Two-Step Synthesis of Stable Dioxadicarbaporphyrins from Bis(3-indenyl)methane**

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The origin of the aromatic characteristics of porphyrins (**1a**) and related systems remains the topic of considerable debate.^[1-3] Although porphyrins and related macrocycles are often considered to be examples of bridged annulenes,^[4-6] some theoretical studies have concluded that the aromatic properties of porphyrinoids are primarily due to the individual pyrrolic rings.^[3,7] However, the extent to which the pyrrolic nitrogens contribute to the aromatic properties of porphyrins has been brought into question.^[1,2] In particular, dideazaporphyrins **2**, which lack these pyrrole-type nitrogens, have been shown to be strongly diatropic and to possess porphyrin-like electronic absorption spectra.^[1,8] In previous



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[**] Part 62 in the series "Conjugated Macrocycles Related to the Porphyrins". This work was supported by the National Science Foundation (NSF) (CHE-0911699 and CHE-1212691) and the Petroleum Research Fund, administered by the American Chemical Society. The authors also thank the Youngstown State University Structure & Chemical Instrumentation Facility's Matthias Zeller for X-ray data collection. The diffractometer was funded by the NSF (0087210), the Ohio Board of Regents (CAP-491), and YSU.

Supporting information for this article is available on the WWW under http://dx.doi.org/10.1002/anie.201206385.

studies, we have probed the aromatic properties of porphyrinoids by replacing one or more of the pyrrolic subunits with carbocyclic rings.^[5,6] Monocarbaporphyrins related to 1b have been investigated in detail,^[9] but far less is known about dicarbaporphyrinoid systems.^[10-16] In principle, these investigations could be extended to quatyrin (1c), the theoretically important hydrocarbon analogue of the porphyrins,^[5] although severe crowding within the macrocyclic cavity may make this system inaccessible.^[17] Syntheses of opp-dicarbaporphyrinoids $3^{[10]}$ and $4^{[11]}$ with two carbocyclic rings interspersed by pyrrolic moieties have been reported, but these porphyrin analogues proved to be rather unstable.^[10–12] Nevertheless, dicarbaporphyrin 3 exhibited strongly diatropic characteristics and the UV/Vis spectrum for this species closely resembled the spectra for true porphyrins.^[10,13,14] More recently, examples of *adj*-dicarbaporphyrinoid systems have been reported^[15,16] including carbaazuliporphyrin **5**.^[15] These porphyrinoids proved to be far more stable than 3 and 4 and exhibited significant macrocyclic aromaticity. Unfortunately, to date no examples of *adj*-dicarbaporphyrins 6 with two adjacent cyclopentadiene rings have been synthesized. In our investigations into mono- and dicarbaporphyrins, indene units have usually been used in place of cyclopentadiene moieties because of synthetic considerations.^[5,6] For this reason, we set out to target dibenzo-macrocycles related to porphyrinoid 6.

It is well known that indene reacts with aldehydes under basic conditions to afford fulvenes (Scheme 1).^[15] We anticipated that bis(3-indenyl)methane $7^{[18]}$ could similarly be used



Scheme 1. Synthesis of fulvenes from indene.

to prepare fulvene derivatives and that this chemistry might be utilized in the synthesis of *adj*-dicarbaporphyrinoids. Owing to the low reactivity of pyrrole aldehydes, this hypothesis was investigated using furan aldehydes. Bis-(3-indenyl)methane was reacted with difurylmethane dialdehyde **8** under basic conditions in an attempt to generate the heterodicarbaporphyrinoid **9** (Scheme 2), but no indication of macrocyclic products could be discerned. This result may be due to an unfavorable geometry of the intermediates for macrocyclic ring closure. Therefore, the formation of fulvene products was examined by reacting **7** with 2 equivalents of a series of aromatic aldehydes and KOH in refluxing methanol. When furfural was reacted **7** under these conditions, a fulvene product was generated as a red precipitate.

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Scheme 2. Proposed syntheses of fulvenes and a porphyrin analogue from bis(3-indenyl)methane.

