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Thin Films of Fluorinated 3d-Metal Phthalocyanines as Chemical Sensors of Ammonia: an Optical Spectroscopy Study

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A comparative study of the sensor response toward gaseous ammonia of hexadecafluorinated 3d-metal phthalocyanine (MPcF₁₆, M=Cu(II), Co(II), Zn(II), Ni(II)) thin films was performed using complementary experimental (viz., surface plasmon resonance, SPR, and IR absorption spectroscopy) along with theoretical (density functional theory calculations, DFT) techniques. SPR measurements revealed changes of both thickness and optical parameters (refraction indices and extinction coefficients) of the MPcF₁₆ films caused by adsorption of NH₃. The MPcF₁₆ species studied exhibited the following order of sensor response: $ZnPcF_{16} > CoPcF_{16} \ge CuPcF_{16} > NiPcF_{16}$. A good correlation was found between the DFT calculated (B3LYP/6-311++G(2df,p)) binding energies, experimentally measured shift of the selected IR bands, and the optical sensor response. Apart from this, we performed a detailed assignment of all intense bands in the vibrational spectra (IR and Raman) of fluorinated 3d-metal phthalocyanines studied.

Keywords: hexadecafluorinated phthalocyanines; optical sensors; SPR; ammonia detection; vibrational spectra; DFT

1. Introduction

Ammonia (NH₃) is a toxic gas presented in large quantities in air, soil, and water. Exposure to low concentrations (25–150 ppm) of ammonia might lead to irritation of human skin, eyes, and respiratory tract. Moreover, exposure to high ammonia concentrations (> 5000 ppm) severely affects human health and can eventually lead to death [1]. Therefore, monitoring of ammonia in the gas phase is a very urgent problem in various applications, from industrial hygiene and environmental protection in chemical industry to clinical diagnostics, etc. [2] So far a huge variety of chemical sensors of ammonia have been proposed, including those based on inorganic, inorganic oxide/dioxide, and conducting polymers [3-7].

Among other materials, phthalocyanines, both unsubstituted (H₂Pc) and metal complexes (MPcs), are organic p-type semiconductors widely used as chemiresistive gas sensors. Moreover, thin films of MPcs are promising for gas sensing applications due to their high chemical and thermal stability and noticeable variations of both conductivity and optical properties upon sorption of analyte gases [8]. It is worth mentioning that both sensitivity and response of MPcs to various analytes were found to depend on the nature of the metal atom and substituents in the aromatic ring as well as on the structure and molecular organization in the thin films [8]. Therefore, the peripheral substitution of the conjugated macrocycle of MPcs by electron-withdrawing or electron-donating groups is a facile way to vary sensitivity and selectivity of their films toward different analytes [9-13]. More specifically, the electron-withdrawing fluorine peripheral substituents alter the sensitivity of MPc species toward reducing gases [10]. For instance, fluorinated zinc phthalocyanine turned out to be less sensitive to oxidizing gases (SO₂ and NO₂) and more sensitive to reducing gases (e.g., NH₃, H₂) [14, 15]. Furthermore, the same trends were observed in the electrical conductivity of ZnPc derivatives [16] in the presence of reducing gases (NH₃ and aliphatic amines), while ZnPcF₁₆ was noticeably more sensitive toward NH₃ and aliphatic amines than the unsubstituted ZnPc. Apart from this, the sensor response of thin films of hexadecafluorinated cobalt phthalocyanine (CoPcF₁₆) toward gaseous ammonia (concentration range from 50 to 1000 ppm) was also studied by total internal reflection ellipsometry [17]. Among other findings, it was demonstrated

that the specific binding of NH₃ molecules causes substantial shift of corresponding bands in the phase difference ($\Delta(\lambda)$) spectrum due to the increase in film thickness and change of its optical parameters.

Note that detailed mechanistic understanding of the sensing action of MPcs at the molecular level is crucial for further development of sensors. To this end, various optical spectroscopy techniques, both electronic (e.g., UV-vis absorption and fluorescence) and vibrational, are widely employed [18-25]. Since the vibrational spectroscopy techniques (IR and Raman) are sensitive to very tiny changes of the structure and environment at the molecular level, they are particularly useful for the study of relatively weak binding interactions between MPc substrate and chemical analytes [26-29]. The IR absorption spectroscopy can also be employed for identification of the MPc polymorphs [30-32].

On the other hand, modern quantum chemical (typically, density functional theory, DFT) calculations complement experiment in a very efficient way: viz., they provide valuable information concerning the structure of complexes of the adsorbed molecules (phthalocyanine-chemical analyte), their vibrational spectra, etc [27, 28, 33]. E.g., DFT calculations were used to calculate energies of formation and charge transfer properties of the complex of an analyte (methanol and ammonia) and ZnPc [27].

At the same time, although the vibrational spectra of metal-free phthalocyanine (H₂Pc) and a series of 3d-metal Pcs (e.g., CuPc, ZnPc, FePc, CoPc, NiPc) were studied in detail [34-38], much less is known about vibrational spectra of their fluorinated counterparts MPcF₁₆. To our knowledge, Raman spectra along with vibrational reorganization energies were reported only for CuPcF₁₆[39].

In the present contribution, we studied the sensor response toward ammonia vapor of hexadecafluorinated 3d-metal phthalocyanine (MPcF₁₆, M=Cu(II), Co(II), Zn(II), Ni(II), Fig. 1) thin films. To this end, complementary experimental techniques (viz., surface plasmon resonance and IR absorption spectroscopy) along with theoretical calculations (DFT) were employed. Note that quantum chemical insight into the structure and energetics of the MPcF₁₆^{...}NH₃ complexes along with IR spectroscopy allows for deeper understanding of the sensing mechanism at the molecular level. Apart from this, we performed a detailed assignment of all intense bands in the vibrational spectra of fluorinated 3d-metal phthalocyanines studied.

