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[CpNi(diselenolene)] Neutral Radical Complexes: Electron Paramagnetic Resonance and Density Functional Theory Investigations

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⁷⁷Se-enriched CpNi(bds) (bds = 1,2-benzenediselenolate), has been synthesized and its *g* tensor and ⁷⁷Se hyperfine tensors have been obtained from its frozen solution electron paramagnetic resonance (EPR) spectrum. These parameters are consistent with those calculated by density functional theory (DFT); it is shown that 10% of the spin is localized on each selenium and that the direction associated to the maximum ⁷⁷Se couplings is aligned along the g_{min} direction, perpendicular to the Ni(bds) plane. EPR measurements and DFT calculations are also carried out on the ⁷⁷Se enriched complex CpNi(dsit) as well on the two dithiolene analogues CpNi-(bdt) and CpNi(dmit). The optimized structures of the isolated CpNi(bds) and CpNi(bdt) complexes have been used to generate the idealized dimers (bds)NiCp···CpNi(bds) and (bdt)NiCp···CpNi(bdt) characterized by Cp···Cp overlap. The exchange parameters *J* calculated at the DFT level for these systems are in reasonable accord with the experimental values. The influence of the geometry of the dimer on its magnetic properties is assessed by calculating the variation of *J* as a function of the relative orientation of the two Ni(diselenolene) or Ni(dithiolene) planes.

Introduction

Nickel dithiolenes constitute an important class of complexes that can easily give rise to paramagnetic intermediates. This property, potentially useful in material sciences, has stimulated many electron paramagnetic resonance (EPR) investigations; this was the case, for example, for a large variety of monoanionic, formally Ni^{III}, d⁷, Ni bis-dithiolene complexes compounds.¹⁻³ Dithiolene complexes, in which Ni is coordinated to a cyclopentadienyl ring, have been considerably less studied.^{4,5} It has been realized only recently that CpNi(dithiolene), as well as their diselenolene analogues, present specific properties that are particularly attractive for the elaboration of magnetic materials. These 17-electron complexes are robust, neutral and paramagnetic.⁶ The absence of counterion in the crystal matrix facilitates interaction between the complexes and is quite appropriate for exchange couplings between the paramagnetic centers, as confirmed for every of these complexes,⁴⁻⁷ where extended onedimensional or three-dimensional magnetic structures were observed, such as uniform spin chains, alternated spin chains or ordered antiferromagnets. On the other hand, for CpNi(bdt), the variation of the temperature dependence of the magnetic susceptibility clearly reveals the presence of antiferromagnetic interactions that cannot be ascribed to an extended set of interactions. For these systems indeed, the crystal structure shows that, in contrast with other CpNi(dithiolene) complexes, the intermolecular metal-sulfur or sulfur-sulfur distances are rather large and suggest that the origin of the exchange coupling lies in an unexpected bimolecular interaction between the cyclopentadienyl rings of two CpNi(bdt) complexes facing each other through a Cp···Cp interaction.⁴ A similar behavior was also recently observed in the CpNi(adt) complex where the faceto-face Cp···Cp interaction of two CpNi(adt) complexes (Chart 1) was held for responsible for the large antiferromagnetic interaction measured in this complex.⁸ Calculation of the spin densities on the Cp ring indicated indeed that up to 20% spin density was indeed delocalized on the Cp ring, in CpNi(bdt)⁴ as well as in CpNi(adt).⁸

However, confirmation of this hypothesis of a strong antiferromagnetic interaction through an intermolecular Cp····Cp interaction requires to experimentally accessing the exact spin delocalization in the isolated complex. Direct information about the spin densities can, a priori, be obtained from hyperfine interactions; for CpNi(dithiolene) complexes, however, no proton coupling is resolved on the EPR spectra and, due to the poor abundance of its magnetic isotope (³³S, $I = \frac{3}{2}$, natural abundance = 0.75%), no sulfur coupling can be detected. Therefore, substitution of sulfur atoms by selenium atoms appears to be the most pertinent method: ⁷⁷Se has a natural abundance of 7.3% and its spin number is only 1/2. Unfortunately, the large line width of the EPR signals of CpNi(bds) precluded the detection of the corresponding satellite lines. We therefore decided to enrich some CpNi(diselenenolene) complexes with ⁷⁷Se isotope and to determine their ⁷⁷Se couplings and g tensor from their frozen solution spectra. In this study, we report the preparation and the EPR study of ⁷⁷Se enriched CpNi (bds) and CpNi(dsit), (Chart 2), as well as the measured g values for the corresponding dithiolene complexes CpNi(bdt) and CpNi(dmit).

