Comprehensive investigation of the triplet state electronic structure of free-base 5,10,15,20-tetrakis(4-sulfonatophenyl)porphyrin by a combined advanced EPR and
Comprehensive investigation on the triplet state electronic structure of free-base 5,10,15,20-tetrakis(4-sulfonatophenyl) porphyrin by a combined advanced EPR and theoretical approach

Cite as: J. Chem. Phys. 152, 000000 (2020); doi: 10.1063/1.5131753
Submitted: 15 October 2019 • Accepted: 19 December 2019 • Published Online: XX XX XXXX

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Note: This paper is part of the JCP Special Topic on Spin Chemistry.

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ABSTRACT
The nature of the photoexcited triplet state of free-base 5,10,15,20-tetrakis(4-sulfonatophenyl)porphyrin (H2TPPS4−) has been investigated by advanced Electron Paramagnetic Resonance (EPR) techniques combined with quantum chemical calculations. The zero-field splitting (ZFS) parameters, D and E, the orientation of the transition dipole moment in the ZFS tensor frame, and the proton hyperfine couplings have been determined by magnetophotoselection-EPR and pulse electron-nuclear double resonance spectroscopy. Both time-resolved and pulse experiments exploit the electron spin polarization of the photoexcited triplet state. Comparison of the magnetic observables with computational results, including CASSCF calculations of the ZFS interaction tensor, provides an accurate picture of the triplet-state electronic structure. The theoretical investigation has been integrated with a systematic analysis on the parent free-base porphyrin molecule to assess the effect of the sulfonatophenyl substituents on the magnetic tensors. Additionally, the magnetophotoselection effects are discussed in terms of tautomerization in the excited singlet state of H2TPPS4−.

I. INTRODUCTION

Porphyrins and their derivatives constitute an important class of biomolecules because of their involvement in several fundamental biochemical processes. To date, the main application of these compounds has been as model biosystems and, interestingly, also in the field of applied sciences. Porphyrin-based materials are good candidates for molecular electronic devices, new solar cells, and they have been shown to function as components of molecular and photonic wires and in nonlinear optical materials, to give only few significant examples. Among the properties that have received much attention in recent years, there is the ability of porphyrinoid molecules to participate in energy-transfer and electron-transfer reactions, playing a central role in these photoinitiated processes in a similar manner as chlorophyll and bacteriochlorophylls act in the primary events of photosynthesis. Inspired by natural photosynthetic antenna complexes and reaction centers, porphyrin-based systems are being designed and investigated for applications in artificial photochemical conversion and storage of solar energy. Water-soluble porphyrins are considered particularly suited for such applications. Among selfassembling chromophores, they are gaining a relevant...
position as building blocks for the design of chlorosome-mimicking antennas due to the possibility of fine-tuning of their spectroscopic and structural properties by conveniently modifying their sidechains. A smart technique to control the aggregation process of tetraphenylporphyrin derivatives involves functionalization of the side chains with charged groups and the fine regulation of the pH of the solvent, as in the case of the anionic free-base 5,10,15,20-tetrakis(4-sulfonatophenyl)porphyrin (H$_2$TPPS$_4^-$), which forms J-aggregates in acidic aqueous solutions. Determining the extent to which the H$_2$TPPS$_4^-$ moiety may or may not be effective as a model compound in artificial photosynthesis requires a detailed understanding, and hence an extensive characterization, of its photophysical properties in terms of excited states. While the singlet excited states have been thoroughly investigated also from the computational point of view, less is known about the triplet state due to the dark nature of this state that precludes the application of most of the optical spectroscopies. Photogenerated triplets are key reaction intermediates in many light-induced processes and they show encouraging features that can be exploited in the field of solar light harvesting.

A time-resolved Electron Paramagnetic Resonance (TR-EPR) investigation on the triplet state of H$_2$TPPS$_4^-$ in different monomeric and aggregated forms has shown variations in the magnetic properties going from one species to the other, which have prompted us to further investigate the free-base H$_2$TPPS$_4^-$ (the molecular structure is shown in Fig. 1) by advanced magnetic resonance and theoretical methodologies.

TR-EPR is a powerful technique for the investigation of photoexcited triplet states and extensive studies employing this spectroscopy have been performed on porphyrins. The key data are the zero-field splitting (ZFS) parameters $D$ and $E$, which are sensitive indicators of the spatial extension and symmetry of the two singly occupied molecular orbitals of the triplet state. In addition, TR-EPR allows us to determine the orientation of this interaction within the molecule, probing the geometrical relation between the optical transition dipole moment (TDM) of the molecule and the

![Figure 1](image-url)

**FIG. 1.** (a) Molecular structure of H$_2$TPPS$_4^-$ with the corresponding TDM directions for the Q$_x$ and Q$_y$ transitions. (b) Energy diagram of the triplet spin sublevels, for $D > 0$ and $E < 0$, as a function of the magnetic field $B_0$, with the field vector parallel to the principal axes of ZFS. The arrows indicate the allowed EPR transitions according to the selection rule $\Delta m_S = \pm 1$. (c) Room-temperature absorption spectrum of H$_2$TPPS$_4^-$ in ethanol/methanol 3:2 solution, with the names of the absorption bands according to the Gouterman model. (d) Isotropically excited TR-EPR spectrum of H$_2$TPPS$_4^-$ (black) and corresponding simulation (light blue). The simulation parameters are reported in Table I.
resulting triplet state by using linearly polarized light for excitation. In the past, such magnetophotoselection experiments were performed on aromatic molecules and were crucial in revealing details of the electronic structure of the main chromophores taking part in early events of photosynthesis. Magnetophotoselection has been recently reproposed to study porphyrin model compounds.

The ZFS parameters are integral properties of the triplet wavefunction and depend on its overall spatial distribution. More specific information about the electron distribution in the HOMO and LUMO orbitals is obtained from the electron-nuclear hyperfine coupling (hfc) tensors, from which the distribution of the unpaired electrons of the triplet state is obtained. In contrast to the large body of EPR and Optically Detected Magnetic Resonance investigations yielding D and E parameters and, in the case of TR-EPR, also the electron spin-polarization pattern, there are only a few studies on the hyperfine structure of the triplet state of porphyrins resolved by Electron-Nuclear Double Resonance (ENDOR) spectroscopy. Pulse ENDOR, combined with repetitive laser excitation, takes advantage of the large spin polarization of the triplet state being present only in the first few microseconds after the laser pulse.

