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Enantioselective Hydrogenation of Activated Aryl Imines Catalyzed by an Iron(II) P-NH-P' Complex

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ABSTRACT: Chiral amines are key building blocks in synthetic chemistry with numerous applications in the agricultural and pharmaceutical industries. Asymmetric imine hydrogenation, particularly with iridium catalysts, is well developed. However, imine reduction still remains challenging in the context of replacing such a precious metal with a cheap, nontoxic, and environmentally friendly substitute such as iron. Here we report that an unsymmetrical iron P-NH-P' catalyst that was previously shown to be effective for the asymmetric hydrogenation of aryl ketones is also a very effective catalyst for the asymmetric hydrogenation of prochiral aryl imines activated with *N*-diphenylphosphinoyl or *N*-tosyl groups. The P-NH-P' abbreviation stands for (*S*,*S*)-PPh₂CHPhCHPhNHCH₂CH₂PiPr₂. DFT results suggest that, surprisingly, the NH group on the catalyst activates and orients the imine to hydride attack by hydrogen-bonding to the PO or SO group on the imine nitrogen, as opposed to the imine nitrogen itself. This may explain why *N*-Ph and *N*-Bu imines are not hydrogenated.

INTRODUCTION

The preparation of enantiopure amines is important for providing synthetic building blocks for bioactive molecules and chiral auxiliaries. The asymmetric hydrogenation (AH) of imines represents one of the most direct and atom economical approaches. The significance of asymmetric imine hydrogenation in the context of pharmaceutical and agricultural industries has been reviewed extensively.¹⁻⁸ To date, large scale AH processes have been predominantly developed with precious metal catalysts based primarily on iridium,⁹⁻¹² and rhodium¹³ while ruthenium complexes are often used in asymmetric transfer hydrogenation (ATH) processes.^{10,14,15}

Imines are generally challenging substrates to reduce in a stereoselective manner. They are prone to isomerization between E and Z forms. As the amine product builds up in the AH process, it can coordinate to the metal and inhibit catalysis by preventing dihydrogen coordination. In order to counter this, placing a bulky sulfonyl or diphenylphosphinoyl substituent on the imine nitrogen can hinder the binding of the substrate and product amine to the metal. These groups also have the advantage of activating the imine to hydride attack by increasing the electrophilicity of the imine carbon. The nitrogen substituent is readily cleaved to produce valuable homochiral amines.¹⁶ Thus activated imine catalytic AH has received attention with complexes based on rhodium,¹⁶ palladium,¹⁷⁻²¹ and ruthenium.²²

Reports of the use of first row transition metal complexes for the catalytic AH of imine are limited. In 2011 Beller and coworkers reported that a range of *N*-aryl substituted imines were hydrogenated to amines with *ee* ranging from 88 to 98% using Knölker's iron complex FeH(CO)₂(Cp'OH) (5 mol%) in combination with the homochiral phosphoric acid (*S*)-TRIP (1 mol%) at 50 bar H₂ and 65 °C.²³ In 2014 our group reported that an iron complex [FeBr(CO)₂(L¹)]BF₄ with the asymmetric ligand (*S*,*S*)-PPh₂CHPhCHMeNHCH₂CH₂PiPr₂ (L¹) when activated with LiAlH₄/*iso*-amylalcohol was active in the presence of 10 mol% KO^tBu for the AH of the *N*-diphenylphosphinoyl imine derived from 1-propiophenone to the amine (90% *ee*) at 20 atm H₂ and 50 °C.²⁴ Bai et al have described the use of homochiral cyclopentadienone)iron complex for the AH of activated imines but the enantioselectivity was poor (ee of the amine up to 40%).²⁵ Otherwise there have been several reports of the ATH of activated imines by catalysts based on iron,²⁶⁻²⁹ cobalt^{30,31} and nickel.³²

RESULTS AND DISCUSSION

In pursuit of a highly enantioselective iron catalyst for direct hydrogenation, our group found that the iron complex FeHCl(CO)(P-NH-P') **1** with the P-NH-P' ligand (*S*,*S*)-PPh₂CHPhCHPhNHCH₂CH₂PⁱPr₂ gave superior results to complex FeHCl(CO)(L¹) for the AH of aryl ketones and was easier to synthesize.³³ Here we report that the iron complex **1** permits the highly enantioselective hydrogenation of a range of activated imines.

We chose the (*E*)-*N*-diphenylphosphinoyl imine derived from acetophenone (**2a**) as the model substrate, and probed various catalytic conditions for hydrogenation (Table 1). An increase in the catalyst loading from 1 mol% to 3 mol% with a KO^tBu loading of 10 mol% in toluene at 80°C under 30 bar of hydrogen pressure resulted in a systematic increase in the yield of product after 20 h.(entries 2-4) Increasing the base to catalyst ratio reduced the yield and caused a degradation in the *ee* (entries 1-2); this suggests that a large excess of base to catalyst may cause undesirable side reactions. The optimal catalytic condition resulted in an 85% yield of the amine in 98.3% *ee* (*S*) (entry 4). The use of 15 bar H₂ gave a slightly lower yield of amine with the identical *ee*, suggesting a minor dependence of hydrogen concentration on the rate of the reaction. There was no degradation of *ee* in the product on going from 10 h of reaction with 64% isolated yield (entry 6) to 20 h with 85% yield (entry 4). The influence of various bases was examined (entries 7-10). Alkali metal *tert*-butoxides gave similar results while catalysis with potassium hydroxide resulted in a product with a slightly degraded *ee*. These results suggest that the Lewis acidity of the alkali metal cation is not a factor. No reaction was observed when potassium carbonate was used. The solvents THF and EtOH were less effective, giving lower yields and *ee* (entries 11 and 12) while iPrOH gave comparable reactivity and stereoselectivity as the toluene system (entry 13).

With the optimized reaction condition in hand, we investigated the scope of the AH with respect to the prochiral imine structures (Table 2). In general, the substrates were reduced with high enantioselectivity (90-99%). A lower yield and *ee* was observed with added bulk on the alkyl group (**2a** versus **2b**). Substrates with a naphthyl group (**2c** and **2d**) were reduced with exceptionally high *ee* of 98% and 99%. The addition of an electron withdrawing chloro substituent at the *para-* or *meta-*position of the aryl ring (**2e** and **2f**) resulted in a higher yield but similar ee compared to that of an electron donating methoxy group at the *para-* or *ortho*-position (**2g** and **2h**). Additionally, the oxygen- or sulfur-containing heterocycle-substituted *N*-phosphinoylimines (**2i-k**) could be reduced to amines with at least 90% *ee*. The thiophene substituent was well tolerated; however the 5-chloro-2-thiophene and furyl derivatives underwent side reactions that resulted in a lower yield of the amine product. Amine **3l** derived from indanone was obtained in low yield and in moderate *ee* (79%), and **3m**, derived from 2-heptanone, in low yield and low *ee*.

