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Extended enantiopure *ortho*-phenylene ethylene (*o*-OPE)-based helical systems as scaffolds for supramolecular architectures: a study of chiroptical response and its connection to the CISS effect†

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A novel synthetic strategy based on a bifunctional stapled chiral nucleus from which segments of different lengths can be added to both ends of *o*-phenylene ethynyls (*o*-OPEs) has been developed to obtain a new type of foldamer and a novel chiral Pd₂L₂ metallacycle. For the first time, an enantiopure fully conjugated helical foldamer having 14 phenyl rings and 13 alkynes is reported. The folded structure has four complete loops and is able to host three Ag(I) cations in their cavity with high binding constants. The complete photophysical and chiroptical (ECD, CPL and VCD) characterization of these foldamers has shown that these molecules show intense chiroptical responses with dissymmetry ratios in the range of 10⁻². Theoretical modeling of these systems reveals the origin of these remarkable responses and points out a potential connection with the chiral induced spin selectivity (CISS) effect. The magnetic dipole moment is proposed as a key physical variable connecting the chiroptical properties and CISS-based spin filtering properties observed in chiral compounds.

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Introduction

Helical structures are widespread motifs in biomolecules and many efforts have been made to imitate such geometries using abiotic components.¹ Thus for example, [*n*]-helicenes have attracted the attention of chemists for almost a century.^{2,3} Particularly appealing is the fact that they present a conjugated system arranged in a helical geometry, which is preserved beyond [6]-helicenes. Their chiroptical properties, derived from their intrinsic chirality, such as optical rotatory dispersion (ORD), electronic circular dichroism (ECD), vibrational circular dichroism (VCD), Raman optical activity (ROA) and more recently, circularly polarized luminescence (CPL),^{4,5} are in fact remarkable, and have been extensively studied. Moreover, the helical geometry is also relevant in other fields. It has been suggested that conductive helical arrangements are of importance in the context of the chiral induced spin selectivity (CISS) effect⁶ once they can be prepared as enantiopure entities. The CISS effect is based on the phenomenon of electron spin polarization in electron transfer, electron transport and bond polarization processes in chiral molecules. Although initial experiments were based on biomacromolecules,⁷ it is

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worth noting that it has also been observed in fully conjugated [7]-helicene and in other helicene-like molecules.⁸ Consequently, fully conjugated helical molecules have been theoretically suggested as interesting candidates for spintronics, quantum sensing and quantum information applications.^{9–11}

Moreover, the remarkable geometry of [6]-helicenes has also been used in the context of chiral metallosupramolecular architectures^{12,13} as bidentate donors with an angle between the binding sites close to 0°. ¹⁴ These systems have emerged as an unique research field owing to their electronic, optical and sensing capabilities. However, the main drawback in [n]-helicene chemistry is the challenging synthesis of members with increasing turns, even in their racemic form. In fact, the longest member of the series, [16]-helicene, composed of two and a half turns, represents an isolated example (Fig. 1, top).¹⁵

Although longer related helicene-like molecules (two and a half turns) have been described, easy access to fully conjugated helical systems continues to be a synthetic challenge.^{16,17} Therefore, easy-to-prepare extended and conjugated helical systems would be highly desirable for important applications like those mentioned above. Within this context, foldamers based on conjugated subunits constitute a privileged scaffold for a number of different reasons: (i) they can be easily prepared by iterative protocols using simple conjugated monomers with different geometries, (ii) the functionality present in such monomers can cooperatively act to strengthen their properties or even create new ones, and (iii) they can also interact dynamically with the environment. In fact, many aromatic foldamers can be switched between different states by different stimuli such as the presence of light,^{18–22} changes in temperature,^{23–26} solvent composition,²⁷ anions,^{28–32} cations,^{33,34} or water binding, and redox processes.^{39,40} Moreover, foldamers are able to generate chirality in a dynamic way even using achiral monomers/repeating units. Therefore, the control of the *P/M* helicity^{41–46} yields systems that are able to act as chiroptical probes^{47,48} or even CPL emitters.^{49–53}

In recent years, we have been working on conjugated helical structures based on *o*-OPEs^{54–58} as an alternative to [n]-helicenes exhibiting, in many cases, better chiroptical responses. Our approach relies on widening the inner cavity present in [n]-helicenes, which translates into an enhancement in the magnetic moment of optical transitions and the corresponding chiroptical response, especially at the longest wavelength. Thus, we have developed two strategies to obtain enantiopure folded fully conjugated helical systems: the first based on the use of chiral sulfoxides^{59,60} and, secondly, the use of chiral staples, demonstrating that enantiopure *P/M* helices can be also prepared by efficient chirality transfer from a chiral staple to the *o*-OPE core, including even double staples to avoid racemization processes.^{61–64} However, the later systems could be synthesized with only one turn and a half.⁶⁵

All the above approaches provide structures with remarkable chiroptical responses in ECD, CPL and VCD. It is worth noting that the response of ECD in terms of molar circular dichroism has been for some time correlated with the magnitude of the chiral-induced spin selectivity (CISS) effect.⁶⁶



Fig. 1 Top: Previous studies with foldamers and helicene-type systems with more than two loops. Bottom: New helical systems developed in this work: chemical structure of stapled nucleus (S,S,P)-1, helical systems (S,S,P)-2–(S,S,P)-6, ligand (S,S,P)-7 and metallosupramolecular complex ((S,S,P)-7)₂Pd₂.

Actually, this phenomenon has been recently discovered, offering a promising change of paradigm for the discrimination and filtering of spin states. The spin of the electron, $S = \frac{1}{2}$, yields two quantum states characterized by its projection on



a given axis, $m_s = \pm\frac{1}{2}$. The two states are degenerate in the absence of either internal or external magnetic fields and, consequently, any electron current under such circumstances will consist of a half-half mixture of both states. The separation of the spin states of the electrons is a milestone of the greatest importance for the development of spintronics.

Although spin states of neutral particles⁶⁷ can be selected passing through an in-homogeneous magnetic field, charged particles, like electrons, suffer from the Lorentz force arising from the interaction of the electric charge with the magnetic field, preventing a Zeeman's effect-based technique to be used for separating electron spin states. Although more complex architectures can be used,⁶⁸ the (preferential) selection of the spin state of electrons travelling through chiral structures is highly appealing. The chemically relevant fact is that now it is the scattering of electrons through the molecular (helical) structure that induces the filtering of the electron spin states. Namely, depending on the helicity of the molecular structure, it preferentially allows the passage of one of the two spin states over the other, because helicity couples the spin of the electron with its own linear momentum, resulting in the selective transmission of electrons with the preferred spin state. This opens the way to great opportunities for "chemical control" in the selection and discrimination of the spin state of electric currents. The key concept resides in the achievement of precise chemical control on the factors affecting the molecular helical structure, *i.e.* length, diameter, pitch, chemical composition, chemical environment, either the solvent selection or its deposition on selected surfaces, including nanoparticles, *etc.* The absence of external electromagnetic fields avoids any interference with the spins in "uncontrollable" ways. The implications of the CISS effect for the control of the electronic spin state have already been documented.⁶⁹ Nevertheless, a long-standing goal, not fully resolved at the time being, is to know beforehand which molecules are best for CISS spin-filtering. Within this context, remarkable effort has been made by Waldeck *et al.*,⁶⁶ who showed conclusive experimental evidence for the direct correlation between CISS induced spin-filtering power and the magnitude of features of the molecule electronic circular dichroism (ECD) spectrum. A direct correlation of the dissymmetry of the electron-transfer rate for photoelectrons produced by circularly polarized light was found with both the sign and intensity of the ECD signal of electron acceptor chiral molecules. This suggests that the larger the ECD signal intensity the better the CISS spin-filtering a molecule can perform. The sign determines which spin is preferentially filtered.

Although the relationship between both phenomena seems plausible, it is not clear which factors are relevant for such a connection. One conceivable hypothesis is that the CISS effect may be associated with a transient magnetic moment in chiral molecules. Indeed, the intrinsic magnetic nature of the CISS effect suggests that induced magnetic transition moments (**m**) could be relevant to characterize the magnetic molecular response associated with the dynamic spin-polarized electron transport through a molecule. The optimal design of struc-

tures in which such a relationship could be validated is, therefore, essential.

