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Microwave-Mediated Synthesis of Bulky Lanthanide Porphyrin–Phthalocyanine Triple-Deckers: Electrochemical and Magnetic Properties

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ABSTRACT: Five heteroleptic lanthanide porphyrin–bis-phthalocyanine triple-decker complexes with bulky peripheral groups were prepared via microwave-assisted synthesis and characterized in terms of their spectroscopic, electrochemical, and magnetic properties. These compounds, which were easily obtained under our preparative conditions, would normally not be accessible in large quantities using conventional synthetic methods, as a result of the low yield resulting from steric congestion of bulky groups on the periphery of the phthalocyanine and porphyrin ligands. The electrochemically investigated triple-decker derivatives undergo four reversible one-electron oxidations and three reversible one-electron reductions. The sites of oxidation and reduction were assigned on the basis of redox potentials and UV−vis spectral changes during electron-transfer processes monitored by thin-layer spectroelectrochemistry, in conjunction with assignments of electronic absorption bands of the neutral compounds. Magnetic susceptibility measurements on two derivatives containing TbIII and DyIII metal ions reveal the presence of ferromagnetic interactions, probably resulting from magnetic dipolar interactions. The TbIII derivative shows SMM behavior under an applied field of 0.1 T, where the direct and Orbach process can be determined, resulting in an energy barrier of $U_{eff} = 132.0$ K. However, Cole–Cole plots reveal the presence of two relaxation processes, the second of which takes place at higher frequencies, with the data conforming to a $1/\tau \propto T^7$ relation, thus suggesting that it can be assigned to a Raman process. Attempts were made to form two-dimensional (2D) self-assembled networks on a highly oriented pyrolytic graphite (HOPG) surface but were unsuccessful due to bulky peripheral groups on the two Pc macrocycles.

INTRODUCTION

Triple-decker complexes containing tetrapyrrolic phthalocyanine (Pc) and/or porphyrin (Por) macrocycles have been used in applications related to their characteristic properties, which are more than the sum of their compositional parts. Examples of applications for these types of compounds include molecular-level information storage,1 field effect transistors,2 and single-molecule magnets.3

In an early synthesis of heteroleptic triple-decker complexes with Pc and Por macrocycles,4 the (Pc)Ln(Pc)Ln(Por) unit was shown to be accessible according to the procedure of Weiss.5 In this synthetic method, a presynthesized double-decker (Pc)Ln(Pc) unit reacted with the in situ (Por)Ln(acac)$_3$ (acac = acetylacetonate) formed by refluxing the porphyrin (Por) with excess Ln(acac)$_3$ in 1,2,4-trichlorobenzene (TCB). However, the yield via this procedure was low due to the harsh conditions of high temperature (bp. TCB, 214 °C) and rather...
long reaction times (ca. 12 h). Moreover, this reaction barely occurred when there was steric congestion resulting from bulky groups on the periphery of the porphyrin and phthalocyanine macrocycles in the targeted triple-deckers.6

Mixed porphyrin/phthalocyanine triple-decker compounds of the type (Pc)₃Ln(Pc)₃Ln(Por) are similar to other types of triple-decker tetrapyrrrole complexes in that they exhibit a large number of oxidation states, reversible electrochemistry, and relatively low oxidation potentials.6 However, to the best of our knowledge, there have been no reports in the literature which have assigned the probable sites of electron transfer for oxidation or reduction of these complexes. The magnetic behavior of heteroleptic (Pc)₃Ln(Pc)₃Ln(Por) derivatives has also not been studied in detail,² as is the case for their double- and quadruple-decker counterparts,³ particularly in terms of the effect of the central lanthanide ions on molecular magnetism.

Both of these points are addressed in the current paper, which describes the design and synthesis of five new triple-decker complexes (Pc)₃Ln(Pc)₃Ln(Por) with bulky substituents on the macrocycles, four of which (TD-1—TD-4) are characterized in terms of their spectroscopic and electrochemical properties. The single-molecule magnetic behavior for two of the compounds is also demonstrated.

