Palladium-Catalyzed Carbonylation/Acyl Migratory Insertion Sequence**

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Migratory insertion is one of the fundamental processes in organopalladium chemistry. In particular, migratory insertion of a CO ligand and the formation of reactive acylpalladium intermediate is a powerful method for introducing a carbonyl functionality into organic molecules. The catalytic cycle involving such a key step has been developed into one of the most important tools to synthesize various carbonyl compounds.[1] However, for a long time, the scope of migratory insertion processes of organopalladium has been limited to those involving carbon monoxide. In view of the similarity between a carbene and carbon monoxide, one may conceive palladium–carbene as another species which may undergo migratory insertion (Scheme 1). Indeed, the migratory insertion process has been reported for stable palladium–carbene species.[2] More recently, catalytic reactions which are proposed to include the migratory insertion of a palladium–carbene intermediate have been reported.[3–6] The migratory group has so far included aryl,[3] benzyl,[3a,4] vinyl,[5] and allyl[6] groups. Herein we report an unprecedented acyl group migration in this type of reaction. This reaction proceeds with a palladium-catalyzed carbonylation of an aryl iodide, palladium–carbene formation, and acyl migratory insertion.

We have previously reported the palladium-catalyzed cross-coupling reaction of diazo compounds with organoboronic acids.[3e] We also studied the coupling of iodides with ethyl diazoacetate (EDA) in the presence of CO.[7] As a continuation of our studies, we investigated the palladium-catalyzed reaction of iodobenzene (1a) with methyl α-diazoacetionate (2a) in the presence of an atmosphere of CO. Methyl 2-methyl-3-oxo-3-phenylpropanoate (4a) was isolated in moderate yield (40%; Scheme 1). The formation of 4a can be rationalized by invoking an acyl group migratory insertion of a palladium–carbene species (from complex a to complex b; Scheme 2).[8]

Optimization of this reaction was attempted (Table 1). Firstly, since the product has a newly incorporated hydrogen atom from an external source, we anticipated that the introduction of a hydrogen source should promote the reaction. To our delight, this was indeed the case; when the reaction was carried out in the presence of triethylsilane (3), the yield of 4a was improved to 88% (Table 1, entry 1).

Furthermore, inspection of the reaction conditions revealed that this reaction proceeded more efficiently in polar solvents such as 1,2-dichloroethane (DCE) and CHCl₃ (Table 1, entries 1 and 3–6), whereas the nonpolar solvent PhMe was found to be unfavorable (Table 1, entry 2). Several other palladium catalysts were then examined. Of the palladium(II) catalysts, Pd(OAc)₂ gave trace amounts of products and [Pd(PPh₃)₂Cl₂] afforded moderate yields (Table 1, entries 7 and 8). [Pd₂(dba)₃] afforded low to moderate product yields under the reaction conditions, both in the presence and absence of phosphine ligands (Table 1, entries 1 and 3–6), whereas the nonpolar solvent PhMe was found to be unfavorable (Table 1, entry 2). Several other palladium catalysts were then examined. Of the palladium(II) catalysts, Pd(OAc)₂ gave trace amounts of products and [Pd(PPh₃)₂Cl₂] afforded moderate yields (Table 1, entries 7 and 8). [Pd₂(dba)₃] afforded low to moderate product yields under the reaction conditions, both in the presence and absence of phosphine ligands (Table 1, entries 9–15). Therefore it was concluded that [Pd(PPh₃)₄] was the best catalyst for this reaction. Notably, the products resulting from a β-hydride elimination were not observed in any case. Finally, a control experiment showed that product 4a could not be detected in the absence of a palladium catalyst (Table 1, entry 16).

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A series of aryl iodides 1a–i were then subjected to the optimal reaction conditions with α-diazoacarbonyl compounds 2a–j. As shown in Table 2, the corresponding 1,3-dicarbonyl products were obtained in moderate to good yields in all cases. The reaction was found to be marginally affected by the steric of the substituents on the aryl iodides, as shown by the product of diazoester 2a with α-iodotoluene (1b), which gave lower yields (Table 2, entry 2). For the scope of diazo substrates, both α-alkyl- and α-aryl-substituted substrates worked well to afford the corresponding products in good yields, except in the case when the α-aryl α-diazoacetate contains strong electron-withdrawing substituents.

