 23 °C Ph₂ Xa (not detected) OAc OAc PPh₃ Pd₃I.OAc Pd₃Br.OAc Exact Mass: 1553.01 1551 1554 1557 1560 1563 (detected as 4a-c) Xc 31P{1H} NMR Xb (not detected) br s 210 200 40 30 20 10 ppm 1596 1598 1600 1602 1604 1606 1608 1610 Detected and characterized X & X' = Br (4a)X & X' = OAc (4b)X = Br, X' = OAc (4c)

Fig. 4 Reactions of dinuclear PdI complex II with organohalides (1a-c), leading to formation of Pd₃ clusters $Pd_3X \cdot OAc$. The X anions in these clusters are likely acetate (mass balance is formally correct using acetate for all Pd species formed), although mixtures of different anions cannot be ruled out for species generated *in situ*. The MS ions are all observed by ESI (+ve mode) as molecular cations, the data for which is presented (measured – in blue; simulated – in red). The $^{31}P^{1}H^{1}$ NMR spectrum for species generated from the reaction of 2-bromopyridine ^{1}C with II illustrates the formation of $Pd_3Br \cdot OAc$ and species $^{4}a-c$ (note a cut//in the ^{31}P NMR spectrum is made between 40 and 190 ppm, due to the wide spectral range, for ease of viewing – full ^{31}P NMR spectra are shown in the ESI†).

stable 16-electron PdII dinuclear complex

respectively) to be compared in THF against a reaction mediated by $[Pd_2(\mu\text{-PPh}_2)(\mu_2\text{-OAc})(PPh_3)_2]$ II. The kinetic profiles for the appearance of 3, with concomitant disappearance of 1c

Xc, putative 14-electron Pd^{II}

(pseudo-zero order in 1c), are shown in Fig. 6. The kinetic profile for the reaction mediated by $Pd_3(OAc)_6/6PPh_3$ (1 : 2, Pd/PPh_3) indicates that the reaction is efficient at 40 °C {Fig. 7(A)} –

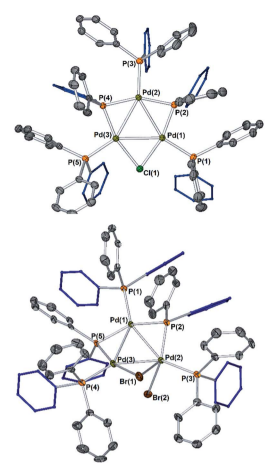


Fig. 5 The single crystal X-ray diffraction structures for [Pd₃(- $X)(PPh_2)_2(PPh_3)_3]X$ (top, X = Cl, $Pd_3Cl \cdot Cl$; bottom, X = Br, $Pd_3Br \cdot Br$); Hatoms and solvent of crystallization are not shown and thermal ellipsoids are set at 50% probability. For Pd₃Cl·Cl the non-coordinating chloride anion is not shown. Selected bond lengths (Å) and angles (°) for: $Pd_3Cl \cdot Cl \ Cl1 - Pd1 = 2.3828(8)$; Cl1 - Pd3 = 2.4002(8); Pd1 - Pd2 =2.9138(3); Pd1-Pd3 = 2.8882(3); Pd2-Pd3 = 2.9127(3); Pd1-Cl1-Pd3= 74.29(2); Pd1-P2-Pd2 = 81.38(3); Pd3-P4-Pd2 = 81.45(3); Cl1-Pd1-Pd2 = 113.303(19); Cl1-Pd1-Pd3 = 53.130(18); Pd3-Pd1-Pd2 = 60.265(8). Pd₃Br·Br Pd1-Pd2 = 2.8355(10); Pd2-Br1 = 2.9423(13); Pd2-Br2 = 2.5698(13); Pd3-Br1 = 2.5490(13); Pd1-Pd3 = 2.8808(10); Pd2-Pd3 = 2.8240(10); Pd2-Pd1-Pd3 = 59.21(2), Pd3-Pd2-Pd1 = 9.8240(10)61.20(2), Pd2-Pd3-Pd1 = 59.60(2); Pd3-Br1-Pd2 = 61.40(3); Br2-Pd2-Pd1 = 101.20(4); Br2-Pd2-Pd3 = 76.91(3); Br2-Pd2-Br1 = 101.20(4)96.67(4).