The structures of the investigated compounds, labeled as TD-1—TD-5, are shown in Scheme 1 and were prepared by microwave irradiation, a practical methodology that minimizes tedious isolation from other scrambled macrocyclic products. By combining spectroscopic and electrochemical methods to characterize the final synthetic products, we are able to discern how the number of undecyl substituents introduced at the meso positions of the porphyrin ring will influence the electronic absorption spectra, redox potentials, and perhaps also the site of electron transfer in the porphyrin–phthalocyanine heteroleptic complexes.

Although we were unable to obtain ordered self-assembled monolayers on a highly oriented pyrolytic graphite (HOPG) surface for imaging by scanning tunneling microscopy (STM) at the liquid–solid interface, we are confident that such complexes may find applications once better grafting protocols have been added to the molecules.

### RESULTS AND DISCUSSION

#### Design and Synthesis

**Design and Synthesis.** +2(5,5'-Trimethyl-1,3-dioxan-2-yl)phthalodinitrile 5,⁶ free-base meso-di- and meso-tetraundecylnaphthalocyanines 7a,b and meso-tetrahexadecylphthalocyanine 7c,¹⁰ were prepared according to published procedures. The conventional Weiss method⁷ was initially employed to prepare the triple-decker complexes TD-1 and TD-2 (Scheme 1). The precursor 5 and Gd(acac)₃·2H₂O were reacted in the presence of dibenzylidenecyclohexane (DBU) to synthesize the double-decker 6a, whose spectrum is shown in Figure S1 in the Supporting Information. The yield of this reaction was comparable to yields reported for similar double-decker complexes.¹⁷

The free-base porphyrin, 7a or 7b, was admixed with Gd(acac)₃·2H₂O and heated to reflux in TCB for 4 h, and 6a was then added after cooling the reaction mixture to room temperature. This was followed by prolonged heating and refluxing for up to 20 h. However, as shown in the low-resolution mass spectra (Figures S2 and S3 in the Supporting Information), only very weak ion peaks attributed to the desired TD-1 or TD-2 products could be detected, while strong ion peaks attributed to the unreacted starting compound 6a are observed. This implies that the reaction occurred but only in a low single-digit yield, making it impossible to collect, purify, and fully characterize these target compounds due to the small amount of the sample obtained by this method.

As mentioned above, heteroleptic triple-decker compounds with bulky groups on the periphery of the porphyrin and/or phthalocyanine macrocycle are obtained in low yield when using the published synthetic method.⁶ For example, Lindsey and co-workers tried but failed to prepare a triple-decker complex utilizing the standard procedure with [(THP)Eu(acac)] (THPH₂ = meso-tetraheptylporphyrin) or [(heptyl)₃Pc]·Eu. The authors invoked steric hindrance of the peripheral alkyl groups on the porphyrin and phthalocyanines to explain the low reactivity.⁶²

To circumvent this drawback, we employed microwave irradiation in a professional oven equipped with a video camera to monitor color changes and/or precipitation, thus avoiding prolonged reaction times. α-Dichlorobenzene (DCB) was chosen as an alternative solvent to trichlorobenzene (TCB) due to its lower boiling point, thus being easier to remove and favoring microwave absorbance. The optimized microwave conditions for stages ii and iii are given in Scheme 1. Under these conditions, TD-1—TD-5 were obtained in relatively larger yields after flash chromatography (38, 33, 28, 36, and 22%, respectively). The double-deckers 6a–c could also be rapidly prepared using microwave irradiation at 140 °C, under the conditions of 15 min, 17 bar, and 250 W. The mass spectrum of the product (see the Supporting Information) shows typical isotopic profiles of the lanthanide atoms (Gd, Tb, and Dy) and clearly confirms the elemental composition of these triple-decker compounds.

**Electronic Absorption Spectra.** Electronic absorption spectra of these triple-decker complexes were recorded in CHCl₃ (Figure 1) and PhCN (Figure S4 in the Supporting Information), and the spectral data are summarized in Table 1.

