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Series of Near-IR-Absorbing Transition Metal Complexes with Redox Active Ligands

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Abstract: New soluble and intensely near-IR-absorbing transition metal (Ti, Zr, V, Ni) complexes were synthesized using a redox non-innocent \( N,N' \)-bis(3,5-di-tertbutyl-2-hydroxy-phenyl) -1,2-phenylenediamine (\( H_4L \)) as a ligand precursor. In all the complexes, ([Ti(\( L_ox \))\(_2\)], [Zr(\( L_ox \))\(_2\)], [V(\( L_{sq1} \))(HL\( L_ox \))]) and [Ni(HL\( L_ox \))\(_2\)], two organic molecules coordinate to the metal center as tri- or tetradentate ligands. The solid-state structures of the complexes were determined using single crystal XRD, and the compounds were further characterized with Electrospray Ionisation Mass Spectrometry (ESI-MS). Thermoanalytical measurements indicated the thermal stabilities of the complexes. All compounds absorb strongly in the near-IR region and show very interesting magnetic and electrochemical properties. Moreover, it was shown that the V and Ni complexes can also convert absorbed near-IR photons to (un)paired electrons, which indicates great promise in photovoltaic applications.

Keywords: metal organic complex; redox-active ligand; non-innocent ligands

1. Introduction

In coordination chemistry, a ligand is referred as ‘non-innocent’ when it forms complexes where the oxidation state of the central atom cannot be absolutely defined [1]. For example, o-aminophenol \( N,N' \)-bis(3,5-di-tertbutyl-2-hydroxy-phenyl)-1,2-phenylenediamine \( H_4L \) (the oxidation states of \( L \), Scheme 1) has a rich redox-chemistry and its complexing abilities towards transition metals, i.e., Ti, Zr, Hf, Mo, W, Mn, Co, Cu and Zn, have attracted substantial interest over the past few years [2–8]. Earlier studies show that this proligand can form complexes with various transition metals in 1:1 and 2:1 stoichiometry. Moreover, the galore electrochemical behavior of \( H_4L \) (from hereon \( L \) corresponds to the ligand in any oxidation state) allows five different oxidation states (Scheme 1). The foremost motivation for research on such redox-active ligands has been in the development of new homogenous catalysts, which could utilize these ligands as electron reservoirs during the catalytic cycle. On the other hand, the redox activity of the ligand may allow the corresponding metal complexes to have multiple low-energy charge transfer processes within, thus rendering the complex able to absorb electromagnetic radiation at the visible and the near-IR regions intensely.

There are many metal-organic transition metal complexes with redox-active ligands which are studied for their potential applicability for strongly near-infrared (NIR)-absorbing dyes [9–19] or as single molecule magnets [20–23] (SMMs). NIR absorption is important for dye-sensitized solar
cells [20–23], camouflaging, optical NIR filtering and for NIR photodetectors. Metal organic NIR dyes are also studied for their high thermal durability and lightfastness compared with organic NIR dyes [24]. As H₄L is a redox-active ligand that can have, in certain protonation states, delocalized \( \pi \)-conjugated systems, it can form stable radicals upon complexation with transition metal ions [8]. Stable radicals, i.e., compounds carrying unpaired electrons, may have strong interactions with photons and therefore exhibit strong absorption at UV-Vis-NIR range. In this study, we have used H₄L as a ligand with the aim of preparing new NIR-absorbing transition metal complexes and have studied their molecular structures as well as their magnetic properties.

![Scheme 1. The different oxidation states of the deprotonated ligand L (adapted from the reference [8]).](image)

2. Results and Discussion

2.1. Synthesis of Complexes

The reaction between metal precursors and H₄L in basic methanol solutions under ambient atmosphere afforded the dark crystals of mononuclear complexes (Scheme 2). All the reactions yielded mononuclear complexes with a 2:1 ligand-to-metal ratio regardless of the equivalent ratios of the starting compounds. The complexes were isolated as dark crystals either directly from the reaction mixture or with subsequent crystallization. The synthesis of [V(L\(^{ox1}\))(HL\(^{ox}\))] was repeated using vanadyl acetylacetonate as a metal precursor to yield a chemically identical product, which crystallized in an orthorhombic crystal system. No further analyses were run utilizing the orthorhombic form. Powder X-ray diffraction (PXRD) was used to establish that the material used for all analyses corresponds to the monoclinic (C2/c) structure (see Supplementary Materials for PXRD analysis and the crystal structure of the orthorhombic polymorph). All complexes are air- and moisture-stable at room temperature. In the DSC-TGA (Differential Scanning Calorimetry-ThermoGravimetric Analysis) measurements, which were carried out with a heating rate of 5 K/min for all complexes in nitrogen atmosphere, it was found that the Ti complex loses ca. 8% of the sample’s weight, which could be attributed to the loss of solvate molecules from the crystal lattice (XRD measurements, see below). The weight then remains stable until ca. 270 °C, where it starts to rapidly decline, probably due to the decomposition of the molecule. The other complexes were found to be stable up to 340 °C, 250 °C and 230 °C, for Zr, V and Ni complexes, respectively, until decomposition occurred.
2.2. Structural Studies

The single crystal X-ray diffraction studies (see Supplementary Materials, Table S1) showed that both Group IV complexes crystallize in a triclinic unit cell with two distinct complex molecules in the asymmetric unit. Ti complex crystallizes as an acetonitrile solvate, whereas Zr complex has a large, ca. 1000 Å³, void space in the crystal lattice with unresolved electron density. This void could potentially be large enough to accommodate an additional uncoordinated ligand. This is, however, not supported by ¹H-NMR spectroscopic evidence which shows a methanol solvent peak instead. Therefore, the electron density in the void was treated as solvent using Olex² solvent mask feature. In both cases, the complexes are formed of neutral molecules in which two nearly identical organic ligands are fully deprotonated (N-H and O-H protons) and coordinated to the metal center as tetradentate ONNO-donors. The coordination geometry around the central atom is best described as a distorted square antiprism with phenolic oxygens coordinated to the central atom in a mer form. The C-N and C-C interatomic distances in the central phenylene rings of each distinct ligand suggest a localized, i.e., a ‘cyclohexadiene diimine’-like, structure rather than a delocalized system (bond parameters are presented in Table 1). The four Ti-N bonds (Table 1) display significant elongation compared to the previously published [Ti(L°red)(py)₂] complex but are very similar to the [Ti(L°ox)(Cl)₂] complex [4]. Although the long Ti-N distances can be caused by ligand-ligand steric effects, the discussed interatomic distances strongly suggest that the ligands in the presented [Ti(L)₂] complex adopt the [L°ox]²⁻ form. Furthermore, according to Brown, it is possible to use the metrical oxidation state (MOS) to estimate the formal oxidation states of the metal-coordinated o-aminophenol moieties based on their geometrical parameters, i.e., the bond lengths of O-C and N-N bonds and the C-C bonds of the phenyl ring [25] or [Ti(L)₂], these calculations gave the MOS values of -1.09(11), -1.22(6), -1.22(9) and -1.37(8), which support the idea that both ligands adopt the partially oxidized [L°ox]²⁻ form. The overall
structure of the \([\text{Zr}(\text{L})_2] \) complex is very similar to that of its Ti analogue (Figure 1), and the bonding parameters (Table 1) likewise indicate that the organic ligands adopt the \([\text{L}^\infty]^{2-}\) oxidation state. Hence, both complexes can be described as \([\text{M}(\text{Lox})_2]\), although the MOS values of the \(\text{o-aminophenol moieties}\) in \([\text{Zr}(\text{Lox})_2]\) are slightly higher \([-1.21(14), -1.39(18), -1.40(14)\) and \(-1.41(12)]\) compared to \([\text{Ti}(\text{Lox})_2]\). However, it is known that complexes of high oxidation state metals may have strong \(\pi\)-donation from the high-lying \(\pi\) orbitals of amidophenoxide ligands, which may cause the metrical oxidation state to differ from the theoretical value \([25]\). It therefore seems obvious that the formal oxidation state of both Ti and Zr centres should be assigned as +4.

