Electron Transfer between Hydrogen-Bonded Pyridylphenols and a Photoexcited Rhenium(I) Complex


Two pyridylphenols with intramolecular hydrogen bonds between the phenol and pyridine units were synthesized, characterized crystallographically, and investigated by cyclic voltammetry and UV-vis spectroscopy. Reductive quenching of the 3MLCT excited state of the [Re(phen)(CO)3(py)]+ complex (phen = 1,10-phenanthroline, py = pyridine) by the two pyridylphenols and two reference phenol molecules was investigated by steady-state and time-resolved luminescence spectroscopy, as well as by transient absorption spectroscopy. Stern-Volmer analysis of the luminescence quenching data provides rate constants for the bimolecular excited-state quenching reactions. H/D kinetic isotope effects (KIEs) for the pyridylphenols are on the order of 2.0, and the bimolecular quenching reactions are up to 100 times faster with the pyridylphenols than with the reference phenols. This observation is attributed to the markedly less positive oxidation potentials of the pyridylphenols with respect to the reference phenols (ca. 0.5 V), which in turn is caused by proton-coupling of the phenol oxidation process. Transient absorption spectroscopy provides unambiguous evidence for the photogeneration of phenoxyl radicals, i.e., the overall photoreaction is clearly a PCET process.

Introduction

The tyrosine Z (TyrZ) / histidine 190 (His190) pair of photosystem II is one of the best-known hydrogen-bonded phenol systems in chemistry.[1] Numerous experimental and theoretical investigations have been geared at understanding the proton-coupled electron transfer (PCET) chemistry of the TyrZ / His190 reaction couple, many of them focusing on simple artificial model compounds in which a phenol unit can form intramolecular hydrogen bonds to a nitrogen base.[2] A wide range of experimental methods has been applied, including EPR,[3] electrochemical,[4] and optical spectroscopic studies.[5] However, in most cases the phenol oxidation process involves oxidants which are in their electronic ground states, and there are comparatively few studies in which the oxidant is an electronically excited molecule.[5c, 6]

Against this background we deemed it interesting to explore the photoredox chemistry between hydrogen-bonded phenol molecules and a photoexcited rhenium(I) tricarbonyl diimine complex which is known to be a potent excited-state oxidant.[6e, 7] The molecular structures of our model systems are shown in Scheme 1. The rhenium(I) complex has a 1,10-phenanthroline (phen) and a pyridine (py) ligand in addition to the three carbonyls, the phenol reaction partners have pendant pyridine units that are connected via a –CH2– group in order to disrupt π-conjugation between the two aromatic subunits. One of the phenols contains no further substituents (PhOH-CH2-py) while the other has tert.-butyl groups at the 4- and 6-positions (tBu2PhOH-CH2-py). We anticipated that when excited to its long-lived 3MLCT state the rhenium(I) complex would be capable of inducing intermolecular electron transfer (ET) with the phenol, and this process should be accompanied by intramolecular proton transfer (PT) between the phenol and the pyridine.

The PCET chemistry of tBu2PhOH-CH2-py with various

Scheme 1. Molecular structures of the photosensitizer/quencher pairs with the two hydrogen-bonded phenol molecules of central interest to this study; hν = light excitation; MLCT = metal-to-ligand charge transfer excitation; ET = electron transfer; PT = proton transfer.
oxidants in their electronic ground states has been found previously to occur via a concerted proton-electron transfer (CPET) mechanism. Here we focus specifically on the excited-state CPET chemistry between [Re(phen)(CO)]^+ and PhOH-CH_2-py or BuPhOH-CH_2-py. As reference phenols without the possibility of forming intramolecular hydrogen bonds we used ordinary phenol (PhOH) and 2,4-di-tert.-butylphenol (BuPhOH).

Results and Discussion

The synthesis of BuPhOH-CH_2-py had been previously described but in our hands a different procedure turned out to be more convenient for obtaining the two pyridylphenols from Scheme 1. Our synthetic strategy is illustrated by Scheme 2 and begins with commercially available 2-bromophenols (1 and 3), which are methylated in order to protect the phenolic function for the subsequent reaction step. The protected phenols (2 and 4) are reacted with pyridine molecule 5 (which is accessible in one step from 2-picoline and disopropyl ketone) using a palladium catalyst. The coupling products (6 and 8) are deprotected using aqueous HBr (in the case of 6) or ethanethiol (in the case of 8) in order to obtain the final pyridylphenols (7, PhOH-CH_2-py and 9, BuPhOH-CH_2-py).

