

Selective Intermolecular Oxidative Cross-Coupling of Enolates

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Supporting Information

ABSTRACT: Selective intermolecular oxidative crosscoupling of enolates, which is a bond-forming reaction between carbanion equivalents, remains as an unsolved issue despite its potential utility for the direct synthesis of unsymmetrical 1,4-diones. The main difficulty derives from the unavoidable homo-coupling. Our strategy depends on the selective one-electron oxidation of one enolate to afford an electrophilic carbonyl α -radical species, followed by trapping with another enolate. The present study demonstrates the selective oxovanadium(V)-induced cross-coupling between boron and silyl enolates.

O xidative cross-coupling of carbon-centered nucleophiles leads to the formation of a new carbon-carbon bond, which is a complementary method to the conventional nucleophile-electrophile coupling (Scheme 1a). However, such cross-coupling of congeneric species is particularly challenging because of the lack of selectivity caused by the concomitant homo-coupling reaction.¹ More specifically, if the reaction is governed by only statistics, the ratio of homo-, cross-, and homo-coupling products is 1:2:1. Therefore, the development of versatile and selective intermolecular oxidative cross-coupling will definitely offer a useful strategy in retrosynthetic analysis of organic synthesis.

Our focus is on the intermolecular oxidative cross-coupling reaction of two different enolates (Scheme 1b). This reaction can provide a direct method to synthesize unsymmetrical 1,4dicarbonyl compounds, which are found in naturally occurring products and medicinal compounds.^{2,3} The first intermolecular oxidative cross-coupling was demonstrated between silvl enolates in 1975.⁴ However, not much work has been done for over four decades, even including not only enolates also enamines as an enolate species.⁵⁻¹⁹ The typical tactic relies on the stoichiometric advantage of one coupling partner (2-10fold), whereas the yields generally remain about 50-80%.^{4-12,14,17,18} We have also reported the oxo- and imidovanadium(V)-induced intermolecular cross-coupling of silyl enolates.^{10,15} Reactivity differences also enable enolate species to undergo cross-coupling. For example, the higher reactivity (lower oxidation potential) of enamines compared with silyl enolates induces their cross-coupling.9 Furthermore, enamine intermediates formed from aldehydes and a chiral amine catalyst oxidatively couple with silyl enolates asymmetrically.^{14,18} The use of an α -stannyl carbonyl derivative in the presence of cerium(IV) oxidant also induces the cross-coupling reaction with silvl enolates.¹¹ The reactivity difference between imide enolates with a chiral auxiliary and ketone or ester Scheme 1. (a) Oxidative Cross-Coupling; (b) Intermolecular Oxidative Cross-Coupling of Enolates; (c) Our Strategy for the Intermolecular Oxidative Cross-Coupling of Enolates



enolates as well as that between amides and ketone enolates provides practical yields (52-73%).^{2,13} It should be emphasized that this research is one of the few examples^{2,13,15,19} of equimolar cross-coupling to date. On the other hand, ketone enolate species are among the toughest combinations to cross-couple because of their similar reactivities,^{4,6-12,16,17,19} where an auxiliary cannot be employed in contrast to the cases of esters, amides, and imides.¹³ Selective cross-coupling of equimolar lithium enolates has been achieved by a strategy utilizing the selective heteroaggregation of lithium enolates by the steric effect, which is another one of the few examples of equimolar cross-coupling.¹⁶ A sophisticated solution with an intramolecular traceless silicon tether of two different enolates has been developed.^{3,20-22} However, the key bis(silyl) enolate

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requires several steps to prepare and the yields are moderate, which hampers convenience.