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**Scheme 1. Synthetic Route for Triple-Decker Complexes TD-1—TD-5**

⁴Reaction conditions: (i) 1-pentanol/DBU, 140 °C, overnight; (ii) DCB, microwave, 180 °C, 10 min, 10 bar, 150 W; (iii) microwave, 220 °C, 30 min, 17 bar, 250 W. Yields are based on the cumulative procedures (ii) and (iii).
The UV–vis absorption characteristics of TD–1–TD–5 are similar to each other due to their similar molecular structures, which are characterized by two phthalocyanines and one porphyrin connected by two lanthanide ions. As seen in Figure 1 and Table 1, the strong bands in the region of spectrum at 341–344 and 400–418 nm in CHCl$_3$ are attributed mainly to the Soret bands of the phthalocyanine and the porphyrin macrocycle, respectively. The Q bands of the compounds at 506–511, 579–584, and 621–634 nm are assigned as arising mainly from the two neighboring Pc ligands in the [(Pc$^2$)$^2$−Ln(Pc$^2$)$^2$−)] part of the molecule. The Q band absorption at 728–749 nm is mainly associated with the Pc ligand, together with some contributions from the Por ligand. These wavelengths are very similar to published wavelengths of analogous monomeric porphyrins and phthalocyanines.

Three of the six bands in Figure 1 and Table 1 vary with the number of meso substituents on the porphyrin macrocycle, while three do not. The Soret and Q bands of TD–1 and TD–2 at 400–406, 621–631, and 728–735 nm in CHCl$_3$ are sensitive to the R$^2$ and R$^3$ substituents on the porphyrin ligand, while the three strong bands at 341–343, 506–507, and 579–580 nm are almost independent of these substituents. This can be seen by comparing the spectra shown in Figure 1 for TD–1 and TD–2, which differ in the number of meso-substituted undecyl groups on the porphyrin macrocycle; TD–1 has two meso-undecyl groups, while TD–2 has four. The three bands of TD–1 at 400, 621, and 728 nm in CHCl$_3$ are shifted to 406, 631, and 735 nm for TD–2 under the same solution conditions. The same trends in electronic absorption spectra are observed in PhCN as well.

A 15 nm difference in the position of the Soret band is seen in Figure S6 in the Supporting Information between monomeric porphyrin $\text{Ta}$ (A$\text{max}$ 404 nm) and $\text{ Tb}$ (A$\text{max}$ 418 nm), which are used to synthesize TD–1 and TD–2 to TD–4, respectively, confirming our assignment of the 400 and 406 nm bands in TD–1 and TD–2 as porphyrin-based. Furthermore, similar positions of the peaks for TD–2 and TD–3 at 343, 406–407, 507–508, 579–581, and 631 nm in CHCl$_3$ indicate a lack of sensitivity of these bands to the lanthanide metal ions of the compounds, in this case Gd and Dy.

From the above analysis, it is thus possible to assign the 400–418 nm band in CHCl$_3$ as being porphyrin-based and all the other bands, at 341–344, 506–511, 579–585, 621–634, and 728–749 nm, as phthalocyanine-based. The same assignments can be made for the absorption bands in PhCN and, in combination with the data from electrochemistry and spectroelectrochemistry, will provide indirect evidence for the site of oxidation or reduction being predominantly porphyrin- or phthalocyanine-based in a specific redox process.

**Electrochemistry.** The electrochemical behavior of TD–1–TD–4 was studied by cyclic voltammetry in PhCN containing 0.1 M TBAP. Under these solution conditions, each compound exhibits four one-electron oxidations and three one-electron reductions (Figure 2), all of which are attributed to ligand-based processes. The measured half-wave potentials are given in Table 2 along with $E_{1/2}$ values taken from the literature for reduction and oxidation of related triple-decker complexes having only phthalocyanine or only porphyrin macrocycles. An illustration showing the structures of these compounds with substituents on the macrocycles is given in Scheme 2.