Synthesis

Figure 1 shows the crystal structure of PhOH-CH_2-py which crystallizes in the monoclinic space group P2_1/n with one molecule in the asymmetric unit. One of the Bu groups happens to be disordered by 5%. This molecule only forms an intramolecular hydrogen bond between H1 and N1. The hydrogen H1 was found in the difference Fourier density. Position and the isotropic vibration were refined freely with a distance restraint of 0.84(2) Å to O1. The O-H distance refined to 0.88(2) Å with a total O-N distance of 2.6956(16) Å with an O-H-N angle of 169(2)°. Thus, there is clear evidence for hydrogen-bonding interactions in the crystal structures of PhOH-CH_2-py and BuPhOH-CH_2-py. A structure of BuPhOH-CH_2-py had been previously published.

Intramolecular hydrogen-bonding in solution

1H NMR spectra of the two pyridylphenols in CDCl_3 exhibit sharp downfield resonances for the phenolic protons, specifically at 11.67 ppm for PhOH-CH_2-py and at 11.40 ppm for BuPhOH-CH_2-py, which is typical for intramolecularly hydrogen-bonded phenols. We conclude that intramolecular hydrogen-bonds are not only present in one of our solid state structures but also in aprotic solution.

Cyclic voltammetry

Figure 2 shows cyclic voltammograms of (a) BuPhOH, (b) BuPhOH-CH_2-py, and (c) PhOH-CH_2-py in dry CH_2Cl_2 in presence of 0.1 M TBAPF_6 electrolyte. The reversible waves at 0.0 V vs. Fc+/Fc (dashed vertical line) are due to ferrocene, which was added in small quantities for internal voltage calibration. The voltammogram of the reference phenol (a) exhibits an irreversible oxidation wave peaking at 1.05 V vs. Fc+/Fc which is typical for ordinary phenols because the O-H proton is lost to the bulk solution in the course of oxidation.

The BuPhOH-CH_2-py molecule, by contrast, exhibits a voltammogram in which the oxidative peak current near 0.5 V vs. Fc+/Fc is roughly 6 times larger than the corresponding reductive peak current. Their voltage separation is 170 mV but depends on voltage sweep rate. The voltammogram in Figure 2b is qualitatively similar to that previously reported for the same compound in CH_2CN solution. The shape of this voltammogram can be explained by the possibility of transferring the phenolic proton to the pendant pyridine base in the course of oxidation and back-transfer to the phenol unit during the subsequent reductive potential sweep. The middle between the oxidative and reductive peak currents in Figure 2b is taken as the oxidation potential of BuPhOH-CH_2-py (Table 1). Importantly, the oxidation potential of BuPhOH-CH_2-py in CH_2Cl_2 is about 0.5 V less positive than the oxidation potential of BuPhOH, a fact that has been previously noted for CH_2CN solution. It has been demonstrated that the unusually low oxidation potential of BuPhOH-CH_2-py and related hydrogen-bonded phenols is a direct manifestation of intramolecular proton transfer accompanying electrochemical phenol oxidation; hydrogen-bonding alone cannot account for the large magnitude of the oxidation potential shift.

The cyclic voltammogram of PhOH-CH_2-py in Figure 2c exhibits an irreversible oxidation wave peaking at 0.66 V vs. Fc+/Fc (Table 1). Despite the presence of an intramolecular hydrogen bond phenol oxidation is clearly irreversible in this case, possibly because of the absence of substituents at the 4- and 6-positions of the phenol. Chemical substituents in ortho- and para-
position to the phenolic function are known to enhance the stability of phenoxyl radicals.[19] By analogy to the other pyridylphenol from Scheme 1 PhOH-CH$_2$-py is oxidized at much less positive potential than the PhOH reference molecule; in this specific case the potential difference amounts to $-0.6$ V (Table 1).

Figure 2. Cyclic voltammograms of the hydrogen-bonded phenols from Scheme 1 in dry CH$_2$Cl$_2$ in presence of 0.1 M TBAPF$_6$. (a) Bu$_3$PhOH; (b) Bu$_3$PhOH-CH$_2$-py; (c) PhOH-CH$_2$-py. The reversible waves at 0.0 V vs. Fc/Fc are due to ferrocene which was added in small quantities for internal voltage calibration; the scan rate was 100 mV/s.

| Table 1. Electrochemical potentials ($E$) for oxidation of the four phenol molecules and for reduction of the photoexcited $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ complex. |
|-----------------|-----------------|
| Redox couple       | $E$ [V vs. Fc$/\text{Fc}^+$/Fc] |
| PhOH/PhOH          | 1.25 [8][9]       |
| PhOH-CH$_2$-py/PhOH-CH$_2$-py | 0.66 [9]     |
| Bu$_3$PhOH/Bu$_3$PhOH | 0.67 [9]    |
| Bu$_3$PhOH-CH$_2$-py/Bu$_3$PhOH-CH$_2$-py | 0.54 [9] |
| $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$/[Re(phen)(CO)$_3$(py)] | 0.77 [10][11] |

[a] From reference [19], converted from V vs. SCE to V vs. Fc$/\text{Fc}^+$/Fc by subtracting 0.38 V as described in ref. [11]. [b] Measured in this work, peak potentials from Figure 2, 0.1 M TBAPF$_6$ electrolyte in CH$_2$Cl$_2$. [c] From reference [9]. [d] In CH$_3$CN. The previously reported value for Bu$_3$PhOH-CH$_2$-py is 0.44 V vs. Fc$/\text{Fc}^+$/Fc in CH$_3$CN.[9] The potential of Bu$_3$PhOH is in line with the value reported in ref. [19] (0.519 V vs. NHE, addition of 0.624 V (according to ref. [19]) gives 1.14 V vs. Fc$/\text{Fc}^+$/Fc).

