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Redox and photocatalytic properties of a Ni^{II} complex with a macrocyclic biquinazoline (Mabiq) ligand†‡

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We present a late, first row transition metal photosensitizer that promotes photocatalytic C–C bond formation. The title compound, [Ni(Mabiq)]OTf, as well as its one-electron reduced form, Ni(Mabiq), were synthesized and molecular structures of both were obtained. The electronic structure of the reduced complex additionally was characterized by spectroscopic and DFT computational methods. Notably, [Ni^{II}(Mabiq)]OTf is photoactive: reduction of the compound was achieved photochemically upon irradiation at $\lambda = 457$ nm and reductive quenching by NEt₃. The performance of [Ni(Mabiq)]OTf as a photoredox catalyst was examined in the cyclization of a bromoalkyl-substituted indole. In this reaction, the first-row transition metal compound is comparable if not superior to [Ru(bpy)₃]²⁺ in terms of efficiency (turnover number) and chemoselectivity. Studies using a series of sacrificial donor amines indicate that the excited state redox potential of [Ni(Mabiq)]^{•+} is ≥ 1.25 V vs. SCE. This value is similar to the excited state potential of commonly employed noble metal based photocatalysts. The Ni-Mabiq compound thus provides a rare example of an earth-abundant photoredox catalyst.

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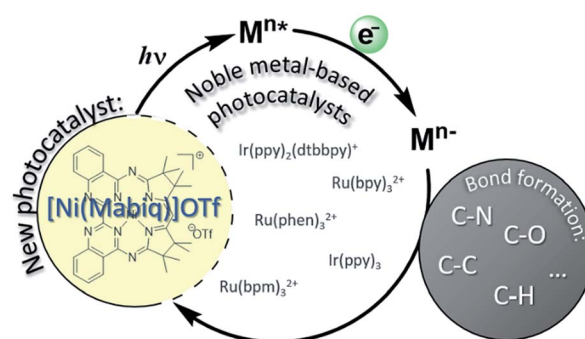
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Introduction

Photoredox catalysis offers nascent opportunities to shift conventional chemical production methods to light-driven processes.¹ The design of new photoactive metal compounds is key to the development of new catalytic transformations. Chromophores can effect a wide array of functional group transformations and C–C bond formation reactions *via* the generation of active radical species, originating from excited state electron transfer processes (Scheme 1).² With very few exceptions, photoredox catalysis relies on the use of noble metal containing photosensitizers, mainly Ru- or Ir-polypyridyl complexes;³ other heavy metal complexes (*e.g.* Os^{II}, Re^I, Mo⁰, W⁰) also have occasionally been employed.^{4,5} [Ru(bpy)₃]²⁺ – with its long lived, charge separated excited state ([Ru^{III}(bpy^{•-})(bpy)₂]^{2+*}) – is the classic and universal photocatalyst for a plethora of applications.²

The use of less expensive, more abundant, late first row transition metal alternatives is generally precluded by their

inherently short excited-state lifetimes. In the first row, nickel complexes have been used in photocatalytic cross coupling reactions,⁶ though the use of an added photosensitizer (commonly Ir) still is required in all of these tandem systems. A class of Ni^{II} ligand-to-ligand charge transfer complexes recently were shown to possess advantageous properties as photosensitizers, but applications have not yet been demonstrated.⁷ In fact, reports describing catalytic applications or reactivity of systems using only non-noble metal photosensitizers are exceedingly scarce.^{8–10}



Scheme 1 Photoredox catalysis commonly relies on noble metal complexes for organic transformations, initiated by single electron transfer upon excitation of Ru or Ir compounds. The [Ni(Mabiq)]OTf catalyst offers an earth-abundant photosensitizer.

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† Dedicated to the memory of Inge Grübel.

