Approaching sub-ppm-level asymmetric organocatalysis of a highly challenging and scalable carbon-carbon bond forming reaction

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The chemical synthesis of organic molecules involves, at its very essence, the creation of carbon-carbon bonds. In this context, the aldol reaction is among the most important synthetic methods, and a wide variety of catalytic and stereoselective versions have been reported. However, aldolizations yielding tertiary aldols, which result from the reaction of an enolate with a ketone, are challenging and only a few catalytic asymmetric Mukaiyama aldol reactions with ketones as electrophiles have been described. These methods typically require relatively high catalyst loadings, deliver substandard enantioselectivity or need special reagents or additives. We now report extremely potent catalysts that readily enable the reaction of silyl ketene acetals with a diverse set of ketones to furnish the corresponding tertiary aldol products in excellent yields and enantioselectivities. Parts per million (ppm) levels of catalyst loadings can be routinely used and provide fast and quantitative product formation in high enantiopurity. In situ spectroscopic studies and acidity measurements suggest a silylium ion based, asymmetric counteranion-directed Lewis acid catalysis mechanism.

hiral tertiary aldols are encountered in a variety of natural products and pharmaceuticals (Fig. 1a)¹. Despite the considerable advances in catalytic asymmetric aldol reactions of aldehydes²⁻⁶, only a few examples using ketones as electrophiles are known so far7-9. Thermodynamics typically favour the corresponding retro-aldol reaction¹⁰⁻¹³, while the lower reactivity of ketones compared to aldehydes further challenges the reaction kinetically. Importantly, however, the energy that is released in the tautomerization of the initial enol product of the reverse aldol reaction to the ketone is absent in the corresponding Mukaiyama variant. Therefore, the forward Mukaiyama aldol reaction¹⁴⁻¹⁶ is favoured, even with ketones. However, the enantiofacial differentiation of ketones is still much more challenging than that of aldehydes¹⁷. The Denmark group disclosed the first Mukaiyama aldol reaction of ketones catalysed by chiral bis-N-oxides acting as a Lewis base^{17,18}. This method requires the use of highly reactive trichlorosilyl enolates of methyl acetate as the nucleophile (Fig. 1b, top). Shibasaki and Kanai developed an enantioselective Mukaiyama aldol reaction of trimethylsilyl enolates to ketones in the presence of a chiral copper(I) fluoride-phosphine complex as the catalyst (Fig. 1b, bottom)¹⁹. Furthermore, the group of Campagne reported that chiral 2,2'-bis(di-p-tolylphosphino)-1,1'-binaphthyl (Tol-BINAP)copper(II) trifluoromethanesulfonate catalyses the vinylogous Mukaiyama aldol reaction of ketones²⁰.

Despite these advances, a more practical and general protocol for the Mukaiyama aldol reaction of ketones was envisioned exploiting the extraordinary reactivity of our recently developed imidodiphosphorimidate (IDPi) catalysts²¹⁻²⁵. In recent years, our group has made progress in the field of silylium-based, asymmetric counteranion-directed Lewis acid catalysis (silylium ACDC)²⁶ of Mukaiyama-type²⁷⁻²⁹ reactions. These transformations invariably require the use of aldehydes or imines as electrophiles, while ketones have remained elusive.

Results and discussion

Optimization of catalyst and silyl ketene acetal. For our initial studies, the addition of silyl ketene acetal **2a** to 2-acetonaphthone (**1a**) was chosen as a model reaction (Fig. 2). Indeed, with 0.1 mol% of chiral disulfonimide (DSI) catalyst $C-1^{27,30,31}$, no desired product was detected even after prolonged reaction times (24h). Full consumption of the ketone was observed, but instead of the desired aldol product, only silyl enol ether **4** was obtained. In sharp contrast, with 0.1 mol% of IDPi catalyst **C-2**, tertiary aldol product **3aa** was formed within 3 min and with an enantiomeric ratio (e.r.) of 69.5:30.5.

With these results in hand, an increase in enantioselectivity through further synthetic manipulations of the four aryl substitutents at the 3,3'-positions of the chiral 1,1'-bi-2-naphthol (BINOL) scaffold of the catalyst was pursued (Table 1, for detailed results on the optimization of solvents and silyl ketene acetals, see Supplementary Table 1). Besides the reported catalysts C-2 and C-3²¹, several new IDPi catalysts were synthesized (C-4, C-5, C-6, C-7 and C-8) and applied to the model reaction. Gratifyingly, elongating the alkyl chains at the 3,5-positions of the four aryl substituents increased the enantioselectivity, ultimately leading to an e.r. of 97:3 with catalyst C-8 (entries 4 to 7).