Detailed information on the structure of the porphyrin triplet state may be derived by comparison of experimental ZFS parameters and proton hfc values (hfc) with the results of quantum mechanical calculations. Nowadays, computational methods are well established for most of the spin Hamiltonian parameters, such as hfc tensors and g tensors. The additional fine structure observed in triplet state spectra due to the electron–electron spin coupling, in contrast to the otherspinHamiltonian terms, has been addressed only for small molecules and there is not a consolidated approach for its quantitative analysis. A limited number of theoretical studies undertaking the calculation of the ZFS tensor of porphyrins has been reported so far and, to the best of our knowledge, no one based on multiconfiguration post HF methods for this important class of heterocyclic tetrapyrrolic compounds.

In this contribution, a comprehensive study of the triplet state of H2TPPS4− is provided by combining the advanced EPR techniques, best-suited for the triplet state investigation, and state-of-the-art computational methods, which permit accurate calculation of the magnetic observables, i.e., ZFS parameters and proton hfc. In parallel, the free-base porphyrin (H2P) molecule, which is the parent compound, will be addressed to stress the similarities and differences due to the substituents in H2TPPS4−. The nice agreement between the experimental and computed hfc and ZFS parameters, including the ZFS tensor orientation with respect to the TDM, allows us to propose a reliable and detailed picture of the triplet wavefunction of a representative member of this important class of molecules. This is essential for designing novel artificial systems and for developing further applications, which exploit the photophysics of the triplet state.

II. MATERIALS AND METHODS

A. Sample preparation

H2TPPS4− was purchased from Sigma Aldrich and used without further purification. All the H2TPPS4− solutions used for spectroscopic characterization were freshly prepared by dissolving the powder in a mixture of ethanol and methanol 3:2. The concentration of the solutions was determined by optical absorption spectroscopy and has a final value of about 60 μM. The choice of using a mixture of ethanol and methanol for the monomeric samples has the important advantage of generating a good glass when frozen.

Samples for EPR were degassed by freeze and thaw cycles, inserted into quartz EPR tubes (2.4 mm inner diameter, 4 mm outer diameter), sealed under vacuum, and immediately frozen in liquid nitrogen. Samples were stored in liquid nitrogen until measurements.

B. EPR measurements

TR-EPR spectra were recorded at X-band on a Bruker Elexsys E580 spectrometer equipped with a dielectric cavity thermostated at 80 K with a cold nitrogen vapor flow inside a CF935 cryostat. The microwave power used for the TR-EPR experiments was about 1.5 mW. TR-EPR spectra were recorded in direct detection mode without applying any field modulation or phase-sensitive detection. The EPR signals were collected from the microwave preamplifier (bandwidth 20 Hz–6.5 MHz) and sampled with a LeCroy 9360 oscilloscope (10 ns per point). The time resolution of the spectrometer has been estimated to be about 900 ns, mainly due to the cavity response. Laser excitation of the samples was performed using an optical parametric oscillator (OPO) pumped by the third harmonic of a Nd:YAG laser (Quantel Rainbow), operating at a repetition rate of 10 Hz. Laser pulses were 5 ns long, with energies of 3.5 mJ, and a wavelength of 520 nm. Two different polarizations of the light were employed for the magnetophotoselection experiments: one with the electric field perpendicular and the other parallel to the static magnetic field of the spectrometer. The rotation of the polarization plane of the light was obtained using a half waveplate; a linear polarizer was added near the optical window of the cavity for a better control of the polarization. To record the isotropic TR-EPR spectrum, a solution frozen to form a bad milky glass (therefore composed of microcrystallites) was used in order to obtain depolarization by multiple scattering events.

X-band pulse EPR experiments were conducted on a Bruker Elexsys E580 equipped with a dielectric ENDOR cavity (EN4118X-MD4) –9.5 GHz. The temperature was maintained at 20 K with a helium cryostat with optical access (Oxford CF935) driven by a temperature controller (Oxford ITC503). The radiofrequency (RF) amplifier was ENI A-500W. Pulsed laser excitation at 532 nm (5 mJ per pulse and repetition rate of 10 Hz) was provided by the second harmonic of a Nd:YAG laser (Quantel Brilliant). Field-swept electron spin echo spectra were recorded using a two-pulse echo sequence according to the scheme: flash–DAF−π/2−τ−τ−echo (DAF = delay after flash). Mims ENDOR experiments were recorded using the microwave pulse sequence, flash–DAF−π/2−τ−π/2−T−π/2−echo, with 16 ns pulse duration, in conjunction with an RF pulse of 6 μs duration located during the delay T and starting 1.2 μs after the second microwave pulse. The delay τ was variable and the time T was 8.8 μs, long enough to accommodate the RF pulse. Mims ENDOR spectra were recorded at three different τ values (120, 180, and 240 ns) and added together to eliminate τ-dependent blind spots. Pulse ENDOR spectra were accumulated for ~5 h.
C. Spectral analysis

TR-EPR data were processed by subtracting: (i) the average baseline before the trigger event and (ii) a time profile taken at an off-resonance position. Spectra were extracted as single time points at about 1.4 μs after pulsed laser excitation.

Simulations of triplet spectra have been performed using a home-written MATLAB program. For the isotropically excited TR-EPR spectrum, the parameters of the simulations include the g and ZFS tensors and the relative triplet sublevel populations. For the magnetophotoselection TR-EPR spectra, in order to take into account the photoselection of specific orientations, a probability function is also included.

TR-EPR spectra recorded for light polarized parallel and perpendicular to the magnetic field direction were analyzed according to Ref. 23. Here, only a brief summary of the procedure is presented.

