Spectroelectrochemical Properties and Catalytic Activity in Cyclohexane Oxidation of the Hybrid Zr/Hf-Phthalocyaninate-Capped Nickel(II) and Iron(II) tris-Pyridineoximates and Their Precursors

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Abstract: The in situ spectroelectrochemical cyclic voltammetric studies of the antimony-monocapped nickel(II) and iron(II) tris-pyridineoximates with a labile triethylantimony cross-linking group and Zr(IV)/Hf(IV) phthalocyaninate complexes were performed in order to understand the nature of the redox events in the molecules of heterodinuclear zirconium(IV) and hafnium(IV) phthalocyaninate-capped derivatives. Electronic structures of their 1e-oxidized and 1e-electron-reduced forms were experimentally studied by electron paramagnetic resonance (EPR) spectroscopy and UV−vis−near-IR spectroelectrochemical experiments and supported by density functional theory (DFT) calculations. The investigated hybrid molecular systems that combine a transition metal (pseudo)clathrochelate and a Zr/Hf-phthalocyaninate moiety exhibit quite rich redox activity both in the cathodic and in the anodic region. These binuclear compounds and their precursors were tested as potential catalysts in oxidation reactions of cyclohexane and the results are discussed.

Keywords: clathrochelates; phthalocyanines; zirconium(IV); hafnium(IV); nickel(II); iron(II); spectroelectrochemistry; homogeneous catalysis; oxidation reactions; DFT calculations

1. Introduction

The covalent and coordination molecular assemblies with several electronically coupled metal centers are intensively studied [1–26] because of their prospective application in molecular electronics and light-harvesting (supra)molecular systems. Hybrid mult centered inorganic and organometallic transition metal arrays with relatively isolated π- and σ-electronic systems, which belong to different metal-centered frameworks, have received
a relatively little attention up to date. These polytopic compounds have proven to be potentially useful building blocks for the design of redox- and photoredox-driven molecular electronic devices and artificial photosynthetic systems with long-lived charge separation states [27–33]. In particular, this class of hybrid transition metal complexes includes zirconium(IV) and hafnium(IV), as well as lutetium(III) phthalocyaninate-capped iron(II) cage compounds 1-4 (so-called “phthalocyanininoclathrochelates”), first prepared about 15 years ago [34,35] (Scheme 1). Their molecules contain a phthalocyanine-centered extensive π-system that is coupled with a quasi-aromatic metal-encapsulating macrobicyclic framework. These compounds have been characterized by single crystal X-ray diffraction, spectroscopic and electrochemical methods. Detailed spectroelectrochemical study of the oxidation of these hybrid complexes is also reported [36]. By comparison of their redox potentials with those of the parent zirconium(IV) and hafnium(IV) phthalocyaninates, the first reversible oxidation and reduction waves in the corresponding cyclic voltammograms (CVs) were attributed to the apical phthalocyanine moiety [37,38]. The second quasi-reversible and the third irreversible anodic waves were assigned to the oxidation of the clathrochelate framework based on similarity to the oxidation processes observed in the case of the parent iron(II) clathrochelates [39,40]. To corroborate the experimental electrochemical assignments and the theoretical density functional theory (DFT) calculations, the spectroelectrochemical experiments for the hybrid complexes 1-3 (Scheme 1) have been conducted [36]. Upon oxidation at their first oxidation potential, the intensities of the initial phthalocyaninate-localized Q- and B-bands decreased, while three new absorption bands appeared in the UV−vis spectra of these complexes. The oxidation was accompanied by several isosbestic points, thus suggesting formation of the corresponding phthalocyaninate-centered mono- and di-cation-radicals [41]. In contrast, upon electrolysis at the second oxidation potential, all the phthalocyaninate-centered UV−vis bands characteristic for their monocationic-radicals forms lost their intensities without appearance of new absorption bands. The initial neutral iron(II) phthalocyaninoclathrochelates 1-3 could not be regenerated from the corresponding doubly oxidized macropolycyclic dications. This fact indicates [36] that the doubly oxidized dications are not stable and degrade in solution. Further evidence provided the CV data [35], which revealed quasi-reversibility of the second oxidation process only at high scan rates.

The redox properties of hybrid di- and trinuclear Fe,Zr- and Fe,Hf-porphyrinoclathrochelates 5-8 shown in Scheme 1, which are formed by capping with Lewis-acidic MIV TPP group(s), where MIV is Zr4+ or Hf4+ cation and TPP2− is 5,10,15,20-tetraporphyrinate dianion, were also investigated [42] by various electrochemical and spectroelectrochemical techniques. The results of the CV and differential pulse voltammetry (DPV) experiments suggested that, in the case of the monoporphyrinoclathrochelates 5 and 6, three redox processes could be clearly identified. Two reversible oxidation waves were assigned to the Fe2+/3+ and the TPP1-/2−/q clathrochelate- and porphyrinate-localized redox processes, respectively, while the cathodic wave was attributed [42] to the TPP2−/3− reduction. In the case of the iron(II) diporphyrinatoclathrochelates 7 and 8, the Fe2+/3+ redox process is substantially shifted to lower potentials due to a more electron-donating effect of its porphyrinate ligand fragment, as compared with that of the capping boron atom in the monoporphyrinato-clathrochelate analogs. The porphyrinate-centered oxidation of these tritopic complexes was observed [42] as two closely separated but clearly recognized redox waves. Analogously, two closely spaced porphyrinate-centered reduction waves have also been detected [42]. In general, the UV−vis spectra of these di- and tritopic hybrid complexes suggest a lack of any electronic communication between the clathrochelate and porphyrinate frameworks. The corresponding (spectro)electrochemical data [42] also indicated very weak, if any, long-range electronic coupling between two macroheterocyclic π-systems in the molecules of trinuclear iron(II) diporphyrinoclathrochelates.
Scheme 1. Chemical drawings of the hybrid iron(II) phthalocyanino- and porphyrinoclathrochelates.