Next, we investigated the catalytic performance of the iron complex **1** with ketimines with various substituents on the imine nitrogen. The AH of the *N*-tosylimine (**4a**) produced the amine in comparable yield and *ee* to *N*-phosphinoylimines. The product amine (**5a**) was recrystallized and characterized by X-ray crystallography to reveal the absolute configuration to be (*S*). The cyclic *N*-sulfonylimine derived from saccharine (**4b**) was resistant to AH and afforded a low yield of amine with low enantioselectivity. The *N*-phenylimine (**4c**) and *N*-butylimine (**4d**) could not be hydrogenated under these conditions.

Table 1. Optimization of Catalytic Conditions for the Iron-catalyzed AH of Imine 2a^a



entry	solvent	Τ (°C)	catalyst	base	base	H ₂ pres-	time (h)	yield (%)	ee (%) ^b	
			loading (mol%)		loading (mol%)	sure (bar)				
1	Toluene	80	1	KO ^t Bu	5	30	20	27	99.3	
2	Toluene	80	1	KO ^t Bu	10	30	20	11	95.6	
3	Toluene	80	2	KO ^t Bu	10	30	20	51	99.0	
4	Toluene	80	3	KO ^t Bu	10	30	20	85	98.3	
5	Toluene	80	3	KO ^t Bu	10	15	20	80	98.3	
6	Toluene	80	3	KO ^t Bu	10	30	10	64	98.3	
7	Toluene	80	3	NaO ^t Bu	10	30	20	82	98.3	
8	Toluene	80	3	LiO ^t Bu	10	30	20	84	98.3	
9	Toluene	80	3	КОН	10	30	20	87	97.7	
10	Toluene	80	3	K ₂ CO ₃	10	30	20	0	-	
11	THF	60	3	KO ^t Bu	10	30	20	81	97.9	
12	EtOH	80	3	KO ^t Bu	10	30	20	73	97.8	
13	iPrOH	80	3	KO ^t Bu	10	30	20	83	98.3	

^a The reactions were performed in duplicate with 30 bar of H₂, 3.87 µmol of catalyst, 6 mL of solvent, 20 h unless specified. ^b Enantiomeric excess was determined by HPLC on a chiral stationary phase.

Table 2. Iron-catalyzed AH of N-diphenylphosphinoyl Ketimines

$ \begin{array}{c} $	$ \begin{array}{c} $	0 1 1 1 1 1 1 1 1 1 1 1 1 1	
entry	Ar/R	Yield of 3 (%)	ee (%) of 3 (S) ^b
1	Phenyl/Me (2a)	85	98
2	Phenyl/Et (2b)	81	94
3	1-naphthyl/Me (2c)	85	99°
4	2-naphthyl/Me (2d)	78	98
5	<i>p</i> -chlorophenyl/Me (2e)	84	98
6	<i>m</i> -chlorophenyl/Me (2f)	81	94
7	<i>p</i> -methoxyphenyl/Me (2g)	76	96
8	<i>o</i> -methoxyphenyl/Me (2h)	74	96
9	2-furyl/Me (2i)	58	92
10	2-thiophene/Me (2j)	82	95°
11	5-chloro-2-thio-	59	90
	phene/Me (2k)		
12	21	42	79
13	2m	14	35

^a The reactions were performed with 3.87 μmol of catalyst (3 mol%), 0.0129 mmol KO^tBu (10 mol%), 0.129 mmol substrate, 30 bar of H₂, 6 mL of toluene, 80°C, 20 h. ^b Enantiomeric excess was determined by HPLC on a chiral stationary phase. ^c The absolute configuration was determined to be (*S*) by single crystal X-ray crystallography.

Table 3. Iron-catalyzed AH of N-tosyl, Cyclic N-sulfonyl and N-aryl and N-Butyl Ketimines^a

entry	substrate	yield (%)	<i>ee</i> (%) ^b
1		82	97°
2	O S N 4b	26	55
3,4	4c R = Ph $4d, R = Bu$	<1	-

^{a-c} See the footnotes of Table 2.

In order to gain insight into the catalytic mechanism, density functional theory (DFT) calculations were performed using the M11L functional with the inclusion of toluene solvation. The favored mechanism, as shown in an abbreviated fashion in Scheme 1, where the *trans*-dihydride **8** transfers a hydride to the imine in the outer coordination sphere. Complex **8** is proposed to form from the catalyst precursor **1** by the heterolytic splitting of dihydrogen to the added base, presumably *via* complex [FeH(H₂)L](OtBu) **7** when KOtBu is used. The heterolytic splitting of the η^2 -H₂ ligand *via* **TS**^E produces the dihydride **8**. The turnover-limiting transfer of a hydride **8** to the imine *via* **TS**^A for the phosphinoylimine (Figure 1) or **TS**^C for the *N*-sulfonylimine *N*-1-phenylethylidene-phenylsulfonamide (**imine**2, Figure 2) results in the formation of an outer-sphere, ion-paired amide complex **9** with an Fe-NH...O hydrogen bond. The NH proton remains on the nitrogen. The amine can form by protonation of the amide by an alcohol complex [FeH(OHtBu)(CO)(P-NH-P')]⁺ (Scheme 2).

A notable feature of this mechanism is the hydrogen bond formed between the N-H proton on the P-NH-P' ligand and the P=O moiety of the *N*-phosphinoyl group in \mathbf{TS}^{A} or a S=O moiety of the *N*-sulfonyl group in \mathbf{TS}^{C} to form an eight-membered transition state. The NH-O interaction helps activate the imine carbon to nucleophilic attack and enhances the enantioinduction by anchoring the substrate to the catalyst. This type of ligand-substrate interaction in activated imine hydrogenation has not been proposed previously. The closest case is the proposed Al-O=P interaction of *N*-phosphinoylimines in aluminum-mediated stoichiometric reductions.³⁴ In the previous studies of FeH₂(CO)(P-NH-P') complexes, the enantiodetermining step in the AH of ketones was proposed on the basis of density functional calculations (DFT) to be hydride transfer to the ketone which is hydrogen-bonded to the NH of the P-NH-P' ligand (Figure 1, **TS**^B).^{33,35} Similarly, The N-H proton is commonly proposed to interact with the imine nitrogen and act as proton donor to unactivated imines.³⁶⁻³⁸ However the nitrogen of these activated imines are sterically hindered and not basic. The TS structure **TS**^{G-S} with the imine nitrogen of **imine2** accepting a hydrogen bond from the catalyst's NH (Figure S5) has an energy that is 7 kcal/mol higher in energy than that of **TS**^{C-S} and a TS frequency of -1150 cm⁻¹. This could be the explanation for the inactivity of the catalyst toward *N*-phenyl and *N*-butyl imine hydrogenation where a comparable activation is not present.

Scheme 1. Proposed Catalytic Cycle for the Asymmetric Imine Hydrogenation Catalyzed by an Iron P-NH-P' Complex.



Similar to the ketone hydrogenation transition state TS^{B} ,³³ the smaller methyl group on the substrate in TS^{A} or TS^{C} is pushed against a phenyl of the PPh₂ group while the larger phenyl substituent on the imine carbon is oriented toward the open side of the catalyst (Figures 1, 2). The hydride addition results in the (*S*) configuration of the amine. For the imine, the aryl of the sulfonyl group or a phenyl of the OPPh₂ group fits into a gap between the PⁱPr₂ group and the axial NH group.



