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rsc.li/chemical-scienceVisible light-promoted ring-opening
functionalization of unstrained cycloalkanols *via*
inert C–C bond scission†‡Dongping Wang,^a Jincheng Mao[Ⓜ] and Chen Zhu[Ⓜ]*ab

Described herein is a novel, useful, visible light-promoted ring-opening functionalization of unstrained cycloalkanols. Upon scission of an inert cyclic C–C σ -bond, a set of medium- and large-sized rings are readily brominated under mild reaction conditions to afford the corresponding distal bromo-substituted alkyl ketones that are hard to synthesize otherwise. The products are versatile building blocks, which are easily converted to other valuable molecules in one-step operation. This protocol is also applicable to the unprecedented ring-opening cyanation and alkynylation of unstrained cycloalkanols.

Introduction

C–C bond activation is always a challenging issue in synthetic chemistry. Over the past few decades, this area has received great attention relying on the cleavage of cyclic C–C bonds.¹ Strained cycloalkanols such as cyclopropanols and cyclobutanols have emerged as privileged precursors for the preparation of β - and γ -substituted ketones through radical-mediated ring-opening functionalization.^{2,3} Recently, we have consecutively reported the ring-opening functionalization of cyclobutanols to construct various chemical bonds, *e.g.*, C–F, C–Cl, C–N, C–S, C–C, *etc.*, based on silver or manganese catalysis.⁴ However, the opening of unstrained rings (in particular, 5–7 and 12–15 membered rings) is always confronted with a formidable challenge. It is mainly attributed to the significantly decreased ring-strain energy (Scheme 1A).⁵ During our research, it was also found that the intramolecular dehydration took place in competition with a ring-opening pathway to suppress the reaction outcome. Therefore, an efficient catalytic protocol should be sought for the ring opening of unstrained cycloalkanols.

Photoredox catalysis has proven to be a powerful tool for the mild generation of an alkoxyl radical that triggers the subsequent ring-opening reactions or other transformations.⁶

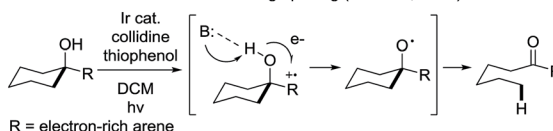
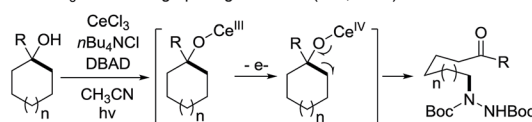
Recently, Knowles *et al.* applied intramolecular proton-coupled electron transfer (PCET) to the photocatalytic ring opening of unstrained cycloalkanols.⁷ In this remarkable process, however, electron-rich tertiary alcohols were required to generate the arene radical cation, a key intermediate for the intramolecular PCET process. This somewhat limited the practicality of the protocol (Scheme 1B). Later, Zuo *et al.* disclosed an elegant

A. Ring strain energy

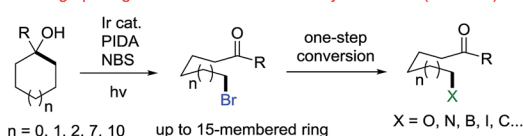
Ring Size	3	4	5	6	7	8	9	10	11	12	...	15
Strain energy (kcal/mol)	27.5	26.3	6.2	0.1	6.2	9.7	12.6	12.4	11.3	4.1	...	1.9

accomplished in this work

B. Intramolecular PCET-enabled ring opening (Knowles, ref. 7)

C. CeCl₃-enabled ring-opening amination (Zuo, ref. 8)

D. Ring-opening bromination of unstrained cycloalkanols (this work)



- Advantages:
1. Opening of medium/ large-sized rings
 2. Mild reaction conditions
 3. Broad functionality tolerance
 4. Useful and versatile products

Scheme 1 Ring-opening functionalization of unstrained cycloalkanols.

