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Employment of a C(sp³)-based nucleophile for photoinduced palladium-catalysed cross-coupling†

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The development of efficient methods for C(sp³)-C(sp³) bond formation remains a longstanding challenge in synthetic chemistry, especially in palladium catalysis employing sterically bulky electrophiles. In this study, we present a novel approach for achieving C(sp³)-C(sp³) cross-coupling via photoinduced palladium catalysis, employing cyclopropanols as masked C(sp³)-nucleophiles. Leveraging the unique reactivity of photoexcited palladium, this protocol enables radical-mediated C(sp³)-C(sp³) coupling across a broad range of substrates including sterically hindered and functionally diverse alkyl halides under mild conditions. This method significantly expands the extent of palladium-catalysed cross-coupling for bond construction between sp³-hybridized carbon units, providing streamlined access to structurally complex C(sp³)-rich frameworks that are crucial for medicinal chemistry.

Introduction

The strategic formation of carbon-carbon bonds is one of the most important processes in the synthesis of valuable organic molecules. In this sense, transition metal-catalysed processes have gained considerable attention due to their convergence in nature and high efficiency.¹ Among many types, palladium-catalysed cross-coupling has been one of the most popularized methods in forging various types of carbon-carbon bonds due to the facile generation of the reactive palladium(0) catalyst, redox-neutrality, and synthetic robustness.² Conventional catalytic methods of this kind have predominantly relied on the use of C(sp²)-based coupling partners as nucleophiles and/or electrophiles.³ However, palladium-catalysed C(sp³)-C(sp³) coupling remains severely limited, especially in terms of employing sterically demanding electrophiles with sp³ hybridization (Fig. 1A).^{3e,f} This is mainly due to the inherent mechanistic restrictions of C(sp³)-based electrophiles, which undergo inefficient oxidative addition and compete with rapid β-hydride elimination (Fig. 1B, left).^{3e,f,4,5} Therefore, extending cross-coupling to the functionalization of C(sp³)-based building blocks has relied on the use of other reactivity categories, such as early transition metal catalysis, photoredox catalysis or electrochemical methods.⁶

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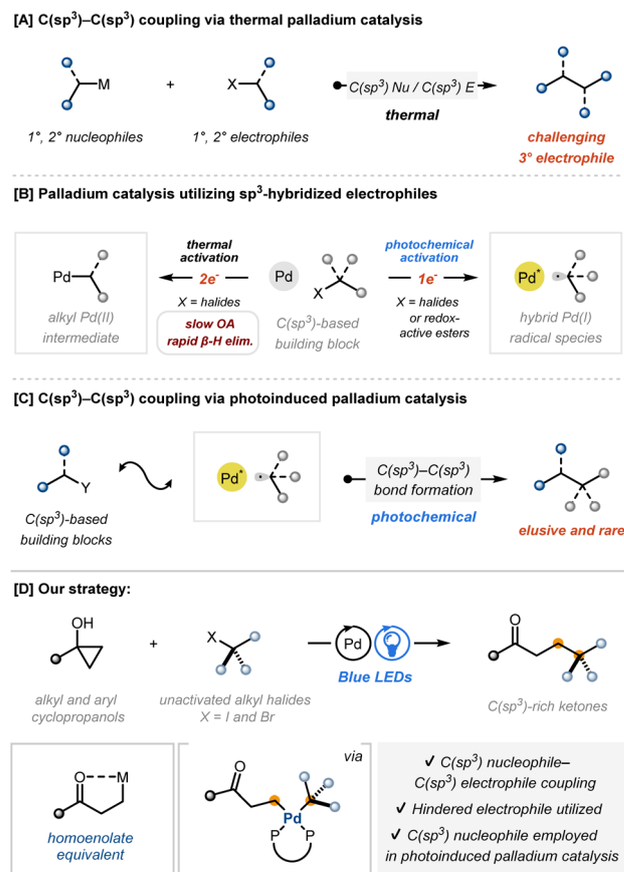


Fig. 1 State of the art for palladium-catalysed C(sp³)-C(sp³) cross-coupling.



