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Development of an imidazole-based N,N-bidentate ligand for the manganese catalyzed direct coupling of nitriles with alcohols†

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3d-Metal catalyzed borrowing hydrogen (BH) reactions represent powerful and environmentally friendly approaches for the direct coupling of alcohols with nitriles to assemble various important branched nitriles. The development of simple and efficient ligands is a crucial issue in this field. In this study, we designed a series of readily available *N*,*N*-bidentate ligands that demonstrated good efficiency in the Mncatalyzed BH reaction of alcohols and nitrile derivatives, yielding the targeted nitriles in moderate to good yields. Remarkably, the mildness and practicality of this protocol were further demonstrated by the successful synthesis of anipamil *via* a two-cascade borrowing hydrogen procedure.

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Introduction

α-Alkylated nitriles are not only widely present in numerous biologically active molecules, pharmaceuticals, and lightemitting diodes but also serve as versatile building blocks for the synthesis of diols, lactones, lactams, amino alcohols, and cyclic amine derivatives. 1-4 Considering the significant importance of these scaffolds, tremendous efforts have been dedicated to developing efficient methodologies for α-alkylated nitriles. Traditionally, a general approach for accessing these compounds involved the condensation of nitriles with ketones, aldehydes, or alkyl halides under strong base-promoted conditions.5,6 However, this traditional strategy faced a significant challenge in overcoming multiple side reactions, such as selfcondensation of the nitriles, the aldol reaction, or the Cannizzaro reaction. It also needed to expand the functional group tolerance towards stoichiometric strong bases while avoiding the generation of copious wastes. Alternatively, the borrowing hydrogen (BH) strategy, using alcohol as an efficient alkylating agent, offers an atom-economical and environmentally benign tool for constructing new C-C bonds with water as the sole byproduct.7 A breakthrough in this field was made by Grigg and co-workers in 1981, who reported a ruthenium-catalyzed alkylation of nitriles with alcohols to deliver α-alkylated nitriles.8

Manganese, the third most abundant transition metal in the earth's crust, has emerged as an efficient catalyst in borrowing hydrogen (BH) reactions, replacing precious metals (Scheme 1). In 2018, Maji and co-workers introduced a phosphine-free bidentate ligand derived from the condensation of thiophen-2-yl ketone and pyridine-containing hydrazine. 16a They applied this ligand in the first Mn-catalyzed α-alkylation of aryl nitriles with alcohols. Soon after, the groups of Reuping and El-Sepelgy demonstrated the utility of the Mn-PNP complex for the same reaction, synthesizing a diverse range of substituted alkylated nitriles.16b Notably, the PNP ligand's instability due to the easy oxidation of P(III) to P(v) was recognized. Recently, Mukherjee's group identified a simple Mn catalyst, generated in situ from Mn(CO)₅Br and 2,2'-bipyridine, for this reaction, albeit with high catalyst loading and a narrow substrate scope.17 Despite the significant advances in Mn-based catalytic systems for αalkylation of aryl nitriles, the discovery of a cheap and efficient Mn catalyst remains a great challenge.

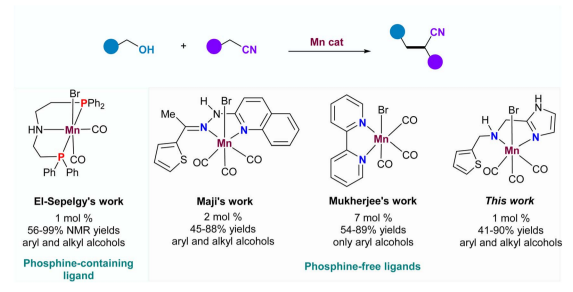
In our previous work, we developed several powerful imidazole-based tridentate ligands for Mn-catalyzed asymmetrical hydrogenation of various ketones, leading to highly enantioenriched alcohols. Recognizing the crucial role of the imidazole moiety in these ligands, we envisioned that combining the imidazole group with another nitrogen component could yield a readily available and efficient ligand for Mn-

Since then, several other noble metals such as Ir, ⁹ Rh, ¹⁰ Os¹¹ and Ru¹² have been successfully employed as efficient catalysts in the direct coupling of nitriles and alcohols. However, considering the toxicity, cost, and limited availability of noble metals, the development of abundant first-row-transition metal catalysts (Fe, ¹³ Co, ¹⁴ Ni, ¹⁵ Mn¹⁶) has become a highly attractive strategy to reduce the use of noble metals in homogeneous catalysis.