The reductions of monomeric phthalocyanines are significantly easier than reductions of monomeric porphyrins having the same central metal ions, and the same trend is observed when the electrochemical data for TD–1–TD–4 are compared with those of the related homoleptic Ln$_2$(Pc)$_3$ and Ln$_2$(Por)$_3$ triple-deckers. For example, the three Ln$_2$(OEP)$_3$ derivatives undergo only a single one-electron reduction in CH$_2$Cl$_2$ at $E_{1/2} = -1.67$ V vs SCE, while the three Ln$_2$(Pc)$_3$ complexes exhibit three one-electron reductions within the same potential range, the first of which is located at $E_{1/2} = -1.38$ to $-1.42$ V vs SCE, as seen in Table 2. The first two reductions of Ln$_2$(Pc)$_3$ (compounds TD–6–TD–8) have $E_{1/2}$ values almost identical with those for compounds in

![Figure 1. UV–vis absorption spectra of TD–1–TD–5 recorded in CHCl$_3$ ($c = 8.5 \times 10^{-3}$ mol/L).](image)

| Table 1. Electronic Absorption Data of TD–1–TD–5 in CHCl$_3$ and PhCN with Assignment of Bands to Specific Macrocycles |
|---|---|---|---|---|---|---|---|
| solvent | triple-decker | Ln | Pc | Por | Pc | Pc | Pc |
| CHCl$_3$ | TD–1 | Gd | 341 (5.16) | 400 (4.92) | 506 (4.33) | 580 (4.44) | 621 (4.80) | 728 (4.43) |
| TD–2 | Gd | 343 (5.20) | 406 (4.86) | 507 (4.55) | 579 (4.60) | 631 (4.87) | 735 (4.43) |
| TD–3 | Tb | 343 (5.18) | 407 (4.86) | 508 (4.45) | 581 (4.59) | 631 (4.85) | 742 (4.39) |
| TD–4 | Dy | 344 (5.21) | 418 (4.93) | 511 (4.58) | 584 (4.63) | 634 (4.86) | 749 (4.41) |
| TD–5 | Tb | 341 (5.23) | 407 (4.87) | 509 (4.59) | 581 (4.64) | 632 (4.90) | 741 (4.45) |
| PhCN | TD–1 | Gd | 343 (5.02) | 402 (4.79) | 504 (4.18) | 581 (4.27) | 622 (4.66) | 734 (4.22) |
| TD–2 | Gd | 343 (5.08) | 410 (4.95) | 513 (4.47) | 582 (4.49) | 633 (4.75) | 744 (4.27) |
| TD–3 | Tb | 342 (5.11) | 409 (4.81) | 514 (4.48) | 584 (4.47) | 633 (4.74) | 745 (4.45) |
| TD–4 | Dy | 344 (5.09) | 416 (4.77) | 514 (4.47) | 585 (4.45) | 636 (4.71) | 752 (4.22) |
| TD–5 | Tb | 344 (5.15) | 410 (4.85) | 511 (4.56) | 583 (4.59) | 634 (4.80) | 745 (4.41) |

*Broad band.*
the series of (Pc)Ln(Pc)Ln(Por) (TD-1–TD-4), which are reduced at −0.64 to −0.66 V in the first step and −1.08 to −1.10 V in the second. The difference in potentials between the first oxidation of monomeric porphyrins and monomeric phthalocyanines with the same central metal ions is not as large as in the case of reductions, and thus one cannot easily assign the site of electron transfer for the first oxidation of the monomeric macrocycles on the basis of $E_{1/2}$ values alone. The same is true for compounds in the triple-decker series, where the first oxidations of the Ln$_2$(OEP)$_3$ derivatives (TD-9–TD-11) are easier than the first oxidation of the Ln$_2$(Pc)$_3$ species by an average of 115 mV; however, as seen in Table 2, much smaller differences in $E_{1/2}$ are observed when potentials are compared for the second, third, and fourth oxidations of triple-decker compounds in the series of Ln$_2$(Pc)$_3$ (TD-6–TD-8) with those of Ln$_2$(OEP)$_3$ (TD-9–TD-11).

Although no significant differences can be seen between $E_{1/2}$ values for the second oxidation of compounds in the Ln$_2$(OEP)$_3$ and Ln$_2$(Pc)$_3$ series (where $E_{1/2}$ values range from 0.59 to 0.67 V), these potentials are all more negative than $E_{1/2}$ values for the second oxidation of TD-1–TD-4, where the half-wave potentials range from 0.82 to 0.87 V.