Figure 1. The proposed transition state structure (TS^{A-S}) for the transfer of a hydride from the dihydride complex to the activated *N*-phosphinoylimine **2a** compared with that to acetophenone (TS^B) to give the (*S*) imine or alcohol, respectively.



Figure 2. The proposed transition state (TS^{C-S}) for the hydride transfer from iron to **imine2** (ChemDraw view and ChemCraft view with most of the hydrogens deleted) on the basis of DFT (Gaussian09/M11L/6-31+G*/SDD for Fe/PCM toluene, ΔG^{\ddagger} 34 kcal/mol relative to the dihydride and free imine, 531*i* cm⁻¹, 80 °C). Selected bond distances around the 8 membered substrate-catalyst interaction: Fe...H 1.66, H...C 1.77, C-N 1.32, N-S 1.60, S-O 1.45, O...H 1.97, H-N 1.03, N-Fe 2.05 Å.

A complete cycle (Scheme 2) was calculated for the AH of **imine2** to the amine *S*-PhMeCHNHSO₂Ph (**amine2**) and the free energy profile of the reaction is shown in Figure 3. The alkoxide complex **6** is the turn-over limiting intermediate, the lowest point in the reaction profile. The splitting of dihydrogen to the alkoxide *via* **TS**^E is a lower energy process than the hydride transfer **TS**^{C-S}. Therefore the energy span of the reaction is 27 kcal/mol, consistent with a reaction requiring a temperature above ambient. The off-cycle amido complex **15** with the amide *N*-bonded to iron, as shown in the center of the cycle, is sterically hindered and is only 4 kcal/mol more stable than the outersphere amide complex **9** and is not a stable species relative to the coordinated tBuOH complex **13** or the alkoxide complex **6** plus the **amine2** product. This is not the case for the catalytic cycle involving the hydrogenation of the cyclic *N*-sulfonylimine **4b** where the amide, when *N*-bonded to iron in the amido complex **16** (Figure 4), is more stable than the alkoxide complex (plus the energies of the imine plus H₂). Thus the most stable species in the cycles are the alkoxide complex **6**, or in the case of the small imine **4b**, the amido complex **16**. The need for the ratio KO'Bu:Fe 3.3:1 may reflect a balance between favoring the dihydrogen splitting process by an increase in concentration of the base and inhibiting dihydrogen coordination by competitive coordination of the alkoxide to the iron in **6**.

Scheme 2. Possible Catalytic Cycle for the AH of imine2 Where the Dihydride Complex 8 Transfers a Hydride to the Imine in TS^{C-}





Figure 3. Reaction coordinate diagram corresponding to the catalytic cycle shown in Scheme 2.



Figure 4. Structure of the amido complex **16**, with the amide form of (*S*)-amine **5b** coordinated to [FeH(CO)(P-NH-P')]⁺. Non-essential hydrogens have been deleted for the sake of clarity.

Attack on the opposite face of the imine *via* TS^{A-R} (Figure 5) or TS^{C-R} (Figure S2) produces a transition state 6-7 kcal/mol higher in energy than that of TS^{A-S} or TS^{C-S} ; the *pro-R* calculated structures have the larger phenyl group against the PPh₂ phenyl. Calculations of the hydride transfer to the cyclic *N*-sulfonylimine **4b** gave a small energy difference between *pro-S* (TS^{F-S}) and *R* (TS^{F-R}) transition states caused by the more compact size of the substrate (Figure S3).



Figure 5. Structure of the proposed *pro-S* transition state (at left) **TS^{A-S}** (522*i* cm⁻¹) and *pro-R* transition state (at right) **TS^{A-R}** (544*i* cm⁻¹) for the hydride transfer to the *N*-phosphinoyl imine **2a** from the FeH₂(CO)(P-NH-P') catalyst **8.** Non-essential hydrogens have been deleted for the sake of clarity. **TS^{A-S}** is 29 kcal/mol relative to complex **8** plus imine or 32 kcal/mol relative to Fe-O'Bu complex **6**; **TS^{A-R}** 38 kcal/mol.

Figure 6 shows the energy span of the reaction from the lowest energy species in the system: from the alkoxide complex **6** or the amido complex FeHL(*S*-amide3) **16** to the enantio-determining hydride transfer to imine transition state (TS) for the imines, **imine1** (**2a**) **imine2** (PhMeC=NSO₂Ph) and **imine3** (**4b**) where [FeHL]⁺(amide)⁻ is **9** in Scheme 1. Thus the net reactions correspond to:

Imine1 + FeH(OtBu)L + H ₂ \rightarrow [FeHL] ⁺ (amide1) ⁻ + HOtBu (via TS ^{A-S} or TS ^{A-R})	(1)
Imine2 + FeH(OtBu)L + H ₂ \rightarrow [FeHL] ⁺ (amide2) ⁻ + HOtBu (via TS^{C-R} or TS^{C-R})	(2)

Imine3 + FeH(amide3)L + H₂ \rightarrow [FeHL]⁺(amide3)⁻ + amine3 (*via* TS^{F-S} or TS^{F-R}) (3)

The *pro-S* transition state for **imine1** is found to be 6 kcal/mol lower in energy than the *pro-R*, representing a theoretical ee of 99.9% (*S*) ion reasonable agreement with the experimental value of 98% ee (*S*). The energy span of 32 kcal/mol is consistent with the experimental conditions where 353 K is required for the reaction to proceed. The alkoxide complex **6** was found to be lower in energy than the dihydride complex **8** plus free HOtBu by 2.7 kcal/mol (also true for **imine2** but not shown in Figure 6). Similarly the *pro-S* transition state for **imine2** is found to be 7 kcal/mol lower in energy than the *pro-R*, consistent with the observed ee of 97% (*S*) for the amine **5a** from imine **4a**. The possibility that *t*BuOH shuttles a proton from the catalyst NH to the imine when the hydride transfers from iron to the imine carbon was investigated but all structures were found to be too crowded to allow the imine to approach to receive the proton. The energy span for the more compact **imine3** (**4b**) is determined starting from the amido complex **16** produced from a previous hydride transfer step and going to the TS^F transition states; the amido complex **16** is lower in energy than alkoxide complex **6** in this case when additional components are added to conserve the number of atoms. No difference was calculated for the *pro-R* and *pro-S* transition states for the AH of **imine3**, consistent with the low ee of 55% (*S*) for amine **5b** which represents a Δ G of 0.8 kcal/mol, within the error range of the calculation. In all cases the hydride transfer occurs before a proton transfer to the imine.





Figure 6. The energy span of the catalysis for three imines, representing also the enantiodetermining hydride transfer transition state.

An alternative route is one where dihydrogen splits heterolytically between the cationic Fe(II) center and the amide of complex **9** to reform the dihydride **8** *via* **TS**^{**D**} (Figure 7) and where the outer-sphere amide form of the substrate is protonated giving the amine product directly. The energetics of this mechanism (Figure 8) are similar to those of alkoxide mechanism (Figure 3) but now the H₂ splitting step is the turnover limiting transition state. After the precatalyst **1** is activated by KOtBu, this route may also be operational.



