

# A Core-Modified Rubyrin with *meso*-Aryl Substituents and Phenanthrene-Fused Pyrrole Rings: A Highly Conjugated Near-Infrared Dye and Hg<sup>2+</sup> Probe\*\*

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Expanded porphyrins<sup>[1]</sup> with five or more heterocyclic rings, such as sapphyrins, rubyrins, and more extended structures, are large conjugated macrocycles that basically preserve key characteristics of porphyrins (e.g. Soret and Q-type absorption bands, photoactivity) yet often show various novel features that have led to applications in medical and pharmaceutical areas, nonlinear optics, and supramolecular photochemistry.<sup>[2]</sup> Expanded-porphyrin structures have been elaborated by macrocycle expansion, the introduction of heterocycles other than pyrrole (core modification), *meso* substitution, and heterocycle inversion ("confused" porphyrins).<sup>[1]</sup> Because such porphyrins are macrocycles, their interesting cation and anion coordination properties render them promising hosts in molecular recognition and chemical sensing.<sup>[3,4]</sup>

Of the large number of expanded porphyrins published in the past 20 years, only few have been employed in recognition and sensing studies, mainly all-pyrrole derivatives for anion

signaling.<sup>[5]</sup> Core-modified expanded porphyrins have been employed even less frequently, for example, thiophene-modified sapphyrins and rubyrins in anion-binding studies.<sup>[6]</sup> Although the cation-coordination chemistry of expanded porphyrins has been investigated,<sup>[7,8]</sup> reports on the use of such compounds in metal-ion sensing are even scarcer.<sup>[9]</sup>

Based on our interest in the optical determination of heavy-metal ions, in particular of Hg<sup>2+</sup>,<sup>[10]</sup> and the development of new red/near-infrared (NIR) dyes,<sup>[11]</sup> we became intrigued by the possibility of equipping expanded porphyrins such as rubyrin, that is, [26]hexaphyrin(1.1.0.1.1.0), with various "soft" donor sites,<sup>[12–14]</sup> to shift metal-ion preferences from, for example, Zn<sup>2+</sup>, Cu<sup>2+</sup>, and Co<sup>2+</sup> in all-pyrrole derivatives<sup>[7]</sup> to more thiophilic target ions. Moreover, since the strategy of core annelation has been successfully employed to shift the optical spectra of porphyrins into the NIR,<sup>[15,16]</sup> we synthesized rubyrin **1** and investigated its spectroscopic properties and Hg<sup>2+</sup>-sensing features.

The synthesis of **1** started from the oxidative coupling of monolithiated thiophene<sup>[17]</sup> and commenced with the condensation of 5,5'-dilithiated **2** with 4-fluorobenzaldehyde to yield the novel bithiophene diol **3** (Scheme 1).<sup>[18]</sup> Condensation of **3** and phenanthrene-annelated pyrrole under modified Lindsey conditions and subsequent oxidation with DDQ yielded rubyrin **1** in 10% yield. In agreement with observations on **7**,<sup>[13]</sup> <sup>1</sup>H NMR measurements at –40°C in CDCl<sub>3</sub> showed two multiplets at  $\delta$  = 9.47 and 9.16 ppm, corresponding to two types of magnetically nonequivalent bithiophene protons that presumably reflect a different degree of distortion from planarity for the two bithiophene units. Since the *ortho* and *meta* protons of the phenyl rings are directed toward the rubyrin macrocycle and are thus deshielded by the rubyrin and phenanthrene ring currents, multiplets are found at  $\delta$  = 8.55 (8H) and 7.95 ppm (8H). The remaining two multiplets at  $\delta$  = 7.39 (8H) and 7.09 ppm (8H) are assigned to the protons of the phenanthrene rings.