In all cases, the catalytic activities were extremely high and the reaction was completed within 30 min with 1.0 mol% of catalyst. Moreover, the catalyst loading can be reduced to 0.05 mol% (500 ppm) without deterioration of yield and enantioselectivity (97:3 e.r., entry 8). After screening of several different silyl ketene acetals (**2a–2d**), *tert*-butyldimethylsilyl (TBS) or triethylsilyl (TES) proved to be the most suitable silyl groups in terms of reactivity and enantioselectivity (entries 8 to 11).

Ketone scope and synthetic utility. Under the optimized reaction conditions, a variety of ketones (1b-1v) were subjected

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Fig. 1 [Representative examples of tertiary aldol-containing natural products and catalytic asymmetric routes to these moieties. a, Natural products containing tertiary aldols. **b**, Previously reported asymmetric Mukaiyama aldol reactions of ketones as electrophiles, using trichlorosilyl enolates of methyl acetate catalysed by a chiral Lewis base (top) and trimethylsilyl enolates catalysed by a chiral copper(I) fluoride-phosphine complex (bottom). Cy, cyclohexyl, L*, chiral ligand.



Fig. 2 | Reactivity comparison of DSI C-1 and IDPi C-2 in the Mukaiyama aldol reaction of ketone as electrophile under identical reaction conditions. Reactions were performed with ketone **1a** (0.5 mmol), silyl ketene acetal **2a** (2.0 equiv.) and catalyst (**C-1** or **C-2**, 0.1 mol%) in Et₂O (2.5 ml, 0.2 M) at 20 °C. The yield was determined after chromatographic purification and e.r. was determined by HPLC analysis. Tf, SO₂CF₃, TBS, SiMe₂t-Bu.

to this protocol (Table 2). Remarkably, a loading of 0.005–0.05 mol% (50–500 ppm) with catalysts C-4, C-7 or C-8 in Et₂O at –20 °C proved to be sufficient to provide the desired products (**3b**–**3v**) in quantitative yields (up to >99%) and with excellent e.r. (up to >99:1). Besides (hetero)aryl-alkyl ketones (**1b**–**1k**), alkyl-alkyl ketones (**1m**–**1o**) were also converted with good to excellent e.r. (88:12 to 95:5). Alkynyl ketone **1p** gave an excellent e.r. of 95:5. Interestingly, benzylideneacetone-type α , β -unsaturated ketones **1q**–**1v** were highly reactive and generally gave good 1,2- versus 1,4-regioselectivity (5:1 to 10:1) and excellent enantioselectivity (e.r.=97.5:2.5 to >99:1). Ketones other than methyl ketones, ethyl ketone **1c** and cyclic ketone **1l** were also

successfully transformed into desired products **3c** and **3l** (e.r. = 94:6 and 86:14), respectively.

To illustrate the synthetic utility of our method, derivatizations of the chiral tertiary aldols **3** were performed. Due to the high volatility of silyl ketene acetal **2** and the extremely low catalyst loading, analytically pure product **3** could be easily obtained after removing the volatiles under reduced pressure. The clean reaction profile of products allowed direct conversion of the material without the need for further chromatographic purification. Several functionalized chiral tertiary alcohols, such as β -silyloxy aldehyde **5** (89%, 95:5 e.r.), diols **6** (two steps, 92%, 95:5 e.r.) and **7** (88%, 95:5 e.r.), carboxylic acid **8** (91%, 95:5 e.r.), a product of transesterification

Table 1 | Optimization of the IDPi catalysts and the silyl ketene acetals 2



Reactions were performed with ketone **1a** (0.5 mmol), silyl ketene acetal **2** (2.0 equiv.) and catalyst (0.05-1.0 mol%) in Et₂O (2.5 ml, 0.20 M) at -20 °C. All yields are those of isolated products and e.r. was determined by HPLC analysis. TBS, *tert*-butyldimethylsilyl; TMS, trimethylsilyl; TES, triethylsilyl; TIPS, triisopropylsilyl. *Reaction was performed at 24 °C.

(9, 84%, 95:5 e.r.), and amide 10 (two steps, 83%, 95:5 e.r.) were successfully obtained. In addition, chiral tertiary aldols **3p** and **3u** were quantitatively transformed to product **3n**, without erosion of e.r. (see Supplementary Sections 7 and 8 for details).