The phthalocyanine π-systems are known to possess photo- and electrochromic properties, as well as photocatalytic activity [43–45]. The phthalocyanine-based catalysts [46] can be easily obtained and possess very high chemical robustness even in harsh media. On the other hand, the designed d-metal clathrochelates were reported [47–49] to be useful for protein sensing or efficient electro- and (pre)catalysts for hydrogen and syngas (H₂ + CO) production. A series of the polyamine cobalt(III) cage complexes with tethered mercaptoethylamide terminated oligopeptide apical substituents, allowing for immobilization on a surface of the working gold electrode, was described [50] to be potent heterogeneous electrocatalysts in both the hydrogen evaluation reaction (HER) and the oxygen reduction to hydrogen peroxide. Photodissociation of water was realized on rhodium-doped strontium
titanate photoelectrode surface modified by a cobalt(II) clathrochelate when using methanol as a sacrificial agent [51]. Polyamine cobalt(III) clathrochelates were reported to be active and robust homogeneous catalysts for the oxidation of styrene by H_2O_2 both in acetonitrile solution and in ionic liquid medium [52]. The mononuclear cobalt(II) cage complex of a heteroditopic N_5O_3-macrocyclic ligand was reported [53] to be an efficient catalyst for oxidation of a number of olefins and benzyl derivatives with molecular oxygen or 2-methylpropanal as the oxidants at room temperature and atmospheric pressure. Photoinduced catalytic oxygen production was realized in an aqueous acetonitrile in the presence of the mononuclear copper(II) cage complex with a fluorescent anthracene-containing polyamine macrocyclic ligand as photocatalyst [54]. The unique water-stable iron(IV) clathrochelate [55] was discovered to catalyze the photochemical oxidation of water to dioxygen with high turnover frequency and with record turnover number, working at relatively low overpotential of this redox process [56].

Herein, we report on spectroelectrochemical studies of metal(IV) phthalocyaninate-capped iron(II) and nickel(II) tris-pyridineoximates along with their precursors shown in Scheme 2. In addition, the catalytic activity of 9–16 in homogeneous oxidation reactions of cyclohexane is also reported.

**Scheme 2.** Compounds 9–12 studied in this work (top) and their precursors 13–16 (lower panel).

### 2. Results and Discussion

**Synthesis.** The heterodinuclear hybrid complexes 9–12 (Scheme 2) were prepared in good yields (65–79%) by transmetallation of the monotriethylantimony(V)-capped nickel(II) and iron(II) tris-pyridineoximates 13 and 14 as the reactive complex precursors with zirconium(IV) and hafnium(IV) phthalocyaninates [Zr(Cl_2)Pc] (15) and [Hf(Cl_2)Pc] (16) (see Scheme 2) as the Lewis acids in methanol–dichloromethane solutions/suspensions at room temperature [57]. The identity of the compounds 9–16 was verified by IR spectra (Figures S1–S6) and ESI MS studies.
**ESI mass spectrometry.** Due to their cationic nature, the Ni- and Fe-containing compounds gave strong and clean single m/z-signals in the mass spectra. Their experimental masses and isotopic distributions corresponding to [M]+ matched the theoretical values on the full MS level (Table 1 and Figure 1). The Zr- and Hf-containing Pc precursors 15 and 16 released the chlorides and were detected as methoxido adducts [M − 2Cl + OCH$_3$]+ (Table 1), which stemmed from methanol used as the solvent.

| Compound | Species Observed | m$_{\text{exp}}$ | m$_{\text{theor}}$ |
|----------|------------------|-----------------|-----------------|
| 9        | [M]$^+$          | 1067.16         | 1067.15         |
| 10       | [M]$^+$          | 1155.18         | 1155.20         |
| 11       | [M]$^+$          | 1063.15         | 1063.16         |
| 12       | [M]$^+$          | 1153.17         | 1153.20         |
| 13       | [M]$^+$          | 673.10          | 673.12          |
| 14       | [M]$^+$          | 669.11          | 669.12          |
| 15       | [M − 2Cl + OCH$_3$]$^+$ | 633.07 | 633.07 |
| 16       | [M − 2Cl + OCH$_3$]$^+$ | 723.12 | 723.11 |

**Figure 1.** The experimental and theoretical isotopic distributions of the four investigated hetero-dimetallic complexes (9–12) and their synthetic precursors (13–16).