Figure 7. An alternative transition state TS^D for the heterolytic splitting of dihydrogen to the outersphere sulfonylamide.



Figure 8. Reaction coordinate diagram (kcal/mol) for a model catalytic cycle for the asymmetric hydrogenation of **imine2** catalyzed by the FeH₂(CO)(P-NH-P') catalyst **8** where dihydrogen splits into a hydride on iron and a proton on the nitrogen of the outer sphere amide form of the substrate. The H₂ splitting transition state structure is shown with non essential atoms deleted for clarity. See also Figure S4 for the complete cycle.

We did not find an energetically reasonable mechanism that involved the dissociation of an arm of the tridentate ligand to allow an inner sphere hydride transfer. Inner sphere mechanisms involving dissociation of a ligand to let the bulky imine coordinate and allow hydride migration to the imine carbon were deemed too costly energetically since they involve the breaking of an iron-phosphorus bond and then the high transition state energy for hydride transfer in a very crowded system. The weaker iron-phosphorus bond would be the one to the homochiral PPh₂CHPh part of the ligand; the loss of this bond would likely result in poor enantioinduction. The possibility of paramagnetic species was ruled out by calculating the energies of the S=1 and S=2 structures for the five coordinate iron complex 9, the most likely to have an open shell. The S=1 is 18.2 kcal/mol higher in energy than 9 with S=0, and the S=2, 30.8 kcal/mol higher in energy than 9.

CONCLUSION

We have found a well-defined base-metal precatalyst FeH(Cl)CO(P-NH-P') **1** for the enantioselective synthesis of a range of *N*-phosphinoyl and *N*-tosyl aryl amines by AH. This catalyst has two phenyl groups on the ligand backbone to lock a five membered ring (*S*,*S*)-PPh₂CHPhCHPhNFe into a rigid chiral array of phenyl groups to interact with the imine substrate. It enters the catalytic cycle more directly than a previously reported [FeBr(CO)₂(L¹)]BF₄ system that does not have a hydride or NH group and **1** is more enantioselective than the previous version that had a more flexible ligand backbone with a smaller methyl and phenyl on the backbone. The small, cyclic imine **4b** is proposed, on the basis of DFT calculations to form a stable amido iron complex after the hydride transfer which limits turnover. The other activated imines are bulky enough to avoid forming such a stable complex so that a *tertiary*-butoxide complex becomes the turnover limiting intermediate according the calculations. The calculations revealed that the role of the oxygen of the substituent on the activated imine nitrogen is to hydrogen bond with the N-H group of the iron catalyst and participate in anchoring the *pro-S* face of the C=N bond over the hydride. The hydride transfer transition state with an eight membered -Fe-H-CNEO-HN- ring structure is turnover- and enantio-determining.



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N-phenyl and *N*-butyl imines were not reduced using **1** possibly because they are not electrophilic enough and cannot form this ring for hydride transfer. This new information provides insight into ligand design for future AH catalysts.

EXPERIMENTAL SECTION

General information. All manipulations that involved air- or moisture-sensitive materials were performed using Schlenk techniques or a glovebox under an argon or nitrogen atmosphere. All solvents were degassed and dried using standard procedures prior to all manipulations and reactions unless stated otherwise. ³⁹ (1S,2S)-2-(diphenylphosphino)-1,2-diphenylethylamine was purchased from ACROS Organics and Iron catalyst **1** was prepared according to the previous literature.³³ The catalyst is air sensitive as a solid and in solution. It is stable in a pure nitrogen or argon atmosphere indefinitely. It should be weighed in a glovebox. (*E*)-1-Phenyl-*N*-tosylethanimine (**4a**) and 3-methyl-1,2-benzisothiazole 1,1-dioxide (**4b**) were prepared using previously reported procedure by Wang and coworkers.¹⁸ (*E*)-*N*-(1-Phenylethylidene)aniline (**4c**) was prepared from acetophenone and aniline in toluene.⁴⁰ *N*-(1-Phenylethylidene)butylamine (**4d**) was prepared using acetophenone, *N*-butylamine and titanium tetrachloride in diethyl ether and dichloromethane.⁴¹ Deuterated solvents were purchased from Cambridge Isotope Laboratories, INC, and dried over activated molecular sieves. Column chromatography was performed with SiliaFlash P60 40-63 µm silica gel from SilicycleTM.

NMR spectra of the samples that were prepared under argon in degassed solvents were recorded at ambient temperature and pressure using a Bruker Advance III 400 MHz, Agilent DD2 500 MHz or Agilent DD2 600 MHz spectrometer. ¹H NMR spectra were internally referenced to tetramethylsilane resonance (TMS, 0 ppm). ¹³C{¹H} NMR spectra were referenced to solvent carbon resonances. Data for ¹H NMR spectra are reported as follows: chemical shift (δ ppm), multiplicity (s = singlet, d = doublet, t = triplet, q = quartet, m = multiplet, br = broad), coupling constant (Hz), integration. Enantiomeric excesses (ee) were determined by HPLC analysis performed on an Agilent HP 1100 Series modular system operated by ChemStation LC 3D software, v.10.02. on chiral column DAICEL CHIRALPAK AD-H. Enantiomeric purity (reported as % ee) was determined by comparison with the racemic assay and calculated as follows:

% ee = $[(Area A - Area B)/(Area A + Area B)] \times 100.$

General procedure for preparation of N-phosphinoylimines. *N*-phosphinoylimines were prepared using modified procedure from previous literature.^{42,43} A representative example is the synthesis of (*S*)-*N*-(1-phenylethyl)diphenylphosphinamide (**2a**). A suspension of hydroxylamine hydrochloride (2.78 g, 40 mmol) and sodium acetate (4.10 g, 50 mol) in 80% aqueous ethanol solution (25 mL) was stirred at 25°C for 30 minutes. Then, acetophenone (2.9 mL, 25 mmol) was added and the reaction mixture was heated to reflux for 3 hours. It was cooled down, evaporated of all volatiles under the vacuum and the residue was put through a short silica pad. The filtrate was concentrated *in vacuo* to yield crystalline oxime. It was used without further purification.

In a Schlenk flask, oxime was dissolved in a mixture of DCM and hexanes (1:1, 50 mL) and the solution was cooled down to -41°C with MeCN/dry ice. Triethylamine (3.8 mL, 27.5 mmol) was added dropwise, followed by chlorodiphenylphosphine (4.9 mL, 27.5 mmol). The reaction mixture was stirred overnight, concomitantly warming to room temperature. All volatiles were evaporated in vacuum and filtered over a short pad of Celite, washed with 2x20 mL of DCM. The filtrate was extracted with 2x20 mL water and 20 mL brine and dried over anhydrous MgSO₄. The organic layer was concentrated *in vacuo* to afford the crude ketimine. It was purified with column chromatography (1:1 = ethyl acetate:hexanes), and recrystallized with benzene/hexanes to yield **2a**.