Photoinduced palladium catalysis has recently emerged as a powerful tool to overcome the limitations of traditional methods that require the activation of C(sp³)-based electrophiles.⁷ Excited-state palladium complexes, generated by visible light irradiation, can activate C(sp³)-based precursors to form hybrid alkyl radical palladium species *via* single-electron oxidative addition (Fig. 1B, right). Subsequently, it serves as a valuable radical-based reaction partner for challenging bond-forming transformations.^{8–11} Nonetheless, the applications of photoinduced palladium catalysis for the formation of C(sp³)-C(sp³) bonds are still limited and the reactivity of hybrid alkyl radical palladium with C(sp³)-based nucleophilic coupling partners remains rare and elusive (Fig. 1C).¹²

Recognizing the critical importance of introducing sp³-hybridized carbon nucleophiles, the potential of cyclopropanols as latent homoenolate equivalents was investigated in the context of photoinduced palladium catalysis. Cyclopropanols serve as three-carbon synthons by undergoing ring-opening processes and have been utilized as valuable synthetic building blocks to accomplish the formation of various carbon-carbon bonds.^{13,14} In the present study, cyclopropanols were utilized as masked C(sp³)-based nucleophiles, facilitating the efficient formation of C(sp³)-C(sp³) bonds in conjunction with photoinduced palladium catalysis to achieve the activation of challenging electrophiles under mild reaction conditions (Fig. 1D). This approach underscores the potential of photoinduced palladium catalysis to induce an unprecedented bond formation event with a formally nucleophilic counterpart.

Results and discussion

We commenced our investigations by cross-coupling of phenylcyclopropanol **1a** with commercially available *tert*-butyl bromide **2a** in the presence of a palladium catalyst and visible light irradiation (456 nm) (Table 1).¹⁵ It has been shown that *rac*-BINAP is the optimal supporting ligand for the palladium catalyst (entries 2–5). In addition, the use of other palladium sources decreased the reaction efficiency, regardless of the initial oxidation state of palladium precatalysts (entry 6). In the case of the base additive, other inorganic bases were not as effective as cesium carbonate (entry 7), and similar observations were made with amine-based organic bases (entry 8). Additional control experiments demonstrated that all reaction components, including palladium precatalyst, base, and the irradiation of light are essential for optimal reactivity (entries 9–11). Moreover, it was found that the use of 10 mol% of palladium catalyst and 20 mol% of ligand is required for the best performance of the system (entry 12). Finally, the protocol was smoothly performed in a larger scale (entry 13).

After establishing the optimal reaction conditions, we began to evaluate the applicability of the developed method (Table 2). Initially, we investigated the functional group tolerance of our system with a variety of substituted cyclopropanols. Cyclopropanols containing an electron-donating group, such as methyl (**3ab**), methoxy (**3ac**), or dimethylamino groups (**3ad**), were found to be effective reaction partners to afford the cross-coupling products. Moreover, cyclopropanols with an electron-

