Pulsed Current Electrodeposition of Silicon Thin Films Anodes for Lithium Ion Battery Applications
Review

Product Selectivity in Homogeneous Artificial Photosynthesis Using [(bpy)Rh(Cp*)X]^{n+}-Based Catalysts

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Abstract: Due to the limited amount of fossil energy carriers, the storage of solar energy in chemical bonds using artificial photosynthesis has been under intensive investigation within the last decades. As the understanding of the underlying working principle of these complex systems continuously grows, more focus will be placed on a catalyst design for highly selective product formation. Recent reports have shown that multifunctional photocatalysts can operate with high chemoselectivity, forming different catalysis products under appropriate reaction conditions. Within this context [(bpy)Rh(Cp*)X]^{n+}-based catalysts are highly relevant examples for a detailed understanding of product selectivity in artificial photosynthesis since the identification of a number of possible reaction intermediates has already been achieved.

Keywords: artificial photosynthesis; product selectivity; reaction mechanism; rhodium

1. Introduction

Based on mankind’s tremendous impact on planet earth, the scientific question of whether a new division of geological time should be defined and consequently be termed “anthropocene” has emerged [1]. Due to the mostly detrimental influences of human activity on our environment, as well as for economic reasons, catalysis has played a major role in the development of industrial chemistry [2]. The continuous enhancement of catalysts is also driven by the intellectual insatiable desire of keeping up with nature’s highly selective operating enzymes, whose activity has been optimized to perfection during millions of years of evolution. Within this context, highly stereoselective catalysis in organic synthesis has been one of the most outstanding achievements during the last decades of catalyst development [3].

On the other side, nature is additionally not only able to perform ground state chemistry, but also uses light energy as a primary resource for several energy demanding chemical conversions. This is accompanied by the challenging handling of excited states and highly reactive intermediates. Due to a precise arrangement of functional proteins within the thylakoid membrane of green plants, the light driven water splitting of natural photosynthesis generates the indispensable ubiquitous oxidant O_{2}, as well as reducing equivalents in the form of NADPH along with the energy carrier ATP, and can therefore be ascribed from a chemist’s point of view as one of nature’s biggest masterpieces [4–6]. Although light nowadays has its application as a key reagent or energy source in selective chemical transformations [7], the knowledge about product selectivity with respect to artificial photosynthesis is still in its infancy.

In this feature article, recent work regarding the product distribution of homogeneous artificial photosynthetic systems capable of generating different reaction products at the catalytic center will be discussed, with a clear focus on RhCp*-based catalysts. In order to pave the way towards developing...
an understanding of the reaction outcome of the different systems, overviews of the underlying reaction mechanisms at the catalytic centers will be provided.

2. General Considerations on Product Selectivity within Artificial Photosynthetic Systems and Selected Aspects of the Homogeneous Photocatalytic CO₂ Reduction

Due to its highly sophisticated architecture and the presence of two catalytic centers, natural photosynthesis is able to drive the overall water splitting reaction. By virtue of the complex interplay of light absorption, electron transfers, and catalysis, typical artificial photosynthetic systems focus on one of the redox half reactions. Besides the photosensitizer (PS), which absorbs the light, and the catalytic center (CC), that drives the substrate conversion after electron transfer reactions with the excited photosensitizer (PS*), a sacrificial agent has to be added in order to close the catalytic cycle. As depicted in Figure 1, an artificial photosynthetic system can be multifunctional, if the activated catalytic center (CCₐ) is able to convert different substrates into different redox products. Therefore, the catalytic cycles have to exhibit a common branching point. However, the branching point of the catalytic cycles does not necessarily need to be the catalytically active species itself; a simple catalytically inactive reaction intermediate could be envisaged instead.

![Figure 1](image-url)

**Figure 1.** General diagram for a multifunctional artificial photosynthetic system capable of transforming two different substrates (A and C) to the appropriate redox products (B and D). The system’s main components are the photosensitizer (PS) and the catalytic center (CC, in its activated/inactive form denoted with the subscript a/i). The regeneration of PS by a reaction with a sacrificial agent is omitted for clarity.

From Figure 1, the general idea of how the product distribution can be principally switched between different reaction outcomes becomes evident. If the amount of product B needs to be maximized, catalytic pathway 2 should be avoided as effectively as possible. The easiest approach to achieve this would be simply not adding substrate C to the system. In cases where this is not possible (usually due to the presence of protons as a substrate for hydrogen evolution catalysis), k₁ needs to be much higher than k₂, which could be accomplished by choosing appropriate reaction conditions, such as the right pH value, light intensity, water content, etc., or by designing a more sophisticated ligand environment around CC, which would allow substrate A to react much faster with CCₐ than substrate C, based on steric or electronic factors.

One prominent example for the possible generation of different redox products is the photocatalytic reduction of carbon dioxide, whose general mechanism will be briefly discussed in the following section for the purpose of illustrating the general catalysis scheme of Figure 1.

Due to the unavoidable depletion of fossil fuels, the conversion of CO₂ into valuable chemicals would establish a sustainable supply of man-made energy carriers. Hence, a lot of work has been done in order to design highly active photocatalysts using solar energy for driving CO₂ reduction reactions.

A broad variety of coordination compounds (bearing Mn, Fe, Co, Ni, Ru, Re, or Ir as the central metal) have been found to function as catalysts for the photocatalytic conversion of CO₂, mainly generating the two electron reduced carbon-based products CO and HCOOH, as well as H₂ as the side
product from the competing proton reduction \[8\]. As described above, the detection of multiple redox products indicates the presence of at least one reactive species, appearing in two or more catalytic cycles. Figure 2 shows a simplified reaction scheme for the key intermediates of the photocatalytic CO\(_2\) reduction processes \[9\].