When a polarized light source is used to photoexcite the sample, the probability to excite each molecule depends on the angle between the TDM and the direction of the light polarization. Assuming that, after the light excitation, an efficient intersystem crossing (ISC) takes place, the orientational distribution of triplet states inside the sample is uneven and depends on the direction of the TDM in the ZFS reference frame. For the analysis of the TR-EPR spectra in the presence of magnetophotoselection, it is convenient to express the probability $p$ of exciting a specific molecular orientation as a function of the three Euler angles, $(\alpha, \beta, \gamma)$, between the laboratory frame and the ZFS axes. For a specific orientation of the TDM in the ZFS frame, defined by the angles $\alpha$, $\phi$, the excitation probability is

$$p_{\text{par}/\text{perp}}(\alpha, \beta, \gamma) = (\hat{u}_{\text{TDM}} \cdot \hat{a})^2/N,$$

(1)

where $\hat{u}_{\text{TDM}}$ and $\hat{a}$ are the unit vectors along the direction of the TDM and the electric field component of the light. This probability function takes into account that molecules with different orientations are excited when the polarization of the light is varied from a parallel to a perpendicular configuration with respect to the static magnetic field.

The spectrum is calculated as

$$I(B_0)_{\text{par}/\text{perp}} = \sum_{i,j} \int_0^{2\pi} \int_0^{\pi} G(B_{\text{res}}(\alpha, \beta) - B_0) \cdot P_+(\alpha, \beta, B_0) \cdot \left[ \int_0^{\pi} p_{\text{par}/\text{perp}}(\alpha, \beta, \gamma) \sin \beta \, d\beta \right] \sin \beta \, d\beta,$$

(2)

where $G(B_{\text{res}}(\alpha, \beta) - B_0)$ is a Gaussian line shape function centered at the resonance field $B_{\text{res}}$, $P_+(\alpha, \beta, \gamma)$ is the non-Boltzmann population difference between the two resonant states, and $p_{\text{par}/\text{perp}}(\alpha, \beta, \gamma)$ is the excitation probability specific for the parallel and perpendicular configuration. The analytical expressions obtained for the integral within the square brackets for the two orientations of the light polarization with respect to the static magnetic field are reported in the supplementary material.

In the simulation of the magnetophotoselection data, depolarization effects are taken into account by allowing a low-weight contribution from isotropic light excitation. These effects are mainly due to multiple reflections inside the cavity and scattering of the light.

For the Mims ENDOR experiments, the values of the proton hfs along the principal axes of the ZFS tensor were determined by Gaussian deconvolution of the ENDOR spectra.

D. Computational details

All calculations were carried out with the Amsterdam Density Functional (ADF) suite of programs. 48-50 The hybrid B3LYP functional was used 48-50 combined with a triple-ξ quality basis set of Slater-type functions (TZ2P) augmented with two sets of polarization functions. The innermost orbitals of each element were kept constant during the SCF cycles (frozen core approximation); up to 1s for carbon, nitrogen, oxygen, and sulfur. Dispersion effects were included using the correction proposed by Grimme, i.e., D3 with a BJ damping function. 45 This level of theory is denoted B3LYP-D3(BJ)/TZ2P. First, geometry optimizations of the archetypal $\text{H}_2\text{P}$ and of $\text{H}_2\text{TPPS}^-$ were performed using D$_{2h}$ symmetry constraint.

TD-DFT calculations were carried out on the optimized geometries using all-electron TZ2P basis sets for all the atoms. The approximate xc potential obtained with the statistical averaging of (model) orbital potentials (SOAP) was used to calculate the excitation energies, which is suitably designed with a correct asymptotic behavior 48,49 and has been employed with success to investigate excited state properties.

For the calculation of the hyperfine tensor, zero field splitting tensor, and spin density, both B3LYP-D3(BJ) and BP86 51-53 functionals were used combined with all-electron TZ2P basis set.

The scalar zero order regular approximation (ZORA) 45-48 was employed to account for relativistic effects. These levels of theory are denoted ZORA-B3LYP-D3(BJ)/TZ2Pae and ZORA-BP86/TZ2Pae, respectively. No appreciable differences were found and so only data computed at the former level of theory will be discussed in the text. In order to investigate the effect of different orientations of the phenyl rings, geometries were manually built starting from the $D_{2h}$ optimized structures with the rings at 30° and 60° with respect to the tetrapyrrolic plane, without altering the porphyrin core. In addition, for these dihedral values (which are explained in detail in Fig. S9), different mutual orientations of the rings were considered. The magnetic observables were computed for these geometries at the ZORA-BP86/TZ2Pae level of theory.

Due to the unsatisfactory results for the D tensor, ascribed mainly to the limitations of the single-determinant DFT approach, CASSCF calculations were carried out to compute the zero field splitting tensor, as implemented in ORCA software package. 59 The structures obtained by D$_{2h}$ constrained geometry optimizations at the B3LYP-D3(BJ)/TZ2P level were used. MP2 natural orbitals from the unrelaxed density matrix were generated and employed as starting guess. The def2-TZVP basis set was used for all the atoms and def2-TZVP/C was used for the RI approximation. In the CASSCF calculations, def2/j in conjunction with RIJCOSX was employed to approximate the Fock operator for $\text{H}_2\text{TPPS}^-$. A CAS(4,4) active space was chosen including the two π and the two π* frontier molecular orbitals of $\text{H}_2\text{P}$ and $\text{H}_2\text{TPPS}^-$. The sets of four MP2 natural orbitals are shown in Figs. S1 and S2. The spin orbit contribution to the D tensor was computed, but, as expected, it resulted to be negligible; thus, the values reported and discussed in the text correspond to the spin-spin contribution.
III. RESULTS

A. Time-resolved EPR and magnetophotoselection

Magnetophotoselection is a phenomenon observed for photosexcited triplet states, generated by light polarized parallel or perpendicular to the magnetic field, making the relative intensities of the EPR lines dependent on the light polarization. It is a consequence of the fact that the EPR intensities depend not only on the population difference between the triplet sublevels, but also on the nonuniform spatial distribution of the molecules due to selective excitation. Magnetophotoselection has been used in a limited number of cases to attain quantitative information on the relative orientation of the ZFS tensor axes with respect to the optical transition moment, if one of the two is known. 

