A METATHESIS MODEL FOR ESTER HYDROGEATION BY A RUTHENIUM OCTAHEDRAL COMPLEX USING DFT CALCULATIONS

by

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A thesis submitted in partial fulfillment of the requirements for the degree of Master of Sciences to the Department of Chemistry of the Faculty of Arts and Sciences at the American University of Beirut

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Electronic structure DFT method had been used to investigate the mechanism of ester hydrogenation by a known octahedral ruthenium hydride catalyst having a PNN pincer type ligand. Our results identified a direct low-energy reaction path connecting the ester and the octahedral ruthenium hydride with an octahedral ruthenium alkoxide and an aldehyde. This path represents a novel ion-pair metathesis mode in which a hydride and an alkoxide are exchanged between a metal and an acyl group in an outer sphere mode. A related path is computed for the reaction of a carboxamide. The given reaction path provides a new perspective to understanding the mechanism of the given catalyst that is radically different from the generally accepted metal-ligand cooperation mechanism invoking a formation and reaction of a hemiacetal. The calculations are also used to identify new complexes that may be used to support the proposed mechanism experimentally.
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CHAPTER I

INTRODUCTION

A. Green Chemistry

The concept of greening chemistry is a relatively new idea in the business and regulatory communities that evolved from pollution prevention initiatives.[1] The concept of Green Chemistry has had this large impact due to the fact that it goes beyond the research laboratory in isolation and has touched industry, education, environment, and the general public.[2] In our efforts to improve crop protection, commercial products, and medicines, we also caused unintended harm to our planet and humans. For this fact, the world has been deviating away from classical chemical processes to ones that would rather cause less harm to the environment. One of the major goals of catalysis today is the design of environmentally benign, atom-economical processes, to replace traditional, waste forming reactions, including some of the most fundamental synthetic methods.[3] In addition, due to the resource depletion and environmental concerns, chemists and chemical industries have been working to discover environmentally benign synthetic methods and sustainable catalytic reactions that avoid toxic reagents and waste productions.

The most important aspect of Green Chemistry is the concept of design. Design is a statement of human intention and one cannot do design by accident. It includes novelty, planning and systematic conception. The Twelve Principles of Green Chemistry are “design rules” to help chemists achieve the intentional goal of
sustainability. Green Chemistry is characterized by careful planning of chemical synthesis and molecular design to reduce adverse consequences. [2]

B. Methods for Ester Hydrogenation

Ester hydrogenation has been performed using stoichiometric amounts of the ester and LiAlH₄. This reducing agent is widely used in organic synthesis and it can hydrogenate all carboxylic acids, carbonic acids and their respective derivatives.[4] However, the main drawback of this reaction is the production of hazardous metal oxides and metal hydrides that pose serious environmental concerns especially on the aquatic life. [5]

\[
\begin{align*}
\text{R'CHO} + \text{LiAlH}_4 (2 \text{ moles}, \text{diethyl ether}) & \rightarrow \text{R'CH}_2\text{OH} + \text{R'CHOH} + 2\text{AlH}_3 + 2\text{LiOH} \\
\end{align*}
\]

In addition, this reagent is not ideal from the perspective of cost, safety and simplicity of work-up procedures. LiAlH₄ is extremely flammable and highly reactive towards water, this could possess hazardous threats to the experimental setup. Also, LiAlH₄ is prepared from the reaction of LiH and AlCl₃ followed by filtration to remove LiCl byproducts. [6]

\[
4 \text{LiH} + \text{AlCl}_3 \rightarrow \text{LiAlH}_4 + 3 \text{LiCl}
\]

The use of hydrogen gas for ester reduction would obviously provide an alternative to LiAlH₄ reduction that will be atom economical and won’t generate any waste.[7]
Considering studying esterification and hydrogenation reactions, which are fundamental and significant chemical reactions in synthesis of organic compounds, various methodologies have been established, developed and are being currently used in chemical synthesis.[8, 9] Classically, esterification has been performed using an acid catalyzed reaction that uses stiochiometric ratios of a carboxylic acid and an alcohol and that produces esters and water.[10]

\[ \text{RCOOH} + \text{R’OH} \rightarrow \text{RCOOR’} + \text{H}_2\text{O} \]

Alternative methods have been investigated and esters have been synthesized without the need of a carboxylic acid or any of its derivatives. An attractive approach was developed which focused on the dehydrogenative coupling of alcohols and amines into esters and carboxamides, such reactions would result in the evolution of Hydrogen gas as an only product.

In an attempt to green this chemistry, in recent years new classes of homogeneous catalysts were developed to conduct the hydrogenation reaction with H₂. Reactions mediated by the transition metal complexes have been applied widely both in synthesis and as catalysts to mediate the hydrogenation of esters and carboxamides to alcohols and amines[3]. These reactions can be facilitated by the cooperation of the non innocent ligands and the metal center by undergoing reversible chemical and structural changes. [7, 11, 12]

Milstein and coworkers have designed a new class of transition metal complexes where the ligand is a type of non innocent pincer ligand that supports metal-
ligand cooperation mechanism which appears to involve an aromatization-dearomatization process of the pyridine or acridine-based PNP and PNN “pincer” ligands. These complexes were reported to lead to the unusual X-H (X=H, C, O, N, and B) activation reactions and to environmentally benign catalysis involving dehydrogenation and hydrogenation reactions. Several reviews by Milstein and coworkers have appeared that give an overview of the experimental studies of these chemical reactions mediated by their designed complexes.[8, 13-15]

C. Pincer Ligands

Pincer ligands are one class of chelating agents that bind tightly to three adjacent coplanar sites, a transition metal in a meridional configuration.[16] Such ligands are categorized by the nature of donor atoms along the ligand scaffold and they are usually referred to as DXD, having D indicating the flanking donors and X being the central anchoring donor. [17]

![Diagram of a tridentate pincer ligand](image)

**Figure 1.1:** An illustration of the general structure of a tridentate pincer ligand

Significantly, the realization that pincer ligands offer both a unique, highly protective environment for the coordinated metal center and opportunities to fine-tune the steric and electronic properties of the metal atom has generated extensive research
into the use of these complexes as catalysts.[18] The research on metal-ligand complexes where the latter are phosphine-based pincer complexes has been an influence on numerous researches.[19, 20] The first metal complexes having pincer ligands have the PCP structure, indicating that the ligand consists of two phosphine arms interconnected by a hydrocarbon chain that could be either aromatic or aliphatic.[19] The phosphine arms can also accommodate alkyl or aryl groups, the general structures of such pincer ligands are given below where R could be either an alkyl or an aryl group:

Metal pincer complexes are typically robust structures with high thermal stability and many researchers are currently investigating pincer complexes for various applications ranging from homogeneous catalysis to the development of chemical sensors and switches.[16]

A key feature about pincer based metal complexes is that such metals provide a special balance between the stability of the complex and its reactivity. Both aspects can be optimized by modifying the pincer or part of it, or in other reported cases, modifying the metal center. Such modifications would eventually lead to an enhancement in the metal complex reactivity in addition to increase in selectivity of the complex and providing a new aspect for fundamental insights for particular reaction systems.[19, 21]
Catalyst designs that depicted Ruthenium (II) as its metal center were reported by Van Koten et. al. Their pincer ligand catalyst had NCN and PCP as the ligands and were reported to catalyze the hydrogenation of ketones to their respective alcohols, such reactions could proceed only in presence of a strong base, e.g. KOH. The significance of such reactions were that they were of great attention since they were one of the leading pincer complexes which showed practical simplicity and potential use at ambient pressure; such conditions are critical for the search for optimized systems for essential chemical reactions.[22, 23]

![Figure 1.2: Three different Ru (II) complexes having NCN (1) and PCP (2 and 3) pincer ligands](image)

**D. Bond Activation by Metal-Ligand Cooperation**

PNN Pincer ligands such as ((2-(di-tert-butyl phosphinomethyl)-6-diethyl amino methyl) pyridine) coordinate with the metal via three sites: phosphine arm, pyridine ring and amine arm. These ligands are electron-rich groups and therefore are also classified as electron donating groups so they would have the ability to stabilize the
unsaturated metal center. The planar geometry of the pyridine ring would provide a suitable space for possible metal-substrate interaction.[11, 12, 24]

![Figure 1.3: An Example of a PNN Pincer ligand in 2-D (left) and 3-D structure optimized using Gaussian 09 software and visualized using GaussView](image)

**Figure 1.3:** An Example of a PNN Pincer ligand in 2-D (left) and 3-D structure optimized using Gaussian 09 software and visualized using GaussView

![Figure 1.4: Milstein’s Square pyramidal Ruthenium PNN pincer catalyst (left) and PNP catalyst (right)](image)

**Figure 1.4:** Milstein’s Square pyramidal Ruthenium PNN pincer catalyst (left) and PNP catalyst (right)

Milstein’s catalyst [Ru(PNN)(CO)(H)] is a square pyramidal metal complex that is a prototype of pincer ligated complexes. Such complexes are known to add H₂ reversibly, with a heterolytic bond cleavage of the hydrogen molecule, via a metal-ligand cooperation mechanism in which the proton adds to the methylene linker neighboring the phosphine arm of the ligand and the hydride adds to the metal to form
octahedral Ruthenium complexes in which the hydrides are trans with respect to each other. [11, 12]

**Figure 1.5:** H₂ addition to square pyramidal Ruthenium PNN catalyst via Metal-Ligand Cooperation

In addition to H₂ addition, 1-Ru is reported to perform various other additions to the initially square pyramidal complex. Of these additions, we report here relevant ones to our study, such reactions include the addition of R-OH, and R-NH₂ additions to the 1-Ru square pyramidal complex:

**Figure 1.6:** H₂, R—OH, and R—NH₂ additions to the square pyramidal complex
Milstein’s reported study indicates an efficient and selective class of catalysts that are capable of transforming primary alcohols into esters with an evolution of H₂ gas, the reactions are also found to be done at relatively mild and neutral conditions, unlike the previously reported chemical synthesis where the reaction either should be acid or base catalyzed or a stochiometric amount of a reducing agent is needed.[3, 12]

E. Catalysis by PNN complexes

Milstein’s group reported that the square pyramidal Ruthenium complex (1-Ru), in presence of H₂ gas, catalyzes the hydrogenation of esters and carboxamides into alcohols and amines according to the following equation:

\[
\begin{align*}
R'COO^{-}R' + 2H_2 &\xrightarrow{1-Ru} RCH_2OH + R'OH \tag{2}
\end{align*}
\]

\[
\begin{align*}
R'CONHR' + 2H_2 &\xrightarrow{1-Ru} RCH_2OH + R'NH_2 \tag{3}
\end{align*}
\]

**Figure 1.7:** Equations (2) and (3): Hydrogenation of Esters and carboxamides into alcohols and amines respectively, catalyzed by 1-Ru under H₂ atmosphere

Such a system developed by Milstein and co. which is based on the formation of ester and carboxamide bonds under neutral conditions involves a dehydrogenative coupling of alcohols and amines using 1-Ru as the corresponding catalyst. Remarkably Milstein’s group was able to adapt the reverse of equations (2) and (3) to promote the production of esters and carboxamides by coupling of alcohols and amines under reflux
and using the same square pyramidal complex with a clean process that would only produce \( \text{H}_2 \) gas as a byproduct.[8]

In addition, the dehydrogenative coupling of alcohols by 1-Ru catalyst catalyzes trans esterification and amidation of esters and amines respectively. Such reactions are also of great importance in chemical synthesis and the mentioned chemical processes occur as well in an eco-friendly manner and is atom economical. [25, 26]

\[
\begin{align*}
2 \text{R'}\text{OH} + \text{ROCH}_2\text{R} & \rightleftharpoons 1\text{-Ru} \rightarrow 2 \text{R'OR} + 2 \text{H}_2 \\
2 \text{R'NH}_2 + \text{ROCH}_2\text{R} & \rightleftharpoons 1\text{-Ru} \rightarrow 2 \text{R'NHR} + 2 \text{H}_2
\end{align*}
\]

**Figure 1.8:** Equations (4) and (5) transesterification processes of esters with alcohols and amines catalyzed by 1-Ru

**F. Mechanistic studies for the potential energy surface for the dehydrogenative coupling of alcohols and amines into esters and carboxamides**

An understanding of the mechanisms can help researchers to improve the current reactions and design new reactions. Thus the mechanism of catalysis by 1-Ru has been studied theoretically. Studies focused on an important role for the ligand in hydrogenation, where the reaction occurs via metal-ligand cooperative mechanism that in turn includes reversible aromatization-dearomatization of ligands.[9]
1. Metal-Ligand Cooperation Mechanism (MLC)

Initially, the mechanism was proposed to follow a Metal-Ligand Cooperative mechanism that imposes 1-Ru and hemiacetals as intermediates for ester hydrogenation, and 1-Ru and hemiaminals for carboxamide hydrogenation has been known to be a generally accepted mechanism.\[27, 28\]

![Diagram](image)

**Figure 1.9:** Equations (6) and (7) showing the formation of the hemiacetals and hemiaminals by the reaction of esters and carboxamides with 1-Ru-H respectively.