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‡ Dedicated to Professor Takahiko Akiyama on the occasion of his 60th birthday.



photocatalytic ring-opening amination of unstrained cycloalkanol using a cerium chloride complex (Scheme 1C).⁸ Beyond this, other types of ring-opening functionalization of unstrained cycloalkanol are still anticipated.

Alkyl bromides are versatile building blocks in synthetic chemistry, which can be easily converted to other valuable molecules through nucleophilic substitution or cross coupling reactions. We considered that the ring-opening bromination of unstrained cycloalkanol would give rise to distally bromo-substituted alkyl ketones which are synthetically valuable but hard to synthesize otherwise. Herein, we provide support for this hypothesis (Scheme 1D). A set of medium- and large-sized rings are readily brominated through the mild cleavage of an inert cyclic C–C σ -bond with the assistance of visible-light irradiation. The newly formed C–Br bonds can function as privileged precursors for many other useful chemical bonds. Moreover, this protocol is also applicable to the unprecedented ring-opening cyanation and alkynylation of unstrained cycloalkanol.

Results and discussion

At the outset, we implemented the reaction parameter survey with the less strained 1-phenylcyclopentanol **1a** as a model substrate and NBS as a bromine source (Table 1). It was found that the use of biphasic solvents was crucial to the reaction. In the presence of a photoredox catalyst (PC) and a hypervalent iodine reagent, the ring-opening bromination readily proceeded under visible-light irradiation, affording the desired δ -bromo alkyl ketone **2a**. A brief evaluation of solvents indicated that $\text{CCl}_4/\text{H}_2\text{O}$ delivered the best yield (entry 4), while the use of $\text{PhCF}_3/\text{H}_2\text{O}$ also resulted in a comparable yield (entry 9). The yield was slightly improved to 76% by using PC 2 instead of PC 1 (entry 13). Other than PIDA, hypervalent iodine reagents such as BI-OH, IBX, and DMP were also suitable for the reaction, but led to lower yields (entries 16–18). The amount of H_2O was important to the reaction outcome. Increasing the volume of H_2O in biphasic solvents decreased the yield (entry 19), whereas using anhydrous CCl_4 only led to trace amounts of product **2a** (entry 20). Both PC and PIDA were not indispensable; removing either one still afforded the product in modest yields (entries 21 and 22). It might be suggested that the overall ring-opening process was the outcome of two different pathways. The reaction did not proceed in the absence of both PC and PIDA or in the dark (entries 23 and 24). Finally, the yield of **2a** was further improved to 78% with the use of $\text{PhCF}_3/\text{H}_2\text{O}$ as the mixed solvent (entry 25).

With the optimized reaction conditions in hand, we set out to investigate the generality of the protocol (Scheme 2). First, a variety of cyclopentanol were tested. Generally, electron-deficient aryl substituents were preferred for the reaction, leading to δ -bromo alkyl ketones in synthetically useful yields. Halides (**2b–2d**), in particular bromides (**2d**), were well tolerated, providing a platform for the late-stage product manipulation. Positional change (*para*-, *meta*-, and *ortho*-) of substituents did not impede the transformation (**2g–2i**). Although highly electron-rich aryl compounds such as anisole

