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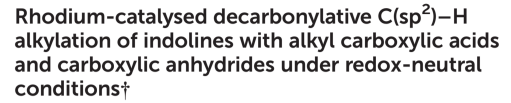


Cite this: *Org. Biomol. Chem.*, 2022, **20**, 2808

Received 4th February 2022, Accepted 16th March 2022

DOI: 10.1039/d2ob00249c

rsc.li/obc



Hirotsugu Suzuki, D Yuya Kawai, Yosuke Takemura and Takanori Matsuda D *

We developed a rhodium-catalysed decarbonylative C(sp²)–H alkylation method for indolines. This reaction facilitates the use of alkyl carboxylic acids and their anhydrides as a cheap, abundant and non-toxic alkyl source under redox-neutral conditions, featuring the introduction of a primary alkyl chain, which cannot be addressed by previous radical-mediated decarboxylative reaction. Through a mechanistic investigation, we revealed that an initially formed C-7 acylated indoline was transformed into the corresponding alkylated indoline via a decarbonylation process.

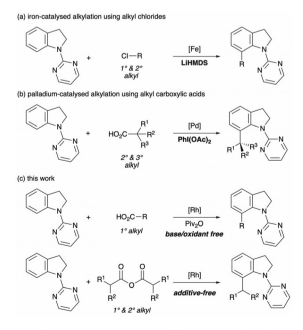
Indoles and their derivatives are ubiquitous structural motifs found in various natural products and pharmaceuticals. As such, efficient synthetic methods to access densely functionalised indoles have been extensively developed. Among them, site-selective C-H functionalisation of indoles allows the forging of new chemical bonds directly from a simple starting material for the rapid assembly of molecular complexity. However, selective activation of the C(7)-H bond is still challenging because the reactivity of the benzenoid moiety is lower than that of the pyrrole one. Consequently, ligand-directed C-7 functionalisation of indolines has emerged as a promising alternative to the corresponding functionalisation of indoles.

The C-7 alkylation has also attracted considerable attention within the field of C–H functionalisation of indolines, utilising a series of alkylating reagents: activated alkenes, diazo compounds bearing an electron-withdrawing group, cyclopropanols and aziridines. However, these alkylating reagents cannot yield indolines bearing a simple alkyl chain without using any specific functional groups. Recently, Punji *et al.* addressed this problem by using simple primary and secondary alkyl chlorides in their iron-catalysed alkylation, but its highly basic conditions might narrow the substrate scope (Scheme 1a). 12

Department of Applied Chemistry, Tokyo University of Science, 1-3 Kagurazaka, Shinjuku-ku, Tokyo 162-8601, Japan. E-mail: mtd@rs.tus.ac.jp

† Electronic supplementary information (ESI) available: Experimental procedures and characterisation data for new compounds. See https://doi.org/10.1039/d2ob00249c

Alkyl carboxylic acids have recently garnered significant attention as an alkylating reagent owing to their advantages, such as being inexpensive, abundant in nature, stable under benchtop conditions and easy to handle. Despite these advantages, only one precedent of C-7 alkylation of indolines has been reported to date: the palladium-catalysed oxidative decarboxylative alkylation of indolines with alkyl carboxylic acids (Scheme 1b). In this method, primary alkyl carboxylic acids could not be coupled with indolines because of the low stability of the primary alkyl radical species. Moreover, the oxidative conditions sometimes reduce the generality of the reaction and complicate the isolation procedure. Considering these limitations, we developed the redox-neutral decarbonylative alkylation of indolines by employing (*in situ*-formed) carboxylic anhydrides (Scheme 1c). Considering the procedure of the redox-neutral decarbonylative alkylation of indolines by employing (*in situ*-formed) carboxylic anhydrides (Scheme 1c).



Scheme 1 Transition metal-catalysed C-7 alkylation of indolines.

Inspired by previous work on decarbonylative alkylation, ¹³ we began our investigation by employing 1-(pyrimidin-2-yl) indoline (1a), propionic acid (2a) and pivalic anhydride with various rhodium salts as a catalyst (Table 1). An initial experiment was performed in 1,4-dioxane at 130 °C for 18 h in the presence of [RhCl(CO)₂]₂ (2.5 mol%), producing the desired 7-ethyl-1-(pyrimidin-2-yl)indoline (3a) in good yield (entry 1). Other rhodium catalysts, namely [RhCl(cod)]₂, [Rh(OAc)(cod)]₂, [Rh(cod)₂]BF₄, [Rh(cod)₂]OTf and RhCl(PPh₃)₃, yielded little of the desired C-7 alkylated product (entries 2-6). Among the solvents tested, 1,2-dichloroethane (DCE) was proved to be the best choice (entries 7-10). Other anhydrides, acetic anhydride, di-tert-butyl dicarbonate (Boc₂O) and the combination of Boc₂O and pivalic acid, decreased the yield (entries 11-13). Lower reaction temperature (110 °C) was inadequate for the reaction to promote (entry 14). Increasing the amount of 2a and pivalic anhydride resulted in an improved yield of the coupling product (entry 15). A control experiment showed that both the rhodium catalyst and pivalic anhydride are essential for the reaction to proceed (entries 16 and 17).

