Rational selection of co-catalysts for the deaminative hydrogenation of amides†

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The catalytic hydrogenation of amides is an atom economical method to synthesize amines. Previously, it was serendipitously discovered that the combination of a secondary amide co-catalyst with (iPrPNP)Fe(H)(CO) results in a highly active base metal system for deaminative amide hydrogenation. Here, we use DFT to develop an improved co-catalyst for amide hydrogenation. Initially, we computationally evaluated the ability of a series of co-catalysts to accelerate the turnover-limiting proton transfer during C=N bond cleavage and poison the (iPrPNP)Fe(H)(CO) catalyst through a side reaction. TBD (triazabicyclodecene) was identified as the leading co-catalyst. It was experimentally confirmed that when TBD is combined with (iPrPNP)Fe(H)(CO) a remarkably active system for amide hydrogenation is generated. TBD also enhances the activity of other catalysts for amide hydrogenation and our results provide guidelines for the rational design of future co-catalysts.

Introduction

The selective hydrogenation of carbonyl complexes is one of the most important and widely used catalytic reactions in organic synthesis.1–3 However, the reduction of electron rich carboxylic acid derivatives, such as amides, is still difficult.4,5 The ubiquity of the amide functional group in biological systems, pharmaceuticals, and industrial chemicals has spurred considerable effort to create efficient catalytic systems for amide hydrogenation. Nevertheless, amides are still typically reduced using waste generating stoichiometric reagents, such as LiAlH₄, and to date only a small number of homogenous catalysts can directly hydrogenate amides to amines.5,7–12 These catalysts, which include both precious and base metal systems, provide proof-of-principle that this atom economic transformation is possible, but can still be improved.1

Current mechanistic models for transition metal catalyzed amide reduction, in particular deaminative hydrogenation to produce an amine and an alcohol, propose a sequential reduction of the amide to an intermediate hemiaminal (step 1, Scheme 1), which then undergoes C=N bond cleavage to yield an amine and an aldehyde (step 2). Subsequent hydrogenation of the aldehyde affords the corresponding alcohol (step 3).12,13 Most well-defined catalysts for deaminative hydrogenation rely on a Noyori-type,14,15 bifunctional pathway whereby a metal-hydride and adjacent ligand based proton are delivered to the carbonyl C=O moiety (Scheme 1). Intriguingly, recent mechanistic studies indicate that while the Noyori-type catalyst structure is essential for facilitating the dihydrogen addition steps of the process (1 and 3, in Scheme 1), the proton transfer between the O- and N-ends of the hemiaminal (step 2), which
calculations to design co-catalysts tailored for the deaminative hydrogenation of tertiary amides. Our best co-catalyst, triazabicyclo[5.4.0]decene (TBD), acts as push–pull proton shuttle for C–N bond cleavage, and leads to significant improvement in iron catalyzed deaminative amide hydrogenation. Importantly, the improvement from TBD also occurs for a number of other transition metal catalysts for deaminative amide hydrogenation, suggesting that the addition of co-catalysts of this type is a general strategy for improving amide reduction.

Results and discussion
Computational co-catalyst design
On the basis of the mechanism shown in Scheme 2, DFT calculations were performed on a series of potential organic co-catalysts for the hydrogenation of DMF using FeN (Table 1; see ESI† for computational details). The co-catalysts assessed included molecules with either single site hydrogen bond donors (entries 4–7 and 10) or with both hydrogen bond donor and acceptor sites which could act as push–pull proton shuttles (entries 1–3, 8 and 11). Various aryl and alkyl substituents (i.e. H, Me, iPr, Bu, Ph) were introduced into the pool of co-catalysts, to sample a wide range of stereoelectronic effects. In molecules with C–O or C–N functional groups, only electron-rich systems were chosen to minimize the hydrogenation of the co-catalyst. Although there are co-catalysts that are hydrogenated when used as reactants (e.g. formanilide and acetamide),† when they are used in catalytic concentrations their consumption is slower than that of the reactants, enabling their co-catalytic effect.† The ability of each potential co-catalyst to assist with the hemiaminal proton transfer‡ involved in the C–N bond cleavage (step 2 in Scheme 1) was quantified by computing the transition state(s) associated with this process (ΔGad, Scheme 2 and Fig. 1), which can be either concerted (with TBD, methanol and morpholine) or step-wise (all other co-catalysts). In the latter, the N-protonation of the hemiaminal is followed by its O-deprotonation, which is rate-limiting for all co-catalysts except urea. The thermodynamic preference of the co-catalyst to trap the iron complex was quantified by computing the free energy for the formation of the off-cycle adducts from FeN (ΔGad; Scheme 2 and Fig. 1). In this framework, all co-catalysts were screened with the aim of finding an optimal balance between a low ΔGad and a thermoneutral or endergonic value of ΔGad. Catalyst hydrogenation (ΔGhyd; Scheme 2) competes with adduct formation and, thus, there is an interplay between the free energies of both reactions. In the case of FeN, ΔGhyd (which will depend on the nature of the catalyst) was calculated to be −10.2 kcal mol⁻¹ under the experimental conditions.¹⁰ The value of ΔGad − ΔGhyd (ΔGp, in Table 1) is therefore a measure of how adduct formation may limit the reaction by catalyst poisoning (i.e. a more positive value is indicative of less deactivation). For example, the production of methanol may be expected to inhibit catalysis by adduct formation with FeN (entry 7; ΔGad = −6.9 kcal mol⁻¹). However, catalyst hydrogenation is even more favorable (ΔGhyd = −10.2 kcal mol⁻¹) making the FeH₂N the likely preferred species (ΔGp = 3.3 kcal mol⁻¹).

