

Ligand-Enabled *Meta*-C–H Alkylation and Arylation Using a Modified Norbornene

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Supporting Information

ABSTRACT: 2-Carbomethoxynorbornene is identified as a more effective transient mediator to promote a Pd(II)catalyzed *meta*- $C(sp^2)$ -H alkylation of amides with various alkyl iodides as well as arylation with previously incompatible aryl iodides. The use of a tailor-made quinoline ligand is also crucial for this reaction to proceed.

D irected C-H activation has been largely associated with the functionalizations of *ortho*-C-H bonds.¹ The development of a U-shaped template to direct *meta*-C-H activation has provided a promising solution to this problem.² To further improve the efficiency and scope of *meta*-C-H activation reactions, we have established a ligand-enabled catalytic system that combines Pd(II)-catalyzed *ortho*-C-H activation with norbornene-mediated Pd(II)/Pd(IV) catalysis developed by Catellani and Lautens (eq 1)^{3,4} to achieve *meta*-selective C-H



alkylation and arylation.⁵ A related *meta*-C–H arylation adopting a similar strategy has also recently appeared in literature.⁶ Prior to these developments, an elegant C-2 alkylation of indole using norbornene as the mediator has also been reported by Bach (eq 2).⁷ In our amide-directed *meta*-C–H activation reactions,⁵ there exist significant limitations. First, alkylation with alkyl iodides containing β -hydrogen is problematic (the use of 6 equiv ethyl iodide is required to give only a 21% yield). Second, aryl iodide coupling partners without ortho-coordinating groups are not compatible except for a single example with highly reactive 3,5-bis(trifluoromethyl)iodobenzene.⁵ Herein we report the development of a more efficient norbornene-type mediator that can promote efficient meta-C-H alkylation with a wide range of alkyl iodides as well as arylation with common aryl iodides in the presence of a quinoline-based ligand. This approach is fundamentally different from other meta-selective C–H activation/carbon–carbon bond forming reactions.^{8–10}

Challenges associated with C–H alkylation reactions are anticipated from the extensive studies on transition-metalcatalyzed C–C coupling involving alkyl coupling partners.¹¹ Although a few *ortho*-C–H alkylation reactions have been developed,^{12,13} directed *meta*-C–H alkylations with simple alkyl iodides have not been reported to date despite the significant synthetic importance. For example, our U-shaped templatedirected *meta*-C–H coupling is not compatible with alkyl boron reagents at this stage of development.^{2d,e} Analysis of the catalytic cycle of our norbornene-mediated *meta*-C–H arylation and methylation reactions (Figure 1) led us to focus on tuning the



Figure 1. Plausible catalytic cycle for norbornene-mediated *meta*-C–H alkylation.

reactivity of intermediates III and IV which can potentially participate in multiple reaction pathways, some of which are nonproductive. Our preliminary experimental result with ethyl iodide also suggests that the formation of the cyclobutane adduct as the major side product largely outcompetes the ethylation pathway. Reductive elimination from either III or IV could be responsible for this observation. Considering that norbornene is intimately involved in these competing steps, we envisioned that it is possible to favor the desired oxidative addition of ethyl iodide with III or aryl-ethyl reductive elimination from IV by systematically modifying the structure of norbornene. Notably, Catellani previously observed that bicyclo[2.1.1]hex-2-ene and bicyclo[2.2.2]oct-2-ene other than norbornene can also react with aryl halides to give benzocyclobutene,¹⁴ albeit without forming the Catellani reaction product.

Prior to the extensive tuning of the norbornene structure, we needed to identify a pyridine- or quinoline-based ligand that will give the optimum reactivity for C-H activation and norbornene

Received: July 14, 2015 **Published:** August 27, 2015 insertion steps leading to the formation of intermediate III. Guided by our previous extensive studies of these ligands in the *meta*-C-H arylation reaction,⁵ a brief survey of a few superior ligands rapidly identified L1 as the most effective ligand as indicated by the formation of the benzocyclobutene adduct in 85% yield (Table 1). Hence we began to prepare various norbornenes and test them in the *meta*-C-H alkylation reaction with EtI in the presence of ligand L1 (Table 1).





^{*a*}Conditions: **1** (0.1 mmol), **2a** (2.5 equiv), Pd(OAc)₂ (10 mol %), **L1** (10 mol %), Norbornene derivative (1.5 equiv), AgOAc (3.0 equiv), DCE (1.5 mL), 75 °C, air, 16 h. ^{*b*1}H NMR yields, using CH₂Br₂ as internal standard. ^{*c*}Possible side product was not observed with or without the addition of **2a**. ^{*d*}Using 0.5 equiv of **N15** instead of 1.5 equiv.