Towards extremely low catalyst loadings. To further investigate the performance of IDPi catalyst C-8, very low catalyst loading³² experiments were implemented on a preparative scale. Deca-grams of products 3i (>99%, 20.5g) and 3v (92%, 20.1g) were quantitatively obtained in very high enantioselectivities (95:5 e.r. and 96:4 e.r.) using 25 ppm or 2.8 ppm of the catalysts, respectively (Fig. 3a; see Supplementary Section 12 for analytical results). Substrate 1u was further explored¹⁷. Due to the existence of a background reaction (Supplementary Fig. 2), conditions suitable for obtaining excellent e.r. in these large-scale syntheses were first identified (at 5°C with 8.0 M; see Supplementary Table 3 for full data). Initially, we fixed the amount of ketone 1u at 1.00g and lowered the catalyst loadings from 8 ppm (0.10 mg) to 2.8 ppm (0.037 mg). In all cases, full conversion of 1u and quantitative yield of tertiary aldol product **3u** (~90% yield) with >95:5 e.r. was observed. Excitingly, with 30.0 g of substrate, only 1.1 mg of IDPi C-8 successfully produced 59.2g of the enantioenriched product (86% corrected yield, turnover number (TON) ~307,000, Fig. 3a and Supplementary Table 4). Ketone 1u was partially soluble at the early stage of the reaction, and slowly dissolved during the course of the reaction, finally leading to a homogeneous solution (Fig. 3d).

We also investigated the repeated addition of starting materials after consumption of the initial reaction mixture with 2.8 ppm catalyst loading (batch **A**, Fig. 3b; see Supplementary Table 5). On completion of the reaction and removal of the volatiles, starting materials and solvent were recharged and subjected to the aldol reaction under the same conditions without the addition of more catalyst (batch **B**; see Supplementary Table 6). A third portion of substrate and reagent was added (batch **C**; see Supplementary Table 7), ultimately leading to a total conversion of 95.2% (82% yield: total TON ~911,000). It is noteworthy that ppm-level loadings in organocatalysis have previously been reported, for instance, in silylative kinetic resolutions of racemic alcohols³³ and in asymmetric transfer hydrogenations³⁴. To the best of our knowledge, however, approaching sub-ppm level loadings (<1 ppm, Fig. 3b,c) of an organocatalyst in a challenging carbon–carbon bond forming reaction with high enantioselectivity is unprecedented.

In contrast to the formidable catalytic activity of our IDPi catalyst, DSI C-1 (0.1 mol%) provided a mixture of 1,4-addition product 11 with tertiary aldol 3u (47:53 e.r., 54% isolated yield, 11/3u = 1.5:1; see Supplementary Fig. 5). In addition, the silyl enol ether of starting ketone 1u (compound 12, 4%) was also generated (Fig. 3e).

Investigation of the reaction mechanism. To gain insights into the reaction mechanism, real-time analytical investigations were conducted using in situ Fourier transform infrared (FT-IR) spectroscopy (Fig. 4a)^{35,36}. The absorption peak of the carbonyl group of ketone **1a** at 1,688 cm⁻¹ was monitored during the reaction in the presence of 0.1 mol% of IDPi catalyst **C-6** at 20 °C (step A). After 7 min, silyl ketene acetal **2a** (1,656 cm⁻¹) was added to the reaction mixture (step B). The reaction proceeded rapidly and was completed in less than 3 min (>99% conversion by ¹H NMR spectroscopy) to give aldol **3aa** (Fig. 4a, left, 1,744 cm⁻¹) as a single product with an e.r. of 95:5. As previously mentioned, DSI **C-1** did not catalyse the desired transformation but exclusively gave silyl enol ether **4** instead

 Table 2 | Substrate scope for the Mukaiyama aldol reaction of ketone 1 with silyl ketene acetal 2a using different IDPi catalysts, and synthetic utilities of product 3



Reactions were performed with ketone 1 (0.5 mmol), silyl ketene acetal 2a (2.0 equiv.) and IDPi catalyst (0.05 mol% = 500 ppm) in Et₂O (2.5 ml, 0.20 M) at -20 °C for 12-72 h. Gram-scale reactions were performed with ketone 1 (1.0 g), silyl ketene acetal 2a (1.1 equiv.) and IDPi catalyst (50 or 100 ppm) in Et₂O under the indicated conditions. All yields are those of isolated products and e.r. was determined by HPLC analysis. *Corrected yields of 1,2-adducts are indicated (9-22% of 1,4-adducts were observed, see Supplementary Section 11 for details). *No 1,4-adduct was observed.



Fig. 3 | Mukaiyama aldol reaction of ketones on a preparative scale with catalyst loadings between 0.9 and 25 ppm. a, Reaction details and results. b, Repeated addition of starting materials. c, Reaction progress of the aldol reaction of 1u to 3u catalysed by C-8 (0.9-2.8 ppm). d, Images of crude reaction mixtures with thin-layer chromatography (TLC) analyses (left and middle images: after 1 day and 13 days; TLC eluent: acetone/hexanes = 1:20, left lane: 1u/centre lane: co-spot/right lane: mixture) and dried product 3u (right image: 250 ml round-bottom flask). e, Reaction outcome using DSI C-1 (0.1 mol%) as catalyst. aNo 1,4-adduct was observed. bThe corrected yield of 1,2-adduct is indicated. TON, turnover number.