For H$_2$TPPS$^{3^-}$, having a D$_{2h}$ symmetry, the two absorption Q bands are not degenerate and this results in two resolved sets of peaks exhibiting a vibronic structure, with both the 0–0 and the 0–1 transitions visible in the absorption spectrum as shown in Fig. 1(c). The Q$_r$ and Q$_l$ transitions are characterized by a TDM aligned along and perpendicular to the NH bonds, respectively, as reported in the literature and confirmed by the present DFT calculations (see Table S2 in the supplementary material). This configuration is opposite to the one reported for H$_2$TPP$^{3^{-}}$. The overall set of optical properties calculated for H$_2$TPPS$^{3^-}$ is summarized in the supplementary material as they are functional to the calculations of the magnetic properties derived from EPR-magnetophotoselection. The magnetophotoselection experiments were performed by exciting the Q$_r$ transition by irradiating at 520 nm, where both the Q$_r$ and Q$_l$ transitions and the J-aggregates have negligible absorption.

At first, the isotropically excited TR-EPR spectrum was obtained from the frozen solution of H$_2$TPPS$^{3^-}$ (see the experimental details in the Materials and Methods section and Fig. S7). Then, the TR-EPR spectra recorded after excitation of H$_2$TPPS$^{3^-}$, in a transparent glass carefully prepared to avoid cracks, with light polarized parallel and perpendicular to the magnetic field, were obtained and they are shown in Fig. 2. Since the experimental set-up assured the same excitation conditions, the spectra are displayed without renormalization and show an evident magnetophotoselection effect, with enhanced X and Y triplet transitions in the parallel spectrum and enhanced Z transitions in the perpendicular spectrum. The spectra are also different from that obtained by isotropic irradiation. To further confirm isotropic excitation of the molecules, we have compared the spectrum lineshape obtained with isotropic irradiation with the one calculated as the sum of the experimental spectrum recorded after excitation with polarization of the light parallel to the external magnetic field and twice the spectrum obtained with perpendicular polarization, in analogy to optical polarization measurements. A nice matching of the lineshape of

![FIG. 2. (a) TR-EPR spectra of H$_2$TPPS$^{3^-}$, recorded at 80 K, using an excitation light polarized parallel (blue) and perpendicular (red) to the magnetic field. (b) Simulations of the TR-EPR spectra of H$_2$TPPS$^{3^-}$ with polarization of the light parallel (light blue) and perpendicular (light red) to the magnetic field. The simulation parameters are reported in Table I. The black arrows represent the enhancement or the reduction of the signal when going from parallel to perpendicular excitation at the canonical positions.]

**TABLE I. ZFS parameters, relative triplet sublevel populations, and TDM orientation in the ZFS axes system determined through simulation of the TR-EPR spectra.**

<table>
<thead>
<tr>
<th></th>
<th>D (10$^{-4}$ cm$^{-1}$)</th>
<th>E (10$^{-4}$ cm$^{-1}$)</th>
<th>$p_x$</th>
<th>$p_y$</th>
<th>$p_z$</th>
<th>$\omega$ (deg)</th>
<th>$\psi$ (deg)</th>
<th>Weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simulation</td>
<td>391</td>
<td>–76</td>
<td>0.3</td>
<td>0.6</td>
<td>0.1</td>
<td>90</td>
<td>0</td>
<td>50</td>
</tr>
<tr>
<td>Computation</td>
<td>208</td>
<td>5</td>
<td>…</td>
<td>…</td>
<td>…</td>
<td>90</td>
<td>90</td>
<td>50</td>
</tr>
</tbody>
</table>

$^a$The D and E parameters and $p_x$, $p_y$, $p_z$ were obtained from simulation of the isotropically excited TR-EPR spectrum. $\omega$, $\psi$, and weight were extracted from simulation of the TR-EPR spectra recorded in the magnetophotoselection experiments. $\omega$ is the angle between the Z axis of the ZFS axes system and the TDM, and $\psi$ is the angle between the X axis of the ZFS axes system and the TDM projection into the XY plane (see Fig. 2). Weight is the contribution of each set of angles to the simulation.

$^b$Level of theory: CASSCF(4,4)/def2-TZVP/B3LYP-D3(BJ)/TZ2P.
the experimental and the calculated spectra was obtained, as shown in Fig. S7. Spectral simulations were performed in order to define quantitatively the geometrical relation between the TDM and the ZFS tensor axes in the magnetophotoselection experiments. The key parameter of the simulation is the orientational distribution function reported in Eq. (1), while D, E, and the relative triplet population rates are fixed parameters and are derived from the isotropically excited spectrum. The simulations exhibit not only the correct lineshape, but also the experimental trend in terms of relative intensities. Unexpectedly, the outcome of the global fitting is that the distribution function is the sum of two contributions, with either the principal axes X and Y of the ZFS system aligned along the TDM with an equal weight and an estimated error of 5° at most (for details see Fig. S8 in the supplementary material). A single contribution is unable to reproduce all the features of the spectra and the intensity variations from the parallel to the perpendicular configuration. No optical TDM is associated with the triplet Z axis, confirming the assignment of this axis to the out-of-plane direction and the positive value of D for H$_2$TPPS$^{4-}$, as found for other free-base analogs.\(^{65}\) Table I summarizes the outcome of the spectral analysis, comparing the results on the ZFS tensor with those calculated using the CASSCF method. The orientation of the ZFS tensor in the molecular frame, as derived from computations, is displayed on top of the optimized geometry of H$_2$TPPS$^{4-}$ in Fig. 3.

B. Triplet state $^1$H ENDOR

Triplet state $^1$H ENDOR permits measurement of the hfc tensor component in the reference frame of the ZFS tensor, since it is characterized by a strong orientation selection. For each EPR canonical transition, there are two ENDOR resonance frequencies at

$$V_{\text{ENDOR}}(\pm 1) = V_i \pm A_i \text{ with } i = X, Y, Z,$$  \hspace{1cm} (4)

where $V_i$ is the free-proton nuclear Larmor frequency, $A_i$ is the sum of the Fermi-contact interaction, $A_{ic}$, and the dipolar contribution.\(^{21,67}\)

It can be seen from these equations that the resulting triplet-state ENDOR spectrum is asymmetric, providing the further advantage of allowing us to determine the sign of the hfc relative to the sign of the D value. The line at resonance frequency $V_i$ is common for all protons and, consequently, attains a high intensity.

