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Titanium-based tetrakis-2,3-[5,6di(substituted)pyrazino]porphyrazine: synthesis and characterization

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Abstract: Tetrapyrazinoporphyrazine (TPysPz) ligands and metal complexes find, generally, application as electronic materials and catalysts. Considering the limited application of Titanium (Ti), we prepared and characterized a family of ligands and Ti-based complexes of tetrakis-2,3-[5,6-di-R₈-pyrazino]porphyrazine (R= H, 2-Py, Ph). UV-Vis measurements in different solvents confirmed molecular aggregation, which resulted more pronounced in the presence of 2-pyridil and phenyl substituents on the macrocycle edge. Because of low solubility, solid state NMR was applied for structure characterization. Additional IR, MALDI-TOF and SEM analyses were carried out to complete the characterization. Cyclic voltammetry in DMSO/Bu₄NBF₄ 0.1 M unveiled that our Ti-complexes can take part in up to five redox events. The first two involved quasi-reversible Ti(IV) reduction followed by two or three reduction at the expense of the TPysPz macrocycle. To test the applicability of our compounds as catalytic materials, we performed a preliminary cyclic voltammetry investigation in the solid-state, which showed typical peaks of hydrogen redox reactions.

Introduction

Azaanalogues of phthalocyanines, tetrapyrazinoporphyrazines (TPysPzs) ligands and metal complexes have seen a rapid development in the last three decades. Ranging from electronic materials to photodynamic therapy and catalysis, a plethora of applications has been documented in the literature for these compounds.^[1-2] Interestingly, the introduction of different substituents in the periphery of the macrocycle is able to tune its

properties, activity and solubility in organic or aqueous media. As reported in the literature, several metals (Mg, Zn, Cu, Ni, Fe, Co, Si) can be coordinated by TPysPzs. Among them, Titanium (Ti) has been seldom employed in the template synthesis of tetrapyrazinoporphrazine metal complexes,^[3] despite its abundance in the Earth's crust and its similarity in ionic radius and coordination chemistry with iron (Fe^{III}).^[4-6] In general, in oxidizing atmosphere, titanium exists as Ti(IV) and it can be coordinated by tetrapyrrole ligands as a titanyl unit (TiO²⁺).^[7-10]

In this paper, we present the synthesis and the characterization of a series of ligands and Ti-based complexes of tetrakis-2,3-[5,6-di-R₈-pyrazino]porphyrazine (R= H, 2-Py, Ph). For all the compounds (**2a-c** and **3a-c**, Scheme 1) synthesized a full characterization from the spectroscopic and electrochemical point of view is presented. Aggregation behavior was highlighted by UV-Vis spectroscopy, while cyclic voltammetry was employed to unveil the presence of differences in the redox behavior between Ti-based complexes **3a-c**, the corresponding ligands **2a-c** and similar metal complexes of TPysPzs previously described in literature.

The interest towards compounds **3a-c** arises from the potentiality of applying these molecules as catalysts in oxygen reduction reaction (ORR),^[11-12] as dyes in dye sensitized solar cells (DSSC),^[13-15] and in the detection of hydrogen peroxide.^[16] Indeed, the increasing energy demand connected with the replacement of fossil fuels and environmental concerns has made necessary the development of advanced and next-generation technologies for energy production. Among the approaches pursued to enhance these processes, the use of catalysts revealed to be the most effective. Although porphyrin, corrole and phtalocyanine-based

ligands can be a valid alternative,^[11-12] their synthesis is frequently characterized by low yields, a feature which can hamper their production on industrial scale. While the proof of concept has been demonstrated, the potentiality of tetrapyrazinoporphyrazines derivatives has still to be explored.

Results and Discussion

Preparative Aspects.

As reported in the Experimental section and in the Supporting Information, ligands **2a-c** and complexes **3a-c** can be prepared from the common starting-materials **1a-c** (Scheme 1). The synthesis of **1a-c** was accomplished by modifying the procedure reported by Mørkved employing diaminomaleonitrile and an α diketone as starting-material in glacial acetic as solvent (**1a**: 76% yield; **1b**: 73% yield; **1c**: 85% yield).^[17] The solids obtained were purified by extraction and washing with water, brine and saturated solutions of NaHCO₃, and crystallized from water/ethanol prior to use. Pyrazinoporphyrazine macrocycles 2a-c were prepared according to the procedure reported by Ercolani, Kadish et al. by autocyclotetramerization of 1a-c at high temperatures (from 175°C to 240°C according to 1a-c melting-point) in the presence of DBU as catalyst (2a: 21% yield; 2b: 20% yield; 2c: 18% yield).^[18-19] While the ligand synthesis could be realized without paying specific attention to the water content, the synthesis of the complex needed the application of Schlenk technique and stringent dry reaction conditions in the presence of an excess of 1a-c in order to avoid the formation of titanium dioxide (TiO2). The Ti-complex was obtained by melting the starting-material 1a-c and subsequently adding titanium (IV) butoxide against a stream of inert gas. The reaction was kept at high temperature for 5 hours, then the greenish/bluish solids 3a-c were purified by washing with MeOH and/or CH₂Cl₂ in order to remove the contaminants. The solids were collected each time by centrifugation and dried under vacuum to constant weight (3a: 78% yield; 3b: 65% yield; 3c: 95% vield).



Scheme 1. Reaction scheme for the synthesis of ligands 2a-c and Ti-based complexes of tetrakis-2,3-[5,6-di-R₈-pyrazino]porphyrazine (R= H, 2-Py, Ph) 3a-c. Numbers in blue will be useful in the assignment of the ¹³C-SSNMR spectra reported in Figure 6.

Solution Studies: UV-Vis Spectra.

The UV-vis spectral behavior of Titanium-based complexes **3a-c** was studied as a function of time in dimethylformamide (DMF), a polar aprotic solvent, and in methylenchloride (CH₂Cl₂), a

nondonor solvent. All complexes yielded clear solution in both solvents, and the formation of colloidal suspensions as described by Donzello *et al.* for similar compounds was not observed.^[18-19] A greater solubility was obtained in DMF than in CH₂Cl₂.