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Scheme 1 Manganese catalyzed α -alkylation of nitriles with alcohols.

catalyzed α-alkylation of aryl nitriles with alcohols. Herein, in our continuous interest in this field,19 we report a simple imidazole-based N,N-bidentate ligand-assisted Mn-catalyzed efficient synthesis of α-alkylated nitriles through a BH process. This protocol demonstrates a broad substrate scope with good functional tolerance at low catalyst loading (26 examples, 41-90% yields) and is applicable in the total synthesis of anipamil.

Results and discussion

The currently developed imidazole-based N,N-bidentate ligands L1-L4 can be easily obtained in yields ranging from 47% to 65% by treating imidazole aldehydes or ketones with amines in

Scheme 2 Synthetic route for the novel Mn catalyst.

MeOH under reductive conditions (Scheme 2). Subsequently, the Mn catalysts Mn-1-Mn-4 are prepared by coordinating Mn(CO)₅Br with L1-L4 in toluene at reflux. With these catalysts in hand, our studies commenced by screening them for the α alkylation of benzonitrile with benzyl alcohol (Table 1). Initially, the screening of Mn catalysts revealed that NH unprotected imidazole ligands provided better yields compared to those

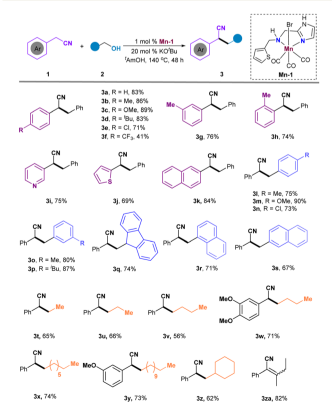
Table 1 Optimization of reaction conditions

Entry	Cat.	Base	Solvent	$Yield^{b}$ (%)
1	Mn-1	KO^t Bu	^t AmOH	75
2	Mn-2	KO^tBu	^t AmOH	35
3	Mn-3	KO^tBu	^t AmOH	70
4	Mn-4	KO^tBu	^t AmOH	41
5	Mn-1	KO^tBu	Toluene	62
6	Mn-1	KO^tBu	Xylenes	64
7	Mn-1	KO^tBu	^t BuOH	63
8	Mn-1	KO^t Bu	THF	21
9	Mn-1	KO^tBu	1,4-Dioxane	43
10	Mn-1	^t BuOLi	^t AmOH	15
11	Mn-1	^t BuONa	^t AmOH	46
12	Mn-1	AcONa	^t AmOH	30
13	Mn-1	KOH	^t AmOH	59
14^c	Mn-1	KO^t Bu	^t AmOH	72
$15^{c,d}$	Mn-1	KO^tBu	^t AmOH	83
16	_	KO^tBu	^t AmOH	Trace
$17^{c,d,e}$	Mn-1	KO ^t Bu	^t AmOH	78

^a General conditions: 1a (1.0 mmol), 2a (0.5 mmol), Mn cat. (2 mol%), ^tBuOK (20 mol%) were dissolved in t-AmOH (1.5 mL) and heated to 140 °C for 36 h. ^b Isolated yield. ^c 1 mol% of **Mn-1** was used. ^d Stirred for 48 h. e 10 mmol scale.

protected by a methyl group, and **Mn-1** emerged as the optimal catalyst for this reaction (Table 1, entries 1–4). A subsequent examination of solvents showed that 'AmOH yielded better results than others (Table 1, entries 5–9). Moreover, bases containing potassium resulted in higher yields than those with lithium and sodium, with KO'Bu proving to be the best base, offering 3a in a 75% yield (Table 1, entries 10–13). Additionally, a slight decrease in yield was observed when reducing the catalyst loading from 2 mol% to 1 mol% (Table 1, entry 14). Fortunately, extending the reaction time to 48 h with 1 mol% of **Mn-1** showed improved conversion, yielding the desired 3a in 83% (Table 1, entry 15). As expected, the reaction failed to yield α -alkylated nitrile without the presence of **Mn-1** (Table 1, entry 16). Notably, the reaction could be successfully conducted on a 10 mmol scale, yielding 3a in 78% (Table 1, entry 17).

