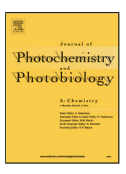
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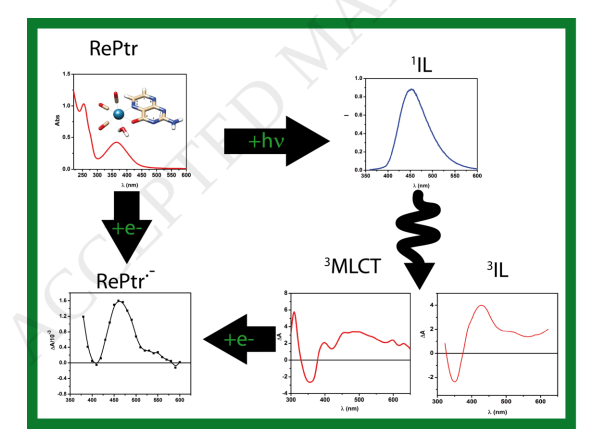
Photophysics, photochemistry and thermally-induced redox reactions of a (Pterin)rhenium(I) complex

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Graphical abstract



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Highlights

- Photophysics and photochemistry study of the (Pterin)rhenium(I) complex
- Nature of the excited states depend on solvent polarity.
- Flash photolysis and Pulse radiolysis experiments generated reactive species
- MCR-ALS analysis and TD-DFT calculations helped to ascertain the nature of these species

Abstract

In this work, we present a whole and deep study on the thermal redox and the photophysical and photochemical reactions of a tricarbonyl Re(I) complex coordinating Pterin, fac-Re^I(CO)₃(pterin)(H₂O) (pterin = 2-amino-4-oxo-3H-pteridine). In aqueous solutions, the fluorescence of the complex is attributed to the emitting ¹IL state ($\tau_{emi} \sim 7.6$ ns). In MeCN, however, the luminescence was ascribed to an overlapping dual emission from ¹IL and ³MLCT states (τ_{emi1} = 8.0 ns and τ_{emi2} = 1.0 µs). Oxygen quenching of the ³MLCT based luminescence occurred with $k_q = 1.6 \times 10^9 \text{ M}^{-1}\text{s}^{-1}$. In glasses at 77K, nevertheless, the ³MLCT prevailed over ¹IL states. Flash photolysis experiments in aqueous solutions showed the spectrum of ³IL while in MeCN the presence of ³MLCT was evident. Pulse radiolysis experiments under oxidizing and reducing environments were performed in aqueous solutions of the Re-Pterin complex. By performing multivariate curve resolution - alternating least-squares (MCR-ALS), two species were identified under reducing conditions: an intermediate related to the semireduced radical of pterin ligand and the dihydrogenated $[Re(CO)_3(7,8-PtrH_2)(H_2O)]$ product. TD-DFT calculations helped to ascertain the nature of these species. Flash photolysis experiments where the excited states were reduced with triethylamine were in good agreement with pulse radiolysis

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experiments under reductive conditions. The oxidized transient spectrum was also obtained by pulse radiolysis, which compares very well with those published for Re(II) species, leading us to propose oxidation in the metal core as the reaction product of the transient under oxidizing conditions.

The combination of different spectroscopic techniques along with the theoretical calculation allowed elucidating the nature, dynamics and reactivity of the excited states prevail in a Re-Pterin complex. This is of the particular importance considering that equivalent studies have not been reported for any other rhenium complex of substituted pterins nor for complexes of other transition metal ions of pterin derivatives.

KEYWORDS

Rhenium complex, photochemistry, excited states, Pterin, TD-DFT.

Introduction

Pterins (2-amino-4-oxo-3H-pteridine) (Ptr) are heterocyclic compounds widely distributed in biological systems. Pterins are involved in various photobiological processes, in the mammal metabolically routes, in sensory organs of vertebrates and invertebrates and pigments in arthropods [1],[2]. The photochemical and photophysical properties of pterins have been extensively studied [3]–[7]. The coordination of an organic ligand to a metal introduces new electronic transitions such as metal-to-ligand charge transfers (MLCTs). These MLCT excited states show a strong dependence on the polarity of the solvent and a

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marked heavy metal effect, the latter being reflected in quantum yields of triplet formation close to unity [8].

Transition metal complexes comprise an interesting and widely extended field for studies related to different areas of scientific research such as electron transfer studies [9], solar energy conversion [10], and catalysis [11]. Possible applications as luminescent sensors [12],[13] and molecular materials for nonlinear optics or optical switching [14],[15] have also emerged. In particular, luminescent transition metal complexes of Re(I) and Ru(II) with polypyridyl ligands have been recognized as good potential candidates for the development of pH sensing devices [16], [17] (and references therein). Moreover, there are potential biochemical and technical applications based on the formation of adducts between transition metal complexes of Re(I), Ru(II), Os(III), Rh(III), or Co(III) and biological macromolecules such as DNA which take advantage of the luminescent properties of the ³MLCT states [18]–[20]. As these complexes show exceptionally rich excited-state behavior and redox chemistry as well as thermal and photochemical stability [8] they have also been used as biological labelling reagents and noncovalent probes for biomolecules and ions [13]. In this regard, some complexes including biologically relevant ligands, such as substituted pterins, coordinated to the Re(CO)₃Cl core have been synthesized and characterized [21]-[23].

As far as we know, there are only a few examples concerning the photophysical properties of metallic complexes coordinating pterin and at present there are not published papers in the literature regarding the photophysical properties of Re(I) complexes that coordinate that ligand [24]. In this work, we present a whole and deep study on the solvent

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dependent thermal redox and the photophysycal and photochemical reactions of a tricarbonyl Re(I) complex coordinating Ptr, *fac*-Re^I(CO)₃(pterin)(H₂O) (**RePtr**).

Experimental Part

Materials.