To test the ability of density functional theory (DFT) calculations to predict the structure of the isolated complexes, the experimental g and ⁷⁷Se coupling tensors are compared with

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CHART 1



those predicted by this theory and the calculated exchange coupling constant J resulting from the interaction between two CpNi(bds) or two CpNi(bdt) complexes is compared with the value previously obtained from magnetic measurements. An important objective of these investigations is also to point to the dependence of the magnetic properties of (bds)NiCp···CpNi(bds) or (bdt)NiCp···CpNi(bdt) on the relative orientation of the two monomers.

Experimental Methods

Synthesis. *General Details.* All reactions were carried out under an argon atmosphere by means of standard Schlenk techniques. All solvents for chemical reactions were dried and distilled by Na-benzophenone (for toluene) or CaH₂ (for methanol) before use. Metallic ⁷⁷Se was obtained from Eurisotop (Saclay, France), a subsidiary of Cambridge Isotope Laboratories, Inc.

Synthesis of ⁷⁷Se-Enriched (NEt₄)₂[Zn(dsit)₂]. A 1.6 M solution of *n*-butyllithium in hexane (0.44 mL, 0.7 mmol) was slowly added from a syringe to a stirred solution of diisopropylamine (0.09 mL, 0.65 mmol) in 2 mL of dry THF under -78 °C. After 1 h, the THF solution (3 mL) of vinylene trithiocarbonate (44 mg, 0.33 mmol) was added over 15 min. The reaction mixture was allowed to react under -78 °C for 3 h. The solid ⁷⁷Se (48 mg, 0.62 mmol, red form) was directly added into the reaction mixture in one portion. The vigorously stirred solution was allowed to warm up to room temperature, and reacted for 18 h. Then a dark-red solution was obtained. The solvent was evaporated under reduced pressure under argon atmosphere. A methanol solution (3 mL) containing tetraethylammonium bromide (137 mg, 0.65 mmol) and zinc chloride (45 mg, 0.33 mmol) was added into the residue. Then a red precipitate was rapidly formed. After stirring for 30 min, water (10 mL) was added to precipitate the salt. The cloudy solution was filtered and the red solid obtained was washed with water. The red solid was dried under reduced pressure. A 211 mg amount of the crude 77Se4-(NEt4)2[Zn(dsit)2] containing 77Serich isotope was obtained. This product might include some impurities (e.g., remaining elemental ⁷⁷Se and vinylene trithiocarbonate), but it can be used directly for the next reaction.

Synthesis of ⁷⁷Se-Enriched [CpNi(dsit)]. The crude ⁷⁷Se₄– (NEt₄)₂[Zn(dsit)₂] (211 mg) with ⁷⁷Se and [Cp₂Ni](BF₄) (130 mg, 0.47 mmol) were reacted in refluxing methanol for 2 h. ⁷⁷Se₂–[CpNi(dsit)] was obtained in 20% overall yield based on elemental ⁷⁷Se. HR-Mass (MALDI⁺) Calcd for C₈H₅NiS₃⁷⁷Se₂: 408.7305. Found: 408.7270.