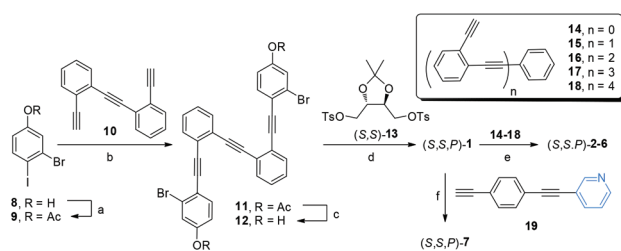
Here we report on a versatile scaffold to explore new helical architectures using simple derivatizations. Chiral stapled nucleus (*S,S,P*)-**1** can be used as a chiral seed for the synthesis of highly extended enantiopure conjugated helical systems (*S,S,P*)-**2**–(*S,S,P*)-**6** (Fig. 1, bottom) containing one up to four complete turns. In the presence of Ag(I) we have obtained the most extended enantiopure systems with a well-defined structure described to date.^{70–73} This situation is optimal to gain insight into the importance of the length of helical systems, the magnitude of the magnetic dipole transition moment (**m**) associated with the less energetic transition, and their potential use as a predictor of CISS capabilities for an organic architecture. We must emphasize that a comprehensive theoretical model coupling optical dichroism with the CISS effect is still missing, but the work reported here may be a step forward in this direction.

Moreover, the usefulness of (*S,S,P*)-**1** is also demonstrated with the preparation of ligand (*S,S,P*)-**7**. This compound increases the toolbox of the supramolecular research area showing that simple helical foldamers can be also used as a building block for the Pd(II)-directed self-assembly of a chiral Pd₂L₂ metallacycle, thus expanding the scarce number of examples of chiral metallosupramolecular architectures reported to date.¹³

Results and discussion

Synthesis

The main objective of our approach is the synthesis of a chiral core able to be further functionalized. Taking into account our previous successful route to enantiopure stapled *o*-OPEs based on an etherification reaction of a simple *o*-OPE and a chiral bis-tosylate derivative,⁵³ we decided to adopt this strategy to access compound (*S,S,P*)-**1**. Hence, this compound could be easily prepared following the route described in Scheme 1. (*S,S,P*)-**1** presents two aromatic bromides suitable for symmetric



Scheme 1 Synthesis of helical systems (*S,S,P*)-**2**–(*S,S,P*)-**6** and ligand (*S,S,P*)-**7**. Reagents and conditions: (a) Ac₂O (1.4 eq.), DMAP (2 eq.), CH₂Cl₂, rt, 5 min, 100%; (b) Pd(CH₃CN)₂Cl₂ (0.03 eq.), PtBu₃-HBF₄ (0.06 eq.), CuI (0.03 eq.), iPr₂NH, rt, 24 h, 70%; (c) K₂CO₃ (7 eq.), THF, MeOH, rt, 30 min, 88%; (d) CsCO₃, CH₃CN, reflux, 24 h, 56%; (e) Pd(CH₃CN)₂Cl₂, CuI, PtBu₃-HBF₄, iPr₂NH/THF, 60 °C, 24 h; (f) Pd(CH₃CN)₂Cl₂, CuI, PtBu₃-HBF₄, iPr₂NH/THF, 60 °C, 24 h; 81%. Yields of step e for each derivative are described in the ESI.†



extension of the conjugated system using different Sonogashira reactions. Enantiomeric (*R,R,M*)-**1** was also prepared resulting in the expected and opposite chiroptical properties (see Fig. S118 and S134†). For simplicity, from now on the discussion is restricted to (*S,S,P*)-**1** enantiomer. A simple coupling reaction affords enantiopure compounds (*S,S,P*)-**2** to (*S,S,P*)-**6** with increasing number of alkynes, enabling the study of systems from three alkynes/one turn, (*S,S,P*)-**1**, to thirteen alkynes, (*S,S,P*)-**6**, reaching in the latter case four complete turns, which has no precedent in the literature. The coupling reaction of the chiral nucleus (*S,S,P*)-**1** with pyridine-containing dialkyne **19** yielded (*S,S,P*)-**7**. These reactions demonstrate that (*S,S,P*)-**1** is in fact a versatile chiral building block that enables access to a variety of structures in a straightforward and modular way.

Photophysics of compounds (*S,S,P*)-**1** to (*S,S,P*)-**7**

Compound (*S,S,P*)-**1** was fully characterized by NMR and mass spectrometry (see the ESI†) but also by means of optical and chiroptical techniques. The absorption spectrum in CH₂Cl₂ exhibits two main bands centered at 305 nm and 290 nm with a smooth shoulder around 350 nm (Fig. 2a). On the other hand, the emission spectrum presents one band at 410 nm (Fig. S110†) with a fluorescence quantum yield of 4.6% (Table S1†).

Circular dichroism of (*S,S,P*)-**1** (CH₂Cl₂) showed a positive Cotton effect at the longest wavelength, $\Delta\epsilon = +47 \text{ M}^{-1} \text{ cm}^{-1}$, which is characteristic of *P* helicity (Fig. 2b). We note that molar circular dichroism ($\Delta\epsilon$) depends on the intensity of absorption (that is, on electric transition dipole moment) and its magnitude cannot be always correlated with the intrinsic chirality of the sample. In that situation, a dimensionless g_{abs} value, $\Delta\epsilon/\epsilon$, is more representative and easy to compare with theoretical values. Nevertheless, extracting reliable experimental g_{abs} values from ECD bands is tricky since calculations predict energetically close transitions with the opposite sign, such that the observed g_{abs} value results from the algebraic sum of g -values for the individual transitions. Being interested in the first electronic transition, the best estimate can be obtained from the tail of the CD band. In that case, the experimental g_{abs} value ($+1 \times 10^{-2}$) is comparable to previously described values for related systems.^{52,53}

The origin of this helical chirality could be clarified using density functional theory (DFT) calculations. After a molecular mechanics (MM) conformational search, all structures within 5 kcal mol⁻¹ have been optimized with both *P* and *M* helicities at the B3LYP/6-31G** level in CH₂Cl₂, treated with the polarizable continuum model (PCM). Only three conformers, corresponding to the relative spatial orientations of the bromine atoms (*out-out*, *out-in* and *in-in*), have been obtained with non-negligible populations (Fig. 2c and Table S10†). They all present *P* helicity and, considering the calculated energy, are nearly equally populated. As a consequence, the *P* epimer dominates the conformational equilibrium and makes possible a definite assignment of the (*S,S,P*)-**1** absolute configuration in solution. This conclusion was strengthened by the similarity

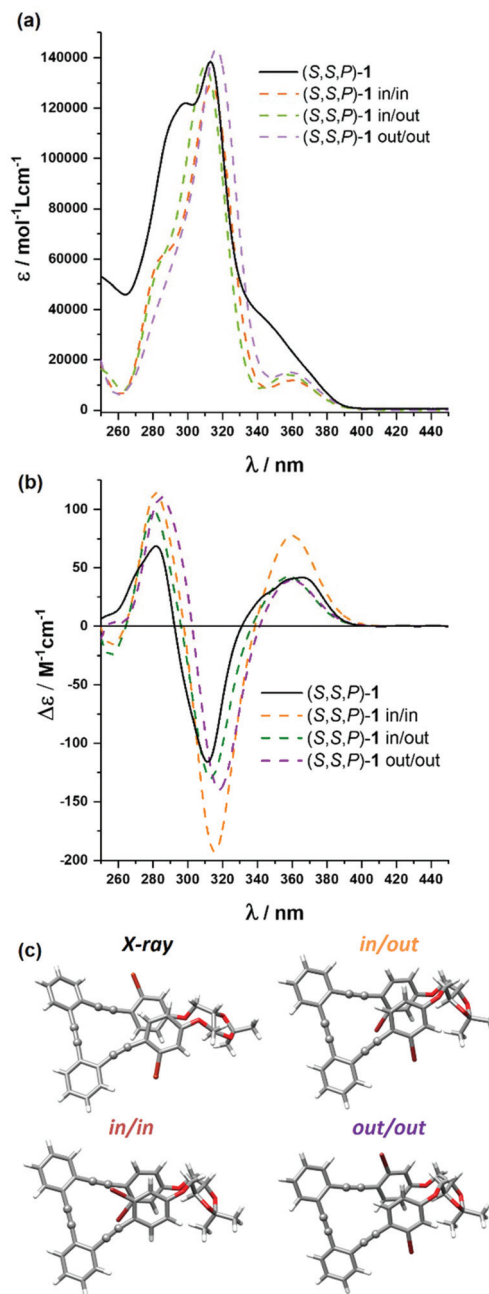


Fig. 2 Comparison of the calculated (dashed orange, purple and green) and experimental (black) (a) absorption and (b) ECD spectra of (*S,S,P*)-**1** in CH₂Cl₂ ($2.5 \times 10^{-5} \text{ M}$); (c) X-ray structure of (*S,S,P*)-**1** and calculated structures of the *in/in*, *in/out* and *out/out* conformers. Calculated spectra are 20 nm blue-shifted and their intensity divided by a factor of two.

between the averaged simulated ECD and absorption spectra of the abovementioned *P* conformers (M06/6-31G** level) and the experimental ones (Fig. 2a and b). The calculated g_{abs} value for the longest wavelength transition is in reasonable agreement with the experimental one.