The **RePtr** complex was available from previous work [25][26]. Spectrograde and HPLC grade acetonitrile (MeCN) (J.T. Baker), methanol (MeOH, Sigma Aldrich) and ethanol (EtOH, Sigma Aldrich) were used without further purification. 2-hydroxybenzophenone, phenalenone, New Coccine, D₂O, NaOH, and HClO₄ were purchased from Sigma Aldrich at the highest purity available and were used as received. Deionized water (> 18 M Ω cm⁻¹, < 20 ppb of organic carbon) was obtained with a Millipore system.

Photophysical Measurements.

UV-vis spectra were recorded on a Shimadzu UV-1800 spectrophotometer. Emission spectra were obtained with a computer-interfaced Near-IR Fluorolog-3 Research Spectrofluorometer, and were corrected for differences in spectral response and light scattering. Solutions were deaereated with ultrahigh-purity O₂-free nitrogen in a gas-tight apparatus before recording the spectra.

Emission quantum yields, ϕ_{em} , were calculated with **eq. 1**, by using solutions of a reference compound with a known quantum yield of emission, ϕ_{ref}

(1)

$$\phi_{\rm em} = \left(\frac{A_{\rm ref}}{A_{\rm s}}\right) \left(\frac{I_{\rm s}}{I_{\rm ref}}\right) \left(\frac{n_{\rm s}}{n_{\rm ref}}\right)^2 \phi_{ref}$$

In eq. 1, ns and nref are the refractive indexes of the solutions containing the sample and the reference compound respectively, A is the absorbance of the sample or reference at the excitation wavelength (A_s and $A_{ref} < 0.1$) and I is the integral of the emission spectrum and was used as a relative measure of the respective intensities of the luminescence. An aqueous solution of pterin at pH=5 was used as a reference with $\phi_{ref} = 0.33$ [27]. Emission lifetime measurements were performed using either Fluorolog-3 with Time-correlated Single-Photon Counting (TCSPC) unit with either 341 and/or 388 nm NanoLEDs. Photoacoustic measurements were performed by using a set-up already described [28]. The resolution time in our experimental set-up, τ_R , was ca. 800 ns [28]. New Coccine or 2hydroxybenzophenone were used as calorimetric reference (CR) compounds in the buffer and in MeCN solutions, respectively [29][30]. Experiments were performed under a controlled atmosphere, bubbling N_2 or O_2 in the solution, for 15 min. In principle, all the excited species with lifetimes $\tau \leq 1/5 \tau_{\rm R}$ release their heat content as prompt heat, whereas excited species with a lifetime of $\tau \ge 5 \tau_{\rm R}$ function as heat storage within the time resolution of the LIOAS experiment.

For the handling of the LIOAS signals, **eq. 2** was used, which relates the peak to peak amplitude of the first optoacoustic signal (H) with the fraction of the excitation laser fluence (F) absorbed by the sample [31].

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(2)

$$H = K \alpha F (1-10^{-A})$$

In this equation the experimental constant *K* contains the thermo-elastic parameters of the solution as well as instrumental factors, *A* is the absorbance of the sample at λ_{exc} and α is the fraction of the energy released to the medium as prompt heat, i.e. within the time resolution of the experiment.

The efficiency of **RePtr** complex toward singlet oxygen sensitization was assessed by the direct measurement of the ${}^{1}O_{2}$ (${}^{1}\Delta_{g}$) near-infrared luminescence. After the irradiation of aerated solutions of the complex the generation of ${}^{1}O_{2}$ (${}^{1}\Delta_{g}$) was evidenced by the appearance of the characteristic ${}^{1}O_{2}$ (${}^{1}\Delta_{g}$) $\rightarrow {}^{3}O_{2}$ phosphorescence at 1270 nm. Time resolved phosphorescence detection was used for singlet oxygen detection. The near IR luminescence of ${}^{1}O_{2}$ (${}^{1}\Delta_{g}$) was detected by Ge-photodiode (Applied Detector Corporation, time resolution 1 μ s). The quantum yield of ${}^{1}O_{2}$ (${}^{1}\Delta_{g}$) formation, ϕ_{Δ} , was determined by measuring its phosphorescence intensity using an optically matched solution of phenalenone ($\phi_{\Delta} = 0.98$) [32] as a reference sensitizer.

Transient absorption spectra were recorded using flash photolysis apparatus described elsewhere [33]. In these experiments, 25 ns flashes of 351nm (ca. 25-30 mJ/pulse) light were generated with a Lambda Physik SLL-200 excimer laser. Decays typically represented the average of 10 to 30 pulses. Edinburgh L980 flash photolysis instrument with Nd:YAG laser Continum (355 nm, laser pulse ~30 ns) excitation sources was also used. The intensity of both emitted and scattered light was subtracted in each trace. Solutions for the photochemical work were deoxygenated with streams of ultrahigh-purity N₂ before

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and during the irradiations. A flow system ensured that the fresh solution was brought to the reaction cell in experiments where the decomposition of the Re(I) complexes and/or formation of products interfered with the optical measurements. The concentration of the complexes were adjusted to provide homogeneous concentrations of the photogenerated intermediates within the volume of the irradiated solution, *i.e.*, optical densities equal to or less than 0.1 (optical length = 1 cm) at 355 nm.

Curve Resolution Methods

For the analysis of the time-resolved difference spectra we have chosen one of the most widely used algorithms namely multivariate curve resolution - alternating least-squares (MCR-ALS), which can provide simultaneous estimates for the concentration and spectral profiles of unknown spectroscopic mixtures [34],[35] (and references therein).

Pulse radiolysis

Pulse radiolysis experiments were carried out with a model TB-8/16-1S electron linear accelerator. The instrument and computerized data collection for time-resolved UV-vis spectroscopy and reaction kinetics have been described elsewhere [36],[37].