The characterization of *N*-phosphinoylimines 2a, ⁴³ 2b, ²⁷ 2c, ²⁷ 2d, ²⁷ 2e, ⁴³ 2f, ⁴³ 2g, ⁴³ 2h, ⁴³ 2j, ²⁷ 2k, ²⁷ 2l, ²⁷ 2l and 2m²⁶ was consistent with those previously reported in the literature. Racemic mixture of *N*-phosphinoylamines were prepared by NaBH₄ reduction of the respective imines.

General procedure for asymmetric reduction of imines. All of the hydrogenation reactions were performed at constant pressures using a stainless steel 50 mL Parr hydrogenation reactor. The reactor was maintained at the appropriate temperature in oil bath. The reactor was flushed several times with hydrogen gas at 10 bar prior to addition of catalyst and substrate solution, and base solution. In an argon glovebox, a vial with stir bar was charged with the catalyst 1 (2.5 mg, $3.87 \times 10^{-3} \text{ mmol}$), substrate ($1.29 \times 10^{-1} \text{ mmol}$) and 3 mL toluene. For base solution, a separate vial was charged with KOtBu (1.4 mg, $1.29 \times 10^{-2} \text{ mmol}$) and 3 mL toluene. The solutions were stirred for 5 minutes and transferred to two 6 mL syringes with 12 in. needles; the needles were then stoppered well. Both syringes were taken out of the glovebox and injected into the prepared Parr reactor against a flow of hydrogen gas. The reaction was stopped after a given amount of time by exposing it to air, and the reactor was allowed to cool down. The catalytic mixture was filtered through a silica plug and toluene was evaporated *in vacuo*.

Column chromatography (1:9:90 = triethylamine:acetone:DCM for *N*-phosphinoylamine, 1:10 = hexanes:ethyl acetate for *N*-tosylamine, DCM for cyclic *N*-sulfonylamine) was performed to purify the crude mixture.

(*S*)-*N*-(1-phenylethyl)-*P*,*P*-diphenylphosphinamide (3a).²⁶ Off-white crystalline solid. Duplicate average of 35.4 mg (85% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.93-7.89 (m, 2H), 7.84 – 7.81 (m, 2H), 7.51 – 7.22 (m, 11H), 4.43 – 4.36 (m, 1H), 3.17 (dd, *J* = 9.8, 5.7 Hz, 1H), 1.58 (d, *J* = 6.8 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 23.69; 98% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R= 8.3 min (minor), 9.2 min (major).

(*S*)-*N*-(1-phenylpropyl)-*P*,*P*-diphenylphosphinamide (3b).²⁶ Off-white crystalline solid. Duplicate average of 35.2 mg (81% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.88-7.85 (m, 2H), 7.77 – 7.73 (m, 2H), 7.50 – 7.14 (m, 11H,), 4.13 – 4.07 (m, 1H), 3.23 – 3.20 (dd, *J* = 9.7, 6.2 Hz, 1H), 2.04 – 1.98 (m, 1H), 1.87 – 1.80 (m, 2H), 0.79 (t, *J* = 7.4 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 23.66; 94% *ee.* Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R= 7.5 min (minor), 9.8 min (major).

(*S*)-*N*-(1-naphthalen-1-yl-ethyl)-*P*,*P*-diphenylphosphinamide (3c).²⁷ Off-white crystalline solid. Duplicate average of 40.9 mg (85% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.94-7.91 (m, 2H), 7.89 – 7.83 (m, 2H), 7.79 – 7.75 (m, 3H), 7.64 – 7.62 (m, 1H), 7.51 – 7.36 (m, 7H), 7.28 – 7.24 (m, 2H), 5.27 – 5.21 (m, 1H), 3.37 (dd, *J* = 9.3, 5.8 Hz, 1H), 1.70 (d, *J* = 6.7 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.89; 99% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R = 9.0 min (minor), 13.2 min (major). Single crystals of this compound suitable for X-ray diffraction analysis were obtained by diffusion of hexanes into a saturated dichloromethane solution at room temperature.

(*S*)-*N*-(2-naphthalen-2-yl-ethyl)-*P*,*P*-diphenylphosphinamide (3d).²⁶ Yellow crystalline solid. Duplicate average of 37.5 mg (78% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.95-7.91 (m, 2H), 7.85 – 7.77 (m, 5H), 7.68 – 7.64 (m, 1H), 7.51 – 7.41 (m, 7H), 7.34 – 7.31 (m, 2H), 4.60 – 4.53 (m, 1H), 3.29 (dd, *J* = 9.8, 5.8 Hz, 1H), 1.67 (d, *J* = 6.7 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.66; 98% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R= 9.1 min (minor), 9.7 min (major).

(*S*)-*N*-(1-(4-chlorophenyl)ethyl)-*P*,*P*-diphenylphosphinamide (3e).²⁶ Off-white crystalline solid. Duplicate average of 38.7 mg (84% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.92-7.88 (m, 2H), 7.82 – 7.78 (m, 2H), 7.52 – 7.36 (m, 6H), 7.28 – 7.21 (m, 4H), 4.41 – 4.34 (m, 1H), 3.19 (dd, *J* = 9.8, 5.7 Hz, 1H), 1.56 (d, *J* = 6.7 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 23.18; 98% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 15:85, 1.0 mL/min, t_R= 12.2 min (minor), 12.9 min (major).

(*S*)-*N*-(1-(3-chlorophenyl)ethyl)-*P*,*P*-diphenylphosphinamide (3f).⁴⁵ Off-white crystalline solid. Duplicate average of 37.0 mg (81% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.93-7.87 (m, 2H), 7.83 – 7.77 (m, 2H), 7.53 – 7.35 (m, 6H), 7.25 – 7.14 (m, 4H), 4.42 – 4.32 (m, 1H), 3.14 (dd, *J* = 9.7, 5.6 Hz, 1H), 1.56 (d, *J* = 6.8 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.64; 94% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R= 6.1 min (minor), 8.7 min (major).

(*S*)-*N*-(1-(4-methoxyphenyl)ethyl)-*P*,*P*-diphenylphosphinamide (3g).²⁶ Off-white crystalline solid. Duplicate average of 34.4 mg (76% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.91-7.88 (m, 2H), 7.84 – 7.81 (m, 2H), 7.50 – 7.34 (m, 6H), 7.24 – 7.18 (m, 2H), 6.86 – 6.80 (m, 2H), 4.38 – 4.32 (m, 1H), 3.15 (dd, *J* = 9.6, 5.9 Hz, 1H), 1.55 (d, *J* = 6.8 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.78; 96% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R= 9.7 min (minor), 10.7 min (major).

(*S*)-*N*-(1-(2-methoxyphenyl)ethyl)-*P*,*P*-diphenylphosphinamide (3h).²⁶ Yellow crystalline solid. Duplicate average of 33.6 mg (74% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.88 – 7.84 (m, 2H), 7.82 – 7.76 (m, 2H), 7.49 – 7.38 (m, 4H), 7.36 – 7.31 (m, 2H), 7.23 – 7.21 (m, 1H), 7.06 – 7.05 (m, 1H), 6.90 - 6.85 (m, 2H), 4.50 – 4.44 (m, 1H), 3.93 (dd, *J* = 11.0, 7.8 Hz, 1H), 3.75 (s, 3H, OCH₃), 1.57 (d, *J* = 6.8 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.94; 96% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 10:90, 1.0 mL/min, t_R= 22.5 min (major), 23.9 min (minor).