**Table 1.** Selected bond lengths (Å) for \([\text{Ti}(\text{Lox})_2]\) and \([\text{Zr}(\text{Lox})_2]\).

|          | Ti1A-O1A | 1.977(2) | Zr1A-O1A | 2.113(4) |
|----------|----------|----------|----------|----------|
| Ti1A-O2A | 2.019(2) | Zr1A-O2A | 2.131(4) |          |
| Ti1A-O3A | 2.016(2) | Zr1A-O3A | 2.096(3) |          |
| Ti1A-O4A | 2.022(2) | Zr1A-O4A | 2.134(3) |          |
| Ti1A-N7A | 2.318(2) | Zr1A-N7A | 2.382(4) |          |
| Ti1A-N14A| 2.198(3) | Zr1A-N14A| 2.325(4) |          |
| Ti1A-N27A| 2.282(3) | Zr1A-N27A| 2.367(4) |          |
| Ti1A-N34A| 2.228(3) | Zr1A-N34A| 2.319(4) |          |
| N7A-C6A  | 1.379(4) | N7A-C6A  | 1.377(7) |          |
| N7A-C8A  | 1.333(4) | N7A-C8A  | 1.333(6) |          |
| N14A-C13A| 1.327(4) | N14A-C13A| 1.346(6) |          |
| N14A-C15A| 1.377(4) | N14A-C15A| 1.379(6) |          |
| N27A-C26A| 1.361(4) | N27A-C26A| 1.399(13)|          |
| N27A-C28A| 1.350(4) | N27A-C28A| 1.342(6) |          |
| N34A-C33A| 1.340(4) | N34A-C33A| 1.340(6) |          |
| N34A-C35A| 1.365(4) | N34A-C35A| 1.387(6) |          |

**Figure 1.** The crystal structures of \([\text{Ti}(\text{Lox})_2]\) and \([\text{Zr}(\text{Lox})_2]\). The hydrogen atoms and tert-butyl groups are omitted for clarity. Thermal ellipsoids are drawn at the 30% probability level.

According to the single crystal XRD analysis, the vanadium complex is formally a neutral species, where two ligands show different coordination modes and protonation states, i.e., \([\text{V}(\text{L})(\text{HL})]\). One of the two distinct ligands is coordinated to the central metal ion as a fully deprotonated tetradentate ONNO-donor, whereas the second ligand is coordinated as a partially deprotonated tridentate ONN-donor. The sole phenolic hydroxyl group that remains protonated and uncoordinated forms a hydrogen bond to the phenolate oxygen of the four-dentate ligand \([d(O1A···O4A = 2.967 \text{ Å})]\). The phenolic oxygens are aligned in a \(\text{H}_{4}f\)-fashion and the geometry of the central V atom is best described as a distorted pentagonal bipyramid with the tridentate ligand in a planar geometry (dihedral V-N-N-O angle is \(-4.6^\circ\)) and the tetradentate clearly twisted (dihedral O-N-N-O angle is \(37.0^\circ\)). Interestingly, the crystal packing of \([\text{V}(\text{L})(\text{HL})]\) is isostructural with the known Mo\(^{6+}\) and W\(^{6+}\) complexes.
complexes which have the formula \([M(L)_{\text{red}}(HL)_{\text{sq1}}]\) \([5,6]\), even if the formal oxidation state in the vanadium complex is apparently lower.

Like the isostructural Mo and W complexes, \([V(L)(HL)]\) consists of two distinct molecular units in the asymmetric unit. The intramolecular bond parameters between these units are very similar and thus the complex \(V1A\) is taken as an example while discussing the bonding characteristics of the complex. The intramolecular distances are shown in (Table 2, Figure 2). The C–C bond distances in the central six-membered \(C_6\) rings indicate that the tridentate ligand bears a cyclohexadiene backbone whereas the tetradeptate ligand shows a more delocalized \(C_6\) system (Table 2). This is also reflected in the C–N bond distances, which for the tridentate ligand are ca. 0.05–0.07 Å shorter compared with the tetradentate ligand and are closer to values that would be expected for an imine. MOS calculations for the two distinct \(\sigma\)-aminophenols of the tetradeptate ONNO ligand gave the values of \(-1.65(12)\) and \(-1.79(11)\), thus yielding an approximate formal oxidation state of \(-3\) for the entire ligand. The fully coordinated \(\sigma\)-aminophenol moiety of the tridentate ONN ligand, on the other hand, yields a MOS value of \(-1.10(6)\). This information, combined with the magnetic measurements (vide infra), indicates that the complex is expected to have one tetradeptate and triply anionic \([L_{\text{sq1}}]^{3-}\) ligand, with an unpaired electron, as well as one tridentate, partly protonated and singly anionic \([HL_{\text{o}}]^{-}\) ligand and \(V^{4+}\) \(d^1\) central metal ion. Hence, although \([V(L_{\text{sq1}})(HL_{\text{o}})]\) shows remarkable structural resemblance to its formerly synthesized Mo \([7]\) and W \([6]\) analogues, the oxidation states of both chelating ligands are different to the group VI metal complexes reported earlier. This reveals that the overall structure of the ligand is not entirely dependent on the oxidation state.

**Figure 2.** The crystal structures of \([V(L_{\text{sq1}})(HL_{\text{o}})]\) and \([Ni(HL_{\text{o}})_2]\) with the intramolecular hydrogen bonds drawn with dashed lines. The C–H hydrogen atoms and tert-butyl groups are omitted for clarity. Thermal ellipsoids are drawn at the 30% probability level.

**Table 2.** Selected bond lengths (Å) for \([V(L_{\text{sq1}})(HL_{\text{o}})]\) and \([Ni(HL_{\text{o}})_2]\).

| Bond         | Value    | Bond         | Value    |
|--------------|----------|--------------|----------|
| V1A-O1A      | 3.571(2) | Ni1-O1       | 2.0616(16)|
| V1A-O2A      | 1.971(19)| Ni1-O2       | 3.548(2) |
| V1A-O3A      | 1.949(19)| Ni1-O3       | 2.0315(14)|
| V1A-O4A      | 2.0544(19)| Ni1-O4      | 3.476(2) |
| V1A-N7A      | 2.071(2) | Ni1-N7       | 1.9924(16)|
| V1A-N14A     | 2.155(2) | Ni1-N14      | 2.1021(17)|
| V1A-N27A     | 2.052(2) | Ni1-N27      | 1.9882(16)|
| V1A-N34A     | 1.989(2) | Ni1-N34      | 2.0824(16)|
| N7A-C6A      | 1.425(3) | N7-C6        | 1.375(3) |
| N7A-C8A      | 1.322(4) | N7-C8        | 1.335(3) |
| N14A-C13A    | 1.323(4) | N14-C13      | 1.323(2) |
| N14A-C15A    | 1.362(4) | N14-C15      | 1.414(3) |
| N27A-C26A    | 1.397(4) | N27-C26      | 1.363(3) |
| N27A-C28A    | 1.392(4) | N27-C28      | 1.308(3) |
| N34A-C33A    | 1.371(4) | N34-C33      | 1.310(2) |
| N34A-C35A    | 1.402(3) | N34-C35      | 1.415(2) |
[Ni(HL)\textsubscript{2}] is formed of neutral molecules, where the two organic ligands are coordinated to the metal centre as three-dentate ONN-donors in a nearly symmetric fashion. In both ligands, the shortest C–N bonds (1.308, 1.310, 1.323 and 1.335 Å) are towards the central phenylene ring of the ligand corresponding to imine double bonds (see Table 2). In general, the bonding parameters indicate that the organic ligands are in a partly deprotonated [HL\textsuperscript{ox}]\textsuperscript{−} oxidation state with each ligand having one dangling phenolic hydroxyl group which engages in intramolecular hydrogen bonding with the coordinated O-atoms. These two coordinated phenolic oxygens (O1 and O3) are aligned in a \textit{cis} form in respect to the central atom. The coordination geometry around the central atom is best described as a distorted octahedron, which matches with the proposed structure for the high-spin Ni\textsuperscript{2+} complex. The calculated MOS values are −0.92(14) and −1.15(12) for the coordinated o-aminophenol moieties. These findings support strongly the idea of two identical [L\textsuperscript{ox}2\textsuperscript{−}] ligands within the complex.