The electrochemistry of the $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ complex and related rhenium(I) tricarbonyl diimines was explored extensively in the past.[7, 19] In Table 1 we merely give the electrochemical potential for one-electron reduction of $^3\text{MLCT}$-excited $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ as reported in the literature.[9]

**Optical absorption**

Figure 3 shows UV-vis spectra of the four phenols and the rhenium(I) complex from Scheme 1 in CH$_2$Cl$_2$ at 25 °C. The important message from Figure 3 is that all four phenols are spectroscopically innocent at wavelengths longer than 330 nm. Between 270 and 280 nm they exhibit weak absorptions as previously reported for other phenols; in presence of covalently attached pyridine units the extinction between 270 nm and 280 nm increases because pyridine has itself weakly absorbing n-$\pi^*$ transitions occurring in this spectral range.[20] As reported previously, the $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ complex exhibits a metal-to-

ligand charge transfer (MLCT) band with maxima at 380 nm and 336 nm, while the absorption maximum at 276 nm has been attributed to phenanthroline-based electronic transitions.[19a, 19b]

The most important observation from Figure 3 is that with light of 410 nm wavelength we can selectively excite the $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ complex even in presence of large excess of any of the four phenols. Furthermore, there is no phenol absorption in the spectral range in which the rhenium(I) complex emits (450 nm – 700 nm); this is why in Figure 3 we show the entire spectral range between 250 nm and 700 nm. We note that phenol has a triplet energy ($E_T$) of 3.55 eV,[21] while the $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ complex has $E_T = 2.75$ eV,[7, 19a] hence we can a priori rule out the possibility of $^3\text{MLCT}$ excited-state quenching by triplet-triplet energy transfer from $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ to the phenols.[22]

![Figure 3. Optical absorption spectra of the four phenols and the rhenium(I) complex from Scheme 1.](image)

**Luminescence quenching experiments**

The solid trace in Figure 4a is the emission spectrum of $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ in aerated CH$_2$Cl$_2$ with 100 mM CH$_3$OH at 25 °C. The excitation wavelength was set at 410 nm. The broad and unstructured luminescence band is due to the typical $^3\text{MLCT}$ emission of rhenium(I) tricarbonyl diimines.[19a] The solid trace in Figure 4b shows the temporal evolution of the $^3\text{MLCT}$ luminescence from Figure 4a after excitation with ~10 ns laser pulses at 410 nm; detection occurred at 530 nm. The luminescence intensity decays in a single-exponential manner over more than two orders of magnitude and one extracts a $^3\text{MLCT}$ lifetime of 1.2 μs, in line with previous reports.[19a, 19b] The dashed lines in Figure 4a/4b were recorded in presence of variable concentrations (1 mM – 10 mM) of PhOH. No significant luminescence quenching is observed with PhOH, neither in intensity (Figure 4a) nor in decay time (Figure 4b).

Likewise, when using deuterated phenol (PhOD), the emission intensity stays virtually unchanged (Figure 4c) and the luminescence decays are no faster than in the absence of PhOD (Figure 4d).

We conclude that the ordinary phenol is unable to quench the $^3\text{MLCT}$ excited state of $[\text{Re(phen})(\text{CO})_3\text{(py)}]^+$ under the experimental conditions chosen here.
Figure 5 shows the results of an analogous series of experiments performed with PhOH-CH$_2$-py. From Figure 5a we learn that the emission intensity of [Re(phen)(CO)$_3$(py)]$^+$ is significantly quenched in the presence of 1 mM to 10 mM PhOH-CH$_2$-py (dotted traces compared to solid trace). Similarly, the luminescence decays are strongly dependent on the PhOH-CH$_2$-py concentration (Figure 5b). When using deuterated PhOD-CH$_2$-py the luminescence decays (Figure 5d) are noticeably slower than for undeuterated PhOH-CH$_2$-py at equal concentration (Figure 5b). Likewise, in the luminescence intensity data of Figure 5c quenching at a given phenol concentration is noticeably weaker than for the undeuterated quencher in Figure 5a. Thus, there appears to be a significant H/D kinetic isotope effect (KIE).