A coordination compound in an oxidized state (M\(_{\text{ox}}\)) represents the starting point of the overall cycle. A light driven electron accumulation onto M\(_{\text{ox}}\) yields the reduced intermediate M\(_{\text{red}}\), which either takes up a proton to generate a metal hydride species or directly binds carbon dioxide, forming a metal carboxylate species. The latter liberates via an initial protonation step of the reduced carbon species CO, as well as water. HCOO\(^-\) is produced by the insertion of carbon dioxide into the metal hydride bond, forming a labile formate complex. In addition, the closing of the catalytic cycle can also be achieved by the reaction of M–H with a second proton, thereby generating molecular hydrogen.

The chemoselectivity of any process will mainly be determined by differences in the reaction kinetics at the branching points \[9\]. Crucial factors herein are predominantly the energy content of the relevant species generated from the common branching point intermediate with the appropriate substrates and the available concentration of the latter. Coherently low energy barriers and high substrate concentrations for a specific dark reaction following the light driven generation of a certain branching point species will increase the amount of a certain reaction product in an artificial photocatalytic system.

With a much more sophisticated ligand environment at the catalytic centers in comparison to the man-made ones, nature drives the transformation of CO\(_2\) into valuable chemicals using earth abundant metals. A key moiety of several enzymes dealing with the incorporation of CO\(_2\) into organic substrates is the metal carbamato species, which can be found in rubisco and biotin-dependent enzymes \[10\]. Its formation can be interpreted similarly to the insertion of carbon dioxide into the M–H bond shown in Figure 2, indicating that the latter represents a very important step for the generation of useful carbon containing chemicals.

![Figure 2](image.png)

**Figure 2.** Simplified reaction scheme for the competing reduction reactions involved in artificial photocatalytic CO\(_2\) reducing systems. The n electrons stem from an excited/reduced photosensitizer.

It should be noted that the above discussed general mechanism is only a highly superficial view on the actual complex sequence of elementary steps underlying the overall photocatalytic CO\(_2\) reduction processes; interested readers are referred to several selected review articles \[8,9,11–14\]. Nevertheless, the simplified reaction scheme depicted in Figure 2 was chosen in order to clarify the role of catalytic intermediates exhibiting multiple reaction pathways.

Although principally at least three different reduction products can be obtained by performing a photocatalytic carbon dioxide reduction, several systems have been found to exhibit a high chemoselectivity for either CO or formate production. Transition metal complexes based on the
[(bpy)Re(CO)_3X]^n+ structural motive mostly form carbon monoxide as the main product in water-free solvent systems, with chemoselectivities reaching almost 100% [15–20]. Interestingly a supramolecular heterodinuclear Ru–Re photocatalyst showed an opposed product selectivity towards formate generation in an aqueous solution [21]. This solvent dependent behavior could be attributed to the different reaction kinetics for the formation of the intermediates M–H and M–CO₂ from the common branching point species M_red, as depicted in Figure 2. As the proton concentration in an aqueous solution is higher than in an organic solution, the formation of M–H will be accelerated and consequently, the yield of HCOO⁻ increases.

Impressive selectivities and TONs (turnover numbers) have also recently been obtained with a cobalt cryptate as the catalyst [22]. In contrast to the mentioned Re complexes, the Ru-based catalyst of the general composition [(NN)Ru(CO)_2Cl_2] or [(NN)_2RuX_2] (X = CO, Cl, DMF; NN = N,N-chelating diimine ligand) shows a selectivity towards formate production in the majority of cases, although sometimes CO can also be obtained as the major product [8,23–25].

Since access to the much more valuable higher reduced carbon species, such as methanol using highly efficient and durable photocatalytic processes, would represent a major turning point in the global energy crisis, further developments in the field of visible light driven CO₂ reduction and an increase in the knowledge of the underlying reaction mechanisms are highly likely to occur.

3. Application of [(bpy)Rh(Cp*)X]^n+‐Like Coordination Compounds as Multifunctional Redox Catalysts

As measured by the amount of possible selective redox transformations, [(NN)Rh(Cp*)X]^n+‐like catalysts hold a leading position within artificial photosynthetic systems (Cp* = pentamethylcyclopentadienyl, X = Cl, OH, H_2O). As depicted in Figure 3, a broad number of substrates can be reduced using [(NN)Rh(Cp*)X]^n+‐like structures as catalysts by means of chemical, electrochemical, or photochemical methods. In addition, imines [26–28] and α,β-unsaturated carbonyl compounds [29] can also be reduced with this type of catalyst, as well as nitrogen containing heterocycles such as quinolines or quinoxalines [30,31]. To illustrate the high potential of this organometallic rhodium-based catalyst, a fully homogeneous system for the total photocatalytic water splitting using [Co₄(H₂O)₂(PW₆O₂₄)₂]^10⁻ as the water oxidation catalyst and [(bpy)Rh(Cp*)(H₂O)]^{2+} as the NAD⁺ reducing catalyst using [Ru(bpy)_3]^2+ as a photosensitizer has been reported [32]. However, only turnover numbers (TONs) below 10 were found for the Rh catalyst and the Ru chromophore.

**Figure 3.** Selected reduction reactions that can be accomplished by [(bpy)Rh(Cp*)X]^n+ under appropriate reaction conditions. Literature: reaction (a) [33–38], reaction (b) [39,40], reaction (c) [26,29,41–44], reaction (d) [45–55], and reaction (e) [48,56–59].

As shown in Figure 4, the generation of a catalytically active species can be achieved on the one hand by the thermal decomposition of a suitable hydride donor in the coordination sphere of the catalyst using formate most frequently [56], but also alcohols [44] and phosphite [57], or on the other hand, by a sequential two electron reduction via an electrochemical or photochemical pathway and a concomitant oxidative addition of a proton.
The molecular structure of the actual active rhodium catalyst has been under debate during the last years. Since the late 1980s, the catalytically active species has been described as a Rh\textsuperscript{III}–H (structure A in Figure 4) [33,45]. The experimental proof of the existence of this species has been performed by Steckhan et al. [51], as well as Fukuzumi et al. [60], using a 6,6′-dimethylated bipyridine as the \textit{N,N}-chelating ligand. However, as for other sterically less demanding \textit{N,N}-chelating ligands, the Rh\textsuperscript{III}–H moiety has so far not been observed. The independent discovery of an alternative possible catalytically active species (see structure B in Figure 4) by Pitman et al. [61], as well as Quintana et al. [62], established the potential presence of multiple reactive species involved in the reductive transformation of several substrates.