The given MLC reactions are related to the bifunctional reactions invoked by Noyori to account for ketone hydrogenation by octahedral and piano stool ruthenium-amino hydride catalysts (HRu-NH) where the amino group serves as the proton donor. Noyori proposed bifunctional hydrogenation to take place in an outer-sphere mode without prior coordination of the ketone to the metal.\[29, 30\]
Figure 1.10: Outline of Noyori’s bifunctional reaction mechanisms

Although 1-Ru is known to add H₂ via metal-ligand cooperation mechanism to form the octahedral 1-Ru-H, yet the exact mechanism of catalysis of 1-Ru has not been well understood (equation 1). Metal-ligand cooperation mechanism (MLC) is not restricted to the H₂ addition to 1-Ru; in fact, it has been reported in the chemical literature that numerous metal complexes, including catalysts, are known to proceed in the reaction pathway via a metal-ligand cooperation mechanism, where a part of the ligand is also involved in the mechanistic flow of the reaction. Such mechanisms involve a limited change in the structure of the ligands for the sake of activating the substrate and in turn forming the desired product. [19, 31]

Milstein and coworkers proposed an MLC mechanism that could be the key mechanism in catalysis. [8] They have assumed that the hydrogenation of esters and carboxamides to proceed by an MLC mechanism where the proton of the phosphine arm is transferred to the carbonyl oxygen of the substrate and the hydride is transferred to the carbonyl carbon resulting in hemiacetals or hemiaminals formations as intermediates in addition, 1-Ru being restored (equations 6 and 7). [32, 33]. They elaborate on the
issue, stating that the pincer complex first undergoes deprotonation at the pyridinyl methylenic carbon which gives rise to the dearomatized pyridine entity. The formed complex can then activate various chemical bonds, including but not limited to H—H, H—OH, H—OR, H—NH₂, H—NR₂, and H—C bonds. This reaction occurs by a cooperation mechanism between the metal and the ligand; at this stage, aromatization at the pyridine ring is regained as the octahedral 1-Ru-H is regenerated. The whole process does not involve any change in the oxidation state of the Ruthenium metal center. [3, 8, 26, 28]

![Cooperative ligand site](image_url)

**Figure 1.11:** Top view of 1-Ru square pyramidal complex indicating the site of ligand cooperation (in red)

Milstein’s suggested mechanism relates to the bifunctional mechanism reported by Noyori. Noyori’s mechanism is known to proceed as a bifunctional hydrogenation in an outer sphere model without having the carbonyl functional group being coordinated to or in any interaction with the metal center. [29, 30]

To account for the complete hydrogenation of esters and carboxamides into alcohols and amines, the process requires a hydrogenolysis step of the C—OR bond or the C—NHR bond at some point in the reaction. Considering esters, and the hemiacetal intermediate would certainly fragment into an alcohol and an aldehyde even in a
complete absence of a catalyst (equation 8).[34] At a later stage, the aldehyde, in the presence of H₂ gas, would be hydrogenated by the present catalyst to give a second alcohol molecule. However, in the reaction of hydrogenation of carboxamides, hemiaminals are expected to undergo dehydrogenation reactions and transform into imines, the latter would be hydrogenated into amines depending on the reaction conditions and on the catalyst being used (equation 9). [10]

![Diagram of chemical reactions](image)

**Figure 1.12:** Equations (8) and (9) that demonstrate the dissociation of hemiacetals and hemiaminals

Depending on the catalyst and the reaction conditions, the imine can in turn be hydrogenated into a final amine product.[35-37] No imine or coupled amine products are observed in catalysis by 1-Ru. In a rather striking effect, however, when the PNP analogue of 1-Ru is used as the catalyst in amine coupling, imines are observed as the only products.[38] Significantly, although 1-Ru is a catalyst for alcohol coupling into esters, when it is used in the coupling of amines with alcohols, only carboxamides are observed as products.

The previously mentioned MLC mechanism has been taken into consideration by various computational studies. In a study by Wang and co., they investigated the
study of the amine coupling into carboxamides and suggest that the reaction would flow via metal ligand cooperation mechanism. Wang’s mechanism begins in two preparatory steps: first steps is described as a dehydrogenation of the alcohol by 1-Ru catalyst to yield an aldehyde followed by an MLC addition of the N-H bond to 1-Ru to produce the octahedral ruthenium amide complex (2-Ru-NHR). [39]

**Figure 1.13:** Outline of MLC mechanism for Amine-Alcohol coupling catalyzed by 1-Ru

After that, the formation of the carboxamide occurs in two stages, starting by a reaction between the aldehyde and Ruthenium-Amide (2-Ru-NHR) via formation of a six-member outer sphere ring transition state (TS_{HemAm}), this ring forms also via Metal ligand cooperation in which the Ru—N bond starts to break and the latter points out towards the carbonyl carbon, on the other hand, the pyridinyl methylenic proton is in the process of being abstracted by the Oxygen of the substrate. The products of this transition state are the separated hemiaminal and restoration of the 1-Ru catalyst. In the second stage of the reaction, 1-Ru and the produced hemiaminal react again in a two step reaction where at first a proton is transferred from the oxygen of the catalyst to the
phosphine arm of the catalyst, where the pyridine ring regains its aromaticity. The last step would involve a hydride transfer from the carbonyl carbon of the substrate to the
Ruthenium metal, giving off the final products: 1-Ru-H octahedral complex and a carboxamide.

2. **Bifunctional Double Hydrogen Transfer Mechanism (BDHT)**

Li and co performed another theoretical study for the same reaction pathway; they considered the formation of hemiacetals and hemiaminals as subsequent to the condensation of aldehydes that was produced from a dehydrogenative coupling of alcohols or of alcohol-amine systems without a need for a catalyst. After that, 1-Ru is thought to catalyze hemiaminals and hemiacetals dehydrogenation via sequential proton and hydride transfers that follow the metal ligand cooperation mechanism. In their studies, and if taken in the direction of esters and carboxamides, both Wang and Li would start by hydrogenation of the carbonyl into a hemiacetal or a hemiaminal; the divergence occurs in how hydrogenation takes place.[9, 39, 40]

For the alcohol dehydrogenation, Li and co. proposed a pathway that they referred to as Bifunctional Double Hydrogen Transfer (BDHT). Such mechanism goes as follows; the BDHT dehydrogenation mechanism is mediated by the bifunctional active site of 1-Ru that consists of a Lewis acidic Ru center and a Lewis basic sp² carbon center neighboring the phosphine arm of the pincer ligand. This step is very similar to the hydrogen transfer step that is thought to proceed by metal-ligand cooperation mechanism. [39, 40]
Figure 1.14: Potential Energy Surface of Alcohol dehydrogenation via bifunctional hydrogen transfer mechanism, values are given as ΔG [ΔH] and are reported in kcal/mol.

The BDHT pathway is shown to be a straightforward pathway in which the Ru metal center and the sp² carbon center serve as the hydride and the proton acceptors respectively. This double hydrogen transfer occurs via two steps, first a proton transfer from the oxygen to the carbon which proceeds via TS-1 (TS5 according to Li) transition state, this state shows the largest energy barrier, with a ΔG value of 24.8 Kcal/mol, rendering it the rate determining stage. This step is followed by a formation of an intermediate complex, Cplx-2 (12 according to Li), in which the Hydride is partially coordinated with the Ruthenium center and the aromatization of the pyridine ring is
gained as the hybridization of the carbon center changes from sp² to sp³ by gaining a proton and the electron pair is donated to the pyridine ring. The second step would be the hydride transfer from the carbonyl carbon to the Ruthenium metal. This is shown on the potential energy diagram as TS-2 (TS6 according to Li). TS-2 has a relatively lower free energy value than TS-1 (22.9 kcal/mol compared to 24.8 kcal/mol, with a ΔΔG value of -1.9 kcal/mol). TS-2 paves the way to form the octahedral dihydride Ruthenium complex 1-Ru-H and a reactive aldehyde.

As for the hemiacetal/hemiaminal formations, Li and co. present a coupling mechanism where the aldehyde that has been formed in the previous stage would couple either with an amine to give a hemiaminal or with an alcohol to give a hemiacetal, both of which can undergo dehydrogenation at a later step to produce an amide and an ester respectively. After that, the hemiacetals/hemiaminals would undergo dehydrogenations to give off the desired esters or carboxamides respectively. As reported in their study, Li and co describe the reaction as being flowing according to Bifunctional Double Hydrogen Transfer mechanism that was described earlier. In a similar way to alcohol dehydrogenation reactions, both the hemiacetal and the hemiaminal (labeled as 14heac and 13heam respectively) undergo dehydrogenation stepwise. Also, the energy barrier, shown in the figure as the transition state TS14 for the 13heam dehydrogenation was found to be 27.3 kcal/mol relative to 1-Ru (1cat in Li’s figure) and 13heam. This value is very similar to the energy barrier for the dehydrogenation of hemiacetal (TS16) which was also reported to be 27.3 kcal/mol relative to 1-Ru and 13heac. The flow of the reaction in both ways starts with a proton transfer from the oxygen of the substrate to the sp³ carbon of the phosphine arm. Again, aromatization is gained and this step is depicted in the diagram below as TS14 for hemiaminal dehydrogenation and TS 16 for
hemiacetal dehydrogenation. Both transition states would lead to octahedral complexes as intermediates, 30 and 31 in the figures, where the hydrogen of the carbonylic carbon is pointed towards the metal center. The step would be followed by a hydride transfer, shown as TS15 and TS17 in the diagram in which both would lead the octahedral Ruthenium dihydride complex 1-Ru-H (5dih according to Li’s diagram) and either carboxamide 4amd or ester 29est.

Figure 1.15: (A) Pathway for the dehydrogenation of the hemiacetals/hemiaminals intermediates via BDHT and (B) the corresponding energy profiles reported as ΔG [ΔH] in kcal/mol[41]
Li and co. have presented a thorough computational study on the role of Ru(II)-PNN pincer ligand in the hydrogenation of alcohols and amines into esters and carboxamides. Their presented bifunctional double hydrogen transfer mechanism is a similar approach to the earlier described Metal-ligand cooperation mechanism in which the ligand would directly interfere and aid the metal in catalysis.[9]

G. Proposed Direct Route Metathesis Mechanism

Herein, and using electronic structure DFT methods, we investigate alternative, more direct H/OR and H/NHR metathesis routes to the alcohol/amine coupling that circumvent the intermediacy of hemiacetals or hemiaminals. The newly proposed mechanism involves formation of hemiacetaloxide and hemiaminaloxide ion-pairs by addition of an aldehyde (from metal-catalyzed alcohol dehydrogenation) to an octahedral ruthenium-alkoxide or ruthenium-amide intermediate (from alcohol or amine addition to 1-Ru), followed by simple rearrangement (slippage) within the intact ion-pairs to transfer a hydride from the hemiacetaloxide or hemiaminaloxide to the metal. We show that the computed potential energy surfaces that are sometimes invoked to support the MLC mechanism correspond to indirect routes to metathesis. Both the ion-pair and the MLC routes predict the dehydrogenative coupling of ethanol and methanol into methyl acetate to be kinetically much more favored than the kinetics of formation of N-methylacetamide from ethanol and methylamine. However, the calculations provide evidence for the accessibility of a low energy NHR/OR metathesis path that would amidate the ester into the experimentally observed thermodynamically more favored carboxamide product.
H. Computational Methods and Solvation Models

This study was done on the dimethyl amino analogue of 1-Ru, but for convenience we use the same notation for both systems. All calculations were carried out using Gaussian 09. [42] Geometry minimization and normal mode vibrational analysis were done in the gas phase at the M06 level,[43] using the 6-31++G(d,p) basis set on the nonmetal elements, and the Hay–Wadt relativistic effective core potential (ECP) on ruthenium[44, 45], along with a double-ζ basis set augmented with one f polarization and one diffuse d function (exponents = 1.24 and 0.015, respectively) [46]. Selected transition states were also used to conduct intrinsic reaction coordinate analysis (IRC) [47]. Final electronic energies were obtained using the gas phase geometries at the M06, M06L, [48] andωB97X-D [49] levels of theory which are among the more popular methods used in computational organometallic chemistry. For this purpose the larger 6-311++G(2d,2p) basis set was used on the nonmetal elements, and an additional f polarization function with exponent = 0.4 was added to Ru. The reported standard state Gibbs free energies were obtained at 298 K and 1 atm using the entropies from the gas phase calculations after scaling them by a factor of 0.5. The reason for scaling the entropies in associative reactions is discussed in the studies by Wang where he based his scaling on experimental results. [38] The scaling factor influences the absolute free energies of the transition states of interest but does not change the main conclusions of the study.