(*S*)-*N*-(1-furylethyl)-*P*,*P*-diphenylphosphinamide (3i).²⁶ Yellow crystalline solid. Duplicate average of 23.4 mg (58% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.97-7.86 (m, 4H), 7.51 – 7.40 (m, 6H), 7.35 (dd, *J* = 1.9, 0.9 Hz, 1H), 6.28 (dd, *J* = 3.17, 1.88 Hz, 1H), 6.13 (dt, *J* = 3.1, 0.8 Hz, 1H), 4.45 – 4.36 (m, 1H), 3.35-3.30 (m, 1H), 1.60 (d, *J* = 6.8 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.85; 92% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R= 6.9 min (major), 7.8 min (minor).

(*S*)-*N*-(1-thiophenylethyl)-*P*,*P*-diphenylphosphinamide (3j).²⁶ Off-white crystalline solid. Duplicate average of 34.6 mg (82% yield).¹H NMR (400 MHz, CDCl₃): δ 7.98-7.93 (m, 4H), 7.52 – 7.41 (m, 6H), 7.21 (dd, *J* = 5.0, 1.3 Hz, 1H), 6.94 – 6.91 (m, 2H), 4.62 – 4.55 (m, 1H), 3.25 (dd, *J* = 10.8, 5.4 Hz, 1H), 1.70 (d, *J* = 6.7 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.44; 95% *ee*. Product ratio was determined

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by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R = 8.6 min (major), 9.4 min (minor). Single crystals of this compound suitable for X-ray diffraction analysis were obtained by diffusion of hexanes into a saturated dichloromethane solution at room temperature.

(*S*)-*N*-(1-(2-chlorothiophenyl)ethyl)-*P*,*P*-diphenylphosphinamide (3k).²⁶ Yellow crystalline solid. Duplicate average of 27.7 mg (59% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.94-7.88 (m, 4H), 7.50 – 7.38 (m, 6H), 6.72 (d, *J* = 3.8 Hz, 1H) 6.66 (dd, *J* = 3.8, 1.1 Hz, 1H), 4.48 – 4.38 (m, 1H), 3.40 (dd, *J* = 10.7, 5.3 Hz, 1H), 1.62 (d, *J* = 6.6 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.71; 90% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R = 6.4 min (minor), 7.4 min (major).

N-(1-indanyl)-*P*,*P*-diphenylphosphinamide (31).²⁶ Off-white crystalline solid. Duplicate average of 17.8 mg (41% yield). ¹H NMR (400 MHz, CDCl₃): δ 8.04-7.95 (m, 4H), 7.70 – 7.72 (m, 1H), 7.53 – 7.44 (m, 6H), 7.24 – 7.18 (m, 3H), 4.73 – 4.63 (m, 1H), 3.24 (dd, *J* = 11.4, 6.4 Hz, 1H), 2.96 (ddd, *J* = 15.8, 8.7, 2.8 Hz, 1H), 2.80-2.72 (m, 1H), 2.65 – 2.56 (m, 1H), 2.04 – 1.94 (m, 1H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.48; 79% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 20:80, 1.0 mL/min, t_R= 8.2 min (minor), 9.4 min (major).

N-(1-(methyl)pentylidene)-diphenylphosphinamide (3m).²⁶ Off-white crystalline solid. Duplicate average of 5.7 mg (14% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.94 – 7.88 (m, 4H), 7.49 – 7.42 (m, 6H), 3.22 – 3.12 (m, 1H), 2.67 – 2.63 (m, 1H), 1.56 – 1.08 (m, 11 H), 0.86 (t, *J* = 7.1 Hz, 3H); ³¹P{¹H} NMR (121 MHz, CDCl₃): δ 22.04; 35% *ee*. Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 15:85, 1.0 mL/min, t_R= 9.7 min (minor), 7.3 min (major).

(*S*)-*N*-(1-phenylethyl)-4-methyl-benzenesulfonamide (5a).¹⁸ Off-white crystalline solid. Duplicate average of 29.1 mg (82% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.64-7.60 (m, 2H), 7.22 – 7.17 (m, 5H), 7.16 – 7.11 (m, 2H), 4.79 – 4.72 (m, 1H), 4.48 – 4.38 (m, 1H), 2.39 (s, 3H) 1.43 (d, *J* = 6.8 Hz, 3H); 97% *ee.* Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 10:90, 1.0 mL/min, t_R= 12.8 min (minor), 14.1 min (major). Single crystals of this compound suitable for X-ray diffraction analysis were obtained by diffusion of hexanes into a saturated dichloromethane solution at room temperature.

3-Methyl-1,2-benzisothiazoline 1,1-Dioxide (5b).¹⁸ Off-white crystalline solid. Duplicate average of 5.5 mg (26% yield). ¹H NMR (400 MHz, CDCl₃): δ 7.78 (d, *J* = 7.8 Hz, 1H) 7.62-7.66 (m, 1H), 7.53 (t, *J* = 7.5 Hz, 1H), 7.40 (d, *J* = 7.8 Hz, 1H), 4.83 – 4.76 (m, 1H), 4.64 (br s, 1H), 1.63 (d, *J* = 6.7 Hz, 3H); 55% *ee.* Product ratio was determined by HPLC, Chiralpak AD-H column, IPA/Hex = 15:85, 1.0 mL/min, t_R= 12.5 min (major), 13.5 min (minor).

DFT calculations. Density functional theory calculations were performed using the Gaussian09 package⁴⁶ and the M11-L hybrid functional. All atoms were treated with the 6-31G* basis set while iron was treated with the Stuttgart effective core potential SDD. A toluene solvation model was applied. Complete details can be found in the SI.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website. A PDF file of HPLC traces of the products. X-ray crystal structures and absolute configuration determinations of 3c, 3j, and 5a. DFT calculation details and supplementary figures. A file of the DFT calculated coordinates and energies and, in the case of transition states, the imaginary frequency. The crystallographic cif files have been deposited as CCDC 1903117-1903119.

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Notes

The authors declare no competing financial interests.

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REFERENCES

(1) Seo, C. S. G.; Morris, R. H., Catalytic Homogeneous Asymmetric Hydrogenation: Successes and Opportunities. *Organometallics* **2019**, *38*, 47-65.

(2) Zhang, Z. F.; Butt, N. A.; Zhang, W. B., Asymmetric Hydrogenation of Nonaromatic Cyclic Substrates. *Chem. Rev.* 2016, *116*, 14769-14827.

(3) Drommia, D.; Arena, C. G. Recent Advances on Chiral Catalysts for Asymmetric Hydrogenation of C=C, C=N and C=O Bonds. In *Curr. Org. Chem.*; Bentham Science Publishers B.V.: 2016; Vol. 20, p 2552-2590.

(4) Hopmann, K. H.; Bayer, A., Enantioselective Imine Hydrogenation with Iridium-Catalysts: Reactions, Mechanisms and Stereocontrol. *Coord. Chem. Rev.* **2014**, *268*, 59-82.