Scheme 2. Reaction mechanism for the deaminative hydrogenation of amides by Noyori-type catalysts. Isomer = Fe N-bound form of the adduct. Color code: hemiaminal formation (blue), C–N cleavage by proton transfer (red), formaldehyde hydrogenation (green) and adduct formation (black).
Table 1  Evaluation of co-catalysts for the hydrogenation of tertiary amides with FeN

| Entry | Co-catalyst | $\Delta G^\ddagger_{HT}$ | $\Delta G_{add}$ | $\Delta G^\ddagger_{P}$ | TON$^d$ | Conv.$^d$ |
|-------|-------------|--------------------------|----------------|----------------------|--------|---------|
| 1     |             | 21.3                     | -1.5           | 8.7                  | 830    | 59%     |
| 2     |             | 25.3                     | -8.3           | 1.9                  | 780    | 55%     |
| 3     |             | 22.6                     | -12.2          | -2.0                 | 630    | 45%     |
| 4     |             | 24.3                     | -11.5          | -1.3                 | —      | —       |
| 5     |             | 21.8                     | -11.4          | -1.2                 | —      | —       |
| 6     |             | 25.5                     | 4.4            | 14.6                 | 560    | 40%     |
| 7     | CH$_3$OH    | 29.6                     | -6.9           | 3.3                  | 510    | 37%     |
| 8     |             | 22.3                     | -9.3           | 1.4                  | 440    | 31%     |
| 9     | No additive | —                        | —              | —                    | 320    | 22%     |
| 10    |             | 34.6                     | 8.0            | 18.2                 | 320    | 22%     |
| 11    |             | 35.3                     | -8.8           | 1.4                  | 90     | 6%      |

$^a$ $\Delta G^\ddagger_{HT}$ (in kcal mol$^{-1}$) corresponds to the calculated energy of the proton-transfer transition state with the highest energy for DMF assisted by the co-catalysts (Scheme 2, Fig. 1). $^b$ $\Delta G_{add}$ (in kcal mol$^{-1}$) corresponds to the calculated energy of the formation of the adduct (isomer with the lowest energy) formed by [FeN] with the co-catalysts (Scheme 2, Fig. 1). $^c$ $\Delta G^\ddagger_{P} = \Delta G_{add} - \Delta G_{hyd}$ (−10.2 kcal mol$^{-1}$ for all co-catalysts). $^d$ Experimental reaction conditions: 30 atm H$_2$, 5 μmol of [FeN] (0.07 mol%), ($\pm$1.75 mol%) of each additive and 7 mmol of 4-formylmorpholine in 5 mL of THF at 100 °C for 2 h. TON and conv. were determined by GC-FID analysis of the products and remaining starting material. Each entry is the average of two or more trials.

Fig. 1 TSSs and adducts obtained from DFT calculations to compute the assisted proton transfer barrier ($\Delta G^\ddagger_{HT}$) and adduct formation free energy ($\Delta G_{add}$) for the co-catalysts shown in Table 1. (-9.3 kcal mol$^{-1}$), likely due to its lower basicity compared to TBD. Acetanilide (entry 2) also afforded promising results, in this case showing that replacement of H by Me in the originally reported formanilide co-catalyst (entry 3) changed $\Delta G_{add} - \Delta G_{hyd}$ from negative to positive, meaning lower competition of the adduct formation towards amide hydrogenation. Among single site hydrogen bond donors, phenols (entries 4–6) exhibited some promise as a proton shuttle, although sterically large substituents were required to alleviate formation of iron adducts (Adduct$^{a}$, Fig. 1). Interestingly, morpholine and urea yield the largest energy barrier of all of the co-catalysts (34.6 and 35.3 kcal mol$^{-1}$, respectively). This result suggests that a purely basic co-catalyst, although beneficial to prevent adduct formation, does not assist with the hemiaminal proton transfer. Overall, the computational results indicate that the best co-catalysts are those which provide spatially separated hydrogen bond donor and acceptor sites which can act as push–pull proton shuttles, together with a basic character and/or sterically bulky groups to prevent the formation of adducts.