Due to the endo orientation of the hydrogen atom, the conversion of structure A into structure B can be regarded as the reductive elimination of a proton from the rhodium center onto a carbon atom of the Cp\textsuperscript{*} ligand. As species A was detected at low temperatures and the conversion into species B occurs upon warming, the thermodynamically favorable formation of the latter most likely proceeds via the initial generation of A [61,62].

\begin{center}
\includegraphics[width=0.5\textwidth]{figure4.png}
\end{center}

\textbf{Figure 4.} Simplified access paths to the catalytically active species.

In a very recent report, the influence of the nature of the \textit{N,N}-chelating ligand on the structure of the catalytically active species was discussed [63]. Similarly to the bulky 6,6′-dimethylated bipyridine, the 6,6′-dihydroxymethyl bipyridine described in this study also shows a \textsuperscript{1}H-NMR signal attributable as Rh\textsuperscript{III}–H.

However, when using 3,3′-, 4,4′-, or 5,5′-dihydroxymethyl bipyridine as the \textit{N,N}-chelating ligand (see Figure 5), the formation of the reduced Cp\textsuperscript{*} ligand (Cp\textsuperscript{*}H) was observed instead. These findings were rationalized by the enlarged Cp\textsuperscript{*}-Rh distance in the case of the 6,6′-dihydroxymethyl bipyridine due to steric interactions with the Cp\textsuperscript{*} ligand, which might finally prohibit hydrogen transfer onto the five-membered ring.

\begin{center}
\includegraphics[width=0.5\textwidth]{figure5.png}
\end{center}

\textbf{Figure 5.} Molecular structure of different regioisomeric dihydroxymethyl bipyridines bound to the Rh(Cp\textsuperscript{*})Cl fragment [63].

As depicted in Figure 4, access to the catalytically active species is not only possible via a thermal process by formally abstracting “H” from a suitable donor, but also by means of photo- and
Detailed investigations demonstrated that the special ligand environment of the \([(\text{N},\text{N})\text{Rh}^{\text{III}}(\text{Cp}^*)X]\^{\text{II}}\)-like catalyst grants it a favorable electrochemistry for applications in homogeneuos photo- and electrocatalysis. Access to the reactive \([(\text{N},\text{N})\text{Rh}^{\text{I}}(\text{Cp}^*)]\) intermediate is accomplished via a two electron reduction at a rather low cathodic voltage (ranging from $-700$ to $-900$ mV against Ag/AgCl depending on the N,N-chelating ligand) \([51]\). After the addition of one electron, the primarily obtained coordinatively unstable Rh\(^{\text{II}}\) species disproportionates extremely rapidly, leading to the desired Rh\(^{\text{I}}\) compound (for the majority of the applied complexes \([64]\)), as this process is merely connected with the easy loss of a weakly bound monodentate ligand \([33,37,65]\). In addition, a crystal structure of the reduced d\(^8\) intermediate \([(\text{bpy})\text{Rh}^{\text{I}}(\text{Cp}^*)]\) revealed that the bpy and the Cp\(^*\) plane are lying perpendicular to each other, as depicted in Figure 4, different to the piano-stool like structure of the d\(^6\) species \([(\text{bpy})\text{Rh}^{\text{III}}(\text{Cp}^*)X]\^{\text{II}}\) \([66]\).

As the thermal, as well as the photo- or electrochemical, route leads to intermediate A, substrate conversions that were to date only reported using the thermal approach should also be realizable using electrochemical or photochemical methods under appropriate reaction conditions. This would further expand the already wide range of applications of \([(\text{bpy})\text{Rh}(\text{Cp}^*)X]\^{\text{II}}\)-like catalysts in homogeneous photocatalytic systems.

### 3.1. Reduction of Carbonyl Compounds to Primary or Secondary Alcohols

A typical substrate for transfer hydrogenations are carbonyl compounds. Within several thermal processes, the use of chiral N,N-chelating ligands coordinated to the Rh(Cp\(^*\)) moiety allowed ketone reductions with high enantioselectivity, indicating that the ligand-induced asymmetry of the whole catalyst is sufficiently preserved at the rhodium center to sterically discriminate between the two different enantiotopic faces of the ketone \([42–44]\). In addition, chromenones, a group of cyclic \(\alpha,\beta\)-unsaturated ketones, were reduced to the fully saturated chiral alcohol with high ee-values, using HCOONa as a hydride donor \([29]\). A number of these stereoselectively reducible ketones are depicted in Figure 6.

![Figure 6](image-url)

**Figure 6.** Range of carbonyl compounds that were reduced stereoselectively to the corresponding alcohols using enantiopure \([(\text{N},\text{N})\text{Rh}(\text{Cp}^*)X]\^{\text{II}}\) catalysts and sodium formate as a hydride source \([29,42–44]\).

In order to evaluate a rational reaction scheme for the rhodium catalyzed reduction of carbonyl compounds to the appropriate alcohols, Leiva et al. performed concentration dependent experiments using a variety of aldehydes and ketones, as well as sodium formate as a hydride donor and \([(\text{bpy})\text{Rh}(\text{Cp}^*)(\text{H}_2\text{O})](\text{OTf})_2\) as a catalyst \([41]\). Although the formation of the catalytically active \([(\text{bpy})\text{Rh}(\text{Cp}^*)\text{H}]^+\) species was proposed to be the rate limiting step, the structure of the substrates had a significant effect on the observable TOFs (turnover frequencies, i.e., turnover number per
time). Generally, aldehydes were converted faster than ketones due to the higher electrophilicity of the carbon atom. In addition, higher reaction rates were observed for substrates with an increased electron density on the oxygen atom, indicating that the binding of the substrate into the coordination sphere of the catalyst is important and may take place in concert with the hydride transfer reaction. From these observations, a simplified reaction mechanism was proposed, as depicted in Figure 7. After the replacement of a ligand L for a hydrido ligand (reaction a) via the decomposition of formate, coordination of the carbonyl compound occurs (reaction b). Following a hydride transfer via a four-membered transition state, the replacement of the alkoxide ligand (or alcohol after protonation) for L regenerates the initial Rh species (reaction c).

