New sterically-hindered o-quinones annelated with metal-dithiolates: regiospecificity in oxidative addition reactions of a bifacial ligand to the Pd and Pt complexes†

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An unusual reactivity of sterically hindered o-quinones with an annelated dithiete ring towards coordination at a dithiolene site has been discovered. New Pd and Pt dithiolate complexes have been synthesized. The reaction proceeds regioselectively, and the quinone site of the parent ligand is not affected even while using an excess of the metal complex. Both Pt and Pd complexes display a square planar surrounding for the metal ion and have very similar NMR, IR and UV/Vis spectra. Surprisingly, being coordinated at the dithiolene site to the metal, the ligand exhibits activity like an o-quinone, it could be reduced with different metals resulting in the corresponding o-semiquinonates which were confirmed by EPR spectroscopy. It was shown that an unpaired electron exhibits HFC with the phosphorus nuclei of phosphine ligands coordinated to the metal ions at the dithiolene site of the molecule.

Introduction

o-Quinones have been a subject of research interest for several decades. Their stable sterically hindered derivatives are utilized in coordination chemistry as redox-active bidentate ligands. Metal o-semiquinonate complexes possess a huge variety of chemical and physical properties, such as valence tautomerism,¹ photo and thermomechanical effects,² etc. Studies of metal dithiolenes have grown enormously in the last few years. Their potential applications are in molecular inorganic conductors and superconductors,³ biomimetic catalysts⁴ and so on. The introduction of heteroatomic functionalities into the o-quinone molecule can drastically change its properties as well as increase the number of coordination sites. In the latter case the quinone becomes a bridging ligand.⁵ The regioselectivity towards binding of the metal ions at different coordination sites of such ligands is of special interest.

Results and discussion

A new sterically-hindered o-quinone 1, annelated with a dithiete ring, has been synthesized recently.⁶ This species is regarded as a ligand comprising 1,2-dithiolene and 1,2-dioxolene sites ready for metal coordination. Previously we have shown that 1 acts in the reactions with alkali metals, thallium, and copper as a typical o-quinone: it involves a dioxolene site in the reduction leading to the corresponding o-semiquinonates (SQ) or catecholates. As far as the dithiolate face is concerned, it was functionalized using an excess of decacarbonyl dimanganese in toluene (⁵), but only after quinone was bound (Scheme 2).⁶ Thus, the regioselective addition to the dithiolene site was not carried out initially.

Herein we present new dithiolate complexes of Pd (2) and Pt (3) with ligand 1. The intriguing feature of these complexes is the vacant o-quinone functionality, therefore the newly synthesized species attract great interest as ligand–precursors of heterobimetallic complexes.

Scheme 1  Synthesis of dithiolate complexes 2–3 from ligand 1.
The dithiolate complexes of transition metals have been thoroughly studied due to their unique properties and application in the field of magnetic materials,\textsuperscript{7} non-linear optics,\textsuperscript{8} dyes\textsuperscript{9} and others. One of the most widely used synthetic methods for the preparation of dithiolate complexes is the treatment of dithiols or their alkali salts with metal halides. 1,2-Dithielenes also have been described as a potential precursor for the synthesis of dithiolate complexes as alternative species. But taking into account their non-availability, the chemistry of oxidative addition of the metals to the benzodithienes has not been studied yet. There is a review dedicated to metal complexes derived from 3,4-bis(trifluoromethyl)1,2-dithiete,\textsuperscript{10} and single examples of Pd and Pt interaction with 3,4-di(adaman-tan-1-yl)-1,2-dithieth\textsuperscript{11} and 4,4-di-tert-butyl-1,2-dithietan-3-one.\textsuperscript{12} From this point of view, oxidative addition of metals with simultaneous cleavage of the S–S bond in the dithiete ring of ligand 1 is a possible path for the preparation of dithiole metal complexes. This synthetic route requires low-valent metal compounds that are easy to be derived for both palladium and platinum.

The reactions of 1 with M[PPh\textsubscript{3}]\textsubscript{4} (M = Pd, Pt) were carried out in the stoichiometric ratio of 1 : 1 in THF at room temperature under vacuum. The colour of the solution turned from violet to green within 30 minutes. Recently, Rawson et al. reported the oxidative addition chemistry of trietioicins (equivalent to a dithiete dimer) to group 10 metals leading to the corresponding dithilates.\textsuperscript{13} But the reaction with tetra-thiocins requires microwave heating for activation.

Products 2 and 3 were separated in the crystalline form, and their structures were established by a single-crystal X-ray diffraction study.

