ChemComm

COMMUNICATION



View Article Online View Journal | View Issue

Open Access Article. Published on 30 June 2014. Downloaded on 8/6/2025 6:21:54 PM.

Cite this: Chem. Commun., 2014, 50, 9270

Received 4th June 2014, Accepted 28th June 2014

DOI: 10.1039/c4cc04271a

www.rsc.org/chemcomm

Merging of inner and outer ruthenium organometallic coordination motifs within an azuliporphyrin framework[†]

Michał J. Białek and Lechosław Latos-Grażyński*

The insertion of ruthenium(II) into an azuliporphyrin (TPAP) has yielded carbonyl ruthenium(II) azuliporphyrin [Ru(TPAP)(CO)] featuring an equatorial CNNN set of donors. Its azulene moiety serves as the π -coordination platform to accommodate the Ru₄(CO)₉ cluster. This chemistry proved to be general giving rise to a series of bimetallic complexes [M(TPAP){Ru₄(CO)₉}] (M = Ru(CO), Ni, Pd, Pt).

The dynamic development of synthetic routes for carbaporphyrinoids allows for the exploration of porphyrin-like or porphyrin-unlike coordination chemistry.^{1–3} The scope of possible coordination modes includes metal–carbon (sp²) and metal–carbon (sp³) σ -bonds, an agostic side-on interaction between the inner C–H and a metal centre, and an intramolecular metal(π)– η^2 –CC interaction inside a porphyrinoid frame. A unique π -coordination was identified in metalloceneporphyrinoids.⁴ More intricate organometallic structures have been reported which include the tripalladium sandwich complex consisting of two dianionic palladium(π) dicarbaporphyrin units surrounding a palladium(π) cation,⁵ and a gable-porphyrintype organization where two N-confused porphyrins are bridged by the rhodium Rh₄(CO)₄ cluster.⁶

Unconventional coordination motifs can be foreseen presuming that the π -surface of carbaporphyrinoids provides several potential metal binding sites by analogy to the π -complexes reported for tetrapyrrolic ligands where the external coordination involves pyrrole fragments.⁷

Facile introduction of an azulene fragment instead of one pyrrolic unit gives an azuliporphyrin^{8,9} – a highly attractive candidate for exploration of organometallic chemistry inside a carbaporphyrinoid core.¹ Until now, *meso*-tetraaryl and β -alkylated azuliporphyrins have formed organometallic compounds where a direct M–C σ -bond [M = Ni(π), Pd(π), Pt(π), Ir(π)] has been solely



Scheme 1 Reaction of ruthenium dodecacarbonyl with an azuliporphyrin.

detected.^{2,10–12} This study aims to explore the diversity of ruthenium toward azuliporphyrin coordination modes.

A ruthenium(u) ion has been readily inserted into azuliporphyrin **1**, following the procedure reported previously for ruthenium(u) regular porphyrins, to give the corresponding complexes **2** and **3** in which the azuliporphyrin acts as a dianionic organometallic ligand (Scheme 1).

The regular complex 2 can be detected and isolated when 1 is reacted with a sub- or equimolar amount of $[Ru_3(CO)_{12}]$. In the presence of a ruthenium source in excess, only the cluster complex 3 is obtained (see ESI†). This step reveals the unique coordination reactivity of the azulene unit, merging two (inner and perimeter) coordination motifs. Namely, the ruthenium(π) azuliporphyrin binds the $Ru_4(CO)_9$ cluster taking advantage of the azulene π -surface.

Observations of reactivity of compound **3** show essentially three kinds of such toward some nucleophilic agents (deactivated alumina, pyridine, acetonitrile), *i.e.* contraction of a seven-membered ring leading to demetalated benzocarbaporphyrins, secondary cluster reactivity connected with its stepwise substitution and complete cluster removal forming **2**.

The effects of π -coordination are highly pronounced in spectroscopy as well as in the solid state. The ¹H NMR spectra of the tripyrrolic unit, which is common for structures **2** and **3** (Fig. 1), resemble the basic pattern of the azuliporphyrin consistent with its borderline aromaticity.^{8,9} The typical spectrum of **2** affords an AB spin system at $\delta = 7.70$, H(7,18), and δ 7.62 ppm, H(8,17), with a coupling constant (³*J* = 4.9 Hz) typical for a pyrrole ring in porphyrinoids, which is accompanied by a singlet at

Department of Chemistry, University of Wrocław, F. Joliot-Curie 14,

⁵⁰⁻³⁸³ Wrocław, Poland. E-mail: lechoslaw.latos-grazynski@chem.uni.wroc.pl † Electronic supplementary information (ESI) available: Synthetic procedures, figures of spectra (¹H, ¹³C NMR; HRMS; UV-Vis), detailed crystallographic description and data. CCDC 1006123–1006129. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c4cc04271a



δ = 7.69 ppm from H(12,13) of the central pyrrole unit. The NMR resonances of an azulene unit have been assigned to 2, namely: ¹H NMR – 6.80 H(2¹,3¹), 6.38 H(2²,3²), 6.64 H(2³) ppm; ¹³C NMR – 131.5 C(2¹,3¹), 136.9 C(2²,3²), 135.1 C(2³) ppm.