With the optimized reaction conditions established, our focus shifted to exploring the substrate scope for accessing α-alkylated nitriles (Scheme 3). Initial examination of aryl rings adjacent to the nitrile group revealed that electron-donating substituents on benzene rings (Me-, CH₃O-, and ^tBu-) provided higher yields (3b, 86%; 3c, 89%; 3d, 83%) compared to the electron-withdrawing group (3f, 41%). Importantly, halide groups like chlorine were retained under the reaction conditions, yielding product 3e in an acceptable yield. The substituent position had minimal impact on the reaction outcomes; *ortho*-substituted aryl nitrile showed lower yield than those with *meta*- or *para*-substituents (3a, 83% νs. 3g, 76% νs. 3h, 74%). Additionally, heteroaryl-substituted nitriles were compatible, delivering products 3i, 3j, and 3k in



Scheme 3 Substrate scope.

Scheme 4 Total synthesis of anipamil.

69–84% yields. A variety of aromatic primary alcohols were screened, showing no significant electron and steric effects and producing the targeted products 31-3p in moderate to good yields. Fluorenyl and naphthyl groups were also tolerated, generating the desired 3q–3s in 67–74% yields. The reaction proceeded smoothly with primary alkyl alcohols, yielding 3t–3z in moderate to good yields. Unfortunately, when the primary alcohol is changed to a secondary alcohol, the hydrogenation product cannot be obtained, but the dehydrogenation product 3za can be obtained in 82% yield.

To showcase the scalability and utility of this method, we applied it to the synthesis of the calcium channel blocker anipamil (Scheme 4). Initially, a 5 mmol scale of 3-methoxyphenylacetonitrile 1y underwent a Mn-1-catalyzed BH reaction with 1-dodecanol to yield secondary nitrile 3y in 72% yield after 48 h. Subsequently, the second BH coupling reaction of 3y with propylene alcohol, enabled by a Ru-MACHO catalyst, proceeded smoothly to generate tertiary nitrile 4y in 95% yield. The bromination of nitrile 4y with PBr₃ led to the formation of 5y in 78% yield. Finally, the nucleophilic attack of 6 on 5y resulted in the generation of the calcium channel blocker anipamil in an 80% yield.

A series of control experiments were conducted to gain a preliminary understanding of the reaction mechanism, as depicted in Scheme 5. Firstly, the reaction of 1a with deuterated benzyl alcohol $2a-d_2$ resulted in the formation of deuterated

Scheme 5 Control experiments.

Plausible reaction mechanism

product 3a-d in 78% yield, with a 63% deuteration at the βposition of the nitrile group. This implies that the hydride originated from the alcohol $2a-d_2$ (Scheme 5(1)). Subsequently, the condensation of 1a with benzaldehyde led to the formation of unsaturated aryl nitrile 8a in 90% yield. This intermediate was successfully reduced to 3a-d in 82% yield, with 91% deuteration under standard conditions using deuterated benzyl alcohol 2a-d2 as a hydrogen transfer reagent (Scheme 5(2) and (3)). These experiments suggested that the aldehyde may be the key intermediate in this reaction, further confirming that the hydrogen source comes from the alcohol.

Based on these observations and previous literature, 20 we propose a plausible mechanism illustrated in Fig. 1. Initially, an active Mn catalyst I is formed with the assistance of a base, initiating the dehydrogenative process of alcohol 2a to produce aldehyde 7a, along with Mn-D species III through a sixmembered Mn complex II. Subsequently, aldehyde 7a and phenylacetonitrile 1a undergo a Knoevenagel-type condensation to yield α,β -substituted acrylonitriles 8a. Finally, the Mn-D species III facilitates the reduction of acrylonitriles 8a via a transient state IV, leading to the formation of nitrile 3a and the regeneration of active Mn catalyst I.

Conclusions

In conclusion, we have developed a series of novel N,Nbidentate ligands for the Mn catalyzed borrowing hydrogenation of alcohols with nitriles. This procedure featured a broad

substrate scope with a good functional tolerance under simple conditions to provide the secondary nitriles in moderate to good yields (26 examples, 41-90% yields). Remarkably, the mildness and practicality of this protocol was further demonstrated by the synthesises of anipamil via two-cascade borrowing hydrogen procedure. Mechanistic studies were also conducted, which confirmed that the hydride came from the alcohol and aldehyde was the key intermediate. The present cheap metal-catalyzed mild, general strategies are expected to be of high interest to scientists in academia and industry.