Experimentally, much of the hydrogenation and coupling chemistry of 1-Ru had been done using toluene as solvent, though some experiments had been also carried out in THF, dioxane, DMSO, and anisole, or without a solvent. [11, 12] In the present study two approximation levels are applied to model the solvent. In one, bulk solvent
effects are included using the SMD model [50], with toluene, THF, or methanol as the solvent continuums. Because alcohols are either reactants or products in the reactions of interest, the second approximation level includes a methanol molecule explicitly in the calculations (along with the solvent continuum) to mimic possible specific H-bonding interactions.
CHAPTER II

DIRECT OUTER-SPHERE METATHESIS MODEL: DFT RESULTS

A. Computed Reaction Energies of 1-Ru

The reactions of the square pyramidal 1-Ru with H$_2$, methanol, and methylamine (eqs 10–12) are of central importance to the present study. In Table 1 we compare the free energies of the three reactions computed using the M06, M06L, and $\omega$B97X-D density functionals and in different solvent continuums. The values given in the equations are for the M06L-toluene results.

\[
\begin{align*}
1\text{-Ru}+H_2 & \rightarrow 2\text{-Ru-H} \\
\Delta H^\circ & = -11.4; \Delta S^\circ = -16.2 \text{ eu} \\
\Delta G^\circ & = -6.6 \text{ kcal/mol} & (10) \\
1\text{-Ru}+\text{MeOH} & \rightarrow 2\text{a-Ru-OMe} \\
\Delta H^\circ & = -11.8; \Delta S^\circ = -22.6 \text{ eu} \\
\Delta G^\circ & = -5.0 \text{ kcal/mol} & (11) \\
1\text{-Ru}+\text{MeNH}_2 & \rightarrow 2\text{a-Ru-NHMe} \\
\Delta H^\circ & = 0.0; \Delta S^\circ = -22.7 \text{ eu} \\
\Delta G^\circ & = +6.8 \text{ kcal/mol} & (12)
\end{align*}
\]

<table>
<thead>
<tr>
<th></th>
<th>H$_2$</th>
<th>MeOH</th>
<th>MeNH$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>M06</strong> (toluene)</td>
<td>-3.9</td>
<td>-1.4</td>
<td>+10.5</td>
</tr>
<tr>
<td><strong>M06L</strong> (toluene)</td>
<td>-6.6</td>
<td>-5.0</td>
<td>+6.8</td>
</tr>
<tr>
<td>$\omega$B97X-D (toluene)</td>
<td>-7.1</td>
<td>-3.5</td>
<td>+8.4</td>
</tr>
<tr>
<td><strong>M06L</strong> (THF)</td>
<td>-7.5</td>
<td>-3.0</td>
<td>+8.7</td>
</tr>
<tr>
<td><strong>M06L</strong> (methanol)</td>
<td>-7.7</td>
<td>-2.4</td>
<td>+10.8</td>
</tr>
</tbody>
</table>

Table 1. Computed $\Delta G^\circ$ of H$_2$, Methanol and Methylamine Addition to 1-Ru, Results from single-point calculations in a polarizable solvent continuum using geometries minimized in the gas phase at the M06 level.
The three methods predict H₂ addition to 1-Ru to be exoergic by 3.9–7.1 kcal/mol (ΔG° at 298.15K and 1.0 atm). The results are qualitatively consistent with the observation that 1-Ru adds H₂ readily to give an isolable ruthenium trans-dihydride product. Solutions of 1-Ru in benzene are also known to react with few equivalents of methanol to give an octahedral ruthenium alkoxide, but this product could not be isolated as a solid. The different methods in Table 1 predict methanol addition to 1-Ru to be approximately 2 kcal/mol less exoergic than H₂ addition. Finally all of the methods in Table 1 agree that the thermodynamics of methylamine addition to 1-Ru should be about 12 kcal/mol less favored than methanol addition.

The energies in Table 1 exhibit small dependencies on the choice of the solvent continuum in the calculations. For H₂ addition, changing the continuum from toluene to THF or methanol favors the reaction by ~1 kcal/mol. In contrast, THF and methanol disfavor methanol and methylamine addition by more than 2 kcal/mol.

For the three reactions in Table 1, the M06 energies are significantly less exoergic than either the M06L or the ωB97X-D ones. As mentioned above, the free energies in Table 1 are obtained after scaling the gas phase entropies by a factor of 0.5. This value is arbitrary and probably exaggerated. If the entropies were scaled by 0.7, ΔG° for methanol addition in toluene at the M06 level would be +1.2 kcal/mol, which would be inconsistent with the experiment. With a scaling factor of 0.7, the M06L and ωB97X-D energies remain exoergic by 2.3 and 0.8 kcal/mol, respectively. Studies by Truhlar[51] and more recently by Gusev[52] (using different basis sets than the ones employed here) showed the M06L functional to be systematically accurate in reproducing experimental enthalpies of organometallic reactions having different
characters. A recent theoretical investigation of agostic bonding in nickel complexes by Pudasani and Jenesko showed the M06L results to be in good agreement with those from the high level correlation consistent composite approach for transition metals.[53] Similarly, a comprehensive benchmark study by Remya and Suresh of noncovalent interactions between polar molecules found the M06L functional to be the “best performer” in a comprehensive pool of density functionals.[54] The latter systems are directly relevant to the type of potential energy surfaces elucidated in the present study. Accordingly, we chose to base the discussion in the present work on the M06L energies (with the 0.5 entropy scaling factor), and at the end we give a table that compares the relative energy of the key transition states at different levels and in different solvents. We note that we had completed geometry minimization at the M06 level before the given method-validation studies were published. We verified for select reactions that the same final energies are obtained if the geometries were minimized at the M06L or ωB97X-D levels.

B. Metathesis in the Hydrogenation Direction

Although the ultimate interest of the present study is the dehydrogenative coupling of alcohols and amines, we find it most useful to approach the problem from the hydrogenation direction starting with the octahedral 2-Ru-H. Because we are largely after supporting the plausibility of a new reaction mode for the C–OR or C–NHR bond-making or -breaking step in a complex multistage reaction system, we chose to work with the simplest ester and carboxamide, namely methyl acetate (3) and N-methylacetamide (4). Due to the asymmetry of the PNN ligand the two hydrides in 2-Ru-H are not equivalent. This allows definition of several stereoisomeric outer-sphere pathways in the reaction of 2-Ru-H with a carbonyl compound[55, 56]. We focus on the
pathway that aligns the carbonyl of 3 or 4 along the ruthenium–amine bond of 2-Ru-H (defined as path a) as illustrated in equation 13 for the precomplex between 2-Ru-H and 3.

On this path the carbonyl oxygen is on the opposite side of the axial methylene proton of the phosphine arm that is needed in metal–ligand cooperation, which we denote as path b and will be discussed later. The structural and energy data for the stationary points identified on path a are given in Figure 2.1.
Figure 2.1. Stationary points on the outer-sphere PES in the reaction of an ester (3) and a carboxamide (4) with 2-Ru-H. Energies are given in kcal/mol relative to the separated reactants (M06L-toluene SMD continuum).

The PES in Figure 2.1 starts with a transition state (3a-TS-H) for hydride transfer characterized by a Ru–H and C–H bond distance of 1.94 and 1.42 Å; respectively, and an imaginary frequency of 506i cm\(^{-1}\). 3a-TS-H leads to an ion-pair minimum between a hemiacetaloxide and a square pyramidal ruthenium cation in which the C–H bond is pointed to the metal at 2.04 Å (3a-IP-H). The C–H bond of the hemiacetaloxide anion in 3a-IP-H is long (1.27 Å), implicating some degree of activation due to metal coordination. The proposed H/OR metathesis pertains to rearrangement of the ion-pair to coordinate the OMe group of the hemiacetaloxide to the metal in place of the \(\alpha\)-hydrogen. The PES has a minimum for the latter species (3a-IP-OMe) characterized by a relatively long Ru–OME bond (2.42 Å), and, more
importantly, a significant stretch of the C–OMe bond to 1.60 Å. The computed stretch implies that coordination of the OMe group to the metal activates the C–OMe bond just as it does for the C–H bond in 3a-IP-H. From 3a-IP-OMe, the gas phase calculations identify a TS for C–OMe cleavage (3a-TS-OMe) that mirrors the hydride transfer TS. 3a-TS-OMe can be thought to “transfer” an alkoxide from the hemiacetaloxyde to the metal to give the octahedral 2a-Ru-OMe and eliminate acetaldehyde.

The net reaction of 3 in Figure 2.1 is a metathesis exchanging a hydride and an alkoxide between the metal center of 2-Ru-H and the acyl group of the ester. The given reaction requires a “mechanism” to alternate the coordination of the C–H and C–OMe groups of the hemiacetaloxyde to the metal. In Figure 2.1 we consider a transition state (3a-TS-Slip-1) that achieves the rearrangement in a direct way within the intact ion-pair by pointing both the α-H and the OMe groups of the hemiacetaloxyde to the metal. The Ru–H (2.47 Å), C–H (1.13 Å), C–OMe (1.51 Å) and Ru–OMe (2.99 Å) parameters in 3a-TS-Slip-1 are all intermediate between the corresponding values in the C–H and C–OMe ion-pair. The 3-D molecular displays given in Figure 2.2 reveal a good structural fit of the hemiacetaloxyde in the catalyst “cavity” in each of the three TSs on the metathesis PES.

![Molecular displays of the three TSs on the metathesis PES of 3](image)

**Figure 2.2:** Molecular displays of the three TSs on the metathesis PES of 3
The computed enthalpy and free energy of the precomplex between the ester and 2-Ru-H in equation 13 are −3.5 and 3.9 kcal/mol, respectively (M06L, in toluene continuum). The barrier for hydride transfer relative to the separated reactants is 17.6 kcal/mol. This is a relatively low barrier, a result that can be attributed to two factors: (i) an electronic one following from the trans-configuration of the two hydrides in 2-Ru-H, [55, 56] and (ii) a structural factor following from the presence of an opening over the amino group which clearly allows the ester to approach the hydride without much steric demand (Figure 2.2). The C–H bound ion-pair minimum has an energy nearly identical to that of 3a-TS-H, and from this point the barrier to rearrangement via 3a-TS-Slip-1 is only 3.8 kcal/mol. The OMe bound ion-pair is 4.5 kcal/mol more stable than 3a-IP-H, which seems consistent with the better coordinating ability of an OMe group compared to a C–H bond. From 3a-IP-OMe the gas phase M06 electronic barrier to C–OMe cleavage via 3a-TS-OMe is just 0.5 kcal/mol. IRC calculations in the gas phase starting from 3a-TS-OMe confirm the connectivity (Figure 2.3) and supports the near absence of a barrier. The barrier for C–OMe dissociation becomes slightly negative when the thermal enthalpic terms and solvent effects are included via single point M06L calculations.
Figure 2.3: Intrinsic Reaction Coordinates plots starting from 3-TS-H and 3-TS-OMe
In fact, for many of the other stereoisomeric paths the calculations failed to identify a TS for C–OMe cleavage due to the flatness of the PES of the C–OMe cleavage step. We recommend that a relaxed scan of the PES as a function of the C–OR bond distance is used before seeking to identify TSs related to 3a-TS-OMe. Complete aldehyde elimination from 3a-IP-OMe is exoergic by 6.1 kcal/mol. The similar energy of each C–H and OMe ion-pair and the corresponding C–H and C–OMe TS indicates that the given ion-pairs themselves can be viewed as activated species on the PES, meaning that once the hydrogen or the OMe group of the α-carbon of the hemiacetaloxide gets into the vicinity of the metal center, it will undergo a barrierless downhill C–H or C–OMe cleavage (corresponding, respectively, to hydride or alkoxide transfer to the metal). Under this condition, 3a-TS-Slip-1 is the highest energy point on the metathesis PES with \( G^e_{\text{rel}} = 21.8 \) kcal/mol relative to the separated reactants. This means the direct ion-pair mediated H/OR metathesis of esters can be a chemically relevant reaction even at ambient temperatures.