Figure 1 shows the X-ray crystal structure of **1**.<sup>[19]</sup> Average bond lengths of 1.354 and 1.395 Å for the N–C<sub>α</sub> and C<sub>meso</sub>–C<sub>α</sub> bonds indicate  $\pi$ -electron delocalization in the macrocyclic core. Double-bond character is also found for the average C<sub>α</sub>–C<sub>β</sub> (1.397 Å) and C<sub>β</sub>–C<sub>β</sub> (1.389 Å) bonds of the thiophene rings. In contrast, C<sub>α</sub>–C<sub>β</sub> and C<sub>β</sub>–C<sub>β</sub> bond lengths of 1.504 and 1.374 Å indicate localized single and double bonds in the pyrrole rings. As a result of steric crowding, the phenanthrene units are largely bent away from the least-squares plane of the macrocyclic core (Figure 1) and the fluorophenyl rings are only tilted by  $\theta$  = 8.5–34.4° out of that plane; the average

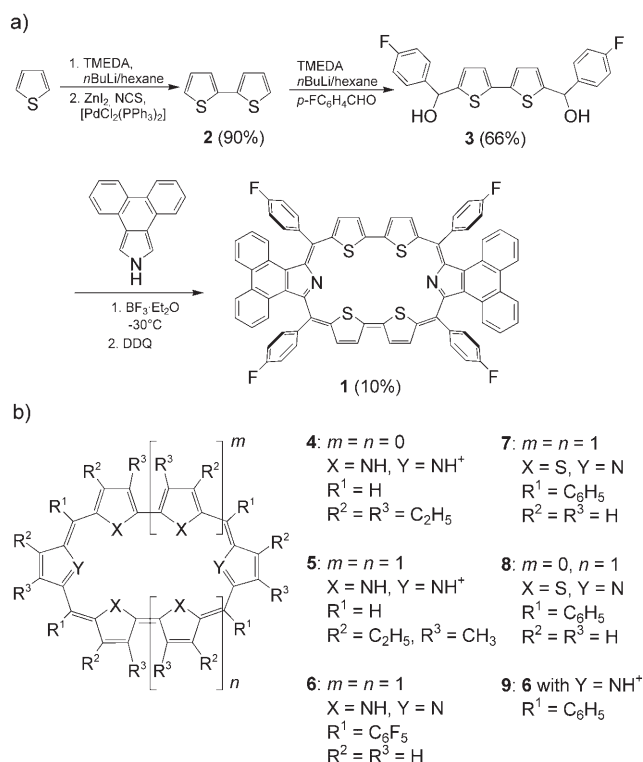
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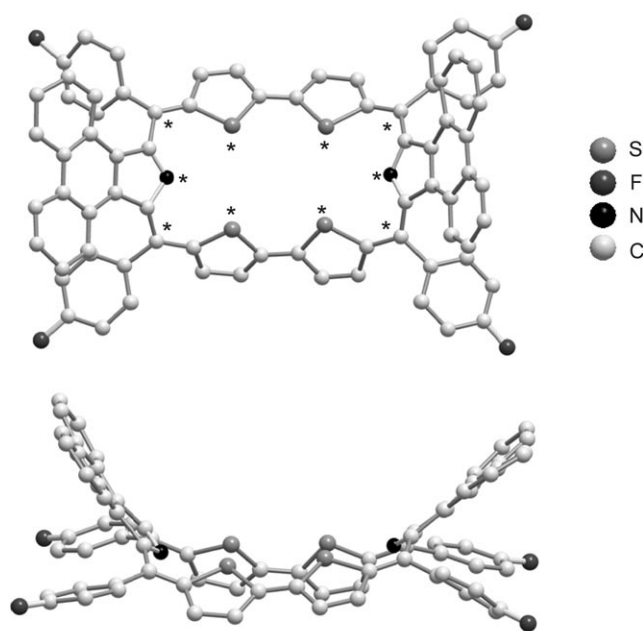
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**Scheme 1.** a) Synthesis of phenanthrene-anellated rubyrin **1**. TMEDA = *N,N,N',N'*-tetramethylethylenediamine, NCS = *N*-chlorosuccinimide, DDQ = 2,3-dichloro-5,6-dicyano-1,4-benzoquinone. b) Known porphyrins discussed in the text.

