Highly efficient conjugate reduction of α , β -unsaturated nitriles catalyzed by copper/xanthene-type bisphosphine complexes[†]

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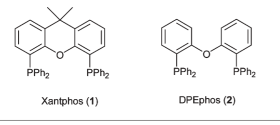
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 α , β -Unsaturated nitriles are chemoselectively reduced to the corresponding saturated nitriles in high yields using a copper-DPEphos or Xantphos complex as catalyst in the presence of polymethylhydrosiloxane (PMHS) as the stoichiometric reducing agent and *t*-butanol as additive.

 α ,β-Unsaturated nitriles are versatile synthetic intermediates in organic synthesis¹ and the selective reduction of conjugated nitriles to their saturated counterparts has long been a synthetic interest. While various reduction methods of α ,β-unsaturated nitriles have been developed by using a stoichiometric amount of metal² and metal hydrides,³ only a few metal-catalyzed conjugate reductions were reported with a relatively nonreactive source of hydride entities such as silanes.⁴ These reducing systems, however, have limitations, as the reduction is very slow, low-yielding, and often β,β-disubstituted- α ,β-unsaturated nitriles cannot be reduced at all.

Despite recent advances of Cu–H catalyzed hydrosilylation methodologies,⁵ an efficient conjugate reduction method of α , β -unsaturated nitriles catalyzed by Cu–H has not been reported. It was observed that unsaturated nitriles reacted more slowly than other related unsaturated functionalities in various metal-catalyzed reductions^{4,6} and the linearity of the CN group prevented the formation of a stable enolate analogue.^{4b} The CN functionality was also reported to inhibit the catalytic activity of Cu–H in other copper-catalyzed reductions.^{5c,7}

Recently, we have shown that Cu–H for the asymmetric reduction of ketones could be generated from the combination of catalytic amounts of copper(II) acetate or copper(II) acetate monohydrate and (*S*)-BINAP in the presence of Ph₂SiH₂ or PhSiH₃.⁸ We, therefore, decided to investigate the conjugate reduction of α , β -unsaturated nitriles with our catalytic system and report here that a complex of copper(II) acetate and Xantphos (1) or DPEphos (2) is quite effective for catalyzing the conjugate reduction of α , β -unsaturated nitriles in the presence of PMHS at ambient temperature.



† Electronic supplementary information (ESI) available: experimental procedures and spectroscopic data of all products (4a-4j). See http:// www.rsc.org/suppdata/cc/b4/b418586b/ *jaesook@ajou.ac.kr

In our initial investigations on the reduction of (E)-cinnamonitrile (3a) with PMHS, we employed the Cu(OAc)₂·H₂O/BINAP catalyst system, which was quite effective for the reduction of aromatic ketones. However, the reaction showed no appreciable conversion even after long reaction times. We screened a series of commercially available bisphosphine ligands and organosilanes with no success. We then turned our attention to the use of additives to promote reaction rates since many hydrosilylation reactions were reported to be accelerated by alcohol additives.9 Addition of 4 equiv of t-BuOH to the reaction mixture led to an increased rate of reduction¹⁰ and **3a** was guite efficiently reduced to its saturated analogue in high yield (Table 1, entry 5). Other achiral bisphosphine ligands were investigated and the results are summarized in Table 1. While bisphosphine ligands that have flexible backbones such as dppb, dppp are either inactive or moderately active (entries 3 and 4), bisphosphines based on rigid aromatic backbones proved quite efficient (entries 5-7). In particular, Xantphos $(1)^{11}$ was quite effective for the reduction of 3a yielding the saturated counterpart 4a in high yield (entries 7 and 8). DPEphos¹¹ showed comparable reactivity, but the isolated yield of 4a was lower for this substrate.

As shown in Table 2, a range of α , β -unsaturated nitriles prepared from the corresponding aldehydes or ketones were chemoselectively reduced smoothly at room temperature in high yields by a Cu(OAc)₂/Xantphos catalyst or Cu(OAc)₂/DPEphos catalyst, in the presence of PMHS and *t*-BuOH.¹² A wide range of aromatic and aliphatic substrates were reduced to the saturated nitrile products in good yields. Functional groups such as MeO, Cl, F, CN were tolerated (entries 2–4 and 9). A heterocyclic substrate, 3-(2-furanyl)-2-propenenitrile (**3e**), was also effectively

