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Cobalt-catalyzed annulation of styrenes with α -bromoacetic acids†

Tung T. Nguyen, *\overline{O} *ab* Bao H. T. Ngo, *ab* Huy X. Le, *ab* Linh N. P. Vu, *ab* Tuong A. To, *\overline{O} *ab* Anh N. Q. Phan* and Nam T. S. Phan* and Nam T. S

We report a method for addition of α -bromophenylacetic acids to vinyl C=C bonds in styrenes to afford γ -lactones. Reactions employed a simple cobalt catalyst $Co(NO_3)_2 \cdot 6H_2O$ in the presence of dipivaloylmethane (dpm) ligand. Many functionalities including halogen, ester, and nitro groups were compatible with reaction conditions. If α -bromoesters were used, vinylacetates were the major products.

γ-Lactones are ubiquitously found in many natural products and bioactive molecules.1 Among the available synthetic methods to obtain these five-membered rings, those using transition metals for functionalization of unsaturated carboncarbon bonds, such as alkenes and alkynes, with oxygencontaining nucleophiles have been substantially developed.² Notable reports have used scarce noble metals and/or photoredox conditions.3 Some examples employing abundant firstrow metals to obtain γ -lactones from α -bromoacetic acids or isosteric esters have been reported.4 A seminal method for copper(0)-promoted hydrocarboxylation of alkenes with α iodocarboxylates was introduced by Metzger and Mahler.4a Following that, copper(1) salts have been developed for catalytic annulation of α -bromoacetamides with styrenes. ^{4b,4c} Methods for using low valent complexes of other non-noble metals such as those of Ni(0) or Fe(0) are known as well.4d,4e It should be noted that the reported transformations were presumably triggered by a homolytic cleavage of carbon-halogen bonds in αhalocarboxylates.4d We envisioned that such an activation could be obtained if a suitable cobalt complex is used.

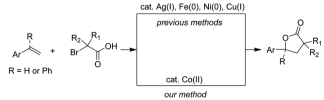
Using cobalt adducts in the organic transformations that included alkyl halides was known. 5 Most of the known examples involved the formation of cobalt($_{\rm I}$) species, thus usually suffering from the presence of strong reductants and/or tailored ligands. Meanwhile, a direct elaboration of cobalt($_{\rm II}$) complexes for incorporation of alkyl electrophiles is much rarer. It was hypothesized that a Co($_{\rm II}$)/Co($_{\rm III}$) catalytic cycle could be exploited since single-electron-transfer oxidation of Co($_{\rm II}$) was feasible. 6 Herein we report our attempts to annulate styrenes with α -bromophenylacetic acids for synthesis of γ -lactones

using a simple cobalt(II) catalyst (Scheme 1). The method has allowed for the use of α -halocarboxylic acids that are not limited to those containing quaternary centers. 4d,4e

We firstly investigated the coupling of diphenylethylene 1a and α -bromophenylacetic acid 2a to afford the γ -lactone 3aa. Brief results and control experiments are shown in Table 1.7 Optimization was carried out with respect to cobalt salt, ligand, base, and solvent. Using Co(acac)₂ as catalyst for the annulation afforded 3aa in 69% yield (entry 1). If a cobalt(III) precursor was employed, a trace amount of the lactone was detected (entry 2). Since Co(OAc)₂ was inferior to Co(acac)₂ (entry 3), our hypothesis was that 1,3-diketone may play an important role in the coupling.8 Running the reaction with Co(NO₃)₂·6H₂O catalyst and acetylacetone ligand (L1) furnished 3aa in a comparable yield with respect to that of Co(acac)₂ (entry 4). A better yield of the product was observed if dipivaloylmethane (dpm) ligand (L2) was used (entry 5). Only 27% yield of 3aa was obtained in case of an electron-poor 1,3-diketone (L3, entry 6). An unsymmetrical ligand bearing a phenyl moiety (L4) was also attempted, furnishing the lactone 3aa in moderate yield (entry 7). Sodium carbonate was an inferior base to potassium carbonate (entry 8). To our surprise, adding a stoichiometric amount of DMF helped to increase the yield of 3aa (entry 9). 1,4-Dioxane could be used as the solvent for this coupling (entry 10). Notably, a small amount of by-product derived from 2a and CH₃CN could be observed. Thus, pivalonitrile solvent was

