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Introduction

Spectrally-resolved third-harmonic generation and the fundamental role of O–H····Cl hydrogen bonding in O_h , T_d -cobalt(II) tetraphenylmethanebased coordination polymers⁺

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The reaction of a phosphonate-diester tetraphenylmethane-based tecton, tetrakis[4-(diethoxyphosphoryl)phenyl]methane, (L) with cobalt(11) chloride afforded a centrosymmetric coordination polymer (CP), $[L\cdot 2Co(H_2O)_4^{2+}\cdot 2CoCl_4^{2-}]_{q_2}$, **2-Cl**, possessing simultaneously octahedral (O_h) and tetrahedral (T_d) metal centers. This material served as a model compound for the demonstration of factors influencing the spectral dependence of one of the nonlinear optical (NLO) phenomena, the third-harmonic generation (THG). The spectrally-resolved THG (SR-THG) measurements in the range from 1125 to 1750 nm revealed that a maximum of THG response is obtained when the fundamental beam is around 1300 nm. The SR-THG study was combined with an analysis of the self-absorption effects of pumping and of thirdharmonic radiation; based on these results, we put forward a hypothesis that the THG action spectrum is influenced more by the ability of the material to self-absorb the third harmonic rather than by the extent of self-absorption of the pumping radiation. Apart from investigations of NLO properties, we have explored coordination and particularly the supramolecular interactions that build up the 2-Cl CP. Despite the tetrahedral, spatial shape of the ligand L, CP 2-Cl has a two-dimensional net. The structure was found to be strongly supported by O-H···Cl hydrogen bonds, since each $CoCl_4^{2-}$ complex anion is an acceptor of eight of such interactions within a distorted square grid layer of cobalt(II) ions. While coordination and hydrogen-bonded nets are both featuring the sql topology when treated separately, the consideration of both of them as topological paths yields a trinodal 4,4,6-connected net, described by the point symbol $(4^2 \cdot 8^4)(4^5 \cdot 6)_2(4^6 \cdot 6^6 \cdot 8^3)_2$. SR-THG and structural studies of **2-Cl** have been also supported by far- and mid-infrared spectroscopy, UV-Vis-NIR solid state absorption analysis, thermogravimetry and preliminary magnetic characterization.

Frequency mixing phenomena such as SHG (second-harmonic generation) and THG (third-harmonic generation) are nonlinear optical parametric processes in which the pumping laser radiation is nonresonantly transformed within the material (that is, without participation of photon absorption events by electron levels), as a result producing the radiation at two and three times shorter wavelengths, respectively.¹ An important feature of the former phenomenon, arising from the properties of the second-order optical susceptibility $\chi^{(2)}$ tensor, is that it can be elicited only in crystalline materials featuring non-centrosymmetric space groups; hence, SHG can be treated as a spectroscopic tool for the determination of the "true" crystal non-centrosymmetry. The THG process, on the other hand, is described by the third-order optical susceptibility tensor $\chi^{(3)}$, which, in contrast to $\chi^{(2)}$, has non-zero components under all symmetries. Accordingly, frequency tripling is achievable in any crystalline solid, thus in all CPs (coordination polymers). However, while the SHG process has been broadly studied by many research groups in CP materials²⁻⁷ over the past few years, the THG phenomenon has not been investigated until recently and to our knowledge there are only a few reports that highlight the THG process in these materials.^{8,9} An in-depth review of CPs as materials not only for frequency mixing but also for other NLO processes such as

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[†]Electronic supplementary information (ESI) available: PXRD traces, crystallographic tables, additional characterization of the THG process, figures showing the solid state absorption spectrum, ¹H, ³¹P, ¹³C NMR spectra, IR spectra, TGA/ DTA traces and additional discussions. CCDC 1541606. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c7dt01518f

two- and multiphoton absorption as well as triplet-triplet upconversion and lasing can be found elsewhere.¹⁰

In our previous contribution we demonstrated, using the spectrally-resolved SHG (SR-SHG) technique for a series of non-centrosymmetric tetraphosphonate diester (dialkyl) CPs¹¹, that measurements of the SHG response in a broad wavelength range provide an approach which is superior to routinely conducted Kurtz-Perry-type^{12,13} single-wavelength measurements since they are able to reveal distinct maxima and minima in SHG action spectra, in particular when the CPs of interest possess absorption bands in the Vis-NIR region. We have also shown that it is essential to compare the solid state absorption spectra with the SHG action spectra to understand why at some wavelengths the SHG response is suppressed due to material's self-absorption of laser radiation excitation or its harmonics.