The H/D KIE mentioned above shows up directly in the Stern-Volmer constants: From the intensity data in Figure 6a 

\[ k_{SV, H} = (5.9 \pm 0.1) \times 10^8 \text{ M}^{-1} \text{s}^{-1} \text{ for PhOH-CH}_2\text{-py, } k_{SV, D} = (2.8 \pm 0.1) \times 10^8 \text{ M}^{-1} \text{s}^{-1} \text{ for PhOD-CH}_2\text{-py.} \]

Based on the "MLCT lifetime of [Re(phen)(CO)$_3$(py)]$^+$" (1.2 $\mu$s in aerated CH$_2$Cl$_2$, see above) we calculate rate constants for bimolecular excited-state quenching of $k_{Q, H} = (5.9 \pm 0.1) \times 10^8$ M$^{-1}$ s$^{-1}$ for PhOH-CH$_2$-py and $k_{Q, D} = (2.8 \pm 0.1) \times 10^8$ M$^{-1}$ s$^{-1}$ for PhOD-CH$_2$-py (fifth and sixth column of Table 2). The H/D KIE is the ratio between $k_{Q, H}$ and $k_{Q, D}$ and amounts to 2.1±0.1 (last column of Table 2). From the luminescence intensity data in Figure 6a one extracts $k_{Q, H} = (5.9 \pm 0.1) \times 10^8$ M$^{-1}$ s$^{-1}$ for PhOH-CH$_2$-py and $k_{Q, D} = (3.0 \pm 0.1) \times 10^8$ M$^{-1}$ s$^{-1}$ for PhOD-CH$_2$-py, yielding a value of KIE (2.0±0.1) in accordance with the lifetime data.

Figure 6a is a Stern-Volmer plot based on the luminescence intensity data from Figures 4/5, and Figure 6b is a Stern-Volmer plot based on the luminescence lifetime data from Figures 4/5.[23] The open circles in Figure 6a/b represent data obtained using PhOH-CH$_2$-py, the open squares represent data obtained using the deuterated analogue PhOD-CH$_2$-py. Linear regression fits yield the Stern-Volmer constants ($k_{SV}$) given in the third (ordinary phenols) and fourth column (deuterated phenols) of Table 2.
Table 2. Results from luminescence quenching experiments

<table>
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<tr>
<th>phenol</th>
<th>Exp. type</th>
<th>$K_{SV,H}$ [M$^{-1}$]</th>
<th>$K_{SV,D}$ [M$^{-1}$]</th>
<th>$k_{Q,H}$ [M$^{-1}$ s$^{-1}$]</th>
<th>$k_{Q,D}$ [M$^{-1}$ s$^{-1}$]</th>
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<tr>
<td>PhOH/D</td>
<td>intensity</td>
<td>3.4±1.2</td>
<td>0.5±3.7</td>
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<td>lifetime</td>
<td>8.4±0.4</td>
<td>13.8±1.0</td>
<td>(7.1±0.3) 10$^6$</td>
<td>(11.7±0.8) 10$^6$</td>
<td>N/A</td>
</tr>
<tr>
<td>PhOH/D-CH$_2$-py</td>
<td>intensity</td>
<td>691±7</td>
<td>355±5</td>
<td>(5.9±0.1) 10$^6$</td>
<td>(3.0±0.1) 10$^6$</td>
<td>2.0±0.1</td>
</tr>
<tr>
<td></td>
<td>lifetime</td>
<td>701±3</td>
<td>334±6</td>
<td>(5.9±0.1) 10$^6$</td>
<td>(2.8±0.1) 10$^6$</td>
<td>2.1±0.1</td>
</tr>
<tr>
<td>Bu$_2$PhOH/D</td>
<td>intensity</td>
<td>391±8</td>
<td>332±8</td>
<td>(3.3±0.1) 10$^6$</td>
<td>(2.8±0.1) 10$^6$</td>
<td>1.2±0.1</td>
</tr>
<tr>
<td></td>
<td>lifetime</td>
<td>437±11</td>
<td>309±10</td>
<td>(3.7±0.1) 10$^6$</td>
<td>(2.6±0.1) 10$^6$</td>
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<tr>
<td>Bu$_2$PhOH/D-CH$_2$-py</td>
<td>intensity</td>
<td>1572±9</td>
<td>892±5</td>
<td>(13.3±0.1) 10$^8$</td>
<td>(7.6±0.1) 10$^6$</td>
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<tr>
<td></td>
<td>lifetime</td>
<td>1648±6</td>
<td>773±8</td>
<td>(14.0±0.1) 10$^8$</td>
<td>(6.8±0.1) 10$^8$</td>
<td>2.1±0.1</td>
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Ster-Stern-Volmer constants obtained from emission intensity or lifetime experiments with normal (k$_{SV,H}$) and deuterated phenols (k$_{SV,D}$). Rate constants for bimolecular excited-state quenching with normal (k$_{Q,H}$) and deuterated phenols (k$_{Q,D}$); calculated from k$_{SV,H}$ and k$_{SV,D}$ values using the lifetime of $^3$MLCT-excited [Re(phen)(CO)$_3$(py)]$^-$ in aerated CH$_2$Cl$_2$ with 100 mM CH$_3$OH (1177 ns). H/D kinetic isotope effect (KIE) calculated from the ratio of k$_{Q,H}$ and k$_{Q,D}$.