The $^1$H Mims ENDOR spectra recorded for H$_2$TPPS$^{4-}$ at two of the canonical field positions, Y$^\prime$ and Z$^\prime$, in protonated and deuterated solvent, are shown in Figs. 4(a) and 4(b). In the corresponding insets, the echo-detected spectrum is reported in order to highlight the working field positions. Since the D value for H$_2$TPPS$^{4-}$ is known to be positive from CASSCF calculations, the sign of the proton hfc is assigned as indicated on top of the ENDOR spectra. The hfc along the Y and Z axes of the ZFS tensor were determined through Gaussian deconvolution, and the results are shown in Table II. The spectral features, which disappear in the spectra of D$_2$TPPS$^{4-}$ in the deuterated solvent, are unambiguously assigned to the NH protons (3) since deuterium exchange takes place for those central protons.

A comparison with the results of the DFT calculations at the ZORA-BP86/YZ2Pae//B3LYP-D3(BJ)/TZ2P level on the excited triplet state of H$_2$TPPS$^{4-}$ has allowed assignment of all the hfc to specific protons. The calculations were performed in the ZFS tensor frame, as the principal hfc tensor components are, in most cases, not collinear with the ZFS tensor. The hfc have been calculated for several different conformers with imposed rotations of the phenyl rings, as shown in Fig. S9. The calculated hfc tensor, reported in Table II, refer to the conformer, in which the phenyl rings have been rotated, counterclockwise-clockwise-clockwise-counterclockwise, by 60° with respect to the tetrapyrrolic plane. The choice of a specific conformer has been guided by the matching between the calculated and experimental hfc. However, it has emerged from the analysis that, while the angle between the tetrapyrrole macrocycle and the phenyl substituents is affecting the hfc significantly, the mutual orientation of the rings is not relevant. The conformations with the phenyl rings at 90° with respect to the tetrapyrrole plane shows a number of ENDOR peaks smaller than that experimentally observed. Finally, the conformers with the rings at 30°, beyond being unstable from an energetic point of view, present strong deviations from the experimental hfc in the Z components of H (4).\(^{68}\) The results for all the conformations are summarized in Table S5 and the comparison between the experimental and calculated hfc is reported in Fig. S10.

DFT calculations predict a single hfc for the H (1) and H (2) protons, while experimentally two hfc can be attributed to the H (1) protons. This can be clearly seen in the ENDOR spectra corresponding to the Z transition and for this reason the deconvolution of the corresponding peaks requires a large Gaussian component or even two components. In the literature, the symmetry lowering from D$_{6h}$ to C$_{3h}$ was ascribed to interactions with the solvent or to the admixture of a second triplet state.\(^{69,70}\)

The trend of the computed $A_z$ components is in excellent agreement with those determined experimentally: the α-protons H (2) have small negative hfc, while the H (1) and the NH (3) have
FIG. 4. X-band pulsed Mims ENDOR spectra of $\text{H}_2\text{TPPS}^-$ (top) and $\text{D}_2\text{TPPS}^-$ (bottom) recorded at 20 K along the $Z^+$ (a) and the $Y^+$ (b) ZFS canonical transitions, as indicated by the red arrows in the field-swept electron spin echo spectra shown in the insets. The ENDOR spectra are displayed together with the best fit (light blue) given by Gaussian functions below the spectra. The signs of the hfcs are shown above the spectra. The attribution of the peaks in the ENDOR spectra is represented according to the numbering scheme shown in (c). The asterisks denote the ENDOR peak arising from residual contributions of other orientations or free-proton lines disappearing after deuteration. (c) Molecular structure of $\text{H}_2\text{TPPS}^-$ and corresponding orientation of the ZFS principal axes and numbering scheme of the protons adopted in this work. (d) Energy of the spin sublevels of a triplet state coupled to a nucleus $I = 1/2$ with positive and negative hyperfine splitting, for the magnetic field parallel to a generic ZFS axis. The gray arrows indicate the EPR transitions for $m_I = -1/2$ (those for $m_I = +1/2$ are omitted), the light blue arrows indicate the ENDOR transitions.
larger negative hfcs. A similar behavior has been observed previously for both the H$_2$P and the free-base tetraphenylporphyrin molecules, indicating that the presence of the (SO$_3$)$_2^-$ substituent does not significantly affect the spin density distribution in the triplet state.

The small hfcs, in the vicinity of the strong Larmor peak, have been assigned to the different types of phenyl protons. Close to $v_{11}$, there is also a contribution from protons with weak dipolar interactions with the unpaired electrons, such as protons in the solvent, which give rise to the "matrix" ENDOR line. These peaks disappear, as expected, in deuterated solvent.

The assignment of the experimental values obtained for $A_Y$, which has been reported in the literature only in the case of H$_2$P and for an incomplete set of protons, is not as straightforward as for the $A_Z$ components. Although the hfc calculated in the Y direction of the ZFS tensor is slightly overestimated for protons (1), even considering different possible conformers, the overall picture of the hfcs is perfectly in line with the experimental results. Small positive values are computed for both the $a$-protons H (2) and the NH (3), while a larger negative contribution is found for the $a$-protons H (1).

Experimentally, the directions of the in-plane principal axes could not be distinguished by magnetophotoselection, as already pointed out in Sec. III A. Furthermore, although CASSCF calculations were performed to improve the accuracy of the ZFS parameters, $E$ remains a critical parameter to be computed as it derives from the difference between two large components of the dipole-dipole coupling tensor. A significant inaccuracy in the estimation of $E$ is accompanied by a potential mismatch of the directions of the associated in-plane principal axes. Indeed, a satisfactory agreement can also be obtained when comparing the ENDOR experimental values with the calculated $A_X$ components, improving the accuracy for protons (1) but at the expense of protons (2). This uncertainty is exacerbated by the impossibility to use the ENDOR data recorded along the $X$ field position for an unequivocal assignment, due to the contamination of residual lines from other orientations. In addition, the orientation selection at the $Y$ field position is not as good as for the $Z$ position, and therefore, hfcs are not as reliable.

From the assigned hfcs, a map of the spin-density distribution of the triplet electrons over the molecule can be obtained. The calculated spin-density plot is depicted in Fig. 3.