In this context, the present research was undertaken to develop the cross-coupling of ketone enolates with equimolar use of each coupling partner. Our working hypothesis depends on the distinction of roles for two enolates using different metals (M and M'). More specifically, this strategy includes two key steps: (1) selective one-electron oxidation of one enolate by a transition-metal oxidant M''^{n+} to give a Lewis acidactivated electrophilic carbonyl α -radical species²³ and (2) its radical addition with silvl enolate $(M' = SiR_3)$, which is known to behave as a radical acceptor^{9,11,14} (Scheme 1c). On the other hand, we have studied the oxovanadium(V)-induced homocoupling of boron enolates.²⁴ Our preliminary study suggested that boron enolate 1a is more reactive than silvl enolate 2a in the homo-coupling reaction and that the employed oxovanadium(V) oxidant leaves silvl enolate 2a intact (Figure S1). Furthermore, stereoselective preparation of boron enolates has been established via trapping of the corresponding enolates²⁵ or hydroboration of α,β -unsaturated carbonyl compounds.²⁶ Therefore, boron enolates were chosen as the partner of silvl enolates. Here we report the intermolecular oxidative cross-coupling between equimolar boron and silyl enolates by VO(OEt)Cl₂ to give unsymmetrical 1,4-dicarbonyl compounds with high selectivity.

The investigation was initiated with the cross-coupling of boron enolate 1a and silyl enolate 2a using $VO(OEt)Cl_2$ (Scheme 2a), which is a Lewis acid with one-electron oxidation capability.²⁷ Boron enolate 1a was prepared by hydroboration

Scheme 2. (a) Oxidative Cross-Coupling of Boron Enolate 1a and Silyl Enolate 2a; (b) Selected Region of the ¹H NMR Spectrum of the Crude Material after the above Cross-Coupling Reaction without THF for 20 h; (c) Gram-Scale Reaction



of chalcone with 9-borabicyclo[3.3.1]nonane (9-BBN) dimer or a tetrahydrofuran (THF) solution of 9-BBN to give the corresponding (Z)-enolate 1a quantitatively. $VO(OEt)Cl_2$ (1.25 equiv based on the sum of 1a and 2a) was added to a CH₂Cl₂/THF solution of equimolar amounts of substrates 1a and 2a under nitrogen at room temperature, and then the reaction mixture was stirred for 30 min. To our surprise, this reaction afforded the desired cross-coupling product 3aa selectively in 94% yield with a small amount of the homocoupling product 4a resulting from boron enolate 1a, while the homo-coupling product 5a resulting from silyl enolate 2a was not observed in the ¹H NMR spectrum of the crude materials (molar ratio of cross-coupling product 3aa and homo-coupling products 4a and 5a = 98:2:0). Furthermore, the same reaction without THF gave 3aa with excellent selectivity in 94% yield. Small amounts of the compounds derived from protonation of both enolates 1a and 2a were observed as byproducts. Tracking the reactions with and without THF revealed that the reaction under the conditions with THF is much faster than that without THF but that the selectivity is a little lower (Figure S2). A longer reaction time (20 h) under the conditions without THF resulted in a yield of 96% with high selectivity. Scheme 2b shows the selected region of the ¹H NMR spectrum of the crude material. The obvious peaks for the homo-coupling product 5a were not observed, but a small amount of the homo-coupling product 4a and protonated species 6 and 7 were found. It should be noted that this reaction is applicable to almost gram-scale reaction without severe losses in yield and selectivity (83% isolated yield, 3aa:4a:5a molar ratio = 99:1:0; Scheme 2c).

On the other hand, the use of lithium enolate **1a-Li** instead of boron enolate **1a** led to contrasting results (Figure S3): only a trace amount of cross-coupling product **3aa** was obtained, but homo-coupling product **4a** was formed in 31% yield. This selective homo-coupling of **1a-Li** in the presence of silyl enolate **2a** may be explained by aggregation of the lithium enolate,¹⁶ which may facilitate its homo-coupling reaction.

This cross-coupling reaction depends on the oxidant (Table S1). The use of VO(OPr-*i*)₂Cl, which has lower oxidation capability and Lewis acidity,²⁸ gave rise to a low yield with low selectivity (51% yield, **3aa:4a:5a** molar ratio = 59:43:1). Other typical oxidants, including $[Ce(NO_2)_6](NH_4)_2$, FeCl₃, and CuCl₂, were not effective for the reaction, which may be partly due to their low solubility in CH₂Cl₂.

