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Palladium-catalyzed reductive electrocarboxylation of allyl esters with carbon dioxide†

Ke-Jin Jiao,^a Zhao-[Min](http://orcid.org/0000-0002-4985-1071)g Li,^a Xue-Tao Xu, D^b Li-Pu Zhang,^b Yi-Qian Li,^b Kun Zhang^b and Tian-Sheng Mei D^{*a}

Palladium-catalyzed regioselective electrocarboxylation of homostyrenyl acetates with $CO₂$ has been successfully developed, providing α -aryl carboxylic acids with good selectivity and yield. In addition, the catalytic asymmetric carboxylation of cinnamyl acetate has been demonstrated, despite moderate

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Introduction

Catalytic carbon–carbon (C–C) bond forming reactions that utilize carbon dioxide $(CO₂)$ to directly access carboxylic acids have received extensive attention in the past decades due to the attractiveness of CO₂ as a C1 synthon.^{1,2} In particular, transition metal-catalyzed allylation of carbon dioxide has emerged as a promising tool to construct $β, γ$ -unsaturated carboxylic acids.3 Most allylic carboxylation reactions that incorporate $CO₂$ require highly reactive substrates such as allylstannanes or allylboron reagents, which is due to both the thermodynamic stability and kinetic inertia of $CO₂$.⁴ Recently, Martin and co-workers elegantly demonstrated Ni-catalyzed carboxylation of allyl acetates with $CO₂$ utilizing Mn as a reducing agent (Scheme 1a).⁵ Similarly, Sato and co-workers have successfully developed Pd-catalyzed carboxylation of allylic alcohols using Et₂Zn as a reducing agent.⁶ More recently, the Martin group⁷ and our group⁸ independently reported Ni-catalyzed carboxylation of allyl alcohols using Mn powder. Unfortunately, the use of super-stoichiometric amounts of reducing agents like Mn or $Et₂Zn$ is a drawback of these reactions. Furthermore, these developed protocols are not suitable for homostyrenyl electrophiles (Scheme 1b). Enantio- and regioselective carboxylations of homostyrenyl electrophiles are needed as an efficient way to deliver α-aryl carboxylic acids, which are found in numerous **PUBLIC CONFINITY CONTROLL CONFILM CONTROLL CONFILM CONTROLL CONFILM CONTROLL CONFILM CONTROLL CONFILM CONTROLL CONTROLL CONTROLL CONFILM CONTROLL CONFILM CONFIDENCIAL CONFIDENCIAL CONFIDENCIAL CONFIDENCIAL CONFIDENCIAL CO**

enantioselectivity.

medicines (Scheme 1c). Toward that end, the development of catalytic carboxylation reactions using more environmentally benign reducing agents is highly desirable.⁹

Electrochemistry has been demonstrated to be an efficient and environmentally friendly approach for organic chemistry that employs electrons as redox reagents.^{10–12} As part of our ongoing interest in using electric currents as redox reagents in palladium catalyzed reactions, 13 we questioned whether catalytic carboxylations of allyl acetates could be achieved using electric current as a reducing agent. 14 Herein, we describe the first Pd-catalyzed regioselective carboxylation of homostryrenyl acetates utilizing electric current as the reductant, thus providing an environmentally benign method for accessing α-aryl

Scheme 1 Catalytic carboxylation of allyl electrophiles.

^aState Key Laboratory of Organometallic Chemistry, Center for Excellence in Molecular Synthesis, Shanghai Institute of Organic Chemistry, University of Chinese Academy of Sciences, Chinese Academy of Sciences, 345 Lingling Lu,

Shanghai 200032, China. E-mail: mei7900@sioc.ac.cn

^bSchool of Chemical & Environmental Engineering, Wuyi University, Jiangmen, 529020, China

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carboxylic acids. In addition, we demonstrate the first catalytic asymmetric carboxylation of cinnamyl acetate, which provides a heretofore unavailable opportunity to construct chiral phenyl acetic acids from homostyrenyl acetates (Scheme 1d).