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Furthermore, changes in the intensities of the UV-vis absorptions could be observed as a function of time (Figure 1). This behavior can be ascribed to molecular aggregation both in the form of intermolecular association or dissociation upon time. Complexes **3a-c** showed a different response depending on the solvent employed. In general, in CH₂Cl₂ the spectra were characterized by an increase of Q and B bands with time (Figure 1) reaching their maximum intensities after one hour. These spectral changes can be attributed to intermolecular dissociation ending with the formation of monomeric macrocyclic units in keeping with previous results.^[18-19]

The same behavior was shown by complex **3a** in DMF. In contrast, the UV-vis spectra of compounds **3b** and **3c** in the same solvent were characterized by a progressive increase in the absorption, respectively at 761 and 759 nm, with a simultaneous decrease in the Q and B band intensities. If disaggregation could be obtained for all compounds in the nondonor solvent, CH₂Cl₂, and for compound **3a** in DMF, the introduction of aromatic substituents such as 2-pyridil (complex **3b**) and phenyl groups (complex **3c**) caused an intensification of the aggregation phenomena. These could be potentially due to an increase in the π - π stacking interactions between the Titanium-based complex monomers reaching its maximum after one hour. The UV-vis spectra in DMF are characterized by intense and sharp peaks in the Q-region respectively at 647 (**3a**), 667 (**3b**) and 665 (**3c**) nm (Figure 1 and Table 1).



solvent from DMF to CH₂Cl₂. A summary of **3a-c** absorptions, together with a comparison with the corresponding ligands **2a-c** is presented in Table 1. In particular, the introduction of eight 2-pyridil (**3b**) or eight phenyl groups (**3c**) caused a bathochromic shift in the Q band respectively of 20 nm (for complex **3b** in CH₂Cl₂ the shift is of 22 nm) and 18 nm (for complex **3c** in CH₂Cl₂ the shift is of 20 nm) in DMF with respect to the non-substituted complex **3a**. These shifts are quite small because only a partial conjugation between the substituents and the macrocycle framework is possible due to intrinsic steric hindrance of 2-pyridil and phenyl groups.

Weak vibrational bands were present between 587 and 605 nm. These peaks became more intense in CH_2Cl_2 for compounds **3b** and **3c** (peaks at 603 and 667 nm), while for complex **3a** the peak at 585 nm remained broad. The splitting in the Q band in CH_2Cl_2 has been observed for other metal complexes in chlorinated solvents and it is probably due the formation of excitonic coupling because of molecular association.^[20]

Finally, as observed for porphyrazines and phthalocyanines, because of HOMO-LUMO π - π * transitions, B bands between 315 and 421 nm are also present (Table 1).

As it can be seen from Table 1, the UV-vis spectra of compounds **3a-c** strictly parallel the ones of the corresponding un-metallated macrocycles, with the exception in the B band region. As shown in Figure 2, ligands were, generally, characterized by an envelope in the 300-400 nm region, whereas the corresponding metallated compound presented more defined and red-shifted absorptions. In contrast to complex **3a**, ligand **2a** presented a very low solubility in both DMF and CH₂Cl₂ (see Figure 2a for a comparison between **2a** and **3a**), while more soluble ligands were obtained by the introduction of 2-pyridil and phenyl groups (Figure 2b).

Table 1. UV-Vis spectral data for ligands 2a-c and complexes 3a-c in different solvents.						
Compound	Solvent	B band region [nm]	Q band region [nm]			
2a	DMF	404 435 576	636			
3a	DMF	326 587	647			
3a	CH_2CI_2	338 585	646			
2b	CH ₂ Cl ₂	355 421 581	617 634 668			
3b	DMF	315	603 667			
3b	CH ₂ Cl ₂	320 577	634 668			
2c	DMF	336	657			
2c	CH ₂ Cl ₂	342	638 667			
3с	DMF	375	605 665			
3c	CH ₂ Cl ₂	362 441	607 637 666			

Figure 1. UV-vis spectral changes of Titanium-based complexes 3a-c in DMF and CH_2Cl_2 solutions as a function of time.

Generally, a negligible solvent effect was observed in the Q band position, while the B band was subjected to a red-shift for complexes **3a-b** and to a blue-shift for complex **3c** changing the



Figure 2. Comparison between the UV-vis spectra of ligands 2a-b and complexes 3a-b in different solvents.

Electrochemical Measurements.

As proposed in literature for tetra-2,3-pyrazinoporphyrazines ligands/metal complexes and for phatholocyanine and diazepinoporphyrazine analogues, the macrocycle is able to give four one-electron reductions in the range of potentials $0.0 \div -2.0$ V vs SCE. In general, cyclic voltammograms (CVs) of ligands present, also, a fifth redox event which has been attributed to the reduction of a dianionic specie in equilibrium with the neutral porphyrazine as confirmed by the measurements in pyridine. In the case of cobalt-based complexes, two reductions on charge of Co^{II} occur prior to the classical four-reduction sequence observed for Cu^{II}, Zn^{II} and Mg^{II} complexes.

Cyclic voltammograms of complexes **3a-c** in DMSO and DMF containing Bu₄NBF₄ 0.1 M as support electrolyte are shown in Figures 3 and 4. Scan rates of 0.5 V/sec were employed for each measurement. Even though, a suspended excess of compound was employed well defined redox processes were observed, only, for complex **3a** in both solvents, particularly in DMSO. The quality of voltammograms in DMF was strongly affected by the poor solubility of compounds **3b-c**. On the other hand, low intensity redox events were clearly identified in DMSO for all three compounds. In general, no oxidations were observed up to a positive potential of 1.1 V *vs* SCE (The complete set of voltammograms recorded is summarized in the Supporting Information).

The voltammogram in DMSO of ligand 2a showed only one reversible peak at -0.77 V (Figure 3), probably due to the reduction of dianion $2a^2$. This result is in agreement with the

literature^[18-19, 21] where tetrapyrazinoporphyrazines are known to undergo deprotonation upon dissolution in DMSO and their dianion TPysPzs²⁻ are in equilibrium with the neutral forms TPysPzsH₂. In contrast, the CV of **2a** in DMF/CH₂Cl₂ 5/1 showed one irreversible peak at -0.89 V for the reduction of the corresponding dianion and a second reversible peak, barely visible, at -1.42 V, which can be probably attributed to the reduction of its neutral form (Figure 3).^[22]

As shown in Figure 3 and 4, complex **3a** exhibits in DMSO five redox couples labeled I - V. In the case of couples I - IV, the cathodic to anodic peak separation ($\Delta E = E^{p}_{c} - E^{p}_{a}$) at 0.5 V/s was in the range 85 – 100 mV; however the cathodic to anodic peak current ratios (I_{pc}/I_{pa}) were near unit, suggesting a *quasi-reversible* behavior. The CV of **3a** in DMF, **3b** and **3c** in DMSO showed a similar electrochemical behavior, although peaks were not cleanly resolved (Figures 3-4).