Each of the ionized compounds was then subjected to fragmentation experiments by collision-induced dissociation (CID). For this purpose, each precursor mass was selected and individually fragmented giving MS$^2$ spectra. The Ni-Sb precursor 13 mass did not yield fragment ions on the MS$^2$ level upon fragmentation. In contrast, the Fe-Sb precursor 14 showed ethyl group loss at the antimony, release of the chelating pyridineoximate ligand and oxidation (Figure S7).

Notably, the fragmentation spectra of 11 and 12 were similar, as were the respective fragment spectra of 9 and 10. This indicates that the nickel(II) and iron(II) tris-
pyridineoximates have a stronger impact on the fragmentation pathways of 9–12 than the respective group 4 transition metals zirconium(IV) and hafnium(IV) (Figures S8 and S9).

The MS² spectra of the Fe-containing 11 and 12 (Figure S8) displayed characteristic fragments suggesting sequential release of the three pyridine ketimine moieties by cleaving the N–O bond and retaining O in the complex, i.e., Δm = 118. Furthermore, the fragment mass of 1082.2 of 12 might be indicative of FeO-release (Δm = 71). This was not observed for the iron analogue 11.

Again, the nickel(II) complexes 9 and 10 yielded less fragments for interpretation. The fragmentation spectra of the nickel-containing 9 and 10 (Figure S9) showed a lower number of fragments, but the mass differences to the precursor ion suggested that the first pyridineoximate ligand was cleaved off intact, Δm = 136. Second, the Δm = 116 indicates the release of a pyridine ketimine ligand accompanied by redox changes on Ni, which was not observed for the Fe complexes. Release of the group 4 metals zirconium and hafnium was not observed. Then, MS³ experiments were performed for the bimetallic complexes by selecting a specific mass of each MS² spectrum, which was again subjected to fragmentation. The fragmentation energy of the MS³ level was slightly higher than at the MS² level. However, the MS³ level did not reveal significant new information about the investigated complexes (Figures S8 and S9).

Cyclic voltammetry, spectroelectrochemistry and DFT calculations. The cyclic voltammograms of complexes 9–12 in acetonitrile (ACN)/n-Bu₄NPF₆ (Figure S10) showed two fully reversible reduction waves and three oxidation events. Similar behavior was recently described for the same complexes in dichloromethane [57]. In the anodic part, the first two nearly reversible redox waves overlap, leading to the double peak shape. The third oxidation step is less reversible. Analogous redox response for all four investigated complexes indicates the ligand based redox locus. The reversibility and redox mechanism in the region of the first two reduction peaks for complex 9 (Ni–Zr) was further investigated by the in situ spectroelectrochemical UV-vis-NIR cyclic voltammetric experiments in ACN/n-Bu₄NPF₆ under argon atmosphere in a thin layer honeycomb spectroelectrochemical cell (Figure 2).

Figure 2. In situ UV-vis-NIR spectroelectrochemistry of complex 9 (Ni–Zr) in acetonitrile (ACN)/n-Bu₄NPF₆ in a thin layer honeycomb spectroelectrochemical cell: potential dependence of UV-vis-NIR spectra with corresponding cyclic voltammogram (scan rate v = 10 mV s⁻¹).
Even by decreasing the scan rate to 10 mV s\(^{-1}\), the cathodic reductions remain reversible and upon scan reversal the products formed upon reduction are fully re-oxidized back to the initial state. The recovery of the initial optical bands upon the voltammetric reverse scan confirms the electrochemical and chemical reversibility of the redox process at the first and the second reduction peak, respectively. It is worth noting that in the region of the first reduction peak, two new optical bands at 585 and 982 nm, with vibronic structure, emerged (Figure 3a).

In addition, a decrease in the intensity of the initial optical bands at 335 and 682 nm, characteristic of the phthalocyaninate unit, indicates that the reduction takes place mainly on this part of the complex. This reduction in the region of the first electron transfer was accompanied by several isosbestic points what is characteristic for the formation of the corresponding phthalocyaninate-centered radicals \[41,58–60\]. This was confirmed by in situ EPR spectroelectrochemistry where a single line EPR spectrum with a \(g\)-value of 2.0049 and a line width \(\Delta H_{pp} = 7.7\) G appeared upon cathodic reduction at the first

**Figure 3.** (a) Evolution of UV-vis-NIR spectra in 2D projection upon forward scan in the region of the first reduction peak for complex 9 (Ni-Zr) in ACN/\(n\)-Bu\(_4\)NPF\(_6\) (black trace: \(-0.85\) V, red trace: \(-0.90\) V, blue trace: \(-0.95\) V, dark cyan trace: \(-1.00\) V, magenta trace: \(-1.05\) V, dark yellow trace: \(-1.10\) V, navy trace: \(-1.15\) V, wine trace: \(-1.20\) V, pink trace: \(-1.25\) V, all vs. Fc\(^+\)/Fc); (b) evolution of UV-vis-NIR spectra in 2D projection upon forward scan in the region of the second reduction peak for 9 (black trace: \(-1.30\) V, red trace: \(-1.35\) V, blue trace: \(-1.40\) V, dark cyan trace: \(-1.45\) V, magenta trace: \(-1.50\) V, dark yellow trace: \(-1.55\) V, navy trace: \(-1.60\) V, wine trace: \(-1.65\) V, pink trace: \(-1.70\) V, olive trace: \(-1.75\) V, royal trace: \(-1.80\) V, all vs. Fc\(^+\)/Fc).
reduction peak (see black trace in Figure 4), confirming large spin delocalization on the phthalocyaninate unit with an extensive \( \pi \)-system. Localization of the unpaired electron on the phthalocyaninate moiety has been further confirmed by the DFT calculations, as can be seen in the corresponding spin density distribution maps (Figure 5).