The scope of both boron and silvl enolates was explored with the above-optimized conditions (Table 1). Here the yields of cross-coupling products 3 are isolated yields (unless otherwise noted), and the selectivities are based on the molar ratios obtained from the ¹H NMR spectra of the crude materials. Cross-coupling of boron enolate 1a and 4-fluorophenylsubstituted silvl enolate 2b instead of 2a took place in 99% yield with high selectivity (entry 1). Substituted boron enolate 1b also selectively cross-coupled in 97% yield (entry 2). Boron enolates 1c-f with a methyl group at the 2-position were employed for the oxidative cross-coupling with 2a and 2b to give good yields and selectivity (entries 3-10). The use of boron 1-cyclohexylprop-1-en-1-olate 1g as an example of an aliphatic-substituted substrate afforded the cross-coupling products in good yields (77% for 3ga and 84% for 3gb) with high selectivity (entries 11 and 12). In the case of 2c as an example of an aliphatic silyl enolate, the oxidative crosscoupling with boron enolate 1a took place to give 3ac in moderate yield and selectivity at -35 °C (entry 13). 1-Styryl1

Table 1. Oxidative Cross-Coupling of Boron Enolate 1 and Silyl Enolate 2^a

	$R^{1} + R^{2} + R^{5} + R^{4} -$	VO(OEt)Cl ₂ (1.25 equiv based on the sum of 1 and 2) CH ₂ Cl ₂ , N ₂ , rt, 20 h	$ \begin{array}{c} $	$R^2 R^3$ Ph + R^4 $R^5 R^4$ 4 5	\mathcal{R}^{5} \mathcal{R}^{4} \mathcal{R}^{6} O
Entry	Boron enolate 1	Silyl enolate 2	Cross-coupling product 3	Yield of 3 (%) ^b	Molar ratio ^c 3 : 4 : 5
1	1a B	OSiMe ₃ 2b F	Ph Bn O O O O O O O O O O O O O O O Ph	3ab : 99	99 : 1 : 0
2	Me 1b OMe	2a	Me MeO	3ba : 97	99 : 1 : 0
3 ^{<i>d,e</i>}	\Box		0	3ca : 97	96:4:0
4 ^e	B-O	20	Ph	3da: 97	93:7:0
5'		28	R ⁷	3ea: 88	93:7:0
6^r				3fa: 79	93:7:0
7 ^{0,0}	R ⁷ 1d : R ⁷ = Me		o r	3CD: 98	95:5:0
of	1e : R ⁷ = Cl	2b		30b : 97	92:8:0
9 10 ^f	1f : $R' = CF_3$		_₽ ⁷ ^U ^O	3fh 72	95.5.0 93.7.0
11 ^f	O'B	2a 2b	H or F	3ga: 77 3gb: 84	99 : 1 : 0 99 : 1 : 0
	1g		<u> </u>		
13 ^{g,h}	1a	OSiMe ₃	O Ph Bn O Bn O	3ac : 71 ^{<i>i</i>}	83 : 17 : 0
14 ^j	B Ph B Bn Bn Bh	2a	Ph Bn O Ph	3ha : 57	98 : 2 : 0
15	O-B Ph 1i	2a	Ph Ph	3ia : 85	99 : 1 : 0
16 ^k	1a	OSi(<i>t</i> -Bu)Me₂ ∕∕OEt	Ph Bn O O	3ad : 91 ^{<i>i</i>}	97 : 3 : 0
17 ^k	1g	2d		3gd : 63 ⁱ	93 : 7 : 0
18	1f	OSiMe ₃	F ₃ C Ph	3fe : 75 (<i>dr</i> = 71 : 29)	98 : 2 : 0 [/]
19 ^{g,h}	1a	OSiMe ₃	Ph Ph Bn O	3af : 85 ⁱ (<i>dr</i> = 56 : 44)	94 : 6 : 0