The half-wave potentials (E_{1/2}) and the potential separation between different stepwise reductions (Δ) are reported in Table 2 for **3a-c** complexes. For comparison, E_{1/2} and Δ of ligand **2b**/metal M^{II} cations complexes (**2b**-M^{II})^[23] and tetra-substituted oxo(phthalocyaninato)titanium(IV) complexes [(PhCH₂O)₄Pc-Ti^{IV}O and (PhOC₆H₄OCH₂)₄Pc-Ti^{IV}O]^[24] are reported in the same Table.

As shown in Table 2, the E_{1/2} and the potential separation between couples III, IV and V for compounds **3a-c** are very similar to those of metal complexes **2b-**M^{II}; as proposed in literature, these stepwise reduction sequences are assumed to be ligand-centered. Moreover, the new titanium-based tetrapyrazinoporphrazine complexes are reduced more easily than their analogues (PhCH₂O)₄Pc-Ti^{IV}O and (PhOC₆H₄OCH₂)₄Pc-Ti^{IV}O thanks to the substitution of benzene rings in phthalocyanines with pyrazine rings. It has been demonstrated that the sequence of reduction process for oxotitanium tetra-substituted phthalocyanines complexes involved two initial metal-centered Ti^{IV} \rightarrow Ti^{III} and Ti^{III} \rightarrow Ti^{III} stepwise reductions, followed by two other ligand-based reduction steps.^[24] From all these observations we propose the following mechanism for the reduction of compounds **3a-c** (Scheme 2):

 $[TPyzPz-Ti^{|1|}] + e \leftrightarrows [TPyzPz-Ti^{|1|}]$ $[TPyzPz-Ti^{|1|}] + e \leftrightarrows [TPyzPz-Ti^{|1|}]$ $[TPyzPz-Ti^{|1|}] + e \leftrightarrows [TPyzPz-Ti^{|1|}]^{-}$ $[TPyzPz-Ti^{|1|}]^{-} + e \leftrightarrows [TPyzPz-Ti^{|1|}]^{2-}$ $[TPyzPz-Ti^{|1|}]^{2-} + e \leftrightarrows [TPyzPz-Ti^{|1|}]^{3-}$

Scheme 2. Suggested mechanism for reduction of complexes 3a-c.

In other words, as suggested by cyclic voltammetry, the electrochemical behavior of complexes **3a-c** can involve a *quasi-reversible* reduction of titanium(IV) in the first two steps followed by two or three tetrapyrazinoporphyrazine-based reduction.



Figure 3. Top and middle: cyclic voltammograms of complex 3a and ligand 2a in DMSO and DMF, 0.1 M Bu₄NBF₄, scan rate 0.5 V/s. A mixture DMF/CH₂Cl₂ 5/1 was applied as solvent in order to increase the solubility of ligand 2a. Bottom: cyclic voltammograms of complex 3b and 3c in DMSO, 0.1 M Bu₄NBF₄, scan rate 0.5 V/s.

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Figure 4. Cyclic voltammograms of complex 3a in DMSO, 0.1 M Bu₄NBF₄, scan rate 0.5 V/s.

Table 2. Half-Wave Potentials^a (E_{1/2}, V vs SCE) and potential separations between different stepwise reductions (Δ) of complexes 3a-c and [2b-M^{II}]

Compound	I	П	Ш	IV	V	Δι-ιι	Δ -	Διν	Δ _{IV-V}
3a ^b	-0.087	-0.39	-1.23	-1.60	-1.89°	0.30	0.84	0.37	0.29
3a ^{c,d}	-0.32	-0.45	-1.33	-1.73	-1.95°	0.13	0.86	0.40	0.23
3b ^{b,c}	-0.062	-0.34	-1.00	-1.20	-1.58	0.28	0.66	0.20	0.38
3c ^{b,c}	-0.062	-0.39	-1.11			0.33	0.72		
[2b- Zn ^{II}] ^{e,f}	-0.34	-0.72	-1.38	-1.66	-1.83	0.38	0.66	0.28	0.17
[2b- Zn ¹¹] ^{f,g}	-0.26	-0.67	-1.38	-1.64	Ŧ	0.41	0.71	0.26	
[2b -Cu ^{II}] ^{f,g}	-0.22	-0.58	-1.22	-1.58		0.36	0.64	0.36	
[2b -Co ^{II}] ^{f,g}	-0.06	-0.76	-1.31	-1.77		0.70	0.55	0.46	
[2b- Mn ^{II}] ^{f,g}	-0.16	-0.89	-1.33	-1.64		0.73	0.44	0.47	
(PhCH ₂ O) ₄ Pc-Ti ^{IV} O ^{h,i}	-0.18	-0.58	-1.51	-1.76		0.40	0.93	0.25	
(PhOC ₆ H ₄ OCH ₂) ₄ Pc-Ti ^{IV} O ^{h,i}	-0.14	-0.55	-1.42	-1.71		0.41	0.87	0.29	

[a]: $E_{1/2} = (E_{P_c} + E_{P_a})/2$; [b]: DMSO, 0.1 M Bu₄NBF₄; scan rate 0.5 V/s; [c]: $E_{P_{cathodic}}$; [d]: DMF, 0.1 M Bu₄NBF₄; scan rate 0.5 V/s; [e] pyridine/0.1M TBAP, scan rate 0.1 V/s; [f]: ref [23]; [g]: DMF, 0.1M TBAP, scan rate 0.1 V/s; [h]: DMF, 0.1 M Bu₄NBF₄; scan rate 0.1 V/s; [i]: ref [24].

Cyclic voltammetry technique was also used to study the response of complex 3a towards electrochemical processes, such as hydrogen and oxygen redox reactions, of practical interest in energy conversion devices. To improve the electronic conductivity of the sample, compound 3a was blended with a carbon support (Vulcan XC72) in weight ratio 20:80 and mixed with 2-propanol and 5% Nafion solution in order to prepare the catalytst ink.^[25] A catalyst ink based on Pt/C (E-TEK, with 20 wt.% Pt content with respect to C) was used as benchmark. A properly designed three electrode cell configuration was used to evaluate the catalytic activity of the sample: Ag/AgCl was used as reference electrode; a glassy carbon of 4 mm diameter disk, where a thin layer of catalyst ink was coated with a loading of 50 µg/cm² of the catalyst (3a or Pt) was adopted as the working electrode (WE). A platinum wire was used as counter electrode and the cell was filled with 0.1 M HCIO₄ electrolyte.

