Cobalt-catalysed alkene hydrogenation: a metallacycle can explain the hydroxyl activating effect and the diastereoselectivity†

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Bis(phosphine)cobalt dialkyl complexes have been reported to be highly active in the hydrogenation of trisubstituted alkenes bearing hydroxyl substituents. Alkene substrates containing ether, ester, or ketone substituents show minimal reactivity, indicating an activating effect of the hydroxyl group. The mechanistic details of bis(phosphine)cobalt-catalysed hydrogenation were recently evaluated computationally (X. Ma, M. Lei, J. Org. Chem. 2017, 82, 2703–2712) and a Co(0)–Co(II) redox mechanism was proposed. However, the activating effect of the hydroxyl substituent and the accompanying high diastereoselectivity were not studied. Here we report a computational study rationalizing the role of the hydroxyl group through a key metallacycle species. The metallacycle is part of a non-redox catalytic pathway proceeding through Co(II) intermediates throughout. The preference for alcohol over ether substrates and the high diastereoselectivity of terpinen-4-ol hydrogenation are correctly predicted in computations adopting the new pathway, whereas the alternative redox mechanism predicts ethers rather than alcohols to be more reactive substrates. Additional experimental evidence supports the role of the hydroxyl group in the metallacycle mechanism. Our work highlights the importance of employing known substrate preferences and stereoselectivities to test the validity of computationally proposed reaction pathways.

Introduction

The majority of homogeneous hydrogenation catalysts employed today are based on precious metals such as rhodium, ruthenium, or iridium. During the last two decades, there has been an increasing focus on developing hydrogenation catalysts involving more earth abundant transition metals. Hydrogenation of alkenes or carbonyl compounds with transition metal complexes based on iron, cobalt, nickel, or manganese has been achieved by several groups, including those of Budzelaar,1 Chirik,2 Casey,3 Milstein,4 Hanson,5 Peters,6 Kempe,7 Morris,8 Fout,9 and Beller.10 Despite these promising results, more efforts are required to develop non-precious catalysts for diastereo- and enantioselective hydrogenation reactions. To date only a limited number of non-precious metal complexes are able to catalyse enantioselective alkene hydrogenations.3,4,11,12

Recently, Chirik and co-workers reported bis(phosphine)cobalt dialkyl complexes for the hydrogenation of alkenes under mild conditions (Fig. 1). A significant activating effect by hydroxyl groups was observed for the cobalt catalysts. Ether, ester, or ketone groups did not provide such an effect. The reported cobalt dialkyl catalyst dppeCo(3CH2SiMe3)2 (C1, dppe = 1,2-bis(diphenylphosphino)ethane) catalyses hydrogenation of...
terpinen-4-ol providing 99% conversion (entry 2, Fig. 1), whereas the corresponding methyl ether displays <5% conversion despite higher catalyst loading and longer reaction time (entry 3, Fig. 1).\(^*\) Interestingly, hydrogenation of terpinen-4-ol gives a high diastereoselectivity with a diastereomeric ratio (d.r.) of 99.8 : 0.2 (entry 2, Fig. 1). Compared to tri-substituted alkenes, di-substituted terminal alkenes could be hydrogenated without a hydroxyl group present (entry 4, Fig. 1).

Recent computational work by Ma and Lei on C1 suggests that hydrogenation of hydroxylated tri-substituted alkenes proceeds through a Co(0)-Co(n) redox mechanism (Fig. 2).\(^*\) The proposed catalytic cycle starts with oxidative addition of H\(_2\) to a Co(0) species generating a Co(n)-dihydride, followed by substrate insertion to give an alkyl intermediate. The inter-mEDIATE may undergo direct reductive elimination to yield the product alkane and regenerate Co(0) (Fig. 2, path (a)) or may proceed via β-hydrogen elimination to form an alkene regioisomer, followed by substrate reinsertion and reductive elimination (Fig. 2, path (b)). Regardless if alkene isomerization occurs, the same product is formed and the overall pathway involves a cycling between Co(0) and Co(n) oxidation states. The role of the hydroxyl group was not considered in the previous analysis.\(^*\)

Here we present computational and experimental results, which provide novel insights into the mechanism of C1-catalysed directed hydrogenation. On basis of our results, we propose that hydrogenation of hydroxylated alkenes occurs through a non-redox reaction pathway proceeding through a metallacycle, which is formed through activation of the hydroxyl group. The metallacycle mechanism correctly predicts the catalyst’s preference for hydroxylated alkenes and the high diastereoselectivity observed in hydrogenation of terpinen-4-ol.