2. Experimental details

2.1. Materials

Copper and cobalt hexadecafluorophthalocyanines (CuPcF₁₆ and CoPcF₁₆) were obtained from commercial suppliers (Aldrich) and were used after purification by gradient sublimation in vacuum (10^{-5} Torr) . ZnPcF₁₆ was synthesized in accordance with the technique described elsewhere [40]. Nickel hexadecafluorophthalocyanine (NiPcF₁₆) was synthesized by heating an equimolar 4:1 mixture of a sublimed 3,4,5,6-tetrafluorophthalo-1,2-dinitrile (Aldrich) and nickel acetate in a vacuum-sealed glass tube (10^{-3} Torr) for a period of 7 hours at 220 °C. The tube was afterward opened, and the mixture was washed by ethanol and acetone to remove soluble organic admixtures. All synthesized MPcF₁₆ species were purified by gradient sublimation in vacuum (10^{-5} Torr) at 440-450 °C.

2.2. Deposition and characterization of thin films

Thin films of $MPcF_{16}$ (M=Cu(II), Co(II), Zn(II), Ni(II)) on various substrates were obtained by organic molecular beam deposition at room temperature under vacuum conditions (10⁻⁵ Torr) with the deposition rate of 0.6 nm·s⁻¹. The substrate types were varied according to particular experimental requirements for characterisation techniques employed. Namely, glass slides coated with about 40 nm thermally evaporated gold films were used as substrates for surface plasmon resonance (SPR) experiments, and (100) silicon wafers were used for vibrational spectroscopy measurements. Raman spectra were recorded with a Triplemate SPEX spectrometer equipped with CCD detector in a back-scattering geometry. The 488 nm, 100 mW line of an Ar-laser was used for excitation. Infrared spectra were recorded with a Vertex 80 FTIR spectrometer.

2.3. Quantum chemical calculations

The IR and Raman spectra of $MPcF_{16}$ and their complexes with ammonia were calculated using density functional theory (DFT) at the B3LYP/6-311++G(2df,p) [41, 42] level of theory. All calculations

were performed using Gaussian 09 suite of programs [43]. The experimental frequencies below 170 cm⁻¹ were not considered because of the strong mixing of external and internal vibrations in this range. The Raman intensities were not calculated because of the resonant nature of the Raman spectra obtained using the excitation wavelength in the visible spectral range.

2.4. Evaluation of sensing properties

Surface plasmon resonance (SPR) measurements in the Kretschmann configuration were performed using MPcF₁₆ thin films deposited on gold coated glass substrates. The latter were brought into optical contact with a semi-cylindrical prism (refraction index n = 1.515) using an index-matching fluid. The prism/sample combination was placed on a θ -2 θ rotation table driven by a microprocessor-controlled stepper motor (angular resolution 0.01°). The accuracy of the optical system corresponds to about 10⁻⁴° shift of the resonance angle. Surface plasmon excitation was achieved by focusing a monochromatic ppolarised light beam of a He-Ne laser (λ =633 nm) onto the prism/sample interface. More detailed description of the SPR technique is given elsewhere [44, 45]. A gas cell sealed by the sample through a rubber O-ring was used to study the gaseous ammonia adsorption onto the MPcF₁₆ films. Kinetic SPR measurements of the intensity of reflected light at a fixed angle of incidence θ^* =44.2°, which was chosen to be close to the SPR minimum on the left side of the SPR curve [44], were performed in situ during exposure to NH₃ vapor of the concentration 100 and 200 ppm. In order to study the adsorption kinetics in detail the gas was diluted by mixing with air to a certain pressure and then injected into the gas cell used for the SPR measurements.

3. Results and Discussion

3.1. Surface plasmon resonance study of sensor properties of the MPcF₁₆ thin films

As the first step of our study, we scrutinized the sensor response of the MPcF₁₆ (M = Cu(II), Co(II), Zn(II), Ni(II)) thin films toward gaseous NH₃. To this end, the surface plasmon resonance (SPR) measurements were employed. Note that qualitatively similar results were obtained for all metal

phthalocyanines (MPc) studied. Fig. 2 shows the typical SPR curves representing the variation of reflected intensity as a function of internal angle of incidence for the pure gold-coated substrate (curve a, black), the thin film of as-deposited CuPcF₁₆ (curve b, red), and the film exposed to NH₃ gas (curve c, blue), respectively. The reflected intensity of the monochromatic p-polarised light of wavelength λ =633 nm has a minimum when the incidence angle meets a resonance condition at θ_{SPR} (Fig. 2, curve a) [46]. It seen from Fig. 2 that after deposition of CuPcF₁₆ film the minimum of SPR curve shifted from 44.4° to 46.4° (Fig. 2, curves (a) and (b)). This shift is directly related to the thickness and dielectric constant of the film [46]. The exposure of the films to ammonia vapor (200 ppm in the particular case) leads to a further shift of the SPR minimum to higher angles (Fig. 2, curve c).

The kinetic response curves upon exposure of the MPcF₁₆ thin films (M=Cu(II), Co(II), Zn(II), Ni(II)) to ammonia vapors of two different concentrations (100 and 200 ppm) are shown in Fig. 3. The SPR response corresponds to the intensity of reflected light detected at a fixed angle of incidence θ = 44.2° as a function of exposure time. This particular angle of incidence was chosen to lie in a region of the SPR curve of maximum slope and close to the resonance angle for all MPcF₁₆ films studied (cf. Fig. 2). The time periods of exposure to lower (100 ppm) and higher (200 ppm) concentrations of NH₃, as well as of intermediate purging with fresh dry air, were chosen to be two minutes.