Cyclic voltammetry was performed under nitrogen atmosphere with a scan rate of 50 mV/sec in the potential range $0.05 \div 1.2$ V (all the potential values for this measurement were reported *versus* the reversible hydrogen electrode RHE).

Figure 5 shows the 50th voltammetric cycle recorded for both **3a**and Pt-based samples. Interestingly, sample **3a** showed typical peaks due to hydrogen redox reactions (*i.e.* adsorption/desorption) visible at low potential, delivering higher current densities with respect to the Pt/C catalyst. Unfortunately, negligible electrochemical activity was observed at higher potential regions, where the oxygen reactions are expected. This result could be explained considering the acidic properties of the sample sustained by the presence of the Ti-O system, which

the sample sustained by the presence of the Ti-O system, which can participate in the hydrogen reactions by the reversible deprotonation equilibrium above explained (Scheme 2).

Furthermore, it is worth noticing pointing out that superior behavior has been obtained for bare **3a**/C composite toward hydrogen redox process with respect to platinum. The latter being the benchmark, commonly used catalyst for both hydrogen oxidation and evolution reactions occurring in a fuel cell or electrolyzer device, respectively.^[26-27] Consequently, due to the intrinsic advantages of compound **3a** with respect to platinum, in terms of cost and scalability, the proposed Ti-based complex is considered of great interest, deserving further investigations to evaluate its applicability as catalytic material in fuel cell or water electrolyzer devices.



Figure 5. Cyclic voltammograms of the investigated 3a sample (green) and of benchmark Pt/C (black) catalyst, analyzed in HClO₄, 0.1 M electrolyte at 50 mV/s scan rate.

IR Spectra.

Pyrazinoporphyrazine macrocycles object of this study exhibited quite similar IR spectral patterns. In general, signals of Ticomplexes were sharper and more intense than those of the corresponding ligands, especially between 900 and 1340 cm⁻¹. The porphyrazine macrocycle which is characterized by the presence of conjugated CC and CN double bonds, gave typical absorptions in the range 1000 – 1600 cm⁻¹ due to skeletal vibrations.

Interestingly, the introduction of the Ti=O group in complexes **3ac** caused the appearance of peaks located near 700 and 900 cm⁻¹ (**3a**: 744 cm⁻¹, 906 cm⁻¹; **3b**: 704 cm⁻¹, 743 cm⁻¹, 993 cm⁻¹; **3c**: 762 cm⁻¹, 938 cm⁻¹).^[10, 28-29] Because of the presence of the NH stretching, broad absorptions above 3000 cm⁻¹ were observed for ligands **2b-c** (**2b**: 3292 cm⁻¹, **2c**: 3053 cm⁻¹), while for compound **2a** this broad peak was shifted to 2958 cm⁻¹. Complex **3c** also showed a broad absorption around 3056 cm⁻¹, which can be attributed to the stretching of aromatic CH (see Supporting Information for full IR spectra).

Solid State Studies: Solid State NMR Studies.

For the characterization of all complexes, Solid State Nuclear Magnetic Resonance (SSNMR) ¹H-¹³C Cross Polarization-Magic Angle Spinning (CP-MAS) spectra were acquired and compared with the spectra of the corresponding ligands. It should be noted that in CP spectra the relative intensities of the different peaks are not quantitative, since signals form carbon-13 nuclei in closest

proximity to protons are enhanced by the cross-polarization process.

For complex **3a**, a single peak centered at 148.9 ppm was observed in the aromatic region, similarly to what already reported in literature ^[3], which is ascribable to carbons 1, 2 and 3 (Figure 6a). In the case of complex **3c**, three main peaks could be identified, which are centered at 128.4 ppm (carbons 5, 6, 7), 137.7 ppm (carbon 4) and at about 151.0 ppm (carbons 1, 2, 3) (Figure 6b). The spectrum of complex **3b** presented four main signals in the aromatic region centered at 123.7 ppm (carbons 5 and 7), 136.3 ppm (carbon 6), 147.9 ppm (carbon 8) and 155.6 ppm (carbons 1, 2, 3, 4) (Figure 6c). The spectra of all complexes and ligands also presented low intensity signals in the aliphatic region between 0 ppm and 70 ppm (Figure 6d-f), probably due to residual solvent molecules (such as acetone) co-crystallized with the complexes and/or to reaction by-products.

The spectra of each ligand presented carbon resonances similar to the ones observed for the corresponding complex; only a small change of their chemical shifts could be observed (ligand **2a**: 146.6 ppm; ligand **2b**:123.7 ppm, 136.9 ppm, 147.4 ppm, 155.1 ppm; ligand **2c**: 128.4 ppm, 137.2 ppm, 150.8 ppm; see Supporting Information), indicating that the coordination with the titanyl group induces only a slight modification of the symmetry and the conformational properties of the tetrapyrazinotetraazaporphyrine macrocycle.



Figure 6. Aromatic (left panel) and aliphatic (right panel) regions of the ¹H-¹³C CP-MAS spectra of complexes 3a (a, d), 3c (b, e) and R=3b (c, f). Spinning side bands are marked with asterisks.

MALDI-TOF Analysis.

Ligands and complexes were further analyzed by MALDI-TOF analysis employing 2,5-dihydroxybenzoic acid (DHB) as matrix. DHB was preferred because it undergoes less fragmentation respect to other reference compounds under laser shot (data not shown). To this end, samples were prepared according to the procedures reported in the following references.^[30-32] In general, for ligands **2a-c**, the precursor ion results from the addition of a proton to form the positively charged molecular ion [M+H]⁺ (see

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Supporting Information for related measurements). Moreover, sodium [M+H+Na]⁺ and potassium [M+H+K]⁺ adducts were also identified. Similarly, complexes **3a-c** were detected as precursor ion [M+H]⁺ as well as in cation adduct clusters. MALDI spectra and a summary of all the ligands and complexes species detected are displayed in Supplemental Information.

Morphological Studies by SEM.

Morphological studies of all Titanium-based complexes were carried by SEM analysis, as shown in Figure 6. Among all

samples, material **3a** displayed unique morphological features, resulting in agglomerates of quasi-spherical particles with spotty flat mats. When the complexity of the molecule structure increases, *i.e.* with the introduction of eight 2-pyridil and eight phenyl groups (samples **3b** and **3c** respectively), the corresponding particles exhibit important changes in terms of both shape and size distributions, as more homogeneous, layered-like structures were observed.



