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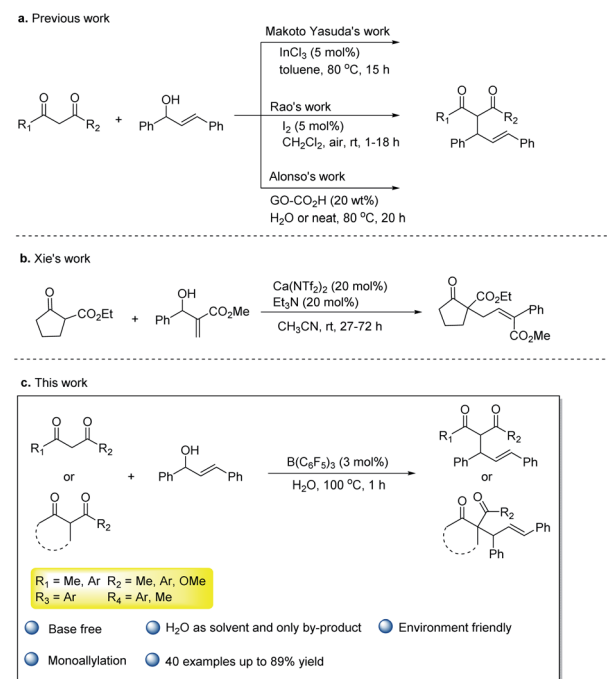
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Introduction

The Tsuji–Trost reaction is an important tool for C–C bond construction. According to the different nucleophiles of the substrates, numerous types of reactions have been reported such as allylic alkylation,¹ allylic arylation,² allylic amination,³ or *O*-allylation.⁴ The allylation of activated methylene compounds is a typical type of the Tsuji–Trost reaction,⁵ and the corresponding allylation products are very useful for the synthesis of a variety of valuable products.⁶ In tradition, the transition metal-catalyzed substitution reaction of activated methylene compounds and allylic alcohols or activated allylic alcohol derivatives with a base was widely reported,⁷ and the diallylated compounds were often detected.⁸ In this type of the reaction, the direct Tsuji–Trost reaction of activated methylene compounds with allylic alcohols would be more atom- and step-economical than activated allylic alcohol derivatives.⁹ In a previous research study, those reactions based on transition metals such as Pd¹⁰ or Pt,¹¹ and the key intermediates were π -allyl metal species. Lewis acid¹² such as InCl₃, FeCl₃, Bi(OTf)₃, LnOTf (Ln = La, Yb, Sc) or Bronsted acid¹³ such as *p*-toluenesulfonic acid have also been reported to catalyze the process efficiently. In addition, iodine,¹⁴ hexafluoroisopropanol,¹⁵ and graphene oxide CO₂H¹⁶ as the metal-free catalysts were used in the allylation of 1,3-dicarbonyl compounds with allylic alcohols (Scheme 1a). Recently, a calcium system¹⁷-catalyzed allylation of the cyclic β -ketone esters with MBH alcohols in MeCN was developed by Xie¹⁷ *et al.* (Scheme 1b). This process can efficiently construct a quaternary carbon. Despite these

developments, some problems such as long reaction time, using metal catalysts or unsafe organic solvents are still remaining. Developing green and economic allylation of activated methylene compounds is of great interest.

Water is a solvent that is more environmental-friendly than organic solvents in green chemistry. However, most of the catalysts used in organic solvents are often incompatible with water, and the low solubility of common organic compounds in

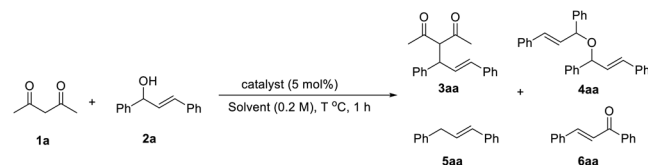


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Table 1 Optimization of reaction conditions^{a,b}