Synthesis of ⁷⁷Se-Enriched (PPh₄)₂[Zn(bds)₂]. Solid sodium (11.9 mg, 0.49 mmol, 1 equiv) and ⁷⁷Se (37 mg, 0.48 mmol, 1 equiv) were weighted under argon in a glovebox in a small test tube, DMF (1 mL) was added, and the mixture was warmed to 100 °C for 2 h. *o*-Dibromobenzene (38 μ L, 0.32 mmol, 0.5 equiv)

was added directly in the test tube and the mixture was stirred at 140 °C for 40 h. The mixture was then cooled to room temperature, centrifuged (3×5 mn at 5000 t/min) and washed with MeOH (3×2 mL) to give an orange solid (32%). NaBH₄ solid (6.5 equiv) was added to a suspension of this poly(*o*diselenobenzene) (2 equiv) in MeOH (2 mL) at 0 °C and stirred for 1 h. ZnBr₂ (1 equiv) was then added to the solution, and after 1 h at 0 °C, PPh₄Br (1 equiv) was added. The mixture was stirred at room temperature for 2 h to give an orange precipitate that was centrifuged (3×5 mn at 5000 t/min) and washed with MeOH (3×2 mL) to give a orange-brown solid (185 mg, 64%). This crude product was used without further purification in the next step.

Synthesis of ⁷⁷Se-Enriched [CpNi(bds)]. The crude ⁷⁷Seenriched [PPh₄]₂[Zn(bds)₂] prepared above was treated under argon with (Cp₂Ni)BF₄ (132.3 mg, 0.48 mmol, 2 equiv) in dry MeOH (5 mL) and warmed to reflux for 2.5 h. After evaporation, the solid is taken in CH₂Cl₂/cyclohexane 1:1 and chromatographied on silica gel to afford the title compound as a dark solid in low yield (15 mg, 16%).

EPR. Measurements were carried out with a Bruker 300 spectrometer (X-band, 100 kHz field modulation). Frozen solution spectra were obtained by very rapidly cooling down the solution to avoid the formation of small crystallites. Frozen solution spectra have been simulated with the XSophe-Xepr-View computer simulation software⁹ or with a program written in our laboratory that uses the Levenberg–Marquardt algorithm.¹⁰

Calculations. Geometries of the isolated complexes were optimized with no constraint using the Gaussian 03 package¹¹ and the hybrid functional B3LYP.¹² The standard 6-311G* ¹³ basis set was used for H, C, S, and Se together with the ECP Stevens-Basch-Krauss basis (SBK)¹⁴ for Ni. Energy minima were confirmed by frequencies analysis. The g tensors and the ⁷⁷Se hyperfine tensors of these complexes were obtained with the B3LYP functional, at the optimized geometries. Three sets of calculations were performed to predict these properties: (1) by using the Gaussian 03 program with the aug-cc-pVDZ basis set for H, C, S, Se atoms¹⁵ and SBK for Ni (set a); (2) by using the ORCA program¹⁶ with the aug-cc-pVDZ for H, C, S, Se atoms and the all-electron Gaussian basis sets triple- ζ quality TZV^{17} for Ni (set b); (3) by using the basis set mentioned for set b together with the scalar relativistic zeroth-order regular approximation program (ZORA)^{18,19} implemented in ORCA (set c). The exchange coupling parameter J has been calculated for the "dimeric" species with the ORCA program using the basis sets mentioned for set b. To assess the role of the functional on the calculated value of J, several types of functionals have been used for the calculations of this parameter: hybrid (B1LYP, B3LYP, PBE0²⁰), generalized gradient approximation (GGA) (BP86, PBE, RPBE, BLYP) and local density approximation (LDA). The variation of J as a function of the dihedral angle SeNiNiSe has been calculated with the BP86 functional. The GaussView program has been used for the representation of molecules and orbitals.