Table 1 Optimization of reaction conditions

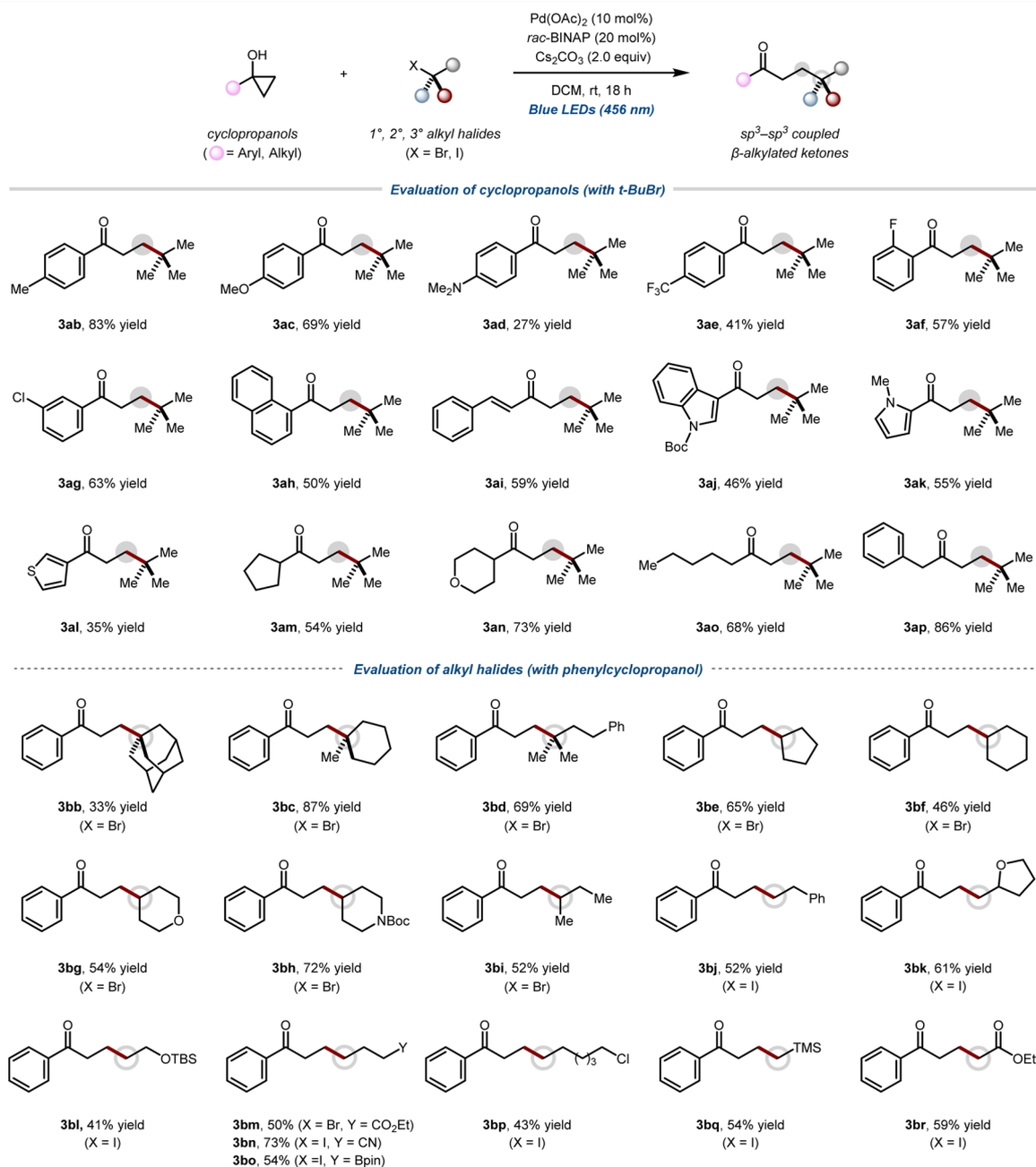
Entry ^a	Deviation from standard conditions	Yield ^b
1	None	90 (85)
2	No <i>rac</i> -BINAP	45
3	XantPhos instead of <i>rac</i> -BINAP	41
4	<i>t</i> -Bu XantPhos instead of <i>rac</i> -BINAP	15
5	DPEPhos instead of <i>rac</i> -BINAP	28
6	PdI ₂ , PdCl ₂ (PhCN) ₂ , PdCl ₂ , Pd ₂ dba ₃	74, 60, 39, 6
7	Li ₂ CO ₃ , Na ₂ CO ₃ , K ₂ CO ₃ , CsOAc, CsOPiv	30, 7, 7, 54, 36
8	TEA, DIPEA	28, 28
9	No Pd(OAc) ₂	<5
10	No Cs ₂ CO ₃	8
11	No light irradiation	<5
12	Pd/L = 5/10 mol%	53
13	2.0 mmol scale	(88)

^a Reaction conditions: **1a** (0.1 mmol), **2a** (2.0 equiv.), Pd(OAc)₂ (10 mol%), *rac*-BINAP (20 mol%), Cs₂CO₃ (2.0 equiv.) in DCM (0.6 mL), rt, N₂, 18 h, irradiated with blue LEDs (456 nm). ^b Determined by ¹H NMR using 1,3,5-trimethoxybenzene as an internal standard; isolated yields given in parentheses.

withdrawing group (**3ae**) also underwent the desired transformation smoothly. Halogen-substituted substrates (**3af**, **3ag**) were also compatible with the reaction conditions, indicating the orthogonal characteristics of the protocol with aryl halides. Cyclopropanol pronucleophiles with an extended π-system, such as naphthyl (**3ah**) or vinyl (**3ai**) moieties, also participated in the reaction with high efficiency. Notably, further addition reaction did not occur to the corresponding α,β-unsaturated carbonyl product, demonstrating the mildness of the developed protocol (**3ai**). Moreover, medicinally relevant heterocycles, including indole (**3aj**), pyrrole (**3ak**) or thiophene (**3al**), were well-tolerated during the transformations. Finally, various aliphatic cyclopropanols bearing cyclopentyl (**3am**), tetrahydropyranyl (**3an**), *n*-pentyl (**3ao**) or benzylic (**3ap**) substituents were also efficiently coupled with electrophiles, affording C(sp³)-rich ketones. In contrast, β-substituted cyclopropanols failed to participate effectively, presumably due to increased steric hindrance (ESI Table S5†).