Entry	Catalyst	Solvent	$T/\text{ }^\circ\text{C}$	3aa (%)	4aa (%)	5aa (%)	6aa (%)
1	—	H_2O	100	—	Trace	—	—
2	$B(C_6F_5)_3$	H_2O	100	86	—	—	—
3	$B(3,4,5\text{-}F_3\text{-}C_6H_2)_3$	H_2O	100	56	22	8	6
4	$B(2,4,6\text{-}F_3\text{-}C_6H_2)_3$	H_2O	100	49	19	10	8
5	$CuSO_4\cdot 5H_2O$	H_2O	100	43	27	—	—
6	$FeCl_3$	H_2O	100	Trace	22	8	5
7	$Fe(OTf)_3$	H_2O	100	Trace	62	11	12
8	$CoCl_2\cdot 6H_2O$	H_2O	100	Trace	16	—	—
9	$SnCl_2\cdot 2H_2O$	H_2O	100	Trace	Trace	—	—
10	$Cu(OTf)_2$	H_2O	100	Trace	52	—	—
11	$TsOH$	H_2O	100	—	64	—	—
12	H_3BO_3	H_2O	100	—	15	—	—
13	$B(C_6F_5)_3$	$MeCN$	100	82	Trace	Trace	Trace
14	$B(C_6F_5)_3$	DCE	100	83	—	—	—
15	$B(C_6F_5)_3$	Cyclohexane	100	79	—	—	—
16	$B(C_6F_5)_3$	$EtOH$	100	Trace	—	—	—
17	$B(C_6F_5)_3$	1,4-Dioxane	100	—	13	—	—
18	$B(C_6F_5)_3$	H_2O	30	39	26	—	—
19	$B(C_6F_5)_3$	H_2O	50	57	12	—	—
20	$B(C_6F_5)_3$	H_2O	70	73	Trace	—	—
21	$B(C_6F_5)_3$	H_2O	90	78	Trace	—	—
22 ^c	$B(C_6F_5)_3$	H_2O	100	89	—	—	—
23 ^d	$B(C_6F_5)_3$	H_2O	100	76	—	—	—

^a Reaction conditions: **1a** (30.0 mg, 0.3 mmol), **2a** (63.0 mg, 0.3 mmol), and catalyst (5 mol%) in solvent (1.5 mL) for 1 h at $T\text{ }^\circ\text{C}$. ^b Yield of the isolated product. ^c The catalyst was 3 mol%. ^d The catalyst was 1 mol%.

water makes its practical use highly problematic. Thus, a catalyst that activates the substrate and is stable in air and water needs to be developed.

Tris(pentafluorophenyl)borane¹⁸ is widely used in the reduction,¹⁹ dehydrogenation,²⁰ addition reaction,²¹ or other reactions because of non-toxicity, stability and good water solubility. Moreover, Stephen²² came up with the concept of frustrated Lewis pairs in 2006. $B(C_6F_5)_3$ also became the darling of the frustrated Lewis pairs with its strong electrophilicity and large space resistance. Piers²³ (hydrosilylation) and Stephen²⁴ (catalytic hydrogenation) have carried out numerous pioneering work on metal-free catalytic reduction to make $B(C_6F_5)_3$ a versatile catalyst. However, the strong Lewis acidity of $B(C_6F_5)_3$ could cause the B–C bond to undergo preconized hydrolysis in water. The interaction between borane with different groups and water has been comprehensively explored.²⁵ The presence of water would affect the catalytic effects of $B(C_6F_5)_3$, and it limits the development of $B(C_6F_5)_3$ or frustrated Lewis pairs chemistry involved. Previously, there have been some developments in frustrated Lewis pairs chemistry that $B(C_6F_5)_3$ could catalyze reactions with a tiny amount of water.²⁶ Recently, some examples were reported about the complex formed by $B(C_6F_5)_3$ combined with water that could catalyze the reactions. For example, Tang *et al.*²⁷ reported that $B(C_6F_5)_3$ catalyzed α -diazo