As found for the ester, the PES in the reaction between N-methylacetamide (4) and 2-Ru-H in Figure 2.1 also starts with a precomplex followed by a hydride transfer TS (4a-TS-H) leading to a C–H bound hemiaminaloxide ion-pair minimum (4a-IP-H). The barrier for hydride transfer to 4 is much larger than in the reaction of 3. As found for the ester, the energy of the TS and the C–H ion-pair minimum of 4 are close: 27.5 and 25.0 kcal/mol, respectively.

A “stepwise” ion-pair mediated metathesis starting with the carboxamide requires coordination of the amino nitrogen of the hemiaminaloxide to the metal. A minimum for the latter species (4a-IP-NHMe; Figure 2.1) is computed to have \( G^e_{\text{rel}} = 19.4 \) kcal/mol, a full 5.6 kcal/mol lower than the preceding C–H bound ion-pair, which
is reasonable, given that the amino group is a classical two-electron donor ligand. In
spite of the favorable thermodynamics, however, the barrier for the direct slippage step
from 4a-IP-H to 4a-IP-NHMe via 4a-TS-Slip-1 is 5.2 kcal/mol, slightly larger than the
matching barrier for hemiacetaloxide slippage (3.8 kcal/mol). The TS structures suggest
the difference could follow from slightly greater steric demands in 4a-TS-Slip-1 due to
the additional hydrogen of the NHMe group. With an increased energy input in both the
hydride transfer and the slippage components of the H/NHMe metathesis, $G^\ast_{\text{rel}}$ of 4a-
TS-Slip-1 comes to 30.2 kcal/mol. This is a quite substantial barrier, but it is still not
prohibitive for the reaction to be chemically relevant under the conditions used in the
hydrogenation of carboxamides, which are typically done at temperatures greater than
100 °C and reaction times in the hours.

Upon slippage from 4a-IP-H to 4a-IP-NHMe the C–H bond of the
hemiaminaloxide contracts from 1.23 to 1.13 Å, whereas the C–NHMe bond lengthens
from 1.48 to 1.56 Å. This suggests that coordination of the amino group to the metal
weakens the C–NHMe bond, but not to the same extent computed for C–OR activation.
However, full C–NHMe bond cleavage starting from 4a-IP-NHMe into the separated
acetaldehyde and the octahedral 2-Ru-NHMe products is calculated to be highly
endothermic ($\Delta H^\circ_{\text{Diss}} = 9.5$ kcal/mol). Thus, even with the favorable dissociation
entropy term (scaled by 0.5) acetaldehyde elimination from 4a-IP-NHMe is uphill by
2.2 kcal/mol. This behavior is fundamentally different from C–OMe cleavage, where
the corresponding $\Delta H^\circ_{\text{Diss}}$ and $\Delta G^\circ_{\text{Diss}}$ starting from 3a-IP-OMe were 1.0 and −6.1
kcal/mol, respectively. The different thermodynamics of the C–OMe and C–NHMe
cleavage reactions raises questions on whether the kinetics of the C–NHMe will
encounter an increased barrier or not. As shown in Figure 2.4, C–NHMe cleavage leads
at first to an adduct between acetaldehyde and the octahedral 2a-Ru-NHMe characterized by an N–C bond distance of 2.91 Å. The transformation from ion-pair 2a-Ru-NHMe to the adduct is uphill by 3.4 kcal/mol. Our extensive attempts to identify a TS for C–NHMe cleavage from 4a-IP-NHMe to the adduct have not been successful. A relaxed PES scan as a function of the C–NHMe bond distance supports the absence of any significant barrier to dissociation beyond the uphill thermodynamics. Thus, in Figure 2.1 we use the energy of the aldehyde adduct \( G^\circ_{\text{rel}} = 22.8 \text{ kcal/mol} \) as a rough estimate for the energy of the missing 4a-TS-NHMe. On the basis of the given results we can conclude that the slippage TS is the highest energy point on the direct H/NHMe metathesis PES in Figure 2.1.

**Figure 2.4:** Comparison of C–NHMe bond cleavage of the amine-bound hemiaminaloxide ion-pair on paths a and b. Energies are given in kcal/mol, and distances in Å.
Further investigation of the question of the barrier of the metal-mediated C–NHMe cleavage starting from a higher energy ion-pair on path b (4b-IP-NHMe; $G^\ddagger_{\text{rel}} = 23.3$ kcal/mol) identified a TS for C–NHMe cleavage characterized by a C–NHMe bond distance of 2.21 Å and an imaginary frequency of 138i cm$^{-1}$ (4b-TS-NHMe; Figure 2.4). As found on path a, C–NHMe cleavage on path b leads to an aldehyde adduct of 2b-Ru-NHMe as a true minimum on the PES but with a slightly higher energy than the adduct on path a ($G^\ddagger_{\text{rel}} = 24.1$ vs 22.8 kcal/mol). Overall, the given data indicate that formation and cleavage of the amine-bound hemiaminaloxide ion-pair on path b should be much less favored than that on path a. Nevertheless, $G^\ddagger_{\text{rel}}$ of 4b-TS-NHMe comes to 26.4 kcal/mol relative to the separated reactants, which is still significantly lower than that of 4a-TS-Slip-1 on path a (30.2 kcal/mol; M06L). We note however that at the ωB97X-D level the energies of the slippage (path a) and C–N cleavage (path b) TSs become equal ($G^\ddagger_{\text{rel}} = 29.6$ kcal/mol). Note that on path b the carbonyl oxygen gets close to the axial proton of the phosphine arm (1.76 Å in 4b-IP-NHMe and 1.96 Å 4b-TS-NHMe), but there is no indication of any active role of the ligand in assisting C–NHMe cleavage. In addition, the significantly higher energy of 4b-IP-NHMe compared to that of 4a-IP-NHMe (3.9 kcal/mol) suggests an absence of a stabilizing bonding CH–carbonyl interaction.

C. Intrinsic Reaction Coordinates

To further characterize the PESs elucidated above we conducted IRC analyses starting from 3a-TS-Slip-1 and 4a-TS-Slip-1. The calculations were carried out in the gas phase at the same (M06) level used in geometry minimization. The M06 electronic energy along the two IRCs relative to the TSs is plotted in Figure 2.5. The figure includes 3-D molecular displays of the C–H and C–OMe ion-pairs extracted from the
actual IRC outputs. The results confirm that TS-Slip-1 mediates a direct connection between C–H and C–OMe or C–NHMe “rotamer” ion-pairs. The structural parameters along the two IRCs starting from the C–H ion-pair side reveals smooth contraction of the C–H bond along with a smooth elongation of the C–OMe or C–NHMe bond as TS-Slip-1 is reached and crossed. Finally, the plots reveal a quite flat region of the PES around each of the slippage TSs, which is consistent with the small magnitude of the imaginary frequency in 3a-TS-Slip-1 and 4a-TS-Slip-1: 106i and 125i cm⁻¹, respectively.

Figure 2.5. IRCs originating from 3a-TS-Slip-1 (green curve) and 4a-TS-Slip-1 (blue curve). M06 gas phase electronic energies are defined relative to the TSs in kcal/mol.

D. Energy Profiles for full Hydrogenation

Figure 2.6 compares the Gibbs free energy profiles for complete hydrogenation of 3 and 4. For convenience, the thermodynamics data for the individual hydrogenation and hydrogenolysis steps are collected in figure 2.7.
Figure 2.6. Gibbs free energy profiles (298 K and 1 atm) for the hydrogenation of 3 and 4 using 2-Ru-H (M06L-toluene; in kcal/mol).

\[
\begin{align*}
\text{MeCHO} + \text{H}_2 & \rightleftharpoons \text{EtOH} & \Delta H^0 = -15.4 & \Delta G^0 = -11.0 \\
\text{2a-Ru-OMe} + \text{H}_2 & \rightleftharpoons \text{2-Ru-H} + \text{MeOH} & \Delta H^0 = 0.3 & \Delta G^0 = -1.6 \\
\text{2a-Ru-NHMe} + \text{H}_2 & \rightleftharpoons \text{2-Ru-H} + \text{MeNH}_2 & \Delta H^0 = -11.4 & \Delta G^0 = -13.3 \\
\text{H}_2 + \text{2} & \rightarrow \text{EtOH} + \text{MeOH} & \Delta H^0 = -8.1 & \Delta G^0 = -5.2 \\
\text{NH} + \text{2} & \rightarrow \text{EtOH} + \text{MeNH}_2 & \Delta H^0 = -5.9 & \Delta G^0 = -2.8
\end{align*}
\]

Figure 2.7. Computed Thermodynamics Data Used in Figure 2.6 (M06L-Toluene; in kcal/mol)
The metathesis in the direction of acetaldehyde formation from 3 and 4 is endoergic by 7.4 and 21.6 kcal/mol, respectively. However, hydrogenation of acetaldehyde is exoergic by 11.0 kcal/mol (eq 15 in figure 2.7), so it provides a major component in driving the hydrogenation thermodynamics of both the ester and the carboxamide.

For complete hydrogenation and catalysis the octahedral 2a-Ru-OMe and 2a-Ru-NHMe products must undergo hydrogenolysis to produce an alcohol or an amine and regenerate 2-Ru-H. These reactions are exoergic by 1.6 and 13.3 kcal/mol, respectively (Table 1 and equations 16 and 17 in Figure 2.7). With these computed energies, the net hydrogenation of methyl acetate and N-methylacetamide comes to be exoergic by 5.2 and 2.8 kcal/mol, respectively (equations 18 and 19 in Figure 2.7).

Mechanistically, hydrogenolysis of the Ru–OR and Ru–NHR bonds can proceed in several routes including initial MLC elimination of the alcohol or the amine followed by H₂ addition to the square pyramidal 1-Ru, direct hydrogenolysis with H₂,[57] and solvent-assisted mechanisms.[58-62] Given the highly favorable thermodynamics and the accessibility of many reaction routes, hydrogenolysis of the Ru–OMe and Ru–NHMe bonds is not expected to be kinetically demanding. Similarly, in the presence of 2-Ru-H (which should be abundant under high H₂ pressure) hydrogenation of the aldehyde is expected to be fast. For simplicity we keep the barriers for aldehyde hydrogenation out of the PESs in Figure 2.6. The important point for our purposes is that Figure 2.6 gives unambiguous evidence that the metathesis route to hydrogenation should be kinetically and thermodynamically much more favored for 3 than for 4. This may sound trivial since it is well-known that carboxamides are
generally much harder to hydrogenate than esters.[63] However, accepting the large
differences in the hydrogenation PESs becomes critical in the coming discussion of the
origin of the observed selectivity to carboxamides over esters in the dehydrogenative
coupling of alcohols and amines.

E. Metathesis via carbonyl insertion and deinsertion

The slippage TSs in Figure 2.1 alternate the coordination of the C–H bond of a
hemiacetaloxide or hemiaminaloxide with an alkoxy or amino group. In Figure 2.8 we
consider another ion-pair rearrangement mode via TS-Slip-2 to coordinate the terminal
C–O⁻ bond of the hemiacetaloxide or hemiaminaloxide to the metal.

![Figure 2.8](image)

**Figure 2.8.** Three ion-pair-mediated reactions. The energies are given in kcal/mol
relative to the separated 2-Ru-H and 3 or 4.