The high co-catalytic activity predicted for TBD (entry 1) was further analyzed by performing microkinetic modelling$^{20–22}$ using the complex reaction network we previously found for amide hydrogenation (see ESI$^{11}$). Under the conditions typically used experimentally (1.4 M of DMF, 0.02 M of TBD, 1 mM of FeN and fixed concentration of 0.162 M of H$_2$, at 100 °C), the microkinetic model yielded a high conversion of 27% over a short reaction time of 2 hours. This conversion is substantially higher than the conversion with formanilide as co-catalyst (12%). The same trend was observed by using 4-formylmorpholine as the substrate (see ESI$^{11}$), a benchmark tertiary amide used in our prior studies on FeN-catalyzed deaminative hydrogenation. In this case, the conversions with TBD and formanilide were 56% and 46%, respectively.

Experimental co-catalyst and catalyst testing

The computational predictions of co-catalyst efficacy were examined experimentally using 4-formylmorpholine (Table 1).
Each potential co-catalyst was tested in catalytic trials with a 1.75 mol% loading, along with 0.07 mol% FeN under previously optimized conditions. Due to the high activity of the catalyst, the variable amount of time required to manipulate the pressure vessel between trials, and the need to equilibrate the vessel at the reaction temperature, it was not possible to acquire reliable initial rate measurements (conversions < 10%). Instead, reaction times were limited to 2 hours to minimize conversion and provide kinetically relevant comparisons. The reaction progress was monitored by amide conversion because of chromatographic issues in quantifying the morpholine product. However, no signals other than starting materials, morpholine and methanol were observed by GC-FID. As predicted by DFT, TBD proved a remarkable co-catalyst, affording a greater than two fold enhancement in TON (compare entries 1 and 9) over the short reaction time. Examination of the influence of TBD loading from 0 to 250 μmol [see ESI; Fig. S7] indicated a strong correlation between TON and TBD, saturating at approximately 200 μmol. The computational results also successfully predicted the relative ability of the other co-catalysts. For additives in which ΔG‡ is positive, the best co-catalysts should be those which lower the ΔG‡R involved in the hemiaminal C–N bond cleavage, as illustrated in entries 1, 2, 6 and 7. In cases where iron deactivation is problematic due to a large negative ΔGadd, then the key barrier to amide hydrogenation is approximated by the total energy difference between ΔGadd − ΔGhyd and ΔG‡R, which explains the superior performance of acetanilide over formanilide (entries 2 and 3). The only discernable variation from this trend is the unexpectedly poor performance of 1,2,3-triphenylguanidine (entry 8), which may react in a different manner as indicated by an immediate color change upon treatment with FeN. The use of urea in the reaction appeared to inhibit the reaction (lower conversion than without additive), which is likely due to other irreversible reactions with the iron species or difficulties in drying the very hydroscopic parent amide. Still, our rational co-catalyst design has led to the identification of a remarkably active catalytic system for selective amide hydrogenation.

The few homogenous transition metal catalysts reported for deaminative hydrogenation are all proposed to follow similar pathways (Scheme 1), with Noyori-type bifunctional mechanisms being prominent.14,15 Given the importance of non-metal mediated hemiaminal cleavage in our computed mechanism, we hypothesized that the co-catalytic enhancements observed here with FeN should be generalizable to other systems. Indeed, highly active ruthenium catalysts recently reported by Beller and Sanford16–20 also exhibit substantial enhancement in activity upon co-catalytic addition of TBD or formanilide (Table 2). The [[(PNP)RuH] (CO)(BH4) (RuBH4)] precatalyst ([PNP] = NH[CH2CH2(PPh2)]2) exhibited a near 4-fold increase in TON for 4-formylmorpholine hydrogenation over a short 2 hour reaction time in the presence of TBD, making it one of the most active systems for hydrogenation of this benchmark substrate. In this case, formanilide inhibits the reaction by forming a stable ruthenium adduct (Fig. S6f). In contrast, with the (PNN)RuH(CO)(BH4) ([RuPNN]) [PNN = 3-di-tert-butylphosphino)-N-[1-methyl-1H-imidazol-2-1]-methyl]propylamine) precatalyst, the relative difference in performance between TBD and formanilide is not as large, likely because the steric bulk of the tert-butyl substituents on the phosphine donors lowers the stability of a formanilide aduct.