ester insertion into the O–H bond in water to afford numerous α -hydroxy esters in 2018. Our group has reported the coupling reaction of naphthoquinones with indoles catalyzed by $B(C_6F_5)_3$ in water,²⁸ and the allylation reaction of electron-rich aromatic rings with allyl alcohols catalyzed by $B(C_6F_5)_3$ and 2,6-lutidine realized the coupling of the sp^2 hybrid C–H bond with allyl alcohols in water.²⁹ These successful examples of $B(C_6F_5)_3$ catalytic reactions in water have greatly aroused our enthusiasm in this field. We have already known that the allyl alcohols can be activated by $B(C_6F_5)_3$ to form allyl carbocation in water. Also, 1,3-dicarbonyl compounds as the classical nucleophiles, which were activated by $B(C_6F_5)_3$, have also been reported in previous literature.³⁰ Based on these studies, we proposed the allylation of 1,3-dicarbonyl compounds and allylic alcohols by $B(C_6F_5)_3$ catalyzed in water, and this approach would be an efficient and environmentally friendly way (Scheme 1c).

Results and discussion

Acetylacetone **1a** and (*E*)-1,3-diphenylallyl alcohol **2a** were selected as the model substrates to investigate the allylic reaction under numerous reaction conditions (Table 1). Initially, the reaction was carried out in different catalysts, $100\text{ }^\circ\text{C}$ for 1 h in water. First, the reaction was largely impossible without any



catalyst (entry 1). Moreover, to our delight, $B(C_6F_5)_3$ has catalyzed the allylic reaction as well as we thought, 86% yield of product **3aa** was obtained (entry 2). $B(3,4,5-F_3-C_6H_2)_3$ or $B(2,4,6-F_3-C_6H_2)_3$ demonstrated activities in this transformation, and the yields of **3aa** were just moderate. Compared with $B(C_6F_5)_3$, the Lewis acidity of $B(3,4,5-F_3-C_6H_2)_3$ or $B(2,4,6-F_3-C_6H_2)_3$ is weaker, which leads to the decrease in the yield of the desired products and the formation of by-products (entries 3 and 4). In addition, $CuSO_4 \cdot 5H_2O$ had a partial catalytic effect in the reaction (entry 5), other metal Lewis acid or non-metallic proton acid had a little catalytic effect on the generation of **3aa** (entries 6–12). Except for $B(C_6F_5)_3$, other catalysts with weaker Lewis acidity could produce the bis(1,3-diphenylallyl) ether **4aa** and **5aa**, **6aa** as by-products in this condition. Further screening of solvents showed that the solvent had some effects on the reaction. When the reaction was performed in CH_3CN , DCE or cyclohexane, **3aa** could be obtained in good yields (entries 13–15). When using EtOH and 1,4-dioxane as solvents, we did not detect any **3aa** in the condition (entries 16 and 17). The water did not affect the reaction. Moreover, when the temperature was adjusted to 30 °C, the yield of the expected product was decreased significantly and that of the by-product bis(1,3-diphenylallyl) ether increased (entry 18). With the increase in temperature, the by-product was reduced (entries 19–21). In addition, we also tried to decrease the amount of the catalyst to 3 mol% and the yield could be maintained (entry 22), and when the amount of catalyst to 1 mol% was decreased, the yield of the product was reduced (entry 23). At last, the best conditions for the reaction were determined: **1a** (0.3 mmol), **2a** (0.3 mmol), $B(C_6F_5)_3$ (3 mol%), H_2O (1.5 mL), 100 °C, 1 h.