2.3. NMR and MS Spectroscopic Studies

The \textsuperscript{1}H-NMR spectra of Ti and Zr complexes display two distinct singlets (ca. 1.2 and 0.9 ppm) for the tert-butyl groups and four multiplets (ca. 6.95–7.55 ppm) for aromatic protons demonstrating the high symmetry of the molecules in the solutions. Although the ligand H\textsubscript{2}L can form stable radicals upon coordination and intramolecular redox reactions, the well-resolved NMR spectra (see Supplementary Information) verify the diamagnetic nature of the Ti and Zr complexes at room temperature in solution. The protonation states of the coordinated ligands were also studied using electrospray ionization (ESI-MS) mass spectroscopy, and the molecular ions [M + H]\textsuperscript{+} were detected at \textit{m/z} = 1073.62 for [Ti(L\textsuperscript{ox})\textsubscript{2}] and at \textit{m/z} = 1115.62 for [Zr(L\textsuperscript{ox})\textsubscript{2}], which indicates that the ligands are fully deprotonated upon coordination.

The NMR studies for the V and Ni complexes provided little information on the molecular structure in solution, indicating the paramagnetic nature of the complex. Therefore, the Evans NMR method \cite{26} for measuring the magnetic susceptibilities in solutions was applied to get \( \mu_{\text{eff}} = 0.66 \mu_B \) for the V complex. This effectively equals to less than half spins on the molecule, thus suggesting that the electronic ground state of the complex is a mixture of singlet and triplet/pentet states, which is also supported by SQUID (Superconducting Quantum Interference Device) results (see below). Although we cannot entirely rule out the possibility of paramagnetic impurity being the source of the non-zero magnetic moment in solution, such impurities are not present in the ESI-MS spectra, which gave \textit{m/z} values of 1077.6760 (ESI\textsuperscript{+}(\textsuperscript{+})) and 1077.6251 (ESI\textsuperscript{−}(\textsuperscript{−})), with the highest intensity. This would thus indicate the detection of [M + H]\textsuperscript{+} and [M + H]\textsuperscript{−} ions, while the calculated exact mass for the complex based on the X-ray studies is 1076.6323. The MS patterns were also rather atypical for a vanadium complex (see Supplementary Information), which may very well be the manifestation of the redox behavior of the complex. In addition, the positive ionization mode gives traces of [M + Na]\textsuperscript{+} and [M + K]\textsuperscript{+} ions that are detected at \textit{m/z} = 1099.6508 and 1115.6145, respectively.

[Ni(HL\textsuperscript{ox})\textsubscript{2}] proved to be paramagnetic in the NMR measurements, so the Evans method was applied to measure \( \mu_{\text{eff}} = 2.84 \mu_B \) at room temperature. The magnetic properties were further studied in the solid state using SQUID experiments (see below). The protonation state of the coordinated ligands was also verified using ESI-MS, while the molecular ion [M + H]\textsuperscript{+} was detected at \textit{m/z} = 1085.6731.

2.4. DFT Studies for the Vanadium Complex

Density functional theory (DFT) was employed to investigate the electronic structure of the [V(L)(HL)] complex. DFT calculations were carried out using PBE0 functional \cite{27-29} and def2-TZVP basis sets \cite{30} (with def2/J auxiliary basis sets \cite{31}) on a modified structure of [V(L)(HL)] with tert-butyl groups replaced with H atoms. Geometry optimizations were carried out within the Orca program (version 4.2.0) \cite{32} for singlet, broken symmetry singlet, triplet and pentet electronic states, using the modified single crystal X-ray structure as a starting point. The optimized structures were then subjected to vibrational analyses to establish that they each correspond to a minimum on the potential energy surface.
From the studied systems, a broken symmetry (BS) singlet ($S = 0$) was found to correspond to the ground state, with the triplet ($S = 1$) only 6 kJ/mol higher in energy, followed by the closed shell singlet ($S = 0; 22$ kJ/mol higher than BS singlet). The optimized structure of the pentet ($S = 2$) was found to be considerably less favorable (51 kJ/mol higher in energy). The coordination sphere bond lengths (Table S2) of the ground state broken symmetry solution fall within ±0.03 Å from the corresponding experimental values (complex A in the asymmetric unit). The triplet structure shows a slightly larger deviation, whereas, interestingly, the closed shell singlet solution shows the best fit for experimental metal-ligand bond lengths. Furthermore, the asymmetry of the V-N bonds is best reproduced by the closed shell singlet optimized geometry. The xyz coordinates of the optimized structures are given in Supplementary Materials, Table S4.

Analysis of the spin density of the broken symmetry singlet shows the localization of the opposite spins to the vanadium center and, respectively, to the tetradeutate ligand with only a minute contribution from the tridentate ligand (Figure 3). The visual information together with the analysis of the Löwdin atomic spin populations strongly point to an antiferromagnetically coupled radical ligand and vanadium(4+) center as the ground state structure. The low-lying excited triplet state shows the spin localized, to a great extent, to the vanadium ion, but also delocalized along the ligand backbones. According to the Löwdin atomic spin populations the triplet contains a vanadium(III) ion, and thus, the ligands should exist at either [L$^{sq1}$]$^{2-}$ and [HL$^{ox}$]$^{-}$ or [L$^{sq1}$]$^{3-}$ and [HL$^{sq2}$]$^{-}$ (or [HL$^{sq1}$]$^{2-}$ and [L$^{sq2}$]$^{-}$) oxidation states, respectively. From these formal oxidation states, the former has no unpaired electrons, whereas the latter oxidation states would require the assignment of half a spin to each of the ligands. As there is significant spin density on both ligands, the results indicate that the ligands of the excited triplet state structure can be assigned as [L$^{sq1}$]$^{3-}$ and [HL$^{sq2}$]$^{-}$ (or [HL$^{sq1}$]$^{2-}$ and [L$^{sq2}$]$^{-}$).

**Figure 3.** Spin density plots for broken symmetry singlet and excited triplet states of modified structure of [V(L)(HL)].

### 2.5. Electrochemical Studies

Cyclic voltammetry was used to study the redox stability and the characteristic redox behavior of [Ti(L$^{ox}$)$_2$] and [Zr(L$^{ox}$)$_2$] by scanning the potential range from $+1.3$ to $-2.3$ V vs Fc$^+/Fc$. A different range, from $+1.0$ to $-2.0$ V vs Fc$^+/Fc$, was applied for [V(L$^{sq1}$)(HL$^{ox}$)] and [Ni(HL$^{ox}$)$_2$]. The reversible electron transfer processes for Ti and Zr and Ni complexes are probably only ligand based $[5,6,8]$, as similar waves can be observed in the voltammograms of each of the compounds. Some of the redox processes are at overlapping potentials, or faint, thus rendering the voltammograms somewhat difficult to interpret. The voltammogram of the V complex, on the other hand, is clearly different.
The voltammogram exhibits two reversible one-electron processes (+0.01 and +0.54 V) and several irreversible processes in the mix. It is therefore difficult to draw definite conclusions regarding which redox event takes place at each of the redox waves of the voltammogram.