In small scale reactions using 50 mg of 7, this product was formed in up to 98% yield, although the yields were lower (73-78%) in larger scale reactions using >1 g of bis-(3-indenyl)methane. Similar results were obtained using thiophene-2-carbaldehyde, benzaldehyde, or 4-substituted benzaldehydes. Initially, it had been anticipated that difulvenes **10** would be formed in these reactions, but the ¹H NMR spectra for these products showed that they lacked the expected plane of symmetry. A possible explanation for these results was that the E,Z isomer **11** had been produced rather than the E,E isomer 10. The ¹H NMR spectrum for the furfural derivative showed the presence of a broad 2H triplet (J = 2 Hz) at 4.11 ppm that was coupled to two 1H resonances at 6.96 and 7.24 ppm. This indicated that a CH=C-CH₂-C=CH unit was present within the structure. While this is consistent with structure 11, detailed analyses using ¹H-¹H COSY, nOe difference ¹H NMR, ¹³C NMR, and HSQC indicated that the fully conjugated species 12a was in actual fact the product of this chemistry (Scheme 3). The thiophene product could similarly be attributed to 12b, whereas the benzaldehydederived series were consistent with structures 13a-d. These unusual conjugated species are structurally related to fully conjugated open-chain tetrapyrroles (bilins) and can therefore be considered to be bilin analogues. The structure of the dithiophene bilin analogue 12b was confirmed by X-ray crystallography, which clearly demonstrated the asymmetry of the two indenyl moieties (Figure 1). There are two crystallographically independent molecules in the asymmetric unit of the structure. The indene containing the apical methylene was identified in both independent molecules. Identification on the basis of difference Fourier location of the hydrogen atoms matched identification of the methylene carbon atoms on the basis of carbon-carbon bond distances.

Bilins 12 and 13 gave orange solutions in dichloromethane, and the UV/Vis spectra showed several broad absorptions between 280 and 450 nm. However, when trifluoroacetic acid (TFA) was added a new species was generated. Addition of TFA to 12a gave a dark green solution that showed strong absorptions at 450, 623, and 819 nm. The ¹H NMR spectrum for 12a in TFA/CDCl₃ showed that the system had gained a plane of symmetry and a 4H singlet was observed at 4.63 ppm. These results indicated



Scheme 3. Synthesis of bilin analogues.



Figure 1. X-ray crystal structure of dithiophene bilin analogue **12b**. Selected bond lengths [Å]: C4–C21 1.388(8), C1–C21 1.457(8), C6–C22 1.466(8), C9–C22 1.487(8). Selected non-bonded separations [Å]: C4---C6 2.487(8), C1---C19 2.501(8), C9---C11 2.492(9), C14---C16 4.57(1), S(23)---S(24) 3.650(3). One of two crystallographically independent molecules is shown for clarity.

that protonation had primarily occurred on an indene unit and that a resonance-delocalized cation, represented by the canonical form $12aH^+$, had been generated (Scheme 3). Similar results were obtained for the dithiophene system 12b. Furthermore, all of the benzaldehyde-derived bilin analogues 13a-d gave rise to the equivalent protonated species $13H^+$. These unusual cations are resonance-delocalized carbenium ions that are formed under very mild conditions. The protonated species derived from 13a and 13b showed a broad absorption in their UV/Vis spectra at 727 and 729 nm, respectively, but this band was bathochromically shifted to 769 nm for the *tert*-butyl substituted version 13f. The dimethoxy bilin analogue gave the largest red shift for this series, showing a long wavelength band at 811 nm and its spectra more closely resembled that of 12a and 12b. The

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cation derived from **12 a** was somewhat unstable and, after an hour or so, the ¹H NMR spectrum of **12 a** in TFA/CDCl₃ showed only broad unresolved absorptions. However, protonated bilins **12b**H⁺ and **13**H⁺ were all reasonably stable and persisted in solution for prolonged periods of time.