The new slippage mode leads to octahedral Ru–hemiacetaloxide and Ru–
hemiaminaloxide complexes, 2a-Ru-HemAc and 2a-Ru-HemAm, respectively, having
Ru–OR bond distances of 2.24 Å, close to the value of 2.21 Å for the Ru–OMe bond in
2a-Ru-OMe (Figure 2.1). A molecular display of 2a-Ru-HemAc is given in equation 20
(without the tert-butyl groups).
The reactions mediated by TS-Slip-2 correspond to insertion of a carbonyl group into a Ru–H bond which has ample experimental precedence. Of particular relevance to the present study are the experiments by Bergens, where cyclic esters were observed to undergo rapid insertion in trans-[Ru(H)$_2$(Binap)(diamine)] complexes even at low temperatures.[64] The more common examples of carbonyl insertion in octahedral complexes involve ketones [65-73] and CO$_2$.[74-80] The reverse of the insertion step in Figure 2.8 is an unconventional β-hydride elimination from an alkoxide taking place via ion-pair rearrangement without the need of a vacant coordination site, which also has experimental precedence.[81-83]

For both 3 and 4 the insertion barrier via TS-Slip-2 is slightly larger than the one for direct metathesis via TS-Slip-1. The thermodynamics of insertion on the other hand are more favored than metathesis, offering $G^\circ_{rel} = 5.4$ or 11.4 kcal/mol for the reaction of 3 and 4, respectively. These results imply that insertion can in practice compete with direct metathesis, but it does not provide a thermodynamics sink. However, from the insertion products it is possible to define a third ion-pair rearrangement mode via TS-Slip-3 that exchanges the coordination of the anionic C–O$^-$ terminal of the alkoxide with the alkoxy or the amine group. The 3-D display in eq 20 shows it should not take many structural rearrangements to reach TS-Slip-3. The latter
slippage TSs mediate aldehyde deinsertion (or elimination) from 2a-Ru-HemAc and 2a-Ru-HemAm. The combined insertion and deinsertion steps provide therefore an indirect ion-pair mediated route to metathesis. $G^\ddagger_{\text{rel}}$ of 3a-TS-Slip-3 and 4a-TS-Slip-3 is 16.4 and 27.5 kcal/mol, respectively, significantly smaller than the other slippage TSs. Thus, even if the kinetics were to favor insertion via TS-Slip-2 over direct metathesis via TS-Slip-1, TS-Slip-3 will still provide an energetically accessible indirect ion-pair-mediated route to metathesis.

IRC calculations starting from 3a-TS-Slip-2 confirm a smooth connectivity between 3a-IP-H and 2-Ru-HemAc. IRC calculations originating in 3a-TS-Slip-3 on the other hand lead to the insertion product 2a-Ru-HemAc in one direction, but it converges after two steps to the same geometry as that of 3a-TS-Slip-3. This behavior may follow from the flat nature of the PES near TS-Slip-3 which is evident from the small magnitude of the imaginary frequency in 3a-TS-Slip-3 and 4a-TS-Slip-3, 98i and 115i cm$^{-1}$, respectively. Animation of these vibrations leaves no doubt they are for a rotation that exchanges the coordination of the terminal O and either the OMe or NHMe group. We have located several stereoisomeric transition states of 3a-TS-Slip-3 and 4a-TS-Slip-3, but we could not identify any minima in the close vicinity of any of them. As shown in equation 20, TS-Slip-3 requires minimal structural arrangement of the insertion intermediate, and thus, the IRC result of a minimum near TS-Slip-3 is most likely a computational artifact.

Figure 2.9 illustrates stereospecific details pertaining to how the hemiacetaloxide or hemiaminaloxide is produced and transformed on the insertion route to metathesis.
Figure 2.9: Stereospecific Details of the Insertion–Deinsertion Path to Metathesis

The carbonyl of 3 or 4 approaches 2-Ru-H from the pyridine side with the alkyl group pointing toward the phosphine group. In TS-Slip-2 the exchanging C–H and C–O⁻ bonds are aligned along the pyridine–Ru–CO axis, so the C–H bond eclipses the Ru–CO bond. As the Ru–alkoxide bond is formed, the three substituents on the tetrahedral carbon of the alkoxide are rotated so as to stagger the Ru–CO bond in between the C–H and C–X bond (X = OMe or NHMe). Continued rotation eclipses the C–X and Ru–CO bonds, providing thereby the correct alignment needed in TS-Slip-3. In the process, the orientation of the alkyl group is flipped toward the amine, and the aldehyde eliminates from the same side used in the insertion direction. Note that the alkoxide oxygen in 2a-Ru-HemAc and 2a-Ru-HemAm is tilted over the pyridyl ring, thus making an O–Ru–N angle of near 80° (eq 20). This structural feature appears to be common in octahedral Ru-alkoxide as was discussed elsewhere.[84]
F. Effect of Explicit Solvent Molecule

Complete hydrogenation of 3 and 4 produces methanol. To investigate possible effects of H-bonding of the alcohol on the metathesis PESs, we conducted calculations that included a single methanol molecule to mimic an H-bond solvent donor. Starting with the separated reactants, H-bond formation between methanol and 2-Ru-H is calculated to be slightly exoergic (ΔG° = −1.4 kcal/mol) so we take this species as the reference point in defining the energies of the other points on the PESs. For each ion-pair the methanol was introduced such that it undergoes an H-bond with the formally “anionic” oxygen center. The results summarized in Figure 2.10 show the methanol molecule stabilizes all of the ion-pair TSs in the reaction of both 3 and 4 by 3–5 kcal/mol. The effect is not unexpected, since the given reactions replace an H-bond between methanol and a metal-hydride in the reactants by a stronger H-bond involving a negatively charged oxygen in the ion-pair minima and TSs.

**Figure 2.10.** Energies of the slippage TSs computed while including a methanol molecule (S1) to mimic an explicit H-bond donor solvent. Free energies are given in kcal/mol relative to the methanol-solvated 2-Ru-H and either 3 or 4.

G. Indirect metathesis via MLC

In this section we examine more closely the metal–ligand cooperative (MLC) route to hydrogenation. For this purpose Figure 2.11 tracks an outer-sphere PES that
allows interaction between the carbonyl oxygen of \( \text{3} \) or \( \text{4} \) with the axial proton of the phosphine arm of \( \text{2-Ru-H} \) (distinguished as path \( \text{b} \); see eq 13).

**Figure 2.11.** Reaction between \( \text{2-Ru-H} \) and \( \text{3} \) or \( \text{4} \) on path \( \text{b} \) involving ligand deprotonation. The energies are given relative to the separated reactants in kcal/mol.

As found in Figure 2.1, hydride transfer from \( \text{2-Ru-H} \) to methyl acetate on path \( \text{b} \) yields a C–H ion-pair minimum (3b-IP-H), but this time the terminal oxygen of the hemiacetaloxide is pointed at the axial C–H proton of the phosphine arm at a distance of 1.94 Å. From 3b-IP-H, proton transfer from the PNN ligand to the terminal oxygen of the hemiacetaloxide via 3b-TS-Prt-1 has a barrier of 5.5 kcal/mol. The immediate product from this step is 1b-Ru-HemAc-1, a loose precomplex between a hemiacetal and the neutral square pyramidal 1-Ru in which the C–H bond of the hemiacetal is pointed to the metal at a distance of 2.5 Å and the alcoholic proton is at 2.2 Å from the
deprotonated carbon of the phosphine arm. $G^\text{rel}$ of 3b-TS-Prt-1 is 23.8 kcal/mol (in toluene), slightly higher than the energy of 3a-TS-Slip-1 in Figure 2.1 (21.8 kcal/mol). Thus, in spite of the very different character of the two reactions, the slippage and ligand deprotonation TSs appear to be competitive. However, for hydrogenation to proceed, the hemiacetal produced after proton transfer would still need to undergo further hydrogenolysis of the C–OR bond. In investigating a possible role of 1-Ru in the latter reaction we considered the MLC mechanism reported in the study by Wang.[39] This led us to identify 3b-TS-Prt-2 in Figure 2.11 having $G^\text{rel}$ = 20.0 kcal/mol. We find the geometry (near linear C–H–O angle), imaginary frequency (1600i cm$^{-1}$), reduced mass (1.1 amu), and the animation of the imaginary frequency of 3b-TS-Prt-2 to be all fully characteristic of a localized motion of the alcoholic proton between an OMe-coordinated hemiacetal precomplex and the carbon of the phosphine arm. This differs from the delocalized description of the same TS in which the C–OR bond is cleaved concomitantly with proton transfer.[39] Consistent with a localized proton transfer step, Figure 2.11 shows 3b-TS-Prt-2 to connect an OMe-coordinated conformer of a hemiacetal precomplex of 1-Ru (1b-Ru-HemAc-2) with an OMe-coordinated hemiacetaloxide ion-pair (3b-IP-OMe). This means that the combined MLC sequence of hemiacetal formation and subsequent reaction with 1-Ru simply switches the C–H coordination of the hemiacetaloxide produced right after hydride transfer into the more favorable mode in which the OMe group points to the metal. In other words the given deprotonation and protonation sequence of the ligand provides an indirect route to interchanging the C–H and C–OMe ion-pairs taking place on path b. Further cleavage of the C–OMe bond from 3b-IP-OMe will complete an MLC-mediated H/OR metathesis. The same is true for the sequence involving the reaction of the carboxamide
in Figure 2.11, where a hemiaminal is formed by ligand deprotonation via 4b-TS-Prt-1, rotated to bind the amine to the metal in 1b-Ru-HemAm-2, and then deprotonated via 4b-TS-Prt-2 to give the amine-coordinated hemiaminaloxide ion-pair (4b-IP-NHMe). The latter ion-pair still needs to undergo aldehyde elimination to complete a metathesis. As discussed in Figure 2.4, C–NHMe bond cleavage starting from 4b-IP-NHMe proceeds by a distinct TS (4b-TS-NHMe) with $G^\ddagger_{rel} = 26.4$ kcal/mol.

IRC calculations confirm the connectivities associated with a localized proton transfer originating from the C–H bound 3b-TS-Prt-1 and 4b-TS-Prt-1. Likewise, for the NHMe-bound deprotonation of the hemiaminal by 1-Ru, the IRC starting from 4b-TS-Prt-2 converges into the expected hemiaminal precomplex in one direction and the NHMe-bound ion-pair of the hemiaminaloxide in the other (blue line in Figure 2.12).

**Figure 2.12.** Intrinsic reaction coordinates originating from 3b-TS-Prt-2 (hemiacetal; green curve) and 4b-TS-Prt-2 (hemiaminal; blue curve). M06 gas phase electronic energies defined relative to the TSs in kcal/mol.
For the reaction of the OMe-bound hemiacetal on the other hand, the IRC originating from 3b-TS-Prt-2 converges to the OMe bound hemiacetal precomplex in one direction, but it reaches a plateau in the direction of the OMe bound ion-pair and then proceeds with elimination of the aldehyde (Figure 2.12). This result enforces the proposition that once the OMe group of the hemiacetaloxide anion gets in the vicinity of the metal there will be no barrier to C–OMe cleavage and Ru-alkoxide formation. In spite of the lack of a distinct TS for C–OMe cleavage, the IRC from 3b-TS-Prt-2 in Figure 2.12 is still most consistent with a dehydrogenation mechanism of the hemiacetaloxide in which the proton is transferred separately from C–OMe cleavage. We emphasize that the calculations do identify a true minimum for the OMe-bound hemiacetaloxide ion-pair on path b (3b-IP-OMe in Figure 2.11). This minimum is very shallow, and we argue it is best viewed as an activated species on the PES that undergoes a barrierless C–OR bond cleavage. To some extent therefore, the 3a-TS-Slip-1 is effectively like a concerted metathesis TS.

H. Energy of the Slippage and MLC TSs

Table 2 compares the energies of the slippage and the C–H-bound proton transfer TSs in the reactions of 3 and 4 using different density functionals and solvent continuums. For the ester, the M06L $G^\dagger_{\text{rel}}$ of 3b-TS-Prt-1 in toluene (23.8 kcal/mol) is slightly higher than 3a-TS-Slip-1 (21.8 kcal/mol). For N-methylacetamide on the other hand, the M06L $G^\dagger_{\text{rel}}$ of 4b-TS-Prt-1 is 28.3 kcal/mol, 1.9 kcal/mol lower than $G^\dagger_{\text{rel}}$ of 4a-TS-Slip-1 (30.2 kcal/mol; in toluene continuum). The oB97X-D or the M06L levels selectively stabilizes the proton transfer TSs (to 24.2 kcal/mol), thus increasing the preference for 4b-TS-Prt-1 over 4a-TS-Slip-1 to 4.6 kcal/mol (in toluene; Table 2). The
results are not changed when full geometry minimization and normal-mode analysis are done at the ωB97X-D level.