Besides the above delineated thermal transfer hydrogenation reactions in which the catalytically active species is present in a significant concentration due to the excess of hydride donors, a selective reduction of aldehydes in the presence of ketones via a homogeneous photocatalytic process was described by Ghosh et al. [26]. Using proflavine as a photosensitizer, [(bpy)Rh(Cp*)Cl]Cl as a catalyst, and TEOA as a sacrificial electron donor, the catalytically active species was generated very slowly, which allowed the maintenance of a continuously low Rh-H amount and an efficient kinetic discrimination of aldehydes and ketones by steric factors. Contrary to the mechanistic investigations of the thermal process described above, the authors suggested that within the photocatalytic process, the hydride transfer from the rhodium center onto the carbonyl compound represents the rate determining step.

![Figure 7. General reaction scheme for the [(bpy)Rh(Cp*)L]n+ catalyzed reduction of carbonyl compounds. Within step a, the catalytically active species can be generated either by a thermal or by a electro- or photochemical process, as shown in Figure 4.](image)

### 3.2. Use of [(bpy)Rh(Cp*)X]n+ Catalysts for Hydrogen Evolution and CO₂ Reduction

Reduced carbon species, as well as molecular hydrogen, are prime targets for the storage of light energy. Due to the rather low volumetric energy density of hydrogen, methods for its incorporation into materials, as well as for its temporary introduction into carrier molecules, are under intensive investigation [67,68]. Within this context, formic acid is one possibility of transiently storing H₂ [69]. In order to use molecular hydrogen as a green fuel, efficient catalytic liberation from the storage materials is an important prerequisite for this technology. It has been shown that [(bpy)Rh(Cp*)(H₂O)]²⁺ is able to selectively decompose formic acid into H₂ and CO₂ via a rhodium formato and a rhodium hydride intermediate without the production of CO [60]. Interestingly, a fast H/D exchange of the RH intermediate was observed, illustrating the bivalent protic and hydridic nature of the metal bound hydrogen. The pKₐ value for the acid base equilibrium of [(NN)Rh(Cp*)H]⁺/[(NN)Rh(Cp*)]⁰ + H⁺ has been determined to lie between seven and eight, depending of the structure of the N,N-chelating ligand [33,58,70].

[(bpy)Rh(Cp*)X]n+ and its derivatives are also efficient catalysts for the electrochemical generation of dihydrogen. In accordance with recently published calculations [62], hydrogen evolution is only observed at low pH values or by the use of strong acids [33−36,38]. The liberation of molecular...
hydrogen from the Rh-H intermediate has been ascribed as the rate determining step of the overall catalysis, whereas the rate constants for the disproportionation reaction yielding the \([\text{bpy}]\text{Rh(Cp*)}\) intermediate and its protonation to form \([\text{bpy}]\text{Rh(Cp*)H}^+\), are multiple orders of magnitude higher [33,65]. Concerning the activity of these Rh-based catalysts, two successive papers reported a faradayic efficiency of 100% for hydrogen formation at \(-0.55\) V vs. SCE by using a polymeric Rh\(\text{Cp*}\) catalyst in an aqueous solution of pH 1 [34,35]. This correlated to a TON of 353 after 14 h.

Therefore, the photocatalytic generation of \(H_2\) using an intermolecular system containing \([\text{Ru(bpy)}_3]^{2+}\) as a photosensitizer, \([\text{bpy}]\text{Rh(Cp*)}(\text{H}_2\text{O})^{2+}\) as a catalyst, and sodium ascorbate as an electron donor showed a strong dependence of the catalytic activity on the proton concentration, with an optimum pH value of 3.6. A negligible formation of hydrogen was observed at pH values higher than five and lower than two, which indicates on the one hand, that a sufficient thermodynamic driving force for the proton hydride reaction needs to be generated, which can be achieved by a high proton concentration, and that on the other hand ascorbate rather than ascorbic acid acts as the electron donor [37]. Additionally, molecular hydrogen was also evolved by thermally produced \([\text{bpy}]\text{Rh(Cp*)H}^+\) in the presence of photoactivated platinum nanoparticles [71]. Herein the rhodium hydrido complexes served as electron donors for the catalytically active particles.

Despite intensive work in the field of catalytic hydrogen evolution using \([\text{bpy}]\text{Rh(Cp*)X}^{n+}\)-based catalysts, a detailed understanding of the important proton hydride reaction liberating \(H_2\) and the closing of the catalytic cycle remains unclear. Besides the possibility of a simple acid base reaction outside of the rhodium coordination sphere, the binding of a second proton to the intermediates A or B of Figure 4 has been hypothesized, resulting in a Rh assisted intramolecular liberation of hydrogen [62].

In addition to the evolution of \(H_2\), the reduction of carbon dioxide has also been achieved by electro- and photochemical methods using \([\text{NN}]\text{Rh(Cp*)X}^{n+}\) catalysts. Besides the direct \(\text{CO}_2\) reduction in the coordination sphere of the catalyst, which will be described here, an indirect pathway by the assistance of an enzyme, powered by the \([\text{NN}]\text{Rh(Cp*)X}^{n+}\) driven regeneration of nicotinamide cofactors will be discussed in the next section.