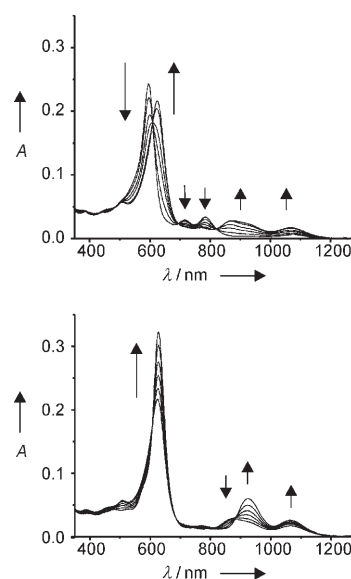


**Figure 1.** Top and side view of the X-ray crystal structure of **1**. Hydrogen atoms are omitted for clarity. Atoms marked with asterisks define the least-squares plane of the central macrocycle.

bond length between the  $C_{\text{meso}}$  atoms and the fluorophenyl rings is 1.4815 Å. The dominant structural features are thus

the bowl-type distortion of the molecule and the existence of a 26- $\pi$ -electron aromatic ring. Moreover, the fluorophenyl rings are much more in plane than those in related dyes such as a benzohexaphyrin ( $\theta = 49.3\text{--}76.3^\circ$ ),<sup>[16c]</sup> the non-anellated sapphyrin **8** ( $\theta = 57.9\text{--}68.6^\circ$ ),<sup>[6]</sup> and the bowl-shaped hexaphyrin **9** ( $\theta = 36.2\text{--}46.7^\circ$ ).<sup>[20]</sup> If one considers porphyrin photochemistry this suggests that the exchange of the 4-substituent of the *meso*-phenyl rings could potentially be used to tune the spectroscopic properties of dyes such as **1**.<sup>[21]</sup>

The absorption spectrum of **1** in its free-base form in  $\text{CHCl}_3$  displays a Soret band at 596 nm ( $\lg \epsilon = 5.17$ ) and three weak and broad Q bands at 714 ( $\lg \epsilon = 4.24$ ), 784 (4.31), and 1076 nm (3.65), the latter well in the NIR region (Figures 2 and 3). These favorable features can be rationalized as

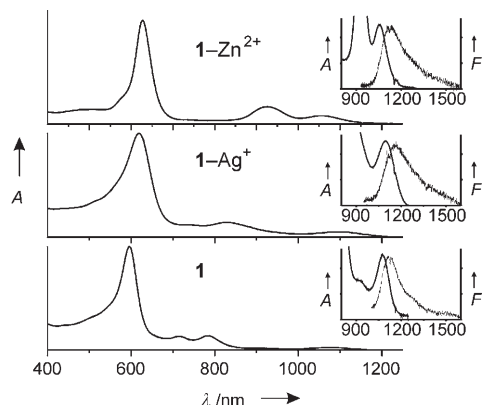


**Figure 2.** Titration spectra of **1** with TFA in  $\text{CHCl}_3$ . Top: no TFA and  $2 \mu\text{M} \leq c_{\text{TFA}} \leq 0.32 \text{ mM}$ ; bottom:  $0.32 \text{ mM} \leq c_{\text{TFA}} \leq 19 \text{ mM}$ .

follows: Extension of the  $\pi$  system of the parent porphyrin **4** (Scheme 1) by two pyrrole units yields rubyrin **5** and shifts the Soret and Q bands by roughly 100 and 270 nm, to 505 and 850 nm.<sup>[22]</sup> Further red-shifts can then result from *meso* substitution.<sup>[23]</sup> Rubyrin **6** displays bands at  $\lambda_{\text{Soret}} = 543$  and  $\lambda_{\text{Q}} = 926 \text{ nm}$ .<sup>[24]</sup> Modification of the rubyrin core by replacing the N atoms with other heteroatoms then alters the cavity size and electronic structure of the ring, leading to moderate hypsochromic shifts in the case of N-for-S exchange; for the tetrathiophenerubyrin **7**,  $\lambda_{\text{Soret}} = 523 \text{ nm}$ .<sup>[13]</sup> Since our aim was to synthesize a dye that absorbs at distinctly longer wavelengths, the strategy of phenanthrene anellation at the  $\beta$ -pyrrole position that we recently employed for tetrapyrrole porphyrins<sup>[25]</sup> also proved very successful here:  $\lambda_{\text{Soret}}^1$  and  $\lambda_{\text{Q}}^1$  are red-shifted by approximately 70 and 120 nm relative to the bands of **7**.