Finally, it should be noted that half-wave potentials for the third and fourth oxidations of TD-1–TD-4 are virtually the same as $E_{1/2}$ values for the third and fourth oxidations of TD-6–TD-8, the latter of which must be Pc-based in the Ln$_2$(Pc)$_3$ series by virtue of the fact that no porphyrin ligands are present in the compounds.

In summary, the above analysis of the electrochemical data is consistent with the first reduction at one or both of the Pc ligands in TD-1–TD-4 and also with the first oxidation at one or both of the Pc ligands in the same series of compounds.

![Figure 2. Cyclic voltammograms of TD-1–TD-4 in PhCN with 0.1 M TBAP.](image)

<table>
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<th>Table 2. Half-Wave Potentials and Assignment of Site of Electron Transfer for Reduction and Oxidation of TD-1–TD-4 in PhCN and of Reference Compounds TD-6–TD-11 in CH$_2$Cl$_2$ <em>a</em></th>
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*The measured redox potentials are given in Table 2. *See structures in Scheme 2. *The first oxidation is not Pc-based for Ln$_2$(OEP)$_3$, which contains only porphyrin macrocycles.

![Scheme 2. Structures for Three Groups of Triple-Decker Complexes: (a) from This Work; (b) from Ref 7b; and (c) from Ref 15*](image)
In this regard, it should be noted that the experimentally measured HOMO−LUMO gaps of TD-1−TD-4 range from 1.15 to 1.20 V in PhCN, values which are about 100 mV larger than the HOMO−LUMO gaps reported for an earlier characterized series of (Pc)Ln(Pc)Ln(Por) compounds\(^{16}\) and 200 mV larger than the HOMO−LUMO gaps for compounds in the Ln\(_2\)(Pc)\(_3\) series TD-6−TD-8, where \(\Delta E_{1/2}\) ranges from 0.97 to 1.03 V, as seen in Table 2.

Significantly, however, the HOMO−LUMO gaps for the three Ln\(_3\)(OEP)\(_3\) compounds (TD-9−TD-11) range from 1.81 to 1.92 V, consistent with a difference in the site of electron transfer between the first-decker derivatives with three OEP macrocycles and those with three Pc macrocycles (TD-6−TD-8) or those with two Pc and one Por macrocycle (TD-1−TD-4).

Thin-layer UV−vis spectroteleocohemistry,\(^{17,18}\) in combination with assignments of electronic absorption bands for the neutral compounds in the series of TD-1−TD-4, was utilized to elucidate and confirm the site of electron transfer proposed above for the first oxidation and first reduction of the triple-decker compounds. Examples of the spectral changes are given in Figure 3 and Figures S7−S9 in the Supporting Information, which show UV−vis spectra obtained during the first one-electron oxidation and first one-electron reduction of TD-2, TD-1, TD-3, and TD-4, respectively.

For TD-2, the band at 408 nm of the neutral compound is attributed to the porphyrin Soret band, in part on the basis of the analysis of electronic absorption spectra of the monomeric porphyrin precursor, 7a or 7b (Figure S6 in the Supporting Information), and in part on previous assignments of the absorption bands in the literature.\(^{19−21}\) This band in the triple-deckers remained almost unchanged in intensity and position after the first one-electron oxidation at an applied potential of 0.70 V (Figure 3a), which suggests that the first oxidation of TD-2 is located in large part on one or both of the phthalocyanine macrocycles in the triple-decker compounds. This assignment is also consistent with the observed changes in the phthalocyanine bands of the neutral compound as the reaction proceeds. The bands of the neutral compound at 344, 512, and 586 nm in PhCN decrease in intensity during the first oxidation, as seen in Figure 3a. The Q band at 633 nm, which is assigned to the Pc macrocycle (see Table 1), also decreases significantly in intensity and is replaced by a new band at 660 nm in the one-electron-oxidized compound.