Fortunately we could also obtain the single-crystal X-ray diffraction structure of (*S,S,P*)-**1**, which confirms its helical



geometry with *P* preference in the solid state (Fig. 2c). This helical arrangement generates a dihedral angle between the central aromatic rings close to 40°. The outer aromatic rings adopt a parallel orientation, although displaced, with a distance between their main planes of 3.5 Å and a distance between centroids of 3.8 Å, suggesting the establishment of π -stacking interactions between them.

Being dynamic systems, foldamers are affected by the environment and, consequently, their properties are usually solvent dependent. In this case, we only observed a single set of signals in NMR, showing that any conformational equilibrium takes place faster than the NMR timescale. The absorbance and fluorescence of (*S,S,P*)-1 were analyzed in different solvents, including apolar, polar and protic ones. Although no significant differences were observed in the emission spectra, a slightly increment of the shoulder in the absorbance spectra was observed in some solvents (Fig. S109†). Fluorescence lifetimes were also measured at the emission maximum using an excitation source of 325 nm. In all tested solvents, decays were fit to a biexponential function where the shortest fluorescence lifetimes were around 0.5 ns in most cases, except in CH₂Cl₂ in which it presented a considerably lower value of 0.2 ns. However, we observed higher differences for the longest fluorescence lifetimes, which varied between 0.7 ns in CH₂Cl₂ to 4.1 ns in toluene (see Table S2† for more details). In view of these results we recorded the time-resolved emission spectra (TRES) of compound (*S,S,P*)-1 in CH₂Cl₂ covering all of the emission range, between 340 and 600 nm. This global analysis also afforded a biexponential decay showing, along the whole spectra, an exclusive contribution of the shortest lifetime species, the presence of the largest being almost negligible (see Fig. S106†). Regarding chiroptical properties and according to the abovementioned data, minor changes in the ECD of (*S,S,P*)-1 were observed. We noticed that the chiroptical response of (*S,S,P*)-1 is maximized in MeOH with an experimental g_{abs} value of $+1.3 \times 10^{-2}$.

Since (*S,S,P*)-1 is fluorescent and chiral, CPL is a valuable tool to extract information of the dynamic system. Although the fluorescence was weak, reliable dissymmetry factors of the emission, g_{lum} values defined as $2(I_L - I_R)/(I_L + I_R)$, could be measured (CH₂Cl₂: $g_{\text{lum}} = +1.1 \times 10^{-2}$, MeOH: $g_{\text{lum}} = +1.0 \times 10^{-2}$, MeCN: $g_{\text{lum}} = +1.0 \times 10^{-2}$, *n*-hexane: $g_{\text{lum}} = +0.9 \times 10^{-2}$). These results suggest that the *P* helicity is preserved in the excited state. Consequently, the similarity of g_{abs} and g_{lum} values also reveals that the structure is highly preserved in the excited state despite the apparent conformational flexibility.

Analogous studies were carried out with compounds (*S,S*)-2 to (*S,S*)-7 (Fig. 3). Again, chiroptical properties, especially ECD and CPL, were highly informative and solvent dependent. The compound presenting five alkynes, (*S,S,P*)-2, showed remarkable molar circular dichroism in MeCN ($\Delta\epsilon = +41 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +1.0 \times 10^{-2}$), being compatible with a relatively ordered structure. Again, we found a correlation between the polarity of the solvent and the g_{abs} value for the longest wavelength transition, with similar values in MeOH ($\Delta\epsilon = +36 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +1.4 \times 10^{-2}$) to those obtained in MeCN, but being

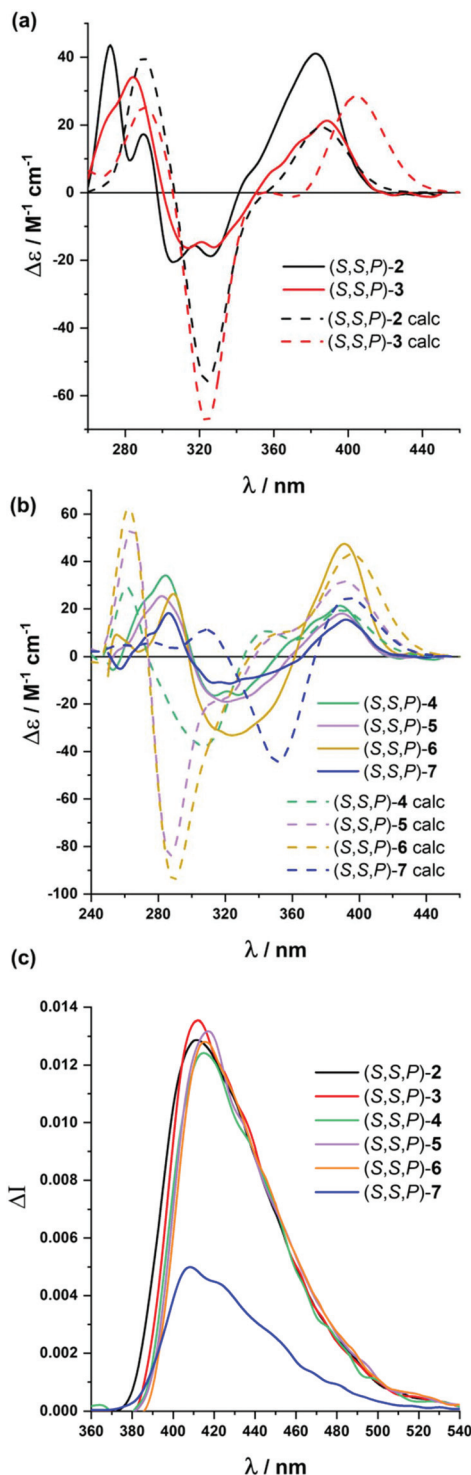


Fig. 3 Experimental (solid line) and calculated (dashed line) ECD spectra of (a) (*S,S,P*)-2 and (*S,S,P*)-3 and (b) (*S,S,P*)-4–(*S,S,P*)-7 in MeCN (calculated spectra are 20 nm shifted) and (c) CPL spectra of (*S,S,P*)-2–(*S,S,P*)-7 in MeOH.

weaker in CH₂Cl₂ ($\Delta\epsilon = +20 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +0.8 \times 10^{-2}$) and hexane ($\Delta\epsilon = +20 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +1 \times 10^{-2}$). If we assume g_{abs} values as the representative parameter of the system helicity,



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polar solvents are apparently favoring the folding process. Although this is a well-known effect for *m*-OPEs,²⁷ this is the first time it is clearly observed for *o*-OPEs.

Theoretical calculations support the prevalence of the fully folded conformations both considering CH₂Cl₂ and acetonitrile in the polarizable continuum model using the integral equation formalism variant (iefpcm) (see the ESI†), showing again a clear preference for the *P* conformation.⁷⁴ The rotational barrier for the unfolding process was theoretically estimated by turning one arm of the folded structure in 10⁰ steps (Fig. S156†), achieving a maximum at 8.1 kcal mol⁻¹. Therefore, the partially disordered situation favored by less polar solvents might explain the observed differences (see Fig. S119, in the ESI†).