**Figure 4.** EPR spectra of the 1e-reduced 9 (Ni–Zr) (black trace) and 1e-oxidized 9 (Ni–Zr) (red trace) in ACN/\( \pi \)-Bu\(_4\)NPF\(_6\).

**Figure 5.** B3LYP/LanL2DZ-calculated spin density distributions in the 1e-reduced forms of the complexes 9–12. The isovalue is set to \( \pm 0.002 \).
In the case of 11 and 12, the spin density is localized exclusively on the phthalocyaninate moiety, while in the case of 9 and 10, the spin density is found to be localized on the phthalocyaninate part as well as on the central atom (Ni) and in its vicinity (Figure 5).

At the second reduction peak, the bands of the 1e-reduced form at 585 and 982 nm decreased and a new band at 534 nm appeared. This can be attributed to the 2e-reduced form of phthalocyaninate moiety (Figure 3b). According to the DFT calculations, the triplet state is energetically preferred over the singlet state in these 2e-reduced forms of 9–12, and the corresponding spin density distribution maps are displayed in Figure S11. A similar UV-vis-NIR spectroelectrochemical response was found for Fe–Zr analogue 11, as shown in Figure S6a,b, and also for 12 (Ni–Zr) (Figure S12c,d)

In the region of the first oxidation double-peak, new optical bands at 544 and 852 nm were observed, as shown for 9 (Ni–Zr) in Figure 6, with a simultaneous decrease in the intensity of the initial optical bands at 335 and 682 nm, again indicating that the oxidation takes place mainly on the phthalocyanine part of the complex.

Figure 6. Evolution of UV-vis-NIR spectra in (a) 2D and (b) 3D projection upon cyclic voltammetry (see inset in (a)) in the region of the first oxidation double-peak for complex 9 (Ni–Zr) in ACN/n-Bu4NPF6 at scan rate v = 10 mV s⁻¹.

Although the anodic oxidation is less reversible, as can be seen in the in situ cyclic voltammogram, it was possible to measure the EPR spectrum of the corresponding 1e-oxidized 9 (Ni–Zr) with a g-value of 2.0018, confirming even more delocalized spin density distribution in comparison to the 1e-reduced state (see red trace in Figure 4). Calculated spin density distribution maps of the oxidized forms of 9–12 (Figure S13) are very similar to the ones of their mono-reduced forms (Figure 5).

To confirm the suggested locus of the redox processes, the corresponding molecular precursors 13–16 (Scheme 2) were studied in detail by cyclic voltammetry (Figure S14) and spectroelectrochemistry. The electronic structure of 1e-oxidized and 1e-reduced states was
studied by electron paramagnetic resonance (EPR), UV–vis–near-IR spectroelectrochemistry, and density functional theory calculations.

One fully irreversible cathodic peak was found for metal(II) tris-pyridineoximate precursors 13 and 14 (Scheme 2) in ACN/n-Bu4NPF6 solutions (Figure S14a,b) at much more negative reduction potentials, as observed for the hybrid complexes (see Figure S10), providing evidence that the first two cathodic waves correspond to the reduction of Zr/Hf-phthalocyaninate units 15 and 16 (Scheme 2), which are reduced at much less negative reduction potentials (Figure S14c,d). Note that in the corresponding hybrids, much more reversible redox behavior was found compared to that of the precursors. So, the incorporation of metal(II) tris-pyridineoximates exerts a stabilization effect on the charged Zr/Hf-phthalocyaninate frameworks. Nevertheless, EPR spectroelectrochemistry confirmed the formation of phthalocyaninate centered both 1e-reduced and 1e-oxidized radicals, as shown in Figure S15 for precursor 15. Interestingly, the corresponding 1e-reduced form of 15 exhibits the hyperfine splitting, while the 1e-oxidized form shows the single peak pattern, indicating larger spin delocalisation over the ligand, as confirmed by DFT calculations (see Figure S15, down).

The reversible anodic peak for Fe(II) tris-pyridineoximate 14 in ACN/n-Bu4NPF6 at +0.36 V vs. Fc+/Fc (Figure S14b) corresponds to the Fe2+/Fe3+ redox couple. During in situ UV-vis-NIR spectroelectrochemistry at 10 mV s⁻¹ at the honeycomb platinum working electrode, a less reversible behavior was found, but upon reverse scan, a recovery of the initial optical band at 540 nm, characteristic for the Fe²⁺ state, was observed, indicating rather high stability of the Fe³⁺ form (Figure 7). This band was assigned to the Fe(d)—Ligand(π*) MLCT in the iron(II)-containing tris-pyridineoximate fragment [57], which indicates a noninnocent character of the ligand. This corresponds well to the shape of HOMO orbitals of 13 and 14 (see Figure S16 and Figure 8, respectively) and is in agreement with the noninnocent character of the tris-pyridineoximate moiety.