Having determined the optimised reaction conditions, we explored the decarbonylative alkylation of various alkyl carboxylic acids 2 with 1a (Table 2). 17 Acetic acid, n-octanoic acid, isovaleric acid and 3-phenylpropionic acid delivered the corresponding C-7 alkylated indolines 3b-e in 73-85% yields. Moreover, the reaction with 3-phenylpropionic acid proceeded smoothly on 1 mmol scale, showing the feasibility for a large

Table 1 Optimisation of reaction conditions

Entry	Rh catalyst	Solvent	Anhydride	Yield (%) ^b
1	[RhCl(CO) ₂] ₂	1,4-Dioxane	Piv ₂ O	69
2	[RhCl(cod)] ₂	1,4-Dioxane	Piv ₂ O	0
3	[Rh(OAc)(cod)] ₂	1,4-Dioxane	Piv ₂ O	8
4	[Rh(cod) ₂]BF ₄	1,4-Dioxane	Piv ₂ O	Trace
5	Rh(cod)2 OTf	1,4-Dioxane	Piv ₂ O	Trace
6	$RhCl(PPh_3)_3$	1,4-Dioxane	Piv ₂ O	Trace
7	$[RhCl(CO)_2]_2$	Toluene	Piv ₂ O	59
8	$[RhCl(CO)_2]_2$	DCE	Piv ₂ O	83
9	[RhCl(CO) ₂] ₂	MeCN	Piv ₂ O	24
10	[RhCl(CO) ₂] ₂	DMF	Piv ₂ O	Trace
11	[RhCl(CO) ₂] ₂	1,4-Dioxane	Ac_2O	50
12	[RhCl(CO) ₂] ₂	1,4-Dioxane	Boc_2O	31
13	[RhCl(CO) ₂] ₂	1,4-Dioxane	$Boc_2O + PivOH$	16
14^c	$[RhCl(CO)_2]_2$	1,4-Dioxane	Piv ₂ O	41
15^d	$[RhCl(CO)_2]_2$	DCE	Piv ₂ O	91(92)
16	- , ,-3-	DCE	Piv ₂ O	0
17	$[RhCl(CO)_2]_2$	DCE		Trace

^a Reaction conditions: 1a (0.2 mmol), 2a (0.24 mmol), Rh catalyst (5 mol% of [Rh]) and Piv2O (0.3 mmol) were reacted in solvent (0.5 mL) at 130 °C for 18 h, unless otherwise noted. b Yield was determined by ¹H NMR analysis using 1,2,4,5-tetramethylbenzene as an internal standard. The value in parentheses indicates isolated yield. ^c At 110 °C. ^d 2a (0.4 mmol) and Piv₂O (0.5 mmol) were used.

Table 2 Substrate scope

indolines						
Me N N	Ph N N N N N N N N N N N N N N N N N N N	Me N N N N N N N N N N N N N N N N N N N	Me N N			
3j, 94% yield	3k , 96% yield	31, 81% yield	3m, 84% yield			
MeO N N N N N N N N N N N N N N N N N N N	Me N N N	F N N N	CI			
3n, 88% yield	3o, 85% yield	3p , 89% yield	3q , 86% yield			
F N N	Me N N N N N N N N N N N N N N N N N N N	N N N N N N N N N N N N N N N N N N N	N N N N N N N N N N N N N N N N N N N			
3r , 88% yield	3s, trace	3t, 73% yield	3u, 52% yield			
Ph N N	N N N	CI CI				

^a Reaction conditions: 1 (0.3 mmol), 2 (0.6 mmol), [RhCl(CO)₂]₂ (2.5 mol%) and Piv₂O (0.75 mmol) were reacted in DCE (0.75 mL) at 130 °C for 18 h. ^b 1 mmol scale. At 150 °C.

scale synthesis (3e). Alkyl substituents bearing phenoxide, phthalimide and methyl ester moieties were tolerated, providing the corresponding alkylated indolines in moderate to good yields (3f-h). 2-(4-Methoxyphenyl)acetic acid afforded an acceptable yield of 3i. Subsequently, the substrate scope of the indoline derivatives was examined. The reaction of 2- and