The facile protonation and unique properties of bilin analogues **12** and **13** demonstrate that these systems are important in their own right. However, our primary goal was to prepare macrocyclic products from these intermediates. These studies focused on the reactivity of the furfural-derived bilin **12a**, as furans are far more reactive towards electrophilic substitution than thiophenes. When **12a** was reacted with paraformaldehyde in the presence of an acid catalyst, low yields of porphyrinoid product **9** were noted (Scheme 4). The best results were obtained when **12a** was reacted with one



Scheme 4. Synthesis of dioxadicarbaporphyrins.

equivalent of $(CH_2O)_n$ in refluxing 20% TFA/1,2-dichloroethane, followed by oxidation with aqueous ferric chloride. When the oxidation step was not carried out, the conjugated macrocycle could still be isolated, but the product was contaminated with by-products that gave rise to broad absorptions in the ¹H NMR spectra. The oxidation appeared to decompose these impurities as well as facilitating the formation of **9**. Even so, the best yields for this reaction were only 5%, and oxidation with 2,3-dichloro-5,6-dicyano-1,4benzoquinone (DDQ) led to complete decomposition. Much better results were obtained when using aromatic aldehydes.

The reaction of **12a** with one equivalent of benzaldehyde in the presence of 0.5 equivalents of $BF_3 \cdot Et_2O$ in CH_2Cl_2 at room temperature, followed by a brief treatment with DDQ, gave 15-phenyldioxadicarbaporphyrin **14a** in up to 10% yield (Scheme 4). When one equivalent of $BF_3 \cdot Et_2O$ was used, much lower yields were obtained, but lower concentrations of the catalyst also gave poor results. The quantity of solvent used was also a factor, and for reactions with 100 mg of **12a** the best results were obtained when using 10 mL of CH₂Cl₂. Refluxing the reaction in 1,2-dichloroethane gave poor results, but reactions in refluxing CH₂Cl₂ raised the yield to 39% for the phenyl substituted macrocycle. The substrates 4chloro- and 4-bromobenzaldehyde (15b and 15c) also gave the best results under these conditions, affording the corresponding aryl substituted porphyrinoids 14b and 14c in 26% and 40% yields, respectively. At room temperature, 14b and 14c were obtained in only slightly lower yields (23% and 38%, respectively) and these results indicated that electronwithdrawing groups may have a beneficial effect on these reactions. However, 4-nitrobenzaldehyde gave only 12% yield of 14d under optimized conditions. In this case, the best results were obtained under more dilute conditions (100 mL of CH_2Cl_2 for 100 mg of **12a**) and at room temperature. Reasonable yields of 14e and 14f could also be obtained under dilute conditions (15% and 10%, respectively) using dialdehydes 15e and 15f. Although linked diporphyrinoids might be formed from dialdehydes 15e and 15 f, this type of product has not as yet been isolated from these reactions. Nevertheless, the ease of introducing diverse functional groups could potentially allow these dicarbaporphyrinoids to be incorporated into supramolecular systems.

Macrocycles **9** and **14a–f** afforded orange solutions that gave porphyrin-like UV/Vis spectra, each with a Soret band between 436 and 440 nm (Figure 2). For **9**, the Soret band



Figure 2. UV/Vis spectra of 14a in CH_2Cl_2 (-----) and 2% TFA- CH_2Cl_2 (cation 14aH⁺; -----).

appeared at 436 nm and Q-type bands appeared at 510 and 547 nm. Moreover, weak absorptions were observed at 668 and 689 nm. Phenyl substituted porphyrinoid **14a** exhibited minor bathochromic shifts, showing these absorptions at 439, 513, 549, 673, and 690 nm (Figure 2). The addition of TFA to solutions of **9** or **14** in CH₂Cl₂ gave the green-colored protonated species **9**H⁺ and **14**H⁺, respectively, which gave complex UV/Vis spectra showing multiple bands in the Soret region and four absorptions between 600 and 810 nm (Figure 2). The ¹H NMR spectra for dioxadicarbaporphyrins

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9 and **14** showed that the macrocycles are highly diatropic. Impure samples of **9** in CDCl₃ gave resonances for the *meso*protons downfield at 10.06 (1 H), 9.58 (2 H), and 9.31 ppm (1 H), and the internal CH groups afforded a 2H singlet at -3.60 ppm. However, pure **9** was poorly soluble in organic solvents and the ¹H NMR spectrum showed the equivalent resonances at 9.91, 9.40, 9.15, and -4.29 ppm, and these shifts are attributed to aggregation effects. The ¹H NMR spectra for aryl substituted porphyrinoids **14** also showed strong diatropic ring currents, and the *meso*-protons for **14a** were observed at 10.05 (1 H) and 9.58 ppm (2 H), whereas the internal CH groups gave a 2H resonance at -3.61 ppm (Figure 3). The NMR spectra for the cations **9**H⁺ and **14**H⁺ in TFA/CDCl₃ demonstrated that the protonated species