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We now report a Ni^{II} complex with a macrocyclic biquinazoline (Mabiq) ligand,^{11,12} $[\text{Ni}(\text{Mabiq})]\text{OTf}$ (**1**), providing a rare example of a non-noble metal based photosensitizer. The divalent complex **1** can be photochemically reduced and the electronic structure of the product, $[\text{Ni}(\text{Mabiq})]$ (**2**), is described herein. Using a series of synthesized sacrificial donor molecules, the photo-excited state redox potential of **1** was assessed and compared to common noble-metal photocatalysts. We demonstrate the photoredox catalytic ability of **1** in a radical-based cyclization of a bromoalkyl-substituted indole. The reaction relies solely on the Ni-Mabiq photocatalyst, without the need for an additional noble metal photosensitizer.

Results and discussion

The yellow diamagnetic **1** (Scheme 2) was readily prepared by complexation of the ligand with $\text{Ni}(\text{OTf})_2$ in ethanol solution (see ESI† for further details). The Ni ion adopts the expected square planar geometry in the solid state (Fig. S1†). The electronic spectrum of **1** in DCM exhibits intense absorption bands ($\epsilon \approx 10^4 \text{ M}^{-1} \text{ cm}^{-1}$) in the visible light region ($\lambda_{\text{max}} = 414, 435$, and 457 nm ; $\epsilon = 13.6, 16.4, 22.3 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$; Fig. 2 inset). Similar features were observed in the spectra of other M-Mabiq complexes, notably in the spectrum of $\text{Zn}(\text{Mabiq})\text{Cl}$.^{12a} The Zn^{II} complex displayed two strong absorption bands at 471 and 502 nm that were assigned as Mabiq $\pi \rightarrow \pi^*$ transitions. However, d-d or metal-to-ligand charge transfer (MLCT) processes may additionally contribute to the corresponding absorptions of **1**. The cyclic voltammogram of **1** in MeCN (Fig. S9†) exhibits a reversible, formally $\text{Ni}^{\text{II/I}}$ redox couple at -1.05 V vs. $\text{Fc}^{+/0}$ (Fc = ferrocene; $\text{Fc}^{+/0} = 0.4 \text{ V}$ vs. SCE). Additional, seemingly reversible, reductive processes appear at potentials $< -1.5 \text{ V}$.

The one-electron reduced $\text{Ni}(\text{Mabiq})$ (**2**) was subsequently generated from **1** using CoCp_2 as the reductant (Scheme 2). The molecular structure of **2** (Fig. 1) reveals shorter Ni-N bond distances ($\text{Ni}-\text{N}_{\text{avg}} = 1.874 \text{ \AA}$ vs. $\text{Ni}-\text{N}_{\text{avg}} = 1.882 \text{ \AA}$ in **1**), as well as the hallmark changes in the diketimate C-N bonds that signify reduction of the Mabiq ligand (Table S8†). The $S = 1/2$ ground state of the complex was verified by EPR spectroscopy. The spectrum is consistent with a ligand-centered radical, with $g_{\text{iso}} = 1.995$ (Fig. S4†). Low energy features at $\lambda_{\text{max}} = 641, 711$ and 801 nm ($1.4, 3.5, 5.4 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$) are apparent in the electronic spectrum of **2** in THF (Fig. 2 inset), accounting for the

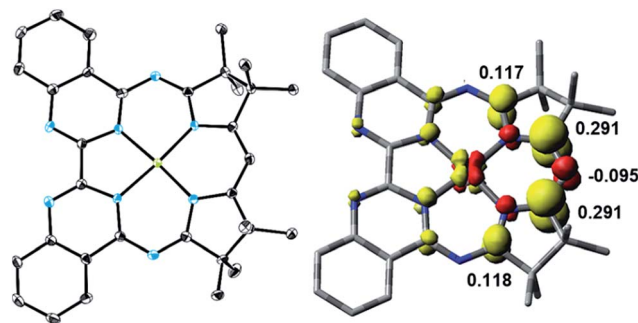


Fig. 1 Left: Molecular structure of **2** (50% probability ellipsoids; hydrogen atoms omitted for clarity). Right: DFT-derived (B3LYP) spin density plot for **2** based on Löwdin population analysis (isosurface value = ± 0.005).