Experimental

General procedure for the synthesis of α -alkylated nitrile

To a mixture of **Mn-1** catalyst (1 mol%), KO^tBu (0.2 eq.), nitrile (1.0 mmol) and primary alcohol (0.5 mmol), 1.5 mL of ^tAmOH was added. Then, the reaction was stirred under Ar in a pressure tube (ACE pressure tube, 15 mL). The reaction was stirred at 140 °C for 48 hours. After cooling to room temperature, the reaction was diluted with ethyl acetate (10 mL) and water (10 mL). The organic layer was separated, and the aqueous layer was extracted with ethyl acetate (10 mL) for three times. The combined organic layers were washed by brine and dried over magnesium sulfate and the volatiles were removed under reduced pressure. The residue was purified by column chromatography on silica gel (petroleum ether/ethyl acetate = 20:1-10:1) to give the desired product.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 F. F. Fleming, L. Yao, P. C. Ravikumar, L. Funk and B. C Shook, *J. Med. Chem.*, 2010, 53, 7902–7917.
- 2 F. F. Fleming, Nat. Prod. Rep., 1999, 16, 597-606.
- 3 X. Bao, Q. Wang and J. Zhu, *Angew. Chem., Int. Ed.*, 2019, **58**, 2139–2143.
- 4 A. D. Dorsey, J. E. Barbarow and D. Trauner, *Org. Lett.*, 2003, 5, 3237–3239.
- (a) D. F. Taber and S. J. Kong, J. Org. Chem., 1997, 62, 8575–8576; (b) D. Savoia, E. Tagliavini, C. Trombini and A. Umani-Ronchi, J. Org. Chem., 1980, 45, 3227–3229; (c) F. L. Cook, C. W. Bowers and C. L. Liotta, J. Org. Chem., 1974, 39, 3416–3418; (d) S. Caron, E. Vazquez and J. M. Wojcik, J. Am. Chem. Soc., 2000, 122, 712–713; (e) B. A. D'Sa, P. Kisanga and J. G. Verkade, J. Org. Chem., 1998, 63, 3961–3967; (f) R. Guillot, A. Loupy, A. Meddour, M. Pellet and A. Petit, Tetrahedron, 2005, 61, 10129–10137.
- 6 (a) B. A. D'Sa, P. Kisanga and J. G. Verkade, J. Org. Chem.,
 1998, 63, 3961–3967; (b) R. Guillot, A. Loupy, A. Meddour,
 M. Pellet and A. Petit, Tetrahedron, 2005, 61, 10129–10137.
- 7 (a) B. G. Reed-Berendt, D. E. Latham, M. B. Dambatta and L. C. Morrill, ACS Cent. Sci., 2021, 7, 570-585; (b) T. Irrgang and R. Kempe, Chem. Rev., 2019, 119, 2524-2549; (c) A. Corma, J. Navas and M. J. Sabater, Chem. Rev., 2018, 118, 1410-1459; (d) R. H. Crabtree, Chem. Rev., 2017, 117, 9228-9246; (e) Q. Yang, Q. Wang and Z. Yu, Chem. Soc. Rev., 2015, 44, 2305-2329; (f) G. Guillena, D. J. Ramón and M. Yus, Angew. Chem., Int. Ed., 2007, 46, 2358-2364; Angew. Chem., 2007, 119, 2410-2424.
- 8 R. Grigg, T. R. B. Mitchell, S. Sutthivaiyakit and N. Tongpenyai, *Tetrahedron Lett.*, 1981, 22, 4107–4110.
- 9 (a) C. Löfberg, R. Grigg, M. A. Whittaker, A. Keep and A. Derrick, J. Org. Chem., 2006, 71, 8023-8027; (b)
 B. Anxionnat, D. Gomez Pardo, G. Ricci and J. Cossy, Org. Lett., 2011, 13, 4084-4087; (c) T. Sawaguchi and Y. Obora, Chem. Lett., 2011, 40, 1055-1057; (d) M. Morita, Y. Obora and Y. Ishii, Chem. Commun., 2007, 2850-2852; (e)
 B. Anxionnat, D. Gomez Pardo, G. Ricci and J. Cossy, Eur. J. Org Chem., 2012, 4453-4456.