Figure 7. Evolution of UV-vis-NIR spectra in (a) 2D (forward scan, black trace: 0.15 V, red trace: 0.20 V, blue trace: 0.25 V, dark cyan trace: 0.30 V, magenta trace: 0.35 V, dark yellow trace: 0.40 V, navy trace: 0.45 V, wine trace: 0.50 V, pink trace: 0.55 V, olive trace: 0.60 V, all vs. Fc+/Fc) and (b) 3D projection during cyclic voltammetry (see the corresponding voltammogram in Inset of (a), scan rate 10 mV s⁻¹) of 14 in the region of the first oxidation peak in ACN/n-Bu4NPF6.
The Ni(II) tris-pyridineoximate precursor 13 in ACN/\textit{n}-Bu$_4$NPF$_6$ showed an irreversible oxidation at $E_{\text{pa}} = 0.53$ V vs. Fc$^+/\text{Fc}$ that could be assigned to the Ni$^{2+}$/Ni$^{3+}$ redox couple. Analogous irreversible intense cathodic peaks at $E_{\text{pc}} = −1.93$ V vs. Fc$^+/\text{Fc}$ found for both M(II) tris-pyridineoximates 13 and 14 can be assigned to the 2e-reduction leading to the irreversible decomposition of the complexes, as evidenced by UV-vis-NIR spectroelectrochemistry for 14 (Figure S17).

**Catalytic studies.** The catalytic activity of dinuclear zirconium(IV) and hafnium(IV) phthalocyaninate-capped nickel(II) and iron(II) tris-pyridineoximate complexes 9–12 and their precursors 13–16 was tested for the peroxidative oxidation of cyclohexane as a model reaction. In fact, cyclohexane oxidation into cyclohexyl hydroperoxide and its deperoxidation into cyclohexanol and cyclohexanone are among the most important processes in the chemical industry, with the KA oil (cyclohexanone (K) + cyclohexanol (A)) mixture being the substrate for subsequent oxidation into adipic acid using nitric acid [61].

The catalytic experiments were initiated at room temperature and then by conventional thermal heating up to 60 °C, using an aqueous solution of tert-butyl hydroperoxide (THBP, 70%) as oxidizing agent (in view of its lower handling risk), and a low catalyst amount in ACN (see experimental).

The heterodinuclear hybrid complexes 9–12 exhibited some activity at room temperature (e.g., up to 9% KA oil yield for the Hf-Fe complex 12 after 6 h reaction), which was significantly enhanced by performing the oxidation reactions at 60 °C for the same time (6 h). Further increase in the temperature was avoided for safety reasons. The KA oil yields obtained under the optimized mild conditions are depicted in Figure 9.
Figure 9. Effect of reaction time on the yield of KA oil obtained by oxidation of cyclohexane catalyzed by complexes 9–16. Optimized mild reaction conditions: catalyst (5 µmol), cyclohexane (2.5 mmol), tert-butyl hydroperoxide (TBHP) aq. sol. 70% (5.0 mmol), temperature = 60 °C (thermal heating) in ACN (3 mL). %Yield = moles of cyclohexanone + cyclohexanol per 100 mol of cyclohexane (gas chromatography (GC) yield after treatment with PPh₃).

The Hf-Fe complex 12 led to the highest KA oil yield (up to 36.9%) at 60 °C after 6 h reaction, suggesting a synergic effect on the catalytic activity of the heterodinuclear species. In fact, per se, the iron and hafnium complexes 14 and 16 yielded maxima of 20.6 and 15.9% of KA oil, respectively.

A similar synergism, although not so much pronounced, is observed for the Zr-Fe complex 11, which exhibits a higher activity than those of the precursor iron (14) and zirconium (15) compounds (e.g., KA oil yields of 26.3, 20.6 and 11.1%, respectively, after 6 h at 60 °C). However, eventual synergic effects are much less significant in the cases of the Hf-Ni (10) and Zr-Ni (9) dimetallic complexes. This suggests that the synergism is mainly based on the combination of the metals and not of the corresponding ligands.

The neutral compounds 15 and 16 (hence without the potentially dangerous perchlorate anion) were further tested for the MW-assisted oxidation of neat cyclohexane and, at the above optimized conditions (60 °C), afforded up to 26% yield of cyclohexanol and cyclohexanone mixture (for complex 16) after 3 h under irradiation (Figure 10). Thus, the microwave irradiation appears to enhance the catalytic activity of both 15 and 16, leading to significantly higher KA oil yields (26.0 vs. 11.6% and 19.1 vs. 7.0%, respectively, for 16 and 15 after 3 h reaction) than those obtained under thermal heating (compare Figures 9 and 10). This is in accord with the previously observed behavior of other catalytic systems [62–64], where the use of MW irradiation provided a much more efficient synthetic method than conventional thermal heating. Due to the perchlorate counteranion of compounds 9–14, their catalytic performance comparison under MW conditions was avoided.
Figure 10. Effect of reaction time on the yield of KA oil obtained by microwave (MW)-assisted oxidation of neat cyclohexane catalyzed by complexes 15 and 16. Reaction conditions: catalyst (5 µmol), cyclohexane (2.5 mmol), TBHP aq. sol. 70% (5.0 mmol), temperature = 60 °C. %Yield = moles of cyclohexanone + cyclohexanol per 100 mol of cyclohexane (GC yield after treatment with PPh₃).