The grey filled circles in Figures 6a/6b represent data obtained for the undeuterated reference phenol PhOH. One extracts K$_{SV,H}$ = 3.4±1.2 M$^{-1}$ from the intensity data in Figure 6a and K$_{SV,H}$ = 8.4±0.4 M$^{-1}$ from the lifetime data in Figure 6b, which in turn yields k$_{Q,H}$ values on the order of 10$^6$ M$^{-1}$ s$^{-1}$. This order of magnitude of k$_{O,H}$ underscores what in principle is already evident from the raw data in Figure 4: Reductive excited-state quenching by PhOH is not kinetically competitive with (radiative and nonradiative) deactivation processes of photoexcited [Re(phen)(CO)$_3$(py)]$^-$. Thus, even though k$_{Q,D}$ values for deuterated phenol are technically available from the data in Figures 4c/4d, calculation of an H/D KIE is not meaningful in the case of the simple reference phenol.

Figures 6c/6d show Stern-Volmer plots based on [Re(phen)(CO)$_3$(py)]$^-$ luminescence quenching experiments with Bu$_2$PhOH (grey filled circles), Bu$_2$PhOD (grey filled squares), Bu$_2$PhOH-CH$_2$-py (open circles), and Bu$_2$PhOD-CH$_2$-py (open squares). The respective raw data are shown in Figures S1 and S2 of the Supporting Information. The bimolecular rate constants for excited-state quenching with Bu$_2$PhOH and its deuterated congener extracted from this data are all around 3±10$^6$ M$^{-1}$ s$^{-1}$ (Table 2), for Bu$_2$PhOH/D-CH$_2$-py the k$_{Q}$ values are about a factor of 3 larger. H/D KIES range from close to 1.0 for Bu$_2$PhOH to ~2.0 for Bu$_2$PhOH-CH$_2$-py.

All luminescence quenching experiments were performed in presence of 100 mM CH$_3$OH / CD$_2$OD to ensure deuteration of the phenol molecules for the KIE studies. Use of pure CH$_3$Cl or CD$_2$Cl$_2$ leads to markedly lower KIES, presumably due to D/H exchange of the deuterated phenols when brought into contact with glassware / cuvettes.

Figure 7 shows a plot of the (average) k$_{Q}$ values versus standard Gibbs free energy of reaction ($\Delta G_{ET}$) associated with electron transfer from the individual phenols to $^3$MLCT-excited [Re(phen)(CO)$_3$(py)]$^-$. The free energies were calculated based on the redox potentials from Table 1, using the previously determined electrochemical potential for one-electron reduction of photoexcited [Re(phen)(CO)$_3$(py)]$^-$ of 0.77 V vs. Fc$^+$/Fc (bottom row of Table 1).$^{59}$

**Transient absorption**

Figure 8a shows the transient absorption spectrum obtained from an acetonitrile solution with 6.7±10$^{-5}$ M [Re(phen)(CO)$_3$(py)]$^-$/CH$_3$OH and 10 mM Bu$_2$PhOH-CH$_2$-py. Selective excitation of the rhenium(I) complex occurred at 355 nm (Figure 2) with laser pulses of ~10 ns width. The data was time-averaged in a window ranging from 0 to 200 ns after the excitation pulse. The spectrum in Figure 8a exhibits the signatures of the reduced rhenium tricarbonyl diimine complex and neutral phenoxyl radical at the same time. The intense narrow band centered around 315 nm and the weaker featureless band extending from 300 to 200 ns after the excitation pulse. The spectrum in Figure 8a exhibits the signatures of the reduced rhenium tricarbonyl diimine complex and neutral phenoxyl radical at the same time. The intense narrow band centered around 315 nm and the weaker featureless band extending from 300 to 550 nm is typical for the one-electron reduced form of the rhenium complex considered here.$^{56, 24}$ On the other hand, the narrow peaks at 390 nm and 409 nm (dashed vertical arrows) are due to the phenoxyl radical as becomes evident from comparison to the spectrum in Figure 8b. The latter spectrum was recorded after 355-nm excitation of a CH$_2$Cl$_2$ solution containing 2 mM Bu$_2$PhOH-CH$_2$-py, 5 mM 1,4-dicyanonaphthalene, and 0.3 M biphenyl. These reaction conditions (making use of 1,4-dicyanonaphthalene as a photosensitizer and biphenyl as a co-donor) represent an efficient means for the photogeneration of neutral phenoxyl radicals.$^{30}$ In presence of phenol the spectral signatures of reduced 1,4-dicyanonaphthalene and oxidized biphenyl have disappeared within 6 μs, and hence when detecting with a delay of 6.6 μs after the 10-nj laser pulse one obtains the spectrum shown in Figure 8b, representing the
absorption spectrum of the phenoxyl radical of \( \text{Bu}_2\text{PhOH-CH}_2\text{-py} \).\cite{25-28} The observation of the same spectral features in Figure 8a is direct evidence for a PCET reaction, since one detects the oxidized and deprotonated form of the phenol in addition to the reduced form of the rhenium(I) reaction partner. The proton acceptor site is most likely the pyridine unit of \( \text{Bu}_2\text{PhOH-CH}_2\text{-py} \) but this cannot be monitored by transient absorption spectroscopy.