Structural studies of 2 show that the palladium center is surrounded by two phosphorus and two sulfur atoms in a slightly tetrahedrally distorted square planar geometry (Fig. 1). The Pd(1) atom deviates from the P(1)P(2)S(1)S(2) plane by 0.065(3) Å. The P–Pd–P, S–P–S and P–Pd–S angles vary in the interval 85.03(2)–98.71(2)°. The Pd–S and Pd–P bond distances are 2.2707(6) Å and 2.3230(5) Å, respectively. These values correlate with those for the complex Pd(td)(PPh\textsubscript{3})\textsubscript{2} (6) (td = 1,2-toluenedithiolate), described by Cao and coworkers (Scheme 3).\textsuperscript{14} It is noteworthy that a series of Pd complexes with the 4,5-dimethoxybenzenedithiolate ligand and chelate phosphines (7–9) also have similar structural parameters of the metal center surrounding.\textsuperscript{15}

As far as a 6-membered carbon ring is concerned, the steric repulsion between bulky tert-butyl substituents on one hand and sulfur and oxygen atoms on the other hand causes its strong distortion. However, an obvious bond alternation as well as the values of bond lengths O(1)–C(5) and O(2)–C(4) (see Table S1 in the the ESI†) confirm that the species 2 exhibits an α-quinone nature. One more crystal polymorph of 2 was obtained by slow cooling of THF solution (2b). The structural parameters of the metal center are close to 2a, but the crystal lattice contains an additional molecule of THF (Fig. S1†).

The crystals of 3 were produced by slow mixing of THF solution with hexane. An X-ray data study reveals a square planar coordination of 3 (Fig. 2 and S2†). The parameters of 2 are close to a square planar geometry and given in Table S1.\textsuperscript{1} In the quinone ring the distortion is strong: the torsion angle OCCO is 37.36°. An analysis of the structural data allows us to conclude that 3 might be regarded as an α-quinone.
The IR spectra of 2 and 3, exhibiting strong absorption, characteristic of C=O groups, are very similar. Quite similar signals appear in both the $^1$H and $^{13}$C NMR spectra for 2 and 3. In $^{31}$P-NMR of 3 heteronuclear spin–spin coupling between $^{31}$P and $^{195}$Pt takes place, and a satellite doublet can be observed apart from the main singlet.

The redox characteristics of 2–3 were assessed by using cyclic voltammetry in DMF (Fig. S12–S13). The measurements were carried out under the same conditions for both Pd and Pt dithiolate as well as for parent $\sigma$-quinone 1, which reversibly reduced at $-0.42$ V (Fig. S11†). Reduction of 2 displays two irreversible waves at $-0.90$ V and $-1.20$ V, respectively ($E_{\text{red}}$ vs. Ag/AgCl). Similarly, reduction of 3 displays two irreversible waves at $-0.80$ V and $-0.94$ V ($E_{\text{red}}$ vs. Ag/AgCl). Compared with 1, such results indicate that coordination of the metal fragment at the dithiolene site significantly lowers the electron-acceptor properties of the dioxolene structure: the first reduction potential shifts by $0.48$ V and $0.38$ V for 2 and 3, respectively. Electrochemical oxidation of 2 and 3 shows two-electron quasi-reversible waves at $0.95$ V and $1.13$ V, respectively.

The UV/Vis spectra of the complexes in THF are depicted in Fig. S12. The redox characteristics of $\sigma$-quinones reactions with alkali metals, thallium amalgam and deca-carbonyl dimanganese. The formation of the corresponding $\sigma$-semiquinonates 10–15 (Scheme 4) was established by EPR spectroscopy (Fig. 3–5 and S16–S18†). The parameters of the spectra are shown in Table 1. It should be noted that all of the reduced derivatives 10–15 were studied only in solution. Their isolation in the crystal phase failed due to their relatively high lability.

EPR studies show that an unpaired electron undergoes hyperfine coupling (HFC) with two equivalent phosphorus nuclei of the phosphine ligands coordinated to the metal ion at the dithiolene site of the molecule. At the same time, the coupling constants of the metal nucleus at the dioxolene coordination site are considerably lower than those for mononuclear $\sigma$-semiquinonates. Thus, the HFC constant of $^{203,205}$Tl nucleus in THF decreases from 52.5 G (for thallium semiquinonate (SQ) with a non-coordinated dithiolene site) to 30.3 G (10) and 31.7 G (13). It is a known fact that the value of the splitting constant of the metal nuclei in semiquinonates depends on the redox potential of the quinone ligand. Taking this into account, its small values confirm that the oxidative ability of the dioxolene structure becomes lower after metal addition to the dithiolene site. This is consistent with the data of electrochemical measurements, which showed a significant shift of the first reduction potential for 2 and 3 compared to parent quinone 1. Moreover, treatment of 2 with additional equivalents of a mild reductant such as Pd(PPh$_3$)$_4$ does not yield binuclear species (Scheme 1), although the formation of catecholates easily proceeds in the case of a better acceptor o-chloranil or 3,6-di-tert-butyl-o-benzoquinone.

