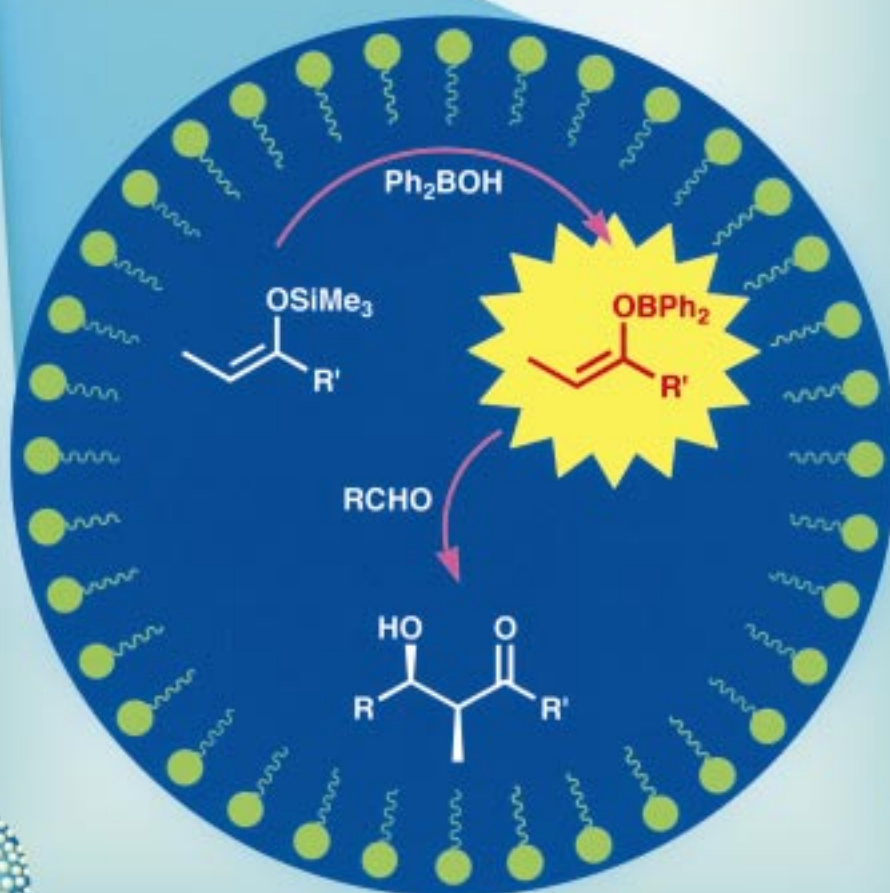


Diastereoselektive Aldolreaktionen mit Borenolaten konnten in Wasser mit einer katalytischen Menge einer Borverbindung erfolgreich durchgeführt werden.



Die Reaktion findet in kolloidalen Partikeln aus den Substraten und einem Tensid statt, sodass die Hydrolyse der Borenolate verhindert werden kann. Mehr dazu auf den folgenden Seiten.

Catalytic Use of a Boron Source for Boron Enolate Mediated Stereoselective Aldol Reactions in Water**

Yuichiro Mori, Kei Manabe, and Shū Kobayashi*

Boron enolate chemistry has been well-established and several useful reactions using boron enolates have been developed. However, stoichiometric amounts of boron sources such as dialkylboron triflates have been required in all cases so far reported. Herein, we disclose the first example of the catalytic use of a boron source in boron enolate mediated stereoselective aldol reactions.^[1] Interestingly, the reactions proceed in water without using any organic solvents.

Recently, we reported the use of Lewis acid surfactant combined catalysts (LASCs) in organic synthesis in water.^[2] LASCs create colloidal particles of approximately 1 μm diameter with organic materials in water, and useful synthetic reactions such as aldol, allylation, Mannich, and Michael reactions were shown to proceed smoothly in the presence of a LASC. In the course of our investigations to improve the reaction system, we tested various compounds which might act as better catalysts in water. We chose the Mukaiyama aldol reaction as a model reaction. After screening various catalysts, we found diphenylborinic acid (Ph_2BOH , **1**),^[3,4] which is stable in water, was an effective catalyst in the presence of sodium dodecyl sulfate (SDS) as a surfactant.^[5] We then examined a reaction of benzaldehyde with the silyl enolate **2** in water (Table 1). While the reaction proceeded very slowly in the presence of 0.1 equivalents of **1** and SDS (Table 1, entry 1), a dramatic improvement of the yield was observed when benzoic acid was added to this system (entry 2). To our surprise, the diastereoselectivity (*syn/anti*) of the product was much higher than that of the previously reported LASC system.^[6] In the presence of SDS only (entry 3), or in the presence of benzoic acid and SDS (without **1**, entry 4), the desired adducts were obtained in low yields with low diastereoselectivities. In the presence of **1** and benzoic acid (without SDS), high *syn* selectivity was obtained, albeit the yield was low (entry 5). The best yield and selectivity were obtained when **1** (0.1 equiv), SDS (0.1 equiv), and benzoic acid (0.01 equiv) were used at 0 °C (entry 6). Note that the use of water as a solvent is essential in this reaction. The reaction proceeded sluggishly in organic solvents such as dichloromethane and diethyl ether (entries 7, 8). Much lower yield than that in water was obtained under neat conditions (entry 9).