Since the high reactivity of the norbornene insertion is largely derived from the ring strain, we prepared a number of norbornenes fused with an aryl group at the 5,6-positions (N2–N4). These structural modifications significantly reduced the formation of the cyclobutane adducts 4. However, the yield of the meta-alkylation product is only moderately improved when the trifluoromethyl-group-bearing norbornene derivative N4 is used. Further tuning the ring strain by replacing the carbon at the 6 position with an oxygen significantly reduced the overall reactivity (N5). We then introduced various electron-withdrawing functional groups at the 5, 6, and 7 positions to study the electronic effects (N6-N10). A noticeable increase in yield was observed with N7 containing two ester groups. The comparison of the results with N6 and N7 is revealing. While the steric hindrance at the 5 and 6 positions of N7 seems to hamper the undesired reductive elimination pathway leading to cyclobutane adduct 4g, the norbornene insertion step (step 2) is presumably also drastically inhibited compared to N6 affording lower reactivity.

Since the preliminary screening of various readily available norbornenes did not afford significant improvement, we decided to synthesize new norbornenes to accelerate the migratory insertion (step 2) of the arylpalladium species with norbornene. Considering the superior reactivity of styrenes and acrylates in Heck coupling and C–H olefination reactions due to the polarization of the double bond,¹⁵ we introduced phenyl, acetyl, sulfonyl, nitrile, and ester groups to the 2-position in conjugation with the double bond (N11–N15). Among these modified norbornenes, N15 was identified to be the most effective mediator affording the *meta*-C–H alkylation product in 97% yield. The use of a catalytic amount of N15 (0.5 equiv) is feasible, albeit affording a lower yield (72%, see Supporting Information (SI)). The low reactivity with N12-N14 indicates that the desired reaction pathway is highly sensitive to the electronic and steric effects of the double bond. For example, the carbonpalladium bond α to keto and nitrile group formed from N12 and N14 seems to retard the meta-C-H activation (step 3). In the presence of DOAc, the fully recovered starting material containing high ortho-D-incorporation and no meta-D-incorporation supports this hypothesis (see SI). It is also important to note that the cyclobutane adduct is not formed with norbornenes N12-N15 even in the absence of EtI; presumably, the substitution of electron-withdrawing groups at the 2-position prevents the reductive elimination through both steric hindrance and electronic effect. In addition to being dependent on the structure of norbornene, this reaction is also significantly influenced by the ligands. Replacement of L1 by other phosphine and carbene ligands also decreases the yield of the alkylated product to 10% and 45% respectively (see SI).

With these high-yielding conditions in hand, we examined the scope of alkyl iodides (Table 2). Alkyl iodides with a long carbon chain afford the *meta*-alkylated products in good yields (3a-3c).





^{*a*}Conditions: 1 (0.1 mmol), 2 (2.5 equiv), $Pd(OAc)_2$ (10 mol %), L1 (10 mol %), N15 (1.5 equiv), AgOAc (3.0 equiv), DCE (1.5 mL), 75 °C, air, 16 h. ^{*b*}Isolated yields. ^{*c*}Using 3.0 equiv of N15 instead of 1.5 equiv. ^{*d*}1 (1 mmol scale). ^{*e*}Ethyl bromide (2.5 equiv), 48 h; ¹H NMR yield measured using CH_2Br_2 as internal standard.

A similar result is obtained with more sterically hindered *i*-butyl iodide (3d). Alkyl iodides containing a trifluoromethyl, aryl, benzyl, silyl protected hydroxyl, and protected amino group as well as chloro, nitrile and ketal functionalities are all excellent coupling partners rendering this *meta*-alkylation method broadly useful (3e-3l). Surprisingly, an alkyl iodide containing an ester group gives the product in a low yield (40%) under standard conditions, which is improved to 74% by using 3.0 equiv of N15 (3m). *Meta*-ethylation of 1 is also performed in 1 mmol scale to give 3a in 91% yield (Table 2). Ethyl bromide is also compatible affording 3a in 40% yield after extending the reaction time to 48 h (Table 2). Under current conditions, secondary alkyl iodides or bromides decomposed without giving the desired alkylation product.