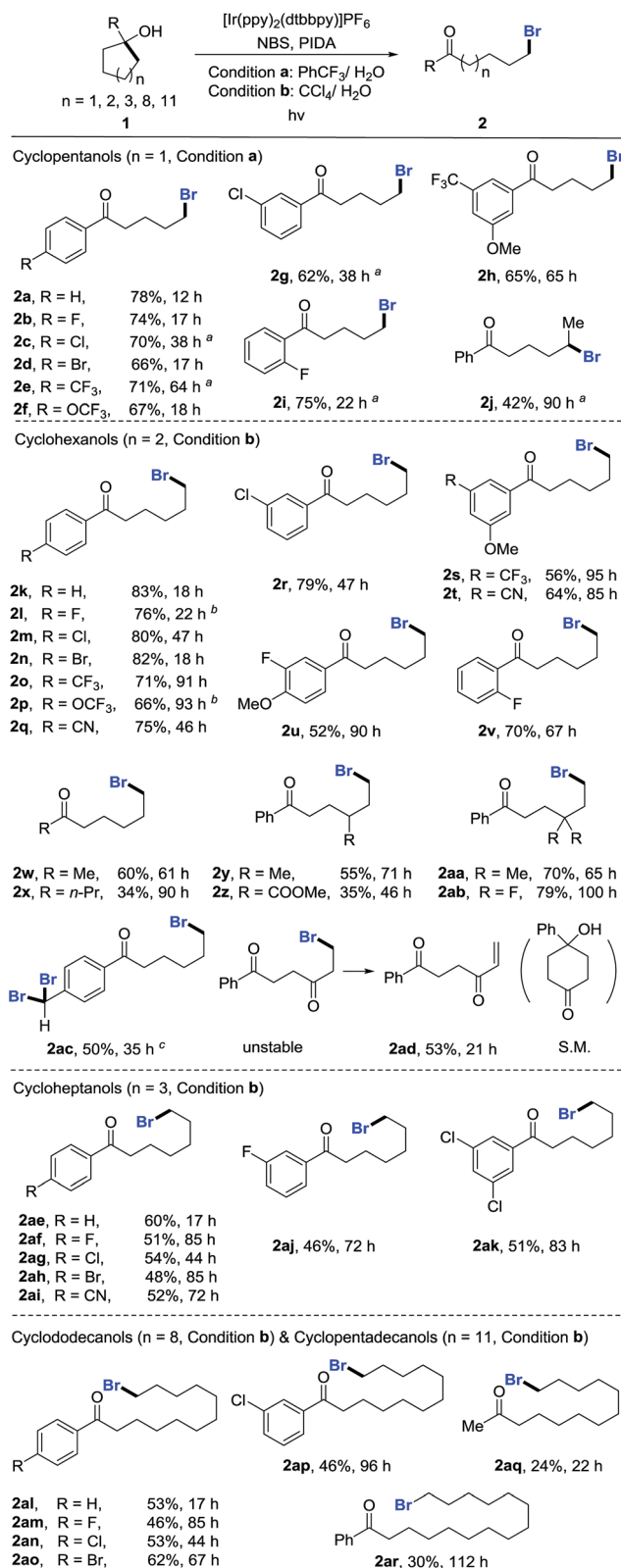
Table 1 Reaction parameter survey^a

1a + NBS $\xrightarrow[\text{Solvent, } h\nu]{\text{PC, Hypervalent iodine}}$ 2a (δ -bromo ketone)				
PC 1: $[\text{Ir}(\text{dF}(\text{CF}_3)\text{ppy})_2(\text{dtbbpy})]\text{PF}_6$; PC 3: <i>fac</i> - $\text{Ir}(\text{ppy})_3$; PC 2: $[\text{Ir}(\text{ppy})_2(\text{dtbbpy})]\text{PF}_6$; PC 4: $\text{Ru}(\text{bpy})_3\text{Cl}_2 \cdot 6\text{H}_2\text{O}$				
Entry	Solvent (v/v 20/1)	Photoredox catalyst	Hypervalent iodine	Yield (%) ^b
1	DCM/ H_2O	PC 1	PIDA	22
2	DCE/ H_2O	PC 1	PIDA	21
3	$\text{CHCl}_3/\text{H}_2\text{O}$	PC 1	PIDA	15
4	$\text{CCl}_4/\text{H}_2\text{O}$	PC 1	PIDA	74
5	DMF/ H_2O	PC 1	PIDA	0
6	DMSO/ H_2O	PC 1	PIDA	0
7	Acetone/ H_2O	PC 1	PIDA	<5
8	Toluene/ H_2O	PC 1	PIDA	<5
9	$\text{PhCF}_3/\text{H}_2\text{O}$	PC 1	PIDA	68
10	$\text{CH}_3\text{CN}/\text{H}_2\text{O}$	PC 1	PIDA	35
11	THF/ H_2O	PC 1	PIDA	0
12	MeOH/ H_2O	PC 1	PIDA	0
13	$\text{CCl}_4/\text{H}_2\text{O}$	PC 2	PIDA	76
14	$\text{CCl}_4/\text{H}_2\text{O}$	PC 3	PIDA	59
15	$\text{CCl}_4/\text{H}_2\text{O}$	PC 4	PIDA	24
16	$\text{CCl}_4/\text{H}_2\text{O}$	PC 2	BI-OH	70
17	$\text{CCl}_4/\text{H}_2\text{O}$	PC 2	IBX	55
18	$\text{CCl}_4/\text{H}_2\text{O}$	PC 2	DMP	65
19 ^c	$\text{CCl}_4/\text{H}_2\text{O}$	PC 2	PIDA	70
20	CCl_4	PC 2	PIDA	<5
21	$\text{CCl}_4/\text{H}_2\text{O}$	PC 2	—	53
22	$\text{CCl}_4/\text{H}_2\text{O}$	—	PIDA	45
23	$\text{CCl}_4/\text{H}_2\text{O}$	—	—	0
24 ^d	$\text{CCl}_4/\text{H}_2\text{O}$	PC 2	PIDA	0
25	$\text{PhCF}_3/\text{H}_2\text{O}$	PC 2	PIDA	78

^a **1a** (0.2 mmol), NBS (0.3 mmol, 1.5 equiv.), hypervalent iodine (0.4 mmol, 2.0 equiv.), and PC (0.006 mmol, 3 mol%) in mixed solvents (2.0 mL/0.1 mL) at rt, 14 W blue LED irradiation. ^b Yields of isolated products. ^c $\text{CCl}_4/\text{H}_2\text{O}$ (2.0 mL/0.2 mL). ^d In the dark.