The electrochemical reduction of \(\text{CO}_2\) has been performed using \([\text{bpy}]\text{Rh}^{\text{III}}(\text{Cp*})\text{Cl}^+\) as a catalyst in aqueous acetonitrile solution [39]. In addition to the formation of formic acid, dihydrogen in a significant and carbon monoxide in a negligible amount were formed. The authors claim that under their experimental conditions, the actual catalytically active species can be described as the neutral, threefold reduced complex \([\text{bpy}]\text{Rh}^{\text{II}}(\text{Cp*})\text{H}\), indicating the necessity of an increased hydricity of the metal bound hydrogen for \(\text{CO}_2\) reduction.

Surprisingly, there is only one report about the direct photocatalytic reduction of carbon dioxide using \([\text{NN}]\text{Rh(Cp*)X}^{n+}\) as a catalyst [40]. \(\text{CO}_2\) has been reduced by \([\text{bpy}]\text{Rh(Cp*)Cl}^+\) in a mixture of \(\text{MeCN:TEOA} = 5:1\), using \([\text{Ru(bpy)}_3]^{2+}\) as a photosensitizer. After 10 h of irradiation, formic acid has been detected as the sole reduced carbon species (TON = 125), besides the significant evolution of hydrogen (TON = 55).

In addition to the above mentioned electrocatalytic and photocatalytic methods of converting \(\text{CO}_2\) into valuable chemicals, the direct hydrogenation of \(\text{CO}_2\) using \(H_2\), thereby generating formic acid, is also feasible using \([\text{bpy}]\text{Rh(Cp*)X}^{n+}\)-based catalysts. In contrast to the Rh complex with the unsubstituted bpy ligand, a marked increase in catalytic activity by the introduction of pH sensitive OH functionalities at the para positions was observed [72]. Deprotonation of the hydroxyl groups under basic conditions generated the oxyanionic species, resulting in an increased electron density at the metal center, which is responsible for the enhancement of catalytic activity (from TON = 216 to 1800). Interestingly, for the iridium analog, the increase of catalytic performance was drastically higher, increasing from a TON of 105 to 190,000 under optimized conditions.

3.3. \(\text{NAD(P)H} \) Formation Using the \([\text{bpy}]\text{Rh(Cp*)X}^{n+}\) Motive

A very interesting property of \([\text{NN}]\text{Rh(Cp*)X}^{n+}\) catalysts is their highly regioselective production of 1,4-dihyro nicotinamide from \(N\)-substituted nicotinamide cations. As the coenzymes \(\text{NAD}^+\) and
NADP$^+$ also exhibit this moiety, [(NN)Rh(Cp*)X]$$^{II^+}$$ complexes allow the construction of semiartificial biocatalytic systems powering enzymatic reduction reactions by transition metal catalyst driven NAD(P)H regeneration. For a detailed overview of the different biochemical applications of these [(bpy)Rh(Cp*)X]$$^{II^+}$$-based cofactor regeneration systems, the interested reader is referred to several review articles [73–75].

It should be noted that other rhodium complexes like [Rh(bpy)$^3$$^{3+}$] and [Rh(terpy)$^2$$^{3+}$], as well as several cobalt complexes, can act as regioselective catalysts for 1,4-dihyronicotinamide formation too [76–81]. However, as for most of these catalysts only very small TONs were observed, [(NN)Rh(Cp*)X]$$^{II^+}$$ has achieved outstanding significance in selective NAD(P)$^+$ reduction catalysis due to its much higher reactivity.

The thermal production of NAD(P)H has most often been performed using formate as a hydride donor [45–47,49,82]. Intensive mechanistic studies performed by Lo et al. using different nicotinamide derivatives, altered in the 1- as well as the 3-position of the pyridine ring, point to a catalytic mechanism, as depicted in Figure 8 [83,84]. As the initial step, formate has to replace the rhodium bound monodentate ligand L. It has been shown that the presence of strong $\sigma$ donor ligands such as NH$_3$ decreases the rate of thermal NAD$^+$ reduction as they compete with formate for the available Rh binding site [46,51,56,59]. A similar but pH dependent behavior was suggested for L = H$_2$O; upon increasing the pH value, the NADH generation slowed down [56]. Since the pK$_a$ value of the metal bound water was determined as 8.2 [70], higher pH values of the solution led to the formation of the hydroxo ligand, which is a stronger $\sigma$ donor than water, and replacement by formate thus becomes more difficult.

![Figure 8. Mechanism of the formate driven nicotinamide cofactor reduction according to Lo et al. [83,84].](image)

However, after the coordination of formate, decomposition affords CO$_2$ and the catalytically active Rh$^{III}$–H species; the latter has been reported to produce hydrogen peroxide in the presence of molecular oxygen with the concomitant reformation the initial Rh$^{III}$–L species [53,56]. The authors proposed a reaction scheme, in which O$_2$ inserts into the Rh–H bond of the electrogenerated [(NN)Rh(Cp*)H]$^+$ species, as depicted in Figure 9 [53]. After protonation and ligand exchange, the initial Rh catalyst gets reformed and H$_2$O$_2$ is liberated.
Whereas a high reduction potential is beneficial in terms of the electrocatalytic activity, the same (see Figure 10). The regioselectivity has been reported to be at least as high as 95% for the 1,4 production varies in the thermal and the electrochemical approach in the opposite direction.

The reaction proceeds finally by the displacement of the reduced nicotinamide by another ligand. Only traces of the 1,6 isomer are produced, which are formed putatively by a subsequent \[\text{[(NN)Rh(Cp*)X]}\] nicotinamide, due to the formation of an energetically favourable six membered transition state (see Figure 10). The regioselectivity has been reported to be at least as high as 95% for the 1,4 product. Only traces of the 1,6 isomer are produced, which are formed putatively by a subsequent \[\text{[(NN)Rh(Cp*)X]}\] catalyzed isomerization of the 1,4-dihydro form. Closing of the catalytic cycle proceeds finally by the displacement of the reduced nicotinamide by another ligand.