Figure 6. SEM images of all Titanium-based complexes.

Conclusion

In summary, this paper presented the synthesis, characterization and the electrochemical behavior of a family of ligands and Ticomplexes based on the tetrakis-2,3-[5,6-di-R8pyrazino]porphyrazine (R= H, 2-Py, Ph) macrocycle. Ligands 2ac were obtained by cyclotetramerization of precursors 1a-c in the presence of DBU as catalyst, while complexes 3a-c could be prepared by precursors melting and addition of Titanium(IV) butoxide under inert gas and dry reaction conditions. Molecular aggregation was confirmed by UV-Vis studies in different solvents. IR spectra were characterized by typical absorptions due to CC and CN skeletal vibrations. Complexes showed additional peaks related to the Ti=O group in the range 700 - 900 cm⁻¹. Broad bands due to NH stretching were also observed for the ligands. MALDI-TOF analysis was performed in order to further investigate the structures. [M+H]⁺ adducts as well cation adduct clusters were identified for both compounds 2a-c and 3a-c. Morphological studies by SEM revealed a change in the morphological features with an increase in homogeneity passing from 3a to 3b-c. The electrochemical characterization was carried out in DMSO/Bu₄NBF₄ 0.1 M. In general, ligands showed only one reversible peak that can be attributed to the dianion reduction. Despite the low solubility, it was possible to observe different redox events of low intensity for compounds 3b-c, while complex 3a gave well defined voltammograms. From the results obtained, we proposed that our Ti-complexes can participate in up to five redox events. The first two involving the step-wise quasireversible reduction of the metal centre (Ti^{IV} \rightarrow Ti^{III} and Ti^{III} \rightarrow Ti^{II}), while the remaining two or three reductions occur on charge of the TPysPz macrocycle. In the light of a future application of compounds **3a-c** in energy conversion devices, such as fuel cell or water electrolyzer, we realized a preliminary study to test the response of these catalysts towards hydrogen and oxygen redox reactions. Therefore, an ink of catalyst blended on a carbon support was used to coat **3a** on a glassy carbon. No activity toward oxygen redaction was observed but, interestingly, the presence of the Ti-O system confers the catalyst the ability to participate in hydrogen reactions delivering higher current densities with respect to Pt/C, introducing this catalyst as a possible alternative to Pt. Evaluation of the applicability of these materials as catalysts in energy conversion devices is ongoing.

Experimental Section

General

Analytical grade solvents and all commercially available reagents were used as received. Reactions were monitored by thin-layer chromatography using silica gel plates TLC Silica gel 60 F254 from Sigma-Aldrich; visualization was accomplished with UV light (254 nm) or KMnO4 stain. Flash chromatography (FC) was carried out using Merck silica gel 60 (230-400 mesh) employing mixtures of petroleum ether and acetone as eluant. ¹H-NMR spectra of starting materials were recorded on a Bruker Avance 300 spectrometer (¹H: 300 MHz at 295 K). Chemical shifts were reported to the solvent residual peak (CDCl3: ¹H = 7.26 ppm). SSNMR spectra of all complexes and their respective ligands were acquired on a Varian

InfinityPlus spectrometer working at a Larmor frequency of 400.34 and 100.67 MHz for ^1H and $^{13}\text{C},$ respectively, using a CP-MAS 3.2 mm probe. ¹H-¹³C CP-MAS spectra were recorded using a contact time of 2 ms, under High-Power Decoupling from ¹H nuclei, at a MAS frequency of 15 kHz. 15000-80000 transients were accumulated, using a recycle delay of 3 s between consecutive transients. ¹³C chemical shifts were referred to hexamethylbenzene and TMS as secondary and primary references, respectively. UV-vis spectra were recorded with a Hewlett Packard 8452 diode array spectrophotometer equipped with an external water bath circulator (Julabo Labortechnik, GmbH) for temperature control (± 0.1 °C), using quartz cuvettes with 1 cm optical path. ATR FT-IR spectra were recorded using a Thermo Scientific Nicolet iS10 spectrometer equipped with a triglycine sulfate detector (DTGS) and acquired with Omnic vers. 8.1.10 software. The spectra were the result of 32 scans with a spectral resolution of 4 cm⁻¹. An ATR correction algorithm and automatic atmospheric suppression were applied to all spectra. Cyclic voltammetry was performed on a homemade electrochemical equipment consisting of a computer-controlled potentiostat with Vernier Software Multi Purpose Laboratory Interface (MPLI) program for Windows, a glassy-carbon disk (diameter of 1 mm) as working electrode; an Ag/AgCl/KCl 3 M as the reference electrode (E vs SCE = E vs Ag/AgCI -0.032), and a platinum disc (surface 1 cm²) as the auxiliary electrode. Cyclic voltammograms were acquired at 0.5 V/sec scan speed, 25°C, in DMF or DMSO containing Bu₄NBF₄ (0.1M) as supporting electrolyte. Concentration of ligands or complexes was 2 mM. Mass Spectrometry Analysis MALDI-TOF-MS spectra were acquired using a 5800 MALDI TOF/TOF Analyzer (Sciex, Ontario, Canada). The spectra were acquired in the positive reflector mode by 20 subspectral accumulations (each consisting of 50 laser shots) in an 50-1500 mass range, focus mass 800 Da, using a 355 nm Nb:YAG laser with a 20 kV acceleration voltage. Peak labelling was automatically done by 4000 Series Explorer software Version 4.1.0 (Sciex) without any kind of smoothing of peaks or baseline, considering only peaks that exceeded a signal-to noise ratio of 10 (local noise window 200 m/z) and a half maximal width of 2.9 bins. Calibration was performed using default calibration originated by five standard spots. 2,5-dihydroxybenzoic acid (DHB) was employed as matrix. SEM images were obtained from AURIGA Zeiss Field Emission.

