Photocatalytic Deoxygenative Alkylation of C(sp³)–H Bonds Using Sulfonylhydrazones

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Abstract

The ability to construct $C(sp^3)-C(sp^3)$ bonds from easily accessible reagents is a crucial, yet challenging endeavor for synthetic organic chemists. Herein, we report the realization of such a cross-coupling reaction, which combines *N*-sulfonyl hydrazones and $C(sp^3)$ -H donors through a diarylketone-enabled photocatalytic Hydrogen Atom Transfer and a subsequent fragmentation of the obtained alkylated hydrazide. This mild and metal-free protocol was employed to prepare a wide array of alkyl-alkyl cross-coupled products and is tolerant of a variety of functional groups. The application of this chemistry further provides a preparatively useful route to various medicinally-relevant compounds, such as homobenzylic ethers, aryl ethyl amines, β -amino acids and other moieties which are commonly encountered in approved pharmaceuticals, agrochemicals and natural products.

Main Text

The synthetic routes of many biologically active molecules have been dominated by the use of $C(sp^2)-C(sp^2)$ cross-coupling reactions, resulting in a significant bias towards flat molecules.¹ To improve the probability of finding new potent drugs and agrochemicals, a new trend has emerged to increase the sp³-character of new leads (so-called Escape-from-Flatland strategy).^{1,2} Consequently, there is a pressing need to develop new synthetic strategies that can realize the creation of $C(sp^3)-C(sp^3)$ bonds.³

Historically, alkyl-alkyl couplings have been challenging to accomplish, but in recent years significant progress has been achieved using transition-metal catalysis, particularly by exploiting nickel-based complexes.^{4–16} These methods rely on the availability of prefunctionalized electrophiles (e.g., alkyl halides) and nucleophilic coupling partners (i.e., organometallic reagents), and often require high catalyst loadings, which necessitate energy-intensive purification strategies to remove any trace metal impurities.¹⁵

Alternatively, an attractive option to enable $C(sp^3)-C(sp^3)$ bond formation would be to use easily accessible reagents in the absence of any transition metals, which would also facilitate applications that require higher regulatory scrutiny, such as the late-stage functionalization of medicinally relevant compounds.^{17–20} To realize this synthetic goal, we were drawn to *N*sulfonyl hydrazones, which are stable, crystalline compounds and are readily synthesized from the corresponding aldehydes (Figure 1a).^{21–24} Notably, these *N*-sulfonyl hydrazones can be exploited as a suitable electrophilic site for both polar and radical addition to yield, upon dinitrogen and sulfinate extrusion, the targeted $C(sp^3)-C(sp^3)$ bonds.^{25–32} To realize a metalfree protocol, we anticipated that a photocatalytic Hydrogen Atom Transfer (HAT) strategy would be appealing to activate aliphatic C–H bonds,^{33–45} thus generating from cheap commodity chemicals the required nucleophilic radicals which can subsequently add to the polarity-matched *N*-sulfonyl hydrazone. Herein, we describe the realization of such a photocatalytic deoxygenative alkylation of *N*-sulfonyl hydrazones using diarylketones as the photocatalyst (Figure 1b). Furthermore, the use of glyoxylate-derived sulfonyl hydrazones enables the formal α -alkylation of esters using cheap aliphatic scaffolds as radical precursors (Figure 1c). It should be noted that these compounds cannot be prepared from common silyl ketene acetals as radical acceptors due to the polarity mismatch between the electron-rich alkene and the nucleophilic alkyl radical (Figure 1c).^{46–50} Finally, we show that the application of this chemistry provides a preparatively useful route to various medicinally-relevant compounds, such as homobenzylic ethers, aryl ethyl amines, β -amino acids and other moieties which are commonly encountered in approved pharmaceuticals and natural products (Figure 1d).^{51–57}