(5) Ager, D. J.; de Vries, A. H. M.; de Vries, J. G., Asymmetric Homogeneous Hydrogenations at Scale. *Chem. Soc. Rev.* **2012**, *41*, 3340-3380.

(6) Fleury-Bregeot, N.; de la Fuente, V.; Castillon, S.; Claver, C., Highlights of Transition Metal-Catalyzed Asymmetric Hydrogenation of Imines. *Chemcatchem* **2010**, *2*, 1346-1371.

(7) Xie, J. H.; Zhu, S. F.; Zhou, Q. L., Transition Metal-catalyzed Enantioselective Hydrogenation of Enamines and Imines. *Chem. Rev.* **2011**, *111*, 1713-1760.

(8) Hogan, P. C.; Chen, C.-L.; Mulvihill, K. M.; Lawrence, J. F.; Moorhead, E.; Rickmeier, J.; Myers, A. G., Large-scale Preparation of Key Building Blocks for the Manufacture of Fully Synthetic Macrolide Antibiotics. *J. Antibiot.* **2018**, *71*, 318-325.

(9) Blaser, H. U.; Pugin, B.; Spindler, F.; Thommen, M., From a Chiral Switch to a Ligand Portfolio for Asymmetric Catalysis. *Acc. Chem. Res.* **2007**, *40*, 1240-1250.

(10) Verzijl, G. K. M.; De Vries, A. H. M.; De Vries, J. G.; Kapitan, P.; Dax, T.; Helms, M.; Nazir, Z.; Skranc, W.; Imboden, C.; Stichler, J.; Ward, R. A.; Abele, S.; Lefort, L., Catalytic Asymmetric Reduction of a 3,4-Dihydroisoquinoline for the Large-Scale Production of Almorexant: Hydrogenation or Transfer Hydrogenation? *Org. Process Res. Dev.* **2013**, *17*, 1531-1539.

(11) Wei, X. D.; Qu, B.; Zeng, X. Z.; Savoie, J.; Fandrick, K. R.; Desrosiers, J. N.; Tcyrulnikov, S.; Marsini, M. A.; Buono, F. G.; Li,
Z. B.; Yang, B. S.; Tang, W. J.; Haddad, N.; Gutierrez, O.; Wang, J.; Lee, H. W.; Ma, S. L.; Campbell, S.; Lorenz, J. C.; Eckhardt, M.;
Himmelsbach, F.; Peters, S.; Patel, N. D.; Tan, Z. L.; Yee, N. K.; Song, J. J.; Roschangar, F.; Kozlowski, M. C.; Senanayake, C. H., Sequential
C-H Arylation and Enantioselective Hydrogenation Enables Ideal Asymmetric Entry to the Indenopiperidine Core of an 11 β-HSD-1
Inhibitor. *J. Am. Chem. Soc.* 2016, *138*, 15473-15481.

(12) Ružič, M.; Pečavar, A.; Prudič, D.; Kralj, D.; Scriban, C.; Zanotti-Gerosa, A., The Development of an Asymmetric Hydrogenation Process for the Preparation of Solifenacin. *Org. Process Res. Dev.* **2012**, *16*, 1293-1300.

(13) Strotman, N. A.; Ramirez, A.; Simmons, E. M.; Soltani, O.; Parsons, A. T.; Fan, Y.; Sawyer, J. R.; Rosner, T.; Janey, J. M.; Tran, K.; Li, J.; La Cruz, T. E.; Pathirana, C.; Ng, A. T.; Deerberg, J., Enantioselective Synthesis of a γ-Secretase Modulator via Vinylogous Dynamic Kinetic Resolution. J. Org. Chem. 2018, 83, 11133-11144.

(14) Nedden, H. G.; Zanotti-Gerosa, A.; Wills, M., The Development of Phosphine-Free Tethered Ruthenium(II) Catalysts for the Asymmetric Reduction of Ketones and Imines. *Chem. Rec.* **2016**, *16*, 2623-2643.

Mangion, I. K.; Chen, C.-y.; Li, H.; Maligres, P.; Chen, Y.; Christensen, M.; Cohen, R.; Jeon, I.; Klapars, A.; Krska, S.; Nguyen, H.; Reamer, R. A.; Sherry, B. D.; Zavialov, I., Enantioselective Synthesis of an HCV NS5a Antagonist. *Org. Lett.* 2014, *16*, 2310-2313.

(16) Spindler, F.; Blaser, H.-U., The Highly Enantioselective Hydrogenation of N-Diphenylphosphinylketimines with Cationic Rh Ferrocenyldiphosphine Catalysts. *Adv. Synth. Catal.* **2001**, *343*, 68-70.

(17) Wang, Y.-Q.; Zhou, Y.-G., Highly Enantioselective Pd-Catalyzed Asymmetric Hydrogenation of N-Diphenylphosphinyl Ketimines. *Synlett* **2006**, *2006*, 1189-1192.

(18) Wang, Y.-Q.; Lu, S.-M.; Zhou, Y.-G., Highly Enantioselective Pd-catalyzed Asymmetric Hydrogenation of Activated Imines. *J. Org. Chem.* **2007**, *72*, 3729-3734.

(19) Yang, Q.; Shang, G.; Gao, W.; Deng, J.; Zhang, X., A Highly Enantioselective, Pd-TangPhos-catalyzed Hydrogenation of *N*-tosylimines. *Angew. Chem. Int. Ed.* **2006**, *45*, 3832-3835.

(20) Yu, C.-B.; Wang, D.-W.; Zhou, Y.-G., Highly Enantioselective Synthesis of Sultams via Pd-Catalyzed Hydrogenation. *J. Org. Chem.* **2009**, *74*, 5633-5635.

(21) Zhou, X. Y.; Bao, M.; Zhou, Y. G., Palladium-Catalyzed Asymmetric Hydrogenation of Simple Ketimines Using a Bronsted Acid as Activator. *Adv. Syn. Catal.* **2011**, *353*, 84-88.

(22) Ma, X.; Qiao, L.; Liu, G.; Huang, Z., A New Phosphine-Amine-Oxazoline Ligand for Ru-Catalyzed Asymmetric Hydrogenation of N-Phosphinylimines. *Chin. J. Chem.* **2018**, *36*, 1151-1155.

(23) Zhou, S. L.; Fleischer, S.; Junge, K.; Beller, M., Cooperative Transition-Metal and Chiral Bronsted Acid Catalysis: Enantioselective Hydrogenation of Imines To Form Amines. *Angew. Chem. Int. Ed.* **2011**, *50*, 5120-5124.

(24) Lagaditis, P. O.; Sues, P. E.; Sonnenberg, J. F.; Wan, K. Y.; Lough, A. J.; Morris, R. H., Iron(II) Complexes Containing Unsymmetrical P-N-P' Pincer Ligands for the Catalytic Asymmetric Hydrogenation of Ketones and Imines. *J. Am. Chem. Soc.* **2014**, *136*, 1367-1380.