Results and discussion

Our initial studies focused on Pd-catalyzed carboxylation of (E) -3-(4-bromophenyl)allyl acetate 1a with $CO₂$ (Table 1). We chose 1a as the model substrate because it contains relatively reactive C–Br and C–O bonds, which may have different reactivities under electrochemical conditions, and the retained bromide may be subsequently elaborated into useful products by employing various metal-catalyzed cross-coupling reactions.¹⁵ To our delight, at 30 °C, using Pd(OAc)₂ as catalyst and 1,2-bis(diphenylphosphino)benzene (DPPPh) as ligand, with Et4NOTs as electrolyte, EtOH as an additive, and DMF as solvent, the reaction of 1a with $CO₂$ gave 81% isolated yield of branched product 2a with 20 : 1 regioselectivity under constant current electrolysis conditions at 8.0 mA $(J = 8.0 \text{ mA cm}^{-2})$ at 3 F mol−¹ (entry 1). Lower regioselectivity was observed when other phosphine ligands were used, including $PPh₃$, and bidendate DPPE, DPPP, DPPB (entries 2–5). Without EtOH, the Organic Chemistry Frontiers

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Table 1 Reaction optimization with substrate $1a^a$

	OAc	$Pd(OAc)_2$ (5.0 mol%) DPPPh (5.3 mol%) EtOH (1.0 equiv)		CO ₂ H Ar 2a	
Br 1a (0.3 mmol)		$CO2$ (1 atm), $Et4$ NOTs (0.07 M) 8 mA, 30 °C, 3 h, DMF (6 mL) undivided cell, Pt-Mg		+ Ar CO ₂ H 2aa	
Entry	Deviation from above conditions	F mol^{-1}	Conv b (%)	$Yield^b$ (9/0)	2a/ $2aa^c$
1	None	3	100	85 $(81)^d$	20:1
2	$PPh3$ (12 mol%) in lieu of DPPPh	3	79	36	2:1
3	DPPE in lieu of DPPPh	3	85	70	15:1
4	DPPE in lieu of DPPPh	3	81	65	18:1
5	DPPE in lieu of DPPPh	3	68	51	6:1
6	No EtOH	3	38	21	2:1
7	$EtOH$ (0.5 equiv.)	3	94	64	20:1
8	$EtOH$ (5.0 equiv.)	3	81	59	11:1
9	no $Pd(OAc)_2$	3	32	19	2:1
10	no DPPPh	3	72	38	3:1
11	no electric current		Ω	NP	
12	Mn or Zn in lieu of electric current PPh ₂		Ω	NP	
	Ph ₂ P PPh ₂	Ph_2P	PP _{h₂} Ph_2F		PPh ₂
DPPPh	DPPF	DPPP		DPPR	

^a Reaction conditions: 1a (0.3 mmol), Pd(OAc)₂ (5.0 mol%), DPPPh (5.3 mol\%) , Et₄NOTs $(0.15 \text{ g}, 0.07 \text{ M})$ and EtOH (1.0 equiv.) , DMF (6 mL) in an undivided cell with a platinum electrode and a mag-
nesium rod as sacrificial anode. b The ratio of B/L products was determined by ¹H-NMR. ^c The yields were determined by ¹H-NMR with $CH₂Br₂$ as the internal standard. ^d Isolated yield of 2a.

regioselectivity and yield were significantly reduced (entry 6). Decreasing or increasing the amount of EtOH to 0.5 or 5.0 equivalents resulted in a slight decrease in yield (entries 7 and 8). Cathodic reduction of EtOH could take place to stabilize the reaction system, although the real role of EtOH is not clear at this stage. Furthermore, both $Pd(OAc)$ and phosphine ligand are required for achieving regioselective carboxylation (entries 9 and 10). Finally, a control experiment gave the results that no reaction was observed in the absence of electric current (entry 11). It is worth noting that chemical reductant such as Mn or Zn is not suitable for this carboxylation reaction (entry 12). This implies that electrochemistry could provide unique reactivity by enhancing the nucleophilicity of resulting palladium species (vide infra).

With the optimized reaction conditions in hand, we next investigated the scope of this reductive electrocarboxylation reaction (Table 2). In addition to Br, our conditions were compatible with numerous other functional groups, including F, Cl, Me, OMe, t -Bu, and CF₃ (2a–2r). Substrates containing electron-withdrawing groups (2a, 2f–2h) deliver better regioselectivity than electronrich ones (2b–2e). Interestingly, this electronic factor is different from Pd-catalyzed allylic alkylation, in which electron-withdrawing groups on aryl rings reduce formation of the branched product.¹⁶ It is worth emphasizing that polysubstituted arenes also deliver high regioselectivity (2n–2r). Finally, substrates bearing polycyclic

Yields of the indicated major product were reported. Regioselectivities were determined by crude 1 H-NMR analysis. b DPPE was used as the ligand.

aromatics such as naphthalene and anthracene were well tolerated in the reaction (2s–2u). Unfortunately, alkyl-substituted allylic esters or 3,3-disubstituted allylic esters are not effective at current reaction conditions.