In the several ns time domain, the radiolysis of H₂O, eq. 3, yields e_{aq}^{-} , H and •OH radicals

[38]:

$$H_2O \longrightarrow e_{aq}^-, \bullet OH, H, H^+$$
(3)

The hydrated electron, the strongest chemical reductant (-2.8 V vs. NHE), will often reduce a substrate S to S^{\bullet -} in a diffusion-controlled reaction, eq. 4:

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S +
$$e_{aq}^{-}$$
 > S⁻ (4)
The oxidizing 'OH radical can efficiently be removed to prevent its reaction with S by the
addition of tert-butyl alcohol or 2-propanol, eqs. 5-6 [38]:
'OH + (CH₃)₃COH → H₂O + ('CH₂)C(CH₃)₂OH (5)
'OH + (CH₃)₂CHOH → H₂O + ('CH₂)C(CH₃)₂OH (6)
The (CH₃)₂C'OH radical, being a more powerful reducing radical (E⁰ = -1.2 V vs NHE) than
the ('CH₂)₂C(CH₃)₂OH radical (E⁰ = -0.1V vs NHE), will usually reduce S to S⁺ (eq. 7)
S + (CH₃)₂C'OH → S⁺ + H⁺ + (CH₃)₂CO (7)
However, the ('CH₂)C(CH₃)₂OH radical in some occasions may also yield S⁺.
Other 'OH radical scavenger is formate, HCO₂. The CO₂⁺ formed radical from eq. 8 will
generally yield the same S⁺ via eq. 9 [38]:
'OH + HCO₂ → H₂O + CO₂⁺ (8)
S + CO₂⁺ → S⁺ + CO₂ (9)
Since 'OH radical is an oxidizing species which reacts with a lack of selectivity, it may be
converted to more selective oxidizing radicals like X₂⁺ (X = halogen or SCN') and N₃⁺, eqs.
10-11, when it is needed to study thermal oxidations of S by pulse radiolysis experiments
[38].
'OH + 2X' → X₂⁺ + OH⁻ (10)
'OH + N₃ → N₃⁺ + OH⁻ (11)
In the study of oxidations reaction 4 is unwanted. It is simple to remove the reducing
radical e_{aq}^{-} after the saturation of the solution with N₂O, eq. 12 [38].

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e_{aq}^{-} + N₂O + H₂O \rightarrow •OH + OH - + N₂

Thiocyanate dosimetry was carried out at the beginning of each experimental session. Details of the dosimetry have been reported elsewhere [37],[39]. The procedure is based on the concentration of $(SCN)_2^{\bullet-}$ radicals generated by the electron pulse in a N₂Osaturated 10^{-2} M SCN⁻ solution (eq. 10). In the procedure, the calculations were made with G = 6.13 and an absorption coefficient ε = 7.58 x 10^3 M⁻¹ cm⁻¹ at 472 nm [37],[39] for the (SCN)₂^{•-} radicals. In general, the experiments were carried out with doses that in N₂saturated aqueous solutions resulted in $(1.7\pm0.1) \times 10^{-6}$ M to $(6.0\pm0.3) \times 10^{-6}$ M concentrations of e^{-}_{aq} . In these experiments, solutions were deaerated with streams of N₂ or N₂O gasses. In order to irradiate a fresh sample with each pulse, an appropriate flow of the solution through the reaction cell was maintained during the experiment.

Computational details

The electronic structure of the rhenium complex was studied by performing Density Functional Theory [40]–[42] (DFT) and Time Dependent Density Functional Theory (TD-DFT) [43]–[45] (TD-DFT) calculations using Gussian 09 software [46]. The ground state structure of the complex was optimized by DFT calculations using B3LYP functional and LanL2DZ basis set. Vibrational frequencies were computed at the same level of theory to confirm that these structures were minima on the energy surfaces. Solvents effects (H₂O) on the optimization of the ground state structure were taken into account by means of the Polarizable Continuum Model (PCM) [47]–[49]. At the optimized ground state geometries, a set of 200 vertical excitations were computed with the PBE0 hybrid

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functional in H₂O. The PBE0 hybrid functional was shown to give a very good performance in the simulation of the electronic spectrum of Re(CO)₃(Ptr)(H₂O) as evidenced by the best match (relative to the performance of other functionals as B3LYP, XB3LYP and CAM-B3LYP) between calculated and experimental electronic transitions [7]. Therefore, all the electronic transitions presented hereafter are those calculated using PBE0 functional for the Re(I) complex. TD-DFT calculations with the PBE0 functional were performed using the 6-311++G(d,p) basis set for C, N, O and H atoms while LanL2TZ(f) [50],[51] (triple zeta basis set designed for an ECP plus f polarization) was used for Re atom. Absorption spectra were simulated with Gaussian distributions with a full-width at half-maximum (fwhm) set to 3000 cm⁻¹ with the aid of GaussSum 2.2.5 program [52]. The calculated electronic spectra of Re(CO)₃(Ptr)(H₂O) are simulated from the theoretical results to ease the comparison with experimental data.

Results and discussion

Photophysical Processes

The luminescence spectrum of **RePtr** in air-equilibrated aqueous solutions upon excitation in the low energy band ($\lambda_{exc} = 350$ nm) consists of a broad unstructured band with λ_{max} centered around 450 nm. When the aqueous solutions were deaerated by bubbling either O_2 -free N₂ or O_2 no significant differences were observed either in λ_{max} , ϕ_{em} and/or luminescence lifetime (τ_{em}). In acidic solutions (pH = 3 and pH = 6) the emission band is centered at $\lambda_{max} = 441$ nm while in alkaline solutions (pH = 10) the luminescence maximum

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experiences a bathochromic shift to $\lambda_{max} = 454$ nm, see **Figure 1**. In previous work [25], protonation studies in aqueous solutions of the RePtr complex showed two acid-base equilibriums with pK_{a1} = 3.9 and pK_{a2} = 8.8. pK_{a1} was assigned to the protonation equilibrium at N3, see **Figure S1**, of the pterin ligand in the complex and pK_{a2} could be ascribed to the deprotonation of a coordinated water molecule. Therefore, the species present are [Re(CO)₃(Ptr)(OH)]⁻ (**RePtr**⁻) at pH = 10 and [Re(CO)₃(Ptr)(H₂O)] and [Re(CO)₃(PtrH)(H₂O)]⁺ (**RePtr**⁺) at pH = 6 and pH = 3, respectively. Nevertheless, neither spectral shifts nor band shape differences can be observed from the comparison of the luminescence spectra of **RePtr** complex at both acidic pHs. The emission quantum yield, however, increases moderately from 6.4 x 10⁻² at pH = 3 to 9.1 x 10⁻² at pH = 6 while at pH = 10 it experiences a dramatic enhancement to $\phi_{em} = 0.25$. Conversely, the luminescence decays, which are monoexponential with lifetimes around 7 ns, experience only minor changes upon pH change (See Table 1).