^{*a*}The 1:2 molar ratio was 1.0 unless otherwise mentioned. ^{*b*}Isolated yields. Yield of $3 = 100\% \times \text{moles}$ of 3/moles of 1 (or 2). ^{*c*}Determined by ¹H NMR analysis of the crude materials. ^{*d*}The reaction time was 17 h. ^{*c*}A small excess amount of 1 was employed (1:2 $\approx 1.15:1$). ^{*j*}The reaction time was 18 h. ^{*g*}A small excess amount of 2 was employed (1:2 = 1:1.1). ^{*h*}The reaction temperature was $-35 \degree \text{C}$. ^{*i*}Determined by ¹H NMR analysis of the crude material based on an internal standard (1,3,5-trimethoxybenzene). ^{*j*}VO(OEt)Cl₂ (1.1 equiv relative to each enolate) was used. The reaction time was 0.5 h, and THF (33 equiv relative to 1h) was added. ^{*k*}VO(OPr-*i*)₂Cl (1.1 equiv relative to each enolate) was used. The reaction time and temperature were 24 h and $-35 \degree \text{C}$, respectively, and THF (33 equiv relative to 1) was added. ^{*k*}VO(OPr-*i*)₂Cl (1.1 equiv relative to 9) was determined by isolation of 3fe and 4f.

substituted boron enolate **1h** showed different reactivity, and the addition of THF was effective in this case to give the product **3ha** in moderate yield with high selectivity (entry 14). Introduction of two methyl groups at the 2-position of the boron enolate did not cause a problem in the yield and selectivity, and the corresponding cross-coupling product **3ia** having a quaternary carbon was obtained (entry 15). Up to here, selective cross-coupling was demonstrated in the combination of substituted boron enolates and unsubstituted silyl enolates at the 2-position. In the oxidative cross-coupling of ketone and ester enolates using VO(OEt)Cl₂, the homocoupling product of the silyl enolate derived from the ester was observed because of the higher reactivity. Therefore, the conditions were reoptimized, and three key points were found: (1) the use of VO(OPr-i)₂Cl, (2) reaction with THF, and (3) low temperature $(-35 \ ^{\circ}C)$. Consequently, the cross-coupling of boron enolate 1a and silvl enolate 2d gave the γ -keto ester 3ad in 91% yield with high selectivity (3ad:4a:5d molar ratio = 97:3:0; entry 16). The use of an aliphatic boron enolate, 1g, was not a problem for the cross-selectivity (entry 17). Finally, as an example of the most challenging combination, crosscoupling between ketone enolates monosubstituted at the 2position (1f and 2e) was carried out. Notably, the crosscoupling proceeded with high selectivity (75% vield, 3fe:4f:5e molar ratio = 98:2:0; entry 18). Such selectivity is considered to be controlled by the difference in the reactivities of boron and silyl enolates. Oxidative cross-coupling of boron enolate 1a and 1-(trimethylsiloxy)cyclopentene (2f) at -35 °C gave 3af in 85% yield with 3af:4a:5f molar ratio = 94:6:0 (entry 19).

To gain insight into the mechanism, a radical clock reaction was investigated using boron enolate 1j with a cyclopropyl group at the 2-position (Scheme 3). Boron enolate 1j was

Scheme 3. Radical Clock Reaction



treated with VO(OEt)Cl₂ in the presence of silyl enolate **2a** and afforded the ring-opened product **8** in 83% yield; the crosscoupling product **3ja** was not detected. This result suggests the formation of carbonyl α -radical species. The detailed mechanism needs more investigation (we try to show a plausible reaction path in Figure S4), but the expected strategy shown in Scheme 1c is considered to operate in this crosscoupling.

In conclusion, the oxovanadium(V)-induced oxidative crosscoupling between boron and silyl enolates was achieved in high yields with high selectivity. One of the keys for this reaction system is the choice of the combination of boron and silyl groups as metals for the enolates, providing a reactivity difference. Scale-up to almost a gram scale was performed. The substrate scope revealed that substituted boron enolates from ketones and silyl enolates can be employed for selective crosscoupling. The developed selective intermolecular oxidative cross-coupling of enolates will bring significant benefit for the straightforward synthesis of unsymmetrical 1,4-dicarbonyl compounds.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.5b05058.

Figures S1–S5, Table S1, synthetic procedures, characterization data, and NMR charts of the products (PDF)

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Notes

The authors declare no competing financial interest.

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