It is seen from Figure 3 that the response values corresponding to the steady state of adsorption were achieved in 20 seconds in all cases except ZnPcF₁₆ (Fig. 3(d)). The recovery times of NiPcF₁₆, CoPcF₁₆ and CuPcF₁₆ sensors are 15-20 seconds, while it is slightly longer (viz., 30 seconds) in the case of ZnPcF₁₆ film (Fig. 3(a-d)). Moreover, the sensor responses were reversible and the films were found to be stable and exhibited no significant loss in sensitivity at least after a few tens of repeated cycles of measurements. Nevertheless, further studies are necessary to determine selectivity and the detection limit of the films. Figure 3 also demonstrates that the values of photodetector response of MPcF₁₆ thin films depend on the nature of the central metal atom in the MPcF₁₆ molecule. Namely, the species studied can be arranged in the decreasing order of response: ZnPcF₁₆ > CoPcF₁₆ ≥ CuPcF₁₆ > NiPcF₁₆ (Fig. 3(a-d)).

In order to obtain more quantitative insight into the ammonia sensing process, we determined the changes of the film thickness d, refractive indices n, and extinction coefficients k (the latter are related to a complex dielectric permittivity via $\varepsilon = (n + ik)^2$) in the two cases: (a) pristine films of all MPcF₁₆ complexes studied and (b) those immediately after exposure to higher concentration of ammonia vapor (200 ppm). To this end, we fit the experimental SPR data by the Fresnel equations [47]. The details of a model employed are described elsewhere [48]. The results are summarized in Table 1.

Note that the adsorption of analyte molecules (NH₃ in the present case) modifies to some extent the UV-vis absorption spectra of the MPcF₁₆ films and, consequently, the extinction coefficient k at a particular wavelength [12, 49]. This, in turn, leads to increase of the refractive index, since the two values n and k are not independent and obey the Kramers-Kronig relations [50] (cf. Table 1 values). The changes of n and k upon exposure to NH₃ in the row of MPcF₁₆ exhibit the same trend as the above discussed response values: the most pronounced effect was observed in the case of ZnPcF₁₆ film (Table 1). Namely, adsorption of the gaseous ammonia results in a decrease of n and k values from 1.64 to 1.51 and from 0.26 to 0.20, respectively (Table 1). Apart from this, the increase of film thickness during NH₃ exposure (Table 1) gives evidence of film swelling.

3.2. Sensor properties at the molecular level: study of binding of ammonia to $MPcF_{16}$ films by IR spectroscopy and theoretical calculations

Among the huge variety of experimental techniques, vibrational spectroscopies (IR and Raman) have been shown to be very useful for study of the surface structure and properties of thin films [51-55]. These techniques are particularly suitable in the case of $MPcF_{16}$ compounds with unknown crystal structure, because the commonly employed X-ray diffraction methods cannot be applied directly in this case.

Note, however, that the vibrational spectroscopy methods can give insight into the structure of thin films only provided that the detailed assignment of the vibrational spectra of the deposited species is available. Unfortunately, in the particular case of fluorinated 3d-metal phthalocyanines and their complexes with ammonia, these spectra have not been discussed in the literature. Therefore, we started from the detailed study of the vibrational spectra of $MPcF_{16}$.

3.2.1. Experimental and theoretical study of vibrational spectra of MPcF₁₆

The MPcF₁₆ complexes are planar molecules of D_{4h} point group symmetry. Assuming a D_{4h} point group for the MPcF₁₆ species comprised of 57 atoms, the corresponding vibrational representation reads as

$$\Gamma_{vib} = 14 \ A_{1g} + 13 \ A_{2g} + 14 \ B_{1g} + 14 \ B_{2g} + 13 \ E_g + 6 \ A_{1u} + 8 \ A_{2u} + 7 \ B_{1u} + 7 \ B_{2u} + 28 \ E_u.$$

The A_{1g} , B_{1g} , B_{2g} , E_g modes are Raman active, and the A_{2u} and E_u modes are IR-active. Note that A_{1g} , B_{1g} , B_{2g} are in-plane and E_g is an out-of-plane vibration. Along with experimental measurements, the vibrational spectra of all compounds under study were calculated at the B3LYP/6-311++G(2df,p) level of theory. The experimental and calculated IR and Raman spectra of MPcF₁₆ species studied are presented in Figs. 4 and 5, respectively. A comparison of the experimental and calculated bands in the IR and Raman spectra of MPcF₁₆ and their assignments are given in the Tables 2 and 3. The experimentally measured vibrational wavenumbers of MPcF₁₆ molecule coincide well with DFT theoretical predictions. The RMS difference between the calculated and experimental wavenumbers was 20 cm⁻¹. Note that in the case of CuPcF₁₆ our results are in a fairly good agreement with recent computations at a comparable level of theory [39].