Results

⁷⁷Se Enrichment of CpNi(diselenolene). The syntheses of non-⁷⁷Se-enriched diselenolene complexes have been published,^{4,21,22} and they implied the reaction of nickel diselenolene complexes, $[Ni(bds)_2]^0$ or PhSb(dsit) with, respectively, Cp₂Ni or (Cp₂Ni)BF₄. However, to be able to carry these reactions with the highest yields on the minimal quantities imposed by the availability of the ⁷⁷Se isotope, we have developed a novel

SCHEME 1



preparation based on the reaction of the stable and easily available zinc bis-dithiolene or diselenolene complexes with $(Cp_2Ni)BF_4$. This reaction was first perfected for the synthesis of CpNi(dmit) from $(Et_4N)_2[Zn(dmit)_2]$ and $(Cp_2Ni)BF_4$, affording CpNi(dmit) in 70% yield. The dsit complex, CpNi(dsit), was similarly obtained from $(Et_4N)_2[Zn(dsit)_2]$ and $(Cp_2Ni)-BF_4$ in 56% yield under the same conditions (2h reflux in MeOH). Based on these strong improvements, it was therefore possible to consider the preparation of the ⁷⁷Se-enriched CpNi(⁷⁷dsit), as outlined in Schemes 1 and 2, respectively.

EPR Results. The X-band EPR spectrum obtained at room temperature with a solution of CpNi(bdt) in THF exhibits a single line centered around g = 2.05, with a line width of ~10 G. A single line is also observed with non-enriched diselenolene complexes CpNi(bds) or CpNi(dsit); in these cases, however, the *g* value is around 2.09 and the line width is close to 25 G. With the ⁷⁷Se-enriched compounds, the line width slightly increases, but no hyperfine coupling is resolved.

As shown by its frozen solution spectrum (Figure 1a), the *g* tensor of CpNi(bdt) is axial with $g_{\perp} < g_{\parallel}$. The spectrum obtained with CpNi(bds) (Figure 1b) exhibits some departure from axial symmetry. The *g* tensors of the two complexes have been determined by a simulation/optimization process; the resulting principal values are given in Table 1 together with the values measured for CpNi(dmit) and CpNi(dsit). For these complexes, the anisotropy ($g_{max} - g_{min}$) is around 0.10; as expected from the room-temperature spectra, the average value for the dithiolene compounds is slightly smaller than for the diselenolene complex.

Enrichment of CpNi(bds) with ⁷⁷Se causes a drastic modification of its EPR response: a well-resolved hyperfine structure is now observed on the frozen solution spectrum (Figure 2a). A similar modification is observed when CpNi(dsit) is enriched in ⁷⁷Se (see Figure 2b).

The spin Hamiltonian used for the simulation of the CpNi-(⁷⁷bds) spectrum takes the *g* tensor and the hyperfine tensors *T* with two ⁷⁷Se nuclei into account. The *g* tensor is, of course, practically identical to the tensor measured for the non-enriched compound. The two ⁷⁷Se *T* tensors exhibit a pronounced axial symmetry, they have the same principal values and their ⁷⁷Se *T*_{II} are aligned along the *g*_{min} direction. The spectra are not very sensitive to the orientation of the "perpendicular" eigenvectors; the very good simulation shown in Figure 2a was performed with *T*_{⊥1} (56 MHz), making an angle of 20° with *g*_{max}. The signs of the hyperfine couplings cannot be directly determined from the spectra. However, a combination that assumes the sign of $T_3 = T_{\text{max}}$ is opposite to the sign of T_1 and T_2 leads to the smallest absolute value for the isotropic coupling constant ($A_{\text{iso}} \sim 26$ MHz); such a small value, together with the large line width of the signal, explains the absence of any hyperfine splitting on the liquid-phase spectrum. Accordingly, with the DFT results (*vide infra*), the combination leading to a positive isotropic coupling constant has been retained. The resulting experimental ⁷⁷Se isotropic and anisotropic coupling constants are reported in Table 1 together with those measured for CpNi-(⁷⁷dsit).

DFT Results on Isolated Complexes. Structure optimizations show that, in accord with preliminary calculations on CpNi-(bdt),⁴ the Ni(bdt) and Ni(bds) moieties are planar and oriented perpendicular to the Cp ring, in good accordance with the crystal structures.⁴ Bond angles and torsion angles agree with those measured by X-ray diffraction; nevertheless, the Cp····Ni distances obtained by DFT are slightly larger than those measured in the crystal (Cp····Ni distances (Å): CpNi(bdt), 1.785 (DFT), 1.721 (crystal); CpNi(bds), 1.789 (DFT), 1.718 (crystal)).