were not suitable due to the electrophilic bromination of the aryl group, it could be overcome by mounting an additional electron-withdrawing group (**2h**). This method also allowed for the preparation of secondary bromides (**2j**). The ring opening of unsymmetric cycloalkanol proceeded in a regioselective manner. However, the synthesis of tertiary bromides failed under the current reaction conditions. The scope of cyclohexanol was investigated more extensively than that of cyclopentanol. Besides the functional groups already examined, other groups such as cyano (**2q** and **2t**) and ester (**2z**) were also compatible to the reaction conditions. In addition to aryl-substituted cyclohexanol, alkyl-substituted cyclohexanol were apt to furnish the elusive distal-bromo dialkyl ketones (**2w** and **2x**). Various substituents could be incorporated into the alkyl chain by using the functionalized cycloalkanol precursors (**2y–2ab**). In the presence of excess NBS, three-fold bromination took place on the toluene-substituted cycloalkanol to give product **2ac** which was brominated at both benzylic and distal





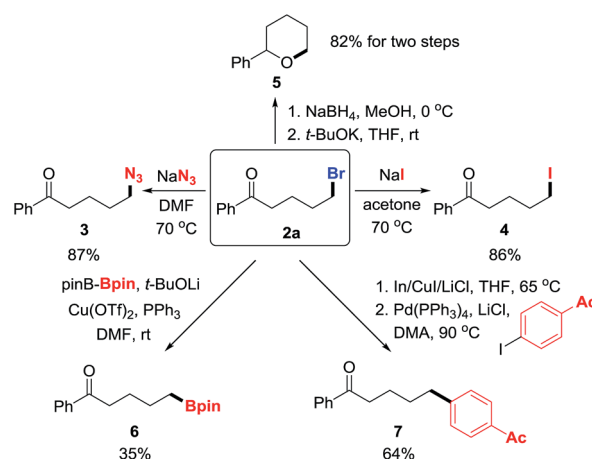
Scheme 2 Scope of unstrained cycloalkanol. Reaction conditions: **1** (0.2 mmol), NBS (0.3 mmol, 1.5 equiv.), PIDA (0.4 mmol, 2.0 equiv.), and PC (0.006 mmol, 3 mol%) in mixed solvents (2.0 mL/0.1 mL) at rt, 14 W blue LED irradiation. Yields of isolated products are given. ^aWith condition b. ^b $[\text{Ir}(\text{dF}(\text{CF}_3)\text{ppy})_2(\text{dtbbpy})]\text{PF}_6$ as PC. ^cNBS (0.8 mmol, 4.0 equiv.).