(25) Bai, X. S.; Cettolin, M.; Mazzoccanti, G.; Pierini, M.; Piarulli, U.; Colombo, V.; Dal Corso, A.; Pignataro, L.; Gennari, C., Chiral (cyclopentadienone)Iron Complexes with a Stereogenic Plane as Pre-catalysts for the Asymmetric Hydrogenation of Polar Double Bonds. *Tetrahedron* **2019**, *75*, 1415-1424.

(26) Zhou, S.; Fleischer, S.; Junge, K.; Das, S.; Addis, D.; Beller, M., Enantioselective Synthesis of Amines: General, Efficient Ironcatalyzed Asymmetric Transfer Hydrogenation of Imines. *Angew. Chem. Int. Ed.* **2010**, *49*, 8121-8125.

(27) Mikhailine, A. A.; Maishan, M. I.; Morris, R. H., Asymmetric Transfer Hydrogenation of Ketimines Using Well-Defined Iron(II)-Based Precatalysts Containing a PNNP Ligand. *Org. Lett.* **2012**, *14*, 4638-4641.

(28) Zuo, W.; Lough, A. J.; Li, Y. F.; Morris, R. H., Amino(imino)diphosphines Activate Iron Catalysts in the Asymmetric Transfer Hydrogenation of Ketones and Imines *Science* **2013**, *342*, 1080-1083.

(29) Bigler, R.; Mezzetti, A., Highly Enantioselective Transfer Hydrogenation of Polar Double Bonds by Macrocyclic Iron(II)/(NH)₂P₂ Catalysts. *Org. Process Res. Dev.* **2016**, *20*, 253-261.

(30) Sugi, K. D.; Nagata, T.; Yamada, T.; Mukaiyama, T., Enantioselective Borohydride Reduction of N-diphenylphosphinyl Imines Using Optically Active Cobalt(II) Complex Catalysts. *Chem. Lett.* **1997**, 493-494.

(31) Yamada, T.; Nagata, T.; Ikeno, T.; Ohtsuka, Y.; Sagara, A.; Mukaiyama, T., Optically Active β -ketoiminato Cobalt(II) Complexes as Highly Efficient Catalysts in Enantioselective Borohydride Reduction of Ketones, Imines, and α , β -unsaturated Carboxylates. *Inorg. Chim. Acta* **1999**, *296*, 86-93.

(32) Xu, H.; Yang, P.; Chuanprasit, P.; Hirao, H.; Zhou, J., Nickel-Catalyzed Asymmetric Transfer Hydrogenation of Hydrazones and Other Ketimines. *Angew. Chem. Int. Ed.* **2015**, *54*, 5112-5116.

(33) Smith, S. A. M.; Lagaditis, P. O.; Lüpke, A.; Lough, A. J.; Morris, R. H., Unsymmetrical Iron P-NH-P' Catalysts for the Asymmetric Pressure Hydrogenation of Aryl Ketones. *Chem. - Eur. J.* **2017**, *23*, 7212-7216.

(34) Graves, C. R.; Scheidt, K. A.; Nguyen, S. T., Enantioselective MSPV Reduction of Ketimines Using 2-Propanol and (BINOL)Al^{III}. Org. Lett. 2006, 8, 1229-1232.

(35) Sonnenberg, J. F.; Sues, P. E.; Wan, K. Y.; Morris, R. H., Ketone Asymmetric Hydrogenation Catalyzed by P-NH-P' Pincer Iron Catalysts: an Experimental and Computational Study. *ACS Catal.* **2017**, *7*, 316–326.

(36) Clapham, S. E.; Hadzovic, A.; Morris, R. H., Mechanisms of the H₂-Hydrogenation and Transfer Hydrogenation of Polar Bonds Catalyzed by Ruthenium Hydride Complexes. *Coord. Chem. Rev.* **2004**, *248*, 2201-2237.

(37) Bellows, S. M.; Chakraborty, S.; Gary, J. B.; Jones, W. D.; Cundari, T. R., An Uncanny Dehydrogenation Mechanism: Polar Bond Control over Stepwise or Concerted Transition States. *Inorg. Chem.* **2017**, *56*, 5519-5524.

(38) Hopmann, K. H., Iron/Brønsted Acid Catalyzed Asymmetric Hydrogenation: Mechanism and Selectivity-Determining Interactions. *Chem. - Eur. J.* 2015, *21*, 10020-10030.

(39) Armarego, W. L. F.; Chai, C. L. L., 2009, Purification of Laboratory Chemicals, 6th Ed. ed.; Butterworth-Heinemann: Oxford, .

(40) Lai, P.-S.; Dubland, J. A.; Sarwar, M. G.; Chudzinski, M. G.; Taylor, M. S., Carbon-carbon Bond-forming Reactions of α-carbonyl Carbocations: Exploration of a Reversed-polarity Equivalent of Enolate Chemistry. *Tetrahedron* **2011**, *67*, 7586-7592.

(41) Abdur-Rashid, K.; Lough, A. J.; Morris, R. H., RuHCl(diphosphine)(diamine): Catalyst Precursors for the Stereoselective Hydrogenation of Ketones and Imines. *Organometallics* **2001**, *20*, 1047-1049.

(42) Bezłada, A.; Szewczyk, M.; Mlynarski, J., Enantioselective Hydrosilylation of Imines Catalyzed by Chiral Zinc Acetate Complexes. J. Org. Chem. 2015, 81, 336-342.

(43) Chen, Y.-J.; Chen, C., Enantioselective Strecker-type reaction of phosphinoyl ketimines catalyzed by a chiral Zr-bipyridyldiol catalyst. *Tetrahedron: Asym.* **2008**, *19*, 2201-2209.

(44) Reingruber, R.; Bräse, S., 1,2-Addition of Trialkylaluminium Reagents to *N*-diphenylphosphinoylketimines in the Absence of Any Additional Reagents. *Chem. Commun.* **2008**, 105-107.

(45) Crampton, R. H.; Hajjaji, S. E.; Fox, M. E.; Woodward, S., Reaction Prospecting by ³¹P NMR: Enantioselective Rhodium-DuPhos Catalysed Addition of ZnMe₂ to Diphenylphosphinoylimines. *Tetrahedron: Asym.* **2009**, *20*, 2497-2503.

(46) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Mennucci, B.; Petersson, G. A.; Nakatsuji, H.; Caricato, M.; Li, X.; Hratchian, H. P.; Izmaylov, A. F.; Bloino, J.; Zheng, G.; Sonnenberg, J. L.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; Montgomery, J. A.; Peralta, J. E.; Ogliaro, F.; Bearpark, M.; Heyd, J. J.; Brothers, E.; Kudin, K. N.; Staroverov, V. N.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Rega, N.; Millam, J. M.; Klene, M.; Knox, J. E.; Cross, J. B.; Bakken, V.; Adamo, C.; Jaramillo, J.; Gomperts, R.; Stratmann, R. E.; Yazyev, O.; Austin, A. J.; Cammi, R.; Pomelli, C.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Zakrzewski, V. G.; Voth, G. A.; Salvador, P.; Dannenberg, J. J.; Dapprich, S.; Daniels, A. D.; Farkas, Ö.; Foresman, J. B.; Ortiz, J. V.; Cioslowski, J.; Fox, D. J., Gaussian 09. 2009, Gaussian 09. Revision D.01.