Compound (*S,S,P*)-2 is also fluorescent, with quantum yield values varying from 19% to 29% depending on the solvent (see Table S3 in the ESI†). The fluorescence lifetime at the emission maximum showed again two values around 4 and 2 ns in all solvents. Interestingly, time-resolved fluorescence allows the deconvolution of the emission spectra, providing the ratio of the different emitting species by using the areas of the species-associated emission spectra (SAEMS). In the four solvents we could detect two main contributions coming from two different species with around 4.6 and 2.6 ns, which cannot interconvert in that timescale. Although relative proportion depends on the solvent nature (see Fig. S107†), two limit situations are present in hexane and MeOH, in which the corresponding populations are reversed. The coexistence of fully folded and partially folded structures in the excited state may be a reasonable explanation, fully folded structures being maximized in MeOH.

Since the chiroptical properties of the abovementioned species are expected to be different, CPL is once again a valuable tool to extract information on the dynamic system. Representative solvents as MeOH, MeCN, CH₂Cl₂ and hexane were evaluated using this technique, obtaining remarkably high *g*_{lum} values (MeOH: +1.3 × 10⁻², MeCN: +1.3 × 10⁻², CH₂Cl₂: +0.88 × 10⁻², hexane: +1 × 10⁻²). Once more, a correlation with the polarity of the solvent was observed, the fully folded conformations with higher *g*_{lum} values being preferred in MeOH or MeCN. The positive sign of CPL is correlated with the positive sign of the ECD signal for the longest wavelength transition, *P* helicity being preserved in the excited state.⁴ The similarity of *g*_{abs} and *g*_{lum} values also reveals that the geometrical structure of the ground state is preserved in the excited state. In fact, the optimized structure for the first excited state starting from the lowest-energy fully folded ground state structure resulted in a nearly superimposable helix in both ground and excited states, showing differences only in the bond length alteration (BLA) of the C–C≡C–C groups (see Fig. 4f and Table S12†).

Similar conclusions were extracted for (*S,S,P*)-3–(*S,S,P*)-5 and (*S,S,P*)-7. Molar circular dichroism and *g*_{abs} values remain high, especially in MeCN ((*S,S,P*)-3: Δε = +21 M⁻¹ cm⁻¹, *g*_{abs} = +0.9 × 10⁻²; (*S,S,P*)-4: Δε = +18 M⁻¹ cm⁻¹, *g*_{abs} = +0.5 × 10⁻²; (*S,S,P*)-5: Δε = +47 M⁻¹ cm⁻¹, *g*_{abs} = +0.9 × 10⁻²; (*S,S,P*)-7: Δε =



Fig. 4 Optimized DFT (B3LYP/6-31G**) structures of: (a) (*S,S,P*)-2; (b) (*S,S,P*)-3; (c) (*S,S,P*)-4; (d) (*S,S,P*)-5; (e) (*S,S,P*)-6. Color coding: C, gray; O, red. H atoms have been omitted for clarity; (f) ground-excited structures and transitions for (*S,S,P*)-2 in absorption (left) and emission (right).

+19 M⁻¹ cm⁻¹, *g*_{abs} = +0.8 × 10⁻²), compared to CH₂Cl₂ ((*S,S,P*)-3: Δε = +6 M⁻¹ cm⁻¹, *g*_{abs} = +0.3 × 10⁻²; (*S,S,P*)-4: Δε = +8 M⁻¹ cm⁻¹, *g*_{abs} = +0.3 × 10⁻²; (*S,S,P*)-5: Δε = +13 M⁻¹ cm⁻¹, *g*_{abs} = +0.3 × 10⁻²; (*S,S,P*)-7: Δε = +19 M⁻¹ cm⁻¹, *g*_{abs} = +0.3 × 10⁻²), although exceptions can be also observed (see the ESI†). These higher *g*_{abs} in MeCN suggest a relatively ordered structure in more polar solvents, in accordance with the stability of the helical structure. Nevertheless, the increasing number of potential conformations did not allow a complete modeling of the conformational space as in previous cases. Even so, it seems reasonable to assume that in the absence of solvophobic interactions many more disordered structures in solution could be present. For compound (*S,S,P*)-6, we found some solubility issues in hexane and MeOH and reliable CD spectra could be obtained only in CH₂Cl₂ (Δε = +13 M⁻¹ cm⁻¹, *g*_{abs} = +0.3 × 10⁻²) and MeCN (Δε = +15 M⁻¹ cm⁻¹, *g*_{abs} = +0.3 × 10⁻²).

Fluorescence lifetimes and quantum yields for such longer oligomers (*S,S,P*)-3–(*S,S,P*)-7 were evaluated in MeOH, MeCN, CH₂Cl₂ and hexane and are summarized in Table S4.† Quantum yields depend on both the solvent and number of alkynes present in the molecule, but no sensible relationships



could be obtained in any of the cases. However, it is remarkable that, except for compound (S,S,P) -3, the intensity-averaged lifetime of longer oligomers is always lower in hexane (around 3 ns) than in MeOH and CH_3CN (around 4 ns), showing again a higher contribution of the shortest lifetime species in apolar solvents, in accordance with the results obtained for TRES of the first member of the series (S,S,P) -2.

All helical systems were also very efficient CPL emitters, especially in MeOH ((S,S,P) -3, $g_{\text{lum}} = +1.3 \times 10^{-2}$, (S,S,P) -4, $g_{\text{lum}} = +1.1 \times 10^{-2}$, (S,S,P) -5, $g_{\text{lum}} = +1.3 \times 10^{-2}$, (S,S,P) -6, $g_{\text{lum}} = +1.3 \times 10^{-2}$, (S,S,P) -7, $g_{\text{lum}} = +0.54 \times 10^{-2}$). Modeling CPL properties has been carried out for compound (S,S,P) -2 in the most stable folded conformer. However, the analysis of larger systems is computationally too demanding. Despite this limitation, the structures were optimized by DFT calculations (Fig. 4b–e). Although, the similarity between g_{abs} and g_{lum} values, which is also observed in experimental trends, is expected, the g_{lum} value is controlled by the $S_1 \rightarrow S_0$ transition, which cannot be assumed to be identical to the $S_0 \rightarrow S_1$ transition because the structural changes in the excited state may affect the ground state transition parameters.

(S,S,P) -1– (S,S,P) -7 Ag(I) complexes

Stapled *o*-OPEs present a perfect arrangement for the coordination of Ag(I) cations (AgBF_4) using carbophilic interactions. In this case, we restricted our study to CH_2Cl_2 owing to the incompatibility of such interactions and polar coordinating solvents.

The first member of the series, (S,S,P) -1, presents evident structural changes after Ag(I) addition as we observed an increased resolution and downfield shielding of the aromatic protons (Fig. S49†). Regarding the ^{13}C NMR spectra, alkyne carbons, which were initially located within 1 ppm to around 92 ppm, evolve to three different signals at 89, 92 and 94 ppm by coordination with the Ag(I) cation (Fig. S50†). The coordination process was also followed by ECD spectroscopy (Fig. S126†). In this particular case, we observed a decrease in the chiroptical response. This fact is in agreement with previous results in related *o*-OPEs with three alkynes owing to a planarization of the helical structure. ECD titration experiments resulted in a relatively strong single binding event, $K_{1:\text{Ag}} = 32\,480 \pm 2\% \text{ M}^{-1}$. Theoretical calculations (M06/6-31G* plus LANL2DZ (valence + ECP) for silver in CH_2Cl_2) showed that the minimum energy structure corresponds to a distorted trigonal bipyramidal geometry (Fig. 6b) in which the bromine atoms participate in the coordination to the Ag(I) cation.