positions. The reaction with the cyclohexanol derived from 1,4-cyclohexandione afforded the unsaturated 1,4-diketone *via* dehydrobromination (**2ad**). Likewise, the ring-opening bromination of cycloheptanol bearing various functional groups also readily proceeded, affording the corresponding distal-bromo alkyl ketones in useful yields (**2ae–2ak**). Unfortunately, the ring opening of eight- and ten-membered cycloalkanol failed, leading to complex mixtures. The reason is unclear so far. Remarkably, the opening of large-sized rings such as cyclododecanol and cyclopentadecanol occurred efficiently, yielding a variety of distally brominated alkyl ketones that are hard to synthesize otherwise (**2al–2ar**).

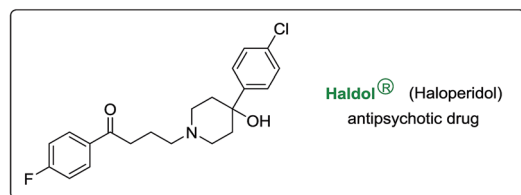
To manifest the utility of the protocol, the products were converted to other valuable molecules *via* nucleophilic substitution or cross-coupling in one or two steps (Scheme 3). First, the bromide in **2a** was easily replaced by azide and iodide, forming new C–N₃ and C–I bonds in good yields (**3** and **4**).⁹ Then, the ketone in **2a** was reduced to alcohol that intramolecularly attacked the alkyl bromide, leading to tetrahydropyran **5** in two steps with high yield.¹⁰ Alternatively, the C–Br bond was readily converted to C–B and C–C bonds *via* transition-metal catalyzed cross-coupling reactions, affording synthetically useful building blocks or molecules (**6** and **7**).¹¹

Haloperidol is a marketed antipsychotic drug extensively used to treat schizophrenia and Tourette's syndrome. After gaining a portfolio of distal-bromo alkyl bromides as precursors, we accomplished the preparation of haloperidol analogues in high yields (**8a–8d**, Scheme 4).¹² Testing of their biological activities is ongoing.

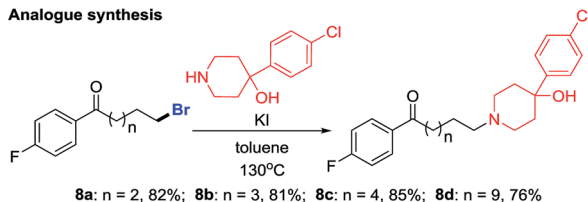
According to the experimental results, the plausible mechanism is depicted (Scheme 5, top). Although the detailed process for the generation of cycloalkoxy radical **I** is not fully understood, the radical **I** might be formed *via* two pathways. First, Stern–Volmer studies disclose that the excited state of Ir^{III} catalyst **PC 2** could be oxidatively quenched by NBS to generate the Ir^{IV} complex (see the ESI†). However, the oxidation potential of this Ir^{IV} complex ($E_{1/2}^{\text{IV/III}} = 1.21$ V in MeCN *vs.* SCE) is insufficient to oxidize cycloalkanol **1** ($E_{\text{p}/2} > 2.0$ V in MeCN *vs.* SCE) to alkoxy radical **I** *via* the SET process (for the cyclic



