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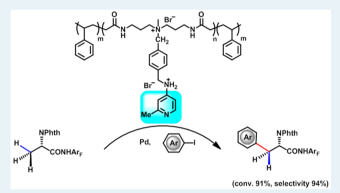
Functionalized Polymer-Supported Pyridine Ligands for Palladium-Catalyzed C(sp³)-H Arylation

Li-Chen Lee,[†] Jian He,[‡] Jin-Quan Yu,*,[‡] and Christopher W. Jones*,[†]

[†]School of Chemical & Biomolecular Engineering, Georgia Institute of Technology, Atlanta, Georgia 30332-0100, United States Department of Chemistry, The Scripps Research Institute, La Jolla, California 92037-1001, United States

Supporting Information

ABSTRACT: The use of ligands to tune the reactivity and selectivity of transition-metal catalysts for $C(sp^3)$ -H bond activation is a current central challenge. One of us previously developed an uncommon example of a homogeneous catalyst that performs controlled C(sp³)-H arylation using pyridine derivatives as ligands, along with Pd [Science, 2014, 343, 1216-1220]. In this work, we report a functionalizable and tunable polymer support used in the immobilization of pyridine derivatives that yields a soluble, polymeric ligand platform facilitating C(sp³)-H activation reactions with good yields, selectivities differing from the homogeneous catalyst, and recovery of Pd. Unlike the homogeneous system, the supported catalysts in Pd-catalyzed C-H monoarylation



reactions respond sensitively to the steric hindrance of the coupling partners.

KEYWORDS: C-H activation, arylation, polymer-supported ligand, immobilized catalyst, palladium

he possibility of direct introduction of a new functionality ↓ (or a new C−C bond) via a direct C−H bond transformation is a highly attractive strategy in organic synthesis. The range of substrates has been expanded to simple hydrocarbons, complex organic small molecules, and synthetic and biological polymers. 1-7 Transition-metal catalysis has been used extensively in recent years to assist in converting unactivated C-H bonds to more reactive carbon-metal bonds that can subsequently be further functionalized to afford the desired products. 8-15

In general, "inert" C(sp³)-H bonds possess low reactivity and high thermodynamic stability, compared to the $C(sp^2)-H$ and C(sp)-H bonds. This makes $C(sp^3)-H$ activation a more challenging endeavor, compared to conventional C-H functionalization routes. Many catalytic systems have been recently developed for Pd-catalyzed C-H activation. 17-22 In particular, directed Pd-catalyzed C(sp³)-H arylation reactions have been developed recently. In 2005, the Pd-catalyzed, quinoline- and pyridine-directed C(sp³)-H arylation with aryl iodides was reported by the Daugulis group.²³ A Pd-catalyzed β -C(sp³)-H arylation of carboxylic acids with aryl iodides subsequently was also achieved by the Yu group.²⁴

Most compounds contain multiple types of C-H bonds and other functional groups, and controlling site selectivity in C-H activation is necessary to help C-H functionalization methods become widely useful. To this end, the use of ligands to tune the reactivity and selectivity of transition-metal catalysts for C(sp³)-H bond activation has been a key recent focus. Recently, one of us reported the first example of ligandcontrolled C(sp³)-H arylation of an amino acid. Alaninederived amide 1 bearing a weakly coordinating aniline auxiliary, 17,25-27 in conjunction with Pd and various pyridine and quinoline-based ligands were used. We found that pyridinebased ligands were capable of lowering the transition-state energy of C(sp³)-H activation, and promoted the selective monoarylation of a C(sp³)-H bond with aryl iodides. Among the various pyridine-based ligands tested, 2-methylpyridine provided the optimal balance between yield and monoarylation versus diarylation selectivity in these homogeneous reactions.