^bVietnam National University Ho Chi Minh City, Linh Trung Ward, Thu Duc District, Ho Chi Minh City, Vietnam

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Scheme 1 Metal-catalyzed synthesis of γ -lactones \emph{via} activation of α -bromoacetic acids.

[&]quot;Faculty of Chemical Engineering, Ho Chi Minh City University of Technology (HCMUT), 268 Ly Thuong Kiet, District 10, Ho Chi Minh City, Vietnam. E-mail: R = H or Ph tungtn@hcmut.edu.vn; ptsnam@hcmut.edu.vn

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Table 1 Studies of reaction conditions^a

Entry	[Co]	Base	Solvent	Yield of 3aa (%)
1	Co(acac) ₂	K ₂ CO ₃	CH₃CN	69
2	Co(acac) ₃	K_2CO_3	CH ₃ CN	Trace
3	Co(OAc) ₂	K_2CO_3	CH ₃ CN	57
4^b	$Co(NO_3)_2 \cdot 6H_2O$	K_2CO_3	CH ₃ CN	70
5 ^c	$Co(NO_3)_2 \cdot 6H_2O$	K ₂ CO ₃	CH ₃ CN	78
6^d	$Co(NO_3)_2 \cdot 6H_2O$	K_2CO_3	CH ₃ CN	27
7^e	$Co(NO_3)_2 \cdot 6H_2O$	K_2CO_3	CH ₃ CN	52
8^c	$Co(NO_3)_2 \cdot 6H_2O$	Na ₂ CO ₃	CH ₃ CN	67
$9^{c,f}$	$Co(NO_3)_2 \cdot 6H_2O$	K_2CO_3	CH_3CN	84
$10^{c,f}$	$Co(NO_3)_2 \cdot 6H_2O$	K_2CO_3	1,4-Dioxane	75
$11^{c,f}$	$Co(NO_3)_2 \cdot 6H_2O$	K_2CO_3	(CH ₃) ₃ CCN	91
Me	Me tBu	O tBu F ₃ 0	CF ₃ M	e Ph
	L1 L	2	L3	L4

 a 1a (0.1 mmol), 2a (0.2 mmol), [Co] catalyst (0.02 mmol), ligand (0.05 mmol), base (0.1 mmol), solvent (1 mL), at 140 °C for 24 h. Yields of 3aa are GC yields using diphenyl ether internal standard. b L1 was used. c L2 was used. d L3 was used. e L4 was used. f DMF (0.1 mmol) was added. Abbreviation: acac = acetylacetonate.

attempted to avoid such addition (entry 11), affording 91% yield of the lactone 3aa.

Scope of the reaction with respect to reagents containing C=C bonds was next studied. The results are shown in Scheme 2. Notably, some products were obtained in excellent diastereoselectivities in comparison with two examples reported by Nishihara and Iwasaki. 4e A triphenyl-substituted γ-lactone 3aa was isolated in 85% yield. The simplest styrene coupled with α bromophenylacetic acid to furnish 3ba in 81% yield. Functionalities including bromo (3ca), chloro (3da), ester (3ea), and nitro (3fa, 3ga) groups were compatible with reaction conditions. Some vinyl arenes containing polyaromatics were also attempted. 2-Vinyl naphthalene was reacted to give the lactone 3ha in 88% yield. Cyclization of 9-vinylanthracene yielded a trace amount of the desired product, somewhat implying the importance of steric effect. Heterocycles, such as thiazole (3ia), were competent substrates. Other α-bromoarylacetic acids such as those containing chloro substituents could successfully annulate with C=C bonds. Thus, the lactones (3hb, 3ac) were obtained in moderate to good yields. At this moment, very electron-poor styrenes (i.e. 3,5-bis(trifluoromethyl)styrene) or internal alkenes (i.e. indene) were not tolerant of reaction conditions.