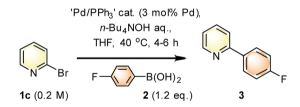
there is an exotherm during initial catalyst turnover (ca. 4 turnovers) which is associated with full dissolution of aqueous 1 M n-Bu₄NOH (into THF - overall concentration equals 0.5 M n-Bu₄NOH). The same reaction mediated by II showed a similar kinetic curve {Fig. 7(B)}, confirming the catalytic competency of this key species isolated earlier. Furthermore, no Pd particles were visibly seen to form during catalysis (the solution appearing completely homogeneous).

Altering the $Pd_3(OAc)_6/12PPh_3$ (1:4, Pd/PPh_3) led to a poor catalyst system for reaction $1c + 2 \rightarrow 3$ at 40 °C (Fig. 7(C), curves illustrated by diamonds}. This catalyst system exhibited higher catalyst efficacy at 70 °C. Thus, additional phosphine slows down catalysis in the reaction of $1c + 2 \rightarrow 3$, at 40 °C, which is

Fig. 6 Schoenebeck's findings¹⁷ on the formation of a Pd₃-type cluster from an electron-rich Pdl dimer species.

an outcome consistent with our previous studies on SMCCs involving 1c.6b

With the finding that $[Pd_2(\mu-PPh_2)(\mu_2-OAc)(PPh_3)_2]$ II reacts with 2-bromopyridine 1c to give [Pd₃(Br)(μ-PPh₂)₂(PPh₃)₃]OAc $(Pd_3Br \cdot OAc)$ and $[Pd(X/X')(N,C^2-pyr)(PPh_3)]_2$ (4a-c), additional catalytic experiments were devised to test the importance of such species in the SMCC reaction $1c + 2 \rightarrow 3$ (Fig. 8). Two control experiments, with different Pd catalysts, were conducted: (a) to establish the catalytic competency of Pd₃Br·Br;¹⁹ (b) to assess the catalytic activity of [Pd(Br)(N,C2-pyr)(PPh3)]2 4a, under comparable reaction conditions. The reaction of $1c + 2 \rightarrow$



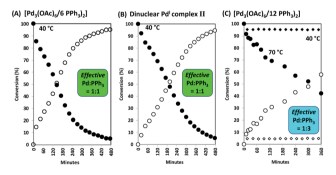


Fig. 7 Kinetic profiles for the SMCC reaction of 1c + 2 to give 3, mediated by $Pd(OAc)_2/nPPh_3$ (n = 2 and 4) and dinuclear Pd^1 complex II. (A) Reaction mediated by $Pd_3(OAc)_6/6PPh_3$; (B) reaction mediated by dinuclear Pd^I complex II; (C) reaction mediated by Pd₃(OAc)₆/12PPh₃. Reactions were monitored by ¹H NMR spectroscopic analysis in a J. Young's NMR tube (spinning). The effective Pd: PPh3 ratio takes into account that one equivalent of PPh₃ is required to reduce Pd II to Pd 0 , with concomitant formation of one equivalent of O=PPh3. For complex II, two PPh3 ligands are present overall, i.e. one per Pd; in this respect the 'PPh2' group was treated as an anionic ligand.

100
90
80
70
\$\frac{\delta}{2}\$ 60
\$\frac{\delta}{2}\$ 40

Fig. 8 Overlay of kinetic curves for the SMCC reaction of $1c + 2 \rightarrow 3$, mediated by $Pd_3Br \cdot Br$ (1 mol%), 4a (0.5 mol%) and II (1.5 mol%, generating $Pd_3Br \cdot OAc$, 4a-c in situ); other reaction conditions identical to Fig. 6. Reactions were monitored by 1H NMR spectroscopic analysis in a J. Young's NMR tube (spinning).

3, mediated by $Pd_3Br \cdot Br$ (1 mol%, giving 1 mol% active Pd – the cluster being treated as a well-defined catalyst species¹¹) gave 3 with full conversion after ca. 7.5 h at 40 °C (Fig. 8). The same reaction mediated by an authentic sample of 4a (0.5 mol% giving 1 mol% active Pd^{6b}) gave 3 with 32% conversion after ca. 7.5 h. These control experiments establish that $Pd_3Br \cdot Br$ is a significantly more active catalysis species than 4a. Thus, when generated $in \ situ$, we expect Pd_3Br^+ species to play a more dominant role in terms of the overall catalyst efficacy $vide \ infra$.