(Fig. 4a, right, 1,751 cm⁻¹)³⁷. The remarkably different reaction outcomes may be explained by considering counteranion basicities. We measured pK_a values of several different chiral Brønsted acid catalysts (developed in our research group) with reference to some achiral acids in acetonitrile (Fig. 4b, see Supplementary Table 10 for pK_a values of our chiral acid organocatalysts and the corresponding references). In line with our previous investigations^{21,23,25}, we propose that our IDPi catalyst is a precatalyst, which is silylated by

ketene acetal **2a** to give the equivalent of a silylium ion pair **13** as the catalytically relevant Lewis acid (Fig. 4c). Silylium transfer onto the ketone then furnishes an oxocarbenium ion pair, the fate of which is dependent on the counteranion; that is, the DSI anion is sufficiently basic to deprotonate the oxocarbenium ion and furnishes the corresponding enol silane (pK_a of DSI **C**-**1**=8.4). In contrast, the IDPi anion is only a very weak base, enabling the desired Mukaiyama pathway to occur (pK_a of IDPi **C**-**3**=4.5, in acetonitrile).

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Fig. 4 | Study of the reaction mechanism. a, Analytical results of the in situ FT-IR investigation of the catalytic Mukaiyama aldol reaction of ketone **1a** with silyl ketene acetal **2a** using IDPi **C-6** (left) and DSI **C-1** (right) as catalysts. **b**, Measured *pK*_a values of chiral acids (DSI **C-1** and IDPi **C-3**) and reported *pK*_a values of achiral acids in acetonitrile (see Supplementary Section 9 for details). BA, Brønsted acid; LA, Lewis acid. **c**, Proposed reaction mechanism of the IDPi-catalysed Mukaiyama aldol reaction.

Conclusions

In conclusion, we have developed a broadly applicable Mukaiyama aldol reaction of ketones 1 with silyl ketene acetals 2. In the presence of IDPi catalysts, a variety of (hetero)aryl-alkyl and alkyl-alkyl ketones are converted into the corresponding tertiary aldols in almost quantitative yields (up to 99% yield) and with very high to excellent enantioselectivities (e.r. up to 99:1). The developed method is operationally simple and scalable, and the obtained products can be readily derivatized. We also show that

only extremely low catalyst loadings are generally required, with 50–500 ppm being sufficient to obtain the desired products quantitatively, typically in less than 12 h. We also report preparative-scale reactions that require only single-digit ppm loadings and in one case, even less than 1 ppm. Although the reactions under these extreme conditions require purified reagents and extended reaction times (2–80 days), to our knowledge the catalyst loadings reported are unprecedented in asymmetric carbon–carbon bond forming reactions.

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Methods

General procedure for the 30 g scale catalytic Mukaiyama aldol reaction of ketones. Ketone 1u (30.0 g, 205 mmol, 1 equiv.) was placed in a flame-dried J. Young Schlenk flask, equipped with a Teflon-coated magnetic stirring bar. IDPi C-8 (1.1 mg, 5.7×10^{-4} mmol, 2.8 ppm) and Et₂O (8.0 M, 25.6 ml) were added at 25 °C and stirred for 30 min. The reaction mixture was cooled to -78 °C and silyl ketene acetal 2a (42.5 g, 226 mmol, 1.1 equiv.) was slowly added. The reaction mixture was stirred for 60 min at -10 °C, then for 60 min at 0 °C, and finally stirred for 13 days at 5 °C. Organic volatiles (boiling points of volatiles: Et₂O=34.6 °C; silyl ketene acetal 2a = 62 °C/12 mbar) were evaporated in vacuo to afford the desired tertiary aldol product 3u (59.2 g, 86% yield, 95:5 e.r.).

Full experimental details and the characterization of new compounds are provided in the Supplementary Information.

Data availability. The data that support the findings of this study are available from the corresponding author upon request.

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Author contributions

H.Y.B. developed the reaction and investigated the substrate scope, derivatizations of the aldol products, and implemented in situ FT-IR study. D.H. first observed the high activity of IDPi catalysts in the described reaction. The IDPi catalysts were developed by H.Y.B., P.S.J.K. and B.L. H.Y.B., P.S.J.K., P.K. and S.L. synthesized the IDPi catalysts used in this study. H.Y.B., C.K.D. and A.D. investigated large-scale and low-catalyst loading experiments. K.K. and I.L. measured pK_x values of acid catalysts. B.L. designed and oversaw the project. H.Y.B. and B.L. wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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