It is seen from Figs. 4 and 5 that almost all vibrational modes of the same nature (both in Raman and IR spectra) have very similar wavenumbers for all four MPcF₁₆ species studied. However, as was pointed out earlier [38], the positions of some bands are dependent on the type of metal atom. More specifically, in the case of unsubstituted MPcs, the fundamentals lying in the spectral region between 1350 and 1500 cm⁻¹ have been identified to be highly sensitive to the nature of the metal ion [36, 56]. In our case, we observed the similar features in the Raman spectra of MPcF₁₆, namely, the position of this band varies from 1507 cm⁻¹ for ZnPcF₁₆ to 1549 cm⁻¹ for NiPcF₁₆ (Table 3). The normal coordinate of this vibration is mostly comprised of displacements of the C_{α} -N_β-C_{α} bridges between the isoindole groups of the

phthalocyanine macrocycle (Tables 2 and 3) [36]. It is also worth mentioning that the variation of wavenumber of this band correlates well with the metal ion size. DFT geometry optimization yields the following values for the cavity size (more precisely, the N_{α}-M-N_{α} distance) of the metal phthalocyanines: Zn (4.00 Å) > Cu (3.94 Å) > Co (3.88 Å) > Ni (3.84 Å). These values are slightly higher (by ~ 0.05 Å) than those for the corresponding unsubstituted metal phthalocyanines [36].

Another group of fundamentals of medium intensity at 1287, 1283, 1273, and 1266 cm⁻¹ for NiPcF₁₆, CoPcF₁₆, CuPcF₁₆, and ZnPcF₁₆, respectively, involves isoindole deformations (Fig. 5, Table 3). Apart from this, the fundamentals in the range 1130-1160 cm⁻¹ are also sensitive to the central metal because they correspond to isoindole ring deformation coupled with C_{α} -N_{α}-C_{α} and C_{α} -N_{α} in-plane vibrations (Table 3). Another group of fundamentals at 937-955 cm⁻¹ was attributed to the isoindole breathing along with stretching of M-N_{α} (Table 3). Finally, there are two groups of fundamentals in the wavenumber regions 230-280 cm⁻¹ and 350-400 cm⁻¹ (Fig. 5, Table 3) dominated by M-N_{α} in-plane and out-of-plane stretching vibrations. In addition, some bands sensitive to the nature of the metal ion are also present in the IR spectra. These fundamentals lie in the wavenumber region 1440-1530 cm⁻¹ (Fig. 4, Table 2).

3.2.2. Vibrational spectra of MPcF₁₆ upon exposure to ammonia vapor

After detailed assignment of vibrational spectra described in the previous section, we studied the IR spectra of the films before and after exposure to ammonia (200 ppm). The frequency shifts of the selected vibrational bands in the IR spectra of all four MPcF₁₆ derivatives upon exposure to NH₃ are listed in Table 4. It is seen from Table 4 that the most pronounced shifts in the IR spectra were observed in the case of ZnPcF₁₆. The typical spectra of ZnPcF₁₆ before (a) and after (b) exposure to NH₃ are presented in Figure 6. Along with experimental data, the calculated vibrational frequencies of the MPcF₁₆^{...}NH₃ complexes are also presented in Table 4. It is worth mentioning that the NH₃ indeed forms the complex with MPcF₁₆ by coordination to the metal center rather than to the peripheral fluorine atoms. For instance, in the case of ZnPcF₁₆, the simple B3LYP estimations yielded the former complex to be ~15 kcal/mol more favourable on the enthalpy scale. This fact also agrees well with existing data on unsubstituted

MPcs [57, 58]. It is seen from Table 4 that the largest shift of IR bands of 5-6 cm⁻¹ is observed in the case of ZnPcF₁₆. The average shift of the corresponding bands decreases to 4, 3 and 1 cm⁻¹ for the CoPcF₁₆, CuPcF₁₆, and NiPcF₁₆ films, respectively (Table 4). The common feature of these vibrations is their sensitivity to the nature of the metal ion (Tables 2 and 3). This fact agrees well with out-of-plane distortion of the Pc core in the complex of metal phthalocyanine and ammonia (vide infra). Moreover, most of the discussed bands are red shifted upon exposure to NH₃, indicating that the corresponding normal coordinates become looser to some extent (Table 2). Note that the observed shifts of these particular IR bands also give evidence that the ammonia forms a complex with MPcF₁₆ moiety at the axial coordination position of the metal ion.

The selected DFT optimized structural parameters of MPcF₁₆ complexes with ammonia along with the calculated energies of complex formation are given in Table 5. It is worth mentioning that the calculated energetics of the complexes correlate well with metal-(ammonia) nitrogen bond lengths and, more importantly, with the sensor response measured by SPR technique (cf. Fig. 3, Table 1 and discussion therein). The values of relative formation enthalpies decrease in the same sequence: $ZnPcF_{16} >$ $CoPcF_{16} > CuPcF_{16} > NiPcF_{16}$. Note that the NH₃ axial coordination also leads to out-of-plane displacement of the central metal ion for all MPcF₁₆ derivatives (Table 5).

To get more insight into the interplay between microscopic structure and sensor response of the films, we compared the MPcF₁₆...NH₃ binding energies, shifts of IR bands, and refraction index changes (Fig. 7). It is clearly seen that the energies of MPcF₁₆...NH₃ complex formation (Table 5), red shifts of wavenumbers of some IR bonds (Table 4), and refractive index changes (Table 1) correlate well with each other (Fig. 7). Thus, on the basis of these facts we propose the variation of wavenumbers of particular IR bonds upon exposure to an analyte to be a useful descriptor for a screening among promising (i.e., those with the most pronounced changes of sensor response upon NH₃ absorption) ammonia sensors.