However, examples exploring the use of heterogeneous catalysts in C-H functionalization reactions are uncommon.²⁸ In the Son and Shin groups, 29 palladium nanoparticles supported on silica nanotubes have been used to catalyze C(sp²)-H arylation. Selective heterogeneous C-H halogenation methods using palladium nanoparticles immobilized into porous metal organic frameworks (MOFs) have been reported. 30,31 Impregnated $Pd(OAc)_2$ on the MOF-5 (O_h) has been employed as a heterogeneous Pd catalyst with ultrahigh surface area. This supported catalyst was applied in the C-H phenylation of naphthalene.³² A Pd/C catalyzed arylation of triphenylene with aryiodonium salts has also been reported recently.3

Herein, we report a functionalizable and tunable polymer used for the immobilization of pyridine ligands that yields a

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Scheme 1. General Preparation of the Polymer-Supported Pyridine Derivative Used as a Ligand for Pd-Catalyzed C(sp³)-H Arylation

Chart 1. Yields Obtained during Ligand Screening in the Pd-Catalyzed $C(sp^3)$ —H Monoarylation of Alanine-Derived Amide 1 and Iodobenzene a,b

"Substrate (0.1 mmol), Pd(TFA) $_2$ (0.01 mmol), ligand (0.02 mmol), and Ag $_2$ CO $_3$ (0.15 mmol) were weighed in an atmosphere open to air and placed in a pressure tube (5 mL) with a magnetic stir bar. Iodobenzene (0.15 mmol), TFA (0.02 mmol), and DCE (0.5 mL) were added. The reaction vessel was sealed, and the mixture was first stirred at room temperature for 10 min and then heated to 100 °C for 20 h with vigorous stirring. The yield percentage and ratios of product 2 to product 3 were determined by 1 H NMR.

soluble, polymeric ligand platform that facilitates $C(sp^3)$ —H activation reactions. When substituted for the homogeneous pyridine ligands developed previously under Pd-catalyzed $C(sp^3)$ —H activation conditions, the resulting catalyst not only offers comparable catalytic activity to the unsupported catalyst system, but it allows for catalyst recovery, new catalytic selectivity not observed in the unsupported system, and some catalyst recyclability. To the best of our knowledge, this represents the first $C(sp^3)$ —H activation of an amino acid derivative using a polymer-supported Pd catalyst.

This new polymer was constructed based on monomer 6, which has two cross-linkable acrylamide and one benzyl bromide functional group (Scheme 1). A low-molecular-weight polymer backbone was formed by radical polymerization

between styrene and monomer $\bf 6$ (Scheme 1). Based on an investigation of the relative rates of disappearance of monomer $\bf 6$ and styrene by in situ $^1{\rm H}$ NMR (see Figure S3 in the Supporting Information), it is apparent that the polymerization proceeded to form a mostly random copolymer. Figure S4 in the Supporting Information shows the integration ratios between styrene ($\rm H_e$) and monomer $\bf 6$ ($\rm H_b$) during polymerization, and it was observed that the ratio increased modestly in the first 4 h and then remained constant during the rest of the reaction. The amide functional groups within the polymer backbone, based on alanine-derived amide 1, were specifically included to help concentrate the polar substrates within the polymeric domains, extracting these species from the relatively nonpolar reaction solutions, 34 to help accelerate the catalytic

reactions. Once the polymeric ligand backbone was prepared, a pyridine-based ligand was post-modified³⁵ onto the polymer by amination of the benzyl bromide functional group within the polymer using commercially available 4-amino-2-methylpyridine, L3 (Scheme 1). The completion of the ligand modification was monitored by ¹H NMR (see Figure S6 in the Supporting Information). As noted above, the coordination environment of the supported ligand within the polymer and metal was adjusted by tuning the ratios of the functionalized monomer and styrene during polymer preparation, altering the spacing between the ligands within the polymer. In this work, the polymer support is abbreviated as "PL", and the ratio of styrene to monomer is added in parentheses as a suffix.