Subsequently, we surveyed the generality of the method in terms of the electrophilic reaction partner to find that a wide range of alkyl halides — primary, secondary, or tertiary — can readily undergo the desired transformation. Various sterically hindered tertiary halides performed well in this transformation



Table 2 Photoinduced palladium-catalysed cross-coupling of cyclopropanols and alkyl halides^a

^a All reactions were performed on a 0.2 mmol scale.

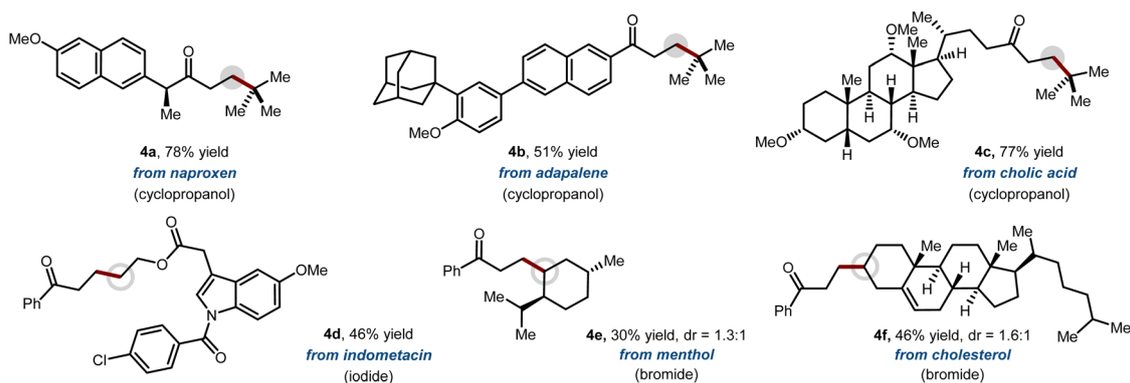
(**3bb–3bd**). Additionally, cyclic secondary alkyl halides containing cyclopentyl (**3be**), cyclohexyl (**3bf**), tetrahydropyranyl (**3bg**) or piperidinyl (**3bh**) moieties provided the cross-coupling products with good to excellent yields. Additionally, an acyclic secondary alkyl halide (**3bi**) was shown to be compatible with the protocol. Lastly, primary alkyl halides were also successfully engaged in the photoinduced palladium catalysis, demonstrating the generality of the method. Primary alkyl halides substituted with arene (**3bj**), tetrahydrofuran (**3bk**) or silyl ether (**3bl**) were viable substrates for the reaction. Also, synthetically

useful functional groups such as carboethoxy (**3bm**), cyano (**3bn**) or boronic acid pinacol ester (**3bo**) groups were shown to be compatible with the reaction conditions. Notably, alkyl substrates containing multiple carbon–halogen bonds afforded the desired product in a chemo-selective manner (**3bp**). Additionally, alkyl halides bearing α -heteroatom (**3bq**) or α -carbonyl (**3br**) substituents were efficiently coupled with cyclopropanols.