As depicted in Figure 10, several nicotinamide-related structural parameters affect the rate of nicotinamide reduction \[83,84\]. Of highest importance is the necessity of a \(\sigma\) donor moiety in the 3-position of the pyridine ring, as with an unsubstituted or 3-methylated derivative, no product formation was observed. Furthermore, an increased \(\sigma\) donor ability accelerates the overall catalysis by making the metal bound hydrogen more hydridic. Substitution patterns which make the carbon atom at C-4 more electrophilic also increase the reaction rates (green arrows in Figure 10).

Similar results have been obtained by the means of electrochemical and photochemical methods \[85\]. Finally, larger \(R\) substituents and the carbamoyl nitrogen inhibit product formation due to steric interactions with the \(\text{Cp}^*\) ligand, as marked in Figure 10.

Depending on the reduction potential of the \(\text{Rh}^{III}/\text{Rh}^I\) couple, the catalytic activity for NADH production varies in the thermal and the electrochemical approach in the opposite direction. Whereas a high reduction potential is beneficial in terms of the electrocatalytic activity, the same
is detrimental when operating thermally by using formate as a hydride donor. As the electron density on the metal increases due to a more basic and electron rich \(N_N\)-chelating ligand, its reduction becomes more difficult, but the subsequent addition of a proton is accelerated [51,54]. The latter property can also be viewed as a higher amount of protonated catalytically active \([(NN)Rh^{III}(Cp^*)H]^+\) at a given pH value, since its deprotonation yields the transiently inactive neutral \([(NN)Rh(Cp^*)]\) species. By operating thermally using formate as a hydride donor, the lower catalytic activity with an increased reduction potential could be interpreted by the different mechanism of generating the catalytically active species. The necessary coordination of the formate ligand, as well as the subsequent hydride abstraction, will be energetically more demanding if the electron density on the rhodium atom will be increased as a consequence of having a more electron rich \(N_N\)-chelating ligand bound to the metal [63].

Inactivation of the rhodium catalyst has been reported for the electrocatalytic reduction of NAD\(^+\) in the presence of amino or thiol groups [86]. Consequently, the mutual inactivation of \([(bpy)Rh(Cp^*)(H_2O)]^{2+}\) and an enzyme bearing cysteine and lysine residues on the accessible protein surface has been observed [87]. A detailed electrochemical investigation on the influence of amino acids on the electrocatalytic activity of \([(bpy)Rh(Cp^*)(H_2O)]^{2+}\) for NADH formation revealed that the strongest inhibitory effect arose from cysteine, histidine, and tryptophane [88]. Although all these detrimental interactions have been observed, a detailed understanding of the underlying molecular processes is still missing. In the case of thermal NADH regeneration using formate as a hydride donor, the lack of catalytic activity can be tentatively explained by a thermodynamically unfavourable ligand exchange from a thiolato ligand to formate [87]. Based on electrochemical studies, it has been revealed that the binding of a chloride ion to the two electron reduced neutral species \([(NN)Rh^{III}(Cp^*)]\) can occur, forming the anionic \([(NN)Rh^{II}(Cp^*)Cl^-]\) species [35]. We hypothesize a similar binding of sulfur or nitrogen containing groups to the rhodium center, which could finally prevent the necessary oxidative addition of a proton as a key step for the generation of the catalytically active species.

As the photochemical generation of NAD(P)H imitates photosystem I of the natural photosynthesis and consequently allows the light driven powering of enzymatic transformations, intensive efforts have been undertaken in order to optimize the catalytic activity of \([(NN)Rh(Cp^*)X]^+\)-based systems (an overview of the working principle is depicted in Figure 11) [55]. A broad variety of photosensitizers have been applied in these systems, ranging from small organic dyes [85,89–91] to functionalized graphenes [92–95], photoredoxactive polymers [96,97], red light absorbing tin porphyrines from Knör and coworkers [98], ruthenium polypyridine complexes [32,97,99], and different solid photoactive materials [100–110].

![Figure 11. Schematic reaction scheme for photobiocatalytic systems.](image-url)
Identically to the photocatalytic generation of hydrogen, the catalytically active rhodium complex is most likely formed via the fast disproportionation of the intermediate Rh$^{III}$ species [33,37]. Since the underlying mechanism for the generation of the protonated active catalyst is similar with respect to the electrochemical and the photochemical approach, the same reactivity trends for different $N,N$-chelating ligands are assumed to be true. This means, in other words, that the more electron rich the catalyst is, the faster the proton uptake becomes, which consequently increases the catalytic activity. However, it should be noted that a higher reduction potential of the Rh$^{III}$/Rh$^{I}$ couple could lead to decreased electron transfer rates from the excited photosensitizer onto the catalyst, thereby overcompensating the positive effect described above. Finally, if the catalytically active species is formed, the light independent transfer of the hydride onto the cationic nicotinamide substrate proceeds either via the mechanism described by Lo. et al. [83,84] or via the reduced Cp* ligand as hypothesized by Pitman et al. [61].

When working with oxidized nicotinamide cofactors as substrates for electrochemical or photochemical reduction processes, special care must be taken in order to prevent the undesirable formation of enzymatically inactive isomers [94], as depicted in Figure 12, since they limit the practical applicability of visible light driven cofactor reductions due to lower enzymatic substrate ($S$) to product ($P$) conversions as a consequence of an inefficient NAD(P)H regeneration.

It has been shown that under photocatalytic conditions and in the absence of the rhodium catalyst, the formation of complex product mixtures is possible [110]. Steckhan and coworkers identified the reduced nicotinamide cofactor product during photoredox catalysis with [Ru(bpy)$_3$]$^{2+}$ as the chromophore in the absence of a rhodium catalyst as the dimeric (NAD)$_2$ species [73]. They propose its formation by radical coupling of the neutral, one electron reduced NAD molecules [111] (see Figure 12), which are formed after electron transfer from the photoreduced chromophore.