**Results and discussion**

**Computational results**

**Redox pathway.** We performed a quantum chemical analysis of a full molecular model of C1 with different substrates using B3LYP-D3 with the solvent model IEFPCM (for computational details see ESI†). Initially, we evaluated if C1 employs a redox mechanism (Fig. 3), similar to the mechanism proposed by Ma and Lei (Fig. 2).\(^*\) Our redox mechanism differs slightly, as we find that substrate coordination precedes addition of H\(_2\), and we conclude that the intermediate formed upon H\(_2\) coordination prefers a Co(0)-H\(_2\) structure over a Co(n)-dihydride configuration (the dihydride is 2.6 kcal mol\(^{-1}\) higher in energy, Fig. S1, ESI†). The overall hydrogenation steps of ours (Fig. 3) and the previous redox mechanism (Fig. 2) are otherwise identical. The rate-determining step is the formation of the alkyl-intermediate (TS\(_{2,3R}\), Fig. 3), involving transfer of a hydride to the methyl-substituted carbon (the pathway involving hydride transfer to the other carbon has a larger barrier, Fig. S2, ESI†). The alkyl-intermediate then undergoes reductive elimination to form the alkane product (a β-H elimination pathway was also evaluated, but is higher in energy, Fig. S3, ESI†).

In order to evaluate the validity of the redox mechanism, we considered the substrate preference with this pathway. As previously determined in experiments, terpinen-4-ol (Fig. 1, entry 2) is the preferred substrate compared to its methoxy derivative (Fig. 1, entry 3). With the redox mechanism, the overall barriers are calculated to be +23.6 kcal mol\(^{-1}\) for terpinen-4-ol and +23.3 kcal mol\(^{-1}\) for the methoxy-derivative, indicating a slight preference for the methoxy-substrate (Fig. 4). This is in disagreement with the experimentally observed strong preference for the alcohol substrate (Fig. 1). Analysis of the substrate selectivity indicates that C1-catalysed hydrogenation may not proceed through the previously proposed redox mechanism.

**Metallacycle pathway.** We explored several alternative mechanistic possibilities (Fig. S2–S4, ESI†), of which only the preferred metallacycle pathway is discussed here (Fig. 5). We propose that reaction between the precatalyst C1 and the substrate terpinen-4-ol under H\(_2\) results in generation of a cobalt(mono)dihydride(alkoxide) species, which we suggest constitutes the active catalyst 1\(_M\) (Fig. 5). Interestingly,
formation of 1M from the precatalyst C1 is preferred over formation of the dihydride species implicated in the redox mechanism (see Fig. S3 ESI† for computed energies). 1M can then undergo a hydride transfer to the double bond of the bound substrate to form the metallacycle 2M (Fig. 5, for optimised geometry see Fig. 6), which is $\sim$9.8 kcal mol$^{-1}$ below 1M (Fig. 4). Subsequent coordination of H$_2$ to the metallacycle appears facile, with a cost of 11 kcal mol$^{-1}$ relative to 2M (TS$_{2\rightarrow3M}$, Fig. 4). The barrier for H$_2$ attack may be slightly underestimated, given that 3M is $\sim$1 kcal mol$^{-1}$ higher in energy than TS$_{2\rightarrow3M}$. Proton transfer from H$_2$ to the alkyl carbon is the rate-limiting step (TS$_{3\rightarrow4M}$, 21.3 kcal mol$^{-1}$ relative to 2M, Fig. 4) and provides the hydrogenated alkane, which still exhibits a deprotonated hydroxyl coordinating to cobalt. In the final step, another substrate molecule transfers its hydroxyl proton to the alkane oxygen (TS$_{5M}$), regenerating 1M and turning over the

Fig. 4 Computed free energies (kcal mol$^{-1}$) for C1-catalysed hydrogenation. The redox mechanism is shown for substrates terpinen-4-ol (R = OH, blue line) and its methoxy derivative (R = OMe, orange line). The metallacycle mechanism is shown for terpinen-4-ol (black line). All energies are referenced to the 1R state of each substrate.