<table>
<thead>
<tr>
<th></th>
<th>ester reactions</th>
<th>carboxamide reactions</th>
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<tbody>
<tr>
<td></td>
<td>3a-TS-Slip-1</td>
<td>3b-TS-Prt-1</td>
</tr>
<tr>
<td>M06 (toluene)‡</td>
<td>16.9</td>
<td>21.8</td>
</tr>
<tr>
<td>M06L (toluene)‡</td>
<td>21.8</td>
<td>23.8</td>
</tr>
<tr>
<td>ωB97X-D (toluene)‡</td>
<td>20.9</td>
<td>20.4</td>
</tr>
<tr>
<td>M06L (THF)‡</td>
<td>23.6</td>
<td>26.3</td>
</tr>
<tr>
<td>M06L (methanol)‡</td>
<td>14.8</td>
<td>24.2</td>
</tr>
<tr>
<td>Opt M06-L (Tol)§</td>
<td>21.6</td>
<td>23.3</td>
</tr>
<tr>
<td>Opt ωB97X-D (Tol)§</td>
<td>20.8</td>
<td>21.3</td>
</tr>
<tr>
<td>(SI) M06L (Tol)§</td>
<td>17.7</td>
<td>19.5</td>
</tr>
</tbody>
</table>

**Table 2.** \(G\)\(_{\text{rel}}\) of TS-Slip-1 and TS-Prt-1 at Different Theoretical Levels (in kcal/mol)

a Results from single-point calculations on geometries minimized in the gas phase at the M06 level.
b Geometries and frequencies computed in the gas phase at the M06L level.
c Geometries and frequencies computed in the gas phase at the ωB97X-D level.

The computed reversal in the energy order of the direct slippage and ligand deprotonation TSs in the reaction of the carboxamide compared to the reaction of the ester follows in part from a significantly larger proton affinity of the hemi-αminal oxide anion compared to the hemiacetal oxide (\(\Delta\Delta\text{G}^\circ = -6.4\) kcal/mol; eq 21; favoring proton transfer in the reaction of 4), and in part from a slightly larger barrier for rearrangement from IP-H to TS-Slip-1 in the carboxamide case (\(\Delta\Delta\text{G}^\circ = 1.4\) kcal/mol; Figure 2.1; disfavoring slippage in the reaction of 4).

\[
\begin{array}{c}
\text{H}^+ \quad \text{OH} \quad \text{OMe} \\
\text{1.40} & \text{1.53} & \qquad \text{NHMe} \\
\text{H} & \text{O} & \text{H}
\end{array}
\begin{array}{c}
\text{H}^+ \quad \text{OH} \quad \text{OMe} \\
\text{1.44} & \text{1.53} & \qquad \text{NHMe} \\
\text{H} & \text{O} & \text{H}
\end{array}
\Rightarrow
\begin{array}{c}
\text{H}^+ \quad \text{OH} \quad \text{OMe} \\
\text{1.40} & \text{1.53} & \qquad \text{NHMe} \\
\text{H} & \text{O} & \text{H}
\end{array}
\begin{array}{c}
\text{H}^+ \quad \text{OH} \quad \text{OMe} \\
\text{1.44} & \text{1.53} & \qquad \text{NHMe} \\
\text{H} & \text{O} & \text{H}
\end{array}
\]

\(\Delta\text{H}^\circ = -6.3\)

\(\Delta\text{G}^\circ = -6.4\) (21)

Given the ion-pair and acid–base nature of the given transformations, the precise contribution of the two paths to metathesis is expected to depend on dynamics.
and solvent effects details.[85] Such effects can be different for the two routes, but they are difficult to compute accurately. Thus, when a THF solvent continuum is used in the calculations, the M06L energies of 4a-TS-Slip-1 and 4a-TS-Prt-1 become nearly identical (Table 2), and in a methanol continuum the slippage TS becomes lower than the proton transfer TS by 2.7 kcal/mol. Furthermore, when a methanol molecule is included as an explicit H-bond donor in the calculations, the M06L 4a-TS-Slip-1 becomes 0.4 kcal/mol lower than 4b-TS-Prt-1 even in the toluene continuum (SI entries in Figures 2.10 and 2.11). We stress that the two TSs being compared are on two PESs leading to the same OR- or NHR-bound hemiacetal oxide and hemiaminal oxide ion-pairs, which we propose to activate the C–OR and C–NHR bonds in preparation for aldehyde elimination to complete an H/OR or H/NHR metathesis. In fact at the oB97X-D level the barrier to C–NHMe cleavage on the MLC path via 4b-TS-NHMe discussed in Figure 2.4 is 29.6 kcal/mol, significantly higher than that via 4b-TS-Prt-1 (24.2 kcal/mol) and identical to that via 4a-TS-Slip-1. The OR- and NHR-bound ion-pairs were not identified in previous computational studies of the MLC mechanisms. Consideration of these ion-pairs gives a fundamentally new perspective to understanding the metal-catalyzed C–OR and C–NHR bond cleavage step in the reaction of 2-Ru-H with 3 and 4 regardless of how the IPs are formed. In addition to the slippage and MLC rearrangement modes, one can envision other routes to obtain the OR and NHR ion-pairs, such as by full dissociation and reassociation of the ion-pairs, formation of solvent separated ion-pairs, or by a protonation and deprotonation sequence of the anion by the alcohol that is produced during hydrogenation. Again, the contribution of such routes will depend on solvent and dynamics effects that may vary as the concentration of the alcohol is changed during the course of catalytic
hydrogenation. The reaction via TS-Slip-1 provides sort of the least action path to rearrange the ion-pairs. The data in Table 2 shows this path to be at least competitive with the MLC mechanism involving ligand deprotonation and reprotonation.

I. Metathesis in the Dehydrogenative Coupling Direction

In this section we discuss the ion-pair PESs in the context of amine dehydrogenative coupling. To be able to address the question of the observed selectivity to carboxamides using the same data for methyl acetate and N-methylacetaamide hydrogenation, we start with 1-Ru in a mixture of ethanol, methanol, and methylamine. The computed free energy of ethanol addition to 1-Ru is −6.9 kcal/mol, compared to −5.0 and +6.8 kcal/mol for methanol and methylamine addition, respectively. According to these results the resting state of the catalyst in the given mixture should be the octahedral ruthenium–ethanoxide (2a-Ru-OEt), so we place this complex along with methanol and methylamine at the zero level on the Gibbs free energy scale in Figure 2.13).
Figure 2.13. Standard state Gibbs free energy profiles (298 K; 1 atm) for dehydrogenative coupling starting with 1-Ru in a mixture of methanol, ethanol and methylamine (M06L-toluene; in kcal/mol)

To produce methyl acetate from the given mixture, the mechanism under consideration requires a preparatory stage in which 2a-Ru-OEt is dehydrogenated into acetaldehyde and H₂ along with the formation of 2-Ru-OMe. The thermodynamics of this stage is uphill by 12.9 kcal/mol. Again, for our purposes we can ignore the barriers in the preparatory stage. Coupling between the aldehyde and 2a-Ru-OMe to give the C–OMe-bound hemiacetaloxide ion-pair is exothermic by 1.1 kcal/mol, but because of the unfavorable entropy term, the step is uphill by 6.0 kcal/mol (Figure 2.1), so $G^\circ_{\text{rel}}$ of 3a-IP-OMe in Figure 2.13 is 19.5 kcal/mol. To eliminate an ester, the hemiacetaloxide must rearrange to orient the C–H bond toward the metal. The rearrangement barrier
from the shallow 3a-IP-OMe via 3a-TS-Slip-1 is 8.3 kcal/mol. 3a-TS-Slip-1 is the highest energy point on the given PES, with a $G^\ddagger_{\text{rel}}$ value of 27.2 kcal/mol. The thermodynamics for transformation from the initial reactants into 2-Ru-H and the ester is uphill by 5.5 kcal/mol.

To form N-methylacetamide from the initial mixture in Figure 2.13, the preparatory stage requires dehydrogenation of 2a-Ru-OEt as well as reaction of the amine to give 2a-Ru-NHMe. Because the thermodynamics of amine addition to 1-Ru is unfavorable (Table 1), the energy input for this stage is large: 24.7 kcal/mol. Coupling of aldehyde with 2a-Ru-NHMe into 4a-IP-NHMe encounters a small barrier estimated at 1.2 kcal/mol in Figure 2.1, but opposite to the reaction of 2a-Ru-OMe, the coupling involving C–N bond formation is exothermic by 2.1 kcal/mol. However, the slippage barrier of the hemiaminaloxide needed to reach the C–H ion-pair is substantial (10.7 kcal/mol), so 4a-TS-Slip-1 comes to 33.3 kcal/mol on the PES. The combined results indicate strongly therefore that, at least for the given substrates, dehydrogenative coupling into an ester should be kinetically much more favored than coupling into a carboxamide (ΔΔ$G^\ddagger_{\text{rel}} = 6.1$ kcal/mol). When the respective MLC routes are considered, the highest energy points on the two coupling PESs will be 3b-TS-Prt-1 and 4b-TS-Prt-1 with $G^\ddagger_{\text{rel}} = 29.2$ and 31.4 kcal/mol, respectively, still in favor of ester formation. Note that 4a-IP-H in Figure 2.13 is 4.6 kcal/mol above 3a-IP-H. These C–H ion-pairs are effectively the TSs for the final hydride transfer steps needed to eliminate the carboxamide or the ester. As such, it is unlikely there can be other metal-catalyzed mechanisms to coupling involving aldehydes that would selectively favor the population of the higher energy 4a-IP-H to give a carboxamide as a kinetic product.
The computed kinetic preference to the ester in Figure 2.13 is opposite to the experimentally observed selectivity to carboxamides. However, the carboxamide is the thermodynamically favored product in the Figure ($\Delta G^\circ_{\text{rel}} = -2.4 \text{ kcal/mol}$). It is possible, therefore, that the ester can in practice be formed as a kinetic product, but it gets amidated under the experimental conditions, a possibility that had been raised in the studies by Milstein. As mentioned in the Introduction, complexes related to 1-Ru are indeed known to catalyze ester amidation (eq 4), an interesting and valuable reaction in its own.[25, 26]

Ester amidation under the alcohol-amine coupling conditions can in principle proceed on the same PESs involving the aldehyde. This will require the ester and 2-Ru-H products to revert back to acetaldehyde and 2a-Ru-OMe so the acetaldehyde may react with 2a-Ru-NHMe to form the thermodynamically more favored carboxamide. In Figure 2.14 we give evidence for the accessibility of an alternative ion-pair NHR/OR metathesis route that achieves ester amidation in a more direct way.

![Figure 2.14](image_url)

**Figure 2.14.** Stationary points on the outer-sphere PES in the reaction of an ester (3) and 2-Ru-NHMe. Energies are given in kcal/mol relative to the separated reactants (M06L-toluene).
J. Ester Amidation via NHR/OR Metathesis

The energies in Figure 2.14 are defined relative to the separated square pyramidal 1-Ru, methylamine and methyl acetate, thus mimicking the initial conditions of an ester amidation experiment utilizing an alkyl amine. From this mixture the preparatory stage for an NHR/OR metathesis of the ester involves only methylamine addition to 1-Ru, so the required energy input is relatively small, 6.8 kcal/mol. Coupling between the ester and 2-Ru-NHMe gives an amine-coordinated ion-pair minimum of the \( \alpha,\alpha-\text{(NHMe)}(\text{OMe})\text{-ethanoxide anion (5a-IP-NHMe).} \) The thermodynamics of the coupling step is computed to be nearly thermoneutral \( (\Delta H_{\text{Add}} = 0.5 \text{ kcal/mol}) \), affording \( \Delta G_{\text{Add}}^0 = 8.5 \text{ kcal/mol} \), but the step has a relatively high barrier via 5a-TS-NHMe with \( \Delta H_{\text{Add}}^\ast \) and \( \Delta G_{\text{Add}}^\ast \) of 8.9 and 17.1 kcal/mol, respectively. 5a-TS-NHMe is characterized by a C–N bond distance of 2.06 Å and \( \nu^\prime = 289i \text{ cm}^{-1} \) for C–N stretching vibration. Subsequent slippage via 5a-TS-Slip-1 in Figure 2.14 rearranges the \( \alpha,\alpha-\text{(NHMe)}(\text{OMe})\text{-ethanoxide within the intact ion-pair to coordinate the methoxy group to the metal (5a-IP-OMe).} \) The switch from NHMe to OMe coordination causes a contraction in the C–NHMe bond from 1.55 to 1.48 Å, and a stretch in the C–OMe bond from 1.50 to 1.60 Å, demonstrating again that metal-coordination activates the respective bonds. From 5a-IP-OMe, cleavage of the C–OMe bond into 2a-Ru-OMe and N-methylacetamide is highly exoergic \( (\Delta G_{\text{Diss}}^0 = -29.7 \text{ kcal/mol}) \), and a PES scan reveals the reaction to be totally barrierless. The net transformation in Figure 2.14 is an ion-pair-mediated metathesis in which an amide and alkoxide are exchanged between a ruthenium amide and an ester. Unlike the metatheses involving H/OR, H/NHR (Figure 2.1) or OR/OR exchange,[55] the highest energy point on the NHMe/OMe metathesis PES in Figure 2.14 is for the TS of the ester–amide coupling step, and not the slippage
TS. The difference between the two TSs is however small (23.9 vs 22.3 kcal/mol).