Table 3. Measured redox potentials for studied complexes (vs Fc⁺/Fc).

| Complex          | Reversible One-Electron Redox Waves (V) |
|------------------|----------------------------------------|
| [Ti(L²ο)₂]       | -1.64 -1.19 -0.17 +0.24 +1.06          |
| [Zr(L²ο)₂]       | -1.95 -1.49 -1.09 -0.02 +0.41 -0.74    |
| [V(L²ο¹)(HL²ο)]  | -1.01 +0.01 -0.54                       |
| [Ni(HL²ο)₂]      | -1.47 -1.07 -0.92 +0.04 +0.36           |

Figure 4. Cyclic voltammogram of [V(L²ο¹)(HL²ο)]. Cyclic voltammetry is performed at RT vs Fc/Fc⁺, in CH₂Cl₂ with 100 mVs⁻¹ scan rate and the rotation of the measurement is clockwise.

2.6. Optical Absorption Studies

One of our motivations for the synthesis of the presented compounds was their potential applicability in photovoltaics as dyes, and thus, their optical absorption properties were studied initially using UV-vis spectroscopy. Accordingly, the UV-vis-NIR spectra in CH₂Cl₂ present distinct absorption peaks for each complex in the vis/NIR range (Table 4, Figure 5). The absorption coefficients of the different complexes range from low to significantly high when compared with the coefficient of the standard ruthenium sensitizer dye (ε ≈ 14 × 10³ M⁻¹ cm⁻¹, λ = 538 nm) [33]. The absorption spectra are rather similar in shape in the solid state, as well in CH₂Cl₂ solution for all complexes (see Supplementary Materials). The absorption coefficients of the different complexes range from low to significantly high when compared with the coefficient of the standard ruthenium sensitizer dye (ε ≈ 14 × 10³ M⁻¹ cm⁻¹, λ = 538 nm) [33]. The absorption spectra are rather similar in shape in the solid state, as well in CH₂Cl₂ solution for all complexes (see Supplementary Materials). The absorption coefficients of the different complexes range from low to significantly high when compared with the coefficient of the standard ruthenium sensitizer dye (ε ≈ 14 × 10³ M⁻¹ cm⁻¹, λ = 538 nm) [33]. The absorption spectra are rather similar in shape in the solid state, as well in CH₂Cl₂ solution for all complexes (see Supplementary Materials). The group IV complexes have similar, ligand-based, π→π* UV absorptions as the ones that have been previously reported [4]. The strong NIR absorptions above 1000 nm are most probably ligand-based transitions, i.e., ligand-to-ligand charge transfer (LLCT) in character, as reported for the related Co complex [7]. The LLCT processes are of both the intra and inter ligand type [34]. As these metal centers are high oxidation state species with empty d-orbitals, there may be some ligand-to-metal charge transfer (LMCT) involved. [V(L²ο¹)(HL²ο)] has a broad absorption at λmax = 680 nm, which reaches the NIR range. This absorption may originate from LLCT with some LMCT or metal-to-ligand CT (MLCT) mixing. As the central metal is not d⁰, also a d-d transition is possible. In addition, the broad and intense absorption at 850 nm for [Ni(HL²ο)₂] is assignable as a LLCT band and/or to the Ni d-d absorptions.
Table 4. The UV-vis-NIR absorption peaks in CH$_2$Cl$_2$ solution and their molar absorption coefficients.

| Complex                | $\lambda_{\text{max}}$ (nm) | $\varepsilon$ ($10^3 \times$ L mol$^{-1}$ cm$^{-1}$) |
|------------------------|-------------------------------|------------------------------------------------------|
| [Ti(L$^{\text{ox}}$)$_2$] | 390                           | 4.3                                                  |
|                        | 570 (sh)                      | 2.6                                                  |
|                        | 710                           | 5.3                                                  |
|                        | 1050                          | 2.2                                                  |
| [Zr(L$^{\text{ox}}$)$_2$] | 285                           | 19                                                   |
|                        | 370                           | 14                                                   |
|                        | 760                           | 13                                                   |
|                        | 1020                          | 8.5                                                  |
| [V(L$^{\text{sq1}}$)(HL$^{\text{ox}}$)] | 315                           | 16                                                   |
|                        | 430                           | 7.8                                                  |
|                        | 540 (sh)                      | 8.1                                                  |
|                        | 680                           | 13                                                   |
| [Ni(HL$^{\text{ox}}$)$_2$] | 460                           | 14                                                   |
|                        | 850                           | 23                                                   |

2.7. EPR Studies for the Vanadium and Nickel Complexes

EPR (Electron Paramagnetic Resonance) studies were conducted to examine the electronic nature of the V and Ni complexes more closely. The Ni complex, having an integer spin $S = \frac{3}{2}$, measured in its powder form, was found to be EPR silent. The spectrum of [V(L$^{\text{sq1}}$)(HL$^{\text{ox}}$)] is described by Hamiltonian for electron spin $S$:

$$H = \beta B g S + S A I,$$

(1)

where $\beta$ is Bohr magneton, $B$ is a magnetic field, $g$ and $A$ are the tensors of axial symmetry of $g$-factor and hyperfine interaction between electron and $^{51}$V (spin $I = \frac{7}{2}$) nuclei, respectively. Two sets of partly resolved lines in spectrum shown in Figure 6 correspond to hyperfine interaction between electron and $^{51}$V nuclei for different values of the parallel and perpendicular components of $A$-tensor: $A_{||} = 42$ G (or 117 MHz, or $39 \times 10^{-4}$ cm$^{-1}$) and $A_{\perp} = 16$ G (or 45 MHz, or $15 \times 10^{-4}$ cm$^{-1}$). The values of $g$-tensor components are $g_{||} = 2.001$ and $g_{\perp} = 1.999$. The estimated hyperfine structure splitting is about two to three times lower compared to previously reported V$^{4+}$ ESR spectra [35–39]. This shows that the localization of the spin density on the $^{51}$V nuclei is lower in the investigated samples. The $g$-factor components are also higher with respect to reported earlier data, which further supports the idea of a delocalized metal-organic radical system.
we propose that the measured magnetic moments, both in a solution and in a solid state, suggest that the Ti and Zr complexes are not able to transform the absorbed NIR photons to free electrons. The measurements suggest that the magnetic moment, and therefore the number of unpaired spins, is decreased within the molecule when the complex \([V(L_{sq1})(HLox)]\) is illuminated. The \(g\) factor of the magnetic susceptibility plot is then due to a low-lying triplet state which is thermally populated, as suggested by the DFT calculations. The field dependence curve of \([V(L^{sq1})(HLox)]\) is typical for a paramagnetic complex and no hysteresis is observed (see Supplementary Materials). By observing the magnetic properties with or without irradiation in the NIR region (a laser diode operating at \(\lambda = 785 \text{ nm}\)), it was confirmed that the Ti and Zr complexes are not able to transform the absorbed NIR photons to paired electrons.

Figure 6. The EPR spectrum of powder \([V(L^{sq1})(HLox)]\) sample.