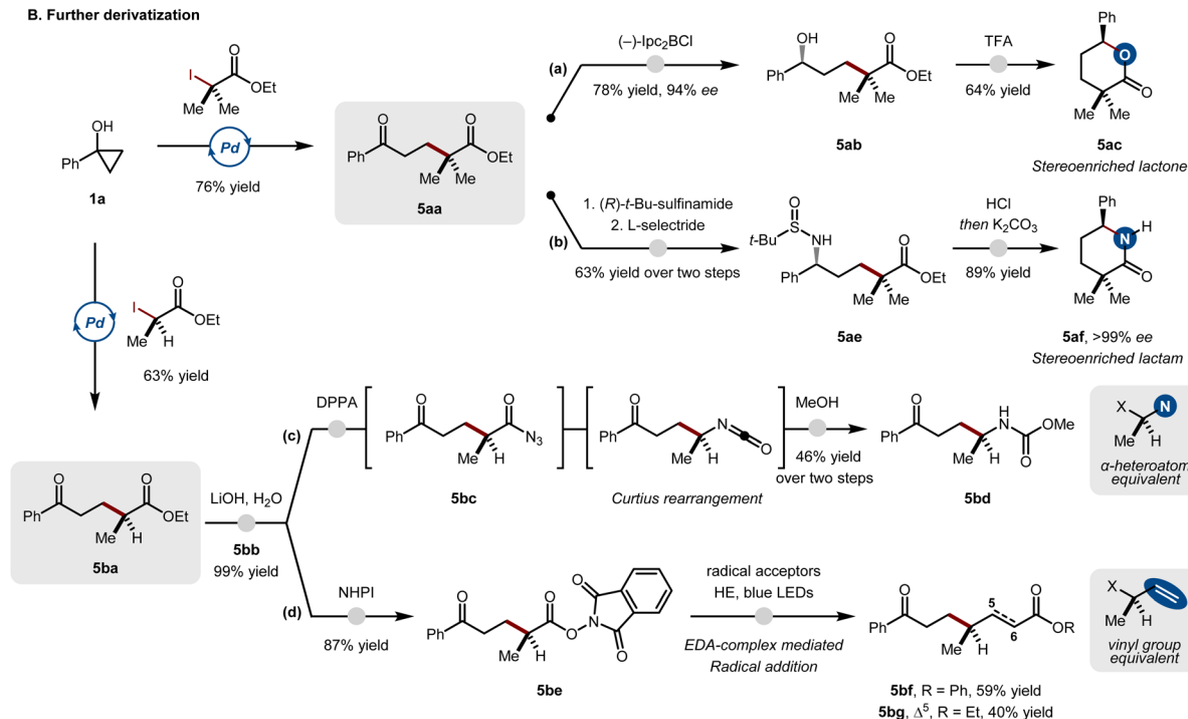
The applicability of the developed method toward complex bioactive architectures was also evaluated (Scheme 1A). A series of drug molecules and natural products, such as naproxen (**4a**),



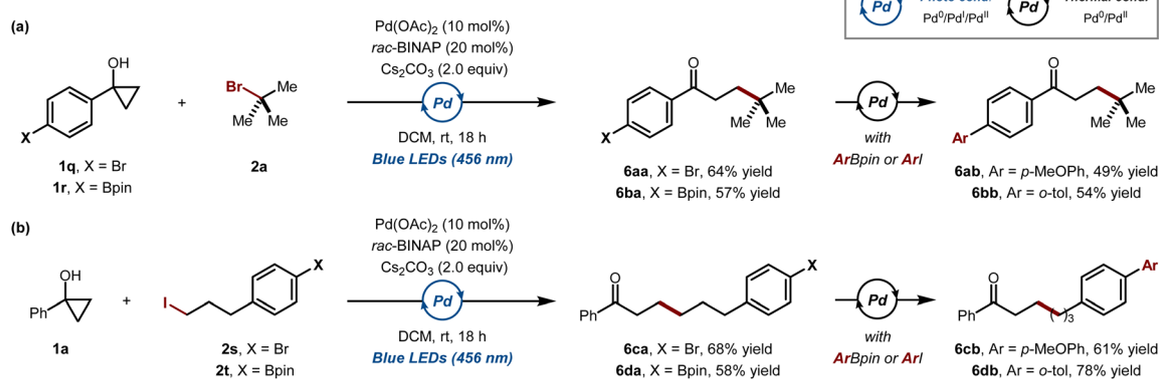
A. Complex bioactive molecule and natural product synthesis



B. Further derivatization



C. Compatibility with traditional palladium(II) cross-coupling



Scheme 1 Synthetic utility.

adapalene (**4b**) and cholic acid (**4c**), could be conveniently converted to the corresponding cyclopropanols. By subjecting the complex cyclopropanol pronucleophiles to the developed

reaction protocol, the extended ketone derivatives of the parent precursors could be successfully prepared. Additionally, complex alkyl halides derived from bioactive molecules, such as



indomethacin (**4d**), menthol (**4e**), or cholesterol (**4f**), were also proved to be competent coupling partners in this transformation, affording C(sp³)-rich architectures.