To date, the use of hafnium or zirconium as catalysts for the oxidation of cyclohexane is scarce. The first mention, in 1990, is a patented [65] process for the production of adipic acid by contacting cyclohexane with air at 80–160 °C and 2–100 bar in the presence of acetic acid, that used Zr and Hf co-catalysts (0.001–1000 atomic ratio Zr:Co, Hf:Co or (Zr+Hf):Co), making it possible to avoid the requirement for expensive halogen resistant construction materials. Two decades later, zirconium-porphyrinic iron-organic frameworks, with tetrakis(4-carboxyphenyl)porphyrin (TCPP), were applied by Zhou et al. [66], as catalysts for the cyclohexane oxidation to KA oil in the presence of TBHP (in decane) at 65 °C, but no catalytic activity was observed for the Zr–porphyrinic (without iron) compound. The reactivity was attributed to the high-density of accessible active porphyrinic iron(III) centres within the porous framework. Transition metal exchanged alpha-zirconium phosphates, alpha-ZrP centre dot M (where, M = Mn(II), Cu(II) or Fe(III)) were also tested [67] for the oxidation of cyclohexane in liquid phase with TBHP. The order of reactivity of alpha-ZrP centre dot M was: alpha-ZrP centre dot Mn(II) > alpha-ZrP centre dot Cu(II) > alpha-ZrP centre dot Fe(III). A maximum of 6% conversion and 100% selectivity for KA oil was observed with the alpha-ZrP centre dot Mn(II)/TBHP system after 5 h of reaction, which could be recycled three times. Very recently [68], the magnetic ZrFe₂O₄@SiO₂-TCPP nanocatalyst afforded a maximum of cyclohexanone and cyclohexanol products of 33.6 and 18.9%, respectively, with the advantage of being recovered by a magnetic field.

In the above cases, Zr is not the sole metal in the catalyst composition which also includes another transition metal with known catalytic activity in cyclohexane oxidation. Remarkably, in our study, besides the Zr and Hf heterobimetallic (9–12) complexes, we were also able to successfully use Zr and Hf monometallic neutral complexes (15 and 16) as efficient catalysts (yields up to 26% under MW irradiation, see Figure 10) for the selective oxidation of cyclohexane to the cyclohexanol and cyclohexanone mixture. Note that the current cyclohexane oxidation is a low-efficiency industrial process, with conversions lower than 10% to ensure a selectivity of 80% for the cyclohexanol/cyclohexanone mixture [69]. Thus, our catalytic outcomes represent a significant achievement in catalyst design for the industrially important cyclohexane oxidation.

The performed catalytic experiments support that cyclohexane oxidation catalyzed by our complexes proceeds through a mechanism of radical nature [70] where the catalyzed decomposition of t-BuOOH with formation of t-BuO and t-BuOO radicals is crucial for the
H-abstraction from cyclohexane, the beginning of its conversion through the radical pathway. Thus, the availability of reducible complexes (proved by the spectroelectrochemical and DFT studies) would be a key requirement for the oxidation to occur.

3. Materials and Methods

General procedures and measurements. All reactions were performed under inert atmosphere using the standard Schlenk techniques. All reagents were obtained from the commercial sources and used without additional purification. Hybrid complexes 9–12 and their precursors 13–16 were prepared as described previously [57]. The IR spectra of 9–14 (see Figures S1–S6) were in accord with previously reported data [57]. Caution: Complexes containing perchlorate should be handled with care.

Electrochemical, spectroelectrochemical and related EPR experiments. A Heka PG310USB (Lambrecht, Germany) potentiostat with a PotMaster 2.73 software package served as the potential control in cyclic voltammetric studies. Cyclic voltammetric experiments in acetonitrile (ACN) in the presence of n-Bu4NPF6 supporting electrolyte (puriss quality from Fluka) were performed under argon atmosphere using a three-electrode arrangement with platinum 1 mm disc working electrode (from Ionode, Tennyson, Australia), platinum wire as counter electrode, and silver wire as pseudo-reference electrode. Ferrocene (from Sigma-Aldrich, St. Louis, MO, USA) served as the internal potential standard. In situ ultraviolet-visible-near-infrared (UV-vis-NIR) spectroelectrochemical measurements were performed on a spectrometer Avantes (Model AvaSpec-2048 14-USB2, Apeldoorn, Netherlands) in the spectroelectrochemical cell kit (AKSTCKIT3) with the Pt-microstructured honeycomb working electrode, purchased from Pine Research Instrumentation (Durham, NC, USA). Halogen and deuterium lamps were used as light sources (Avantes, Model AvaLight-DH-S-BAL). The cell was positioned in the CUV-UV cuvette holder (Ocean Insight, Ostfildern, Germany). Optical spectra were processed using the AvaSoft 7.7 software package. Amperostatic electrochemical in situ EPR experiments were performed in acetonitrile solutions containing 0.5 mM of a given complex and 0.2 M of n-Bu4NPF6. The freshly prepared solutions were carefully purged with argon and inserted in a Varian electrolytic cell equipped with a large platinum mesh. The EPR spectra were measured in situ using an X-band EPR spectrometer EMX (Bruker, Karlsruhe, Germany) with 100 kHz field modulation.