3.3. Comparison with similar metal phthalocyanines

Note that the sensor performance towards ammonia of several metal phthalocyanine and porphyrin species have been reported [59-61]. More specifically, the effects of the central metal atom on gas sensor performance were measured electrically (resistivity changes were monitored) for thin films of tetra-(tert-butyl)-5,10,15,20-tetraazaporphyrins (MTAP(t-Bu)₄, M = Pd, Ni, Co) [60], 1,8,15,22-tetra-iso-pentyloxyphthalocyanine (MPc(iso-PeO)₄, M = Cu, Pb, Ni) [61] and etioporphyrins (MEP) of Cu, Zn, Ni, Co, Pt, Pd [59]. It was shown that the sensor response varied in the following order: Pd > Ni > Co in the case of MTAP(t-Bu)₄ [60]; and Cu > Pb > Ni in the case of MPc(iso-PeO)₄, respectively [61]. This order has been rationalized in terms of the central ion size: the larger ion radius is, the more d-electrons participate in binding with ammonia. Note, however, that a different order of sensor response was observed in the case of MEP: CoEP > NiEP > CuEP > ZnEP [59]. It is seen that entirely different order of the sensor response was observed in the present work for strongly electron-withdrawing fluorosubstituted Pcs (ZnPcF₁₆ > CuPcF₁₆ > NiPcF₁₆).

Probably, more relevant to our case are the gas sensing properties of 2,9,16,23-tetrakis(2,2,3,3-tetrafluoropropoxy) metal(II) phthalocyanines (TFPMPc, M = Co, Zn, Cu, Pb, Pd, Ni), which have also been recently scrutinized [62]. It was shown that the electrical response of TFPMPc to ammonia vapor decreases in the row Co > Zn > Cu > Pb > Pd ~Ni, which correlates well with the DFT calculated binding energies of TFPMPc^{...}NH₃ complexes: more specifically, the higher binding energy is, the more pronounced is the sensor response [62]. It is therefore instructive to compare these results with the findings of the present work: the sensor response order $ZnPcF_{16} > CoPcF_{16} > CuPcF_{16} > NiPcF_{16}$ also exhibits a good correlation with the MPcF₁₆^{...}NH₃ binding energies (cf. Table 5 and Figure 7).

However, it should be clearly emphasized that the sensing response of a material is not solely determined by the bond strength between the central metals of the molecules of thin film and the analyte. On the contrary, many other factors can contribute noticeably: e.g., the number of active sites in a layer, tiny details of sorption-desorption process and, particularly, thickness and morphology of the film.

4. Conclusion

In the present work, adsorption of ammonia vapor to the thin films of hexadecafluorinated 3d-metal (Cu(II), Co(II), Ni(II), Zn(II)) phthalocyanines was studied in situ using surface plasmon resonance (SPR) technique. SPR measurements revealed changes of both thickness and optical parameters (refraction indices and extinction coefficients) of the MPcF₁₆ films caused by adsorption of NH₃. The MPcF₁₆ species studied exhibited the following order of sensor response: ZnPcF₁₆ > CoPcF₁₆ ≥ CuPcF₁₆ > NiPcF₁₆. IR spectroscopy along with DFT calculations provided evidence of coordination of electron-donating NH₃ species to the metal center of MPcF₁₆ molecule. The most pronounced spectral shift of 5-6 cm⁻¹ upon exposure to ammonia vapor was observed in case of ZnPcF₁₆ for the IR bands at 603, 654, 937, 960, 1148 and 1266 cm⁻¹. These shifts are mainly due to axial coordination of NH₃ to the metal ion leading to out-of-plane distortion of the MPc core. A good correlation was found between the DFT calculated binding energies, experimentally measured shift of the selected IR bands and the optical sensor response.

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	Pr	istine films		NH ₃ exposed films				
	n	k	d nm	$\Delta n/n = 0/2$	A 1z/1z 0/	\0		
	(λ=633 nm)	(λ=633 nm)	u, IIII	Δ11/11, %	Δ K / K , %	∆u/u, %		
ZnPcF ₁₆	1.64	0.26	34	7.9	23	18		
CoPcF ₁₆	1.52	0.20	22	6.1	19	16		
CuPcF ₁₆	1.49	0.22	27	5.2	13	11		
NiPcF ₁₆	1.59	0.28	30	2.5	4	4		

Table 1. Thickness (d), Indices of Refraction (n), and Extinction Coefficients (k) for Pristine and NH_3 Exposed MPcF₁₆ Films (200 ppm).