2.8. Magnetic Properties

The magnetic nature and thermal dependency of the complexes’ magnetic properties were investigated using a SQUID magnetometer in the temperature range of 2–300 K. The Ti and Zr complexes exhibit diamagnetic properties in the SQUID measurements, as expected. The possibility of utilizing synthesized complexes in photovoltaic applications was ascertained by observing the magnetic properties with or without irradiation in the NIR region (a laser diode operating at \(\lambda = 785 \text{ nm}\)), and it was confirmed that the Ti and Zr complexes are not able to transform the absorbed NIR photons to free electrons. The measurements suggest that the magnetic moment, and therefore the number of unpaired spins, remains unchanged within the molecule when the complexes \([TiL^{ox})_2\] and \([ZrL^{ox})_2\] are illuminated.

\([V(L^{sq1})(HLox)]\) gave a paramagnetic signal in SQUID measurements. The temperature dependence of the magnetic susceptibility was also measured, and the data were fitted to the Curie law of localized moments. The \(1/\chi_{\text{mol}}\) curve does not follow the Curie law, which indicates that there is some electron spin density delocalized on the ligand rather than on the metal center. The measured effective magnetic moment is low (Figure 7) and similar in magnitude to the \(\mu_{\text{eff}}\) measured in solution using the Evans method. Although we cannot unambiguously show that no magnetic impurities are present, we propose that the measured magnetic moments, both in a solution and in a solid state, suggest that \([V(L^{sq1})(HLox)]\) consists prevalently of an antiferromagnetically coupled radical ligand and \(V^{4+}\) metal ion with one unpaired electron. The paramagnetic signal, which is shown in the molar magnetic susceptibility plot, is then due to a low-lying triplet state which is thermally populated, as suggested by the DFT calculations. The field dependence curve of \([V(L^{sq1})(HLox)]\) is typical for a paramagnetic complex and no hysteresis is observed (see Supplementary Materials). By observing the magnetic properties with or without irradiation in the NIR region (a laser diode operating at \(\lambda = 785 \text{ nm}\)), it was possible to confirm that the complex is able to transform the absorbed NIR photons to paired electrons. The measurements (Figure 8) suggest that the magnetic moment, and therefore the number of unpaired spins, is decreased within the molecule when the complex \([V(L^{sq1})(HLox)]\) is illuminated. The effect was not permanent at any temperature, suggesting that the complex could be used to continuously convert IR quanta to paired electrons.
illuminated. The effect was not permanent at any temperature, suggesting that the complex could be used to continuously convert IR quanta to paired electrons.

**Figure 7.** The measured effective magnetic moments for [V(Lsq1)(HLox)] (bottom) and [Ni(HLox)2] (top) in the function of temperature. The signal observed in the right figure at ca. 70 K is an instrumental artefact.

**Figure 8.** The measured virgin curves of [V(Lsq1)(HLox)] on a SrTiO₃ disk. The red curves result from the measurements done under irradiation in the NIR region, and the black curves are from the measurements done without irradiation.

The [Ni(HLox)₂] complex, as expected, also gave a paramagnetic signal. The temperature dependence of the susceptibilities was also measured, and the data were fitted to the Curie law of localized moments. The 1/χmol plot followed the Curie law, and the effective magnetic moment for the molecule was calculated from the slope of the plot. The result was µeff = 2.88 µB, which is in good agreement with an octahedral high-spin Ni²⁺ moiety. The field dependence curve of [Ni(HLox)₂] is typical for a paramagnetic complex. The Ni complex was found to react under NIR radiation, but slightly differently than the V complex. The Ni complex converted photons to free electrons, thus increasing its magnetic moment (Figure 9). As in the studies with the V complex, the effect was not permanent at any studied temperature.
In conclusion, a series of transition metal complexes was made using a redox non-innocent \(N,N'\text{-bis}(3,5\text{-di-tert-butyl}-2\text{-hydroxy-phenyl})\text{-1,2-phenylenediamine} \text{H}_4L\) as a ligand precursor. \text{H}_4L\ can form complexes with a 2:1 ligand-to-metal ratio with different transition metal precursors. In these complexes, two organic molecules coordinate to the metal centre as tri- or tetradentate ligands and display a range of different oxidation states.

Several group IV metal complexes with L ligand were previously prepared by the reaction of metal tetrachlorides with L to yield molecular complexes \([\text{M}^{2+}(\text{L})_{n}]\) \((\text{M} = \text{Ti, Zr, Hf}; \text{S} = \text{pyridine, THF}; \ n = 2,3)\), which can be further oxidised by PhICl\(_2\) to yield corresponding chlorides \([\text{MCl}_2(\text{L}^{n=3})_{n}]\) \([3,4]\). Furthermore, the reaction of \text{H}_4L\ with \([\text{M}(\text{O-i-Pr})_{4}]\) \((\text{M} = \text{Ti, Zr})\) leads to the formation of either \([\text{Ti}(\text{LOx})(\text{O-i-Pr})_2]_2\) or \([\text{Ti}(\text{O-i-Pr})_2]\text{Cl}_2\), depending on the stoichiometry \([2]\). In the present study we show that under ambient conditions, where the organic proligand is abundantly available, rather than being added dropwise to metal precursor solution, 2:1 ligand-to-metal complex is formed.

The essence of the electronic structure of the V complex remained somewhat unclear after many different analyses. However, preliminary DFT calculations, carried out on a model complex with tert-butyl substituents replaced by H atoms, predict a broken symmetry singlet ground state for \([\text{V}(\text{L})(\text{HL})]\). Analysis of spin density shows a localization of opposite spins on the V center and the tetradentate ligand, respectively, with negligible contribution from the tridentate ligand. This corresponds to the antiferromagnetically coupled \([\text{V}(\text{L}^{\text{sq}})(\text{HL}^{\text{ox}})]\) structure with V\(^{4+}\), suggested also on the basis of metrical oxidation states. However, the broken symmetry singlet ground state and the lowest triplet state are separated only by ca. 6 kJ/mol, with a closed shell singlet a further 16 kJ/mol higher in energy. The existence of several electronic states close in energy suggests that multiconfigurational methods would be required to depict the electronic structure of \([\text{V}(\text{L}^{\text{sq}})(\text{HL}^{\text{ox}})]\) accurately. From the DFT results, it can be deduced, however, that the experimentally observed magnetism is likely due to mixing of several low-lying singlet and triplet electronic states.

All compounds used in this study have strong NIR absorptions and some show very interesting magnetic and electrochemical properties. While strong NIR absorbers, the Ti and Zr complexes regrettfully did not produce any magnetic behavior when illuminated. This was probably due to the diamagnetic nature of the complexes. Nevertheless, the Ti and Zr complexes still pose an interesting comparison with the V and Ni complexes that generate (un)paired electrons under NIR radiation. This indicates great promise in the photovoltaic applications and furthers our knowledge regarding NIR-absorbing complexes, their magnetic behavior and their possible photovoltaic applications. Further investigation of these compounds is currently ongoing in our group.
3. Materials and Methods

The starting complex [Ni(acac)₂·2H₂O] was synthesized by the well-known reaction of Ni(NO₃)₂·6H₂O with the excess of 2,4-pentanedione in an aqueous solution using sodium acetate as a base. The product was filtered and washed with water. The ligand precursor was synthesized using literature procedures [4,8]. Other chemicals were used as purchased from commercial sources. The solvents used were of HPLC grade. All syntheses were done under an ambient atmosphere.