Thus we conclude that in the PhOH-\( \text{CH}_2\text{-py} / \text{rhenium(I)} \) and \( \text{Bu}_2\text{PhOH-CH}_2\text{-py} / \text{rhenium(I)} \) reaction couples considered here the overall quenching process is a CPET reaction involving electron transfer from the phenols to the photoexcited metal complex and transfer of the phenolic proton to the pyridine base. Transient absorption data support this conclusion because they provide direct evidence for the photogeneration of neutral phenoxyl radicals. H/D kinetic isotope effects on the order of 2 point to the involvement of concerted proton-electron transfer in the excited-state chemistry, in line with the prior notion that the oxidation potentials of the pyridylphenols can only be that much lower than those of ordinary phenols (\( -0.5 \text{ V} \)) because electron release is inherently coupled to deprotonation.\cite{29, 5d, 5e}

Interestingly, excited-state quenching with \( \text{Bu}_2\text{PhOH} \) by simple electron transfer is already quite efficient (\( (3.3 \pm 0.1) \times 10^8 \text{ M}^{-1} \text{ s}^{-1} \)), yet \( \text{Bu}_2\text{PhOH-CH}_2\text{-py} \) quenches the rhenium \( \text{MLCT} \) state even more rapidly (\( (13.3 \pm 0.1) \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \)) and with an H/D KIE of \( ~2 \), suggesting that CPET stays kinetically highly competitive with an electron transfer / proton transfer sequence even in a situation in which the initial electron transfer step is thermodynamically possible.

**Experimental Section**

A suspension of NaH (60% in mineral oil, 1.20 g, 30 mmol) in anhydrous THF (6 ml) was cooled to 0°C and a solution of 2-bromophenol (1) (3.44 g, 20 mmol) in anhydrous THF (6 ml) was added dropwise over 1 hour under N\(_2\) atmosphere. After stirring for an additional 10 minutes iodomethane (9.2 g, 64.8 mmol) was added, and the mixture was allowed to warm up to room temperature prior to refluxing for 19 hours. After cooling back to room temperature water was added (160 ml), and the product was extracted with pentane (3 \( \times \) 100 ml). The combined organic phases were dried over anhydrous MgSO\(_4\) and the solvent was removed on a rotary evaporator. Subsequent purification on silica gel with CH\(_2\)Cl\(_2\) as an eluent (R\(_s\) = 0.3) gave 1-bromo-2-methoxybenzene (2) in 97% yield.\cite{21} \( ^1\text{H} \) NMR (300 MHz, CDCl\(_3\)), \( \delta \) (ppm) = 3.89 (s, 3 H), 6.79-6.95 (m, 2 H), 7.27 (td, \( J = 8.2, 1.6 \text{ Hz}, 1 \text{ H} \)), 7.54 (dd, \( J = 7.8, 1.6 \text{ Hz}, 1 \text{ H} \)).

The same procedure using identical molar quantities of starting materials was employed for the synthesis of 1-bromo-3,5-di-tert-butyl-2-methoxybenzene (4) from 2-bromo-4,6-di-tert-butylbenzophenone (3). On silica gel using CH\(_2\)Cl\(_2\) as an eluent, product 4 has R\(_s\) = 0.6, the yield was 97%. \( ^1\text{H} \) NMR (300 MHz, CDCl\(_3\)), \( \delta \) (ppm) = 1.30 (s, 9 H), 1.40 (s, 9 H), 3.91 (s, 3 H), 7.28 (d, \( J = 2.4, 1.6 \text{ Hz}, 1 \text{ H} \)), 7.41 (d, \( J = 2.4, 1.6 \text{ Hz}, 1 \text{ H} \)).