Next, the applicability of the protocol for a broad range of substrates was explored. As shown in Table 2, for almost active methyl compounds of 1,3-diketones and β -ketone esters, they could perform the allylic reaction with corresponding product yields of more than 80% (**3aa**–**3ca**, **3ja**–**3oa**). The yield of the ring of active methylene compounds such as 1,3-cyclohexanedione was 75% (**3da**), and the yield of the product was not significantly affected by the ring tension. When the substrates were ring active methine compounds, the yield of the allylic product was moderate to high (**3ea**–**3ha**, **3pa**–**3sa**). The steric hindrance of the active methine compounds did not affect the yield of the allylic reaction in these substrates interestingly. In contrast, (*E*)-3-cinnamylpentane-2,4-dione was reacted with **2a** to give the allylic product in moderate (**3ia**), but the yield of methyl-(4*E*)-2-acetyl-5-phenyl-

4-Pentenoate was only 23% (**3ta**). It may be that the presence of two ketone carbonyl in (*E*)-3-cinnamylpentane-2,4-dione increases the reaction activity. Interestingly, 3-cyano-4-chromanone **3ua** afforded the corresponding product in good yield. When the compound **3aa** was employed, the corresponding diallylated product was not obtained. Moreover, the compound **3aj** also failed. It may be caused by the steric hindrance. Most of the products have two chiral centers, and we obtained a mixture of diastereoisomer. The dr value was obtained by NMR. However, unfortunately, the dr value was not high.

The reaction between numerous substituted allylic alcohols and acetylacetone was then examined. As depicted in Table 3,