Although the porphyrin Soret band of the neutral compound at 408 nm increases in intensity and shifts to 419 nm during the first reduction (Figure 3b), the overall spectral changes during the first reduction are attributed to a phthalocyanine-based reaction, in large part on the basis of the fact that the two bands at 344 and 633 nm, which are assigned to the Pc macrocycles, both decrease in intensity. This is consistent with conclusions from the above analysis of the electrochemical data and indicates that one or both phthalocyanine macrocycles are the predominant site of the first electron addition. Similar spectral changes are observed for TD-1, TD-3, and TD-4 during the first oxidation and first reduction in PhCN (see Figures S7−S9 in the Supporting Information).

Magnetism. The direct current (dc) magnetic properties of the isostructural triple-deckers Tb\(_2\) (TD-3) and Dy\(_2\) (TD-4) were measured on microcrystalline powder samples and were studied between 1.8 and 300 K under an applied field of 1000 Oe. Plots of \(\chi_M T\) vs \(T\) are given in Figure 4. The \(\chi_M T\) value of 20.06 cm\(^3\) K/mol for the Tb\(^{III}\) containing complex TD-3 is in good agreement (\(S = 6/2\), \(g = 3/2\), \(C = 11.82\) cm\(^3\) K/mol)\(^{22}\) with the expected value of 23.64 cm\(^3\) K/mol. The \(\chi_M T\) value of 29.35 cm\(^3\) K/mol for the Dy\(_2\) complex (TD-4) at room temperature is in good agreement with what is expected for two independent Dy\(^{III}\) (\(S = 5/2\), \(g = 4/3\), \(C = 14.17\) cm\(^3\) K/mol)\(^{22}\) ions (28.34 cm\(^3\) K/mol). It is interesting to compare the magnetic behavior of these two isostructural compounds which contain non-Kramers (TD-3) versus Kramers ions (TD-4). The behavior of the Dy\(^{III}\)-containing compound TD-4 is discussed first in this comparison.

When the temperature is lowered, the \(\chi_M T\) product of TD-4 decreases steadily until it reaches a minimum value of 15.30 cm\(^3\) K/mol at 6.65 K. On further cooling, the \(\chi_M T\) value sharply increases to reach a value of 17.47 cm\(^3\) K/mol at 1.8 K. For the complex TD-3, the decrease of the \(\chi_M T\) product when the temperature is lowered is much smaller and the minimum appears much earlier, at 28.0 K, with a value of 14.16 cm\(^3\) K/mol.
Upon further cooling, the $\chi_M T$ product for TD-3 sharply increases, as also seen for TD-4, and reaches a value of 20.06 cm$^3$ K/mol at 1.8 K. This behavior is indicative of ferromagnetic interactions between the Ln$^+$ ions, as previously shown by Ishikawa and co-workers through a comparative study of PcLnPcYPc-type and homodinuclear PcLnPcLnPc-type triple-decker complexes, mediated not by exchange but rather by magnetic dipolar coupling.3b

The field dependence of the magnetization (Figure 5) was measured between 2.0 and 5.0 K. For compound Dy$_2$ (TD-4) the lack of saturation in the magnetization values indicates the presence of significant magnetic anisotropy and/or low-lying excited states. The magnetization measurements for compound TD-4 show no saturation up to 7 T, and the values of the isotherms rapidly increase at low field before following a more gradual linear increase after 1.0 T without saturation. The anisotropy present in this compound is seen from the reduced magnetization plots shown as $M$ vs $H/T$ (Figure 5 right, inset) which clearly do not superpose onto a single master curve. On the other hand, for compound TD-3, the isostructural Tb$^+$ complex, the magnetization shows a rapid increase in the magnetization at low magnetic fields, reaching a clear saturation above 1.0 T and the reduced magnetization shown as $M$ vs $H/T$ (Figure 5 left, inset), essentially shows a superposition of the three isotherms onto one master curve.

The behavior of both compounds is very similar to what was observed previously for (Pc)Ln(Pc)Ln(Pc)-type homo- and heterodinuclear Tb triple-decker systems.3e However, the magnetization behavior for the Tb analogue was interpreted in terms of non-negligible magnetic anisotropy and/or low-lying excited states, but we prefer to regard the current Tb system as essentially isotropic in terms of its magnetization behavior. The values of the magnetization at 7 T and 2 K are 5.71 $\mu_B$ for Tb$_2$ (TD-3) and 8.41 $\mu_B$ for Dy$_2$ (TD-4).