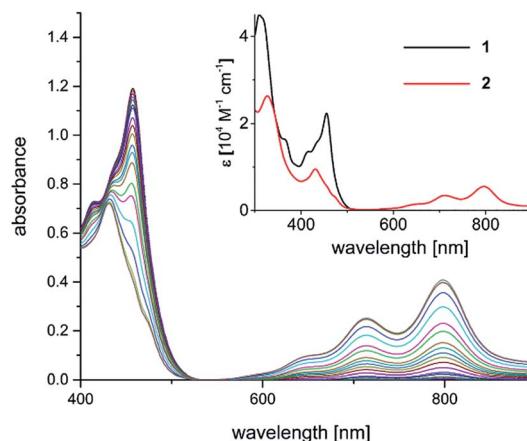
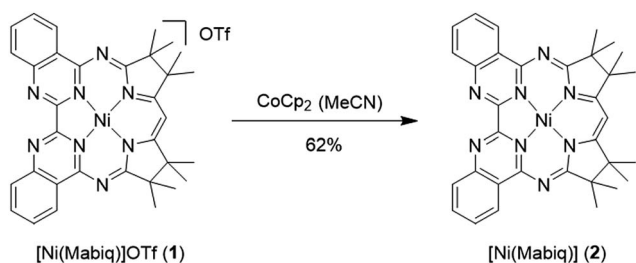


Fig. 2 Spectral evolution during photoconversion of **1** to **2** [c (**1**) = 0.05 mM ; c (NEt_3) = 1.4 M ; $\lambda = 457 \text{ nm}$, DMF]. Inset: electronic spectra of **1** (black trace; CH_2Cl_2) and **2** (red trace; THF).

vibrant green color of the complex in solution. The spectrum again closely resembles that of the one-electron reduced Zn complex, $\text{Zn}^{\text{II}}(\text{Mabiq}^{\cdot-})$.^{12a} The spectroscopic data thus point to ligand-centered reduction of **1**, such that the electronic structure of **2** corresponds to $\text{Ni}^{\text{II}}(\text{Mabiq}^{\cdot-})$. Indeed, DFT calculations (B3LYP) on **2** further support this conclusion.

The DFT-derived (B3LYP) spin density plot (Fig. 1) describes a diamagnetic $d^8 \text{ Ni}^{\text{II}}$ center with an unpaired electron localized primarily on the diketimate unit of the Mabiq ligand. Four doubly occupied d-orbitals can be identified, while the SOMO possesses only *ca.* 4% d-orbital character and otherwise depicts a ligand π^* orbital (Fig. S5†). The latter molecular orbital is antibonding with respect to the diketimate C-N p-orbital interactions, which explains the lengthening of these bonds in the structure of **2**. It is noteworthy that ligand-centered reduction appears to prevail across the series of metal-Mabiq compounds we have examined thus far.¹²

The well-behaved redox chemistry of $[\text{Ni}(\text{Mabiq})]\text{OTf}$ (**1**), its high absorbance in the visible region and its relatively high reduction potential warranted a study of its photoredox properties. As mentioned above, the compound exhibits a strong



Scheme 2 Reaction of **1** with CoCp_2 (Cp = cyclopentadienyl) yields the one-electron reduced **2**.



multi-structured absorption band with a maximum at $\lambda = 457$ nm ($\epsilon = 22\,300$ M⁻¹ cm⁻¹), which invites excitation with a visible light source and quenching studies with a suitable reductant. Gratifyingly, it was indeed found that irradiation of a DMF solution of **1** at $\lambda = 457$ nm, in the presence of NEt₃ (7.5 mM to 1.4 M), leads to a color change from yellow to green and to the formation of complex **2**.