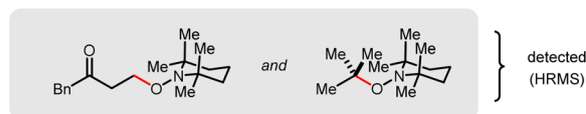
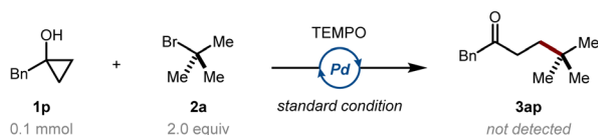
To further highlight the synthetic utility of the protocol, various downstream transformations of the β -alkylated ketone products were attempted (Scheme 1B). A substituted δ -keto ester (**5aa**), which could be conveniently prepared using the developed method, was subjected to either enantioselective carbonyl or imine reduction conditions to yield chiral δ -hydroxy (**5ab**) or δ -amino ester products (**5ae**), respectively (paths a and b). Subsequent acid- or base-mediated cyclization afforded lactone (**5ac**) and lactam (**5af**) products in enantiopure form. Through this divergent synthetic sequence, pharmaceutically important scaffolds could be readily synthesized, which would otherwise be difficult to access with conventional reactivity of ester enolates. Moreover, the carboxyl group of the product could be further exploited as a functional handle to furnish more complex molecular architectures. Another substituted δ -keto ester product (**5ba**) underwent straightforward hydrolysis to give the corresponding δ -keto acid (**5bb**). The reactive intermediate could be advanced to the amino ketone product (**5bd**) by Curtius rearrangement (path c). Alternatively, the carboxylic acid intermediate could be converted to its redox-active ester form (**5be**) to undergo a further electron-donor-acceptor (EDA) complex-mediated Giese addition (**5bf**, **5bg**) (path d).¹⁶ Importantly, through the use of carboxylate as a secondary functional handle, it was possible to construct product structures that are either difficult to access using a halide-based precursor, due to

their susceptibility to elimination (**5bd**), or resistant to undergo the desired transformation under the developed conditions (**5bg**).

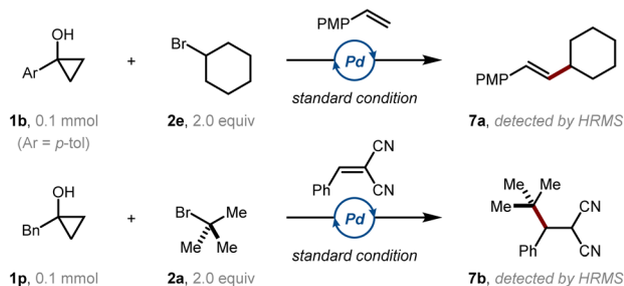
The synthetic versatility of the developed method was further examined by evaluating its chemical orthogonality (Scheme 1C). Cyclopropanols bearing either a bromine (**1q**) or a boronic acid ester (**1r**) on its aryl substituent were subjected to the developed reaction conditions to furnish the cross-coupling products (**6aa**, **6ba**) (Scheme 1C, a). The coupling products with an existing functional handle could be successfully converted into more complex biaryl ketones (**6ab**, **6bb**) by the thermally-induced palladium-catalysed C(sp²)-C(sp²) cross-coupling. Analogously, alkyl halide electrophiles containing identical functional handles could be further elaborated in a similar manner (Scheme 1C, b). These examples demonstrate that the developed method is an orthogonal strategy that can be coupled with the conventional cross-coupling reactions based on the Pd(0)/Pd(II) cycle.

To gain mechanistic insight into the developed synthetic strategy, a number of control experiments were conducted (Scheme 2). The presence of 2,2,6,6-tetramethylpiperidine 1-oxyl (TEMPO), a radical scavenger, significantly inhibited reactivity, and the cross-coupled product (**3ap**) was not formed at all (Scheme 2A). Instead, TEMPO adducts derived from both cyclopropanol and alkyl halide were observed. At this point, it can be speculated that both the formally nucleophilic homo-enolate, originating from the cyclopropanol precursor, and the alkyl counterpart participate in the catalytic cycle in the form of

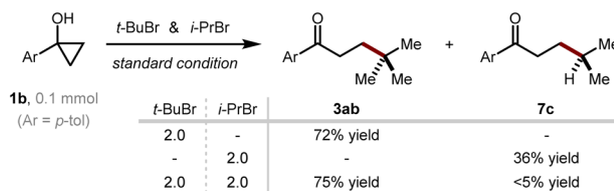
A. Control experiments with radical scavenger