Figure 1. 4-trifluoromethyl-phenyl sulfonyl hydrazones as practical radical acceptors to promote $C(sp^3)-C(sp^3)$ coupling. (a) Synthesis of 4-trifluoromethyl-phenyl sulfonyl hydrazones from easily to access reagents. (b) Photocatalytic generation of carbon-centered radicals add to *N*-sulfonyl hydrazones. Subsequent base-induced fragmentation delivers the targeted $C(sp^3)$ – $C(sp^3)$ cross-coupled product. (c) Our approach to realize the challenging α -alkylation of esters. (d) Bioactive compounds that bear homobenzylic ethers, arylethylamines or β -amino acids.

Our experimental investigation began with the optimization of the photocatalytic HAT step. After extensive screening of all relevant reaction parameters (see SI, section 5.1), we found that when a solution of 4-trifluoromethyl-phenylsulfonyl hydrazone 1a (1.0 equiv.) as radical acceptor, tetrahydrofuran (THF) as C-H donor in the presence of 4, 4'-dichlorobenzophenone (4-Cl₂-BP, 20 mol%) as a HAT photo-organocatalyst in trifluorotoluene (0.1 M) was subjected to 40W Kessil PR160L-390 nm irradiation, the targeted alkylated hydrazide 2a could be isolated in excellent yield (Table 1A, Entry 1, 90% yield). In the absence of either the photocatalyst or light, no product formation was observed, thus verifying the photocatalytic nature of this transformation (Table 1A, Entries 2-3). As expected, heating the reaction mixture in the dark also resulted in the quantitative recovery of **1a** (Table 1A, Entry 4). When dichloromethane was evaluated as solvent, reduced yields for 2a were observed (Table 1A, Entry 5). A high yield for hydrazide 2a was obtained when equal volumes of THF and trifluorotoluene were used (Table 1A, Entry 6, 91% yield). However, lowering the catalyst loading of 4-Cl₂-BP from 20 to 10 mol% resulted in lower yields (Table 1A, Entry 7). Also the electronic properties of the aryl sulfonyl group of the hydrazone were investigated: while the commonly used tosyl (Ar₂) and mesityl (Ar₄) variants resulted in lower yields (Table 1A, Entries 8 and 10),⁵⁸ the presence of a more electron-withdrawing ring completely shuts down the reaction (Table 1A, Entry 9). Of particular note is the practical one-pot protocol in which the hydrazone is generated in situ from 4-(trifluoromethyl)benzaldehyde (70% yield, Table 1A, Entry 11).

Having established suitable conditions for the photocatalytic event, we turned our attention to optimize the ensuing fragmentation of intermediate 2a, which would ultimately yield the targeted cross-coupled product 3^{30} A one-pot, two-step protocol was developed, in which direct addition of 3 equivalents of triethylamine and heating the reaction mixture to 80° C afforded 3 in good isolated yield (Table 1B, Entry 1). Control experiments revealed that both

the presence of a base and the supply of thermal energy were crucial (Table 1B, Entries 2-4). Moreover, decreasing the amount of base led to a reduced reaction efficiency (Table 1B, Entry 5). Surprisingly, when using the corresponding tosyl hydrazone, the denitrogenative cleavage proceeded in lower yield, further highlighting the key role of 4-trifluoromethyl-phenylsulfonyl group in both reaction steps. (Entry 6).

$\begin{array}{c} 130 \\ 1a \\ 20 \text{ equiv.} \end{array} \begin{array}{c} 130 \\ 2a \\ 1a \end{array} \begin{array}{c} 130 \\ F_{3}C \\ 2a \\ 2$	3
Entry Deviation Yield of 2 ^a Entry Deviation	rield of 3 ª
1 none 90% 1 none	90% (80%)
2 No 4-Cl ₂ -BP n.d (95% of 1a) 2 no base	traces
3 dark n.d (98% of 1a) 3 60 °C	15%
4 dark, 80 °C n.d (98% of 1a) 4 room temperature, 3 h	traces
5 DCM as solvent 55% 5 TEA (1 equiv.)	50%
6 THF:TFT 1:1 as solvent 91% 6 Ar ₂	18%
7 4-Cl ₂ -BP (10 mol%) 64% O	
	ме
9 Ar ₃ n.d. CI	
10 Ar ₄ 25% 0 02N	
11 one pot protocol (from the aldehyde I) 70% $Ar_3 = H Ar_4 = H $	