Next, this palladium-catalyzed reaction was extended to diazo substrates not bearing a carbonyl substituent. These nonstabilized diazo substrates can be generated in situ from N-tosylhydrazones. Therefore, phenyl iodide (1a) and N-tosylhydrazone 5a were subjected to similar reaction conditions in the presence of LiOBu. A mixture of the expected ketone 6a and enone 7a (93:7) was isolated in 67% yield (Table 3, entry 1). The product 7a results from a β-hydride elimination in the last step of catalytic cycle (see below). Additional studies revealed that both the yield and ratio of the products were drastically affected by the type of palladium catalyst (Table 3, entries 2–7). After extensive optimization,[9] two sets of reaction conditions were identified, which could afford either 6a or 7a selectively, both in good yields [Table 3, entry 8 (conditions I) and entry 9 (conditions II)].

With the two sets of optimized conditions, the reaction scope was examined (Table 4). Both reaction conditions I and II worked well, affording the either the ketone (6b–e) or the enone (7b–e), respectively; however, in the case of 5c the β-
Palladium-catalyzed reaction of CO with \( \text{ArC} = \text{O} \) generates a carbene intermediate which migrates to the carbon monoxide insertion affords the Pd–acyl intermediate \( \text{B} \). Interaction of the \( \alpha \)-diazo compound with \( \text{B} \) produces palladium–carbene intermediate \( \text{C} \), and migratory insertion of the acyl group into the carbenic carbon atom of the palladium–carbene generates C-bound enolate \( \text{D} \), which equilibrates with the corresponding O-bound enolate \( \text{E} \). In the reaction with \( \alpha \)-diazacarbonyl compound, \( \eta^1\)-O, O-bound intermediate \( \text{E} (\text{R} = \text{COR}^\prime) \) is predominant, from which transmetalation with \( \text{Et}_3\text{SiH} \) and subsequent reductive elimination affords 1, 3-dicarbonyl compound as the only product. For the reaction with a nonstabilized diazo compound as a substrate, the equilibrium between \( \text{D} \) and \( \text{E} \) would be influenced by the phosphine ligands.\(^{[11]} \) \( \beta \)-Hydride elimination from \( \text{D} \) affords enone product.

Since the possibility exists that the primary product of the reaction is a silyl enolate rather than a ketoester, we prepared silyl enolate from ketoester \( \text{4a} \) for comparison.\(^{[9]} \) The silyl enolate was found to be stable to silica gel column chromatography. Careful inspection of the crude mixture of the palladium-catalyzed reaction of \( \text{1a} \) and \( \text{2a} \) under standard reaction conditions could not detect any silyl enolate. This experiment rules out the possibility of silyl enolate as primary product, which is evidence supporting the palladium hydride species \( \text{G} \) as the intermediate in the final step of the catalytic cycle.

To substantiate the mechanistic rationale, we examined palladium-catalyzed reactions of methyl \( \alpha \)-diazoacetate \( \text{2a} \) and benzoyl chloride \( \text{8} \); Scheme 4). The reaction gave the acyl group migration product \( \text{4a} \) in 21\% yield. In the presence of an atmosphere of CO, the yield was slightly higher. A control experiment suggests that no reaction occurs in the absence of the palladium catalyst. These results are consistent with the mechanistic rationale which involves acyl migratory insertion as the key step.

In summary, we have reported the first palladium-catalyzed tandem migratory insertion with both CO and a carbene. This reaction builds a connection between a palladium–carbene process and palladium-catalyzed carbonylation, which may open new possibilities for the exploration of the potential of palladium-catalyzed carbene transformations.

### Experimental Section

Typical procedure for \([\text{Pd(PPh}_3)_4]\)-catalyzed reactions of CO with \( \alpha \)-diazacarbonyl compounds and aryl iodides: Under a nitrogen atmosphere, \([\text{Pd(PPh}_3)_4]\) (17.3 mg, 0.015 mmol) was added to a flame-dried round-bottomed flask. The flask was then sealed and evacuated to a vacuum of 15 mmHg, and fitted with a CO balloon. A solution of iodobenzene (1a: 61 mg, 0.3 mmol), methyl \( \alpha \)-diazoacetate \( \text{2a} \) (68 mg, 0.6 mmol), triethylsilane (3; 38 mg, 0.33 mmol), and triethylamine (61 mg, 0.6 mmol) in 4 mL of DCE was added using a syringe. The mixture was stirred at 60\°C until \( \text{2a} \) disappeared as judged by TLC. The solution was removed in vacuo to yield a residue, which was purified by flash chromatography (silica gel) to afford pure \( \text{4a} \) as a pale yellow oil (51 mg, 88\%).

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For details, see Supporting Information.