Several recent reports have shown that the DFT method is now able to predict not only the hyperfine couplings but also the *g* tensors with a good accuracy.^{23–26} The *g* tensors, obtained with the basis sets mentioned as sets a, b, c in the experimental section, are given in Table 2. For the four complexes, the g_{min} eigenvector is oriented perpendicular to the Ni-containing ring (*z*-axis), and the two other principal directions are oriented parallel and perpendicular to the chalcogen–chalcogen direction of this ring (Figure 3).

For CpNi(⁷⁷bds) and CpNi(⁷⁷dsit), the three calculation sets (sets a, b, c), lead to the ⁷⁷Se hyperfine coupling constants shown in Table 2. For each complex, the two ⁷⁷Se A_{iso} constants are practically equal as well as the corresponding anisotropic coupling constants ⁷⁷Se τ_{aniso} . The two ⁷⁷Se dipolar tensors have their maximum component (τ_{max}) aligned along the g_{min} direction (*z*-axis); the minimum component (τ_1) makes an angle of ~12° with the corresponding Se–Ni bond direction.

DFT Results on Interacting Complexes. Recently, several reports have shown that the broken symmetry approach proposed by Noodleman^{27,28} leads to very satisfactory predictions of the exchange coupling parameter $J^{.29-33}$ In this method, J is given by $J_{\rm N} = [E_{\rm HS} - E_{\rm BS}]/S^2_{\rm max}$, where $E_{\rm HS}$ corresponds to the energy of the triplet (high-spin state) and $E_{\rm BS}$ to the energy of the singlet (low-spin state, where the spin-up electron quasi localizes on the first monomer and the spin-down electron quasi localizes on the second one) calculated in the broken symmetry unrestricted formalism. Another equation for J was proposed by Yamaguchi³⁴ ($J_{\rm Y} = [E_{\rm HS} - E_{\rm BS}]/[\langle S^2 \rangle_{\rm HS} - \langle S^2 \rangle_{\rm BS}]$. Although this DFT–BS method has been criticized,³⁵ it enables us to calculate our rather large system, (bds)NiCp····CpNi(bds) without having to approximate it to a simpler model.

First, the value of *J* has been calculated for the idealized dimer shown in Figure 4. In this system each CpNi(bds) "monomer" adopts the geometry optimized for the isolated complex (Ni···Cp distance = 1.789 Å). To mimic the crystal structure, the two monomers are related by a rotation of $2(2\pi/5)$ around the *x*-axis, oriented perpendicular to the two Cp rings and bearing the two nickel atoms and the centers of the two cyclopentadienes; this leads to SeNiNiSe torsion angles equal to ~36°, and in the real system two slightly different values are found for these angles: 31.2° and 24.8°. The distance between the two cyclopentadiene rings is equal to 3.37 Å as measured in the crystal.

The J_N and J_Y values calculated for this idealized dimer by using various functionals are reported in Table 3 together with



Figure 1. Experimental (top) and simulated (bottom) EPR spectra obtained at 77 K with a THF solution of CpNi(bdt) (a) and CpNi(bds) (b).



Figure 2. Experimental (top) and simulated (bottom) EPR spectra obtained at 77 K with a THF solution of ⁷⁷Se enriched CpNi(⁷⁷bds) (a) and CpNi(⁷⁷dsit) (b).

		g tensor			⁷⁷ Se coupling tensor (MHz)			77 Se $ au_{aniso}$ (MHz)			
complex	g_1	g_2	g 3	g_{av}	T_1	T_2	T_3	$ au_1$	$ au_2$	$ au_3$	⁷⁷ Se A _{iso} (MHz)
CpNi(bdt) CpNi(⁷⁷ bds) CpNi(dmit) CpNi(⁷⁷ dsit)	2.121 2.146 2.096 2.161	2.024 2.110 2.028 2.137	2.018 2.028 2.014 2.050	2.054 2.095 2.046 2.116	(-)56 (-)59	(-)33 (-)37	(+)168 (+)188	(-)82 (-)90	(-)59 (-)68	(+)142 (+)157	(+)26 (+)31

TABLE 1: Experimental EPR Tensors

the values predicted for the geometry directly obtained from the crystal structure (Ni····Cp distance = 1.7 Å, angle between the Cp mean planes = 4.17°).