![Figure 12](image-url)  
**Figure 12.** Schematic overview over the productive and nonproductive light driven photoredox processes within the context of [(NN)Rh(Cp*)X]$^{2+}$ catalyzed NAD(P)H formation. Rate constants for the electron transfer from the excited chromophore to the Rh catalyst or the oxidized cofactor are expressed as $k_1$ and $k_2$. 

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As these dimeric species and the bioactive NADH all show a characteristic UV absorbance band at 340 nm, unambiguous identification of the biologically consumable product is not possible using only UV/Vis spectroscopy [112].

Since the quantity of bioinactive reduced NAD species is commonly small, it can be rationalized that the electron transfer from a highly energy rich excited chromophore to the catalyst is faster than to the oxidized nicotinamide, probably due to the easier reduction of the Rh^{III} center. Although the light driven redox reactions of Figure 12 are depicted to proceed via the excited chromophore, they could also take place via the reduced photosensitizer due to a reductive quenching of the excited state by a suitable electron donor.

However, the immanent problem of the direct one electron reduction of the nicotinamide cofactors using intermolecular homogenous photocatalytic systems might be solved by oligonuclear coordination compounds. These could exhibit, due to their molecular preorganisation of the chromophore and catalyst, much faster collision independent electron transfers onto the latter, which would consequently decrease the amount of bioinactive NAD(P) dimers, even at low catalyst concentrations. An example of these catalysts is depicted in Figure 13, where a photoredox active ruthenium polypyridine chromophore is linked to a [(NN)Rh(Cp*)Cl] catalyst via a tetrapyridophenazine bridging ligand [99]. The molecular catalyst exhibits suitable redox potentials for intramolecular electron transfers [113].

Figure 12 also shows that reduced nicotinamide cofactors are exposed to oxidative decomposition during a photocatalytic process since they are stronger reductants than the commonly used tertiary amines such as triethylamine or triethanolamine [114]. Ishitani and coworkers successfully used artificial NADH derivatives (BNAH = N-benzyl nicotinamide) as sacrificial electron donors for the photocatalytic reduction of CO2. They observed that under the utilized conditions, BNAH acts as a simple one electron donor. Therefore, they hypothesized that the dimeric species (BNA)2, despite its higher reduction potential in comparison to the monomeric BNAH, represents the endpoint of the photooxidation pathway. This was ascribed to fast back electron transfers as a consequence of the rather slow decomposition kinetics of (BNA)2 [23,24,115].

![Figure 13. Molecular structure of a heterodinuclear photocatalyst for the generation of NAD(P)H [99].](image)

In contrast to the findings of the Ishitani group, where BNAH and (BNA)2 act as competing excited state quenchers during the photocatalysis, several other reports indicate that the dimeric species can also serve as an effective electron donor. Irradiation of C_{60} and C_{70} fullerenes led to the generation of the reduced carbon allotropes with quantum yields exceeding unity, showing the ability of (BNA)2 to act as a two electron reductant with the concomitant formation of the oxidized BNA^{+} [116–118]. Even the generation of a two electron reduced tetranuclear Ru complex under visible light irradiation has been reported using the dimer as a sacrificial electron donor [119].

Since enzyme catalyzed reactions are highly interesting due to aspects such as stereoselectivity or high substrate conversion rates at low temperatures, the photocatalytic production of NAD(P)H is often coupled to a biocatalytic dark reaction, as schematically drawn in Figure 11. Interestingly,
simple control experiments almost always reveal that the highly effective substrate conversion is only possible in the presence of the enzyme. This indicates that the catalytically active [(NN)Rh(Cp*)H]⁺ species exhibits high chemoselectivity for NAD(P)H generation over ketone [93,94] or CO₂ [95,97,104,106] reduction under the applied conditions, although the latter substrates are added in high excess. In terms of enzymatic stereoselective alcohol formation, this chemoselectivity is highly beneficial, as shown in Figure 14. The achiral rhodium catalyst is not able to produce significant amounts of the racemic product mixture and therefore does not decrease the enantiomeric excess of the overall system. However, the lack of product formation in the absence of an enzyme in the case of the CO₂ reduction shows that direct carbon-based solar fuel production using [(NN)Rh(Cp*)X]ⁿ⁺ catalysts could be accompanied by significant obstacles.

**Figure 14.** General reaction scheme for the photobiocatalytic enantioselective alcohol formation using [(NN)Rh(Cp*)L]ⁿ⁺ catalysts for the regeneration of NAD(P)H (E = enzyme).

### 3.4. Hypothesis of an Overall Reaction Scheme for the Competing Generation of Molecular Hydrogen and Reduced Nicotinamide Cofactors: Application of [(bpy)Rh(Cp*)X]ⁿ⁺ as a Catalytic Center for Selective Homogeneous Photocatalysis

As described above, [(NN)Rh(Cp*)X]ⁿ⁺ catalysts exhibit different reaction rates for the reduction of various substrates, which finally leads to a high chemoselectivity under appropriate conditions. This behavior apparently has to be correlated with the underlying intertwined catalytic cycles and the energies of important intermediates and transition states, which allows the rhodium catalyst to efficiently discriminate between various reaction partners by virtue of thermodynamic reasons.

A lot of effort has been focused on elucidating the catalytic mechanisms of [(NN)Rh(Cp*)X]ⁿ⁺ complexes in the presence of several substrates. The following discussion intends to create a coherent picture of the intricate mechanisms of hydrogen and NADH production from the available literature. Special attention will be given to slightly basic conditions, since the TONs for NAD⁺ reduction have been observed to be higher by a factor of 100 in contrast to the TONs for hydrogen production using a heterobimetallic photocatalyst [99]. Based on other literature reports, this finding could be anticipated, since for the photocatalytic formation of hydrogen, an optimal pH value of 3.6 has been found [37]. Although under electrochemical conditions the lowering of the pH value was reported to increase the rate of hydrogen generation [34], for the formation of NADH, a maximum of catalytic activity was observed at neutral pH conditions [56].