Compound (S,S,P) -2 is expected to behave as a ligand for only one Ag(I) cation taking into account the previous studies of related *o*-OPE systems.⁶¹ The binding process can be again properly followed by ^1H NMR spectroscopy. NMR titrations showed that the original signals of *ortho* hydrogen atoms of the inner phenyl rings experience the characteristic deshielding (*i.e.* from 6.45 ppm to 6.64 ppm, Fig. 6a) attributed to Ag(I) coordination. In addition, significant changes in the chemical shift of alkyne carbons were observed in the ^{13}C NMR spectrum (Fig. S52†). The ECD response at the longest wavelength

is intense ($\Delta\epsilon = +85 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +2.5 \times 10^{-2}$) in agreement with a well-defined helical structure upon the formation of the complex (Fig. 6). It is worth noting that such a dissymmetry ratio is huge compared with usual values for organic molecules at the longest wavelength. The corresponding ECD titration showed a high binding constant $K_{2:\text{Ag}} = 3.67 \times 10^5 \pm 5\% \text{ M}^{-1}$. As expected, the DFT-based conformational analysis (see the ESI†) of complex (S,S,P) -2:Ag is in agreement with experimental findings, showing a stabilization of the fully folded structure. The calculated ECD is also in good agreement with the experimental one, thus confirming the folding process with Ag(I) (Fig. 5a and 6c). In this case, the distorted trigonal bipyramidal geometry is generated by coordination with five alkynes. Ag–alkyne bond distances are predicted to lay within 2.51–2.69 Å, with the Ag–C (alkyne) distances ranging from 2.57 to 2.76 Å. The distance between the terminal aromatic rings of the side arms and those on the corners on the central core of the helix, which display a quite displaced parallel arrangement, is *ca.* 3.5–3.6 Å, which suggests some extent of π -interactions between them.

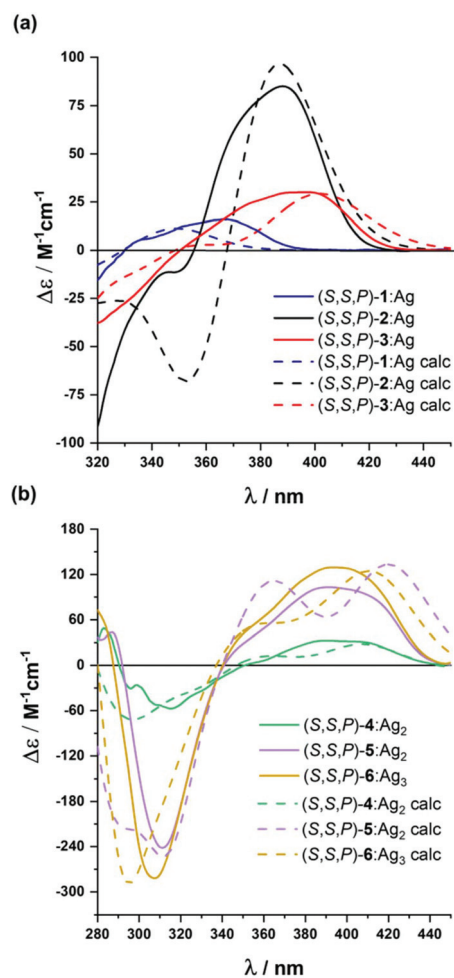


Fig. 5 Experimental (solid line) and calculated (dashed line) ECD spectra of: (a) (S,S,P) -2– (S,S,P) -3 and (b) (S,S,P) -4– (S,S,P) -6 in CH_2Cl_2 in the presence of an excess of Ag(I).





Fig. 6 (a) ^1H NMR spectra (400 MHz, CD_2Cl_2) of compound (*S,S,P*)-2 in the presence of 0, 1, 2, 3 and 4 equivalents of AgBF_4 . (b) Calculated structures (M06/6-31G* plus LANL2DZ for silver) in CH_2Cl_2 for: (b) (*S,S,P*)-1:Ag; (c) (*S,S,P*)-2:Ag; (d) (*S,S,P*)-3:Ag; (e) (*S,S,P*)-4:Ag₂; (f) (*S,S,P*)-5:Ag₂; (g) (*S,S,P*)-6:Ag₃. Color coding: C, gray; O, red; Br, wine; H, white; Ag, light blue. H atoms have been omitted for clarity.

On the other hand, the increase to seven potential binding sites for $\text{Ag}(\text{i})$ cations in compound (*S,S,P*)-3 did not result in a significant increase of the chiroptical properties ($\Delta\epsilon = +21 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +1 \times 10^{-2}$). In principle, a single $\text{Ag}(\text{i})$ cation does not need all the alkynes in the binding event, making the remaining ones useless to stabilize a second $\text{Ag}(\text{i})$ cation. In this situation the presence of loosely or non-bound side arms can result in a more disordered structure compared to (*S,S,P*)-2, even in the presence of $\text{Ag}(\text{i})$. Unfortunately, the complex conformational space makes it difficult to perform any theoretical prediction. As a consequence of the previous assumption,

ECD titrations showed that the compound is only bound to one $\text{Ag}(\text{i})$ cation with a binding constant $K_{3:\text{Ag}} = 3.7 \times 10^6 \pm 15\% \text{ M}^{-1}$. Minimum energy calculated structures showed that $\text{Ag}(\text{i})$ interacts with five alkynes. Only two alkynes remain free to bind a new cation. Our previous studies⁶⁰ indicated that two alkynes are unable to provide an efficient coordination to another $\text{Ag}(\text{i})$ cation. The different spatial arrangement for these two additional unbound side chains and even in-cavity $\text{Ag}(\text{i})$ displacements could explain the weaker ECD response.

The presence of two additional alkynes in (*S,S,P*)-4 should be in principle beneficial for the coordination process. Nevertheless, the ECD response remains weak and similar to the previous case ($\Delta\epsilon = +24 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +0.6 \times 10^{-2}$). This fact suggest that at the concentrations used for the ECD experiments there is only a relevant binding constant, although subsequent $\text{Ag}(\text{i})$ coordination processes can take place with a modest binding constant. Accordingly, in the titration experiments a higher overall binding constant, $K_{4:\text{Ag}_2} = 2.4 \times 10^{10} \pm 12\% \text{ M}^{-2}$, was obtained (see Fig. S129–S131†). It is worth noting that the corresponding NMR studies were carried out at much higher concentrations. Under such experimental conditions more than two equivalents of the $\text{Ag}(\text{i})$ cation are required to fully resolve the signals, suggesting the existence of a symmetric dimetallated structure.

Compound (*S,S,P*)-5, presenting eleven alkynes, is able to accept two $\text{Ag}(\text{i})$ cations, thus yielding a longer helical structure. The first evidence came from the ECD spectra, showing a remarkable molar circular dichroism ($\Delta\epsilon$) at the longest wavelength ($\Delta\epsilon = +85 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +1.7 \times 10^{-2}$) in the presence of an excess of $\text{Ag}(\text{i})$. ECD titrations also showed strong coordination of two $\text{Ag}(\text{i})$ cations ($K_{5:\text{Ag}_2} = 18 \times 10^9 \pm 1.6\% \text{ M}^{-2}$). This is the first example of a pure and well-defined *o*-OPE system presenting an enantiopure helical structure with three and a half turns. This fact shows that the chiral nucleus can be useful with the aid of $\text{Ag}(\text{i})$ cations in the preparation of long and conjugated helical systems. Although different situations are possible, DFT calculations showed that the energetically favored case is that in which the $\text{Ag}(\text{i})$ cations are separated by a distance of approximately 5.2 Å (Fig. 6f).

The best chiroptical response was observed for compound (*S,S,P*)-6, which showed an even higher molar circular dichroism ($\Delta\epsilon = +130 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +2.2 \times 10^{-2}$). Again, this result strongly suggests a fully folded helical structure. ECD titrations also showed three strong binding events with three $\text{Ag}(\text{i})$ cations ($K_{6:\text{Ag}_3} = 2.7 \times 10^{17} \pm 19\% \text{ M}^{-3}$). As it can be seen in Fig. 6g, DFT calculations predict a highly symmetric structure with four complete loops. The interatomic distances between consecutive $\text{Ag}(\text{i})$ cations are 4.4 and 4.5 Å. Interestingly, on the basis of the Ag –alkyne distances, calculations predict that the metal centers are coordinated to five alkynes. As a result, there are two alkynes that coordinate simultaneously to two $\text{Ag}(\text{i})$ centers, which results in longer Ag –alkyne distances (2.74–2.89 Å) in comparison with those involving triple bonds coordinated just to one metal center (Ag –alkyne distances: 2.44–2.65 Å).



VCD studies

The existence of the helical structures was also confirmed by VCD spectroscopy, particularly evident when considering longer oligomers (see the ESI†). Concerning IR measurements, the presence of the Ag(I) complex results in a band shift and broadening of the feature observed at 1500 cm^{-1} and in the appearance of a band at 1315 cm^{-1} (Fig. 7). In VCD an intensification of the triplet signal at 1445 , 1482 and 1493 cm^{-1} (+, -, +) is also characteristic of the presence of interacting Ag(I) cations, particularly considering (S,S,P)-4 to 6 compounds. This hints at the possibility of designing good silver switches based on IR range measurements. This (+, -, +) triplet shows similar intensity in short and long chains in the absence of silver, suggesting that the central portion of the backbone rigidified by the staple is the one responsible for this signal in all compounds.