### IV. DISCUSSION AND CONCLUSIONS

In this work, we elucidate the electronic structure of the lowest excited triplet state of H$_2$TPPS$^{4-}$. The interpretation requires a combination of magnetic resonance and accurate quantum chemistry methods in order to yield precious information on the triplet wavefunction. This is essential if a deeper understanding of the nature of this excited state has to be attained.

The approach here adopted is the combination of multiple EPR spectroscopic observables and computational results. The experiments allow both ZFS and hfc tensors to be determined, the former with respect to the TDM in the magnetophotoselection experiment and the latter in the ZFS frame using pulsed ENDOR spectroscopy, as presented in the Results section. The existing literature on the EPR of the triplet state of porphyrinoids is complete in terms of ZFS parameters and spin polarization pattern but few EPR-magnetophotoselection experiments are available and no simulations are reported, allowing only a qualitative interpretation of the spectra, which contain important structural information.

For triplet states, in general, due to technical difficulties of performing ENDOR on transient species, only sparse experimental data have been reported on molecules of biological interest, including porphyrinoids. In most cases, including the H$_2$P triplet state, only the $A_Z$ component of the hfc tensor has been measured.

This prompted us to perform this comprehensive investigation. The interest for H$_2$TPPS$^{4-}$ derives from the relevant applications of water-soluble porphyrins due to their ability to form J-aggregates. The accuracy in the study of porphyrin monomers appears as an important prerequisite for their use in aggregates.

The computational investigation on the electronic and magnetic features of the lowest excited triplet state of H$_2$TPPS$^{4-}$ was preceded by a systematic analysis on the parent H$_2$P, which is the basic building block of the porphyrins, to assess the role of the

### TABLE II. Experimental and computed proton hfc tensors of H$_2$TPPS$^{4-}$ in the ZFS frame.

<table>
<thead>
<tr>
<th>Hyperfine coupling (MHz)</th>
<th>H (1)$^a$</th>
<th>H (2)$^b$</th>
<th>H (4)$^c$</th>
<th>H (4)$^d$</th>
<th>H (5)$^e$</th>
<th>H (5)$^f$</th>
<th>NH (3)$^g$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A_X$</td>
<td>-2.17/ -2.75</td>
<td>+0.84</td>
<td>-1.40</td>
<td>-1.00</td>
<td>+0.57</td>
<td>+0.57</td>
<td>+0.28</td>
</tr>
<tr>
<td>$A_Y$</td>
<td>-1.86/ -2.69</td>
<td>+0.67</td>
<td>-1.15</td>
<td>-0.94</td>
<td>+0.67</td>
<td>+0.67</td>
<td></td>
</tr>
<tr>
<td>$A_Z$</td>
<td>-3.96</td>
<td>-0.40</td>
<td>-0.80</td>
<td>-1.28</td>
<td>+0.43</td>
<td>+0.67</td>
<td>-3.24</td>
</tr>
<tr>
<td>$A_{iso}$</td>
<td>-3.93</td>
<td>-0.50</td>
<td>-0.86</td>
<td>-1.30</td>
<td>+0.40</td>
<td>+0.40</td>
<td></td>
</tr>
</tbody>
</table>

$^a$Two values are reported for the $A_X$ component of H (1), because its ENDOR signal has been deconvoluted employing two Gaussian functions.

The only hyperfine coupling available for the NH proton is that of the protonated porphyrin H$_2$TPPS$^{4-}$.

$^c$For each proton, two experimental hfc values are reported: one for the fully protonated porphyrin, H$_2$TPPS$^{4-}$ (top value), and the other for D$_2$TPPS$^{4-}$ (bottom value).

$^d$The computational investigation on the electronic and magnetic features of the lowest excited triplet state of H$_2$TPPS$^{4-}$ was preceded by a systematic analysis on the parent H$_2$P, which is the basic building block of the porphyrins, to assess the role of the
substituents on the magnetic tensors, while the electronic absorptions have been more extensively discussed in the literature, as commented in the following.

For $\text{H}_2\text{P}$, the four-level Gouterman model works well, since the frontier orbitals consist of two almost degenerate filled levels well separated from two almost degenerate empty levels. Based on different DFT methods, these orbitals span the irreducible representation $\text{a}_1$ (HOMO–1), $\text{b}_2$ (HOMO), $\text{b}_3$ (LUMO), $\text{b}_8$ (LUMO+1), with some changes in the relative order of the filled and the empty levels depending on the potential. CASCCF calculations provided this order: $\text{b}_8$ (HOMO–1), $\text{a}_1$ (HOMO), $\text{b}_3$ (LUMO), $\text{b}_2$ (LUMO+1). The lowest computed dipole allowed absorptions have been reported to have $\text{b}_8$, $\text{b}_3$, and $\text{b}_2$ irreducible representations both with TDDFT methods as well as with CASPT2. This assignment is also consistent with the Gouterman four-orbital model, according to which the Q (and higher energy B) bands derive primarily from transitions from the two highest n occupied molecular orbitals into the two n–1 lowest unoccupied orbitals. The spectrum, computed by us in gas phase using the SAOP potential, shows the Q-bands at 2.188 eV (B2u) and 2.312 eV (B3g) with oscillator strengths of 0.17 × 10–4 and 0.12 × 10–3, polarized along the NH (3) pyrrole rings and the N pyrrole rings. The energy values are in rather good agreement with the experimental data, which fall in the range of 1.98–2.02 and 2.33–2.42,60–62 but the splitting is too low if compared to the gas-phase measured value (0.44 eV). The same issue was reported by Baerends and co-workers and was ascribed to the geometry sensitivity. Also, the values of the computed oscillator strengths are geometry-dependent and similar very small values have been reported in the literature. Finally, although Q3 is commonly considered the most intense transition, different computational studies identify Q1, as stronger,74,81 in agreement with our findings. Q1 and Q2 absorptions are ascribed to HOMO–LUMO (54%) and HOMO–1-LUMO+1 (45%) and to HOMO–LUMO+1 (58%) and HOMO–1-LUMO (42%), clearly denoting the mixed character of the lowest singlet excitons, which involve the four Gouterman orbitals. This is in agreement with the analysis of the spin density distribution of the lowest triplet state of $\text{H}_2\text{TPPS}^4$–, which very well describes $\text{H}_2\text{P}$, but represents only one conformer of $\text{H}_2\text{TPPS}^4$– due to the orientation of the phenyl rings, which are forced to be at 90° with respect to the porphyrin plane. Calculations of the optical properties, and in particular the polarization of the Q-bands, are relevant for the magnetophotoselection experiment, because they are employed to assign the relative orientation of the ZFS tensor axes within the molecule.

