UV–Vis Spectra and Aggregation of Hybrid Binuclear Lacunar Complexes

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Dedicated to Academician Aslan Yu. Tsivadze on the occasion of his 75th Birthday

Four hybrid binuclear metal phthalocyaninate-capped iron(II) and nickel(II) tris-pyridineoximates have been studied both in solutions and the fluoroplastic thin films by UV-Vis absorption technique. The studied polymer-immobilized cage metal complexes form the H-type dimers within the polymer matrix.

Keywords: Macrocycles, zirconium phthalocyanine, hafnium phthalocyanine, iron complexes, nickel complexes, aggregation.

Электронные спектры и агрегация гибридных биядерных лакунарных комплексов

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Методом электронной спектроскопии охарактеризованы четыре гибридных биядерных комплекса трипирдиноксиматов железа(II) и никеля(II) с металлофталоцианинами в растворах и тонких фторопластовых пленках. Показано, что в полимерной матрице исследованные металлоложекомплексы образуют димеры H-типа.

Ключевые слова: Макроциклы, циркониевый фталоцианин, гафниевый фталоцианин, комплексы железа, комплексы никеля, агрегация.
Hybrid molecular systems based on the functionalized tetrapyrrolic compounds are intensively studied during last decade due to their emergent and attractive physicochemical properties caused by intramolecular interactions between their electronically coupled fragments. Nowadays, three main strategies of designing and preparation of hybrid molecular systems have been developed. They include: (i) the direct covalent modification of the tetrapyrrolic macrocycle, (ii) supramolecular self-assembly of the pre-organized building blocks as well as (iii) the modern approaches of the coordination chemistry. The latter pathway has been used recently to obtain of the hybrid binuclear metal phthalocyaninate-capped tris-pyridineoximates, the molecules of which combine the optical and chemical properties of their macro(bi)cyclic precursors, tetrapyrrolic chromophores and main transition metal clathrochelates. Such hybrid coordination compounds with relatively independent \( \pi \)- and \( \sigma \)-electronic systems can be considered as promising components of molecular platform and scaffold of smart materials for molecular electronics, photovoltaic devices and single-molecular magnets.

In this paper we are report the UV-Vis spectral characterization of four hybrid binuclear metal phthalocyaninate-capped tris-pyridineoximates (Figure 1a) and their aggregation in various media. Metal(II)-hafnium(IV)- and metal(II)-zirconium(IV)-binuclear complexes with apical macroheterocyclic-capping fragment have been obtained by transmetallation of mono(triethylantimony)-capped iron(II) and nickel(II) \( \text{tris}\)-pyridineoximates with zirconium(IV) or hafnium(IV) phthalocyaninates as Lewis acids under mild reaction conditions.

The UV-Vis spectra of the above hybrid complexes (Figure 1b) and the initial zirconium and hafnium(IV) phthalocyaninates (M(Cl\(_2\))Pc) in DMF solution are rather similar in the most intensive fragments bands which assigned \( \pi \rightarrow \pi^* \) transitions in the single tetrapyrrolic ligand \( \lambda_{\text{em}}=335-340 \text{ nm}, \Delta \lambda_{\text{em}}=686 \text{ nm, } \lambda_{\text{em}}=465 \text{ cm}^{-1} \). A low-intense broad band near 530 nm in the spectra of iron(II)-centered binuclear complexes (1, 2) assigned as the metal-to-ligand charge transfer (Fed\( \rightarrow \text{L}\pi^*\) band. The additive character of the above spectra, which are practically suggest the superposition bands of their precursors for both components reveals weak interactions between their Pc \( \pi \)-electronic system and that of the pseudomacrobicyclic metal(II)-centered fragment. A 10-nm bathochromic shift of the MPcs absorption bands upon formation of the hybrid complex can be explained by small distortion of this macroheterocycle.

Despite an unsufficient effect of such covalent hybridization on the UV-Vis spectra of 1-4, their fluorescence is completely quenched as compared to the parent Zr(Cl\(_2\))Pc \( \lambda_{\text{em}}=700 \text{ nm, } \Delta \lambda_{\text{em}}=15 \text{ nm} \).

Single crystal X-ray diffraction data show the formation of the “base-to-base” oriented dimers in the X-rayed crystal with the interbase distance of approximately 3.35 Å caused by the cofacial \( \pi \)-\( \pi \)-stacking interactions between the adjacent tetrapyrrolic fragments. The same type of supramolecular self-assembly is observed in polar aqueous solutions, while in pure DMF solution both of MPcs and the hybrid binuclear complexes 1-4 are monomeric. An addition of water to these solutions resulted in their dimerization with a substantial decrease in intensity of the characteristic absorption bands of the monomeric species with a simultaneous appearance of a new band near 650 nm which is hypsochromically shifted as compared with Q-band of initial monomeric species (Figure 2a). This suggests the formation of the \( H \)-type of dimeric species.

As it can be seen from data of the Table 1, the obtained monomer-to-aggregate ratios in aqueous solutions of the complexes 1-4 with DMF (1 % vol.) and those for their MPcs precursors are substantially different (Figure 2b). Hybrid iron(II)-centered complexes predominantly exist in the form of \( H \)-type dimers in polar media, while nickel(II)-centered complexes consist of equable amounts of monomeric and aggregated forms. The initial Zr(Cl\(_2\))Pc and Hf(Cl\(_2\))Pc in the same solutions exist mainly in monomeric forms.

A similar aggregation behavior of the above hybrid binuclear compounds has been observed in the polymer films produced from poly(vinylidene fluoride-co-hexafluoropropylene) copolymer (PVDF-HFP) (Table 1). Polymer thin films (d=50 μm) were obtained by slow evaporation of the acetone PVDF-HFP solutions (10 % wt.) containing hybrid complexes 1-4 and initial MPcs in dry air. All of them exist predominantly in a monomolecular form in such polymer matrix with lowest monomer-to-aggregate ratios in the case of the iron(II)-centered complexes.
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Because of the excellent thermal and chemical resistance, ferroelectric, piezoelectric and pyroelectric properties of PVDF-HFP matrix, these metal-containing composites seem to be prospective for the design of magnetic smart materials, as well as for the molecular electronics and sensing applications.

Thus, the UV-Vis spectra of the hybrid compounds 1-4 are rather similar in their monomolecular forms, while their aggregation degree in polar media is slightly affected by the nature of the central metal(II) ion. All the complexes under study form the $H$-type dimers both in their aqueous solutions and within the corresponding PVDF-HFP matrix.

The polymer-immobilized hybrid complexes 1-4 exist in the systems described above, predominantly in their monomolecular forms and thus are promising smart materials for the various practical applications.

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