As depicted in Figure 15, the different proposed mechanistic cycles for H₂ formation and selective NAD⁺ reduction are unified in one scheme, together with some additional possible intermediates, which could also represent transition states on the road to the actual involved species. The coloration of Figure 15 is aimed to discriminate reaction steps and intermediates that refer to both catalytic cycles (black), the NAD(P)H formation (blue), and to the evolution of hydrogen (red).
where the preformation of $S_4$, product as described above. However, as $S_5$ possibly Cp* bound hydrogen in $H$ been obtained [61,62]. Therefore, hydrogen evolution might proceed via a metal assisted intramolecular thermodynamically favourable [62] and crystal structures of the 18 valence electron species simple acid base reaction.

force such as a very low pH value or an alternative pathway of hydrogen evolution via a valence electron count of 16, $S_7$ the coordination sphere yielding species $S_3$ lower formation of species $S_4$ from seven to 10, the reversibility of the Rh $\text{III}/\text{I}$ couple increases, which could be ascribed to the lower formation of species $S_4$ [51].

Direct hydrogen evolution from species $S_4$ could be achieved by an acid base reaction out of the coordination sphere yielding species $S_7$. As the latter is coordinatively unsaturated and exhibits a valence electron count of 16, $S_7$ might be relatively high in energy, and either a powerful driving force such as a very low pH value or an alternative pathway of hydrogen evolution via $S_6$ occurs, where the preformation of $S_1$, as well as the rearomatization of the Cp* ligand, could facilitate the simple acid base reaction.

Although these reactions might occur, the transformation of $S_4$ to $S_5$ has been reported to be thermodynamically favourable [62] and crystal structures of the 18 valence electron species $S_8$ have been obtained [61,62]. Therefore, hydrogen evolution might proceed via a metal assisted intramolecular $H_2$ elimination from either $S_9$ or via the intermediate $S_{10}$, which could facilitate the release of the product as described above. However, as $S_4$ and $S_5$ should stay in equilibrium with each other, it might

![Figure 15](image-url)
be possible that hydrogen evolution only occurs via the repopulation of $S_4$ and an intermolecular acid base reaction.

Based on the work of Lo et al., it is clear that for the generation of the 1,4-reduced dihydronicotinamides, binding of the carbamoyl moiety via the oxygen atom to the rhodium catalyst is essential [83,84]. As for the evolution of hydrogen, the generation of NAD(P)H could proceed via $S_4$ or $S_5$, yielding either the ring slipped intermediate $S_{11}$ as proposed by Lo et al. or species $S_{12}$ as proposed by Pitman et al. [61]. The subsequent selective hydride transfer onto the bound nicotinamide could generate intermediate $S_{13}$ in both cases, which again could lose the reaction product directly via $S_7$ or ligand assisted via $S_{14}$, where the latter could represent a transition state between $S_{13}$ and $S_1$.

As $S_5$ has a lower energy content than $S_4$, the high chemoselectivity of NAD(P)H formation over H$_2$ generation at neutral (i.e., cofactor and enzyme compatible) pH values might be caused by the electronic situation of the intermediate $S_5$. Its HOMO is a Rh$^{I}$ centered d$_{z^2}$ orbital and the high electron density on the metal is stabilized by $\pi$ backbonding into the $\pi$-accepting Cp*H ligand. The preference for $S_5$ over $S_4$ therefore seems to arise from an energetically well balanced electron density distribution, although such an interplay could have also be assumed for the interaction of the electron poor Rh$^{III}$ center in $S_4$ and the electron rich $\sigma$-donating Cp* ligand.

However, the crucial difference between hydrogen evolution catalysis and nicotinamide reduction catalysis might be the disruption of the well balanced electron density distribution in $S_5$. Binding of the nicotinamide to $S_5$ yields $S_{12}$, where an electron rich Rh$^{I}$ metal is still bound to the $\pi$-accepting Cp*H ligand, indicating that the distribution of the electron density is still efficiently ensured. On the contrary, the interaction of a proton with the d$_{z^2}$ HOMO of $S_5$ would lead to the formation of $S_9$, where a formally electron poor Rh$^{III}$ center is bound to the $\pi$-accepting Cp*H ligand. Due to the possibly much less effective interplay of mutual stabilizing effects in $S_9$ compared to $S_{12}$, the energy barrier for the formation of the latter would be lower, which could explain the different observed reaction rates for hydrogen production and NAD(P)H formation at neutral pH values [99].

Finally it should be noted that the hypothesized reasons for the observed chemoselectivity of [(NN)Rh(Cp*)X]$^{n+}$ catalysts are based on literature reports and would need further theoretical calculations as support. Moreover, the low substrate (i.e., H$^+$ or H$_3$O$^+$) concentration for the hydrogen evolution catalysis at pH 7 in comparison to commonly used NAD(P)$^+$ concentrations, which are generally in the $\mu$M to mM range, might additionally play a significant role in the observed reaction rate differences.

4. Conclusions

Our aim of this feature article was to summarize the broad and interesting chemistry of [(NN)Rh(Cp*)X]$^{n+}$ catalysts, as well as to raise the readers’ awareness for general concepts and examples of chemoselective catalysis within homogeneous artificial systems. By comparing the results of different catalytic approaches for the reductive transformation of similar substrates and by providing a mechanistic overview over the mostly independently discussed but intertwined catalytic cycles of hydrogen evolution catalysis and NAD(P)H formation, we hope to contribute to a fruitful development of these multifunctional rhodium-based catalysts in order to create pathways for the effective solar driven generation of sustainable energy carriers.

Due to the high selectivity of the catalytically active Rh species towards NAD(P)H formation at neutral pH conditions, the practical application of these photocatalytic systems could be possible by their integration into biotechnological processes. Within this context, the availability of “cheap” electrons for the regeneration of the oxidized photosensitizer is crucial. Herein the immobilization of fully active molecular systems at suitable electrode surfaces could be one of the methods employed to achieve this critical prerequisite.

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