Synthesis of starting materials 1a-c: Compounds 1a-c were obtained by a modification of the procedure reported in ref. [17]. A solution of diaminomaleonitrile (18.5 mmols; 1 Equiv.) and the corresponding α diketone (20.4 mmols; 1.1 Equiv.) in glacial acetic acid (35 mL) was heated under reflux for 3 hours. When the diaminomaleonitrile was totally consumed, the reaction mixture was cooled down and 50 mL of water were added. The aqueous layer was extracted with 100 mL of ethyl acetate and then washed according to the following sequence with water (3 x 50 mL), a saturated solution of NaHCO₃ (4 x 50 mL; exothermic reaction), water (2 x 50 mL) and brine (2 x 50 mL). The organic layer was dried over sodium sulfate. After filtration, the solvent was removed under reduced pressure and the solid residue was purified by recrystallization from ethanol/water at 4°C.

1a: m.p. 180°C; ¹H NMR (300 MHz, CDCl₃): δ= 8.94 (s, 2 H); 76% yield.

1b: m.p. 175°C; ¹H NMR (300 MHz, CDCl₃): δ = 8.31 ppm (d, J= 6 Hz, 1H), 8.06 ppm (d, J= 9 Hz, 1H), 7.96-7.84 (m, 1H), 7.40-7.28 (m, 1H); 73% yield.

1c: m.p. 240°C; ¹H NMR (300 MHz, DMSO-d₆): *δ*= 7.58-7.45 (m, 3H), 7.45-7.33 (m, 2H); 85% yield.

Synthesis of ligands 2a-c: Pyrazinoporphyrazine macrocycles 2a-c were prepared by autocyclotetramerization of starting materials 1a-c as follows: the precursor (400 mg; 3.1 mmols) was placed in a thick test tube and heated in an oil or sand bath until the solid melted (temperature of oil/sand bath employed: compound 1a: 180°C; compound 1b: 175°C; compound 1c: 240°C). A few drops of DBU were added as catalyst. The mixture was manually stirred until a black solid was formed. After complete solidification, the solid was cooled down, finely grounded and purified by Soxhlet

extraction with methanol (16 h) followed by acetone (10 h). The green or bluish powders were dried under vacuum until they reach a constant weight

2a: ¹³C (SSNMR): 146.6 ppm. FTIR-ATR: v^{-} = 722, 725, 857, 890, 1150, 1118, 1193, 1356, 1429, 1599, 1868, 2111, 2936 cm⁻¹; UV/Vis (DMF): λ = 636, 576, 435, 404 nm; MS (MALDI-TOF): *m/z* calc for C₂₄H₁₁N₁₆: 523,1353 [M+H]⁺; found 523.0859; *m/z* calc for C₂₄H₁₁N₁₆Na: 546.1250 [M+H+Na]²⁺; found 546.2062. Elemental analysis calc for **2a**·2**C**₂H₆**O**: C 56.43%, H 3.47%, N 35.08%. Found: C 56.67%, H 3.42%, N 35.66%; 21% yield.

2b: ¹³C (SSNMR): 123.7 ppm, 136.9 ppm, 147.4 ppm, 155.1 ppm. FTIR-ATR: v^{-} = 648, 696, 744, 759, 801, 950, 992, 1092, 1105, 1144, 1225, 1360, 1431, 1470, 1500, 1585, 1566, 1864, 2096, 3285 cm⁻¹; UV/Vis (DCM): λ = 668, 634, 617, 605, 581, 421, 355 nm; MS (MALDI-TOF): *m/z* calc for C₆₄H₃₅N₂₄: 1140.3229 [M+H]⁺; found 1139.3477; *m/z* calc for C₆₄H₃₅N₂₄Na: 1162.3374 [M+H+Na]²⁺; found 1162.3042. Elemental analysis calc for **2b·2H₂O**: C 65.42%, H 3.26%, N 28.60%. Found: C 65.13%, H 3.56%, N 27.44%; 20% yield.

2c: ¹³C (SSNMR): 128.4 ppm, 137.2 ppm, 150.8 ppm. FTIR-ATR: $v^{=}$ 689, 765, 1023, 1173, 1340, 1523, 1860, 2103, 3053 cm⁻¹; UV/Vis (DMF): λ = 657, 336 nm; UV/Vis (DCM): λ = 667, 638, 342 nm; MS (MALDI-TOF): *m/z* calc for C₇₂H₄₃N₁₆: 1131.3857 [M+H]⁺; found 1131.2971; *m/z* calc for C₇₂H₄₃N₁₆Na: 1154.3754 [M+H+Na]²⁺; found 1154.2932. Elemental analysis calc for **2c**: C 76.45%, H 3.74%, N 19.80%. Found: C 77.27%, H 4.04%, N 18.36%; 18% yield.

Synthesis of complexes 3a-c: Titanium-based pyrazinoporphyrazines 3a-c were prepared by cyclotetramerization of starting materials 1a-c in the presence of titanium (IV) butoxide as follows: the precursor **1a-c** (400 mg; 3.1 mmols; 8 Equiv.) was placed under argon in a dry Schlenk flask and heated in an oil or sand bath until the solid melted (temperature of oil/sand bath employed: compound 1a: 180°C; compound 1b: 175°C; compound 1c: 240°C). Against a flux of inert gas, titanium (IV) butoxide (0.39 mmols; 1 Equiv.) was added. The evolution of a white smoke was generally observed. The reaction mixture was stirred at high temperature (temperature of oil/sand bath employed: starting-material 1a: 180°C; starting-material 1b: 175°C; starting-material 1c: 240°C) for 5 hours, then the mixture was cooled down, 30 mL of MeOH were added and the crude product was left under stirring overnight. The solid obtained was separated by centrifugation and brought to constant weight under vacuum. In order to dissolve the contaminants, the complex 3a-c finely grounded, were then resuspended in CH₂Cl₂ and left under stirring for 30 minutes. After this time, the product **3a-c** was collected by centrifugation and dried under vacuum until constant weight

3a: ¹³C (SSNMR): 148.9 ppm. FTIR-ATR: v^- = 744, 906, 909, 913, 1208, 1359, 1507, 1508, 1664, 1864, 2110 cm⁻¹; UV/Vis (DMF): λ = 647, 587, 326 nm; UV/Vis (DCM): λ = 646, 614, 585, 338 nm; MS (MALDI-TOF): *m/z* calc for C₂₄H₉N₁₆OTi: 585.0625 [M+H]⁺; found 584.9780. Elemental analysis calc for **3a**·2H₂O: C 46.47%, H 1.95%, N 36.12%. Found: C 46.23%, H 3.03%, N 33.03%; blue powder obtained with 78% yield.