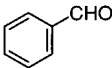
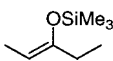
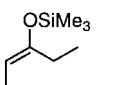
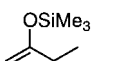
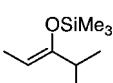
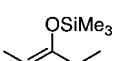
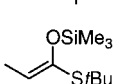
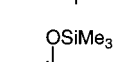
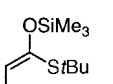
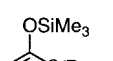
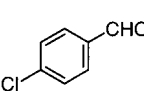
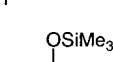
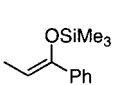
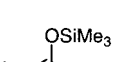
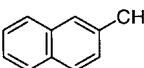
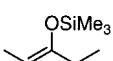
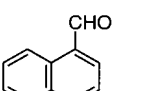
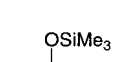
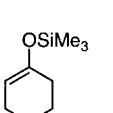
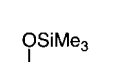
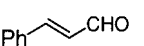
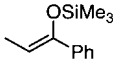
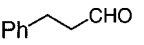
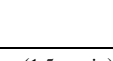
Table 1. Mukaiyama aldol reaction using **1** in water.

Entry	Solvent	PhCO_2H [equiv]	Yield [%]	<i>syn/anti</i>
1	H_2O	–	trace	–
2	H_2O	0.01	90	92/8
3 ^[b]	H_2O	–	2	53/47
4 ^[b]	H_2O	0.01	27	58/42
5 ^[c]	H_2O	0.01	4	91/9
6 ^[d]	H_2O	0.01	93	94/6
7	Et_2O	0.01	trace	–
8	CH_2Cl_2	0.01	trace	–
9	–	0.01	24	90/10

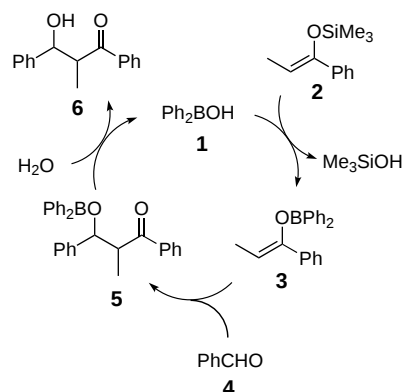
[a] Sodium dodecyl sulfate. [b] Without **1**. [c] Without SDS. [d] 0 °C.

Several examples of the aldol reactions are shown in Table 2. In all cases, the reactions proceeded smoothly in the presence of **1** (0.1 equiv), SDS (0.1 equiv), and benzoic acid (0.01 equiv) to afford the corresponding adducts in good yields. When (*Z*)-3-trimethylsiloxy-2-pent

Table 2. Mukaiyama aldol reactions using **1** in water.^[a]

Entry	Substrate	Yield [%]	syn/anti
1 ^[b]		 ^[e] 60	96/4
2 ^[b]		 ^[f] 72	53/47
3 ^[b]		 ^[g] 72	95/5
4 ^[b,c]		 ^[h] 62	96/4
5 ^[b,c]		 ^[i] 84 ^[k]	39/61
6 ^[b]		 ^[j] 51	95/5
7		 ^[j] 92	90/10
8 ^[b]		 ^[j] 74	97/3
9		 ^[j] 80	92/8
10		 ^[j] 79	47/53
11 ^[d]		 ^[j] 76	91/9
12		 ^[j] 61	92/8

[a] Conditions: silyl enolate (1.5 equiv), **1** (0.1 equiv), PhCO₂H (0.01 equiv), SDS (0.1 equiv), H₂O (aldehyde: 167 mM), 30 °C, 4–24 h. [b] Silyl enolate (3.0 equiv). [c] 0 °C. [d] PhCO₂H (0.1 equiv). [e] Z/E = >99/ <1. [f] Z/E = 19/81. [g] Z/E = 96/4. [h] Z/E = 2/98. [i] Z/E = 97/3. [j] Z/E = >99/ <1. [k] 72 h.



Experimental Section

A typical experimental procedure: aldehyde (0.25 mmol) and silyl enolate (0.375 mmol) were added successively at 30 °C to a stirred white suspension of diphenylborinic acid (0.025 mmol), benzoic acid (0.0025 mmol), and SDS (0.025 mmol) in water (1.5 mL). After 24 h, saturated aq. NaHCO₃ (5 mL) and brine (5 mL) were added, and the mixture was extracted with ethyl acetate, dried over Na₂SO₄, and concentrated. The aldol product was purified by preparative thin layer chromatography (TLC; SiO₂, ethyl acetate/hexane = 1/3).

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- [4] Ph₂BOH was used for the generation of a boron enolate from ethoxyacetylene, see: M. Murakami, T. Mukaiyama, *Chem. Lett.* **1982**, 241–244.
- [5] Anionic surfactants are effective, see the Supporting Information.
- [6] When a Lewis acid (for example, Sc(O₃SOC₁₂H₂₅)₃) was used in water, the diastereoselectivity was *syn/anti* = 49/51 (substrates: aldehyde **4** and silyl enolate **2**), see: refs. [2a, i].
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