B. Control experiments with radical acceptor

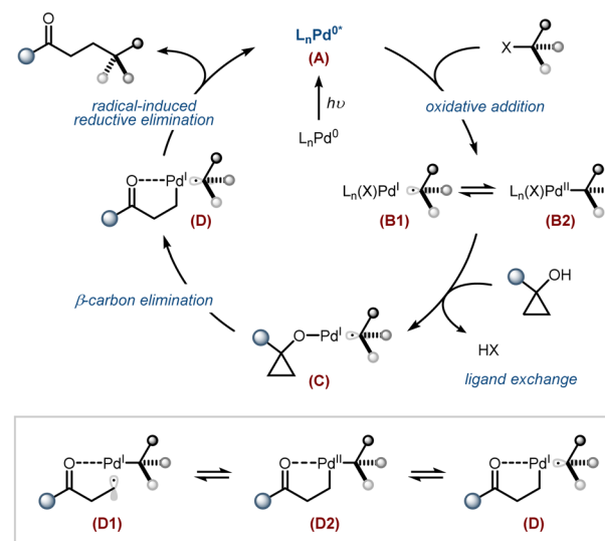


C. Competition experiment (3° vs 2° alkyl halide)^a



^a Yields were determined by ¹NMR analysis with an internal standard.

D. Plausible reaction mechanism



Scheme 2 Mechanistic studies and proposed mechanism.



radical species. Furthermore, the incorporation of electron-rich or -deficient olefins, such as *p*-methoxystyrene or 2-benzylidenemalononitrile, as radical acceptors led to the formation of radical addition products from the alkyl halide precursors (**7a** and **7b**) (Scheme 2B). In addition, competition experiments based on the simultaneous usage of the tertiary and secondary alkyl bromide substrates yielded only the cross-coupled product derived from the tertiary halide (**3ab**), suggesting the involvement of a thermodynamically more stable tertiary radical (Scheme 2C). These results reinforced the involvement of intermediary alkyl radical species.

Based on the mechanistic experiments and existing literature precedents,^{14c,d,17} a plausible reaction mechanism is depicted (Scheme 2D). Initially, alkyl halides undergo a single electron transfer mediated by a photoexcited Pd(0) complex **A** to generate hybrid alkyl Pd(i) radical **B1**, which is in equilibrium with ionic oxidative addition intermediate **B2**. Then, ligand exchange of the halide ligand with cyclopropanol results in the formation of a Pd(i) cyclopropoxide species **C**. The subsequent β -carbon elimination step produces palladium(i) homoenolate intermediate **D**. Finally, C(sp³)-C(sp³) bond-forming radical-induced reductive elimination occurs, yielding the β -alkylated ketone product and regenerating the Pd(0) catalyst. Indeed, alkyl palladium(i) complex **D** leads to the formation of alkyl palladium(ii) complex **D2** and hybrid β -keto radical palladium(i) species **D1**, which are in equilibrium with the corresponding palladium(i) intermediate, as supported by mechanistic investigations (Scheme 2A). The presence of the coupled product was detected when a bicyclic cyclopropanol was used as a mechanistic probe; nevertheless, the possibility of a radical ring-opening pathway cannot be completely ruled out at this stage (ESI Table S5†).

Conclusions

In conclusion, we have developed a photoinduced palladium-catalysed C(sp³)-C(sp³) cross-coupling reaction using cyclopropanols as masked C(sp³) nucleophiles. By strategically utilizing a photoinduced hybrid palladium alkyl radical, the challenging palladium-catalysed C(sp³)-C(sp³) coupling could be accomplished under mild conditions using a diverse range of cyclopropanols and alkyl halides, including sterically hindered tertiary halides. Notably, the method offers a streamlined route to the synthesis of structurally diverse β -alkylated ketones, including the derivatives of complex natural products and drug molecules. We anticipate that this strategy would be a milestone for further advancement in C(sp³)-C(sp³) cross-coupling in palladium catalysis.

Data availability

All data supporting this article have been included in the ESI.†

Author contributions

H. G. L. supervised and provided guidance to the project. B. R. initiated research project and B. H. K. performed the reaction

optimisation. B. H. K. and D. J. K. carried out the synthetic experiments and B. R. and B. H. K. conducted the mechanistic investigations. All the authors analysed and discussed the experimental data. H. G. L., B. H. K. and B. R. prepared the manuscript, and B. H. K. and B. R. prepared the ESI.†

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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