Table 1 Conjugate reduction of (E)-cinnamonitrile under variousconditions

\sim	CN 3 mo	l% Cu(OAc)₂∙H₂C		
3	4 eq	uiv PMHS, toluer	4a	
Entry	Ligand	Additive	Time/h	Yield/% a
1 ^{<i>b</i>}	(S)-BINAP		no rxn	
2	Xantphos		no rxn	_
3 ^c	dppb	t-BuOH	no rxn	_
4^d	dppp	t-BuOH	21	70
5	(S)-BINAP	t-BuOH	1.5	90
6	DPEphos	t-BuOH	<1	83
7	Xantphos	t-BuOH	<1	92
8 ^e	Xantphos	t-BuOH	<1	96

^{*a*} Isolated yields. ^{*b*} No reaction occurred with Ph₂SiH₂ and PhSiH₃ either. ^{*c*} dppb = 1,4-Bis(diphenylphosphino)butane. ^{*d*} dppp = 1,3-Bis(diphenylphosphino)propane. ^{*e*} Cu(OAc)₂ was used.

R ₁	CN	1. 3 mol% Cu(OAc) 4 equiv PMHS/t-E	₂ ·H ₂ O/ BuOH,	toluene, RT		
^{N1} 3		2. 2.5 M-NaOH wor	k up			4
Entry	Substrate	3		Ligand	Time/h	Yield/% a
1 ^{<i>b</i>}		CN	3a	1	<1	92
2	MeO	CN CN	3b	1	1	89
3 ^b	\bigcap	CN	3c	1	1	90
4	F	CN	3d	1 2	1 1	92 87
5		Lu CN	3e	1	1	85
6	H ₃ C	CN	3f	1	7	80
7	CN		3g	1	3.5	87
8 ^c		CN CH3	3h	2	10	87
9 ^c	NC	CH ₃	3i	2	9	84
10 ^{b,c,d}	NC		3j	2	12	94

1. 3 mol% Cu(OAc)₂·H₂O/Ligand (1 or 2)

 R_2

^{*a*} Isolated yield. ^{*b*} Pure (*E*)-isomer was used. ^{*c*} Cu(OAc)₂ was used instead of Cu(OAc)₂·H₂O. ^{*d*} Pure (*Z*)-isomer was used.

reduced in 85% isolated yield (entry 5). Conjugation of the double bond with an aromatic ring is not a requirement for reduction. Simple aliphatic substrates, **3f** derived from 2-octanone and **3g** from cyclohexanone afforded products in hours (entries 6 and 7).¹³ Reduction of the sterically more hindered substrates (**3h–3j**) prepared from the corresponding aromatic ketones were conducted more efficiently by the Cu/DPEphos system than by the Cu/Xantphos system, leading reaction to completion in reasonable reaction times.

Copper complexes of Xantphos or DPEphos are thermally more stable and more efficient for the hydrosilylation of nitriles than a Cu/BINAP complex. While the Cu/Xantphos-type ligands system reduces β , β -disubstituted- α , β -unsaturated nitriles effectively, the Cu/BINAP system results in slow conversion of the substrate and the reaction mixture eventually turns to a black color in hours (*ca.* 12 h), which is presumably resulted from decomposition of Cu–H at room temperature.¹⁴

It seems that in our reducing system employing a Cu(II) salt as the catalyst precursor, the active catalyst is Cu(I)–H, which is the same species normally generated from copper(I) chloride, sodium *t*-butoxide, and a reducing agent.¹⁵ We postulate that 1,4-addition of the Cu–H to α , β -unsaturated nitriles takes place and the resulting organocopper species¹⁶ reacts with *t*-BuOH to yield the protonated product rapidly and a copper alkoxide. The latter then regenerates the active catalyst Cu–H with PMHS.

In conclusion, we have developed an efficient method for the conjugate reduction of α , β -unsaturated nitriles based on Cu(I)–H generated in the presence of the xanthene-based ligands **1** and **2**. The active Cu(I)–H was generated from copper(II) acetate with an organosilane, and was thermally stable enough to carry out the reduction at ambient temperature. The use of *t*-BuOH as additive was a key to success and inexpensive polymeric hydrosilane PMHS could be used as the stoichiometric reducing agent. An asymmetric version of this reaction is actively under investigation in our group with chiral bisphosphine ligands based on a xanthene or ferrocenyl framework.

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aid of Et₂O (10 mL), and then NaOH (2.5 M, 1.2 mL) was added. The biphasic mixture was stirred vigorously for 0.5 h. The layers were separated and the aqueous layer was extracted with Et₂O (3×20 mL). The combined organic layers were washed with brine, dried over MgSO₄, and concentrated. The product was purified by Kugelrohr distillation or silica gel chromatography.

- 13 Acrylonitrile was also a good substrate for the reduction and its conversion could be followed by GC analysis (1.5 h, 100% conversion with 2 equiv. *t*-BuOH). However, the nitrile substrate derived from *t*-butyl methyl ketone (pinacolone) was not reduced under the reaction conditions, probably due to its steric effect.
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