Despite the wealth of information available on the absorption properties of extended porphyrins, only few studies of their emission features have been published.<sup>[8,9,24,26,27]</sup> This can be explained by the difficulty of measuring emission spectra of compounds with low fluores-

cence quantum yields— $\Phi_f \approx 10^{-4}$  have been reported<sup>[26,27]</sup>—at  $\lambda_{em} > 900$  nm with conventional fluorimeters. For the emission spectra shown in Figure 3, we used a different approach and employed an FTIR spectrometer equipped with a



**Figure 3.** Absorption spectra of **1**, **1-Ag<sup>+</sup>**, and **1-Zn<sup>2+</sup>** in  $\text{CHCl}_3$ . Insets: normalized  $S_0$ – $S_1$  transitions in absorption and emission ( $\lambda_{exc} = 532$  nm).

Nd:YAG laser and a photoluminescence detector operating in the conventional mode (see the Supporting Information for details). The fluorescence of **1** was observed between 1050 and 1350 nm with  $\lambda_{max} = 1118$  nm,  $\Phi_f = 3.3 \times 10^{-4}$ , and a fluorescence lifetime  $\tau_f = 90$  ps.<sup>[28]</sup> The Stokes shift of  $355\text{ cm}^{-1}$  indicates that the conformational relaxation in the excited state is not very pronounced. However, the Stokes shift also suggests that the conformation of **1** deviates considerably from planarity because Stokes shifts of only  $20$ – $90\text{ cm}^{-1}$  have been reported for largely planar hexaphyrins.<sup>[8,26,27]</sup> This assumption is supported by the comparatively intense Q bands. When Q bands lose their ideal forbidden character, as in *meso*-substituted and anellated tetrapyrrole porphyrins that adopt a saddlelike conformation owing to internal steric strain, their intensity increases.<sup>[29]</sup>

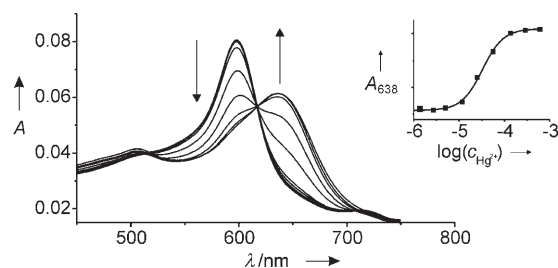
The increased  $\pi$  conjugation of **1** by fusion with phenanthrene rings is also evident from cyclic voltammetric studies. Two quasi-reversible reduction peaks were observed at  $-1.39$  and  $-1.64$  V (vs.  $\text{Fc}^+/\text{Fc}$ ), while four quasi-reversible and one irreversible oxidation peaks appeared at  $-0.34$ ,  $-0.26$ ,  $0.08$ ,  $0.23$ , and  $0.50$  V. The HOMO–LUMO gap for **1** is estimated to be  $1.05$  V, which is significantly smaller than those of, for example, **7-H<sub>2</sub><sup>2+</sup>** ( $1.64$  V) and meso-tetraphenylporphyrin ( $2.26$  V), and reflects the pronounced red-shift of the absorption spectrum of **1**.<sup>[13]</sup>