7nPcF		CoDoF		CuPcE		NiPcF		Sum	Assignments
Evet	Colod	Evet	Calad	Evet	Calad	Evet	Calad	Sylli-	Assignments
196	198	227	205	228	203	229	207	E	C - Co-C N - M-N IP
229	220	275	268	252	245	280	272	A _{2u}	$M-N_{\alpha}$ OOP, F out-of-plane motions
262	277	314	280	280	279	320	282	Eu	C_{δ} - C_{δ} - F , C_{γ} - C_{δ} - F IP
281	284		296		288		298	Eu	C_{δ} - C_{δ} - F , C_{δ} - C_{ν} - F IP
293	325		338	313	329		339	Eu	$M-N_{\alpha}, C_{\beta}-C_{\gamma}-F, C_{\gamma}-C_{\delta}-F$
329	329	341	366	337	340		374	A _{2u}	N_{α} and N_{β} OOP motions, M- N_{α}
352	372	367	407	350	381	369	411	Eu	$M-N_{\alpha}, N_{\beta}-C_{\alpha}-C_{\beta}, C_{\alpha}-C_{\beta}-C_{\gamma}$
370	390	389	388	381	387	392	392	A _{2u}	N_{β} out-of-plane motions, C_{α} - C_{β} - C_{γ} OOP, isoindole OOP deformations
428	461	428	462	428	462	428	465	E _u	$C_{\beta}-C_{\gamma}-C_{\delta}, M-N_{\alpha}$
496	497	499	499	497	498	498	502	E _u	benzene def., N_{β} - C_{α} - C_{β}
603	603	605	605	603	603	606	606	Eu	macroring breathing, M-N $_{\alpha}$
654	658	663	663	657	659	674	667	E _u	$\begin{array}{l} C_{\alpha}\text{-}N_{\alpha}\text{-}C_{\alpha}, M\text{-}N_{\alpha}, C_{\beta}\text{-}C_{\beta}\text{-}C_{\gamma}, \\ C_{\alpha}\text{-}C_{\beta}\text{-}C_{\gamma} \end{array}$
746	764	754	769	749	768	754	774	E _u	$\begin{array}{l} C_{\alpha}\text{-}N_{\beta}\text{-}C_{\alpha},C_{\alpha}\text{-}C_{\beta}\text{-}C_{\beta},C_{\beta}\text{-}C_{\gamma}\text{-}F,\\ C_{\gamma}\text{-}C_{\gamma}\text{-}F,C_{\gamma}\text{-}C_{\delta}\text{-}F \text{ IP} \end{array}$
766	782	769	788	768	784	770	792	A _{2u}	C_{α} and N_{α} out-of-plane motions
835	836	844	846	839	843	846	852	E _u	$C_{\beta}-C_{\alpha}-N_{\beta}, C_{\beta}-C_{\beta}-C_{\gamma}, C_{\alpha}-N_{\beta}-C_{\alpha}$ IP
937	937	953	952	947	944	955	957	E _u	C-F, isoindole breathing, C_{α} -N _B -C _{α} , C_{α} -C _B , M-N _{α} IP
960	963	966	969	963	967	968	976	E _u	Macroring breathing, C-F, M- N_{α} IP
1072	1074	1075	1080	1073	1077	1076	1086	E _u	C_{α} - N_{α} , C_{α} - N_{α} - C_{α} , C_{β} - C_{β} - C_{γ} IP
1130	1142	1142	1145	1141	1145	1142	1150	E _u	Benzene def., C-F, C_{α} - N_{α} - C_{α} , C_{α} - N_{α} IP
1148	1150	1157	1156	1152	1154	1158	1162	E _u	Benzene def., C-F, C_{α} - N_{α} - C_{α} , C_{α} - N_{α} IP
1266	1268	1283	1285	1273	1278	1287	1292	Eu	isoindole def., C_{α} -N _{β} , C_{β} -C _{β} IP
1312	1311	1325	1325	1320	1321	1328	1329	E _u	C_{β} - C_{β} , C_{β} - C_{γ} , C_{γ} - C_{δ} , benzene def. IP
1334	1327	1350	1334	1337	1334	1356	1332	Eu	Pyrrole def., C_{α} - N_{α} , C_{α} - C_{β} IP
	1343		1347		1345	1407	1357	E _u	macroring def.
1449	1451	1469	1468	1460	1462	1472	1477	E _u	C_{α} - N_{β} , C_{α} - C_{β} , C_{δ} - C_{δ} , C_{β} - C_{α} - N_{α} IP
1479	1483	1497	1500	1489	1490	1500	1512	E _u	C_{α} - N_{β} , C_{β} - C_{β} , C_{δ} - C_{δ} , C_{γ} - C_{δ} IP
1490	1495	1505	1516	1507	1512	1508	1526	Eu	C_{α} - N_{β} , C_{γ} - C_{δ} , C_{α} - N_{α} IP
1613	1611	1529	1531	1526	1521	1533	1547	Eu	$C_{\beta}-C_{\gamma}, C_{\gamma}-C_{\delta}, C_{\alpha}-N_{\beta}$
1639	1639	1622	1618	1617	1617	1625	1627	Eu	$C_{\beta}-C_{\beta}, C_{\delta}-C_{\delta}$
1639	1639	1639	1639	1636	1636	1640	1640	E _u	$C_{\beta}-C_{\gamma}, C_{\gamma}-C_{\delta}$

Table 2. Experimental and Calculated (B3LYP/6-311++G(2df,p)) IR Wavenumbers (cm⁻¹) of MPcF₁₆. The Atomic Labels Are in Accordance with Fig. 1.