Electrospray Ionization Mass Spectrometry. The compounds 9–12, as well as their precursors 13–16, were dissolved in methanol to a concentration of 1–5 µM and directly injected into the mass spectrometer. The compounds were analyzed intact and after fragmentation by collision-induced dissociation (CID) on MS2 and MS3 levels. The analysis was performed on a Bruker AmaZon Speed ETD using trapControl Version 7.1 and Data Analysis 4.0 SP5 (Bruker Daltonics GmbH, Bremen, Germany). The compounds were infused at a flow rate of 3 µL min−1. The data were acquired in the positive ion mode employing the following parameters: capillary voltage −4.5 kV, nebulizer 3 bar, dry gas 5 L min−1, dry temperature 180 °C. The mass range was 100–2200. Fragmentation experiments by collision-induced dissociation (CID) were performed using amplification energies of 1.5–2.2 on the MS2 level and MS3 experiments using amplification energies of 2.8–4.5. The low-level cut-off was 20% of the precursor mass by default.

Computational details. Geometry optimizations of the studied complexes (and their precursors) were performed at the B3LYP [71–74] /LanL2DZ [75,76] level of theory using the Gaussian09 program package [77]. The vibrational analysis was employed to confirm that the optimized geometries correspond to the energy minima (i.e., no imaginary vibration presented). Visualization of the optimized structures and the spin density distributions was performed using the Molekel software suite [78].

Catalytic studies. The catalytic tests were performed at room temperature or at 60 °C in a thermostated Pyrex round bottom flask and in open atmosphere, or under microwave (MW) irradiation. The MW experiments were undertaken in a focused Anton Paar Monowave 300 microwave incorporating a rotational system and an IR temperature detector (Anton
Paar GmbH, Graz, Austria), using a 10 mL capacity reaction tube with a 13 mm internal diameter. Complexes 9 to 16 were used. (Caution: the combination of perchlorates with many oxidizable substances may be explosive; dry perchlorates at elevated temperatures may be explosive!).

The peroxidative oxidation reactions were carried out as follows: 1–10 µmol of the complexes 9–16 were added to ACN (3.00 mL) with vigorous stirring, whereafter 2.50 mmol of cyclohexane (270 µL) and 5.00 mmol (690 µL) of tert-butyl hydroperoxide (TBHP, 70% aqueous solution) were added and the reaction solution was stirred for 0.5–3 h at the desired temperature (from room temperature to 60 °C). Blank tests were performed in metal complex-free conditions.

The products analysis was performed as follows: 90 µL of cycloheptanone (internal standard), 10.00 mL of diethyl ether (to extract the substrate and the organic products from the reaction mixture) and an excess of triphenylphosphine were added. The obtained mixture was stirred for ca. 10 min and then a sample was taken from the organic phase and analysed by gas chromatography (GC).

GC measurements were carried out using a FISONS Instruments GC 8000 series gas chromatograph with a flame ionization detector and a capillary column (DB-WAX, column length: 30 m; internal diameter: 0.32 mm) and the Jasco-Borwin v.1.50 software (Jasco, Tokyo, Japan). The temperature of injection was 240 °C. The initial temperature was maintained at 100 °C for 1 min, then raised at 10 °C/min to 180 °C and held (at this temperature) for 1 min. Helium was used as the carrier gas. All products obtained from the catalytic oxidation reactions were identified by their retention times (confirmed with those of commercially available samples) and their quantification was attained by the internal standard method.

4. Conclusions

The ESI mass spectrometric study of 9–12 revealed pronounced similarity in fragmentation behavior of 9 and 10 as well as 11 and 12. In particular, a sequential release of the three pyridine ketimine moieties was typical for the last two complexes.

The investigated hybrid molecular systems 9–12 exhibit quite rich redox activity both in the cathodic (two reversible reductions) and in the anodic region (two reversible and one quasi-reversible oxidations). EPR spectroelectrochemistry and DFT calculations indicate that the first reduction is due to the formation of phthalocyaninate-centered radical, while the second can be attributed to the 2e-reduced form of phthalocyaninate moiety. The dinuclear zirconium(IV) and hafnium(IV) phthalocyaninate-capped nickel(II) and iron(II) tris-pyridineoximate complexes 9–12 and their precursors 13–16 were found to catalyze the peroxidative oxidation of cyclohexane into cyclohexanol and cyclohexanone. The catalytic activity of 9–12 was low at room temperature, but increased significantly at 60 °C. The Hf-Fe complex 12 showed the highest KA oil yield (up to 36.9%) after 6 h reaction time (Figure 9).