The co-catalytic effect of TBD with FeN across different classes of amides was also investigated experimentally (Table 3). Examples of dialkyl and diaryl formamides (entries 1 and 2) exhibited significant enhancement in TON in the presence of TBD compared to the reaction without co-catalyst. N-Phenylacetamide (entry 3), a substrate that previously proved challenging for FeN, was also hydrogenated with greater productivity in the presence of TBD. However, no enhancement was observed upon TBD treatment of the corresponding benzamide (entry 4). This may be due to steric limitation at the carbonyl moiety created by the larger phenyl substituent. In this case, substituting TBD for a smaller co-catalyst provided a modest increase in TON. These results suggest the co-catalytic effect of TBD and related shuttles may be effective with more diverse amides. Admittedly, the enhancement observed with diphenylformanilide was initially unexpected because a mechanism involving the iron-catalyst instead of formanilide, was previously proposed for the hemiaminal C–N bond cleavage using aryl amide substrates.21 However, the calculated ΔG‡R using the

Table 2 Comparison of co-catalysts for amide hydrogenation with pincer supported group 8 catalysts

| Catalyst          | Co-catalyst | TON§ | Conv.§ |
|-------------------|-------------|------|--------|
| [PNP]RuH(CO)(BH4) | None        | 320  | 23%    |
|                   | TBD         | 830  | 59%    |
|                   | HCONNHPh    | 630  | 45%    |
| FeN               | None        | 310  | 22%    |
|                   | TBD         | 1200 | 86%    |
|                   | HCONNHPh    | 0‡   | 0‡     |
| RuBH4H           | None        | 440  | 31%    |
|                   | TBD         | 1170 | 84%    |
|                   | HCONNHPh    | 1040 | 74%    |

* Reaction conditions: 30 atm H2, 5 μmol of [Fe or Ru] (0.07 mol%), 125 μmol of co-catalyst, and 7 mmol of 4-formylmorpholine in 5 mL of THF at 100 °C for 2 h. For [Ru] co-catalysts 10 μmol of NeH2 was added to activate the catalyst. b Determined by GC-FID analysis of the products and remaining starting material. Each entry is the average of two or more trials. c Formanilide reacts irreversibly with this Ru catalyst to form an adduct, see ESI for details.
diphenylformanilide hemiinalmid intermediate and TBD is lower ($\Delta G^\ddagger = 21.8$ kcal mol$^{-1}$) than the barrier for the iron-assisted mechanism ($\Delta G^\ddagger = 28.6$ kcal mol$^{-1}$, see Fig. 2). This result is in agreement with the enhanced reactivity observed for the hydrogenation of diphenylformanilide using TBD as co-catalyst (see Fig. 2).

**Table 3 Co-catalytic enhancement of amide hydrogenations using TBD**

| Entry | Substrate | [TBD] | TON$^a$ |
|-------|-----------|-------|---------|
| 1     | Me        | 1.75  | 300     |
| 2     | Me        | 0.45  | 5180    |
| 3     | Ph        | 1.75  | 230     |
| 4     | Ph        | 1.75  | 120     |

$^a$ Reaction conditions: 60 atm H$_2$, 5 $\mu$mol of [Fe] (0.07 mol%), x $\mu$mol of TBD, and 7 mmol of substrate in 5 mL of THF at 120 $^\circ$C for 16 h. $^b$ TON was determined by GC-FID and NMR analysis of the products and remaining starting material. Each entry is the average of three or more trials. $^c$ TBD was substituted by N-phenylacetamide (Table 1; entry 6).

Fig. 2 Gibbs energies associated with the C–N bond cleavage TSs for diphenylformamide assisted by Fe$^N$ and the TBD co-catalyst.

Conclusions

In conclusion, this work establishes the basis for co-catalyst optimization in amide deaminative hydrogenation reactions using Noyori-type catalysts. Key factors in the co-catalyst design include a push–pull motif of hydrogen bonding sites to assist the C–N bond cleavage of the hemiinalmid and controlled acidity and steric hindrance to prevent catalyst poisoning. Notably, these design principles yielded co-catalysts enhancing the activity of systems based on different transition metals. The generality of the co-catalyst effect and its mechanistic understanding provide new opportunities for the catalytic hydrogenation of challenging electron-rich carbonyl compounds.

Conflicts of interest

There are no conflicts to declare.

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