The formation of the reduced compound was complete after 15 minutes (using 1.4 M NEt₃; c (**1**) = 0.05 mM), as verified spectroscopically (Fig. 2). The photoconversion of **1** to **2** occurs on a much faster timescale in MeCN/THF or DMF/THF mixtures (the solvent combination solubilizes both forms), under identical conditions. The quantum yield for the photoconversion is $\approx 10^{-4}$ (THF : DMF 4 : 1), and correlates with the Et₃N concentration (14–56 mM; Table S3 and Fig. S29†). From the data, the lifetime of the excited state form that reacts with the Et₃N can be estimated as $\approx 1 \times 10^{-8}$ s (see ESI† for details). Steady-state emission spectra recorded at ambient temperature and at 77 K did not reveal any luminescence. Thus, we currently cannot comment in detail on the nature of the excited state processes involved in the photo-reduction of **1**. If one takes the longest wavelength absorption [$\lambda \approx 510$ nm, $E_0 \leq 235$ kJ mol⁻¹ (2.4 V)] of compound **1** to estimate the redox potential of photoexcited complex **1**^{*}, a value of $\leq +1.35$ V (vs. Fc^{+/0}) is obtained.¹³

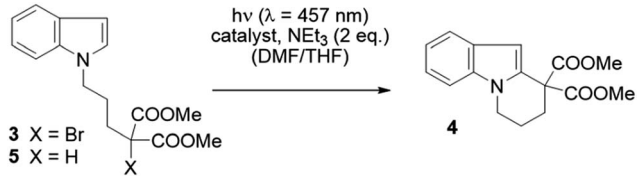
TDDFT (B3LYP) computational studies provide some insight into the nature of the absorptions in the visible region. The calculated transitions correlate well with the experimentally obtained absorbance spectrum of **1** (Fig. S6†). The absorptions at 400–500 nm include a prominent LL/CT transition that corresponds to the HOMO to LUMO transition. The HOMO is localized on the bipyrimidine moiety of the MabiQ ligand, while the LUMO is a diketiminate based π^* orbital (Fig. S7†). Other, less intense, transitions in the vicinity possess d–d (Ni d_{z²} → Ni

d_{x²-y²}) and MLCT (Ni d_{z²} → L π^*) character. These states may contribute to the unique photochemical properties of **1**. However, a detailed investigation regarding the photochemistry and excited state kinetics of this compound is warranted, and will be the subject of future investigations.

We examined whether the photochemical properties of **1** might render it a suitable photoredox catalyst for C–C bond formation reactions. The radical cyclization of the *N*-(ω -bromoalkyl)-substituted indole **3** was chosen as a test reaction.¹⁴ The reaction had been previously studied by the Stephenson group and was found to produce mainly product **4** by C–C bond formation under optimized conditions.¹⁵ Under non-optimized conditions, hydro-de-bromination was a competing side reaction and varying product ratios of **4** and **5** were observed. Optimal conditions were reported to include the use of [Ru(bpy)₃]Cl₂ as the catalyst (1 mol%) and NEt₃ (2 equiv.) in DMF solution and gave product **4** in 60% yield.¹⁵

Given the limited solubility of **2** in DMF, the reaction was initially attempted in a DMF/THF mixture (v/v = 1/2) with 2 mol% of catalyst **1** and 2 equiv. NEt₃ as the quencher (Table 1, entry 1).¹⁶ We were pleased to find that the desired cyclization proceeded smoothly and delivered with high chemoselectivity the desired product **4**. The inseparable hydro-de-brominated by-product **5** was detectable in minor quantities but the ratio of products was 95/5 in favor of cyclization product **4**. When increasing the relative volume of THF in the solvent mixture both conversion and yield improved slightly (entry 2). The selectivity towards the desired reaction was high with a yield of 86% at 94% conversion, *i.e.* 91% yield based on conversion. For comparison, the Ru^{II} complex [Ru(bpy)₃](PF₆)₂ was employed under identical conditions (entry 3). Although the ratio 4/5 was identical with this catalyst, the reaction suffered from a lower conversion and a lower chemoselectivity (63% yield based on conversion). Similar observations were made when the catalyst