We extended the calculations to derive the ZFS parameters and the spin density distribution of the lowest triplet state of $\text{H}_2\text{TPPS}^4$–, evaluating in parallel the corresponding magnetic observables for $\text{H}_2\text{P}$, for which experimental and theoretical data are already available. While in the main text the focus is on $\text{H}_2\text{TPPS}^4$–, the outcome of the calculations for $\text{H}_2\text{P}$ is reported in the supplementary material.

The analysis of the frontier molecular orbitals and of the lowest excited states of $\text{H}_2\text{P}$ and $\text{H}_2\text{TPPS}^4$– pointed out that, in both cases, a single Slater determinant description of the triplets might be inadequate and this evidence must be taken into account when calculating the magnetic properties. Particularly, if the DFT approach gives spin density values and hyperfine tensors in nice agreement with the ENDOR experiment, the ZFS tensor is more sensitive to the wavelength. We started our analysis employing a DFT approach. The geometries of $\text{H}_2\text{P}$ and $\text{H}_2\text{TPPS}^4$– were both optimized at the B3LYP-D3(BJ)/TZ2P level of theory in the triplet state. The hyperfine and ZFS tensors were also computed at the ZORA-B3LYP-D3(BJ)/TZ2P level as well as at the ZORA-BP86/TZ2P level, but no significant differences were found when comparing the results of the two approaches for $\text{H}_2\text{P}$.

The magnetic results were first obtained for D3h constrained geometries. In order to investigate the effect of different orientations of the phenyl rings, two structures were obtained manually by imposing orientations of 30° and 60° with respect to the tetrapyrrolic plane while maintaining the geometry of the porphyrin core frozen. Importantly, the rings can be displaced in different manners and thus more conformations can be explored. We considered selected cases generated by rotating the rings clockwise (conformers A30 and A60), alternated clockwise and counterclockwise (conformers B30 and B60), and counterclockwise-clockwise-clockwise-clockwise (C30 and C60). Based on the energetics, the conformers with the phenyl rings rotated by 30° were discarded. The conformers with the phenyl rings rotated by 60° are the lowest energy structures, although the D3h constrained geometry with the rings at 90° lies only 1 kcal mol–1 above. Among the 60° structures, C60 is the most stable, although by less than 0.5 kcal mol–1 at the employed level of theory. The hfs of all the three conformers with the phenyl rings at 60° are found to be in excellent agreement with the experimental ENDOR data, especially for the Z component, which is also the most reliable direction due to the strong orientation selection that can be achieved. On the other hand, comparison with the ENDOR data provided further evidence that all the 30° conformations are unlikely to be present.

The combination of DFT calculations and ENDOR experiments allowed us to draw an accurate picture of the spin density distribution of the $\text{H}_2\text{TPPS}^4$– triplet state. Compared to the $\text{H}_2\text{P}$ parent compound (Fig. S4), even if a small amount of spin is the investigation of the magnetic properties and their relation to the triplet wavefunction. As an efficient approach, we imposed D3h symmetry to our systems, which very well describes $\text{H}_2\text{P}$, but represents only one conformer of $\text{H}_2\text{TPPS}^4$– due to the orientation of the phenyl rings, which are forced to be at 90° with respect to the porphyrin plane. Calculations of the optical properties, and in particular the polarization of the Q-bands, are relevant for the magnetophotoselection experiment, because they are employed to assign the relative orientation of the ZFS tensor axes within the molecule.
density is moved to the phenyl rings, the overall distribution on the
tetrapyrrole macrocycle is conserved.

While the values of the $A$ tensors are in good agreement with
the experiment, in the case of $\text{H}_2\text{P}$ even slightly improving the
results reported in the literature, the computed ZFS tensor ele-
ments were unsatisfactory. Thus, we considered CASSCF as the
method of choice to improve the wavefunction. For the CASSCF
multiconfigurational approach, the inclusion of the whole $\pi$ system
of the parent $\text{H}_2\text{P}$ is already computationally demanding (24 orbitals
and 26 electrons), as extensively commented by Werner. As the
Gouterman four orbitals model has proved to be robust, we decided
to define an active space formed by the four frontier orbitals, ranging
from HOMO–1 to LUMO+1, which in $\text{H}_2\text{P}$ as well as in $\text{H}_2\text{TPPS}^\pi$
have strong contribution from the $\pi$ porphyrin core. MP2 natural
orbitals were generated as guess for the CASSCF calculation. For
$\text{H}_2\text{P}$, we computed two triplet states, i.e., $^3\text{B}_2u$ and $^3\text{B}_4u$, and the former was found to be more stable, in agreement with the liter-
ature. For $\text{H}_2\text{TPPS}^\pi$, we computed $^3\text{B}_2u$ and $^3\text{B}_4u$, and the latter
turned out to be more stable. The ZFS tensor was calculated using
the CASSCF wavefunction, showing that the spin-orbit component
is negligible, as already pointed out.

To the best of our knowledge, this is the first example of
calculation of the ZFS tensor based on a CASSCF calculation for
porphyrins. Further improvement might be obtained including
dynamic correlation, but the multiconfigurational approach itself
using the small Gouterman active space has provided good semi-
quantitative results.