To assess the dependence of J with the topology of the "dimer", we have also calculated the variation of this parameter as a function of the dihedral angle ξ formed by the two diselenolene or dithiolene planes. The resulting curves, obtained for (bds)NiCp···CpNi(bds) by rotating one of the two Ni-(diselenolene) units around its symmetry axis and by keeping fixed the two cyclopentadienyl rings, are shown in Figure 5a. A similar variation is obtained by rotating one of the two CpNi-(diselenolene) complexes around this axis. The angular dependence of $E_{\rm HS}$ (triplet energy) and of $E_{\rm BS}$ (singlet energy) is reported in Figure 5b.

Discussion

The large size of the studied complexes as well as the presence of several atoms with a high spin—orbit coupling constant (one Ni, two or five chalcogens) is not particularly favorable to an accurate calculation of the g tensor. As shown

by Table 2, this difficulty is reflected by the incidence of the choice of the basis set on the calculated data. Nevertheless, most of the *g* characteristics measured on the spectra are reproduced by DFT: the g_{average} values, rather close to the experimental ones, are larger for the diselenolenes than for the dithiolenes and the calculated anisotropies ($g_{\text{max}} - g_{\text{min}}$) are consistent with EPR measurements.

The spin delocalization on the two selenium atoms can be roughly estimated by comparing the experimental ⁷⁷Se couplings with the atomic constants.³⁶ This leads, for CpNi(⁷⁷bds), to a very small spin density in the s-orbitals of Se ($\rho_s = 26/20120 = 0.1\%$) and to an appreciable p-character ($\rho_p = 142/982 = 14\%$). Similar values are found for CpNi(⁷⁷dsit): ($\rho_s = 0.1\%$, $\rho_p = 16\%$). The DFT calculations confirm these results: for the two Ni selenolene complexes, the Fermi contact interaction, probably due to inner shell polarization, is small and agrees with the A_{iso} value obtained from the frozen solution spectra using the pertinent sign combination (*vide supra*); the maximum anisotropic coupling component τ_{max} is practically equal to the experimental value. Consistent with the EPR results, for each

TABLE 2: Calculated g and ⁷⁷Se-coupling Tensors

			g tensor				77 Se $-\tau_{aniso}$ (MHz)			
	basis set	g_1	g_2	g ₃	$g_{ m av}$	$ au_2$	t_1	$ au_{ m max}$	⁷⁷ Se A _{iso} (MHz)	
CpNi(bdt)	set a	2.1156	2.0442	2.0318	2.0638					
• • •	set b	2.1119	2.0462	2.0245	2.0608					
	set c	2.1225	2.0442	2.0300	2.0655					
CpNi(bds)	set a	2.1383	2.1103	2.0480	2.0988	-109	-25	+135	+23	
	set b	2.1506	2.1433	2.0402	2.1113	-102	-40	+142	+29	
	set c	2.1535	2.1338	2.0489	2.1123	-94	-31	+125	+21	
CpNi(dmit)	set a	2.0754	2.0574	2.0129	2.0485					
	set b	2.0703	2.0554	2.0095	2.0450					
	set c	2.0799	2.0554	2.0131	2.0482					
CpNi(dsit)	set a	2.1413	2.1167	2.0309	2.0963	-106	-55	+161	+11	
	set b	2.1812	2.1155	2.0234	2.1067	-101	-66	+167	+26	
	set c	2.1660	2.1262	2.0309	2.1077	-93	-55	+147	+19	

^{*a*} Set a, G03, B3LYP, basis: C, H, S, Se, aug-cc-pVDZ; Ni, SBK. Set b, ORCA, B3LYP, basis: C, H, S, Se, aug-cc-pVDZ; Ni, TZV. Set c, ORCA, B3LYP, basis: C, H, S, Se, aug-cc-pVDZ; Ni, TZV, ZORA.