To better understand the origin of the spectroscopic changes upon silver addition, we examined two representative cases with the aid of DFT calculations: the shortest oligomer (S,S,P)-2 and a longer case, (S,S,P)-5. For the first case, we considered the possible conformers and compared the calculated spectra for the most stable structures in the absence or presence of silver. The details of this analysis are given in the ESI (conformer characteristics and spectra).† Due to the staple, the structure of the short oligomer is rigid enough to present a VCD signal typical of the ordered OPE structures already observed.^{60,61} In particular, the configuration of the staple dictates the prevalent helicity sense and the most stable conformer presents already the correct shape to host a silver ion; however, also partially unfolded structures of (S,S,P)-2 maintain the spectroscopic pattern. In the presence of Ag(I), the calculated spectrum reproduces the wavenumber shift of the two higher energy components of the previously cited triplet (+, -, +) and the 1315 cm^{-1} feature intensification (Fig. 7). Also the IR spectra differences are correctly predicted.

In the case of the longer oligomer, Ag(I) cations give rise to the enhancement of the triplet at 1450 – 1500 cm^{-1} . The phenomenon can be attributed to normal modes delocalized along the whole molecule, consisting of in-plane bending of the phenyl CH bonds. Structures with various Ag contents have been optimized: one central Ag ion, two Ag ions symme-

trically or non-symmetrically disposed and three Ag ions (see Fig. S150† for the optimized structures and Fig. S151† for the corresponding calculated spectra). Overall, there is a good correspondence between calculated and experimental data. In fact, compound (S,S,P)-5 revealed to be quite efficient in incorporating silver ions and has sufficient length to show an enhancement of VCD signals when it is guided to form a regular helix in the presence of silver. In conclusion, in the presence of silver, the longer backbone assumes an ordered structure able to show intensification of the triplet; the increase in intensity is more regular considering VCD, while ECD shows a non-monotonic dependence on the alkyne number. A complete theoretical representation of all the situations and of the two spectroscopic responses (ECD and VCD) is not easy due to the complexity of the conformational landscape and of the evident solvent effect (see Fig. 2) which is not simple to model.

Chiroptical properties and CISS effect for compounds (S,S,P)-2 to (S,S,P)-6 and their corresponding Ag(I)-complexes

In this vein, compounds denoted as (S,S,P)-2–6, appear to be ideal for examining the influence of the number of turns of helical molecules on their chiroptical properties and its eventual connection to the CISS effect-based spin selection power. The chiroptical properties are customarily quantified by the dissymmetry factor, g_{abs} , of the electronic transitions. The latter quantity is expressed in terms of the extinction coefficients for left-, ϵ_- , and right-polarized, ϵ_+ , incident exciting light and reads as, $g_{\text{abs}} = 2(\epsilon_- - \epsilon_+)/(\epsilon_- + \epsilon_+) = 4(|\boldsymbol{\mu}||\mathbf{m}|)\cos(\theta)/(|\boldsymbol{\mu}|^2 + |\mathbf{m}|^2) \approx 4R/D$, where R and D are the rotational and dipole strengths respectively for the corresponding $S_0 \rightarrow S_n$ transition. This equation expresses the dissymmetry factor in terms of the electric dipole transition moment, $\boldsymbol{\mu}$, and the magnetic dipole transition moment, \mathbf{m} , and the angle subtended by these two vectors, θ . Since the magnetic dipole transition moment is normally much smaller than the corresponding electric one, one can expand $(|\boldsymbol{\mu}|^2 + |\mathbf{m}|^2)^{-1}$ in terms of powers of $(|\mathbf{m}|/|\boldsymbol{\mu}|)$ to obtain, at the first order approximation, $g_{\text{abs}} = 4(|\mathbf{m}|/|\boldsymbol{\mu}|)\cos(\theta)$. On the other hand, the CISS effect translates into an additional component of the magnetic moment associated with a Rashba-like term: $m_{\text{CISS}} = \alpha(\mathbf{E} \cdot (\boldsymbol{\sigma} \times \mathbf{p}))$, where α is a coupling constant, \mathbf{E} the molecular electric field, $\boldsymbol{\sigma}$ the Pauli matrix vector and \mathbf{p} the linear momentum of the electron involved in the transport. This additional term has to be included in the optical response, generating, in principle, a direct connection between ECD and the CISS effect. Namely, as noted above, an electron moving along the electric field in a chiral structure defined by the fixed nuclei and not by an external field, will experience in its rest frame a magnetic field, which will interact equally as above with the electron magnetic moment. Hence, the electron magnetic moment appears to be directly connected to the CISS effect. Given the nature of the optical activity response, one could hypothesize the transition magnetic vector \mathbf{m} as a descriptor for enhanced chiroptical properties, which could explain the observed relationship between the chiroptical properties and enhanced

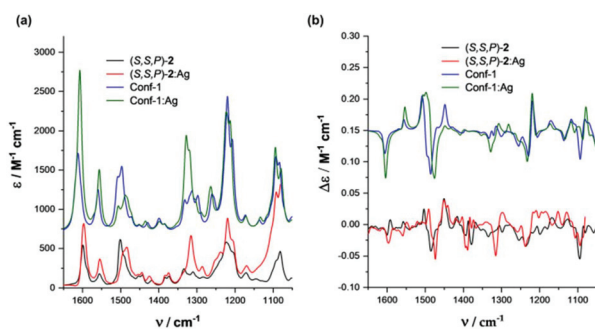


Fig. 7 Comparison of theoretical and experimental (a) IR and (b) VCD spectra of compound (S,S,P)-2 in CH_2Cl_2 solvent.



CISS spin selectivity power.⁷⁵ Despite the fact that g_{abs} values can be high when $|\mathbf{m}|$ and $|\boldsymbol{\mu}|$ are similar in magnitude and aligned as possible, that fact does not guarantee the absolute value of $|\mathbf{m}|$ to be high. That is, compounds can present high g_{abs} values simply by diminishing $|\boldsymbol{\mu}|$. We hypothesize that conductive helical structures, which supposedly are ideal candidates for relevant CISS the effect, must show a high $|\mathbf{m}|$, higher than 1×10^{-20} cgs units,⁷⁶ for transitions capturing the helical structure of the involved molecular orbitals. Moreover, the CISS effect is directional and must be analyzed in the direction of electron transport, which is difficult using scalar quantities like g_{abs} or rotational strength ($R = (|\boldsymbol{\mu}||\mathbf{m}|\cos(\theta))$). The lower energy transitions are of special relevance since the most accessible orbitals for electron transport at low voltages are involved.

We have analyzed the relevant chiroptical parameters, such as electric and magnetic dipole transition moments $|\mathbf{m}|$ and $|\boldsymbol{\mu}|$, dipole (D) and rotational strengths (R), and angle between electric dipole transition moment and magnetic dipole transition moment, obtained by TD-DFT calculations (M06/6-31G**) of molecules (S,S,P)-2 to (S,S,P)-6 and their corresponding Ag(I) complexes. It is important to note that the studied systems are quite large and cannot be fully modelled at a high level of theory. Consequently, only tendencies and not precise values can be provided by the calculations. It is worth noting that theoretical calculations overestimate the predicted g_{abs} values.⁷⁷ In this case, we use CH_2Cl_2 as the solvent for a better comparison with the parent compounds and Ag(I) complexes derived.