A number of relevant aspects about the electronic structure of
the triplet state were revealed based on the comparison between
$\text{H}_2\text{TPPS}^\pi$ and its parent compound $\text{H}_2\text{P}$. As a main result, while
the ZFS parameters vary significantly, the ZFS tensor orientation is
conserved and the triplet electron spin density distribution is found
to be similar. The $D$ parameter is about 10% smaller in $\text{H}_2\text{TPPS}^\pi$
than in $\text{H}_2\text{P}$. In contrast, the $A_2$ hfcs show differences of about
3%, as already pointed out in analogous comparison between the
free-base tetraphenyl porphyrin and $\text{H}_2\text{P}$. The excellent agreement
between the experimental and calculated hfcs provides a valida-
tion of the triplet wavefunction. Calculations and experiments also
agree in indicating that the presence of the sulfate groups does
not significantly alter the electronic features, as already highlighted
when introducing phenyl substituents to $\text{H}_2\text{P}$. This outcome is
important, since the polar groups are introduced to provide solu-
bility and should not cause unwanted variations of the electronic
structure.

When comparing ZFS parameters, although they differ quite
significantly in absolute values from the calculated ones, the 10%
decrease in the $h^\text{i}_{\pi}\text{TPPS}^\pi$ with respect to the parent com-

pound is also fulfilled, proving that CASSCF calculations, describing
correctly the multiconfigurational nature of porphyrins, are more
suitable than DFT to predict the ZFS interaction tensors. The discre-
pancies in the absolute values are likely due to the limited active
space and to the lack of dynamic correlation. The difference in the
sensitivity of the ZFS parameters and the hfcs to mesophenyl sub-
stitution finds confirmation in this work. The ZFS tensor is very
sensitive, because the frontier orbitals are mainly contributing to this
interaction and upon substitution part of the electron density gets
slightly delocalized on the phenyl rings. Conversely, the mechanism
of spin polarization governing the isotropic contribution to the hfcs
derives from spin density on the core orbitals, which are unaffected
by substitution effects.

In the magnetophotoselection experiments, an important
aspect regarding the structural properties of $\text{H}_2\text{TPPS}^\pi$ has also
emerged. As already pointed out in the Results section, the analy-
sis of the corresponding EPR spectra has shown that the principal
axes $X$ and $Y$ of the ZFS system are aligned along the TDM but
unexpectedly the best fit has been obtained with an equal con-
tribution of both ZFS axes, although optical excitation was performed
on the main vibronic band of the $Q_1$ transition. The apparent excit-
tation of both $Q_1$ and $Q_2$ transitions with an equal contribution
definitely does not match to the relative absorption of the two trans-
itions at the excitation wavelength, and therefore, the result cannot
be associated with a direct absorption process. In the quest for the
mechanism that accounts for this effect, we first considered and
later excluded energy transfer processes in fixed-geometry assem-
bles, like in the typical J-aggregates aggregation. The energy transfer
from one molecule to another, within the stack, and with a 90° rela-
tive rotation of the ZFS system is expected to that generated by a pseudo $Q_1$-excitation.

This occurrence was excluded, first because there is no excitation of
any absorption band of J-aggregates at the excitation wavelength, which
was carefully selected to avoid any contribution; then, because the
both the polarization and the lineshape of the spectra did not
change with concentration in diluted solutions; finally, because a
population equilibration of the $X$ and $Y$ states is not observed
experimentally.

Alternatively, a pseudorotation of the molecule may be
obtained by a $N\rightarrow H$ tautomerism between two equivalent trans-
structures involving donation of the two $N\rightarrow H$ protons to the unproto-
nated nitrogen atoms, with the cis tautomer only present as a tran-
sitory intermediate. We have interpreted our experimental obser-
vations invoking a fast phototautomerization process occurring in
the excited singlet state, which is slowed down in the triplet state
precluding its observation in the EPR time-scale. The model we
have introduced to explain the magnetophotosellection experiments
is represented in Fig. 5. Photoexcitation in correspondence of the $Q_1$
band generates a distribution of the excited $\text{H}_2\text{TPPS}^\pi$ molecules with respect to the polarization of the light (see the correspond-
ing orientational distribution function in Fig. 5) with a maximum
when the molecules are oriented with the $Q_1$ TDM parallel to the
light polarization. The phototautomerization exchanges the TDM
directions and also the distribution function because of the molecu-
lar pseudorotation; this is equivalent to the excitation of the $Q_1$
band through the tosylate $Q_1$ TDM. When singlet excitation is followed
by ISC to the corresponding triplet state, the two tautomers are
equivalent from the microscopic point of view, namely, the relative
population rates of the three triplet sublevels are the same but their
orientational distribution function is different. This is the reason
why simulation of the spectra obtained by magnetophotoselection
requires that both the in-plane ZFS canonical orientations are ori-
ented along the optical polarization direction. This model accounts
for the presence of two overlapping orientational distribution func-
tions taking into account the fictitious excitation of both $Q_1$ and $Q_2$
bands produced by the phototautomerization in the excited singlet
state. The process should be fast enough to explain the 1:1 popu-
lation of the singlet excited states polarized along X and Y. At the same
phyrins show mainly rhombic symmetry (\(|E| \neq 0\)). Phototautomerization of \(\text{H}_2\text{TPPS}_4\) is highlighted in the present paper. Slower in the triplet state as compared to the excited singlet state, this behavior of this important class of molecules, which must be much more complex, is underway in order to clarify the complex phototautomerization process. Magnetophotoselection experiments, including wavelength, temperature dependence, and deuteration effects, might be envisaged. Magnetophotoselection experiments and direct comparison of the triplet-state EPR and ENDOR data with accurate theoretical methods including CASSCF calculations of the ZFS interaction tensor. This systematic study, providing the overall picture of the triplet state and including the structural issue of tautomerization, paves the route to further triplet EPR studies on acidified and aggregated water-soluble porphyrin molecules.

**SUPPLEMENTARY MATERIAL**

See the supplementary material for computational details, spectral analysis of the magnetophotoselection experiments, and \(^1\)H-ENDOR spectra analysis.

**ACKNOWLEDGMENTS**

This work was supported by the University of Padova (Grant No. PRAT 2014-CPDA145097/14), CINECA (Casalecchio di Reno, Italy) is gratefully acknowledged for providing the supercomputing facilities used for the quantum chemistry calculations through the ISCRA Grant REBEL2 (Redox state role in Bio-inspired EElementary reactions 2), P.I.: L.O. The authors thank E. Collini and C.W.M. Kay for helpful discussions and M. Dalla Tieza and M. Bortoli for software support.

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