The luminescence parameters of **RePtr** complex in alkaline media shows similar characteristics to the fluorescence of the free ligand at pH = 10.5, i.e., both λ_{max} and ϕ_{em} are similar [6]. On the other hand, the emission quantum yield of **RePtr** complex in acidic media is one order of magnitude smaller than that of the Ptr ligand. After irradiation of the **RePtr** complex at λ_{exc} = 350 nm in aqueous solution, the ¹IL and ¹MLCT excited states are simultaneously populated. These excited states later populate via intersystem crossing processes, ISC, the respective ³IL and ³MLCT. Population of the latter is favored by the strong spin-orbit coupling induced by the metal ion in the complex. TD-DFT predicts that

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the low energy absorption band of both **RePtr** and **RePtr**⁻ consist of a H-1 \rightarrow L transition [7]. **Figure S2** shows density of states (DOS) plot for those rhenium species. It is observed that H-1 \rightarrow L transition in **RePtr** has a marked MLCT (Re(CO)₃ \rightarrow Ptr) character while in **RePtr**⁻ that transition has, in addition, a considerable IL Ptr ($\pi \rightarrow \pi^*$) character. Since population of MLCT is more important than population of IL states, a lower luminescence quantum yield is expected for **RePtr** than pterin. On the other hand, when the prevalent specie is **RePtr**⁻ (basic solution) the contribution of MLCT to the overall excited state is less important prevailing the IL character, and as consequence, the luminescence quantum yield of **RePtr** complex is similar to that of free pterin.

Luminescence properties were also determined in acetonitrile solutions of **RePtr** complex, **Figure 2**. In that condition, we can assume that the coordinated water molecule is replaced by one acetonitrile molecule being the [Re(CO)₃(Ptr)(MeCN)] complex the prevalent species in this solvent [53].

Two emission bands were observed in the luminescence spectrum of **RePtr** complex upon excitation with $\lambda_{exc} = 350$ nm in acetonitrile. The most intense emission band at $\lambda_{max} = 418$ nm is nearly 30 nm blue shifted relative to the emission in aqueous solutions. The emission band centered at $\lambda_{max} \sim 570$ nm has a lesser intensity than the band at $\lambda_{max} = 418$ nm and it is quenched by the presence of oxygen in the solution, **Figure 2B**. When the excitation wavelength is $\lambda_{exc} = 400$ nm the low energy band is clearly observed while the luminescence band centered at $\lambda_{max} = 418$ nm decreased in intensity, **Figure S3A**. In MeOH/EtOH glasses at 77 K, the high energy band has almost disappeared and only a luminescence peaking at 575 nm is observed, **Figure S3B**.

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The same luminescence centered at 418 nm is observed with the **RePtr** complex and the free pterin in MeCN, pointing to the IL character of this emission. However, the additional feature centered at 570 nm, present in the luminescence of the **RePtr** complex, is not observed in acetonitrile solution of free pterin, **Figure S4**.

A sum of two exponentials was required to achieve a satisfactory fit of the emission decay profiles from acetonitrile solutions of **RePtr** complex at λ_{em} = 570 nm. The fast component decay is $\tau = 8.0$ ns under N₂ atmosphere and it is insensitive to the presence of oxygen in the solution. On the other hand, the slow component decay is $\tau = 1.0 \ \mu s$ under N₂ atmosphere and it was shortened by the presence of molecular oxygen, Table 1. The bimolecular rate constant for the quenching of the **RePtr** complex luminescence by oxygen (k_q , Table 1) has been determined from the slope of the linear Stern-Volmer plot $(\tau_0/\tau = 1 + k_a \tau_0[O_2])$ calculated with the values of the emission lifetimes in the absence and in the presence of oxygen (τ_0 in N₂ and τ in air and in O₂, respectively). The oxygen solubility at 1.013 bar of air was calculated according to $[O_2] = 0.21(P_A - P_v)[O_2]_{P=1}[54]$, where P_A and P_v are the atmospheric pressure and the vapor pressure of the solvent, respectively and $[O_2]_{P=1}$ is the oxygen concentration in the solvent at a 1.013 bar partial pressure of O₂. The saturated oxygen solubility at 1.013 bar in the organic solvents and water were taken from literature values [28], [55]. A bimolecular rate constant of $k_q = 1.6 \text{ x}$ 10⁹ M⁻¹s⁻¹ was calculated from the Stern-Volmer plot., **Figure S5**.

Singlet oxygen generation by the **RePtr** complex in both D_2O (at pD 4, 6 and 10) and MeCN solutions was analyzed by time resolved phosphorescence measurements (1270 nm). The phosphorescence of acetonitrile solutions showed clear evidence of singlet oxygen