Table 2 Substrate scope for 1,3-diketones and β -ketone esters^{a,b}

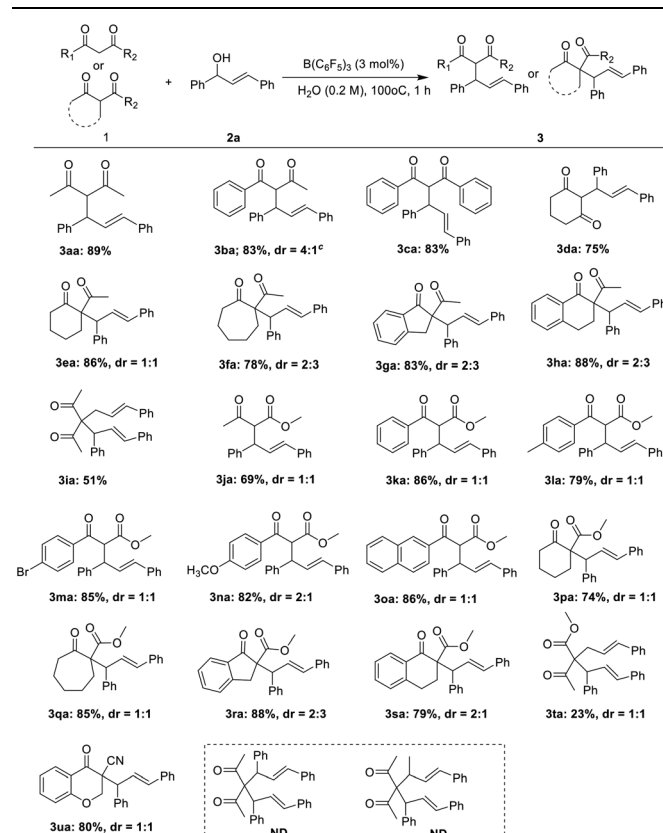
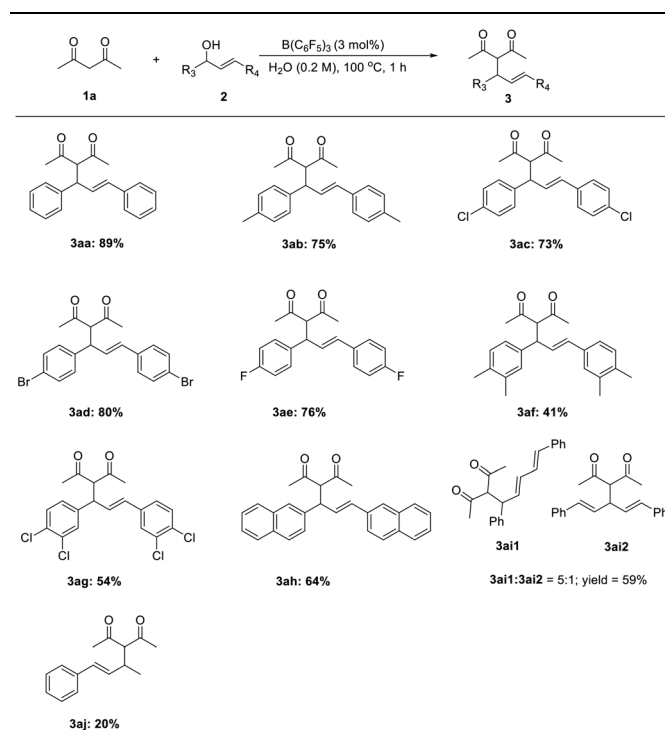
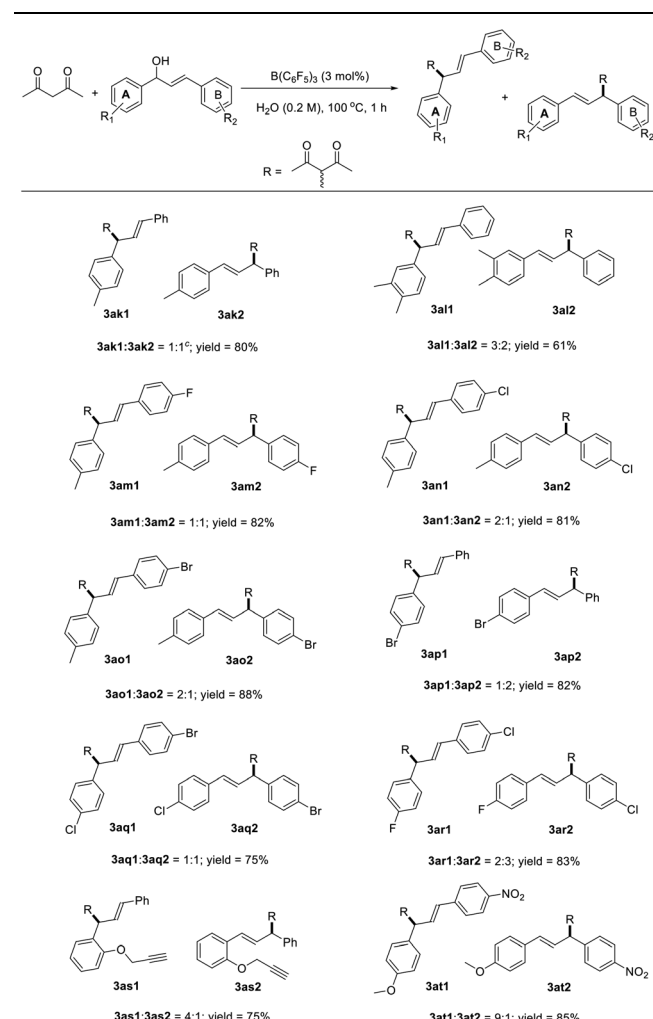


Table 3 Substrate scope for allylic alcohols^{a,b}

^a Reaction conditions: **1a** (0.3 mmol), **2a** (0.3 mmol), $\text{B}(\text{C}_6\text{F}_5)_3$ (3 mol%), H_2O (1.5 mL) at 100°C for 1 h. ^b Yield of the isolated product.