In accordance with changes measured for other rubyrins upon protonation,<sup>[14,30]</sup> addition of an excess of trifluoroacetic acid (TFA) to **1** in  $\text{CHCl}_3$  led to an additional bathochromic shift of the Soret band to  $628$  nm (Figure 2). Moreover, just as was observed for **7** and **7-H<sub>2</sub><sup>2+</sup>**,<sup>[14]</sup> the two high-energy Q bands at  $716$  and  $785$  nm disappear upon formation of **1-H<sub>2</sub><sup>2+</sup>**, and the low-energy Q bands at  $913$  and  $1068$  nm strongly gain in intensity. Such behavior reflects a further distortion of the conformation from planarity. Titration of **1** with TFA revealed two well-separated protonation steps with apparent

$pK_a$  values of  $4.62 \pm 0.03$  and  $2.68 \pm 0.02$  (see the Supporting Information).

When divalent metal ions such as  $\text{Cu}^{2+}$ ,  $\text{Zn}^{2+}$ ,  $\text{Pb}^{2+}$ , or  $\text{Hg}^{2+}$  were added to an equimolar amount of **1** in  $\text{CHCl}_3$ , the observed effects were similar to those upon protonation (Figure 3). The Soret band is shifted to  $627$  nm and two Q bands are observed at  $925$  and  $1056$  nm ( $1058$  nm for  $\text{Hg}^{2+}$ ). In the case of  $\text{Ag}^+$ , the absorption maxima have a less pronounced shift, appearing at  $616$  (Soret band) and  $828/1085$  nm (Q bands), most probably because of the weaker interaction of the monovalent thiophilic ion with the pyrrole nitrogens. Fluorescence studies of **1-Ag<sup>+</sup>**, **1-Zn<sup>2+</sup>**, and **1-Hg<sup>2+</sup>** revealed a 2.2- and 3.3-fold increase in emission upon complexation of the lighter diamagnetic metal ions  $\text{Ag}^+$  and  $\text{Zn}^{2+}$ , and a 1.5-fold quenching for  $\text{Hg}^{2+}$  with band maxima at  $1162$ ,  $1124$ , and  $1190$  nm. The comparatively weak influence of the heavy-atom effect of  $\text{Hg}^{2+}$  is tentatively ascribed to the short fluorescence lifetime  $\tau_f$  of **1**, because intersystem crossing is much less probable than internal conversion for such red-emitting porphyrinoids.<sup>[27]</sup>

As mentioned above, **1** was designed as a  $\text{Hg}^{2+}$  probe. Although the public is alert to the dangers of mercury contamination,<sup>[31]</sup> there is still a need for new or improved sensing methods that are applicable on site and for screening purposes.<sup>[32]</sup> The development of  $\text{Hg}^{2+}$  indicators that can be principally employed in optical-sensing devices is thus an active research field.<sup>[10,33]</sup> Since **1** is not soluble in water, a common strategy was employed to work under realistic conditions: The probe was incorporated into a polymer membrane.<sup>[34]</sup> For most polymer-based  $\text{Hg}^{2+}$  sensors, PVC membranes are used.<sup>[34]</sup> However, the polyurethane hydrogel D4, which was recently identified as an excellent support for pH sensors,<sup>[35]</sup> gives a better performance with **1**. Figure 4

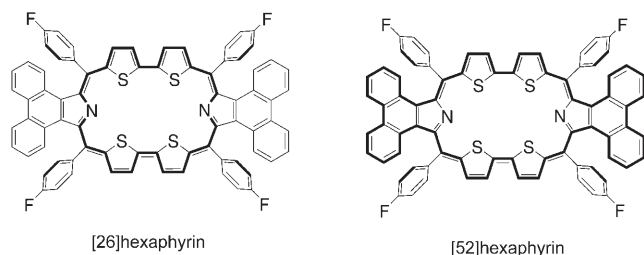


**Figure 4.** Changes in absorption (Soret-band region) of a glass slide coated with a  $35\text{-}\mu\text{m}$  thick membrane of D4–**1** after it was dipped into water samples containing increasing amounts of  $\text{Hg}^{2+}$ . Inset: Titration curve of  $A_{638}$  vs.  $\text{Hg}^{2+}$  concentration.