The Yu group previously developed reaction conditions for the target reaction between alanine-derived amide 1 and iodobenzene (10 mol % Pd(TFA)2, 20 mol % ligand, 20 mol % TFA, Ag₂CO₃ (1.5 equiv), iodobenzene (1.5 equiv) in 1,2dichloroethane (DCE) at 100 °C) to prepare compound 2 in good yield and selectivity. Chart 1 reports data under identical reaction conditions from tests of a library of homogeneous pyridine-based ligands that had similar chemical structures to the optimal ligands previously identified by the Yu group, but were also immobilizable via reaction with a benzyl bromide group onto our polymeric support (see Scheme 1). Ligands L1 and L2 were highly selective for monoarylation, but neither enhanced the conversion substantially, relative to the ligandfree catalyst. Ligand L3 enhanced the reactivity relative to the ligand-free conditions, while offering good selectivity to the monoarylation product. The N-alkylated ligand, L4, also showed similar reactivity, suggesting that incorporation of L3 into a polymer backbone (Scheme S1 in the Supporting Information) would not negatively affect catalytic reactivity, from an electronic perspective. Interestingly, the molecular ligands L3 and L4, which had chemical structures that were similar to that of the polymer-supported ligand, PL(4), showed slightly lower yields of the monoarylated product than the polymer-supported ligand. PL(4) seemed to possess a useful balance of steric and electronic properties that provided compound 2 in high yield and with excellent selectivity for monoarylation.

We hypothesize that the polar amide functional groups in the polymer backbone helped to accelerate the reaction by concentrating the substrate in the polymer domains, partitioning from the relatively hydrophobic solution, while maintaining high selectivity for monoarylation. An array of polymer-supported ligands were thus prepared to have ratios of styrene to functionalized monomer in the range of 2–8, and, subsequently, the impact of the polymer composition on the performance in the target arylation reaction was evaluated. Figure 1 shows that the polymer-supported ligand constructed with a styrene/monomer ratio of 4 ratio yielded the highest reactivity and product selectivity (PL(4)). Based on our observations, polymers with a lower ratio of styrene to functionalized monomer showed poor solubility in dichloroethane, hampering their utility.

Previously, to gain insight into the coordination of the substrate and ligand at the Pd(II) center, the Yu group prepared and characterized a C-H insertion intermediate formed via primary $C(sp^3)-H$ arylation in the absence of aryl iodides. This intermediate was demonstrated to be a viable precatalyst for primary $C(sp^3)-H$ arylation (Scheme 2). We hypothesized that some of the polymers prepared in this work might not easily facilitate the formation of this C-H insertion

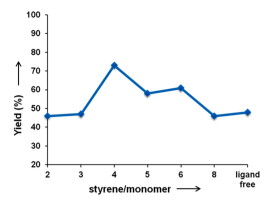


Figure 1. Yield of monoarylation product 2 from reactions using polymers with various ratios of styrene to functional monomers in the Pd-catalyzed $C(sp^3)$ -H monoarylation. The experimental conditions were the same as those described in Chart 1.

intermediate, because of spatial constraints within the polymer. For example, the polymers that had a lower ligand density (see Table S3 in the Supporting Information, styrene/monomer = 8 (PL(8))) might inhibit the proper coordination between two pyridine ligands and Pd during the reaction, giving lower product yields (Figure 1, styrene/monomer = 8 (PL(8))). Interestingly, the polymers that had a relatively high concentration of ligand (Table S3, styrene/monomer = 2 and 3 (PL(2) and PL(3), respectively)), did not facilitate the reaction as effectively either (Figure 1, styrene/monomer = 2 and 3 (PL(2) and PL(3), respectively)). These ligands may have dramatically lowered the energy of the palladacycle intermediate via a chelation effect, thus impeding the subsequent functionalization step of this $C(sp^3)$ -H arylation reaction. We hypothesize that the best polymer provided an optimal balance of solubility and correct spatial coordination with the Pd species.

During the reaction, we hypothesize that the polar amide substrate would be concentrated within the polymer, because of the relatively polar amide-based backbone when the reactions were conducted in relatively nonpolar solvents. In such a scenario, the substrate and Pd-ligand complexes would be in close proximity. This hypothesized preferential interaction of the amide substrate with the functionalized polymer backbone was supported by observed differences in reactivity using various organic solvents (see Table 1). The reaction proceeded smoothly in nonpolar solvents (entries 1 and 2, Table 1), whereas more polar solvents lowered the reaction yield progressively (entries 5 and 6, Table 1). To obtain the optimal balance of selectivity and yield, we used a cyclohexane/DMF mixed solution as the reaction solvent for further studies. In Figure 2, various volume ratios of cyclohexane and DMF were studied in the monoarylation reaction. When the cyclohexane/ DMF volume ratio was 8, both outstanding selectivity (94%) and yield of the monoarylation product (91%) were obtained. In Yu's previous work, 2-methylpyridine was successfully used as the ligand in conjunction with Pd to synthesize phenylalanine derivatives (2a-2f) with electron-rich or electron-poor groups at the ortho-, meta-, or para-positions in high yields.