Further reduction of semiquinonates 10 and 13 with an excess of thallium amalgam doesn’t lead to formation of the
corresponding catecholates: the process proceeds with destruction of the dithiolate metal center. Thus, the preparative path for the synthesis of SQTl species by preliminary exhaustive reduction of the quinone with an excess of thallium amalgam to catecholate followed by addition of an equimolar amount of o-quinone is not applicable for 2 and 3 (Scheme S1 in ESI†).

Relatively high natural abundance of $^{195}$Pt allows for observation of satellite splitting. The EPR spectrum demonstrating these signals is registered for potassium o-semiquinonate 14 (Fig. 5). Apart from the main triplet attributed to HFC with phosphorus, a less intensive satellite doublet of triplets appears due to splitting of nucleus $^{195}$Pt (33.8%, $I = 1/2$, $g_{Pt} = 7.4$ G). The lines in the right component of doublet are wider because of anisotropy, which is typical of platinum complexes. Comparing the HFS constant value at the nucleus of platinum in the binuclear complex 14 with those in the corresponding semiquinonate complexes, where the platinum nucleus is situated in close proximity of the unpaired electron containing dioxolene coordination site, one can easily see that the constant differs by less than two times (12.7–13.2 G in Pt semiquinonates). This confirms a substantial delocalization of the spin density to the periphery of the binuclear complex.

It’s known that the interaction of quinones with Mn$_2$(CO)$_{10}$ in toluene leads to the formation of SQMn(CO)$_4$, and the subsequent addition of triphenylphosphine to the starting mixture gives the species SQMn(CO)$_3$PPh$_3$. The reaction of Mn$_2$(CO)$_{10}$ with 2 and 3 results in the formation of SQMn(CO)$_3$PPh$_3$ regardless of the fact whether PPh$_3$ is present in the starting mixture or not. We suppose that the presence of some amount of phosphine molecules in the solution could be explained by partial dissociation of parent complexes (Scheme 5). Then this phosphine interacts with manganese carbonyl resulting in the corresponding adducts 12 and 15.

To summarize the coordination properties of the bifacial ligand 1, the regioselectivity of oxidative addition depends on the metallic nature of the reducing agent. Strong oxidants, in which the ions possess high Lewis acidity and are singly-charged, attack presumably at the dioxolene coordination site, whereas divalent cations with mild Lewis acidity form complexes at the dithiolene site of the ligand. The dioxolene coordination site in the dithiolate Pt and Pd complexes...
exhibits o-quinone properties and remains ready for the metal coordination. Thus, both the coordination sites of the ligand could be involved in the binding of the metal ions. An unpaired electron localized mainly on the semiquinone site of the molecule acts as a spin label reporting information about the arrangement of the coordination sphere of the metal ion at the periphery. This may be useful in the study of dynamic processes in the coordination sphere of dithiolen complexes. The potential application of such a bifacial ligand might be to construct of an electric junction between the metal surface and coordination molecules in the solution: the dithiolen coordination site could be used for anchoring to the metal cluster or even surface or whereas the dioxolene node is suitable for building the functional coordination polymeric chain. Other areas of potential application for such combined dithielen and dioxole systems are molecular devices, non-linear optics, biological sensors and so on.

Experimental

Reagents and solvents were used as purchased or purified by standard procedures. All synthetic procedures were carried out under vacuum. 2,5-Di-octa-1,5-diene-3,4-dione was prepared according to a literature standard procedures. Reagents and solvents were used as purchased or purified by distillation or sublimation.

Instrumentation

IR spectra were recorded on a FSM-1201 FTIR spectrometer. UV/Vis spectra were recorded on a Perkin-Elmer Lambda 125 spectrometer in 0.1 cm quartz cells. 1H, 13C and 31P NMR spectra were recorded on a Bruker EMX spectrometer. The samples for EPR study were prepared in a quartz ampoule equipped with a three-electrode cell, C3. Conditions: 100 mV s−1 scan rate, 0.1 M Bu4NBF4, glassy carbon working electrode.

Bis(triphenylphosphine) palladium(n) 3,6-di-tert-butyl-4,5-dioxocyclohexa-2,6-diene-1,2-bis(thiolate) (3). A THF solution of 2,5-di-tert-butyl-7,8-dithiabicyclo[4.2.0]octa-1,5-diene-3,4-dione (1, 0.56 g, 2 mmol) and Pt(PPh3)4 (2.47 g, 2 mmol) was stirred for 30 minutes at 25 °C under vacuum. The colour of solution turned from violet to yellow-green. The resulting solution was filtered and evaporated. The crude product was crystallized by slow mixing of THF solution with hexane and isolated (1.70 g, 1.7 mmol, 85% yield) as green crystals.

Notes and references

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