shows the titration spectra obtained after dipping D4–**1** into aqueous samples with increasing amounts of  $\text{Hg}^{2+}$ . In the absence of  $\text{Hg}^{2+}$ , the Soret band is found at  $597$  nm, virtually the same as for **1** in  $\text{CHCl}_3$ , indicating that the matrix does not influence the spectroscopic behavior properties of **1**. The response to  $\text{Hg}^{2+}$  is evident from a shift of the Soret band to  $638$  nm with a clear isosbestic point at  $617$  nm. By monitoring the intensity changes at  $638$  nm, a typical titration curve is obtained with a limit of detection of approximately  $1$  ppm. Quantitative analysis of several titration curves yielded

acceptable fits for a 1:1 complex and a formation constant  $K = (3.0 \pm 0.3) \times 10^4 \text{ M}^{-1}$ . Complexes with a ligand-to-metal ratio of 1:2<sup>[7a,8]</sup> have been found in neither H<sub>2</sub>O nor CHCl<sub>3</sub>, presumably because of the lack of preferred coordination sites for aminophilic cations like Cu<sup>2+</sup> and Zn<sup>2+</sup> and the rather large ionic radius in the case of Hg<sup>2+</sup>. Competition studies with Zn<sup>2+</sup>, Cd<sup>2+</sup>, Cu<sup>2+</sup>, Co<sup>2+</sup>, Ni<sup>2+</sup>, Pb<sup>2+</sup>, Ag<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>, K<sup>+</sup>, and Na<sup>+</sup> underscore the selective response of D4-1. Only at concentrations of > 100  $\mu\text{M}$  does the band at 638 nm increase 1.03- (for Ni<sup>2+</sup>), 1.08- (for Cd<sup>2+</sup> and Pb<sup>2+</sup>), and 1.15-fold (for Cu<sup>2+</sup>); the other ions have no effect. Moreover, as would be expected for free-base expanded porphyrins, different counterions do not influence the performance.<sup>[36]</sup>

In conclusion, we have synthesized a core-modified, expanded porphyrin with polycyclic aromatic units fused to the pyrrole rings and demonstrated its protonation and complexation features. Embedding **1** into a polyurethane membrane produced simple test strips for rapid Hg<sup>2+</sup> screening. This is one of the very few examples—and the first one for cations—where expanded porphyrin probes can be used directly in aqueous samples without complications arising from aggregation.<sup>[37]</sup> Most examples so far use such indicator dyes in organic or mixed organic/aqueous solutions or in combination with solvent extraction and transport protocols.<sup>[5,6,9]</sup> Compound **1** shows exceptionally red-shifted absorption bands with the Soret band appearing at almost the same position (596 nm) as that of a 54- $\pi$ -electron-containing dodecaphyrin (604 nm).<sup>[38]</sup> Although **1** formally represents a [26]hexaphyrin, the spectroscopic and redox properties suggest that ring fusion leads to a dye with cross-conjugated [26]- and [52]hexaphyrin chromophores.<sup>[39]</sup> This



tremendous bathochromic shift results in an absorption spectrum that is compatible with solar radiation in the respective wavelength range<sup>[40]</sup> and shows appreciable molar absorptivities of  $\lg \epsilon > 4.17$  at all common laser wavelengths between 322–780 nm, including HeNe, Ar<sup>+</sup>, many diode lasers, and Nd:YAG laser harmonics. For example, in the case of D4-1 and Hg<sup>2+</sup>, the ratiometric response can be analyzed with a yellow (594 nm) and a red (633 nm) HeNe laser. Compounds such as **1** are thus promising candidates for photonic applications.<sup>[2–5]</sup> With the aid of FTIR–photoluminescence instrumentation, such dyes might make it possible to use the second optical window of telecommunication (O-band, 1260–1310 nm)<sup>[41]</sup> for fiber optical fluorometry and sensing.

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