For the synthesis of molecule 5 a solution of 2-picoline (200 mmol) in anhydrous THF (200 ml) was cooled to -30°C and 1.6 M n-butyllithium in hexane (200 mmol) was added slowly. After stirring at this temperature for 30 minutes disopropyl ketone was added slowly, and the reaction mixture was stirred for another 2 hours at room temperature. Subsequently water (300 ml) was added and the resulting mixture was extracted with ethyl acetate (3 \( \times \) 200 ml). The combined organic phases were dried over anhydrous MgSO\(_4\), and the solvent was removed using a rotary evaporator. The raw product was purified by column chromatography on silica gel using a 3:1 (v:v) mixture of pentane and ethyl acetate as an eluent (R\(_s\) = 0.6). This procedure afforded pure molecule 5 in 83% yield.\cite{21} \( ^1\text{H} \) NMR (300 MHz, CDCl\(_3\)), \( \delta \) (ppm) = 0.88 (dd, \( J = 6.9, 4.9 \text{ Hz}, 12 \text{ H} \)), 1.90 (m, 2 H), 2.98 (s, 2 H), 6.29 (s, 1 H), 7.08-7.13 (m, 1 H), 7.15 (d, \( J = 7.8, 1 \text{ Hz}, 1 \text{ H} \)), 7.58 (td, \( J = 7.7, 1.9 \text{ Hz}, 1 \text{ H} \)), 8.43 (ddd, \( J = 4.9, 1.8, 0.9 \text{ Hz}, 1 \text{ H} \)).

Using a heat gun Cs\(_2\)CO\(_3\) (27.90 g, 85.6 mmol) was dried under vacuum. Subsequently, palladium trifluoroacetate (1.18 g, 3.6 mmol), tricyclohexylphosphine (2.00 g, 7.1 mmol), dry xylene (150 ml), 1-bromo-2-methoxybenzene (2) (16.00 g, 85.6 mmol), and pyridine 5
(14.70 g, 71.3 mmol) were added under nitrogen. The reaction mixture was refluxed under N2 overnight. After cooling to room temperature the mixture was filtered, and the solvent was evaporated under reduced pressure. Column chromatography on silica gel using a 5:1 (v/v) mixture of pentane and ethyl acetate (Rf = 0.1) afforded the coupling product 6 as a yellow liquid in 73% yield.\(^1\) H NMR (300 MHz, CDCl3): δ (ppm) = 3.80 (s, 3 H), 4.17 (s, 2 H), 6.90 (ddd, J = 8.1, 6.3, 2.6 Hz, 2 H), 7.03-7.14 (m, 2 H), 7.14-7.26 (m, 2 H), 7.54 (td, J = 7.7, 1.9 Hz, 1 H). 8.49-8.58 (m, 5 H).

Molecule 6 (0.15 g, 0.78 mmol) was dissolved in aqueous HBr (47%, 4 ml) and the mixture was refluxed for 19 hours.\(^2\) After evaporation of excess acid, water was added to the solid residue, and the solution was neutralized by addition of an aqueous KO2CO3 solution. The product (7) was extracted with CH2Cl2, and the combined organic phases were dried over anhydrous MgSO4 prior to solvent removal on a rotary evaporator. The raw product was purified by column chromatography on silica gel using CH2Cl2 as an eluent (Rf = 0.4) affording product 7 (PhOH-CH3-py) as a white solid in 57% yield.\(^1\) H NMR (300 MHz, CDCl3): δ (ppm) = 4.10 (s, 2 H), 6.82 (td, J = 7.4, 1.3 Hz, 1 H), 6.95-7.02 (m, 1 H), 7.13-7.21 (m, 3 H), 7.32 (d, J = 7.8 Hz, 1 H), 7.69 (td, J = 7.7, 1.8 Hz, 1 H), 8.46 (ddd, J = 5.0, 1.8, 0.8 Hz, 1 H), 11.67 (s, 1 H).\(^1\) C NMR (75 MHz, CDCl3): δ (ppm) = 41.7, 118.5, 119.9, 121.9, 122.8, 126.2, 128.7, 130.2, 138.2, 147.8, 156.7, 161.0. MS (EI): m/z = 186.0921 ([M+H]+). Anal. calc.: C: 80.35, H: 9.97, N: 7.56; found: C: 77.63, H: 9.57, N: 7.47.

For the synthesis of molecule 8 from 1-bromo-3,5-di-tert-butyl-2-methoxybenzene (4) and pyridine 5 the same procedure as described above for molecule 6 was used.\(^3\) The quantities of reagents used in this case were as follows: Cs2CO3, 22.81 g, 70.0 mmol; palladium trifluorocetate: 0.97 g, 2.9 mmol; tricyclohexylphosphine: 1.63 g, 5.8 mmol. Using the same chromatography conditions as described above for molecule 6, the Rv value is ~0.3, and molecule 8 was obtained in 69% yield as a yellow oil.\(^4\) H NMR (300 MHz, CDCl3): δ (ppm) = 1.26 (s, 9 H), 1.43 (s, 9 H), 3.75 (s, 3 H), 4.27 (s, 2 H), 6.98-7.17 (m, 3 H), 7.26-7.31 (m, 1 H), 7.56 (td, J = 7.7, 1.9 Hz, 1 H), 8.57 (ddd, J = 4.9, 1.8, 0.9 Hz, 1 H).