The ¹H and ¹³C-NMR spectra were recorded with 500 MHz Bruker AVANCE-III NMR system. ESI-MS spectra for complexes were measured in the positive-ion mode with a Bruker microOTOF-Q spectrometer. The samples were injected as dichloromethane-acetonitrile solutions. Cyclic voltammetry (CV) for complexes were recorded at ambient temperature using a platinum working electrode, a 1 mm diameter platinum counter electrode, and a Ag/AgCl reference electrode. Samples were dissolved in CH₂Cl₂ containing 0.1 M (Bu₄N)ClO₄ as the supporting electrolyte. The voltammograms were recorded at a scan rate of 100 mV/s, while the potentials were measured in volts versus the Fc⁺/Fc couple. The thermal changes of the complex were studied with TA Instrument SDT Q600 simultaneous TGA-DSC apparatus between 23 and 500 °C in flowing nitrogen gas using an aluminum oxide pan as sample holder and reference. A flow rate of 100 mL/min and a heating rate of 5 °C/min were applied. Amounts of [Ti(L³)₂], [Zr(L³)₂], [V(L⁴)(HL)⁵⁺] and [Ni(HL)⁶⁺] were 7.94, 10.89, 4.77 and 4.48 mg, respectively. UV-Vis spectra in CH₂Cl₂ solution were measured with an Agilent Cary 60 UV/vis spectrophotometer in a 10 mm quartz glass cuvette. UV-Vis-NIR spectra were measured of KBr pellets with a Varian Cary 50 spectrophotometer, and a blank KBr pellet was used as reference. The solid-state powder of the [V(L⁴)(HL)⁵⁺] sample ESR spectrum was recorded at 18 K using a ESR spectrometer operating at the microwave frequency of 9.030 GHz and 100 kHz modulation of magnetic field. A look-in amplifier was used for observing the first derivative shape of ESR absorption lines under magnetic field scan. The temperature of samples was controlled using a He-gas flow cryostat. The magnetic properties were measured in a SQUID magnetometer. The temperature dependence of the zero-field-cooled (ZFC) and field-cooled (FC) magnetization was measured between temperatures of 2 and 300 K with a Quantum Design SQUID magnetometer MPMS XL with external magnetic field of B = 1.0 T on 7.01, 7.67, 35.6 and 10.7 mg samples of [Ti(L³)₂], [Zr(L³)₂], [V(L⁴)(HL)⁵⁺] and [Ni(HL)⁶⁺], respectively, sealed in plastic nonmagnetic straws. The field dependence was measured at 2 K between ~2.5 and 2.5 T using 50 mT steps. Virgin magnetizations as a function of B and magnetic hysteresis curves were recorded in magnetic fields up to 5 T at temperatures of 2, 30, 50 and 100 K. Although the magnetic particles were randomly oriented on the surface of the substrate, the external magnetic field B was always oriented along the out-of-plane axis of the substrate. The photoinduced magnetization measurements were performed in dark or under illumination through an optical fiber with a home-made fiberoptic sample holder attached to the SQUID magnetometer. The samples of [Ti(L³)₂], [Zr(L³)₂], [V(L⁴)(HL)⁵⁺] and [Ni(HL)⁶⁺] were drop casted as diethyl ether solutions on circular SrTiO₃ plates (diameter 5 mm). The light source was a Fabry-Perot laser diode by Thorlabs operating at λ = 785 nm (1.58 eV) with a maximum output power of 10 mW measured at the end of the optical fiber, and hence, the laser fluence on the sample surface was ca. 0.5 mW/mm². The single crystal X-ray diffraction (XRD) analyses were carried out using an Agilent SuperNova microfocus dual source (Cu/Mo) diffractometer equipped with an Atlas detector. The data collection and reduction, including multifaceted crystal model-based analytical absorption correction, were carried out using the CrysAlisPro program [40]. The structures were solved with ShelXS [41] using direct methods and refined on F² using full matrix least squares techniques with ShelXL [42] within the Olex² (v. 1.2) [43] program package. All non-hydrogen atoms were refined anisotropically. The hydrogen atoms were refined using a riding model with fixed thermal parameters 1.2–1.5 times the values of the corresponding host atoms (the O–H distances of the O₂ and O₄ hydroxyl groups were refined freely).
3.1. Preparation of \([\text{Ti}(L^{\text{ox}})_2]\)

A total of 44.2 mg (0.276 mmol) of TiOSO\(_4\)-H\(_2\)O with the ligand H\(_2\)L (156 mg, 0.300 mmol) were dissolved in methanol (30 mL), and a base, Et\(_3\)N (0.170 mL), was added. The reaction mixture turned slowly to turn dark green at room temperature and it was left to stand for 2 days. The reaction mixture was evaporated to dryness, washed with hexane and then filtered. The filtrate was evaporated and filtered through a silica plug using toluene as an eluent. The toluene filtrate was evaporated to dryness and the amount of dry product was 43.1 mg (yield 27%). \([\text{Ti}(L^{\text{ox}})_2]\) dissolves moderately in alcohols and acetonitrile, whereas it is readily soluble in diethyl ether, halogenated organic solvents and hydrocarbon solvents. X-ray diffraction quality crystals of \([\text{Ti}(L^{\text{ox}})_2]\)CH\(_3\)CN were obtained from acetonitrile solution with slow evaporation. IR: 2951 (s), 2904 (s), 2867 (s), 1582 (w), 1446 (s), 1379 (m), 1359 (m), 1322 (s), 1290 (w), 1256 (vs), 1193 (w), 1167 (vs), 1152 (s), 1131 (s), 1113 (vs), 1073 (s), 1023 (m), 998 (m), 948 (m), 885 (m), 794 (w), 665 (m), 647 (m), 602 (w), 586 (w), 545 (m), 506 (m), 463 (m), cm\(^{-1}\). 1H-NMR (CDCl\(_3\)): \(\delta\) 7.59 (dd, \(J_{1HH} = 3.30\) and 7.39 Hz, 4H, Ph-H), 6.97 (dd \(J_{1HH} = 3.33\) and 7.33 Hz, 4H, Ph-H), 5.82 (m), 565 (m), 538 (s), 504 (s) ppm. 13C-NMR (CDCl\(_3\)): \(\delta\) 166.91, 150.95, 140.23, 137.10, 130.21, 126.10, 124.25 (Ph-C) 34.41, 34.24 (C(CH\(_3\)_3)), 31.37, 29.15 (C(CH\(_3\)_3)), ppm. ESI(+)-MS: \(m/z\) 1073.62 ([M + H]\(^+\) calcd. \(m/z\) 1073.64). m.p. 270 °C (decomp., measured for \([\text{Ti}(L^{\text{ox}})_2]\)CH\(_3\)CN).