Having this experience in hand we asked ourselves whether those observations are also relevant for the THG action spectra of CPs. In particular, we wished to establish whether the self-absorption of pumping and emitted radiation is also the dominant factor shaping the SR-THG (spectrally resolved third-harmonic generation) action spectra of CPs.

To provide a demonstration of these issues, we prepared a CP, called herein 2-Cl, constructed from tetrakis[4-(diethoxyphosphoryl)phenyl]methane (L) and cobalt(II) ions of both tetrahedral (T_d) and octahedral (O_h) geometries. Noteworthily, this CP is a centrosymmetric analogue related to SHG-active phosphonate-ester non-centrosymmetric CPs.¹¹ As a result of its peculiar structure, apart from NLO studies, CP 2-Cl gave us the possibility to investigate unusual for most of the phosphonates14-23 and phosphonate monoester24-28 CPs structural aspects, important from the viewpoint of crystal engineering. It is the moderate strength of dialkylphosphonate binding via phosphoryl (P=O) donor coordination combined with the bulkiness of this functional group that induces monodentate axial binding of metal ions.²⁹⁻³² Moreover, the lack of the deprotonation process of dialkyl phosphonates, in analogy to complexes with N-donor bases, enforces the creation of halidemetal anions (e.g. $CoCl_3^-$, $CoCl_4^{2-}$) in order to maintain crystal electroneutrality.^{11,33,34} Nonetheless, participation of the triand tetrahalide complexes of metal ions in framework construction via noncovalent forces such as hydrogen-bonding (HB) is often neglected in favour of coordination analysis, thus in the present paper we place a structural focus on the crucial role of O-H…Cl HB bonds in sustaining the 2-Cl network, also highlighting the implications on the topology of the supramolecular framework.

Results and discussion

Synthesis

In this work, we present the improved synthesis of ligand L, which previously was obtained by us using nickel-catalyzed phosphonylation in boiling *tert*-butylbenzene.³⁵ Here, mild high yield phosphonylation has been realized at 85 °C using



Scheme 1 The structural formula of ligand L and comparison of preparation routes of the noncentrosymmetric and centrosymmetric CPs from L.

C–P coupling with diethyl phosphite, catalyzed with the generated *in situ* dppf (1,1'-bis(diphenylphosphino)ferrocene) complex of palladium acetate, adapting Belabassi *et al.*'s synthetic protocol.³⁶ Moreover, we took advantage of the Kalek *et al.*³⁷ approach, by adding potassium acetate to the reaction mixture, which has been shown to significantly shorten the reaction time. Indeed, the use of the palladium acetate/dppf/potassium acetate system afforded the ligand L in 84% yield (chromatographic purification), after 14 hours of reaction.

The synthesis of **2-Cl** follows the diethyl ether vapour diffusion approach, used by us previously to obtain related non-centrosymmetric CPs based on ligand L (Scheme 1). We found that to obtain a pure phase of **2-Cl**, free of **1-Cl**, a molar excess of metal salt is required (cobalt(π) salt to ligand L ratio equals 6:1). Addition of chloroform is advantageous, due to the shifting of the equilibrium towards the formation of tetrahedral cobalt(π) species. When a smaller cobalt(π) salt excess is used (4:1), without the addition of chloroform, crystallization of both phases (**1-Cl** and **2-Cl**) is observed, as judged by PXRD (Fig. S1, ESI[†]).