Figure 3. Orientation of the g_{\min} and ⁷⁷Se τ_{\max} principal directions for CpNi(bds).



Figure 4. Idealized dimer formed by two CpNi(bds) units.

 TABLE 3: Calculated^a J (cm⁻¹) Values for the Dimeric

 Structures of CpNi(bdt) and CpNi(bds)

		(CpNi	$(bdt))_2$			(CpNi(bds)) ₂			
	crystal structure		idealized structure		crystal structure		idealized structure		
	$J_{ m N}$	$J_{ m Y}$							
LDA GGA	-350	-282	-317	-257	-394		-371		
BP86 PBE RPBE BLYP	-239 -237 -217 -253	-213 -211 -196 -223	-215 -212 -192 -225	-192 -190 -175 -200	-270 -266 -242 -286	-237 -234 -217 -248	-252 -248 -224 -265	$-222 \\ -220 \\ -202 \\ -232$	
hybrid B1LYP B3LYP PBE0 exptl	-78 -89 -71 -279	-77 -88 -70	-67 -80 -61	-66 -79 -60	-83 -99 -76 -313	-83 -97 -75	-73 -92 -66	-72 -90 -65	

^a ORCA, aug-cc-pVDZ basis sets for H, C, S, Se and TZV for Ni.

complex, the two selenium atoms have their τ_{max} eigenvector aligned along the principal direction of g_{min} . This reflects the appreciable contribution of the selenium p_{π} orbitals to the SOMO shown in Figure 6.

The spin densities calculated by DFT are given in Table 4. The values predicted for the selenium atoms very well agree with the EPR spectra; they confirm, for example, that spin delocalization on each selenium atom is slightly smaller for CpNi(bds) than for CpNi(dsit). The same decrease in the spin densities on the two chalcogen atoms is predicted when passing from the dmit to the bdt complex; it is accompanied by an increase in the Ni spin density. A similar spin transfer (\sim +10%) from the chalcogens to the metal occurs when passing from the dithiolene to the corresponding diselenolene complex. The total spin densities on the cyclopentadiene ring, are close to 20% for both CpNi(bds) and CpNi(bdt).

Because DFT calculations are consistent with EPR measurements, at least for the g tensors and the selenium spin densities, it is worthwhile using optimized structures of the isolated CpNi-(bds) and CpNi(bdt) complexes to understand the magnetic properties of these compounds in the crystal state. Consistent with magnetic measurements, the DFT results shown in Table 3 for the idealized dimer or for the exact crystal structure geometry, indicate an antiferromagnetic coupling. All the values of $J_{\rm N}$ obtained with GGA functionals are between -190 and -290 cm^{-1} and are quite different from the values obtained with hybrid functionals. The dependence of J upon the type of functional used for the calculations has been reported in many recent articles.^{30,35,37} The absolute value of J calculated using generalized gradient approximation (GGA) are generally found to be larger than those calculated with hybrid functionals but smaller than with local functionals. Similar variations are observed in Table 3. However, in contrast with the values given in this table, in most of the previous studies a rather good agreement between experiment and calculation was obtained with hybrid functionals, whereas GGA was found to overestimate the antiferromagnetic behavior.^{37,38} For the [CpNi(bds)]₂ and the [CpNi(bdt)]₂ systems, the better accord between measured and predicted J values is obtained by using the BLYP functional. It is difficult to know, for the moment, if this accord is due to some specificities of these systems (e.g., bridge formed by two rings without covalent bonds) or if magnetic interactions with other neighboring CpNi(bds) or CpNi(bdt) units, contributing to the measured J value. However, due to the large intermolecular chalcogen ··· chalcogen distances measured on the crystal,⁴ this last interpretation seems less probable.