Thus, ester amidation is highly exoergic, and the direct metal-mediated metathesis route provides a surprisingly low energy barrier for the transformation.

In studying ester amidation, we also considered the MLC route to an NHR/OR ion-pair rearrangement. The transition state for ligand deprotonation on path b via the NHMe-coordinated α,α-(NHMe)(OMe)-ethanoxide (5b-TS-Prt-1; eq 22) has $G_{\text{rel}}^\ddagger = 22.6$ kcal/mol, similar to 5a-TS-Slip-1 (22.3 kcal/mol; M06L-toluene). However, the TS for ligand reprotonation via the OMe-coordinated anion (5b-TS-Prt-2; eq 22) is 26.3 kcal/mol, indicating the MLC route is overall less favored than ion-pair slippage.

When considered in the context of amine-alcohol coupling discussed in Figure 2.13, ester amidation has to start from 2-Ru-H, the ester, and the free amine. The energy of the latter species on the scale in Figure 2.13 is 5.5 kcal/mol. The preparatory stage for ester amidation requires H$_2$ elimination and 2-Ru-NHMe formation, with an energy input of 13.4 kcal/mol. This puts 5a-TS-NHMe and 5a-TS-Slip-1 at 35.9 and 34.4 kcal/mol, slightly higher than 4a-TS-Slip-1 (33.3 kcal/mol) for ester amidation via the (reversible) aldehyde route. However, the energy scale in Figure 2.13 applies to a condition where each species, including H$_2$, is present under standard state conditions (1

54
atm and 298 K). Experimentally, coupling is possible only because H₂ is liberated from the reaction mixture under reflux, so taking the energy of the “final” ester product as +5.5 kcal/mol on the amidation NHR/OR PES in Figure 2.13 is clearly not representative of the catalytic system, and the contribution of the direct NHR/OR route may in practice be much greater than the one suggested by the standard state energy profile. To put it differently, because H₂ is removed from the reaction mixture under dehydrogenative coupling conditions, rehydrogenolysis of the ester into an aldehyde and an alcohol becomes less likely, leaving the ester/amine coupling route as the “only” remaining option for carboxamide formation. Interestingly, if esters are produced and amidated in the dehydrogenative coupling of amines into carboxamides, esters can in turn be produced and hydrogenated in the reverse hydrogenation of carboxamides starting with 2-Ru-H. This most unusual inference should of course be valid independent of the true mechanism of ester amidation. In the ion-pair slippage framework, carboxamide hydrogenation would initially proceed by the H/NHR metathesis route described in Figure 2.1. As the reaction proceeds, the concentration of the alcohol will increase. If the equilibrium concentration of 2a-Ru-OR under H₂ becomes significant, it can undergo an OR/NHR metathesis with the carboxamide to give an ester and 2a-Ru-NHR. 2a-Ru-NHR will give the amine hydrogenation product. Figure 2.1 establishes that hydrogenation of the ester using 2-Ru-H should be comparatively fast, so this will give the second hydrogenation (alcohol) product of the initial carboxamide.
K. Conclusion

Based on detailed electronic structure calculations, the present study proposes that the dehydrogenative coupling of amines with alcohols or esters into carboxamides that can be catalyzed by Milstein’s d⁶-[Ru(PNN)(CO)(H)] catalysts (1-Ru) can be understood in terms of simple metal/acyl metathesis transformations. The study assumes that alcohol dehydrogenation into aldehydes is an accessible transformation in the given systems. We start with the observation that complexes related to 1-Ru undergo characteristic metal–ligand cooperative (heterolytic) addition of H₂, ROH and NH₂R to give octahedral d⁶-ruthenium complexes (2-Ru-X) in which a hydride, alkoxide, or an amide is positioned trans to a hydride (Figure 2.16).

![Diagram of complexes](image)

**Figure 2.16.** X/Y - metal/acyl metathesis

Hydride transfer from a metal complex such as 2-Ru-H to a carbonyl group is familiar. Our idea is that the coordinated alkoxide and amide anions (X⁻) of the other octahedral complexes can undergo similar reactions with an aldehyde or an ester.
(RCYO). The immediate product from such reactions on the PES is an uphill contact ion-pair of an \( \alpha \)-substituted alkoxide anion (RCXYO\(^-\)) in which the newly formed C–X bond is coordinated to the metal (X-ion-pair). The coordination of the X group appears to activate the corresponding C–X bond. At first there appears to be nothing unusual or remarkable about the given “transfer” or “coupling” steps. However, if we accept that X can move from 2-Ru-X to an organic acyl-Y molecule, it becomes evident that the same reaction can take place in reverse from the \( \alpha \)-substituted alkoxide via Y. This produces a new octahedral ruthenium complex (2-Ru-Y) and a new organic acyl-X product. The net reaction is a novel metathesis in which X and Y are interchanged between a metal and an acyl group. This simple reaction can be readily used to rationalize much of Milstein’s extraordinary chemistry. All is needed here is a trivial rotation of the \( \alpha,\alpha-\)

\((X)(Y)\)-substituted alkoxide to orient Y in the direction of the metal. The calculations show the process can take place in a least action mode via TS-Slip-1. The characterization of the individual X or Y transfer regions on the PES proved to be a tedious exercise in locating shallow ion-pair minima and elusive flat TSs that can easily be absent from the PES. At the end, the results reveal the H/OR and H/HNR slippage TSs to be the highest energy point on the metathesis PES. For these cases the full metathesis PES is in effect like a single concerted step taking place via TS-Slip-1. For the NHR/OR metathesis on the other hand, the highest energy point is the TS for C–N coupling between the ester and metal-coordinated amide, but this is only slightly higher than the slippage TS. If it occurs without a barrier, ion-pair dissociation into free ions and reassociation by a different group would provide a variation to the ion-pair rearrangement mechanism that would circumvent the slippage TS.
The present study shows that the hydrogenation MLC mechanism involving hemiacetal and hemiaminal formation followed by a metal mediated fragmentation is simply an indirect multistep route to the same metathesis that can be mediated by ion-pair rearrangement. The MLC and ion-pair slippage routes to metathesis are computed to be competitive. A definitive theoretical comparison of the possible role of the different routes to the same reaction in catalysis requires dynamics effects and accurate estimates of both the basicity of the $\alpha$-substituted alkoxides and solvation of the ion-pair, which are not easy to evaluate computationally. We demonstrate for example that changing the solvent continuum from toluene to THF or methanol, or including a methanol molecule explicitly in the calculations shift the results in favor of the ion-pair slippage route. These computed results cannot rule out a role for MLC in catalysis, but at the same time they do indicate that the more direct slippage route is also plausible. Taken at face value, our results based on conventional transition state theory and implicit bulk treatment of solvation indicate that the dehydrogenative coupling of methyl amines with ethanol using 1-Ru proceeds by initial formation of an ester via an ion-pair OR/H metathesis followed by ester amidation via a low energy ion-pair NHR/OR metathesis.
CHAPTER III
DIRECT METATHESIS FOR HYDROGENATION OF ESTERS WITH POTENTIAL CATALYSTS

A. Overview

The MLC and ion-pair mediated mechanisms provide fundamentally different routes to achieve the same H/OR metathesis reaction. As stated in the conclusion, given the very different nature of the elementary reactions on the two pathways, the computed small preference for the ion-pair mediated pathway cannot rule out a role of MLC in catalysis. As such it becomes of interest to be able to identify possible experiments that may distinguish between the two. One such possibility is to identify target compounds that would lack the acidic proton. However, the methylene linker can in fact exert structural or electronic effects that facilitate the kinetics of the H/OR metathesis in one way or another. The calculations can be particularly useful in comparing the reactivity of related complexes.

In this section we consider two analogs of 1-Ru-H that lack a methylene unit of the phosphine arm namely $1^{\text{CF}_2}$-Ru-H and $1^\text{O}$-Ru-H.

![Chemical structures]

Figure 3.1: Milstein’s octahedral catalyst (left), and after replacing the CH$_2$ linker by CF$_2$ (center) and by O (right).
$^{1\text{CF}_2}$-Ru-H gets rid of the acidic protons in 1-Ru-H by simply substituting the tetrahedral methylene linker of the phosphine arm by another tetrahedral CF$_2$ one. $^{1\text{O}}$-Ru-H on the other hand, substitutes the CH$_2$ linker by an oxygen atom. As shown in the following figure and Table, the optimized geometries of the two complexes have significant differences. Taking the dihedral angle to be measured between the phosphorous, the two carbons of the methylinic linkers and the nitrogen, values of the angles for 1-Ru-H and $^{1\text{CF}_2}$-RuH are around 48°. Upon substitution of the linker by oxygen, the degree of tilting is decreased to 38. The change in the dihedral changes the spacing over the catalyst where slippage takes place.

<table>
<thead>
<tr>
<th>Octahedral Complex</th>
<th>Dihedral Angle (degree)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Ru-H</td>
<td>47.6</td>
</tr>
<tr>
<td>$^{1\text{CF}_2}$-Ru-H</td>
<td>47.7</td>
</tr>
<tr>
<td>$^{1\text{O}}$-Ru-H</td>
<td>38.4</td>
</tr>
</tbody>
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**Figure 3.2:** 3-D side view of 1-Ru-H (left), $^{1\text{CF}_2}$-Ru-H (center), and $^{1\text{O}}$-Ru-H (right) along with their corresponding dihedral angles.
B. Ester Hydrogenation with $^{1}\text{CF}_2$-Ru-H

We follow the same approach as the main reaction pathway that has been discussed in the previous chapter, taking the simplest ester, methyl acetate, as the reference for calculations.

\[
\begin{align*}
\text{methyl acetate} & \quad \xrightarrow{\text{H}} \quad \text{acetaldehyde} \\
^{1}\text{CF}_2\text{-Ru-H} & \quad + \quad \text{MeO}^+\text{Me} \quad \rightarrow \quad ^{1}\text{CF}_2\text{-Ru-O-Me}
\end{align*}
\]

We compare the results of the potential catalyst ($^{1}\text{CF}_2$-Ru-H) with the one previously calculated (1-Ru-H), under the same levels of theory that was used in the previous chapter; we find out that the energy values are comparable. In the previous 1-Ru-H reported data, the reaction path with the most thermodynamically favored conditions were found to be using toluene as a solvent and under the M06L basis set in SMD continuum.

First of all, comparing the two catalysts, 1-Ru-H and $^{1}\text{CF}_2$-Ru-H, we can find out that for the reported data for the Ru-H bond lengths are 1.73 and 1.71 Angstrom respectively while the hydrides in the $^{1}\text{CF}_2$-Ru-H are of bond lengths 1.74 and 1.70 Angstroms respectively; this could be an indicator of relatively different activations of the trans hydrides of the potential catalyst as well as the original 1-Ru-H catalyst. In addition, both complexes exhibit a puckered octahedral geometry; a factor that would certainly enhance exposing the hydride to the carbonyl.
Figure 3.3: Comparison of the potential energy surfaces of the catalysis of 1-Ru-H (top) and 1-Ru-CF2-H (bottom) under M06 level using toluene as a solvent.

In both cases mentioned, the reaction under study can be broken down into two consecutive parts, the hydride transfer from the Ruthenium to the carbonyl which will
result in a positively charged metal complex and a negatively charged carbonyl. The second step involves the transfer of the Alkoxide from the carbonyl to the Ruthenium center of the catalyst ending up with an alkoxy group coordinated with the metal and an aldehyde. Those two steps are mediated by a transition state that is thought of to be as a rotation of the carbonyl bond where the hydride, originally pointing at the metal center, would exchange positions with the alkoxide, rendering the oxygen pointing to
Ruthenium. In addition, the focus in this part of the study is to compare the energies of the slippage mechanism, for this reason, the focus was only on the ion pair formation and rearrangements and its respective transition states and not on the hydride transfer or the alkoxide transfer transition states.