ZnI	PcF_{16}	Col	PcF_{16}	Cul	PcF_{16}	Nił	PcF_{16}	Sym-	Assignments
Expt	Calcd	Expt	Calcd	Expt	Calcd	Expt	Calcd	metry	
120	116	144	139	127	126	133	138	B_{1g}	M-N $_{\alpha}$, macrocycle breath.
176	172	174	173	175	173	174	174	A _{1g}	macrocycle breath.
201	198	205	205	203	200	208	207	B _{2g}	N_{α} -M- N_{α} , C_{α} - N_{β} - C_{α} IP
279	279	281	282	279	280	281	283	B _{2g}	$C_{\beta}-C_{\gamma}-F, C_{\gamma}-C_{\delta}-F, N_{\alpha}-M-N_{\alpha}$
322	316	326	320	322	317		322	B _{1g}	$C_{\beta}-C_{\gamma}-F, M-N_{\alpha}$
332	320	332	326	332	323	331	329	B _{2g}	N_{α} -M- N_{α} , C_{β} - C_{γ} -F, M- N_{α}
369	330	368	330	368	331		332	A _{1g}	$C_{\beta}\text{-}C_{\gamma}\text{-}F,$ macrocycle breath., $M\text{-}N_{\alpha}$
426	437	421	444	424	440		447	B_{1g}	C_{β} - C_{γ} - C_{δ} , M- N_{α} , macrocycle breath. asym.
468	465	466	466	467	467	469	469	A _{1g}	C_{β} - C_{γ} - C_{δ} , M- N_{α} , macrocycle breath. sym.
512	513	509	514	510	514	511	516	B_{2g}	benzene def.
554	555	553	569	559	560	556	571	B _{1g}	M-N $_{\alpha}$, macrocycle breath.
585	583	584	585	583	583	585	587	A_{1g}	M-N _{α} , inner ring breath., C _{α} -N _{α} -C _{α}
728	724	734	733	733	729	739	739	A _{1g}	$C_{\alpha}-N_{\beta}-C_{\alpha}, N_{\alpha}-C_{\alpha}-N_{\beta}, M-N_{\alpha}$
743	753	747	756	746	751	752	758	\mathbf{B}_{1g}	N_{β} - C_{α} - N_{β} , inner ring def., M- N_{α} IP
768	773	767	772	768	773	770	777	\mathbf{B}_{2g}	$\begin{array}{c} C_{\alpha}\text{-}N_{\beta}\text{-}C_{\alpha},C_{\alpha}\text{-}C_{\beta}\text{-}C_{\beta},C_{\beta}\text{-}C_{\gamma}\text{-}F,\ C_{\gamma}\text{-}\\ C_{\gamma}\text{-}F\ IP \end{array}$
811	813	815	819	814	817	819	825	B _{2g}	$C_{\beta}-C_{\alpha}-N_{\beta}, C_{\beta}-C_{\beta}-C_{\gamma}, C_{\alpha}-N_{\beta}-C_{\alpha}$ IP
946	959	967	962	964	961	965	967	A _{1g}	Macroring breathing, C-F IP
1059	1067	1062	1070	1065	1069	1069	1077	\mathbf{B}_{2g}	C_{α} - N_{α} , N_{α} - M - N_{α} , isoindole def. IP
1135	1139	1133	1142	1135	1142	1112	1137	A_{1g}	Benzene def., C-F, C_{α} -N $_{\alpha}$ -C $_{\alpha}$ IP
1193	1196	1189	1196	1193	1197	1193	1201	\mathbf{B}_{2g}	C_{α} -N _{α} , isoindole def., C _{δ} -F IP
	1295		1307		1303		1313	B_{1g}	C_{α} - N_{α} - C_{α} , C_{α} - C_{β} - C_{β}
1202	1297	1212	1316	1209	1309	1202	1328	A _{1g}	$C_{\beta}-C_{\beta}, C_{\gamma}-F$ IP
1505	1308	1512	1314	1508	1312	1505	1322	B_{2g}	C_{α} - N_{α} , benzene def. IP
	1310		1328		1322		1341	B _{1g}	C_{β} - C_{β} , C_{δ} - C_{δ} , benzene def. IP
1342	1346	1331	1342	1339	1345	1335	1349	A _{1g}	macroring def.
1354	1355	1363	1362	1360	1359	1377	1372	B_{1g}	C_{α} - C_{β} , C_{β} - C_{γ} , benzene def. IP
1399	1398	1396	1405	1404	1404	1411	1416	A _{1g}	macroring def.
1441	1472	1479	1480	1460	1480	1488	1490	A _{1g}	Benzene breathing, C_{α} -N _{β}
1474	1479	1492	1481	1483	1481	1505	1489	\mathbf{B}_{1g}	$C_{\beta}-\overline{C_{\beta}, C_{\delta}-C_{\delta}, C_{\beta}-C_{\gamma}, C_{\alpha}-N_{\beta}}$
1507	1511	1524	1541	1522	1528	1549	1560	A _{1g}	$C_{\alpha}-N_{\beta}, C_{\alpha}-C_{\beta}$
1536	1550	1536	1580	1536	1567	1560	1595	B _{1g}	C_{α} - N_{β}
1612	1614	1615	1620	1616	1620	1623	1630	A _{1g}	$C_{\beta}-C_{\beta}, C_{\delta}-C_{\delta}, C_{\gamma}-C_{\delta}$
1630	1630	1633	1633	1633	1633	1637	1637	B _{2g}	$C_{\beta}-C_{\gamma}, C_{\gamma}-C_{\delta}$

Table 3. Experimental and Calculated (B3LYP/6-311++G(2df,p)) Raman Shifts (cm⁻¹) of MPcF₁₆. The Atomic Labels Are in Accordance with Fig. 1.

ZnPcF ₁₆				CoPcF ₁₆			CuPcF ₁₆				NiPcF ₁₆				
e	xpt	ca	lcd	ех	kpt	ca	lcd	ех	xpt	Ca	alcd	exj	pt	Ca	alcd
Pristine	NH ₃	MDoE	MPcF ₁₆	Pristine	NH ₃	MDoE	MPcF ₁₆	Pristine	NH ₃	MDoE	MPcF ₁₆	Pristine	NH ₃	$MPcF_{16}$	MPcF ₁₆
film	expos.	MPCF ₁₆	NH ₃	film	expos.	WIF CI 16	NH ₃	film	expos.	WIF CI 16	NH ₃	film	expos.		NH ₃
603	597	603	597	605	603	605	602	603	601	603	600	606	606	606	605
654	648	658	653	663	658	663	661	657	655	659	657	674	674	667	667
937	926	937	931	953	948	952	948	947	945	944	941	955	954	957	956
960	956	963	957	966	962	969	966	963	960	967	964	968	967	976	975
1148	1142	1150	1144	1158	1154	1156	1153	1152	1149	1154	1151	1158	1158	1162	1161
1266	1261	1268	1264	1283	1278	1285	1281	1273	1272	1278	1275	1287	1287	1292	1291

Table 4. Experimental Wavenumbers (cm⁻¹) of the Selected IR Bands of $MPcF_{16}$ Thin Films Before and After Exposure to Ammonia Vapor and DFT Calculated (B3LYP/6-311++G(2df,p)) Wavenumbers of the Axially Coordinated $MPcF_{16}$...NH₃ Complexes.