To investigate the influence of the halide counter-ion on spectroscopic and THG properties, we also tried to synthesize 2-Cl analogues using $CoBr_2$ and CoI_2 salts; however, we consistently obtained non-centrosymmetric phases (1-Br and 1-I, respectively) instead of isomorphous 2-Br and 2-I. We suppose that larger lengths of Co–Br (2.41–2.43 Å at room temperature)³⁸ and Co–I bonds (2.59–2.62 Å, at 150 K)³⁹ result in a much bigger size of $CoBr_4^{2-}$ and CoI_4^{2-} which can induce the mismatch of those anions within the hydrogen-bonded cavity. Moreover, larger ionic radii of Br⁻ and I⁻ compared to Cl⁻, combined with lower electronegativity may cause O–H…Br and O–H…I hydrogen bonds to be weaker and therefore will not be strong enough to sustain the structure of 2-Cl (a detailed analysis of hydrogen-bonding below).

Coordination, topology and hydrogen-bonding

The coordination polymer 2-Cl crystallizes in the centrosymmetric space group *Ccca* of the orthorhombic system (no. 68). The asymmetric unit of 2-Cl comprises one-fourth of the ligand residue, half of the tetrahedral cobalt(π) ion (Co1) with two chloride anions and half of the octahedral cobalt(π) ion (Co2) with three coordination water molecules (Fig. S2, ESI†). The composition of 2-Cl can be described by the formula $[L\cdot 2Co(H_2O)_4^{2+}\cdot 2CoCl_4^{2-}]_n$, in which L stands for tetrakis [(diethoxyphosphoryl)phenyl]methane.

As shown in Fig. 1a, the tetrahedral cobalt(π) ion (Co1) is coordinated by four chloride anions (Cl1, Cl1', Cl2, and Cl2'), forming a complex anion CoCl₄²⁻. Co–Cl bond distances are almost identical (for Co1–Cl1 and Co1–Cl2 equal to 2.2728(5) Å and 2.2716(6) Å, respectively). The Co1 coordination sphere is not an ideal tetrahedron, since Cl1–Co1–Cl2 and Cl1–Co1– Cl1 angles take values of 106.98(2)° and 111.49(2)°, respectively. The Co2 cation is coordinated by four coordination water molecules (O1W, O1W', O2W, O3W) in equatorial positions, while the axial positions are occupied by oxygen atoms of phosphoryl groups (O1 and O1'), forming octahedral geometry. Here, the coordination sphere is distorted with respect to both bond lengths and angles. The shortest distance to the oxygen donor equals 2.052(2) Å (Co2–O3W bond), while the longest one is equal to 2.083(2) Å, for the Co2–O1W bond.

Each ligand molecule coordinates to four Co2 cations, giving large coordination rings parallel to the *ab* plane. Recurrence of this motif leads to the formation of a cationic metal–organic corrugated sheet; indeed, a topological simplification⁴⁰ of the **2-Cl** coordination net indicates that this CP is

featured by two-periodic **sql** topology (Fig. 1b), in which the center of the ligand is a four-connected node.

One can distinguish a layer of cobalt(π) ions parallel to the *ac* plane. Every tetrahedral cobalt(π) ion is surrounded by four octahedral cobalt(π) ions (Fig. 2a) and *vice versa*. It follows that the CoCl₄²⁻ anion is an acceptor of eight O–H…Cl hydrogen bonds. Among them, four are crystallographically independent (O1W–H1W1…Cl2, O1W–H2W1…Cl2, O2W–H1W2…Cl1, O3W–H2W3…Cl1), with bond distances ranging from 3.130 Å to 3.242 Å. An array of O–H…Cl hydrogen bonds, according to the graph set notation,⁴¹ can be resolved into two basic ring motifs: R_2^2 (8), denoted in Fig. 2b as **A** and **A**', which incorporate cobalt(π) ions of both geometries, chloride anions, and water molecules, and R_4^2 (8), named **B** and **B**', which involves two water molecules bridged by chloride anions.