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formation. Nevertheless, singlet oxygen generation in D₂O solutions could not be detected under our experimental conditions. Linear correlations were obtained from the plots of the dependence of the singlet oxygen phosphorescence intensity emission at zero time, S(0), as a function of the laser energy for the complex and the reference. From these slopes (Figure S6) and the usual procedure described elsewhere [30],[56] the determined quantum yield of $O_2({}^1\Delta_g)$ production was $\Phi_{\Delta} = 0.06$ in MeCN and $\Phi_{\Delta} < 10^{-3}$ in D_2O . LIOAS experiments carried out with both acetonitrile and aqueous solutions of Re-Ptr complex showed the same behavior: no time shift or changes of shape of the photoacoustic signal occurred with respect to the calorimetric reference (CR) signal (Inset of Figure 3). Linear relationships in both solvents were obtained between the amplitude of the first optoacoustic signal (H) and the excitation fluence (F) for samples and references at various A, in a fluence range between 1 and 20 J/m². The ratio between the slopes of these lines for sample and reference yielded the values of α for the samples. For the complex in MeCN solution the slopes were independent on the specific atmosphere and were the same that the CR measured at the equal experimental conditions, Figure 3. From these plots, considering that $\alpha_{\rm R} = 1$ for CR, the α values were calculated for the **RePtr** complex in both solvents. Therefore, a value of $\alpha = 1.00 \pm 0.04$ was obtained in aqueous and in MeCN solutions (either under N₂ or O₂ atmosphere). Consequently, this complex released to the medium all the absorbed energy as prompt heat (integrated by the transducer) in processes faster than $\tau_R/5$. These values combined with fluorescence data and singlet oxygen quantum yield production fit the energy balance [28],[31].

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Photochemical Reactions.

Figure 4 shows the absorption spectra of the transient generated when deaerated aqueous solutions of **RePtr** ([Re]=1 x 10⁻⁴ M at pH=6) were irradiated at 355 nm in flash photolysis experiments. An absorption maximum near 430 nm overlapped with another band at 400 nm and a long tail stretching up to $\lambda \sim 650$ nm were observed. The ground state depletion in the 300-400 nm region and the presence of isosbestic points makes it possible to determine ε_T (330 nm) = 2100 and ε_T (370 nm) = 4600 M⁻¹cm⁻¹ (ε_T = absorption coefficient of transient). A 430 nm trace could be fitted by a mono-exponential function with a lifetime $\tau_T \sim 2 \mu s$, Figure S7. Since the transient spectra bear a strong resemblance with those of the pterin triplet [57][4], the transient spectra can be assigned to the IL triplet excited state in the **RePtr** complex decaying by radiationless processes, **Figure 4**.

When 2 x 10⁻⁴ M solutions of **RePtr** complex in MeCN were flash irradiated at 355 nm, the generated transient shows an absorption spectrum with maximum near 400 nm, a broad absorption band spanning from 420 to 650 nm and ground state depletion in the 300-400 nm region, **Figure 5a**. **Figure 5b** shows a representative oscilographic trace of the transient at $\lambda_{obs} = 560$ nm, which could be well fitted by a mono-exponential function with a lifetime of $\tau_T = 1.0 \ \mu s$.

The transient spectrum obtained in acetonitrile medium can be assigned as a MLCT triplet excited state based on a couple of characteristics: *i*. the spectral features which differ from that one obtained in aqueous solutions (Figure 4) and *ii*. the transient lifetime is the same to that of the slow component of the luminescence decay, i.e. $\tau_T = \tau_{emi}$ (Table 1)

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Flash photolysis experiments were carried out also in the presence of the sacrificial donor triethylamine, TEA 0.1 M, under similar experimental conditions to those described in **Figure 5**. The excited state of **RePtr** reacted with TEA within the laser flash pulse to yield a transient whose spectral features, $\lambda_{max} = 460$ nm, **Figure 6A**, were different from those of the spectra in **Figure 5**. In addition, this transient decays whit $\tau_T = 132 \,\mu$ s, Figure **6B**. The nature and lifetime of this transient, generated by reaction between ³MLCT excited state of **RePtr** and an electron donor as TEA, can be assigned to a reduced radical of the complex.

Flash photolysis experiments of free pterin solutions in the presence of a sacrificial reductant like guanosine have shown the generation of the reduced Ptr radical [4]. The similarity between the spectrra of **Figure 6** and that of the semireduced pterin radical is consistent with the assignment of a reduced radical in the **RePt**r complex to the transient obtained in our experimental conditions.

Pulse Radiolysis

Pulse Radiolysis (PR) experiments were carried out with **RePtr** solutions under five different experimental conditions: N_3Na/N_2O ; $NaHCO_2/N_2$; $NaHCO_2/N_2O$, 2-propanol/ N_2 and 2-propanol/ N_2O .

Oxidation of RePtr by the N₃^{•-} radical

Transient spectra were recorded when a deaerated **RePtr** solution ([Re] = 4.0×10^{-5} M in H₂O) reacted with the pulse radiolysis generated N₃^{•-} radical according to eqs. 9, 10. The

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transient spectrum, **Figure 7**, shows absorption bands at 340 nm and 430 nm with a shoulder at 470 nm and other less intense absorption bands between 550 and 680 nm. Because of the transient spectral shape, the oxidized product can be assigned to a **Re(II)Ptr** species [58].

Reduction of RePtr by e_{aq}^{-} , (CH₃)₂C[•]OH and CO₂^{•-} radicals

The reactions of e_{aq}^{-} , (CH₃)₂C*OH and/or CO₂* radicals, eqs. 4, 7 and 9, with **RePtr** were studied in aqueous solutions of the complex ([RePtr] = 1.78 x 10⁻⁵ M at pH = 6, phosphate buffer) by means of the UV-Vis spectroscopic changes. In N₂ deaerated solutions, the reaction between e_{aq}^{-} and the **RePtr** complex was completed within the first µs after the radiolytic pulse. The transient spectrum generated by reaction with e_{aq}^{-} , exhibited bands at $\lambda_{max} \sim 375$ nm and 450 nm, **Figure 8A**. It evolved to the final product in the ms time scale. A bleach of the solution at ~ 380 nm can be observed in the spectrum of the product. The same spectral features were observed in the reaction of the complex with CO_2^{*} (**Figure 8B**) and (CH₃)₂C*OH radicals (**Figure S8**). Compared to the e_{aq}^{-} reaction, the pseudo first order rate constants for the build-up of the intermediates generated by the Ccentered radicals are an order of magnitude smaller than the case of the reduction with e_{aq}^{-} (see below).

Analyses of the time resolved spectra.