The remarkable relocations of ¹H and ¹³C azulene resonances of 3 (¹H NMR: 3.74 H(2¹,3¹), 4.84 H(2²,3²), 1.84 H(2³) ppm; ¹³C NMR: 63.2 C(2¹,3¹), 66.7 C(2²,3²), 32.3 C(2³) ppm) reflect a cluster attachment. Actually, the change in the azuliporphyrin coordination mode determined for 3 can be readily appreciated by an analysis of coordination shifts defined here as ¹H or ¹³C chemical shift differences between 2 and 3 at chosen positions (Fig. 1). These values: ¹H NMR – 3.06 H(2¹,3¹), 1.54 H(2²,3²), 4.80 H(2³) ppm; ¹³C NMR – 68.3 C(2¹,3¹), 70.2 C(2²,3²), 102.5 C(2³) ppm, clearly point toward the azulene moiety acting as a coordinating unit. Analogous coordination shifts were previously detected in the ¹H NMR and ¹³C NMR spectra of the ruthenium cluster – regular azulene complexes.^{13,14} Coupling constants for azulene protons have also decreased from about 10 Hz, typical for azulene derivatives, to a range of 6–8 Hz.

Further investigation of metal(II) azuliporphyrin coordination abilities against ruthenium has led to a series of bimetallic complexes with the general formula $[M(TPAP){Ru_4(CO)_9}]$, where M = Ni (4), Pd (5), Pt (6). Chemical shift patterns of these complexes resemble those of 3 (Fig. S3, ESI†), and the azulene unit proved to be an unambiguous spectroscopic probe which could be used to identify the perimeter binding.

The structure of azulene-uncoordinated regular complex 2 has been determined for two forms that differ in the sixth axial ligand: 4-(*N*,*N*-dimethylamine)pyridine – DMAP (2-1) or acetonitrile (2-2) (Fig. 2a and b). These two structures show the specific flexibility of the azulene fragment (Fig. 3). Thus, the azulene moiety is notably prone to conformational changes enforced by external stimuli. To describe the situation quantitatively, the value of the angle between the planes generated by the coordination centre (RuCNNN) and the tropylium ring of azulene can be compared. The deflection of azulene from the macrocyclic plane is significant and equals $15.73(13)^{\circ}$ for 2-1 and $28.0(3)^{\circ}$ for 2-2. This much higher deformation for structure 2-2 is consistent with the involvement of



Fig. 2 Presentation of the geometry of the complexes: (a) 2-1, (b) 2-2, (c) 3, (d) 4. Displacement ellipsoids drawn at 30% probability.



Fig. 3 Ball-and-stick representations for fragments of 2-1, 2-2, 3 showing peculiar azulene flexibility.

the seven-membered ring into strong C–H··· π interactions as revealed by analysis of the Hirshfeld surface (Fig. S26, ESI†) as well as the higher deformation of the CNNN coordination centre (Table S2, ESI†). The discussed angle is even higher for structure **3** since it amounts to 43.3(3)°. This is the result of cluster-azulene interactions (Fig. 2c).

Compound **3** features an interesting metallic skeleton that can be described as a distorted tetrahedron (Ru2, Ru3, Ru4, Ru5) with close contact to the ruthenium(II) cation (Ru1) (Fig. 4a). Three out of



Fig. 4 Cluster geometry and porphyrin conformations for (a) **3** and (b) **4**. Phenyl rings removed to preserve clarity of the figure.

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the four ruthenium atoms (Ru2, Ru3, Ru4), which construct the tetrahedron skeleton, are directly coordinated to the azulene moiety and also bonded to two terminal carbonyl ligands.

The fourth ruthenium atom (Ru5) is linked to three basal ruthenium atoms and its coordination sphere is completed by three terminal carbonyl groups. To enable efficient facial coordination to the base of the tetrahedron the azulene moiety is strongly bent as described above.