Fig. 5 Non-redox metallacycle mechanism proposed for bis(phosphine)cobalt-catalysed hydrogenation of terpinen-4-ol. For details on the activation of the precatalyst, see Fig. S5, ESI†
cycle. The cobalt center has a formal oxidation state of +2 throughout (Fig. 5). Although a Co(0) species in principle can be formed in one step from the Co(II) species 1M via the redox mechanism thus raises from 23.6 kcal \( \text{mol}^{-1} \) to 32.6 kcal \( \text{mol}^{-1} \) \((\text{TS}_{2-3M}, \text{Fig. 4})\). The barrier for the metallacycle mechanism is instead 21.3 kcal \( \text{mol}^{-1} \) \((\text{TS}_{3-4M}, \text{Fig. 4})\). Although DFT protocols may exhibit an error of some kcal \( \text{mol}^{-1} \), we consider a difference of 11.3 kcal \( \text{mol}^{-1} \) between the redox and the metallacycle mechanism to be more than significant to conclude that the metallacycle pathway is preferred for hydrogenation of terpinen-4-ol. Thus, a Co(II) open shell complex is transformed to the Co(I) open shell complex, see ESI, Fig. S6, ESI†.

The calculated free energies show that the metallacycle 2M is facile to form and that it is the lowest-lying intermediate computed here (Fig. 4), implying that other reaction pathways must be referenced to 2M, even if the metallacycle constitutes an off-cycle species to these. The barrier for hydrogenation of terpinen-4-ol via the redox mechanism thus raises from 23.6 kcal \( \text{mol}^{-1} \) to 32.6 kcal \( \text{mol}^{-1} \) \((\text{TS}_{2-3M}, \text{Fig. 4})\). Although DFT protocols may exhibit an error of some kcal \( \text{mol}^{-1} \), we consider a difference of 11.3 kcal \( \text{mol}^{-1} \) between the redox and the metallacycle mechanism to be more than significant to conclude that the metallacycle pathway is preferred for hydrogenation of terpinen-4-ol. Thus, a Co(II) open shell complex is transformed to the Co(I) open shell complex, see ESI, Fig. S6, ESI†.

**Analysis of the substrate preference.** The metallacycle mechanism (Fig. 5) is only accessible for substrates containing a hydroxyl group. For other substrates, an alternative hydrogenation mechanism must operate, which may be the redox pathway (Fig. 3). We have compared the computed barriers for three substrates (Fig. 7) to validate our mechanistic proposals.

Hydrogenation of terpinen-4-ol preferentially occurs via the metallacycle pathway, with a computed barrier of 21.3 kcal \( \text{mol}^{-1} \). Redox hydrogenation of the methoxy-derivative of terpinen-4-ol has a computed barrier of 23.3 kcal \( \text{mol}^{-1} \), whereas the lowest computed barrier for redox hydrogenation of \( \alpha \)-methylstyrene is 19.9 kcal \( \text{mol}^{-1} \) (four possible pathways were modelled, see ESI, Fig. S7†). These results are in excellent agreement with experiment (Fig. 1), which showed 99% conversion of terpinen-4-ol and \( \alpha \)-methylstyrene (after 4 and 2 hours, respectively), whereas the methoxy substrate gave <5% conversion after 16 hours. The barrier difference of 2 kcal \( \text{mol}^{-1} \) between terpinen-4-ol (barrier 21.3 kcal \( \text{mol}^{-1} \)) and its methoxy derivative (barrier 23.3 kcal \( \text{mol}^{-1} \)) translates roughly to a ratio of 97 to 3 (Table S1, ESI†), in good agreement with the experimental result of 99% to <5% conversion.

**Analysis of the diastereoselectivity.** The product formed from hydrogenation of terpinen-4-ol can exist as two different diastereomers, which, respectively, have the iPr and methyl substituents cis or trans to each other (Fig. 8). In our computations, the trans-diastereomer is 1.9 kcal \( \text{mol}^{-1} \) lower in energy than the cis-diastereomer, yet in experiment, the cis-diastereomer is predominantly formed, with a very high diastereoselectivity of 99.8 : 0.2 (Fig. 1). This indicates that the hydroxyl group may have a directing effect that favours formation of the cis-isomer. We have here evaluated if the two discussed mechanisms, the metallacycle mechanism and the alternative redox pathway, are able to reproduce the experimentally observed diastereoselectivity.