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The SVD and MCR-ALS methods were used for the analyses of the time-resolved difference spectra of pulse radiolysis experiments. Three contributions were required to explain with accuracy the trends observed in the spectra generated with each radical. In the fitting procedure non-negative and closure constraints were used. The individual contributions resolved by MCR-ALS method, which allow explaining the entire set of time resolved difference spectra recorded for the experiments performed in the presence of formate/N₂ or /N₂O, are given as supplementary material (**Figure S9**). The spectral contribution labeled as Sp1 (N₂) was ascribed to the solvated electron, e_{aa} , whereas the contribution labeled as Sp1 (N₂O) was assigned to the carbon dioxide anion radical, CO₂[•]. Similar results were obtained for Sp1 (N_2, N_2O) in the presence of 2-propanol. On the other hand, the other two spectral contributions, labeled in Figure S9 as Sp2 $(N_2)/Sp2$ (N_2O) and Sp3 $(N_2)/Sp3$ (N_2O) , were independent of the experimental condition tested, i.e., of the reducing radical present (NaHCO₂ (N₂); NaHCO₂ (N₂O), 2-propanol (N₂) and 2-propanol (N₂O)). Furthermore, these two contributions (from now on labeled as Sp2 and Sp3) showed their main spectral features in the wavelength range associated with the lowest energy absorption band of the **RePtr** complex and therefore were ascribed to its reductive transformation, Sp2 being a transient species and Sp3 the stable product within the timescale of PR experiments. Figure 9 compares the spectrum of the RePtr complex with the absolute spectra for the transient and the product calculated using the relative difference spectra resolved by MCR-ALS analysis, the radiation dose measured with the SCN⁻ actinometer and the radiation chemical yield of the 2-propanol radical.

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In the PR reduction of free pterin solution either by e_{aq}^{-} or CO₂[•], several pterin radicals have been suggested as initial intermediates in the process of reduction. Since rapid protonation after the electron attack occurs, protonated (neutral) radicals at N8 or N3, and either the diprotonated (positive) radical at both N8 and N3 have been proposed [59],[60]. However, as the coordination to the metal undoubtedly should affect the relative basicity of the N atoms in the pterin ligand, we have used DFT and TDDFT calculations to support the proposed species for the transient intermediate observed in PR experiments. Scheme I shows the optimized structures of RePtr, of the protonated radicals at N8 (RePtr(N8)) or N3 (RePtr(N3)), as well as the optimized structure of the diprotonated radical at both N3 and N8 (RePtr(N3,8)**), at the B3LYP/LanL2DZ/PCM (water) level of theory. Figure 10 shows the calculated UV-vis absorption spectra for RePtr, RePtr(N8)*, RePtr(N3)*, RePtr(N3,8)*+ and for the protonated reduced product Re(CO)₃(7,8-PtrH₂)(H₂O) (**RePtrH₂**, see below). From the comparison of **Figures 9** and **10** it can be inferred that radical RePtr(N8)* seems to describe better than radical RePtr(N3)* the overall spectral features of differences Sp2. However, radical **RePtr(N3,8)**^{•+} cannot be completely ruled out and radicals RePtr(N8)[•] and RePtr(N3,8)^{•+} may both be contributing to the spectrum of Sp2.

The analysis of the concentration profiles obtained by MCR-ALS analysis suggested two consecutive processes for all the experimental conditions tested. The concentration profiles were adequately described by an initial first order process, followed by a second order reaction. Since the concentration of radicals initially produced by the pulse was

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much lower than that of **RePtr**, the following kinetic model, eqs. 13 and 14, was proposed for the radiolysis experiments:

$$Sp1 + RePtr \xrightarrow{k_1} Sp2$$
(13)
$$2 Sp2 \xrightarrow{k_2} Sp3$$
(14)

Where **Sp1** stands for e_{aq}^{-} , CO₂⁻⁻ or (CH₃)₂C[•]OH

Hence a Hard-Soft MCR-ALS model [34] was developed for improving the analysis of time resolved spectra. The model included a deterministic part, since the concentration profiles during the iterative procedure were simulated by using only two rate constants (k_1 and k_2) associated with the two consecutive steps.

Therefore, the computational model allowed obtaining not only the spectral shapes and the concentration profiles of each species involved, but also the optimal values for the rate constants associated with each reaction step. **Figure 11** compares the concentration profiles obtained by the HS-MCR-ALS model with the concentration profiles simulated using the optimal values for k₁ and k₂ for the experiment performed in the presence of 2propanol and N₂O. Similar results were obtained for the other three experimental conditions tested. **Table 2** shows the optimal rate constant values obtained for the four experimental conditions tested.

Based on the what is known in the literature regarding pulse radiolytic, chemical and electrochemical reduction of free pterin [59]–[62] the reduction product **Sp3** can be proposed as the **RePtrH**₂ complex. **Figure S10** shows the optimized structure of the **RePtrH**₂ at the B3LYP/LanL2DZ/PCM (water) level of theory. The calculated spectrum of

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RePtrH₂ reproduces with reasonable accuracy the resolved spectrum of **Sp3** of pulse radiolysis experiments (see **Figure 10**). Given that 7,8-PtrH₂ ligand is a two electrons reduction of pterin, a disproportionation reaction between two **RePtr(N8)**• radicals have to be invoked to account for the formation of **RePtrH₂**, eq. 15.

2 RePtr(N8)• \rightarrow RePtrH₂ + RePtr (15)

This disproportionation reaction was proposed in the literature regarding reduction products of reduced pterin by PR [59],[60]. Disproportionation reactions of protonated reduced radicals of Re(I) complexes are common and well reported in the literature [35],[63]–[65].