Figure 3. 500 MHz ¹H NMR spectrum of phenyl substituted dioxadicarbaporphyrin **14a**. Proton assignment numbering follows the convention used for compound **9** in Scheme 4.

retained a powerful diamagnetic ring current. For **14**H⁺, the interior CH₂ group gave a singlet at -4.74 ppm, whereas the indene CH group appeared at -2.96 ppm. The *meso*-protons were significantly shifted downfield to give singlets at 10.54 (1H) and 10.62 ppm (2H). ¹³C NMR spectra for **9** and **14** in CDCl₃ confirmed that the macrocycles have a plane of symmetry, but this is lost for the protonated species **9**H⁺ and **14**H⁺.

The X-ray crystal structure of **14a** has also been obtained (Figure 4). The solid state structure has the molecule centered on a crystallographic two-fold axis of rotation which contains H5, C5, C15, C15a, C15d, and H15d. The macrocycle is



Figure 4. X-ray crystal structure of dicarbaporphyrinoid **14**a. Selected bond lengths [Å]: C1–C2 1.476(2), C3–C4 1.478(2), C4–C5 1.398(2), C4–C21 1.404(2), C21–C1 1.392(2), C15–C16 1.398(2), C16–C17 1.406(2), C17–C18 1.373(2), C18–C19 1.398(2), C19–C20 1.388(2), C20–C1 1.396(2), C16–O24 1.383(2), O24–C19 1.390(2). The molecule is centered on a crystallographic two-fold axis of rotation which contains H5, C5, C15, C15a, C15d, and H15d.

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remarkably planar, as evidenced by the 0.0619 Å root-meansquare distance the framework atoms lie from the plane defined by the 24 framework atoms. The largest deviations from the plane are C5 (0.086(7) Å), C15 (-0.094(9) Å), C17 (0.11(1) Å), and C18 (0.090(8) Å), but these deviate by less than 0.06 Å from the aforementioned plane. The structure exhibits framework bond distances consistent with a nearly idealized delocalized π -bonding model with the π -systems for the indenyl benzo-units decoupled from the [18]annulene core, as evidenced by the more single-bond-like C1–C2 (1.476(2) Å), and C3–C4 (1.478(2) Å) bond distances. As would be expected, the phenyl group is canted 77.10(4)° relative to the mean macrocyclic plane.

In conclusion, a unique series of bilin analogues has been prepared in excellent yields by reacting aromatic aldehydes with bis(3-indenyl)methane in the presence of potassium hydroxide in refluxing methanol. The furfural-derived bilin analogue condensed with aromatic aldehydes in the presence of BF₃·Et₂O to give aryl substituted dioxadicarbaporphyrins in good yields (10–40%). An unsubstituted dibenzodioxadicarbaporphyrin was also obtained by reacting the bilin analogue with paraformaldehyde. The new series of porphyrin analogues are stable compounds and are the first examples of porphyrinoids that incorporate two adjacent indene units. The ease of formation of these conjugated macrocycles will no doubt facilitate detailed investigations into this new class of aromatic bridged annulenes.

Received: August 8, 2012 Published online: ■■■, ■■■

Keywords: aromaticity \cdot bridged annulenes \cdot carbaporphyrins \cdot fulvenes \cdot indenes

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Communications

Dicarbaporphyrinoids

T. D. Lash,* A. D. Lammer, G. M. Ferrence ______ IIII - IIII

Two-Step Synthesis of Stable Dioxadicarbaporphyrins from Bis(3indenyl)methane



Porphyrin without N: Bilin analogues and related aromatic dicarbaporphyrinoids have been prepared from bis(3-indenyl)methane. Even though all four pyrrole rings from the porphyrin macrocycle have



been replaced by two furan and two indene subunits, the system retains porphyrin-like UV/Vis spectra and highly diatropic characteristics.

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