group, the ratio of the double bond migration product to the non-migrated product was about 1 : 1 by NMR (**3ak**); when a methyl group was added at the *meta* position of the allyl alcohol A ring, the yield of the product was slightly lower and the ratio increased to 3 : 2 (**3al**); and when numerous halogen groups were introduced at the *para* position of the allyl alcohol B ring, the ratio changed (**3am**–**3ao**). In the allyl carbanions formed during the reaction, due to the electron-donating effect of the methyl group on the A ring, the proportion of products near to the A ring in the mixtures might be larger. The *para* position of the allyl alcohol A ring was replaced by bromine, and the ratio of the double bond migration product to the non-migrated product was about 1 : 2 by NMR (**3ap**) due to the electronegativity of bromine, which is stronger than that of hydrogen, and the changes of the proportion about **3aq** and **3ar** also conformed to the law of the electronegative size of the atom, and the reaction is also suitable for the reduction of terminal alkenes (**3as**). When A ring was attached to a strong electron donating substituent and the B ring was connected to the strong electron withdrawing group, the uneven distribution of the allyl cation charge occurred, resulting in a product ratio of 9 : 1 (**3at**). Unfortunately, the huge limitation was shown in this reaction concerning the allylic alcohol substrates (Fig. 1).

To illustrate the synthetic utility of the developed process, an amplification scale reaction was implemented. A 1 g scale reaction was carried out for acetylacetone **1a** and (*E*)-1,3-diphenylallyl alcohol **2a**, and the product **3aa** was isolated in

Table 4 Substrate scope for allylic alcohols^{a,b}

^a Reaction conditions: **1a** (0.3 mmol), **2a** (0.3 mmol), $\text{B}(\text{C}_6\text{F}_5)_3$ (3 mol%), H_2O (1.5 mL) at 100°C for 1 h. ^b Yield of isolated product. ^c The ratio of 3 was determined by ^1H NMR.

85% yield. The result showed that the product yield was not affected after the reaction was scaled up, further indicating that the reaction has great potential for application. We derived the product of the allylic reaction obtained, and the reaction of **3ca** and phenylhydrazine in acetic acid at 100°C for one hour could produce tetrasubstituted pyrazole compound **4** in 69% yield (Fig. 2A). The compound **3ja** could be converted to a methyl (*E*)-2-(1-hydroxyethyl)-3,5-diphenylpent-4-enoate with NaBH_4 (Fig. 2B). In addition, the ester group could be removed with NaOH to form α,δ -unsaturated ketone (Fig. 2C). The compound

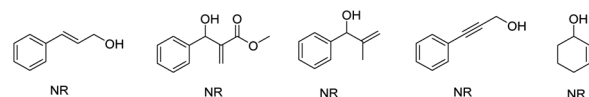


Fig. 1 The allyl alcohols of the corresponding allylated products cannot be obtained in this reaction.



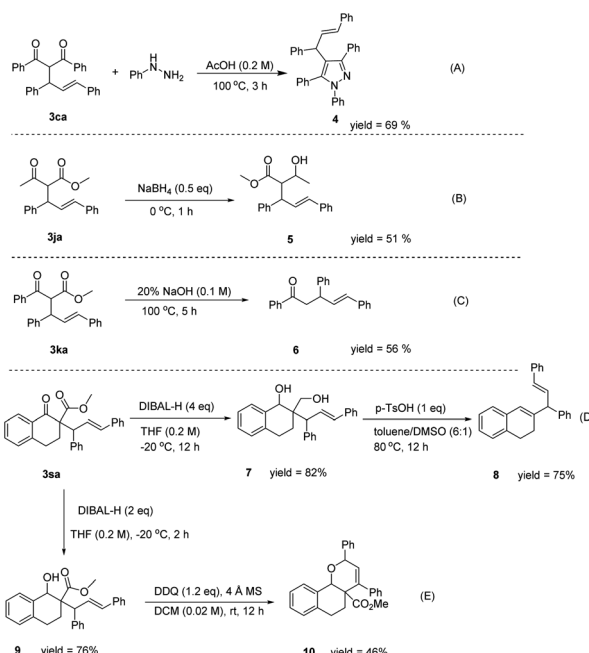
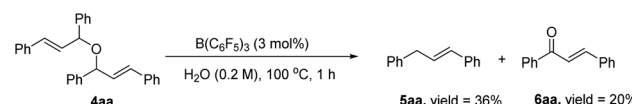