Conclusions

The photochemical and thermal study realized in this work allowed us to identify the nature and characteristics of **RePtr** complex main excited states along with some intermediates and photo-products. In aqueous solutions, the complex luminescence is attributed to the emitting ¹IL state. In MeCN, however, both ¹IL and ³MLCT states contribute to the complex luminescence, the latter state being deactivated by molecular oxygen. Flash photolysis experiments, either in aqueous or MeCN solutions, helped to identify the nature (¹IL and/or ³MLCT) of the generated excited states. In glasses at 77K, nevertheless, the ³MLCT prevailed over ¹IL states. TD-DFT calculations performed on the ligand and the complex are in good agreement with the fact that ϕ_{em} of **RePtr** is much

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lower than that of the free ligand. The deactivation of ${}^{3}MLCT$ by O₂ leads to the formation of ${}^{1}O_{2}$ with a low quantum yield. No formation of ${}^{1}O_{2}$ in D₂O was detected, possibly due to a very low ${}^{3}IL$ formation quantum yield. When **RePtr** ${}^{3}MLCT$ is quenched by a sacrificial reductant, the species generated in flash photolysis experiments can be attributed to the radical of the reduced ligand. This intermediate was also observed in pulse radiolysis experiments. The stable product in the millisecond time scale could be assigned to the dihydrogenated complex [Re(CO)₃(7,8-PtrH₂)(H₂O)]. Under oxidative conditions a transient spectrum was obtained by pulse radiolysis which compares very well with those published for Re(II) species, suggesting that N₃^{•-} radical attack to the **RePtr** complex promotes the oxidation in the metal core.

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Figure captions

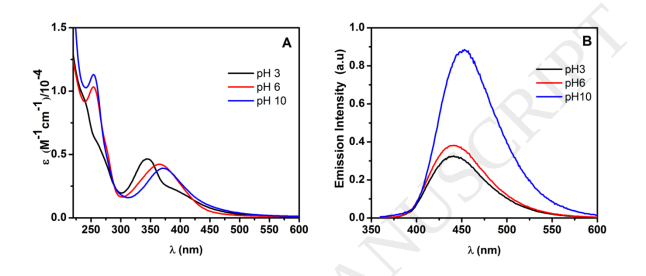
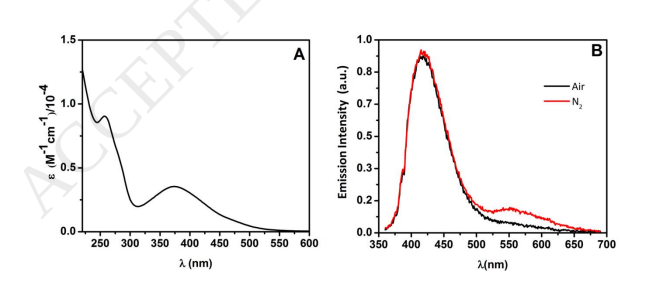


Fig. 1. (A) Absorption and (B) Emission spectra of the three forms acid base of the RePtr complex in aqueous solution at pH = 3, 6 and 10. λ_{exc} = 350 nm



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Fig. 2. (A) Absorption spectrum of the **RePtr** complex in deaerated MeCN and (B) emission spectra of the **RePtr** complex in deaerated and aerated MeCN solution. λ_{exc} = 350 nm

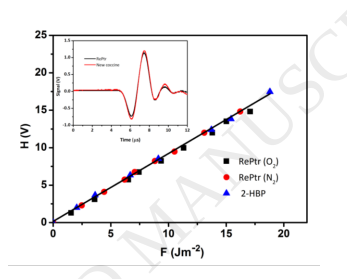


Fig. 3. Amplitude of the photoacoustic signals as a function of laser fluence for acetonitrile solutions: 2-hydroxybenzophenone (2-HBP) and **RePtr** complex (in N₂ and O₂). Inset: Normalized photoacoustic signals of aqueous solutions for New Coccine and the sample with matched absorbances (0.118 \pm 0.002).

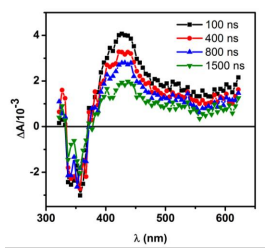
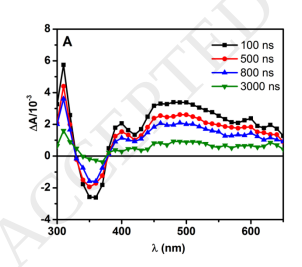
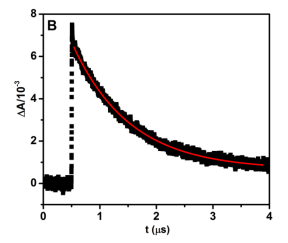


Fig. 4. Absorption spectra of the transient obtained in the λ_{exc} = 355 nm flash irradiation of the **RePtr** complex in a pH = 6 aqueous solution. Delays from the laser flash are reported

in the figure.





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Fig. 5. (A) Absorption spectra of the **RePtr** complex transient in MeCN obtained by flash photolysis experiments at different times after pulse. λ_{exc} = 355 nm (B) Trace of the transient at λ_{obs} = 560 nm.

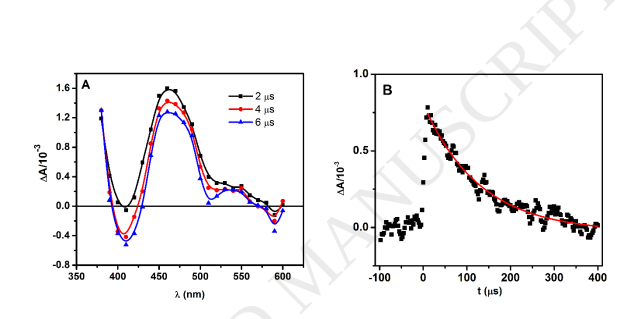


Fig. 6. (A) Absorption spectra of the **RePtr** complex transient in MeCN with TEA 0.1 M obtained by flash photolysis experiments at different times after pulse. λ_{exc} = 355 nm (B) Trace of the transient at λ_{obs} = 440 nm.