With the kinetic profiles for the SMCC reaction of $1c + 2 \rightarrow 3$, mediated by either $Pd_3Br \cdot Br$ or 4a, established, we could then qualitatively compare the catalytic activity mediated by $Pd_3-Br \cdot OAc$ and $[Pd(X/X')(N,C^2-pyr)(PPh_3)]_2$ (4a-c) species, formed *in situ* from the reaction of II with $1c.^{20}$ The observed catalyst activity sits between the high reactivity of $Pd_3Br \cdot Br$ and comparatively lower activity of $[Pd(Br)(N,C^2-pyr)(PPh_3)]_2$ 4a.

Conclusion

Edge Article

20

In conclusion we have demonstrated that reaction of Pd₃(OAc)₆ with 6 equivalents of PPh3, that is in a Pd/PPh3 ratio of 1:2, gives an intriguing dinuclear Pd^I complex, [Pd₂(μ-PPh₂)(μ₂-OAc)(PPh₃)₂], II. Species II is relatively unstable, but characterizable, and we propose it is this species that Amatore and Jutand detected in their early studies, which resisted characterization at that time.4a An important discovery was the finding that II reacts relatively cleanly with the organohalides, CH₂Cl₂ (1a), pfluoroiodobenzene (1b) and 2-bromopyridine (1c) to afford Pd₃ cluster complexes containing bridging halide ligands, i.e. $[Pd_3(X)(PPh_2)_2(PPh_3)_3]X$, carrying an overall 4/3 oxidation state. Use of 2-bromopyridine 1c was critical in understanding that a putative 14-electron mononuclear 'Pd^{II}(R)(X)(PPh₃)' is released on forming [Pd₃(X)(PPh₂)₂(PPh₃)₃]X clusters from II. Altering the Pd/PPh₃ ratio from 1:2 to 1:4 forms $Pd^{0}(PPh_{3})_{3}$ quantitatively, generally in-keeping with Amatore's and Jutand's original studies.4 It has been established further that the Pd/ PPh3 ratios are important in an exemplar SMCC reaction, Pd/ PPh₃ ratio Pd/PPh₃ reaction, $1c + 2 \rightarrow 3$. Near identical catalytic efficacy was seen for a reaction mediated by either 1Pd(OAc)₂/

2PPh₃ or II, whereas the $1Pd(OAc)_2/4PPh_3$ catalyst system was significantly less effective, requiring a higher temperature (70 rather than 40 °C) for reasonable conversion to product 3 to be observed.

An important take home message from our study is that where $[Pd_2(\mu\text{-PPh}_2)(\mu_2\text{-OAc})(PPh_3)_2]$ II can form, *i.e.* when a ratio of Pd/PPh₃ ratio is 1:2 employed in catalysis, reactions with organohalides (common starting materials for cross-coupling catalysis) afford catalytically competent Pd₃ cluster complexes *in situ*, in addition to other known Pd^{II} species (*i.e.* oxidative addition products). If the relative amount of PPh₃ ligand to Pd is low, then Pd clustering tends to occur, to afford either particles (where Pd/PPh₃ = 1:1), or 'ligated clusters', whereas well-defined dimers and trimers are formed where Pd/PPh₃ = 1:2 (the major finding of this study) and when there is enough PPh₃ ligand around, mono-nuclear Pd⁰(PPh₃)_n, *i.e.* n > 2, can be stabilised, aligning with a general understanding of ligated Pd⁰ species in text book mechanisms.

Understanding how [Pd₃(X)(μ-PPh₂)₂(PPh₃)₃]X clusters activate aryl/heteroaryl halides and organometallic coupling partners, *e.g.* aryl boronic acids,¹¹ will no doubt be important going-forwards, which will enable their catalytic properties to be fully delineated and exploited in chemical synthesis. To emphasize this point further, similar Pd₃-type clusters have been studied by Maestri and Malacria in catalysis, particularly hydrogenation.²¹ Our results, taken together with contributions made by others, show that Pd₃-clusters are ripe for exploitation in applied catalysis.

Conflicts of interest

There are no conflicts to declare.

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