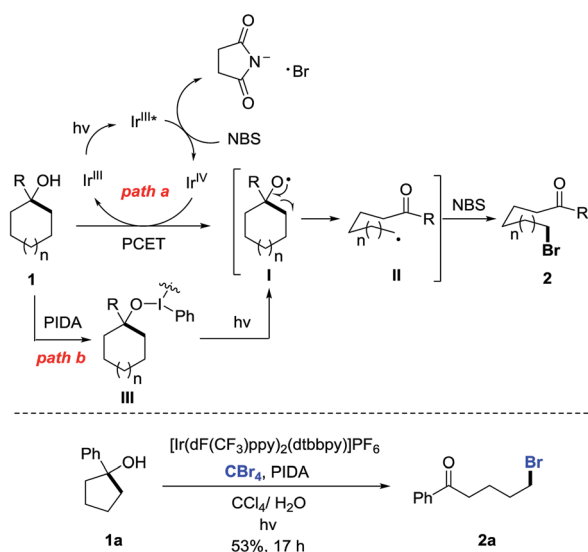
Scheme 3 Transformation of products to other useful molecules.



Analogue synthesis



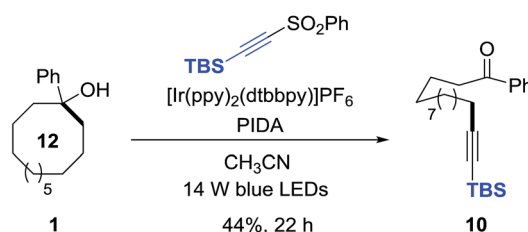
Scheme 4 Production of haloperidol analogues.



Scheme 5 Plausible pathways of ring-opening bromination.

voltammetry studies, see the ESI†). Thus, we postulate an intermolecular proton-coupled electron transfer (PCET) process for the generation of alkoxy radical **I** in the presence of an Ir^{IV} complex and weak base such as a succinimide anion (path a).¹³ Alternatively, homolysis of the O–I bond *in situ* formed by the reaction of cycloalkanol **1** with PIDA might also lead to the alkoxy radical **I** (path b).¹⁴ In either way, the visible-light irradiation is indispensable. The subsequent β -C scission of **I** leads to the ring-opened alkyl radical **II**, which is intercepted by NBS to give the final product **2**. Notably, CBr_4 is also a competent bromine source *in lieu* of NBS to afford the same brominated product, suggesting the formation of intermediate **II** during the reaction (Scheme 5, bottom).

This protocol is also applicable to the challenging ring-opening cyanation and alkynylation of unstrained cycloalkanol.¹⁵ Under the similar reaction conditions, 1-phenyl cyclododecanol was readily converted to the distally cyano- or alkynyl-substituted alkyl ketones in synthetically useful yields (**9** and **10**), providing a non-trivial approach to construct remote C–C bonds (Scheme 6).

A. Ring-opening **cyanation** of large-sized cycloalkanolB. Ring-opening **alkynylation** of large-sized cycloalkanol

Scheme 6 Ring-opening cyanation and alkynylation of unstrained cycloalkanol.

Conclusions

We have described a novel and efficient visible light-enabled ring-opening functionalization of unstrained cycloalkanol *via* the mild cleavage of cyclic C–C σ -bonds. Ring opening of a variety of cyclopentanol, cyclohexanol, cycloheptanol, cyclododecanol, and cyclopentadecanol readily proceeded to furnish the distally brominated alkyl ketones that often function as precursors of complex molecules in organic and medicinal synthesis. As shown above, this method provides important feedstocks for the production of haloperidol analogues. The newly formed C–Br bonds in products are easily transformed into other valuable chemical bonds, thus illustrating the utility of this method. This protocol is also applicable to the elusive ring-opening cyanation and alkynylation of unstrained cycloalkanol, providing an unusual strategy for the construction of remote C–C bonds.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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