It seems that such an extensive $O-H\cdots$ Cl hydrogen bonding system, in which $CoCl_4^{2-}$ resides in supramolecular cavities, is a crucial element for the formation of the CP **2-Cl** that provides an explanation for our inability to obtain **2-Br** and **2-I** isomorphs.

Taking into account the precise geometrical criteria, the obtained supramolecular layer is not an ideal 2D square grid,





Fig. 1 (a) Coordination environments of the Co1 cobalt(II) ion (left), and of the Co2 cobalt(II) ion (dark grey rhombuses). Displacement ellipsoids are drawn at 50% probability level. (b) A coordination layer projected along the *c*-axis, with overlaid topological representation of the **sql** net. Symmetry codes: (i) 1 - x, y, 0.5 - z; (ii) 1.5 - x, 1 - y, z.

Fig. 2 (a) A hydrogen-bonded layer of Co1 and Co2 cobalt(II) ions with shown coordination environments. Topological representation of the **sql** net is overlaid on that layer (light grey squares). Co1 ions are drawn as dark blue polyhedra, Co2 ions are drawn as purple polyhedra. With letters A, A', B, B' are indicated regions of identified hydrogen-bond graph sets. (b) Definitions of graph sets and structural formulae highlighting participating atoms.

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since one can find two unique Co1···Co2 distances: $d_1 = 5.653(3)$ Å and $d_2 = 5.680(3)$ Å (Fig. 2a). The distance between these metal ion layers, d_3 , is equal to 12.735 Å (Fig. 3a). Noteworthily, the angle between situated in row Co1-Co2-Co1 ions is lower than 180° (178.3°). The square grid arrangement of cobalt(II) ions translates into the **sql** topology of the hydrogen-bonded network, with four-connected nodes centered at Co1 and Co2 ions (Fig. 2a).

Consideration of both coordination bonds and hydrogenbonds as topological paths leads to determination of the novel trinodal 4,4,6-c net of point symbol $(4^2 \cdot 8^4)(4^5 \cdot 6)_2(4^6 \cdot 6^6 \cdot 8^3)_2$, in which point symbol notations $(4^2 \cdot 8^4)$, $(4^5 \cdot 6)$ and $(4^6 \cdot 6^6 \cdot 8^3)$ belong to the center of the ligand, Co1, and Co2, respectively. Graphical representation of this topology is shown in Fig. 3b. While the Co2 node was not a topologically unique node when considering only coordination bonds, here, due to participation in the hydrogen-bonded subnet, it became a six-connected node.

There are known cobalt(n) CPs in which Co^{2+} ions are located in layers. Within a layer those ions were found to arrange in dimers,⁴² trimers⁴³ or even can form hexagonal



Fig. 3 (a) A projection showing the organic and metal ion layers in the structure of **2-Cl**, projected along the *a*-axis. (b) A corresponding topological representation of three-nodal 4,4,6-c net of $(4^2 \cdot 8^4)(4^5 \cdot 6)_2(4^6 \cdot 6^6 \cdot 8^3)_2$ point symbol, projected along the *a*-axis.

structures.⁴⁴ An analysis of structures of already known CPs of the square grid type of cobalt(II) arrangement clearly indicates that usually small polytopic ligands were required to obtain such a net, *e.g.* terephtalic acid,⁴⁵ *p*-carboxycinnamic acid,⁴⁶ pyromellitic acid,⁴⁷ 4-(*N*-imidazolo)benzoic acid,⁴⁸ or 3,5dinitrobenzoic acid.⁴⁹ In this context, the presented **2-Cl** distinguishes itself in two respects; on the contrary to the listed above ligands, **2-Cl** is constructed from a spatial, bulky ligand. Moreover, **2-Cl** is the first CP in which tetrahedral and octahedral ions are placed alternately within layers.