As for the formation of the first ion pair, 1-Ru-H shows to form a relatively lower ion pair formation in toluene under M06 theory level than 1\textsuperscript{CF2}-Ru-H; the value of ΔΔG (i.e. the difference between the energies of the similar states at the potential energy diagrams of the two reaction pathways) was found to be +0.6 kcal/mol which indicates a small relative stability of the ion pair of the Milstein catalyst over our proposed catalyst. The values of the bond lengths of C—H in both ion pairs are very similar, with a slight difference in lengths (0.06 Angstroms shorter bond in case of 1\textsuperscript{CF2}-Ru-H). Similarly, both ion pairs exhibit similar coordination bonds between Ruthenium and the Hydrides, with values of 2.04 and 2.06 Angstroms for 1-Ru-H and 1\textsuperscript{CF2}-Ru-H respectively, indicating similar activations of the Ruthenium hydride bonds.

\[ \text{Figure 3.4: Slippage transition state for } 1\textsuperscript{CF2}-\text{Ru-H system} \]
The highest points on the potential energy surfaces for both reaction pathways is a slippage transition state in which the C—H and C—OMe bonds on the carbonyl rearrange in a way that the latter would be pointing towards the metal center of the octahedral catalysts. Comparing the transition states of both catalysts, we find out that both show similar values of imaginary frequencies of 106 cm⁻¹ and 101.15 cm⁻¹ respectively for 1-Ru-H and 1^{CF2}-Ru-H. The value of ΔΔG was calculated to be +1.8 kcal/mol which indicates that the slippage for the 1-Ru-H is slightly more favorable; this could be due to the fact that we replaced the methylene linker by CF₂, since fluorides are more electronegative, they would affect the electronic distribution on the whole catalyst and especially on the Ruthenium atom and its surrounding, which in turn will reduce the negative charge on the hydride so that its transfer would be a bit less favored. Yet, considering the value of ΔΔG (+1.8 kcal/mol) we would infer that the change in electronic properties did not have the drastic effect on the proposed mechanism and the slippage would still be probable for the 1^{CF2}-Ru-H The geometries of the slippage transition states for both 1-Ru-H and 1^{CF2}-Ru-H are quite similar and comparable; with around 0.04 Angstroms difference in Ru—H bonds and Ru—OMe bonds we can still consider both stages to be trending in a similar manner.

The Slippages in both reactions will lead to the formation of the second ion pair in which the Ruthenium will be coordinated to the Methoxy group. Comparing the two ion pairs, we find out that the value of ΔΔG is equal to +3.8 Kcal/mol which is quite significant and in fact being almost double the value of ΔΔG for the slippage (we induce this comparison since we consider the free energy values of the slippage to be the barrier for this reaction). Another parameter to be considered over here is the difference in the calculated bond lengths between the two ion pairs.
In fact, the calculated minimum for the $^{1}$CF$_{2}$-Ru-H system is not ion pair, unlike the ion pair identified in the previous chapter, this identified complex is an aldehyde adduct. The two ion pairs show a significant difference in the C—O bond lengths, while it was around 1.6 Angstroms in 1-Ru-H, it increased by around 0.83 Angstroms to reach a value of 2.43. Such a value of a C—O bond length is considered to be very high compared to the general values reported for the carbon-oxygen bonds by Allen et al. in their determination of bond lengths by X-ray and neutron diffractions (reported values were in the range of 1.4 Angstroms). This large deviation from the standard values of bond length could be an indicator of the dissociation of the C—O bond and more tendency to form the products. In other words, we can say that the ion pair in $^{1}$CF$_{2}$-Ru-H would dissociate as soon as it is formed due to first the relatively higher free energy than 1-Ru-H and secondly due to the existence of the long C—O bond. In fact, we can also assume that the calculated minimum does not correspond to an actual ion pair due to the high C—O bond, this effect is due to the smoothness of the potential energy surface, in which the system goes directly into dissociation as soon as it
overcomes the slippage energy barrier, thus the calculated minima is more of a precomplex to the cleavage of the C—OR bond that would eventually lead to the aldehyde and Ru—OMe products. This assumption can be also validated by considering the final products of the reaction.

$$\text{1-Ru-H} + \text{MeO} \xrightarrow{\text{methyl acetate}} \text{H} \xrightarrow{\text{MeO}} \text{1-Ru-O-Me}$$

$$\Delta G = +7.4 \text{ Kcal/mol}$$

$$\text{1CF}_2\text{-Ru-H} + \text{MeO} \xrightarrow{\text{methyl acetate}} \text{H} \xrightarrow{\text{MeO}} \text{1CF}_2\text{-Ru-O-Me}$$

$$\Delta G = +3.8 \text{ Kcal/mol}$$

**Figure 3.6:** Thermodynamics of the two metathesis reactions indica products in addition to the values of Gibb’s Free Energy for the whole process.

Significantly, the value of $\Delta G$ of the second reaction came up to be +3.8 kcal/mol much less than the value of the free energy for the first reaction which was found to be +7.4 kcal/mol. This significant difference in energies ($\Delta \Delta G = -3.6 \text{ kcal/mol}$) can be a factor to enhance the efficiency of the $1^{\text{CF}_2}\text{-Ru-H}$ potential catalyst as a smaller overall free energy would certainly be thermodynamically more favored. Another factor that could play a role in stabilizing the product of the second reaction over that of the first reaction could be the presence of the two fluorides. As we discussed it previously, the two fluorides affect the partial charge on the Ruthenium atom and thus affect the way that the metal would interact with various electropositive and electronegative atoms. In 1-Ru-OMe, and according to the Mulliken population analysis, the charge on Ru was +0.658 while in $1^{\text{CF}_2}\text{-Ru-OMe}$ the charge was found to be +0.750, being more
electropositive; thus our potential catalyst will have more tendency to attract and an increased probability in bonding to the electronegative oxygen of the methoxy group.

C. Ester Hydrogenation via 1°-Ru-H

The second potential system we study is the one involving 1°-Ru-H as the potential catalyst and following the same direct metathesis route as before:

The energy comparison between the previously calculated 1-Ru-H system and the potential 1°-Ru-H system is done under the same level of theory, M06L, and using Toluene as a solvent in SMD continuum. The energy values on different points of the potential energy surface were comparable.

For the initial catalysts, the values of the Ru—H bonds are comparable, with 1.69 and 1.73 Angstroms in the case of 1°-Ru-H, which is slightly different than the bond lengths in 1-Ru-H (1.71 and 1.73 Angstroms). This indicates, in a similar manner for 1CF2-Ru-H, that the trans hydrides are not equivalent and eventually one is more activated and can be transferred to the substrate. In addition, the puckered geometry of the two catalysts would be another factor that enhances the exposing of the hydride to the carbonyl and provide the suitable space for the substrate to fit into the catalyst.
Figure 3.7: Comparison of the potential energy surface of the catalysis using 1-Ru-H (top) and 1^0-Ru-H (bottom) under M06 level using toluene as a solvent.

As for the formation of the first ion pair, 1-Ru-H shows to form a relatively lower ion pair formation in toluene under M06 theory level than 1^0-Ru-H; the value of ΔΔG (i.e. the difference between the energies of the similar states at the potential energy diagrams of the two reaction pathways) was found to be +1.8 kcal/mol
which indicates a small relative stability of the ion pair of the Milstein catalyst over our potential catalyst. The values of the bond lengths of C—H in both ion pairs are very similar, 3a-IP-H has a C—H value of 1.27 Angstrom while 3a-O-IP-H shows a value of 1.21 Angstroms, with a slight difference in lengths (0.06 Angstroms shorter bond in case of 1O-Ru-H). Also, both ion pairs exhibit similar coordination bonds between
Ruthenium and the Hydrides, with values of 2.04 and 2.02 Angstroms for 1-Ru-H and 1⁰-Ru-H respectively, indicating that the activation of the Ru—H bonds are occurring in a very similar manner.

**Figure 3.8:** Slippage transition state for 1⁰-Ru-H system

Next we take a look at the transition state of the chemical reaction. The slippage transition state occurs similarly here where the C—H and C—OMe bonds on the carbonyl rearrange in a way that the latter would be pointing towards the Ruthenium metal center of the octahedral catalysts. Comparing the transition states of both catalysts, we find out that both show similar values of imaginary frequencies of 106 cm⁻¹ and 110.52 cm⁻¹ respectively for 1-Ru-H and 1⁰-Ru-H. The value of ΔΔG was calculated to be +0.8 kcal/mol, 1 kcal/mol less than the difference between slippages of 1-Ru-H and 1⁰-Ru-H. Also, the geometries of the slippage transition states for both 1-Ru-H and 1⁰-Ru-H are quite similar and comparable; the Ru—H bond in 1⁰-Ru-H system is 2.41, being 0.06 Angstroms shorter than the slippage in 1-Ru-H, similar values of Ru—O bond lengths are obtained, with 2.92 for 1⁰-Ru-H compared to the very slightly longer value (2.99 Angstroms) for 1-Ru-H.
The Slippages in both reactions will lead to the formation of the second ion pair in which the Ruthenium will be coordinated to the Methoxy group. Comparing the two ion pairs, we find out that the value of $\Delta \Delta G$ is equal to -5.7 Kcal/mol which is quite considerable. Unlike the $1^\text{CF}_2$-Ru-H system, in $1^\text{O}$-Ru-H; calculated minimum shows a significant stability relative to the corresponding slippage transition state. Another parameter to be considered over here is the difference in the calculated bond lengths between the two ion pairs.

\[ \text{Figure 3.9: The aldehyde adduct on the potential energy surface of } 1^\text{O}$-Ru-H \]

The two ion pairs show a significant difference in the C—O bond lengths, while it was around 1.6 Angstroms in 1-Ru-H, it increased by around 0.88 Angstroms to reach a value of 2.48. Also in this case, the bond length of C—O is considered to be very high compared to the general values reported for the carbon-oxygen bonds by Allen et al. in their determination of bond lengths by X-ray and neutron diffractions (reported values were in the range of 1.4 Angstroms). This large deviation from the standard values of bond length could be an indicator of the dissociation of the C—O
bond and more tendency to form the products. In other words, we can say that the second ion pair in $1^\text{O}$-Ru-H, same as in the case of $1^\text{CF}_2$-Ru-H, would dissociate as soon as it is formed due to first the relatively higher free energy than that of 1-Ru-H and secondly due to the existence of the long C—O bond. In fact, we can also assume that the calculated minimum does not correspond to an actual ion pair due to the high C—O bond, this effect is due to the smoothness of the potential energy surface, in which the system goes directly into dissociation as soon as it overcomes the slippage energy barrier. Similar to the $1^\text{CF}_2$-Ru-H case, the calculated minimum on the PES corresponds to an aldehyde adduct that is a precomplex to the dissociation of the C—OR bond cleavage.

![Chemical reaction](image)

**Figure 3.10:** Thermodynamics of the two metathesis reactions indicating initial and final products in addition to the values of Gibb’s Free Energy for the whole pathway.

Significantly, the value of $\Delta G$ of the second reaction came up to be $+3.9$ kcal/mol, which is much less than the value of the free energy for the first reaction which was found to be $+7.4$ kcal/mol. This significant difference in energies, same as the case of the previous potential system, which is found to be $\Delta \Delta G = -3.5$ kcal/mol,
can be a factor to enhance the efficiency of the $1^\circ$-Ru-H potential catalyst as a smaller overall free energy would certainly be thermodynamically more favored.

**D. Conclusion**

Upon building up potential catalyst systems by replacing the methylinic linker by one that lacks the acidic protons that are the highlight of the generally accepted Metal-Ligand Cooperation, we were still able to identify similar potential energy surfaces to our proposed mechanism. This would imply that the presence of the methylinic protons is insignificant in the hydrogenation of esters.
CHAPTER IV

CONCLUSION

In conclusion, we present in this study and using electronic structure DFT methods, an alternative metathesis mechanism to the dehydrogenative coupling of alcohols and amides into esters and carboxamides. The identified route is a more direct, low energy route to the coupling that involves formation of two ion pairs mediated by a simple rearrangement within the ion-pairs to transfer the hydride from the metal center to the substrate. Two key features of the proposed mechanism include a direct metathesis route and a lower potential energy system compared to the generally accepted Metal-Ligand Cooperation mechanism.

In addition, DFT calculations on potential systems showed that the absence of the methylinic proton, which plays an essential role in MLC mechanism, would not have an effect on the calculated energies for the slippage mechanisms of systems involving two different potential catalysts for ester hydrogenation.

The study paves the way to reconsidering the Metal-Ligand Cooperation mechanism that has been thought of to be as a generally accepted mechanism and to start pointing out more direct metathesis mechanisms that do not involve the modification of the ligand or part of it. Moreover, Milstein and co. are still in the process of synthesizing new catalysts and reporting new transformations, so theoretical studies will still be necessary to explain the mechanisms of these systems. Considering a newly proposed mechanism could be helpful in directing the mechanistic studies into more vivid conclusions.
REFERENCES


42. Frisch, M.J., et al., *Gaussian 09.* 2009, Gaussian, Inc.: Wallingford, CT, USA.