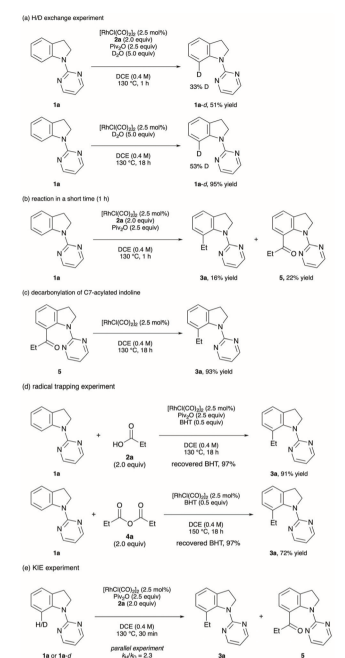
3x. 70% yield

3-substituted indolines produced the desired alkylated indolines 3j-l in 81-96% yields. 4-Methylindoline rendered the desired product 3m in high yield. Indolines bearing an electron-donating and -withdrawing group at the 5-position, including methoxy, methyl, fluoro and chloro groups, underwent the alkylation with high yields (3n-q). 6-Fluoroindoline was compatible with the reaction conditions (3r), while 6-methylindoline failed to react with 3-phenylpropionic acid. Notably, the reaction was not restricted to indolines; a 2-substituted indole and a carbazole reacted with an alkyl carboxylic acid in a similar fashion to yield 3t and 3u in 73% and 52%, respectively. Further, C-2 alkylation proceeded with 1-(pyrimidin-2-yl)indole, furnishing 73% of 3v. The decarbonylative arylation of 1a proceeded smoothly by subjecting benzoic acids to the optimised reaction conditions (3w and 3x).

Alkyl carboxylic anhydrides underwent this decarbonylative alkylation reaction in a similar manner (Table 3). The alkylation of propionic and acetic anhydrides with 1a proceeded smoothly to provide the corresponding products in moderate to high yields (3a and 3b, respectively). To our delight, branched carboxylic anhydrides participated in the reaction to furnish the products 3y-A in 71-85% yields. Application of the reaction conditions to benzoic anhydride derivative successfully led to the formation of the C-7 arylated indoline 3w.

A series of control experiments were performed to gain insight into the reaction mechanism (Scheme 2). Initially, a set of H/D exchange experiments was conducted in the presence/ absence of 2a and pivalic anhydride (Scheme 2a). The addition of D2O to the standard reaction conditions resulted in the incorporation of deuterium into the recovered starting material. Moreover, H/D exchange of 1a was observed by heating 1a, [RhCl(CO)₂]₂ and D₂O (5.0 equiv.) in DCE at

Decarbonylative alkylation using carboxylic anhydrides^a



Scheme 2 Control experiments.

130 °C. These results indicate that the C(sp²)-H bond activation step is reversible. When the alkylation reaction was halted after 1 h, a significant amount of 7-acylated indoline 5 was detected (Scheme 2b). As ketone 5 appears to be a putative intermediate in the formation of 3a, 5 was subjected to the standard reaction conditions without an alkyl carboxylic acid and pivalic anhydride (Scheme 2c). The clean conversion of 5 into 3a was confirmed, indicating that 3a arose from the catalytic decarbonylation of 5.18 In the radical trapping experiment with BHT, no significant decrease of the yield was observed in both alkyl carboxylic acid and anhydride cases, 19 suggesting

^a Reaction conditions: 1a (0.3 mmol), 4 (0.6 mmol) and [RhCl(CO)₂]₂ (2.5 mol%) were reacted in DCE (0.75 mL) at 150 °C for 18 h.

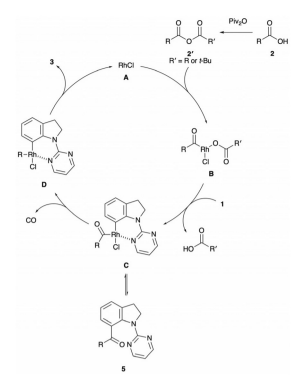


Fig. 1 Proposed reaction mechanism.

the involvement of an alkyl radical species is unlikely (Scheme 2d). Moreover, the kinetic isotope effect experiment revealed that the C-H bond cleavage step is likely to be the rate-determining step (Scheme 2e).

Based on the aforementioned experimental results and previous reports, 13,14,16 we propose the reaction mechanism depicted in Fig. 1. The reaction starts with the formation of mixed anhydride 2' followed by oxidative addition of the C-O bond to rhodium(I) A to furnish acylrhodium(III) carboxylate B. Coordination of the indoline 1 to B promotes a C-7 selective C-H activation, which might proceed via an electrophilic mechanism, 13c yielding six-membered rhodacycle C. Reductive elimination from C affords the acylated product 5. Besides, acylrhodium(III) C undergoes deinsertion of CO and subsequent reductive elimination, leading to the formation of the C-7 alkylated indole 3 along with the regeneration of active Rh (I) catalyst A.

Conclusions

In conclusion, we developed a rhodium-catalysed decarbonylative alkylation approach for indolines under redox-neutral conditions. Alkyl carboxylic acids, which are cheap, abundant and stable, function as ideal alkylating reagents with the assistance of pivalic anhydride to provide the C-7 alkylated indolines in good to high yields. Additionally, a related decarbonylative alkylation reaction with symmetrical carboxylic anhydrides is achieved under acid/base- and redox-neutral conditions. Our reaction facilitates the use of primary carboxylic

acids, which are difficult to use in the previous radicalmediated decarboxylative process, as alkylating reagents. Further investigation of decarbonylative alkylation is currently underway in our laboratory.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by JSPS KAKENHI Grant Numbers JP21K14633 and JP21K05061.

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