Table 5. The M-NH₃ (M=Co, Cu, Ni, Zn) Bond Lengths (in Å), N_{α} -M-N_{α} Angles (degrees), Relative Electronic Energies (ΔE), Enthalpies ($\Delta (\Delta H^{0})$), and Gibbs Free Energies ($\Delta (\Delta G^{0})$) of Complex Formation (at 298 K in the Gas Phase) of the Various MPcF₁₆^{...}NH₃ Compounds Calculated at the B3LYP/6-311++G(2df,p) Level of Theory.

Compound	ΔE, ^a kcal/mol	$\Delta (\Delta H^0),^a$ kcal/mol	$\Delta (\Delta G^0)$, ^a kcal/mol	d(M-NH ₃), Å	N_{α} -M- N_{α}^{b} angle, deg.
ZnPcF ₁₆ NH ₃	-13.5	-11.9	-5.2	2.21	157
CoPcF ₁₆ NH ₃	-10.4	-8.9	-1.5	2.29	173
CuPcF ₁₆ NH ₃	-9.7 ^c	-8.3 ^c	-1.8 ^c	2.46	173
NiPcF ₁₆ NH ₃	-1.9	-1.3	4.1	3.37	179

^a The MPcF₁₆+NH₃ asymptote is a reference for calculations of the relative thermodynamic properties. ^b The atomic labels are in accordance with Fig. 1. ^c Calculated at the B3LYP/6-311G(d,p) level of theory.

Figure captions

Figure 1. Chemical structure of MPcF₁₆.

Figure 2. Surface plasmon resonance curves for (a) pure gold-coated substrate; (b) gold/as-deposited $CuPcF_{16}$ film; and (c) gold/CuPcF₁₆ exposed by NH₃ gas (200 ppm).

Figure 3. Surface plasmon resonance (SPR) response curves of MPcF₁₆ thin films toward gaseous NH₃: (a) NiPcF₁₆, black; (b) CuPcF₁₆, red; (c) CoPcF₁₆, blue; (d) ZnPcF₁₆, green. The SPR response curves were recorded using the wavelength of λ =633 nm (He-Ne laser) at a fixed angle of incidence θ = 44.2°.

Figure 4. The experimental IR spectra of MPcF₁₆: (a) NiPcF₁₆, black; (b) CuPcF₁₆, red; (a) CoPcF₁₆, blue; (d) ZnPcF₁₆, green.

Figure 5. The experimental Raman spectra of MPcF₁₆: (a) NiPcF₁₆, black; (b) CuPcF₁₆, red; (a) CoPcF₁₆, blue; (d) ZnPcF₁₆, green.

Figure 6. Experimental (upper part) and calculated (lower part) IR spectra of $ZnPcF_{16}$ films before (a, black curves) and after (b, red curves) exposure to gaseous ammonia (200 ppm).

Figure 7. Refractive index change of the MPcF₁₆ films after exposure to ammonia vapor vs calculated relative electronic energies (ΔE) (a, blue); the shift of the position of the "inner ring breathing" IR band (at 955, 953, 947, 937 cm⁻¹ for NiPcF₁₆, CoPcF₁₆, CuPcF₁₆ and ZnPcF₁₆, respectively) vs calculated relative electronic energies (b, red).



Figure 1. Chemical structure of $MPcF_{16}$.



Figure 2. Surface plasmon resonance curves for (a) pure gold-coated substrate; (b) gold/as-deposited $CuPcF_{16}$ film; and (c) gold/CuPcF₁₆ exposed by NH₃ gas (200 ppm).



Figure 3. Surface plasmon resonance (SPR) response curves of MPcF₁₆ thin films toward gaseous NH₃ (100 ppm and 200 ppm, respectively): (a) NiPcF₁₆, black; (b) CuPcF₁₆, red; (c) CoPcF₁₆, blue; (d) ZnPcF₁₆, green. The SPR response curves were recorded using the wavelength of λ =633 nm (He-Ne laser) at a fixed angle of incidence $\theta = 44.2^{\circ}$.



Figure 4. The experimental IR spectra of MPcF₁₆: (a) NiPcF₁₆, black; (b) CuPcF₁₆, red; (a) CoPcF₁₆, blue; (d) ZnPcF₁₆, green.



Figure 5. The experimental Raman spectra of MPcF₁₆: (a) NiPcF₁₆, black; (b) CuPcF₁₆, red; (a) CoPcF₁₆, blue; (d) ZnPcF₁₆, green.



Figure 6. Experimental (upper part) and calculated (lower part) IR spectra of $ZnPcF_{16}$ films before (a, black curves) and after (b, red curves) exposure to gaseous ammonia (200 ppm).



Figure 7. Refractive index change of the MPcF₁₆ films after exposure to ammonia vapor vs calculated relative electronic energies (ΔE) (a, blue); the shift of the position of the "inner ring breathing" IR band (at 955, 953, 947, 937 cm⁻¹ for NiPcF₁₆, CoPcF₁₆, CuPcF₁₆ and ZnPcF₁₆, respectively) vs calculated relative electronic energies (b, red).