We have shown the importance of the coordination and hydrogen-bonded net, but in the case of tetraphenylmethanebased compounds, phenyl embraces (PEs) have to also be considered as synthon interactions.^{50–53} PEs are cooperative supramolecular interactions which rely on the multiple edge-to-face C-H··· π contacts between phenyl groups. In 2-Cl, however, no phenyl embraces can be found. Indeed, the shortest distances between central carbon atoms, d(Ccore...Ccore), of tetraphenylmethane moieties are equal to 11.333 Å and 14.948 Å; such values are far beyond the upper limit required for PE (around 7.4 Å).⁵³ The lack of PEs can be attributed to the separation of ligand molecules by octahedral cobalt(II) ions. Also, the relative bulkiness of the diethoxyphosphoryl group needs to be considered. The latter factor, however, does not seem to be a major obstacle since PE analysis of the crystal structure of ligand L, recently published by Perez et al.,⁵⁴ shows that ligand molecules are themselves capable of forming fourfold PE $(4\text{PE}, d(\text{Ccore} \cdots \text{Ccore}) = 7.661 \text{ Å}).$

Spectrally-resolved third-harmonic generation

In most reports describing the SHG performance of CPs using the powder Kurtz-Perry method, the SHG response of a powder of a compound is expressed as relative to SHG intensity of a reference material (*e.g.* KDP, potassium dihydrogen phosphate). In this work, the THG intensities are also expressed as relative values, as a ratio of the integrated THG signal collected from the sample to the integrated THG signal collected from the standard (KDP), normalized to the same integration time. Special care was taken to ensure that geometry of the experimental setup was strictly the same during the measurement of the CP sample and reference at a given wavelength.

This way of presenting the spectral dependence of THG avoids problems with normalization or adjusting the power density of the laser to the constant power value throughout the measurement region. Since the parameters influencing the THG intensity of the reference, *i.e.* $\chi^{(3)}$ and the refractive indices are not supposed to vary markedly within the wavelength range of the measurements, the observed wavelength dependence of the relative THG intensity should mostly reflect the variation of the properties of the investigated compound.

Compound 2-Cl is not emissive upon one-photon excitation (investigated range 250–650 nm), which is probably due to the fluorescence quenching abilities of cobalt(II) ions. Nonetheless, in order to exclude two-photon (and multiphoton) induced fluorescence as a possible constituent of emitted radiation, we investigated closely the emission spectra. As presented in Fig. S3, ESI[†] the only origin of emitted radiation, for both 2-Cl and KDP, appears to be the third harmonic of the fundamental laser radiation (with broadening due to instrumental factors), with no detectable bands of fluorescence.

Additional evidence proved the character of the collected radiation as THG is a measurement of the power dependence of the emitted radiation intensity. Indeed, as seen in Fig. S4, ESI,[†] varying the laser power from 53 to 84 mW at 1500 nm leads to the slope of the linear regression of the $\log(I_A) = f(\log(P))$ relationship equal to 3.1, confirming the presence of a THG process.

The relative intensities of THG of **2-Cl** expressed *versus* KDP (SR-THG action spectrum) are presented in Fig. 4. The data were collected in the range from 1125 to 1750 nm. Starting from 1125 nm at which the THG response is the lowest (0.2 that of KDP), the relative THG strength steeply rises to reach the maximum at around 1300 nm (1.1–1.2 that of KDP). For longer wavelengths, up to 1750 nm the relative THG response decreases, at 1750 nm THG of **2-Cl** *vs.* KDP is equal to 0.38. Measurements at wavelengths shorter than 1125 nm and longer than 1750 nm were unsuccessful due to either decomposition of the sample upon laser radiation, or due to the inability to capture the THG signal with a reasonable signal-to-noise ratio.

In our recent work, we showed that the SHG action spectra of CPs exhibit a complex structure, with maxima at wavelengths at which the self-absorption of both pumping and emitted radiation by the material is the lowest. Thus, to predict the position of SHG maxima the measurement of the solid-state absorption spectrum is needed. The spectrum of 2-Cl with analysis and assignments of absorption bands is provided in the ESI (Fig. S5†).