Next, we examined the influence of the leaving group on the reaction, since many allylic compounds with appropriate leaving groups can form π-allylpalladium intermediate via oxidative addition to $Pd(0)$ under mild conditions.¹⁷ Unsurprisingly, the different leaving groups influence reaction performance under the optimized conditions (Table 3). Among the esters we tested, cinnamyl isobutyrate (3a) gave the best regioselectivity $(10:1)$. However, cinnamyl pivalate $(3b)$ provided the best yield (81%) with slightly lower regioselectivity $(8:1)$. We also investigated the electronic influence of substituted benzoate leaving groups (3d–3g), but we did not observe any correlation between Hammett values and regioselectivity.

The enantioselective carboxylation of allylic acetates is a major challenge.¹⁸ To our delight, chiral bidentate triarylphosphine ligands were found to be efficient for this catalytic asymmetric carboxylation, and moderate enantioselectivity (67% ee) was achieved (Scheme 2). Interestingly, chiral bidentate phosphine ligands are known to be inactive in Pd-catalyzed allylation of aldehydes.¹⁹

To gain insight into the reaction mechanism, a cyclic voltammogram (CV) was conducted. The CV of $[(R)$ -BINAP]Pd

 $(OAc)_2$ in DMF reveals one irreversible reduction wave at -0.60 V vs. Ag/AgI, corresponding to the reduction of $Pd(n)$ to $Pd(0)$ (b, Fig. 1). The addition of homostyrenyl acetate 1b causes an increase in the peak current, which indicates oxidative addition of $Pd(0)$ to allyl acetate **1b** (c, Fig. 1). Furthermore, we also observed the second reduction wave at −0.90 V, which is tentatively assigned to the reduction of the resulting allyl $Pd(n)$ species (c, Fig. 1). The further increase of peak currents (both around -0.65 V and -1.05 V) after the addition of CO₂ suggests that the carboxylation takes place (d, Fig. 1).

Based on Wendt's and Hazari's elegant studies on η^1 . allylPd species reacting with $CO₂$ and our observation, a plausible mechanism for this Pd-catalyzed carboxylation reaction *via* electrochemical reduction is proposed in Scheme $3⁴$ Initially, oxidative addition of Pd(0) to the allyl electrophile generates a cationic π -allylpalladium(π) complex (A) with leaving group (OAc) as the counteranion,²⁰ which is in equilibrium to the terminal η^1 -allylpalladium(π) species (**B** and **C**).⁴ The formation of species C, which leads to linear carboxylation product, is less favored due to the steric hindrance between catalyst and aryl group. Next, complex B is directly reduced at the cathode to form an anionic η^1 -allylpalladium(0) species (D) by overall two-electron transfer, which is consistent with the studies on carboxylation of aryl halides by Amatore, Jutand and co-workers. 21 Finally, the nucleophilic complex **D** reacts with $CO₂$ at the γ -position to deliver product E in the presence **Research Article**
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 α Yields and regioselectivities were determined by crude $\rm ^1H\text{-}NMR$ ana-

Fig. 1 Cyclic voltammograms recorded on a Pt electrode at 100 mV s^{-1} in: (a) DMF containing 0.07 M Bu_4NPF_6 ; (b) solution (a) after addition of 2 mM $[(R)-BINAP]Pd(OAc)_{2}$; (c) solution (b) after addition of 20 mM 1b; (d) solution (c) saturated with $CO₂$.

Scheme 2 Enantioselective carboxylation.

lysis. ^b Isolated yield in parentheses.

Scheme 3 Proposed reaction mechanism.

of Mg^{2+} derived from anodic oxidation, thus regenerating Pd(0). Since carboxylation did not occur in the absence of electric current, the terminal η^1 -allylpalladium(II) species (B) attack $CO₂$ directly to deliver the palladium(π) carboxylate is not likely in our reaction system.²²

Conclusions

In summary, we have demonstrated the first example of Pd(0) catalyzed regioselective carboxylation reaction of homostyrenyl acetates under electrochemical reduction. Compared to established catalytic methods, we employed raw electrons to facilitate this transformation. In addition, an enantioselective carboxylation was also realized via Asymmetric Organometallic Electrochemistry (AOE). These findings provide a new avenue for transition metal-catalyzed electrochemical reactions using chiral ligands. Investigations aimed at understanding the reaction mechanism and further increasing the scope of these reactions are currently underway in our laboratory.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

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