Table 1. Optimization of reaction conditions of both photocatalytic HAT (A) and fragmentation (B) steps

^{*a*}Yields were determined by ¹H NMR using trichloroethylene as external standard. 0.2 mmol scale (0.1 M). In parenthesis yields after column chromatography. See SI for experimental details.

With optimal conditions in hand (see Table 1B, Entry 1), we tested the generality of the onepot deoxygenative alkylation protocol (Figure 2). First, we combined tetrahydrofuran as C–H donor with a diverse set of aromatic aldehyde-derived 4-trifluoromethyl-phenylsulfonyl hydrazones, as shown in Figure 2. Electron-poor, -neutral and -rich aromatic aldehydes successfully served as radical acceptors, affording the corresponding homobenzylic ether products in good isolated yields (3-20, 43-86% yield). Notably, our method tolerates well the presence of halogen groups (7-12), which are useful synthetic handles for further manipulation with classical transition-metal-promoted cross-coupling reactions.¹⁵ Next, we found that a diverse range of heteroaryl aldehydes, such as picolinaldehyde (21), nicotinaldehydes derivates (22-24), indazole (25) and indole (26) were compatible with our reaction protocol. Also, several aliphatic aldehyde-derived sulfonyl hydrazones were subjected to our reaction conditions and afforded the targeted C–H alkylated products, albeit with diminished yields (34–45%, 27–29). To further demonstrate the synthetic value of our method, 4-trifluoromethyl-phenylsulfonyl hydrazones of several drugs and natural products, such as probenecid acid (32, 65%), estrone (33, 38%) and ataluren (34, 77%), were successfully engaged as coupling partners.



Figure 2. Scope of the deoxygenative alkylation of tetrahydrofuran using aromatic and aliphatic aldehyde-derived sulfonyl hydrazones. For further experimental details see the SI. Reaction conditions: hydrazone (0.4 mmol, 1 equiv.) in 4 mL of a 1:1

v:v THF:TFT (0.1 M). "Reaction time for the photochemical step: 48h. ^b1:1 d.r. ^cThe photochemical reaction was carried out in THF (4 mL).

We subsequently examined the scope of C–H donors in our synthetic process and found that a diverse set of readily available saturated heterocycles were viable for coupling with our model aryl sulfonyl hydrazone **1a** (Fig. 3). Different cyclic and linear ethers were efficiently alkylated to deliver the corresponding homobenzylic ether products in good yields (**35–42**, 41-58%). Interestingly, a masked diol (**39**, 50%) was successfully alkylated and a formal one-carbon homologation reaction of 4-trifluoromethylbenzaldehyde (**40**, 55%) was accomplished as well. Next, a series of cyclic and acyclic Boc-protected secondary amines were examined, yielding different medicinally relevant phenyl ethyl amine cores in moderate to good yields (**43–49**, 34-56%). We also found that unprotected and protected cyclic lactams as well as protected urea were viable substrates under the developed reaction conditions (**50–53**, 45-60%).