Table 1 Photoredox-catalyzed cyclization of bromide **3** to tricyclic product **4** and reduction to hydro-de-brominated product **5**; influence of the catalyst and the reaction parameters on the yield and chemoselectivity

						
Entry ^a	Catalyst ^a	mol%	DMF/THF [v/v]	Conv. ^b [%]	Yield ^c [%]	4/5 ^d
1	1	2	1/2	93	84	95/5
2	1	2	1/4	94	86	95/5
3	[Ru(bpy) ₃](PF ₆) ₂	2	1/4	73	46	95/5
4	1	1	1/4	95	84	95/5
5	[Ru(bpy) ₃](PF ₆) ₂	1	1/4	59	23	95/5
6	1 ^e	2	1/4	n.d.	—	—
7	— ^f	—	1/4	12	12	55/45
8	1 ^g	2	1/4	<5	<5	—

^a All reactions were performed on a scale of 0.08 mmol ($c = 25$ mM) with a LED lamp (3 W power output) as light source. Irradiation time: 13 h. ^b The conversion was calculated from recovered starting material. ^c Total yield of isolated products **4** and **5**. ^d Ratio of cyclized to hydro-de-brominated product as determined by ¹H-NMR. ^e Attempted reaction without irradiation. ^f No catalyst was added. ^g No NEt₃ was added. n.d. = not detected.



loading was further decreased to 1 mol%: while the performance of Ni^{II} catalyst **1** remained unchanged (entry 4) the reaction with the Ru^{II} catalyst was sluggish and a decrease in yield was observed (entry 5).

The above reaction is induced by visible light as no conversion occurs without irradiation (entry 6). In the absence of the Ni^{II} catalyst,¹⁷ only 12% of a product mixture was obtained, which was composed of the cyclized product **4** and the reduced product **5** in a 55/45 ratio (entry 7). In the absence of the reductant, no reaction was observed (entry 8). The free HMabiq ligand is not photocatalytically active. The quantum yield for the [Ni(Mabiq)]OTf catalysed cyclization reaction was determined to be $\Phi = 0.006$.

Mechanistically, it is suggested that the photoreduction of complex **1** by NEt₃ (reductive quenching cycle)¹ generates complex **2**, which may engage in SET to the bromide **3** (Scheme 3). Complex **2** was shown to be competent to reduce **3**. We note that modification of the Ni-Mabiq complex during the cyclization reaction was not observed, as verified by ESI-MS and ¹H-NMR (Fig. S8 and S41†).

To experimentally verify the estimated photoexcited state redox potential of **1**^{*}, we synthesized a series of amines with oxidation potentials in the range of 0.78–1.59 V (vs. SCE; Tables 2 and S1†), as determined by CV. The amines were employed as sacrificial donors in the cyclization reaction. Excellent yields of **4** were obtained using the donor molecules with oxidation potentials up to 1.25 V (Table 2, amines **6a–6c**), whereas a drastic decrease in yield was observed using those with higher redox potentials. Only 40% and 20% product yields were obtained with **6d** ($E_{\text{ox}} = 1.41$ V vs. SCE) and **6e** ($E_{\text{ox}} = 1.59$ V vs. SCE), respectively. The product yield in the control reaction using **6e** in the absence of photocatalyst **1** was 7%, a result that is comparable to that obtained using NEt₃ (Table 1, entry 7). The results confirm that the excited state redox potential of **1**^{*} is at least 1.25 V (vs. SCE). Thus, our new Ni-Mabiq complex is a more powerful oxidant than [Ru(bpy)₃]²⁺ ($E_{1/2}$ [Ru^{2+*/}Ru⁺] 0.78 V vs. SCE),¹⁸ and comparable to [Ir(dF(CF₃)ppy)₂(dtbbpy)]⁺ ($E_{1/2}$ [Ir^{3+*/}Ir²⁺] 1.21 V vs. SCE).¹⁹