When subjecting molecule 8 to the same methoxyl-deprotection reaction with aqueous HBr as described above for molecule 6, not only the methoxyl-group but also the tert-butyl substituents of molecule 8 were cleaved off and the reaction afforded molecule 7. It was therefore necessary to apply the following procedure in order to obtain molecule 9: To a suspension of NaH (60% in mineral oil, 14.70 g, 71.3 mmol) were added under nitrogen. The reaction mixture was refluxed under N2 overnight. After cooling to room temperature ethanethiol (0.69 g, 11.2 mmol) was added dropwise. Then molecule 8 (0.50 g, 1.6 mmol) was added and the reaction mixture was stirred at 100 °C overnight. After cooling to room temperature, H2O (4 ml), 1 M aqueous HCl (13 ml) and phosphate buffer (0.5 M, pH = 7) were added, and the mixture was extracted with diethyl ether (3 × 50 ml). The combined organic phases were dried over anhydrous MgSO4, and the solvent was removed on a rotary evaporator; this procedure was accomplished twice in order to ensure high isotope purity. Luminescence quenching experiments in CH3Cl2 occurred in presence of 100 mM CBDOD (99.99%) in order to avoid significant D/H exchange through contact of the phenols with glassware and cuvettes. For experiments with undeuterated phenols, 100 mM CH3OH was added.

The [Re(phen)(CO)3(py)][\[8\]] complex was available from prior studies in the form of its hexafluorophosphate salt.\(^{[6, 9]}\)

\(^1\)H and \(^13\)C NMR spectra were measured on a Bruker B-ACS-120 instrument. Electron-impact mass spectrometry was made with a Finnigan MAT 95 spectrometer. Elemental analyses were performed with a Vario EL3 instrument. For cyclic voltammetry a Versastat3-200 potentiostat from Princeton Applied Research was used. A glassy carbon working electrode, a platinum counter electrode, and a silver quasi-reference electrode were employed. 0.1 M tetrabutylammonium hexafluorophosphate (TBAPF6) was used as an electrolyte; prior to voltage sweeps at rates of 0.1 V/s the solvent was flushed with nitrogen. Optical absorption spectra were recorded on a Cary 5000 instrument from Varian, steady-state luminescence experiments were performed on a Fluorolog-3 apparatus from Horiba Jobin-Yvon. Luminescence lifetime experiments were performed in Geneva using a home-built setup comprised of a Quantel Brilliant Nd:YAG laser with an integrated magic prism OPO as an excitation source and a detection system consisting of a Spex 270M monochromator, an R928 photomultiplier from Hamamatsu, and a Tektronix TDS 540B digital oscilloscope. Transient absorption experiments were performed in Göttingen using an LP920-KS instrument from Edinburgh Instruments equipped with an ICCD camera from Andor and a Quantel Brilliant b laser as an excitation source.

For the X-ray structures crystals of 7 and 9 were grown by slow evaporation of the solvent at room temperature. They were selected and mounted with inert oil on a kapton MicroMount. The data of 7 were collected at 100 K on a Bruker Apex II Ultra with mirror optics. The data of 9 were collected on a Bruker smart Apex II Quazar with an Inotecat IpS.\(^{[28]}\) Data reduction was performed with SINAUT,\(^{[29]}\) and an empirical absorption correction with SADABS\(^{[30]}\) was applied. The structures were solved by direct methods (SHELXL-97)\(^{[31]}\) and refined by full-matrix least-squares methods against F2 with SHELXL-97\(^{[32]}\) and the shelixe\(^{[33]}\) GUI. All non-hydrogen atoms were refined with anisotropic displacement parameters. The hydrogen atoms were refined isotropically on calculated positions using a riding model with their Ueq values constrained to 1.5 times the Ueq values of their pivot atoms for terminal sp2 carbon atoms and 1.2 times for all other carbon atoms. Crystallographic data (excluding structure factors) for the structures reported in this paper have been deposited with the Cambridge Crystallographic Data Centre. The CCD numbers, crystal data and experimental details for the X-ray measurements are listed in the supporting information. CCDC 887369 (7) and 887370 (9) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

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ARTICLES

Two pyridylphenols with intramolecular hydrogen bonds react with a metal complex via concerted proton-electron transfer: Upon photoexcitation of the complex, there is electron transfer from the phenol to the metal, and the phenolic proton is transferred to the pyridine moiety.

William Herzog, Catherine Bronner, Susanne Löfler, Bice He, Daniel Kratzert, Dietmar Stalke, Andreas Hauser, Oliver S. Wenger*

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Electron Transfer between Hydrogen-Bonded Pyridylphenols and a Photoexcited Rhenium(I) Complex