3.2. Preparation of \([\text{Zr}(L^{\text{ox}})_2]\)

Methanol solution (20 mL) of ZrOCl\(_2\)-8 H\(_2\)O (104 mg, 0.323 mmol) was introduced to the solid ligand H\(_2\)L (144 mg, 0.279 mmol) and Et\(_3\)N (0.160 mL). The reaction mixture gradually changed its color from neutral to very intense blue. The closed reaction vessel was left to stand overnight at room temperature to obtain large, X-ray diffraction quality single crystals of \([\text{Zr}(L^{\text{ox}})_2]\). The crystals were isolated by filtration and washed with methanol to obtain 69.6 mg (44%) of solid product. \([\text{Zr}(L^{\text{ox}})_2]\) is insoluble in hydrocarbon solvents. X-ray diffraction crystals of \([\text{Zr}(L^{\text{ox}})_2]\)CH\(_3\)CN were obtained from methanol solution with slow evaporation. IR: 2953 (s), 2905 (s), 2868 (s), 1531 (s), 1461 (s), 1439 (s), 1391 (m), 1265 (vs), 1193 (w), 1167 (vs), 1153 (s), 1131 (w), 1099 (w), 1077 (m), 1023 (m), 998 (m), 948 (m), 885 (m), 794 (w), 665 (m), 647 (m), 602 (w), 586 (w), 545 (m), 506 (m), 463 (m), cm\(^{-1}\). 1H-NMR (CDCl\(_3\)): \(\delta\) 7.57 (dd, \(J_{1HH} = 3.18\) and 7.45 Hz, 4H, Ph-H), 7.28 (d, \(J_{1HH} = 1.89\) Hz, 4H, Ph-H), 7.00 (s, 4H, Ph-H), 6.97 (dd \(J_{1HH} = 3.13\) and 7.33 Hz, 4H, Ph-H), 5.82 (m), 565 (m), 538 (w), 504 (s) ppm. 13C-NMR (CDCl\(_3\)): \(\delta\) 166.11, 152.02, 140.95, 137.10, 130.21, 126.10, 124.25 (Ph-C) 34.41, 34.24 (C(CH\(_3\)_3)), 31.37, 29.15 (C(CH\(_3\)_3)), ppm. ESI(+)-MS: \(m/z\) 1115.62 ([M + H]\(^+\) calcd. \(m/z\) 1115.59). ESI(+)-MS: \(m/z\) 1153.57 ([M + K]\(^+\) calcd. \(m/z\) 1153.55). m.p. 340 °C (decomp.).

3.3. Preparation of \([\text{V}(L^{\text{III}})(HL^{\text{ox}})]\)

A total of 152 mg (0.294 mmol) of H\(_2\)L was added into a methanol solution (8 mL) which contained 79.4 mg (0.314 mmol) of [VO\(_2\)(H\(_2\)O)] and 50 μL of NEt\(_3\). The reaction mixture turned dark purple quite swiftly and the reaction was left to run in a closed vessel for 3 days at room temperature. The reaction mixture was evaporated to dryness and the product was separated by silica column chromatography using diethyl ether as eluent. The yield was 106 mg (67%). The isolated solid dissolves in alcohols and acetonitrile to some extent, while halogenated organic solvents, diethyl ether and hydrocarbon solvents dissolve the complex with ease. X-ray diffraction quality single crystals were obtained directly from the MeOH reaction mixture. IR: 2951 (s), 2904 (s), 2867 (s), 1582 (w), 1446 (s), 1415 (m), 1391 (m), 1360 (s), 1334 (m), 1249 (vs), 1196 (s), 1160 (s), 1136 (s), 1112 (s), 1025 (m), 989 (m), 911 (m), 862 (w), 821 (w), 770 (w), 740 (s), 665 (m), 647 (m), 602 (w), 586 (w), 545 (m), 506 (m), 463 (m), cm\(^{-1}\). 1H-NMR (CDCl\(_3\)): \(\delta\) 1.53, 1.50, 1.29, 1.01 ppm (4s, 72H, C(CH\(_3\)_3)). ESI(+)-MS: \(m/z\) 1077.65 ([M + Na]\(^+\) calcd. \(m/z\) 1076.63). ESI(+)-MS: \(m/z\) 1099.65 ([M + Na]\(^+\) calcd. \(m/z\) 1099.62). ESI(+)-MS: \(m/z\) 1115.61 ([M + Na]\(^+\) calcd. \(m/z\) 1115.59).
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References
1. Jørgensen, C.K. Differences between the four halide ligands, and discussion remarks on trigonal-bipyramidal complexes, on oxidation states, and on diagonal elements of one-electron energy. Coord. Chem. Rev. 1966, [CrossRef]
2. Zelikoff, A.L.; Kopilov, J.; Goldberg, I.; Coates, G.W.; Kol, M. New facets of an old ligand: Titanium and zirconium complexes of phenylenediamine bis(phenolate) in lactide polymerisation catalysis. Chem. Commun. 2009, 6804–6806. [CrossRef] [PubMed]
3. Blackmore, K.J.; Lal, N.; Ziller, J.W.; Heyduk, A.F. Catalytic reactivity of a zirconium(IV) redox-active ligand complex with 1,2-diphenylhydrazine. J. Am. Chem. Soc. 2008, 130, 2728–2729. [CrossRef] [PubMed]
4. Blackmore, K.J.; Lal, N.; Ziller, J.W.; Heyduk, A.F. Group IV coordination chemistry of a tetratetradate redox-active ligand in two oxidation states. Eur. J. Inorg. Chem. 2009, 735–743. [CrossRef]
5. Hänninen, M.M.; Paturi, P.; Tuononen, H.M.; Sillanpää, R.; Lehtonen, A. Heptacoordinated molybdenum(VI) complexes of phenylenediamine Bis(phenolate): A stable molybdenum amidophenoxide radical. Inorg. Chem. 2013, 52, 5714–5721. [CrossRef]
6. Hossain, M.K.; Haukka, M.; Hänninen, M.M.; Lisensky, G.C.; Paturi, P.; Nordlander, E.; Lehtonen, A. An experimental and theoretical study of a heptacoordinated tungsten(VI) complex of a noninnocent phenylenediamine bis(phenolate) ligand. Inorg. Chem. Commun. 2018, 93, 149–152. [CrossRef]
7. Lesh, F.D.; Lord, R.L.; Heeg, M.J.; Schlegel, H.B.; Verani, C.N. Unexpected formation of a cobalt(III) phenoxazinylate electron reservoir. Eur. J. Inorg. Chem. 2012, 463–466. [CrossRef]
8. Chaudhuri, P.; Hess, M.; Müller, J.; Hildenbrand, K.; Bill, E.; Weyermüller, T.; Wieghardt, K. Aerobic Oxidation of Primary Alcohols (Including Methanol) by Copper(II)− and Zinc(II)−Phenoxyl Radical Catalysts. J. Am. Chem. Soc. 1999, 121, 9599–9610. [CrossRef]
9. Amb, C.M.; Heth, C.L.; Evenson, S.J.; Pokhodnya, K.I.; Rasmussen, S.C. Thiophene-Fused Nickel Dithiolenes: A Synthetic Scaffold for Highly Delocalized π-Electron Systems. *Inorg. Chem.* 2016, 55, 10978–10989. [CrossRef]

10. Aragoni, M.C.; Arca, M.; Devillanova, F.A.; Isaia, F.; Lippolis, V.; Mancini, A.; Pala, L.; Verani, G.; Agostinelli, T.; Caironi, M.; et al. First example of a near-IR photodetector based on neutral [M(R-dmet)2] bis(1,2-dithiolene) metal complexes. *Inorg. Chem. Commun.* 2007, 10, 191–194. [CrossRef]

11. Mueller-Westerhoff, U.T.; Vance, B.; Ihl Yoon, D. The synthesis of diolene dyes with strong near-IR absorption. *Tetrahedron* 1991, 47, 909–932. [CrossRef]

12. Bai, J.F.; Zuo, J.L.; Tan, W.L.; Ji, W.; Shen, Z.; Fun, H.K.; Chinnakali, K.; Razak, I.A.; You, X.Z.; Che, C.M. Synthesis, structure and optical limiting effect of two new nickel complexes containing strongly bound geometrically fixed multisulfur 1,2-dithiolene ligands showing remarkable near-IR absorption. *J. Mater. Chem.* 1999, 9, 2419–2423. [CrossRef]

13. Cho, J.-Y.; Fu, J.; Padiha, L.A.; Barlow, S.; Van Stryland, E.W.; Hagan, D.J.; Bishop, M.; Marder, S.R. Synthesis of a Nickel Bis(dithiolene) Complex with Strong Near-Infrared Two-Photon Absorption. *Mol. Cryst. Liq. Cryst.* 2008, 485, 915–927. [CrossRef]