In order to investigate the presence of slow relaxation of the magnetization which may originate from SMM behavior, ac magnetic susceptibility measurements were performed on compounds TD-3 and TD-4. In attempts to suppress any quantum tunneling of the magnetization (QTM), the frequency-dependent ac susceptibility was measured for both complexes with a varying applied field (Figure S10 in the Supporting Information). Although other Tb$^+$ triple-decker compounds can show ac signals without the need for applying a field,3d in this case, for TD-3, the optimum dc field where the quantum tunneling is smallest was found to be 1000 Oe. In the case of the Dy$_2$ triple-decker (TD-4) an optimum field could
not be determined, as evidenced by the very weak out-of-phase signal, which suggests that suppression of QTM is not possible using the available parameters of the SQUID magnetometer.

The ac measurements, carried out under a dc field of 1000 Oe for Tb$_2$ (TD-3) (Figure 6), reveal out-of-phase maxima in the range of 1.8–6.0 K. These maxima, highlighted with a purple arrow, were used to extract the relaxation time as a function of the temperature. Additionally, temperature-dependent ac measurements were performed (see Figure S11 in the Supporting Information), to investigate the dynamic magnetic processes. Temperature-dependent signals are visible for frequencies higher than 100 Hz in the temperature range from 9.0 to 13.0 K. The maxima could be extracted and were used to calculate the energy barrier (Figure 7). Both extracted maxima were plotted as ln $\tau$ versus $1/T$ in order to gain insight into the dynamics of the relaxation processes. There are two distinctly visible regimes, which can be fitted by using eq 1 which takes only a direct and an Orbach process into account.

\[
\tau^{-1}(T, H) = A H^{4/3} T + \tau_0^{-1} \exp\left(\frac{U_{\text{eff}}}{k_B T}\right) + \text{QTM}
\]

In eq 1, the $AH^{4/3}T$ term defines the direct relaxation process for Kramers ions and the second exponential term defines the Orbach process, where the magnetization of the system decays through the thermally activated population of the Kramers doublet. The last term originates from QTM between the two sides of the magnetic anisotropy barrier and is normally only used for zero-static-field data. In order to reduce over-parametrization, this contribution has not been estimated. The presence of an Orbach process is more important in characterizing an SMM, because it allows a spin to reorient by climbing over the anisotropy barrier, $U_{\text{eff}}$.

The high-temperature regime is dominated by an exponential process, that is, a two-phonon Orbach process via a real intermediate state, or thermally assisted quantum tunnelling of the magnetization (TA-QTM), resulting from spin–lattice interactions. At low temperatures a direct process dominates. The fit of the data in Figure 7 gives $A = 1.65 \times 10^{-3}$ m$^{-1}$ K$^{-1/2}$ T$^{-4}$, $U_{\text{eff}}/k_B = 132.0$ K, and $\tau = 5.0 \times 10^{-3}$ s.

The Cole–Cole plots of $\chi''$ versus $\chi'$ for Tb$_2$ (TD-3) between 1.8 and 6.0 K (Figure 8) reveal that there are actually two relaxation processes in the measured temperature range, with profiles typical for the presence of these. The first process, which we have already identified from the Arrhenius analysis of the data, is related to the set of complete semicircles. Additionally, at small $X_{\text{M}}$ and small $f_{\text{M}}$ values the beginning of a second relaxation process appears, which is also in line with the data obtained by Holmberg et al. on their Tb compound, although they chose to concentrate only on the first process (see the Supporting Information of ref 3e). Here we were able to fit the $X_{\text{M}}$ and $f_{\text{M}}$ data simultaneously for the two processes in order to constrain the variable parameters using a generalized Debye model.