We additionally generated a sterically hindered amine, N(CH₂Me)₂Cy₂ (**6f**), to assess whether coordination of the sacrificial donor molecules influences the reactivity of **1**. In contrast to the bpy- and phen-based photosensitizers, the Ni-Mabiq complexes are coordinatively unsaturated and intra-molecular electron transfer from a coordinated amine to the

Table 2 Comparison of the oxidation potentials for different sacrificial donors **6** with the yields obtained for the catalytic reaction of **3** to **4** and **5**

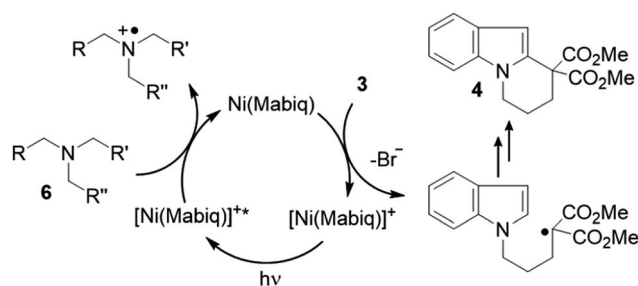
Amine ^a	E_{ox} (vs. SCE)	Yield ^b [%]
Et ₃ N (6a)	0.83	84
6b	1.05	97
6c	1.25	95
6d	1.41	40
6e	1.59	20 ^c
6f	0.78	84

^a All reactions were performed on a scale of 0.08 mmol ($c = 25$ mM) with a 457 nm LED lamp (3 W power output) as the light source. Irradiation time: 13 h. ^b Total yield of isolated products **4** and **5**, with a 95 : 5 ratio of cyclized to hydro-de-brominated product as determined by ¹H-NMR. ^c Average of two runs. $\text{Fc}^{+/0} = 0.4$ V vs. SCE. Cy = cyclohexyl.

divalent metal center could potentially occur. The molecular structure of **6f** (Fig. S3†) suggests that coordination of this amine group to the Ni center is unlikely; the cyclohexyl and mesityl groups encapsulate the nitrogen atom rendering the lone pair inaccessible. Whereas noticeable changes in the absorption spectrum of **1** were observed upon addition of Et₃N (400 equiv.) to a solution of the complex in THF/DMF, the addition of **6f** has no effect (Fig. S30 and S31†). However, the yield of the catalytic reaction using **6f** as sacrificial donor was found to be 84%, a value that is comparable to the one obtained using Et₃N (**6a**) and **6b–6e** as sacrificial donors.

Conclusions

In summary, we have discovered a new photoredox catalyst, [Ni(Mabiq)]OTf (**1**) that is based on the earth-abundant metal nickel. The diamagnetic, bench-stable compound was readily prepared, its redox properties were studied and the one-electron reduced form Ni(Mabiq) (**2**) likewise was isolated. Further studies to elucidate the detailed photophysical properties of **1**^{*} are warranted. However, we have already demonstrated that the photoexcited complex is a strong oxidant, with the capacity to induce C–C bond formation in an initial test reaction. The Ni-Mabiq compound may offer an alternative to noble metal photosensitizers for other synthetic transformations in organic photoredox chemistry, as well as for energy conversion processes. The Mabiq ligand also features a second metal binding site that could be exploited for tandem catalysis. Thus, the macrocycle represents a new type of platform for the development of photoactive systems. With evidence of the ability of **1** to act as a photosensitizer and photoredox catalyst, the broader photocatalytic applications of our system subsequently will be investigated.



Scheme 3 Proposed catalytic cycle for the cyclization of **3** to give **4**.



Conflicts of interest

There are no conflicts to declare.

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