3b: ¹³C (SSNMR): 123.7 ppm, 136.3, 147.9, 155.6 ppm. FTIR-ATR: v^{-} = 705, 744, 790, 958, 993, 1092, 1147, 1359, 1432, 1471, 1566, 1584, 1738, 2113 cm⁻¹; UV/Vis (DMF): λ = 667, 603, 315 nm; UV/Vis (DCM): λ = 668, 634, 605, 577, 320 nm; MS (MALDI-TOF): *m*/*z* calc for C₆₄H₃₃N₂₄NaOTi: 1224.2646 [M+H+Na]²⁺; found 1224.3872; Elemental analysis calc for **3b**·6H₂O: C 58.73%, H 3.39%, N 25.67%. Found: C 58.43%, H 4.06%, N 24.19%; green powder obtained with 65% yield.

3c: ¹³C (SSNMR): 128.4 ppm, 137.7 ppm, 151.0 ppm. FTIR-ATR: v^{-} = 690, 768, 938, 1024, 1104, 1143, 1180, 1231, 1341, 1444, 1540, 1631, 1885, 2102, 3051, 3288 cm⁻¹; UV/Vis (DMF): λ = 665, 601, 375 nm; UV/Vis (DCM): λ = 666, 637, 607, 445, 362 nm; MS (MALDI-TOF): *m/z* calc for C₇₂H₄₁N₁₆OTi: 1193.3129 [M+H]⁺; found 1193.2933; Elemental analysis

calc for $3c{\cdot}2H_2O{:}$ C 70.36%, H 3.61%, N 18.23%. Found: C 70.05%, H 3.88%, N 17.47%; green powder obtained with 95% yield.

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Keywords: Tetrapyrazinoporphyrazine • Titanium complex • Electrochemical Behavior • Spectroscopic Properties

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Entry for the Table of Contents



Tetrapyrazinoporphyrazines are privileged frameworks present in ligands and metal complexes. Choosing titanium as metal center, herein, we unveil the synthesis, the characterization and the electrochemical behavior of tetrakis-2,3-[5,6-di-R₈-pyrazino]porphyrazine (R= H, 2-Py, Ph) ligands and Ti-complexes.

Supplemental Material for:

Titanium-based

tetrakis-2,3-[5,6-di(substituted)pyrazino]porphyrazine: synthesis and characterization

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Scanning electron microscopy (SEM) analysis was performed using an AURIGA Zeiss Field Emission.

2. General procedures and characterization

2.1 General procedure for the preparation of starting material **1a-c**



Compounds **1a-c** were obtained by a modification of the procedure reported by Mørkved *et al.* (Mørkved, E. H.; Ossletten, H.; Kjøsen, H.; Bjørlo, O. J. Prakt. Chem. **2000**, 342, 83).

A solution of diaminomaleonitrile (18.5 mmols; 1 Equiv.) and the corresponding α -diketone (20.4 mmols; 1.1 Equiv.) in glacial acetic acid (35 mL) was heated under reflux for 3 hours. When the diaminomaleonitrile was totally consumed, the reaction mixture was cooled down and 50 mL of water were added. The aqueous layer was extracted with 100 mL of ethyl acetate and then washed according to the following sequence with water (3 x 50 mL), a saturated solution of NaHCO₃ (4 x 50 mL; exothermic reaction), water (2 x 50 mL) and brine (2 x 50 mL). The organic phase was dried over sodium sulfate. After filtration, the solvent was removed under reduced pressure and the solid residue was purified by recrystallization from ethanol/water at 4°C. Products were obtained with the following yields: compound **1a** 76% yield; compound **1b** 73% yield; compound **1c** 85% yield.

2.2 Characterization of starting materials 1a-c

N CN N CN 1a	1a . ¹ H-NMR (300 MHz, CDCl ₃): 8.94 ppm (s, 2 H).
	1b . ¹ H-NMR (300 MHz, CDCl ₃): 8.31 ppm (d, 1H; J= 6 Hz); 8.06 ppm (d, 1H; J= 9 Hz); 7.96-7.84 (m, 1H); 7.40-7.28 (m, 1H).
N CN N CN 1c	1c . ¹ H-NMR (300 MHz, DMSO-d ₆): 7.58-7.45 (m, 3H); 7.45-7.33 (m, 2H).

2.3 <u>¹H-NMR spectra of starting materials</u> **1a-c**





-2×10⁶



2.4 General procedure for the preparation of ligands 2a-c



Ligands **2a-c** were obtained according to the procedure reported by Ercolani, Kadish *et al.* (Donzello, M. P.; Zhongping Ou, Z.; Monacelli, F.; Ricciardi, G.; Corrado Rizzoli, C.; Ercolani, C.; Kadish, K. M. *Inorg. Chem.* **2004**, *43*, 8626).

Pyrazinoporphyrazine macrocycles **2a-c** were prepared by autocyclotetramerization of starting materials **1a-c** as follows: the precursor (400 mg; 3.1 mmols) was placed in thick test tube and heated in a oil or sand bath until the solid melted (temperature of oil/sand bath employed: compound **1a**: 180°C; compound **1b**: 175°C; compound **1c**: 240°C). A few drops of DBU were added as catalyst. The mixture was manually stirred until a black solid was formed. After complete solidification, the solid was cooled down, finely grounded and purified by Soxhlet extraction with methanol (16 h) followed by acetone (10 h). The green or bluish powders were dried under vacuum until they reach a constant weight (Compound **2a**: blue powder, 21% yield; compound **2b**: green powder, 20% yield; compound **2c**: green powder, 18% yield).



2.5 <u>UV-Vis spectra of ligands 2a-c</u>





2.6 General procedure for the preparation of Titanium-complexes 3a-c



Titanium-based pyrazinoporphyrazines **3a-c** were prepared by cyclotetramerization of starting materials **1a-c** in the presence of titanium (IV) butoxide as follows: the precursor **1a-c** (400 mg; 3.1 mmols; 8 Equiv.) was placed under argon in a dry Schlenk flask and heated in a oil or sand bath until the solid melted (temperature of oil/sand bath employed: compound **1a**: 180°C; compound **1b**: 175°C; compound **1c**: 240°C). Against a flux of inert gas, titanium (IV) butoxide (0.39 mmols; 1 Equiv.) was added. The evolution of a white smoke was generally observed. The reaction mixture was stirred at high temperature (temperature of oil/sand bath employed: starting-material **1a**: 180°C; starting-material **1b**: 175°C; starting-material **1c**: 240°C) for 5 hours, then the mixture was cooled down, 30 mL of MeOH were added and the crude product was left under stirring overnight. The solid obtained was separated by centrifugation and brought to constant weight under vacuum. In order to dissolve the contaminants, the complex **3a-c** finely grounded, then resuspended in CH₂Cl₂ and left under stirring for 30 minutes. After this time, the product **3a-c** was collected by centrifugation and dried under vacuum until constant weight (Compound **3a**: blue powder; compound **3b**: green powder; compound **3c**: green powder). Products were obtained with the following yields: compound **3a** 78% yield; compound **3b** 65% yield; compound **3c**