Inspection of the data of Table 1 shows that the magnetic dipole transition moment for the $S_0 \rightarrow S_1$ transition increases monotonically with the number of windings for the whole set of helical molecules (S,S,P)-2 to (S,S,P)-6. This is in agreement with the naïve view that holds that increasing the helicity of the compound should translate into an increase of the magnetic moment. It had been suggested that when the magnetic dipole transition moment stands for the power absorbed from the exciting light magnetic field in a conductive helix, a linear

relationship between the transition magnetic dipole moment and the number of loops n exists.⁷⁸

In that case, the classical electromagnetic power dissipated by light absorption of intensity E_0^2 and wavelength λ by the magnetic field alone in a helix of conductivity σ , cross section a , radius r , pitch length d , and pitch angle Φ , is dictated by $P = a\sigma(E_0 r \sin \Phi/4\lambda)^2(d^2 + 4\pi^2 r^2)^{1/2}n$. Such a linear relationship is nicely paralleled by the plot in Fig. 8 of the magnetic dipole transition moment $|\mathbf{m}|$ versus N , where N , the number of alkynes displayed on the x -axis, is related to the number of loops (n) through $n = N/3$. This is in agreement with a seminal paper about the CISS effect by Zacharias *et al.*⁷⁹ in which the spin polarization increases as a function of the number of turns in helical polypeptide molecules adsorbed on a gold surface, a result that runs parallel with the observed dependence of the transition magnetic moment.

This nice result is based on the alignment of the magnetic transition vector and the helix axis for the $S_0 \rightarrow S_1$ transition. At this point we hypothesized that the projection of the magnetic transition vector on the helix axis could be even a better predictor owing to the fact that spin filtering implies some sort of directionality, implicit in the idea that the scattered electron should pass through the helix. In this way, transitions with magnetic transition vectors misaligned with respect to the direction of the electron transport (helix axis) would not be relevant for the desired relationship between CISS and chiroptical properties. Although the helix axis is not fully conserved in all the structures (see Fig. S157 and S158†) we could observe again an almost perfect alignment between \mathbf{m} and the defined helix axis. Consequently, the linear relationship is preserved (Fig. S159†).

A similar analysis can be carried out with the Ag(I) complexes, although in this case the geometrical variations along the series are higher resulting in a worse linearity (see Fig. 8, pink circles). In any case, a clear increase in the magnetic

Table 1 Electric and magnetic dipole transition moments $\boldsymbol{\mu}$ and \mathbf{m} , dipole (D) and rotational strengths (R), angle between the electric dipole transition moment and the magnetic dipole transition moment $E-M$ and dissymmetry ratio g_{abs} obtained by TD-DFT calculations (m06/6-31g**) for the first electronic transition. The unit for electric and magnetic dipole transition moments is esu cm and that for D and R is esu² cm²

Compound	$\boldsymbol{\mu}$ (10 ⁻²⁰)	\mathbf{m} (10 ⁻²⁰)	D (10 ⁻⁴⁰) length	R (10 ⁻⁴⁰) length	$E-M$	g_{abs}
(S,S,P)-2	204	3.74	41 568	645	32	0.062
(S,S,P)-3	382	3.97	146 279	1407	22	0.03
(S,S,P)-4	426	4.19	181 829	1606	26	0.03
(S,S,P)-5	442	4.20	195 208	1727	22	0.03
(S,S,P)-6	524	4.42	275 017	2237	15	0.03
(S,S,P)-2:Ag	215	3.73	46 231	705	29	0.06
(S,S,P)-3:Ag	249	3.45	62 283	811	20	0.05
(S,S,P)-4:Ag ₂	216	3.84	46 991	815	12	0.06
(S,S,P)-5:Ag ₂	185	3.94	34 572	734	0	0.08
(S,S,P)-6:Ag ₃	289	4.28	83 865	1190	17	0.05

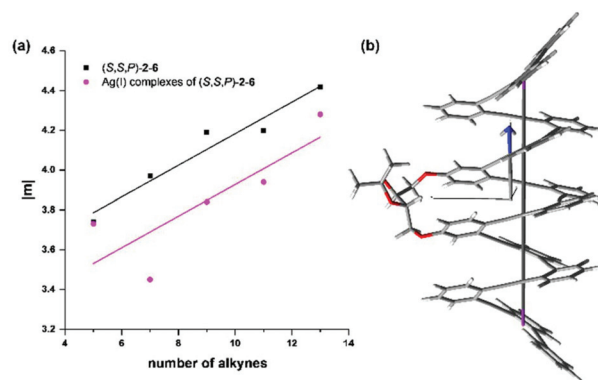


Fig. 8 (a) Linear fitting of $|\mathbf{m}|$ with the number of alkynes of compounds (S,S,P)-2–6 in CH_2Cl_2 solutions in the absence of Ag(I) (black squares, $y = 3.38 + 0.8x$, $R^2 = 0.955$) and silver complexes (S,S,P)-2:Ag, (S,S,P)-3:Ag, (S,S,P)-4:Ag₂, (S,S,P)-5:Ag₂ and (S,S,P)-6:Ag₃ (pink circles, $y = 3.13 + 0.08x$, $R^2 = 0.69$). (b) Calculated structure including helix axis and magnetic dipole transition moment of (S,S,P)-6 (for the rest of compounds see Fig. S157 and S158†).



moment of the transition (**m**) can be observed for the Ag(I)-saturated structures (*S,S,P*)-2:Ag, (*S,S,P*)-3:Ag, (*S,S,P*)-4:Ag₂, (*S,S,P*)-5:Ag₂, (*S,S,P*)-6:Ag₃.

As suggested above we speculate that the magnetic transition vector or its projection in the electron transport direction of a selected transition could be the missing link between the CISS effect and chiroptical properties. This can be supported by the interpretation of the CISS effect as an additional contribution to molecular magnetic response that is triggered in the presence of electron transport through a chiral (helical) structure. A simple theoretical explanation for this dependence of the spin polarization with molecular length in helices can be given in terms of a scattering model including both coherent and incoherent contributions to electron transport.⁸⁰ All these assumptions, although promising, require further theoretical inquiries.

Metallosupramolecular complex ((*S,S,P*)-7)₂Pd₂

Finally, we have checked the coordination capabilities of homochiral ligand (*S,S,P*)-7 towards Ag(I) and Pd(II) cations. Despite the presence of five alkynes, this ligand has two additional pyridine rings that can coordinate to non-carbophilic metal centres as Pd(II). The *meta* substitution of these pyridine rings results in an almost parallel orientation of the binding sites (with an angle between them close to 0°). This geometry allows the assembly of well-defined metallosupramolecular entities with metal centres with coordination sites in a 180° orientation, resulting in either linear Ag(I) complexes or square planar Pd(II) complexes with two additional ligands in *trans* positions.¹²

In the case of the Ag(I) cations, two different binding sites are possible, the alkynes and the pyridines. ECD spectroscopy was highly informative to discriminate between the two alternatives. As in the case of related compound (*S,S,P*)-3, intensification of the ECD signal at the longest wavelength ($\Delta\epsilon = +41 \text{ M}^{-1} \text{ cm}^{-1}$, $g_{\text{abs}} = +0.45 \times 10^{-2}$) is observed upon the addition of a Ag(I) salt to a solution $2.5 \times 10^{-5} \text{ M}$ of (*S,S,P*)-7, compatible with an increase of the helicity of the system (Fig. S121†). Moreover, titration experiments also showed that this increase was related to the coordination of the ligand to one Ag(I) cation ($K_{7:\text{Ag}} = 8.21 \times 10^5 \pm 13\% \text{ M}^{-1}$) (Fig. S132†). Therefore, the interaction of Ag(I) cations with the alkynes seems to be the most favorable interaction for the first added equivalent. Nevertheless, in the presence of an excess of Ag(I) cation the ECD signals drastically decrease suggesting the formation of insoluble material in CH₂Cl₂, probably by subsequent coordination of the cation with the available pyridine groups. All these results allow us to conclude that carbophilic metals cannot easily direct the self-assembly of metallosupramolecular structures for our systems.

On the other hand, the addition of a solution of *trans*-PdCl₂(MeCN)₂ (10 mM) to solutions of (*S,S,P*)-7 in CD₂Cl₂ or DMSO-*d*₆ (10 mM) resulted in the appearance of new signals in the corresponding ¹H NMR spectra. This fact indicates the presence of new species in solution and suggests the formation of a metallacycle as a result of the coordination of the

pyridine rings to the Pd(II) centers (see ESI, Fig. S92–102†). The DOSY NMR spectrum in DMSO-*d*₆ of this mixture (Fig. S103†) agrees with the presence of two species as two diffusion coefficients are observed for the ¹H signals. Thus, the new set of signals that appeared after Pd(II) addition shows a lower diffusion coefficient, supporting the coordination of (*S,S,P*)-7 to the metal center and the self-assembly of the larger metallacycle. By increasing the amount of the added Pd(II) complex, this new set of signals became the main one in the mixture, being practically the only one present when 1–1.2 equiv. of *trans*-PdCl₂(MeCN)₂ were added (Fig. S93 and S98†). In this case, the DOSY NMR spectrum confirms that all signals correspond to a single species that diffuses as a whole, as expected, as only one diffusion coefficient is observed (Fig. S65 and S70†).