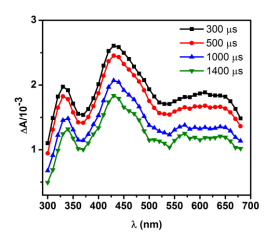
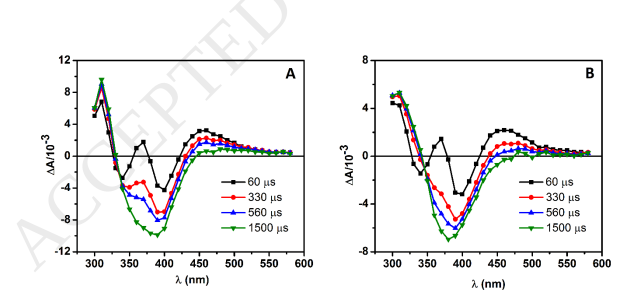


Fig. 7. Absorption spectra of RePtr complex oxidized by $N_3^{\bullet-}$ radical. (aqueous solutions,

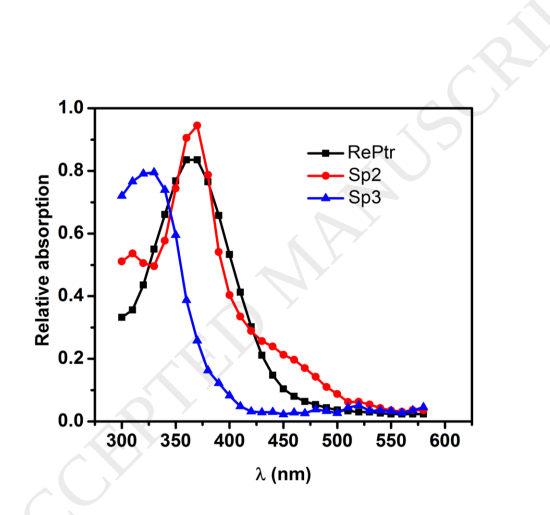
phosphate buffer pH 6, N₂O)

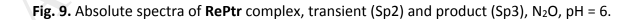


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Fig. 8. Transients absorption spectra of the reduction of RePtr complex complex by (A)

 e_{aq}^{-} and **(B)** CO₂^{•-} (aqueous solutions, phosphate buffer pH = 6, [Na(HCO₂)] = 10⁻² M)





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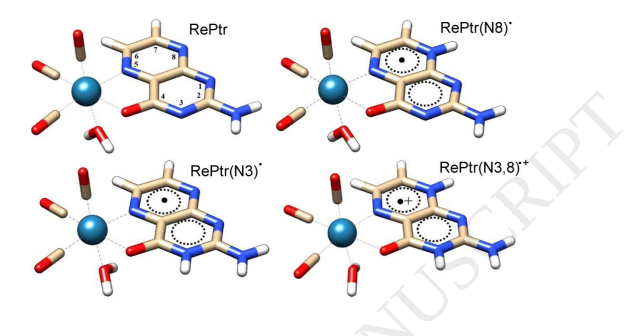


Fig. 10. Absorption spectra obtained by DFT and TDDFT calculations for the RePtr

complex, the protonated radicals RePtr(N8)*; RePtr(N3)* and RePtr(N3,8)*, and RePtrH2.

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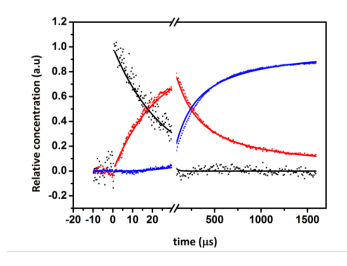
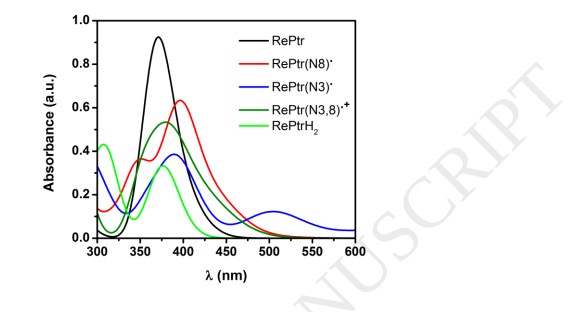


Fig. 11. HS-MCR-ALS concentration profiles and concentration profiles simulated of PR experiment (N_2 or N_2O).

Scheme I. Optimized structures of RePtr, of the protonated radicals at RePtr(N8)[•] and RePtr(N3)[•], and diprotonated radical RePtr(N3,8)^{•+}.

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Solvent	$\lambda_{\max}(nm)$	τ (N ₂)/ns	au (air)/ns	τ (O ₂)/ns	k _q /10 ⁹ M ⁻¹ s ⁻¹	$\phi_{\rm em}({\rm air})(\pm 10\%)$
H ₂ O						
pH = 3	441	7.7	7.4	7.4		6.4 x 10 ⁻²
pH = 6	441	8.7	8.7	8.3		9.1 x 10 ⁻²
pH = 10	454	6.9	6.6	6.6		2.5 x 10 ⁻¹
	418	8.0	8.0			1.7x10 ⁻²
CH₃CN	565	8.0 1.0x10 ³	8.0 2.8x10 ²	 70	1.6	1.9x10 ⁻² (N ₂)

Table 1. RePtr luminescence properties: λ_{max} , lifetimes (λ_{exc} = 341 nm) and luminescence quantum yields in aqueous and acetonitrile solutions. Estimated lifetimes uncertainties

are ± 5%.

	Gas	NaHCO ₂	2-propanol
k1 / s	N ₂	3.67 (± 0.08) x 10 ⁵	3.42 (± 0.07) x 10 ⁵
	N ₂ O	3.70 (± 0.07) x 10 ⁴	3.94 (± 0.05) x 10 ⁴
k ₂ / M.s	N ₂	2.08 (± 0.04) x 10 ⁸	2.59 (\pm 0.03) x 10 ⁸
	N ₂ O	1.60 (± 0.03) x 10 ⁸	4.34 (± 0.05) x 10 ⁸

Table 2. Absolute rate constants obtained from a Hard-Soft MCR-ALS model applied to PR experiments carried out with **RePtr** complex under different experimental conditions. See text for details.

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