The extracted $\tau$ values were used to calculate the energy barrier for both processes. Using an Arrhenius fit for the first process leads to an energy barrier of $U_{\text{eff},1} = 5.7$ K and $\tau = 5.1 \times 10^{-3}$ s ($R = 0.999$) (Figure S12), at a temperature range between 3.0 and 6.0 K. These results are consistent with the fact that at low temperatures, in the case of the non-Kramers ion TbIII, only the $T_{\text{g}}$ ground state is populated, which is well separated from the first excited state, leading to QTM as the major relaxation process when no field is applied. Upon application of a dc field of 1000 Oe, barrier-dependent processes, which are faster than the QTM and direct process, take place. The first process at low frequencies in the range 1–200 Hz where the linear curve fit of the ln $\tau$ versus $1/T$ data suggests that a direct process is operative in this range. The second relaxation process takes place at higher frequencies, and the energy barrier and relaxation time were extracted from the Cole–Cole fit, in the range 3.0–6.0 K, where the data conform to a $1/\tau \propto T^x$ behavior typical for a non-Kramers system (Figure S13 in the Supporting Information), which is in line with a dominant Raman process.

**STM Imaging.** The triple-decker complexes TD-3 with four undecyl groups (C$_{11}$H$_{23}$) and TD-5 with four hexadecyl groups (C$_{16}$H$_{23}$) on the macrocyclic porphyrin were chosen to study their self-assembly behavior on a highly oriented pyrolytic graphite (HOPG) surface for imaging by scanning tunneling
microscopy (STM) at the liquid–solid interface. For comparison, self-assemblies of the free porphyrins 7b,c were also imaged under similar conditions.

In both cases, no stable supramolecular networks could be observed under the imaging conditions, as shown in Figure S14 in the Supporting Information for the two triple-decker complexes. However, with regard to the two free base porphyrins, stable supramolecular networks could be imaged, identical with those previously reported.9

Sandwich double- and triple-deckers can be observed in a straightforward way on a metal surface by STM at low temperature,30 but there have been few reports for room-temperature imaging at the HOPG/liquid interface,31 although under these latter conditions, no particular care was required for sample preparation and imaging conditions. Supramolecular self-assembly on surfaces is usually the result of a delicate balance between intermolecular and molecule–substrate interactions. Janus tectons2 can in principle adsorb in various lying-up or upside-down configurations, which can result in very different adsorption energies.33 The currently examined triple-deckers, TD-3 and TD-5, were specially designed to adsorb on the porphyrin side due to the presence of the long alkyl chains that are usually most favorable for adsorption on HOPG.19 Nevertheless, we cannot exclude here the possibility of other adsorption configurations, with the upper phthalocyanine lying down or standing up on the side, thus prohibiting the formation of stable ordered networks.

**CONCLUSION**

In summary, five novel rare-earth heteroleptic porphyrin phthalocyanine triple-decker complexes with bulky peripheral groups were successfully prepared by a microwave-mediated synthesis in medium yields and characterized by electrochemical and spectroelectrochemical techniques. On the basis of the measured redox potentials, the electronic absorption spectra of the initial compounds, and the measured spectral changes during the first oxidation/reduction reactions, the initial sites of oxidation and reduction were both assigned as being phthalocyanine-based. The magnetic properties of the Tb1 (TD-3) and Dy2 (TD-4) compounds reveal ferromagnetic interactions shown by the susceptibility measurements which are caused by magnetic dipolar interactions. The Tb analogue, TD-3, shows SMM behavior under an applied field of 1000 Oe, where the direct and Orbach processes can be determined, resulting in an energy barrier of $U_{\text{eff}} = 132.0$ K. Cole–Cole plots reveal typical profiles for the presence of two relaxation processes, where the second relaxation process takes place at higher frequencies and the energy barrier and relaxation time were extracted from the Cole–Cole fit. The data conforms to $1/\tau \propto T^\gamma$, which suggests that the second relaxation process is dominated by a Raman process. Furthermore, these compounds could not form 2D self-assembled networks on an HOPG surface due to their bulky peripheral groups. Meanwhile, we were unable to construct in silico stable triple-deckers which could be energy minimized. All of our attempts, either at semiempirical levels (which lack an adequate parametrization for lanthanides) or at the DFT level, resulted in nonstable minima. In the absence of accurate geometries we were thus unable to perform reliable calculations to gain insight into the mixing of the Pn and Pc transitions within the triple-deckers and depending on the Ln ion.

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