The yield was higher than that achieved in the presence of iron and hafnium precursors 14 and 16 (20.6 and 15.9% of KA oil, respectively), implying a synergistic effect of the two metals in 12 on its catalytic activity. Less pronounced synergism was observed for other heterodinuclear complexes (9–11), but to a markedly lesser extent. Further temperature increase was avoided because of safety reasons. The synthesis of similar complexes to 9–12 with other than perchlorate counteranions and their testing at higher temperatures deserves attention. In this context, it should be stressed that the neutral compounds 15 and 16, in which the counteranion is absent, have shown significantly higher KA oil yields in the MW-assisted oxidation of neat cyclohexane at 60 °C for 3 h (26.0 vs. 11.6 and 19.1 vs. 7.0%, respectively, for 16 and 15) compared to those obtained by conventional thermal heating (Figure 10). The performed catalytic studies are in agreement with the mechanism of radical nature and availability of reducible complexes confirmed by spectroelectrochemical and theoretical DFT calculations.
Supplementary Materials: The following are available online, IR spectra Figure S1: IR spectrum of 9, Figure S2: IR spectrum of 10, Figure S3: IR spectrum of 11, Figure S4: IR spectrum of 12, Figure S5: IR spectrum of 13, Figure S6: IR spectrum of 14, the additional mass spectra with associated fragmentation experiments Figure S7: The full mass spectra (top) and the associated fragmentation experiment MS2 (bottom) upon isolation of the given precursor are shown for the synthetic precursor

(Fe–Sb). The corresponding Ni–Sb complex 13 did not yield interpretable fragmentation spectra, Figure S8: The full mass spectra (top) and their associated fragmentation experiments MS2 (middle) and MS3 (bottom) upon isolation of the given precursor are shown for 11 (Zr–Fe) and 12 (Hf–Fe), Figure S9: The full mass spectra (top) and their associated fragmentation experiments MS2 (middle) and MS3 (bottom) upon isolation of the given precursor are shown for 9 (Zr–Ni) and 10 (Hf–Ni). CVs for 9–12 Figure S10: Cyclic voltammograms for complexes (a) 9, (b) 10, (c) 11 and (d) 12, in ACN/nBu4NPF6 (Pt-disc working electrode, scan rate 100 mV s⁻¹), B3LYP/LanL2DZ spin density distributions in the double-reduced forms of the 9–12 Figure S11: B3LYP/LanL2DZ Spin density distributions in double-reduced forms of complexes 9–12. The isovalue is set to ±0.002. Energy differences between the singlet and triplet electronic states (ΔEgs) are 0.035, 0.034, 0.0035 and 0.0034 a.u. for complexes 9, 10, 11 and 12, respectively. The in situ UV-vis-NIR spectra of 11 and 12 in the region of the first and the second reduction waves Figure S12: In situ UV-vis-NIR spectroelectrochemistry of complex 11 (Fe–Zr) in the region of (a) the first (and b) the second cathodic peak, as well as of complex 12 (Fe–Hf) in the region of (c) the first and (d) the second cathodic peak, all in ACN/nBu4NPF6 in a thin layer honeycomb spectroelectrochemical cell, B3LYP/LanL2DZ spin density distributions in the monooxidized forms of 9–12 Figure S13: B3LYP/LanL2DZ spin density distributions in 1e-oxidized forms of complexes 9–12. The isovalue is set to ±0.002, CVs of 13–16 Figure S14: Cyclic voltammograms for the corresponding molecular precursors 13–16 of investigated hybrid Zr/Hf-phthalocyaninate-capped metal(II) tris-pyridineoximates 9–12 in ACN/nBu4NPF6 solutions (Pt-disc working electrode, scan rate 100 mV s⁻¹): (a) 13, (b) 14, (c) 15 and (d) 16, EPR spectra and the spin density distributions for the mono- and di-reduced forms. Figure S15: Top: EPR spectrum of 1e-reduced precursor 15 (black trace) and 1e-oxidized 15 (red trace) in ACN/nBu4NPF6. Bottom: B3LYP/LanL2DZ spin density distributions in 1e-reduced (left) and 1e-oxidized (right) precursor 15. The isovalue is set to ±0.002, frontier orbitals of 13 Figure S16: B3LYP/LanL2DZ HOMO–1, HOMO, LUMO, and LUMO+1 orbitals of the precursor 13, the corresponding relative orbital energies in a.u. are given in parenthesis. The isovalue is set to ±0.02, evolution of UV-vis-NIR spectra of the precursor 14 upon the corresponding CV experiment in the region of the first reduction wave Figure S17: Evolution of UV-vis-NIR spectra in (a) 2D (forward scan) and (b) 3D projection during cyclic voltammetry (see the corresponding voltammogram in Inset of (a), scan rate 10 mV s⁻¹) of 14 in the region of the first reduction peak in ACN/nBu4NPF6.

Author Contributions: Y.Z.V., S.V.D. and S.A.B. prepared the complexes and have written the introduction part, S.M.M.–M., D.G., D.S., C.-M.M., M.–A.L. performed and interpreted the ESI MS measurements, P.R. and D.D. performed electrochemical and spectroelectrochemical investigations, M.M. performed DFT calculations, A.J.L.P. and L.M.D.R.S.M. carried out the catalysis, V.B.A. supervised the project and was involved in writing the manuscript. All authors have read and agreed to the published version of the manuscript.

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