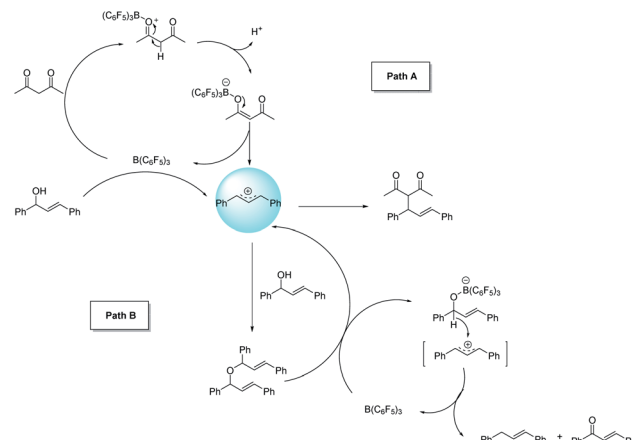
Fig. 2 Preliminary protocol applications.

3sa could be converted to compound 7 or compound 9 by DIBAL-H under different reaction conditions. Also, compound 7 reacted with PTSA in toluene and DMSO at 80 °C for 12 h could be generated from compound 8 (Fig. 2D). Compound 7 might have formed from a four-membered cyclic ether³¹ by *p*-toluenesulfonic acid catalyzed in the reaction, and then the ring was cleaved³² to produce compound 8. The skipped dienes are common structural motifs in numerous biologically active molecules due to the stereochemical identity of the diene unit.³³ However, the direct dehydrogenative cycloetherification of compound 9 promoted by DDQ afforded compound 10 in moderate yield (Fig. 2E). The functionalized dihydropyrans are also an important class of heterocyclic target molecules, serving as the common core of the structure in natural products or precursors to diverse C-glycosides.³⁴

In the next study, we found that the etherification product (4aa) reacted with acetylacetone could give 3aa in good yield under the standard conditions. Also, the yield of the product could be comparable to the template reaction (Scheme 2). Simultaneously, the by-product 4aa would be decomposed into 5aa and 6aa under the standard conditions (Scheme 3). Based on the reported literature,³⁵ we proposed the mechanism of this reaction (Scheme 4). The reaction is a nucleophilic attack about the enol form of 1,3-dicarbonyl compounds and allyl carbocation. First, the carbonyl group of 1,3-diketone can be extremely



Scheme 3 The decomposition of 4aa.



Scheme 4 Plausible reaction mechanism.

activated by $B(C_6F_5)_3$. Second, there are two possible ways in the formation of allyl carbocation: one is that due to the high temperature and strong acidity for the protonated product produced by the catalyst, and the other is that the allyl carbocation in the reaction attacked by the hydroxyl oxygen of the allylic alcohol produced the by-product bis(1,3-diphenylallyl) ether (4aa), and the allyl carbocation can also be formed when the etherified product is cleaved by a protonated catalyst.

Conclusions

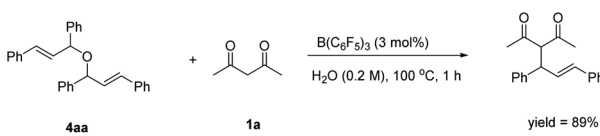
In summary, a metal-free catalyzed allylation of 1,3-diketones or β -ketone esters with allyl alcohols in water has been reported. The allyl alcohols with 1,3-diketones and β -ketone esters are dehydrated in water, and a quaternary carbon center can be well constructed for active methine of 1,3-dicarbonyl compounds in this method. The products can be further converted into other compounds such as tetrasubstituted pyrazole or 1,4-dienes and functionalized dihydropyrans. This reaction has a very good atom economy and water is only the by-product and a green solvent simultaneously. The protocol would offer new insight into the reaction catalyzed by $B(C_6F_5)_3$.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Scheme 2 The allylization of 4aa.



Notes and references

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