In fact two modes of cluster coordination have been recognized. The five-membered ring of azulene prefers η^5 -coordination to one Ru2 atom in the standard half-sandwich fashion. The Ru_4CO_9 cluster coordinates all seven carbon atoms of the tropylium moiety (Fig. 2 and Fig. S28, ESI†). The five carbon atoms are bound to the other two ruthenium atoms in a η^4 -coordination mode, with the central carbon atom $C(2^3)$ bonded to two ruthenium atoms. The fundamental features of the azulene–Ru_4(CO)_9 unit encompassed in 3 are reminiscent of the structure seen for the Ru_4(CO)_9 adduct to 1,3-bis(3-methylthienyl)azulene used as the fundamental building block in studies on post-coordination of multinuclear transition metal clusters to azulene-based polymers.^15

In bimetallic complexes [Ni(TPAP){Ru₄(CO)₉}] **4**, [Pd(TPAP){Ru₄-(CO)₉}] **5**, and [Pt(TPAP){Ru₄(CO)₉}] **6**, the azulene moiety reveals the same coordination mode (Fig. S22–S24, ESI[†]). The deflection angle between the coordination core and the tropylium ring is $36.19(9)^{\circ}$ and $32.8(2)^{\circ}$, for **4** and **5**, respectively, and less pronounced for **6** – 24.69(15)° (Table S3, ESI[†]). The changes are significant in comparison to geometry of [Pt(TPAP)] (7) where this angle is $21.8(3)^{\circ}$ and the azulene fragment is twisted in respect to a macrocyclic frame.

Azulene π -coordination results in the folding of a tripyrrolic brace (Fig. 2d and 4b). This can be expressed by the C_{meso}-M–C_{meso} angle between relevant C_{meso} atoms and a central metal cation. The highest degree of folding can be noticed for 4, which results in C5–Ni1–C15 and C10–Ni1–C20 angles equal to 167.28(8)° and 168.24(9)°, respectively. This can be compared to the nearly planar [Pt(TPAP)] (7) (Fig. S25, ESI†), where the analogous angles are close to 180° (178.4(3)° and 178.4(2)°, respectively).

Within the tetrahedron of ruthenium atoms the Ru–Ru distances fall distinctly into two groups (Table S2, ESI[†]). In the first set Ru2–Ru4, Ru2–Ru5, Ru2–Ru3, Ru3–Ru4 distances vary from 2.8749(15) to 2.898(2) Å. The two other Ru–Ru distances are evidently shorter and amount to 2.714(2) (Ru3–Ru5) and 2.6857(16) Å (Ru4–Ru5). These values are within the limits of ruthenium–ruthenium bond lengths reported for a variety of ruthenium clusters, which diverge in the wide range of 2.65–3.20 Å (Fig. S29, ESI[†]). The suitable folding of azulene moiety forces the distinctive adjacency of the terminal Ru2 atom of the cluster and the azuliporphyrinic metal(π) cation. This distance equals 3.1598(18) Å in the case of Ru1–Ru2 and gradually grows in the series Ni1–Ru1 3.3113(5) Å, Pd1–Ru1 3.4074(12) Å, Pt1–Ru1 3.5883(16) Å.

As established by a detailed CSD database search (Fig. S29, ESI†) the Ru1-Ru2 value is within the limits of the longest

distances considered for Ru-Ru bonding interactions detected in ruthenium clusters. The Ru2 atom definitely approaches the Ru1 cation at a distance much shorter than that expected for van der Waals contact. The Ru1-Ru2 distance of 3 can be compared with the longest bonding Ru-Ru contact detected in the first bis(metallabenzene) sandwich complex (bis(η^{6} -3,5-dimethyl-1tricarbonyl-ruthenabenzene)ruthenium) comprising two ruthenabenzene ligands whereas the Ru-Ru distance between the ruthenabenzene rings equals 3.38 Å.¹⁶ In terms of metal-metal bonded metalloporphyrins the structure of 3 provides an unprecedented structural motif as the sixth position is occupied by the ruthenium of the cluster, feasibly involved in a weak Ru1-Ru2 interaction. A distinct feature of the structure is the very pronounced bending of the Ru1-Ru2 axis from the perfect axial position (ca. 45°) as a consequence of strain induced by the incorporation of Ru2 into a Ru₄(CO)₉ scaffold.

The azulene moiety in the ruthenium(II) azuliporphyrin provides the suitable π -surface to bind the Ru₄(CO)₉ cluster. Two conceptually different organometallic motifs are merged in a unique three-dimensional architecture. We proved that these unique coordination abilities of azulene introduced into a carbaporphyrinoid frame are preserved in other suitable systems producing a series of bimetallic complexes [M(TPAP){Ru₄(CO)₉}], where M = Ni, Pd, Pt.

We thank Professor Tadeusz Lis for valuable discussions. Financial support from the NCN (Grant 2012/04/A/ST5/00593) is kindly acknowledged.

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