To show the generality of our method, annulation of styrenes with α -bromoisobutyric acid 4 was attempted. The conditions

Scheme 2 Annulation of styrenes and α -bromo arylacetic acids. Conditions: 1a-1j (0.5 mmol), 2a-2c (1.0 mmol), Co(NO₃)₂·6H₂O (0.1 mmol), L2 (0.25 mmol), K₂CO₃ (0.5 mmol), DMF (0.5 mmol), (CH₃)₃CCN (5 mL), at 140 °C for 24 h. Yields are isolated yields. Diastereomeric ratios (d.r.) were determined using ¹H NMR spectrum.

were relatively milder than those of α -bromophenylacetic acids, presumably due to the presence of the quaternary carbon that help cyclization step. Scope of styrenes is presented in Scheme 3. An excellent yield of the lactone (5a) was obtained in case of diphenylethylene substrate. Reactions of molecules containing bromo (5c), ester (5e), and nitro (5f) substituents occurred without any difficulties. Similar to the iron-catalyzed method, ^{4e} 2-vinyl naphthalene was suitable for the annulation (5h). Vinyl heteroarenes were competent substrates (5i). Electron-rich arenes were also attempted, furnishing the lactones in good yields (5k, 5l).

Using α -bromoacetates to couple with styrenes could yield lactones. ^{4d,4e} However, only vinylacetates **7a–7c** were obtained in our conditions if similar substrates were employed (Scheme 4). The results were indicative of a slow hydrolysis during reaction course. It should be noted that the presence of quaternary carbons in esters was crucial to afford the addition products.

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Scheme 3 Annulation of styrenes and α -bromoisobutyric acid. Conditions: styrenes (0.5 mmol), 4 (1.0 mmol), Co(NO₃)₂·6H₂O (0.1 mmol), L2 (0.25 mmol), K₂CO₃ (0.5 mmol), (CH₃)₃CCN (5 mL), at 100 °C for 24 h. Yields are isolated yields.

Scheme 4 Scope of α -bromoacetates. Conditions: 1a (0.5 mmol), α -bromoacetates 6a-6c (1.0 mmol), Co(NO₃)₂·6H₂O (0.1 mmol), L2 (0.25 mmol), K₂CO₃ (0.5 mmol), DMF (0.5 mmol), (CH₃)₃CCN (5 mL), at 140 °C for 24 h. Yields are isolated yields.

Based on previous studies, ^{4d,4e} a plausible mechanism was proposed as that shown in Scheme 5. Homolytic cleavage of C(sp³)-Br bond in 2a would afford the benzyl radical 8. Radical addition to 1a generated 9 followed by an oxidation that could afford a benzylic cation 10. The resulting species underwent

cyclization, in the presence of base, to eventually furnish the lactone **3aa**. To support the hypothesis, some control experiments were carried out. Firstly, addition of radical quenchers such as TEMPO or BHT stopped the annulation, somewhat confirming the involvement of radical species during the

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Scheme 5 Mechanistic considerations

reaction course. Secondly, running the reaction in absence of **1a** gave an anhydride **11** (detected by GC-MS). Such the dead-end intermediate could be obtained *via* a homo-combination of the radical **8** followed by a dehydration.

Conclusions

In conclusion, we have developed a general method to couple styrene C=C bonds with α -bromoarylacetic acids to afford γ -lactones. The transformation employed a simple cobalt precursor $\text{Co(NO}_3)_2 \cdot 6\text{H}_2\text{O}$ facilitated by dipivaloylmethane ligand. Reaction conditions were well compatible with a diversified array of functional groups. Preliminary control experiments supported a radical-involved mechanism that could triggered by the cobalt-catalyzed activation of carbon-bromide bond.

Conflicts of interest

There are no conflicts to declare.

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