This reaction was also demonstrated to be tolerant of halide substituents and a wide range of polar functional groups. Herein, we evaluated the catalytic activity of PL(4) with aryl iodides bearing both monosubstituted and disubstituted electron-donating or electron-withdrawing groups under our optimized reaction conditions. Overall, the electron-with-

Scheme 2. Catalytic Reactivity of Intermediates in the C(sp³)-H Arylation Reaction

NPhth
H
CONHAr_F

10 mol%
Intermediate A

TFA, Ph-I

Ag₂CO₃, DCE
100
0
C, 20h

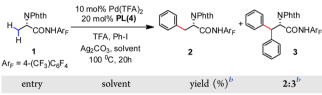
NPhth
CONHAr_F

TRA, Ph-I

Ag₂CO₃, DCE
100 0 C, 20h

Intermediate A

Table 1. Solvent Effect in Pd-Catalyzed C(sp³)-H Monoarylation of Alanine-Derived Amide 1 and Iodobenzene^{a,b}



entry	solvent	yield (%) ^b	2:3 ^b
1	cyclohexane	77	68:9
2	DCE	73	70:3
3	toluene	57	56:1
4	1,4-dioxane	55	54:1
5	DMF	40	40:0
6	CH ₃ CN	7	7:0

"Substrate (0.1 mmol), $Pd(TFA)_2$ (0.01 mmol), PL(4) (0.02 mmol), and Ag_2CO_3 (0.15 mmol) were weighed out in an atmosphere open to air and placed in a pressure tube (5 mL) with a magnetic stir bar. The iodobenzene (0.15 mmol), TFA (0.02 mmol), and solvent (0.5 mL) were added. The reaction vessel was sealed and the mixture was first stirred at room temperature for 10 min and then heated to 100 °C for 20 h with vigorous stirring. ^bThe yield percentage and ratios of product 2 to product 3 were determined by 1H NMR.

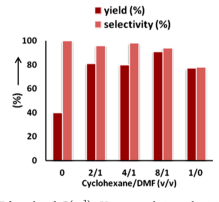


Figure 2. Pd-catalyzed $C(sp^3)$ —H monoarylation selectivity and yield in cyclohexane/DMF mixed solutions. The experimental conditions are the same as those described in Table 1.

drawing substituted aryl iodides were less reactive, compared to the electron-donating substituted aryl iodides. Interestingly, the polymer-supported ligand showed steric selectivity of aryl iodide partners in monoarylation reactions, which has not been observed in the previous homogeneous Pd-ligand catalyzed reaction. With the supported catalyst, the yields were decreased significantly when the substituents on the aromatic ring were close to the active site (2a, 2b, and 2e; see Chart 2). Apparently, Pd ligated to the immobilized ligand was impacted by steric constraints associated with the polymer support. For example, when methoxy-substituted iodobenzenes were used as the coupling partners in this $C(sp^3)$ —H monoarylation, *ortho-*

Chart 2. Substrate Effects in the Pd-Catalyzed C(sp³)-H Monoarylation Using PL(4)

methoxyiodobenzene showed significantly lower conversion than para-methoxyiodobenzene ($2a_1$ and $2a_3$; see Chart 2). This phenomenon was also observed in ester and fluorosubstituted iodobenzenes as well (see $2b_1$ and $2b_3$, as well as $2e_1$ and $2e_3$, in Chart 2), where iodobenzenes bearing substituents at the meta-position showed more moderate conversions ($2a_2$, $2b_2$ and $2e_2$; see Chart 2). This unique property could be potentially exploited in protocols aimed at selective $C(sp^3)-H$ monoarylation with aryl iodides that contain multiple iodides.