0,2 -0,1 -0,0 -

λ[nm]



3. Cyclic voltammograms of ligands 2a-c and complexes 3a-c

Cyclic voltammetry was performed on a homemade electrochemical equipment consisting of a computer-controlled potentiostat with Vernier Software Multi Purpose Laboratory Interface (MPLI) program for Windows, a glassy-carbon disk (diameter of 1 mm) as working electrode; an Ag/AgCl/KCl 3 M as the reference electrode (E *vs* SCE = E *vs* Ag/AgCl -0.032), and a platinum disc (surface 1 cm²) as the auxiliary electrode (Friis, E. P.; Andersen, J. E. T.; Madsen, L. L.; Bonander, N.; Moller, P.; Ulstrup, J. *Electrochim. Acta* **199**8, *43*, 1114). Cyclic voltammograms were acquired at 0.5 V/sec scan rate, 25°C, in DMF or DMSO containing Bu₄NBF₄ (0.1M) as supporting electrolyte. Concentration of ligands or complexes was 2 mM.

Titanium-based complexes **3a-c** were dissolved in 10 mL of DMSO/Bu₄NBF₄ 0.1 M or DMF/Bu₄NBF₄ 0.1 M, the solution was degassed with Argon then the voltammograms were recorded.

3.1 Cyclic voltammograms of ligands



Cyclic voltammograms of ligand 2b and 2c in DMSO,0.1 M Bu4NBF4; scan rate 0.5 V/s



Cyclic voltammograms of complex **3b** and **3c** and their ligands **2b and 2c** in DMF, 0.1 M Bu₄NBF₄; scan rate 0.5 V/s







3.5 Cyclic voltammograms of complex 3c in DMSO



3.6 Cyclic voltammograms of complex 3c in DMF



4. Solid State NMR: ¹H-¹³C CP-MAS spectra of ligands 2a-c



Aromatic (left panel) and aliphatic (right panel) regions of the ${}^{1}\text{H}{-}{}^{13}\text{C}$ CP-MAS spectra of ligands **2a** (a, d), **2c** (b, e) and **2b** (c, f). Spinning side bands are marked with asterisks.

5. MALDI-TOF analysis of ligands 2a-c and complexes 3a-c

In order to determine the mass of ligands and complexes a MALDI-TOF analysis was performed employing 2,5-dihydroxybenzoic acid (DHB) as matrix. The samples were prepared according to the following procedure (see references: Lidgard, R.; Duncan, M. W. *Rapid Commun. Mass Sp.* **1995**, *9*, 128; Canlica, M.; nyokong, T. *Polyedron* **2011**, *30*, 1975):

- A standard solution of DHB matrix was prepared dissolving 0.14 mmols of DBH in 1 mL of CH₃CN 100% (standard solution A);
- A standard solution for each ligand and complex was prepared dissolving 2*10⁻³ mmols of ligand/complex in 1 mL of CH₃CN 100% (standard solution B);
- Standard solution A was diluted 1:100 with CH₃CN 100% (standard solution C);
- Standard solution B was diluted 1:10 with CH₃CN 100% (standard solution D);
- $6 \mu L$ of standard solution C were mixed with 9 μL of standard solution D (solution E);
- 0.5 μL of solution E were applied on the MALDI plate. The solvent was let to evaporate in order to form a crystal.
- The MALDI plate was transferred inside the instrument for the analysis.
- The spectra were acquired in the positive reflector mode by 20 subspectral accumulations (each consisting of 50 laser shots) in an 50-1000 mass range, focus mass 600 Da, using a 351 nm Nb:YAG laser with a 20 kV acceleration voltage.

5.1 Mass spectrum of the DHB matrix (full spectrum)



5.2 Mass spectrum of ligand 2a



5.3 Mass spectrum of ligand 2a: enlargement and comparison with the matrix DHB



5.4 Mass spectrum of ligand 2b



5.5 Mass spectrum of ligand 2b: enlargement and comparison with the matrix DHB



5.6 Mass spectrum of ligand 2c



5.7 Mass spectrum of ligand 2c: enlargement and comparison with the matrix DHB



5.8 Mass spectrum of complex 3a



5.9 Mass spectrum of ligand 3a: enlargement and comparison with the matrix DHB





5.11 Mass spectrum of ligand **3b**: enlargement and comparison with the matrix DHB





5.13 Mass spectrum of ligand 3c: enlargement and comparison with the matrix DHB



Sample name	MALDI-TOF adduct	Mass found	Mass calculated
Ligand 2a	$[M+H]^+$ $[M+H+Na]^{2+}$ $[M+H+K]^{2+}$	523.0859 546.2062 561.2230	523.1353 546.1250 561.0990
Ligand 2b	$[M] \\ [M+H]^+ \\ [M+H+Na]^{2+} \\ [M+H+K]^{2+}$	1138.3590 1140.3229 1162.3042 1179.3101	1138.3398 1139.3477 1162.3374 1179.2218
Ligand 2c	[M+H] ⁺ [M+H+Na] ²⁺	1131.2971 1154.2932	1131.3857 1154.3754
Complex 3a	$[M+H]^+$	584.9780	585.0625
Complex 3b	[M] [M+H+Na] ²⁺	1201.4016 1224.3872	1200.2670 1224.2646
Complex 3c	$[M] \ [M+H]^+ \ [M+Na]^+$	1192.2972 1193.2933 1215.2549	1192.3050 1193.3129 1216.3026

6. IR spectra of ligands 2a-c and complexes 3a-c

6.1 IR spectra of ligands 2a



6.2 IR spectra of ligands 2b





6.4 IR spectra of complex 3a



6.5 IR spectra of complex 3b



6.6 IR spectra of complex 3c



7. SEM images of complexes 3a-c

7.1 <u>SEM images of complex 3a</u>





7.2 SEM images of complex 3b









Mag = 100.00 K X

Sample ID =

+

WD = 5.1 mm

+

7.3 SEM images of complex 3c