In the redox mechanism as depicted in Fig. 3, the diastereoselectivity is determined in the insertion step \((\text{TS}_{3-3M})\), where the OH substituent is oriented towards cobalt, leading to the cis-diastereomer. For formation of the trans-diastereomer, we have analysed four pathways (see ESI, Fig. S8–S11†), and find that the lowest pathway proceeds via \( \beta \)-hydrogen elimination (Fig. S10, ESI†). The overall barriers for formation of the cis and trans-diastereomers via a redox pathway differ by 0.6 kcal \( \text{mol}^{-1} \), which corresponds to a predicted d.r. of 75\(^{\text{cis}}\) : 25\(^{\text{trans}}\) (Fig. 8, Table S2 ESI†).

In the metallacycle mechanism, the diastereoselectivity is determined in the hydride transfer step \((\text{TS}_{3-2M})\) (Fig. 5). Due to the strong coordination of the alkoxide oxygen, hydride addition inevitably has to lead to formation of only the cis-product (Fig. 8), in excellent agreement with the experimentally observed high d.r. of 99.8\(^{\text{cis}}\) : 0.2\(^{\text{trans}}\). Formation of the trans-diastereomer would require cleavage of the cobalt-alkoxide bond, which energetically is very costly (Table S3 ESI†). We propose that in the experimental reaction, the cis-
diastereomer is formed via the energetically preferred metallacycle mechanism, whereas the very small amount of observed trans-diastereomer (0.2%, Fig. 1) must be formed via other mechanisms. Our computational results provide a rationale for the high diastereoselectivity of the cobalt complex C1. Interestingly, the Crabtree iridium catalyst provides the same diastereoselectivity with terpinen-4-ol. However, the mechanistic details of said system are not known so far.

Experimental results

Hydrogenation of deuterium-labelled terpinen-4-ol (C_{10}H_{8}-OD) afforded 1,2-H\textsubscript{2} alkane with no deuterium incorporation into the alkene double bond (Fig. S12, ESI†), consistent with the proposed metallacycle mechanism, where proton transfer from the alkene double bond (Fig. S12, ESI†) supports the formation of a mono-alkoxide intermediate, with terpinen-4-ol (Fig. S13, ESI†). This experimental observation was distilled and analysed by 1H-NMR and afforded 1,2-H\textsubscript{2} alkane with no deuterium incorporation into the alkene double bond (Fig. S12, ESI†), consistent with the proposed metallacycle mechanism, where proton transfer from the alkene double bond (Fig. S12, ESI†).

Our attempts to obtain the metallacycle intermediate 2\textsubscript{M} from different synthetic routes and to characterize it by X-ray crystallography were unsuccessful due to strong interference from the thermodynamically more accessible bis(ligand)cobalt species formed during isolation of cobalt complexes. However, indirect experimental evidence suggests that a reaction between the precatalyst dppeCo(CH\textsubscript{2}SiMe\textsubscript{3})\textsubscript{2} mass of Fig. S12, ESI†) and the proposed redox mechanism is further supported by experimental investigations providing indirect evidence of a cobalt-terpinen-4-ol alkoxide intermediate (Co-OR, Fig. 9).

Our computational analysis further shows that a previously proposed redox mechanism\textsuperscript{48} may be valid for non-hydroxylated substrates, but is unable to explain the activating effect and diastereoselectivity of hydroxylated alkenes. As also reported for iron-catalysed hydrogenation of carbonyl substrates,\textsuperscript{19} we have here shown that known substrate selectivities provide a straightforward tool for testing the validity of proposed mechanisms, and we suggest to always assess these important parameters in computational studies of reaction pathways.

Conclusions

We have here reported a new mechanism for bis(phosphine)-cobalt-catalysed hydrogenation of hydroxylated alkenes, proceeding through an energetically low-lying metallacycle intermediate (2\textsubscript{M}, Fig. 5). In the computational analysis, the metallacycle mechanism correctly predicts the preference for hydroxylated alkenes (Fig. 7) and the high diastereomeric ratio observed in hydrogenation of terpinen-4-ol (Fig. 8). The mechanism is further supported by experimental investigations providing indirect evidence of a cobalt-terpinen-4-ol alkoxide intermediate (Co-OR, Fig. 9).

Conflicts of interest

There are no conflicts to declare.

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Notes and references

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