It was interesting to know whether a maximum of the THG response can be rationalized in the same manner. Accordingly,



Fig. 4 An overlay of the THG action spectrum (red points through which the red line is led, just to guide the eyes) with the solid state absorption spectrum plotted against wavelength (black line) and the solid state absorption spectrum plotted against the tripled wavelength (grey line).

using the analogous approach, we compared the obtained THG action spectrum with the solid state absorption spectrum of 2-Cl plotted in a usual manner (vs. λ , black line) and also plotted against a tripled wavelength (vs. 3λ , grey line), as presented in Fig. 4. The absorption spectrum plotted against the wavelength shows the ability of the material to self-absorb the fundamental frequency radiation, while the same absorption spectrum plotted against thrice the wavelength represents the extent of the THG self-absorption. In the case of 2-Cl, an overlay of these absorption spectra in Fig. 4 does not predict the THG response in a quantitative way. For example, at 1100 nm the absorbance of the fundamental is near zero, and the absorbance of the emitted radiation is moderately strong, while at 1400 nm the absorbance values of both the fundamental and the third harmonic are comparable. It might seem intuitive that the THG response should be higher at 1100 nm than at 1400 nm, while the opposite behaviour is observed. This suggests that self-absorption of the emitted radiation may have a more important influence on the THG action spectrum than self-absorption of the fundamental. Indeed, the maximum of the THG response coincides with the minimum of the absorption spectrum plotted vs. 3λ . In the same vein, in our previous work we observed that the presence of an additional absorption band in the UV-blue region in noncentrosymmetric iodide-containing CP (1-I) caused almost complete suppression of SHG activity.¹¹

An important question is thus what might be the physical reasons for bigger importance of the third harmonic selfabsorption (per analogy also of second harmonic) compared to that of the fundamental. It is clear that pumping light will penetrate deeply into the specimen, due to relatively low scattering of the long-wavelength radiation (which is proportional to λ^{-4}). On the other hand, the generated third harmonic will experience much higher scattering than the fundamental beam, and, since it will be emitted essentially in all directions, it will also be self-absorbed at longer optical paths than that for the pumping beam. In addition to absorption and scattering, an important factor here is also that of the relatively short coherence length expected for the THG process, which limits the optical paths from which the THG is collected.

Preliminary magnetic characterization

The magnetic measurement was conducted in a constant field (300 Oe) in the 1.7–300 K temperature range. Values of magnetization and of magnetic moment are provided per Co ion. In order to determine the fundamental magnetic parameters of compound 2-Cl, the $1/\chi_{\rm M}$ and $\chi_{\rm M}T$ plots as a function of temperature have been drawn (Fig. 5).

As shown in Fig. 5, the experimental $\chi_{\rm M}T$ value at room temperature for CP **2-Cl**, calculated per cobalt(II) ion is equal to 2.63 cm³ K mol⁻¹. This value decreases by lowering the temperature to 55 K (2.35 cm³ K mol⁻¹), where an inflection point is noticeable. From 45 K one can see a strong decrease of $\chi_{\rm M}T$ product to 1.8 cm³ K mol⁻¹ at 1.8 K. The $\chi_{\rm M}T$ value at room temperature is much bigger than that theoretically



Fig. 5 Variation of χT product (triangles) and of χ^{-1} (squares) with temperature. Red line corresponds to the fit in the 50–300 K temperature range with the Curie–Weiss law, while the blue line corresponds to the fit in the 50–300 K temperature range with eqn (1).

calculated for the magnetically isolated high-spin (S = 3/2) cobalt(II) ion (1.875 cm³ K mol⁻¹, 3.87 B.M.) for g = 2.0.⁴⁴