This newly established *meta*-alkylation protocol is also extended to other amide substrates derived from phenyl acetic

Table 3. Meta-Alkylation of Substituted Phenylacetamide^{*a,b*}



^aConditions: **5** (0.1 mmol), **2b** (2.5 equiv), $Pd(OAc)_2$ (10 mol %), **L1** (10 mol %), **N15** (1.5 equiv), AgOAc (3.0 equiv), DCE (1.5 mL), 75 °C, air, 16 h. ^bIsolated yields. ^cUsing 3.0 equiv of **N15** instead of 1.5 equiv. ^dUsing **2b** (5.0 equiv), $Pd(OAc)_2$ (20 mol %), **L1** (20 mol %), **N15** (3.0 equiv), AgOAc (6 equiv), DCE (3.0 mL). ^eUsing **2b** (2.0 equiv), **L1** (20 mol %), 95 °C.

acids using butyl iodide as the coupling partner (Table 3). Substrates containing ortho-methoxy, fluoro, and chloro groups give the desired *meta*-butylated products in good yields (6a-c). Meta-methoxy, methyl, fluoro, chloro, and trifluoromethyl are also well-tolerated (6d-6h). The butylation of the para-fluoro and nonsubstituted phenylacetamides (5i, 5j) under the standard conditions give a mixture of di- and monobutylation products in a ratio of 1:1. However, the dibutylated phenylacetamides 6i-di and 6j-di are obtained as the major products in 77% and 65% yields respectively by using 20 mol % Pd catalyst and 5 equiv of butyl iodide. Reducing the butyl iodide to 2 equiv gives the monobutylated phenylacetamide 6j-mono as the main product in 44% yield. Naphthalene substrate 5k is selectively butylated at the 3-position in excellent yield (92%). Meta-alkylation of tetralone, dihydrobenzofuran, and indoline substrates also proceeds in moderate yields (6l-n), even though the amide directing group is out of plane with the ortho and meta-C-H bonds. However, extending this method to other heterocycles such as pyridines and thiophenes was unsuccessful. Our meta-alkylation protocol is also compatible with mandelic acid and phenylglycine substrates affording the desired products 60-di and 6p-di in good yields. Unfortunately, substantial racemization occurred during the installation of our amide directing group using previously reported conditions (50 is obtained in 67% ee). However, no racemization was observed in this meta-alkylation reaction when using mandelic acid derived amide **50** with 67% ee as the substrate (see SI).

The ability of this modified norbornene to enable *meta*-C-H alkylation encouraged us to revisit our previously reported *meta*-arylation reaction (Table 4).⁴ This previous protocol is only compatible with aryl iodides bearing an *ortho*-coordinating group or multiple electron-withdrawing substituents. Using norbornene **N15**, *meta*-arylation with a broad range of aryl

Table 4. *Meta*-Arylation with a Variety of Aryl Iodides^{*a,b*}



^{*a*}Conditions: 1 (0.1 mmol), 7 (3.0 equiv), $Pd(OAc)_2$ (10 mol %), L1 (20 mol %), N15 (3.0 equiv), AgOAc (3.0 equiv), $PhCF_3$ (1.5 mL), 90 °C, air, 24 h. ^{*b*}Isolated yields.

iodides proceeded smoothly, overcoming previous limitations. Phenyl iodide and other aryl iodides containing a chloride, trifluoromethyl, and ester, as well as methyl, amino, and methoxy groups, all give good to excellent yields. Notably, 5-iodoindole is also compatible affording the desired product in 57% yield.

The multiple elementary steps involved in this catalytic cycle add complexity to the investigation of the origin of the beneficial effect of this modified norbornene (Figure 1). A number of mechanistic aspects, however, are worthy of comment based on our preliminary studies. First, the stereochemistry of the carbopalladation step (step 2) is mostly likely to be *exo* as established in the stoichiometric reaction of Ph–Pd–I with norbornene.¹⁶ Second, the protonolysis of the arylpalladium bond is crucial for regenerating the Pd(II) catalyst and closing the catalytic cycle. To support our hypothesis, a reaction of 1 is performed in the presence of 10 equiv of deuterated acetic acid (see SI). The observed 70% D-incorporation at the *ortho*-position is consistent with the protonolysis pathway. The proton source



from the amide substrate and the HOAc generated from the C–H activation step could account for the 30% H-incorporation at the *ortho*-position. Trifluoromethylated amide **5h** was also subjected to the same deuterium labeling experiment to gain better separation of the aromatic protons in the NMR spectrum, which gave a similar D-labeling pattern (see SI). Finally, the impact of this modified norbornene **N15** is also illustrated in the Catellani arylation reaction. It is well-known that the Catellani *ortho*-substituents.^{3c,17} Other aryl iodides only give a series of side products containing a bicyclic structure.¹⁸ In contrast, the Catellani *ortho*-arylation followed by a Heck reaction proceeds with **N15** to give the standard product **9a–b** in good yield (eq 3).

In summary, we have developed a *meta*-alkylation reaction and significantly improved the scope of the *meta*-arylation reaction of phenyl acetamides. The design of a more reactive norbornene

analogue and appropriate choice of a quinoline-type ligand are crucial for the success of this development. The acquired insight from this study will guide further development of this emerging *meta*-C-H functionalization strategy.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.5b08914.

Experimental procedures and spectral data for all new compounds (PDF)

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Notes

The authors declare no competing financial interest.

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