After having established this one-pot C(sp³)–H alkylation procedure, we questioned whether the instalment of an ester moiety would be feasible through use of an ethyl glyoxylate-derived sulfonyl hydrazone **1aj** (Figure 3, **54-61**). For the cleavage step, a solvent switch to ethanol proved to be essential to ensure reproducible results (see SI, section 5.2). Capitalizing on our two-step process, we discovered that various aliphatic ethers and amines could be readily coupled, delivering the corresponding products in good yields (**54–61**, 35–85%). Notably, our method provides also immediate access to various β -amino acids, common moieties in many marketed therapeutics (**56–61**, 35–76%).^{54,55}



Figure 3. Scope of the deoxygenative alkylation of different $C(sp^3)$ -H donors using 4-CF₃-benzaldehyde-, 2pyridinecarbaldehyde- or ethyl glyoxylate-derived 4-trifluoromethyl-phenyl sulfonyl hydrazones. Reaction conditions: hydrazone (0.4 mmol, 1 equiv.) in 4 mL of TFT (0.1 M). For further experimental details see the SI. ^{*a*}20 equiv. of C-H donor used. ^{*b*}1:1 *v/v* C-H donor:TFT. ^{*c*}5 equiv. C-H donor used. ^{*d*}10 equiv. of C-H donor used. ^{*e*} ¹H NMR using trichloroethylene as external standard due to volatility of **54**. ^{*f*}Solvent switch to ethanol required for the cleavage step.

Next, we performed some experiments aimed at elucidating the mechanism of this photochemical alkylative process. The involvement of carbon-centered radicals was confirmed by the presence of adduct **62**, obtained upon addition of TEMPO as radical scavenger (Figure 4A). The observed kinetic isotope effect (KIE) value of 2.5 indicates that HAT is most likely the rate determining step (Figure 4B).⁵⁹ Quantum yield measurements revealed that the reaction is photocatalytic in nature and that a radical chain is either absent or inefficient ($\phi = 0.45$, see SI, section 9.4).⁶⁰ From this experimental evidence, a plausible catalytic cycle can be derived as presented in Figure 4C. Upon absorption of a 390 nm photon, the excited state of **4-Cl₂-BP** is responsible for the cleavage of the C(sp³)–H bond, thus generating the nucleophilic alkyl

radical **63**. This transient species can undergo a polarity-matched addition to the electrophilic site of **64**, delivering a putative hydrazinyl radical **65**. At this stage, a back-HAT event from the reduced form of **4-Cl₂-BP** to **65** is envisioned, yielding the Csp³–H alkylated product **66** and closing the photocatalytic cycle.

Finally, we demonstrated the synthetic potential of ethyl glyoxylate-derived **1aj** as a useful denitrogenative ester transfer agent through the post-functionalization of compounds **57** and **58** on a 5 mmol scale (Figure 4D). Given the high relevance of therapeutic peptides containing β -amino acids,⁶¹ we showcase the straightforward preparation of dipeptide **69** through a four step sequence, starting from abundant and cheap precursors. Additionally, important motifs in medicinal chemistry, such as β -lactam **71**,⁶²⁻⁶⁴ can be prepared through deprotection of **58** and subsequent Mukaiyama reagent-induced cyclization.⁶⁵

A Radical Trapping





Figure 4. Mechanistic investigation and post-functionalization (see SI for experimental details).

In summary, we describe herein an efficient protocol to forge $C(sp^3)-C(sp^3)$ bonds from easily accessible *N*-sulfonyl hydrazones and $C(sp^3)$ -H donors via a two-step synthetic strategy,

comprising a photocatalytic Hydrogen Atom Transfer and a subsequent fragmentation reaction. We demonstrate the utility of this methodology by combining a wide range of hydrazones and $C(sp^3)$ –H bond-containing reaction partners, displaying tolerance to many functional groups. The value of this synthetic method was further shown for the preparation of medicinally-relevant compounds, such as homobenzylic ethers, aryl ethyl amines, and β -amino acids. Due to its practical and metal-free nature, we anticipate that this reaction will be of added value to those working in academia and industry. Additional studies to extend the application of this $C(sp^3)$ – $C(sp^3)$ bond-forming strategy to new synthetic contexts are underway and will be reported in due course.

Author Contributions

[†]These authors contributed equally.

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