It is important, for the design of new materials, to detect the structural parameters that drive the magnetic properties of a molecular complex. These properties are the outcome among several factors such as hybridization of the various atoms, bond lengths, molecular conformation,³⁹ Due to the axial nature of the Cp···Cp bridge linking the two magnetic moieties in



Figure 5. (a) Variation of the exchange coupling parameter J_N with the SeNiNiSe dihedral angle ξ for (bds)NiCp···CpNi(bds). ξ_0 corresponds to the idealized dimer; J_{exp} is the experimental value. (b) Corresponding potential energy surfaces (singlet and triplet) as a function of the SeNiNiSe dihedral angle ξ . Curves a and b calculated with the BP86 functional.



Figure 6. Representation of the SOMO for CpNi(bds).

	basis set ^a	Se(1)	Se(2)	Ni	C ₆ H ₆	C_2S_2CS	C ₅ H ₅
CpNi(bdt)	set a	0.15	0.15	0.49	0.01		0.20
	set b	0.16	0.16	0.48	0.02		0.18
	set c	0.14	0.15	0.52	0.02		0.17
CpNi(bds)	set a	0.10	0.09	0.60	0.01		0.20
	set b	0.11	0.11	0.58	0.02		0.18
	set c	0.10	0.09	0.62	0.02		0.17
CpNi(dmit)	set a	0.23	0.23	0.29		0.10	0.15
	set b	0.23	0.22	0.29		0.13	0.13
	set c	0.22	0.21	0.33		0.11	0.13
CpNi(dsit)	set a	0.17	0.17	0.45		0.04	0.17
	set b	0.18	0.18	0.42		0.07	0.15
	set c	0.16	0.15	0.48		0.06	0.15

TABLE 4:	Calculated	Spin	Densities
	Curcuracea	N PILL	L'UNDIVIU.

^{*a*} Set a, G03, B3LYP, basis: C, H, S, Se, aug-cc-pVDZ; Ni, SBK. Set b, ORCA, B3LYP, basis: C, H, S, Se, aug-cc-pVDZ; Ni, TZV. Set c, ORCA, B3LYP, basis: C, H, S, Se, aug-cc-pVDZ; Ni, TZV, ZORA.

RNiCp···CpNiR (R = bdt or bds), these "dimers" are specially appropriate to assess the role of the relative orientation of the two Ni(diselenolene) (or dithiolene) planes. The dependence of *J* upon the corresponding dihedral angle ξ is shown in Figure 5 for the bds containing dimer; the value ξ_0 corresponding to the idealized structure is also visualized, together with the experimental value J_{exp} . As expected, the maximum of the antiferromagnetic coupling is obtained when the two chalcogen-Ni-chalcogen units (the fragments that, in the monomer, bear the most important part of the spin) are oriented parallel $(J_{\rm min} = -430 \text{ cm}^{-1}).^{40}$ However, for a dihedral angle between 70° and 110°, the system is predicted to be ferromagnetic with $J_{\rm max}$ equal to ~60 cm⁻¹. It is worthwhile mentioning that similar results are obtained for the dimeric structure of CpNi(bdt); in this case; however, the minimum and maximum values of J are equal to -381 and +54 cm⁻¹, respectively.

Conclusion

Owing to ⁷⁷Se enrichment of CpNi(diselenolene), the *g* tensor as well as the ⁷⁷Se hyperfine tensors of these complexes could be measured from their frozen solution EPR spectra. They reasonably agree with those calculated by DFT. This accord corroborates the spin distribution found by DFT (~10% on each selenium and ~19% on the cyclopentadiene ring). This spin delocalization on the cyclopentadienyl rings is expected to give rise to unresolved proton couplings and explains, at least in part, the broadening of the EPR lines. The exchange coupling parameter *J* calculated for the idealized dimeric structure is consistent with the experimental value, but it also strongly depends upon the dihedral angle between the two diselenolene planes.

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Supporting Information Available: Variation of the exchange coupling parameter J_N and of the singlet and triplet energies with the SNiNiS dihedral angle for (bdt)CpNi···NiCp-(bdt). This material is available free of charge via the Internet at http://pubs.acs.org.

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