The clear solution obtained in DMSO-*d*₆ was monitored over time by ¹H NMR spectroscopy, showing no significant changes in the spectrum over 6 days (see ESI, Fig. S102†). In CD₂Cl₂ the evolution of the system was more complicated since a precipitate was formed soon after the addition of the metal when a final concentration of 5 mM was used. After filtration, the ¹H NMR spectrum of the remaining solution showed two sets of signals, one corresponding to the free ligand and the other one to the newly formed species, postulated as the ((*S,S,P*)-7)₂Pd₂ complex. However, the ratio between species was not maintained over time, as a decrease of the proportion of the coordination complex was observed by ¹H NMR monitoring. Better results were obtained when the concentration was lowered to 2.5 mM, as no precipitation was observed. In this case, the set of signals attributed to the coordinated species was present almost exclusively after 24 h and good stability was observed over time (Fig. S96†).

Comparison of the ¹H NMR of (*S,S,P*)-7 before and after the addition of Pd(II) both in DMSO-*d*₆ and CD₂Cl₂ shows a downfield shift ($\Delta\delta_{\text{Ha}} = 0.23 \text{ ppm}$; $\Delta\delta_{\text{Hb}} = 0.32 \text{ ppm}$ in CD₂Cl₂ and $\Delta\delta_{\text{Ha}} = 0.22 \text{ ppm}$; $\Delta\delta_{\text{Hb}} = 0.23 \text{ ppm}$ in DMSO-*d*₆) of the H atoms of the pyridine ring at positions 2 and 6 (Fig. 9a). This is the expected behavior for the coordination of the pyridine ring to the Pd(II) center. To lend further support to the fact that the observed shift corresponds to the binding event, 3-(phenylethynyl)pyridine was prepared as a model ligand. The similarities between the chemical shift for the signals of the pyridine nuclei and their variation in the two complexes ($\Delta\delta_{\text{Ha}} = 0.23 \text{ ppm}$; $\Delta\delta_{\text{Hb}} = 0.25 \text{ ppm}$ in CD₂Cl₂ and $\Delta\delta_{\text{Ha}} = 0.23 \text{ ppm}$; $\Delta\delta_{\text{Hb}} = 0.19 \text{ ppm}$ in DMSO-*d*₆, see Fig. S93 and 104†) support the same coordination pattern in both cases.

Dilution experiments down to 0.5 mM both in CD₂Cl₂ or DMSO-*d*₆ show that the self-assembled complex is the only species at 2.5 mM in CD₂Cl₂ and in the 5–2.5 mM range in DMSO-*d*₆ and the major one down to 1 mM in both solvents. This fact strongly suggests that the more entropically favored complex is being obtained. At 0.5 mM in CD₂Cl₂, the complex is still the major species in solution, although the free ligand is clearly observed. In contrast, a considerable amount of the bare (*S,S,P*)-7 ligand is observed in DMSO-*d*₆ at the same concentration. This is not surprising taking into account the





Fig. 9 (a) Partial ¹H NMR (500 MHz, CD₂Cl₂) spectra of ligand ((*S,S,P*)-7) (top) and metallocyclic complex ((*S,S,P*)-7)₂Pd₂ (bottom). The lettering corresponds to that shown in Fig. 1; (b) observed (top) and calculated (bottom) HRMS (MALDI⁺) isotopic pattern for the [M - Cl]⁺ ion of metallocyclic complex ((*S,S,P*)-7)₂Pd₂; (c) ECD spectra of complex ((*S,S,P*)-7)₂Pd₂ (black line) and its enantiomer ((*R,R,M*)-7)₂Pd₂ (red line).

higher coordinating character of DMSO compared to CH₂Cl₂, which leads to a higher self-assembly concentration.³³

The self-assembly of a ((*S,S,P*)-7)₂Pd₂ complex was further demonstrated by MALDI mass spectrometry of an evaporated sample of the complex. The high-resolution mass spectrum shows a signal at *m/z* = 2193.3, whose exact mass and isotopic distribution (Fig. 9b) nicely match those calculated for the [M - Cl]⁺ ion, resulting from the loss of one chloride ligand by one of the Pd centers.

Chiroptical properties were not particularly informative owing to just a moderate distortion of the electronic states after coordination. Therefore, very similar UV-Vis and almost identical ECD spectra to those of the free ligand were obtained (Fig. 9c and S122[†]). To confirm that we were really observing the signal associated with the complex, the ¹H NMR spectra of

the sample were recorded before and after the UV-CD measurements, performed at the NMR concentration (Fig. S123 and S124[†]).

Finally, the structures of metallocyclic complex ((*S,S,P*)-7)₂Pd₂ and its ((*R,R,M*)-7)₂Pd₂ enantiomer were investigated by means of DFT calculations at the ωB97XD/def2SVP level of theory both in DMSO and CD₂Cl₂ (Fig. 10 and Tables S25–28[†]). The optimized structures gave structural arrangements which were in good agreement with the experimental data. They showed the coordination of the pyridine rings to the Pd(II) metal centers and the generation of a chiral cavity defined by the helical cores.

Conclusions

In this work, we have synthesized a new family of enantiopure helical foldamers with an increasing number of turns. Moreover, in the presence of the Ag(I) cation well defined structures containing up to three ions and four turns are formed. As a consequence of such a helical structure intense chiroptical properties associated with electron transitions both from the ground (ECD) and excited state (CPL) have been observed. The versatility of this approach has been further validated by preparing an enantiopure metallocyclic structure upon coordination with Pd(II), thus enriching this chemistry with a new class of chiral ligands. We have observed that *g*_{abs} does not monotonically depend on the number of turns, in contrast to the trend observed in the ¹B_u band in helicenes.⁸¹ While the magnetic dipole transition moment seems non progressive with increasing number of rings in helicenes, we have found a lineal dependence of this vector in these helical foldamers. This fact can also be related to the potential use of these systems as spin filters, the magnetic dipole transition moment being a potential predictor of the CISS-based spin filtering capabilities of these Ag(I)-complexes.

Author contributions

A. M. O.: formal analysis, investigation, validation. P. R.: formal analysis, investigation, validation. S. R.: formal analysis, investigation. L. A. C.: conceptualization, formal analysis, writing-original draft. V. B.: investigation, formal analysis, writing-original draft. J. M. P.: investigation, formal analysis. A. J. M.: formal analysis, methodology, writing-original draft. G. M.: methodology, formal analysis. S. A.: formal analysis, investigation, validation. J. U.: methodology, conceptualization, formal analysis, writing-original draft. V. M.: methodology, conceptualization, formal analysis, writing-original draft. G. L.: conceptualization, formal analysis, writing-original draft. D. M.: funding acquisition, methodology, supervision, project administration, writing-original draft, writing - review. J. M. C.: conceptualization, funding acquisition, methodology, project administration, supervision, writing-original draft, writing - review and editing.

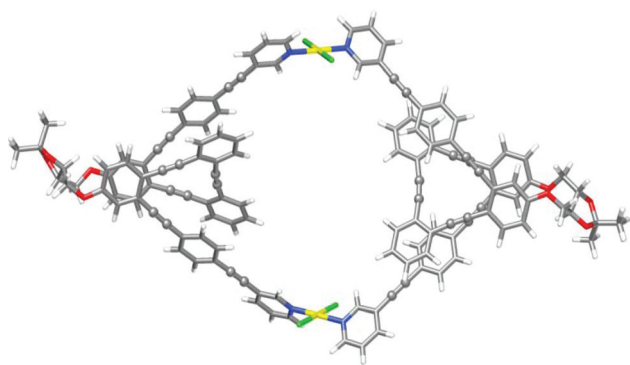


Fig. 10 Optimized DFT (ωB97XD/def2SVP, DMSO) structure of ((*S,S,P*)-7)₂Pd₂. Color coding: C, grey; H, white; O, red; Pd, yellow; Cl, green.



Conflicts of interest

There are no conflicts to declare.

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