The supported ligand, when combined with Pd, yielded a highly effective catalyst, providing altered selectivities and improved yield of the monoarylated product, compared to the homogeneous catalyst under identical reaction conditions, which, by itself, is a useful advance. Supported catalysts offer the potential to also recover and recycle the valuable and expensive Pd. To investigate the potential recyclability of the polymer-supported ligand in Pd-catalyzed C(sp³)—H monoarylation reactions, we used PL(4) and iodobenzene as the model ligand and substrate with DCE as the reaction solvent for ligand recovery and recycle studies (see Table 2). As expected, as a control, it was noted that the C(sp³)—H monoarylation reaction failed in the absence of Pd¹ (Run 1, Table 2). Subsequently, the reaction was run under standard conditions, using the polymeric ligand PL(4), in conjunction

Table 2. Recycling PL(4) in Pd-Catalyzed $C(sp^3)$ –H Monoarylation a,b,c

$$\begin{array}{c} \text{NPhth} \\ \text{H} \\ \text{CONHAr}_F \\ \text{1} \\ \text{Ar}_F = 4\text{-(CF}_3)\text{C}_6\text{F}_4 \\ \end{array} \begin{array}{c} \text{10 mol% Pd(TFA)}_2 \\ \text{20 mol% PL(4)} \\ \text{TFA, Ph-l} \\ \text{Ag}_2\text{CO}_3, \text{DCE} \\ \text{100 } ^0\text{C, 20h} \\ \end{array} \begin{array}{c} \text{NPhth} \\ \text{CONHAr}_F \\ \text{2} \\ \text{3} \\ \end{array} \\ \text{3} \\ \end{array}$$

Run	ligand	$Pd(TFA)_2$	yield (%) ^b	$2:3^{b}$
1	PL(4)	0	NR^c	ND^d
1*	PL(4)	10 mol %	73	70:3
2	recycled PL(4) from run 1*	0	72	70:2
2*	recycled PL(4) from run 1*	10 mol %	73	71:2
3	recycled PL(4) from run 2	0	<5	ND^d
3*	recycled PL(4) from run 2	10 mol %	40	40:0

"Substrate (1.0 equiv), $PL(TFA)_2$ (0.1 equiv), PL(4) (0.2 equiv), and PL(4) (0.2 equiv) were weighed out open to air and placed in a pressure tube (5 mL) with a magnetic stir bar. The iodobenzene (1.5 equiv), PL(4) (0.2 equiv), and PL(4) DCE were added. The reaction vessel was sealed and the mixture was first stirred at room temperature for 10 min and then heated to 100 °C for 20 h with vigorous stirring. The yield percentage and ratios of product 2 to product 3 were determined by PL(4) H NMR. PL(4) NMR = no reaction. PL(4) NDD = not determined.

with Pd. Interestingly, the used PL(4) that was collected after the first run reaction gave an identical yield of the coupled product as the fresh PL(4) in the absence of additional Pd in a recycle test (Runs 1* and 2; see Table 2). Based on the elemental analysis and 1H NMR of the reused material, we verifed that we successfully recycled PL(4) and Pd with high conversion (see Table 2, as well as Figure S7 in the Supporting Information) after recovery from the first reaction. This is, therefore, an uncommon example of ligand and metal recycling in a $C(sp^3)-H$ functionalization reaction. Although the recovered PL(4) catalyst was not robust after multiple runs (Runs 3 and 3*; see Table 2), the recovery of the ligands and Pd metal is attractive and the potential of developing more robust recyclable systems is being explored.

In conclusion, a functionalizable and tunable polymer was synthesized and used in the immobilization of pyridine ligands for use in an important Pd-catalyzed $C(sp^3)$ —H monoarylation reaction. Such $C(sp^3)$ —H activation reactions have the potential to become useful tools to construct and functionalize complex organic molecules. Notably, this is the first example of polymer-supported catalyst that selectively promotes $C(sp^3)$ —H monoarylation. The immobilized ligand showed selectivity toward less-hindered aryl iodides and was reusable with an identical catalytic yield in a second cycle.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscatal.6b01774.

Experimental procedures, ¹H NMR spectra, ¹³C NMR spectra, and other characterization data for the materials, reaction yields under additional reaction conditions (PDF)

AUTHOR INFORMATION

Corresponding Authors

*E-mail: yu200@scripps.edu (J.-Q. Yu)

*E-mail: cjones@chbe.gatech.edu (C. W. Jones).

Notes

The authors declare no competing financial interest.

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