14. Garreau-de Bonneval, B.; Moineau-Chane Ching, K.I.; Alary, F.; Bui, T.T.; Valade, L. Neutral d8 metal bis-dithiolene complexes: Synthesis, electronic properties and applications. *Coord. Chem. Rev.* 2010, 254, 1457–1467. [CrossRef]

15. Marshall, K.L.; Painter, G.; Lotito, K.; Noto, A.G.; Chang, P. Transition Metal Dithiolene Near-IR Dyes and Their Applications in Liquid Crystal Devices. *Mol. Cryst. Liq. Cryst.* 2006, 454, 47–449. [CrossRef]

16. Liu, Y.; Zhang, Z.; Chen, X.; Xu, S.; Cao, S. Near-infrared absorbing dyes at 1064 nm: Soluble dithiolene nickel complexes with alkylated electron-donating groups as Peripheral substituents. *Dye. Pigment.* 2016, 128, 179–189. [CrossRef]

17. Miao, Q.; Gao, J.; Wang, Z.; Yu, H.; Luo, Y.; Ma, T. Syntheses and characterization of several nickel bis(dithiolene) complexes with strong and broad Near-IR absorption. *Inorganica Chim. Acta* 2011, 376, 619–627. [CrossRef]

18. Aiken, D.C.; Ramsey, S.; Mayo, T.; Bellemare, J.; Lambrakos, S.G.; Peak, J. Inverse analysis of near infrared transmission spectra for triarylamine, tetraaryldiamine, nickel dithiolene and indium-iodide dyes. *J. Near Infrared Spectrosc.* 2015, 23, 123–132. [CrossRef]

19. Thorley, K.J.; Hales, J.M.; Anderson, H.L.; Perry, J.W. Porphyrin dimer carbocations with strong near infrared absorption and third-order optical nonlinearity. *Angew. Chemie - Int. Ed.* 2008, 47, 7095–7098. [CrossRef]

20. Demir, S.; Jeon, I.R.; Long, J.R.; Harris, T.D. Radical ligand-containing single-molecule magnets. *Coord. Chem. Rev.* 2015, 289–290, 149–176. [CrossRef]

21. Atanasov, M.; Aravena, D.; Suturina, E.; Bill, E.; Maganas, D.; Neese, F. First principles approach to the electronic structure, magnetic anisotropy and spin relaxation in mononuclear 3d-transition metal single molecule magnets. *Coord. Chem. Rev.* 2015, 289–290, 177–214. [CrossRef]

22. Gómez-Coca, S.; Aravena, D.; Morales, R.; Ruiz, E. Large magnetic anisotropy in mononuclear metal complexes. *Coord. Chem. Rev.* 2015, 289–290, 379–392. [CrossRef]

23. Liu, K.; Shi, W.; Cheng, P. Toward heterometallic single-molecule magnets: Synthetic strategy, structures and properties of 3d-4f discrete complexes. *Coord. Chem. Rev.* 2015, 289–290, 74–122. [CrossRef]

24. Fabian, J.; Nakazumi, H.; Matsuoka, M. Near-Infrared Absorbing Dyes. *Chem. Rev.* 1992. [CrossRef]

25. Brown, S.N. Metrical oxidation states of 2-amidophenoxy and catecholate ligands: Structural signatures of metal-Ligand π bonding in potentially noninnocent ligands. *Inorg. Chem.* 2012, 51, 1251–1260. [CrossRef]

26. Evans, D.F. Paramagnetic Susceptibility, etc. 2003 400. The Determination of the Paramagnetic Susceptibility. *J. Chem. Soc.* 1959, 2003–2005. [CrossRef]

27. Perdew, J.P.; Ernzerhof, M.; Burke, K. Rationale for mixing exact exchange with density functional approximations. *J. Chem. Phys.* 1996, 105, 9982–9985. [CrossRef]

28. Adamo, C.; Barone, V. Toward reliable density functional methods without adjustable parameters: The PBE0 model. *J. Chem. Phys.* 1999, 110, 6158–6170. [CrossRef]

29. Ernzerhof, M.; Scuseria, G.E. Assessment of the Perdew-Burke-Ernzerhof exchange-correlation functional. *J. Chem. Phys.* 1999, 110, 5029–5036. [CrossRef]
30. Weigend, F.; Ahlrichs, R. Balanced basis sets of split valence, triple zeta valence and quadruple zeta valence quality for H to Rn: Design and assessment of accuracy. *Phys. Chem. Chem. Phys.* 2005, 7, 3297–3305. [CrossRef]

31. Weigend, F. Accurate Coulomb-fitting basis sets for H to Rn. *Phys. Chem. Chem. Phys.* 2006, 8, 1057–1065. [CrossRef] [PubMed]

32. Neese, F. The ORCA program system. WIREs Comput. Mol. Sci. 2012, 2, 73–78. [CrossRef]

33. Nazeeruddin, M.K.; Zakeeruddin, S.M.; Humphry-Baker, R.; Jirousek, M.; Liska, P.; Vlachopoulos, N.; Shklover, V.; Fischer, C.-H.; Grätzel, M. Acid–Base Equilibria of (2,2’-Bipyridyl-4,4’-dicarboxylic acid)ruthenium(II) Complexes and the Effect of Protonation on Charge-Transfer Sensitization of Nanocrystalline Titania. *Inorg. Chem.* 1999, 38, 6298–6305. [CrossRef] [PubMed]

34. Vogler, A.; Kunkely, H. Ligand-to-ligand and intraligand charge transfer and their relation to charge transfer interactions in organic zwitterioners. *Coord. Chem. Rev.* 2007, 251, 577–583. [CrossRef]

35. Paul, A.; Assabghy, F. Optical and esr spectra of vanadium (IV) in different simple germanate, phosphate and borate glasses. J. Mater. Sci. 1975, 10, 613–620. [CrossRef]

36. Gallay, R.; Van Der Klink, J.J.; Moser, J. EPR study of vanadium (4+) in the anatase and rutile phases of TiO2. *Phys. Rev. B* 1986, 34, 3060–3068. [CrossRef]

37. Smith, T.S.; LoBrutto, R.; Pecoraro, V.L. Paramagnetic spectroscopy of vanadyl complexes and its applications to biological systems. *Coord. Chem. Rev.* 2002, 228, 1–18. [CrossRef] [PubMed]

38. Gilinskaya, L.G. EPR spectra of V(IV) complexes and the structure of oil porphyrins. *J. Struct. Chem.* 2008, 49, 245–254. [CrossRef]

39. Magon, C.J.; Lima, J.F.; Donoso, J.P.; Lavayen, V.; Benavente, E.; Navas, D.; Gonzalez, G. Deconvolution of the EPR spectra of vanadium oxide nanotubes. *J. Magn. Reson.* 2012, 222, 26–33. [CrossRef]

40. CrysAlisPro program, version 1.171.38.43 2015. Rigaku Oxford Diffraction. 2018. Available online: https://www.rigaku.com/products/sm/crystalix (accessed on 28 April 2020).

41. Sheldrick, G.M. A short history of SHELX. *Acta Crystallogr. Sect. A Found. Crystallogr.* 2008, 64, 112–122. [CrossRef]

42. Sheldrick, G.M. Crystal structure refinement with SHELXL. *Acta Crystallogr. Sect. C Struct. Chem.* 2015, 71, 3–8. [CrossRef] [PubMed]

43. Dolomanov, O.V.; Bourhis, L.J.; Gildea, R.J.; Howard, J.A.K.; Puschmann, H. OLEX2: A complete structure solution, refinement and analysis program. *J. Appl. Crystallogr.* 2009, 42, 339–341. [CrossRef]

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