- 10 (a) F. Li, X. Zou and N. Wang, Adv. Synth. Catal., 2015, 357, 1405–1415; (b) J. Li, Y. Liu, W. Tang, D. Xue, C. Li, J. Xiao and C. Wang, Chem.–Eur. J., 2017, 23, 14445.
- 11 M. L. Buil, M. A. Esteruelas, J. Herrero, S. Izquierdo, I. M. Pastor and M. Yus, *ACS Catal.*, 2013, 3, 2072–2075.
- 12 (a) S. Huang, X. Hong, Y. Sun, H.-Z. Cui, Q. Zhou, Y.-J. Lin and X.-F. Hou, *Appl. Organomet. Chem.*, 2020, 34, e5451; (b)
 T. Kuwahara, T. Fukuyama and I. Ryu, *Chem. Lett.*, 2013, 42, 1163–1165; (c) S. Thiyagarajan and C. Gunanathan, *ACS Catal.*, 2017, 7, 5483–5490.
- 13 W. Ma, S. Cui, H. Sun, W. Tang, D. Xue, C. Li, J. Fan, J. Xiao and C. Wang, *Chem.-Eur. J.*, 2018, 24, 13118.
- 14 (a) P. Chakraborty, N. Garg, E. Manoury, R. Poli and B. Sundararaju, ACS Catal., 2020, 10, 8023–8031; (b) K. Paudel, S. Xu and K. Ding, J. Org. Chem., 2020, 85, 14980–14988; (c) A. Singh and M. Findlater, Organometallics, 2022, 41, 3145–3151.
- 15 (a) R. Saha, S. Panda, A. Nanda and B. Bagh, J. Org. Chem., 2023, DOI: 10.1021/acs.joc.3c00859; (b) S. Bera, A. Bera and D. Banerjee, Chem. Commun., 2020, 56, 6850–6853.
- 16 (a) A. Jana, C. B. Reddy and B. Maji, ACS Catal., 2018, 8, 9226–9231; (b) J. C. Borghs, M. A. Tran, J. Sklyaruk, M. Rueping and O. El-Sepelgy, J. Org. Chem., 2019, 84, 7927–7935.
- 17 B. Krishanu and A. Mukherjee, Chem.-Asian J., 2023, 18, e202300157.
- 18 (a) F. Ling, H. Hou, J. Chen, S. Nian, X. Yi, Z. Wang, D. Song and W. Zhong, Org. Lett., 2019, 21, 3937–3941; (b) F. Ling, J. Chen, S. Nian, H. Hou, X. Yi, F. Wu, M. Xu and W. Zhong, Synlett, 2020, 31, 285–289; (c) Z. Wang, X. Zhao, A. Huang, Z. Yang, Y. Cheng, J. Chen, F. Ling and W. Zhong, Tetrahedron Lett., 2021, 82, 153389; (d) J. He, W. Mao, J. Lin, Y. Wu, L. Chen, P. Yang, D. Song, P. Zhu, W. Zhong and F. Ling, Org. Chem. Front., 2023, 10, 3321–3327.
- 19 (a) D. Song, L. Chen, Y. Li, T. Liu, X. Yi, L. Liu, F. Ling and W. Zhong, Org. Chem. Front., 2021, 8, 120–126; (b) X. Yi, Y. Chen, A. Huang, D. Song, J. He, F. Ling and W. Zhong, Org. Chem. Front., 2021, 8, 6830–6836; (c) S. Wang, D. Song, F. Shen, R. Chen, Y. Cheng, C. Zhao, Q. Shen, S. Yin, F. Ling and W. Zhong, Green Chem., 2023, 25, 357–364; (d) T. Liu, W. Zhang, C. Xu, Z. Xu, D. Song, W. Qian, G. Lu, C.-J. Zhang, W. Zhong and F. Ling, Green Chem., 2023, 25, 3606–3614; (e) D. Song, S. Wang, W. Huang, R. Chen, F. Hu, L. Chen, X. Zhao, F. Ling and W. Zhong, Org. Chem. Front., 2023, 10, 5908–5915.
- 20 (a) R. Ferrari, R. Raddino, C. Ceconi, S. Curello, S. Ghielmi and O. Visioli, *Cardiovasc. Drugs Ther.*, 1989, 3, 403–412; (b) D. Lenke and C. D. Mueller, Use of Anipamil, *US Pat.*, US4777183A, 1988.