Zero-field splitting as well as the spin-orbital coupling occurring in cobalt(II) ions essentially complicate the interpretation and fitting of the magnetic data of complexes and coordination polymers possessing those ions.55 While the latter phenomenon is present to a lesser extent in the case of tetrahedral geometries (singlet term ⁴A₂)⁵⁶ it is pronounced in the case of distorted octahedral cobalt(II) complexes. These phenomena are the origin of the abnormally increased $\chi_{M}T$ product observed for CP 2-Cl. Fitting of the magnetic data in the 50-300 K temperature range with the Curie-Weiss law gives Curie constant $C = 2.71 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$ and Weiss constant θ = -7.83 K, but it should be kept in mind that a negative θ value for cobalt(II) compounds, apart from the occurring antiferromagnetic interaction between metal centers, may have the origin in the strong spin-orbit coupling as well. Thus, to more accurately quantify the strength of the antiferromagnetic exchange, a fitting of the data to the following phenomenological equation^{57,58} has been conducted:

$$\chi T = A \, \exp\left(-\frac{E_1}{kT}\right) + B \, \exp\left(-\frac{E_2}{kT}\right) \tag{1}$$

In eqn (1), A + B is equal to the Curie constant, E_1/k is the activation energy of spin–orbit coupling, and E_2/k stands for the activation energy of the antiferromagnetic exchange.

The best fit in the 50–300 K temperature range gave $A + B = 2.70 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$, $E_1/k = 74.60 \text{ K}$, and $-E_2/k = -6.07 \text{ K}$. A + B value matches well with the Curie constant. The E_1/k value is in the range expected for distorted Co(π) magnetic centers, while E_2/k confirms small antiferromagnetic interactions. We stress that, due to the presence of cobalt(π) ions of both geometries, the calculated parameters should be treated as intermediate values; in other words these can be viewed as an "average" of magnetic participations of tetrahedral and octahedral cobalt(π) ions. A much more accurate modelling of the magnetic properties of mixed-geometry cobalt(π) systems

would require *e.g.* the use of matrix-diagonalization techniques, as demonstrated elsewhere.^{33,34,59–61}

Experimental

Crystallography

The obtained single crystal of 2-Cl was used for data collection on a four-circle KUMA KM4 diffractometer equipped with a two-dimensional CCD area detector. Graphite monochromatized Mo K_a radiation (λ = 0.71073 Å) and the ω -scan technique $(\Delta \omega = 1^{\circ})$ were applied, while additional data collection and reduction, along with absorption correction, were performed using the CrysAlis software package.⁶² The structure was solved by direct methods using SHELXS-2012⁶³ revealing the positions of all or almost all non-hydrogen atoms. The remaining atoms were located as a result of subsequent difference Fourier syntheses. The structure was refined using SHELXL-2012 with anisotropic displacement parameters. Hydrogen atoms connected to carbon atoms were constrained as a riding model. The positions of the H atoms of water molecules were located on a difference Fourier map, where possible, and constrained. The collection parameters data, crystallographic data and final agreement parameters are listed in Table 1. Selected geometrical parameters are listed in Table S1.† The hydrogen bonding geometry for 2-Cl is listed in Table S2.[†] Visualization of the structures was conducted using the Diamond software,⁶⁴ while a topological simplification of coordination networks was performed with the ToposPro package.40 Geometrical features of intermolecular interactions were determined with the help of Platon software.65

Materials

Starting materials were of reagent grade purity and were obtained from commercial sources and used without further purification, except for dioxane, which was kept over 3 Å

 Table 1
 Crystallographic collection and refinement data for 2-Cl

Formula	$C_{41}H_{72}Cl_8Co_4O_{20}P_4$
$M[\text{g mol}^{-1}]$	1528.14
Crystal system	Orthorhombic
Space group	<i>Ccca</i> (no. 68)
a, b, c [Å]	15.6558(5), 25.4691(8), 16.3904(5)
α, β, γ [°]	90, 90, 90
$V[Å^3]$	6535.5(4)
Z	4
Density calc. $[g \text{ cm}^{-3}]$	1.553
$\mu(MOK_{\alpha}) [mm^{-1}]$	1.485
F(000)	3128
Crystal size [mm]	0.22 imes 0.28 imes 0.32
Temperature [K]	295
Radiation [Å]	MoK _α 0.71073
$\theta_{\min}, \theta_{\max}$	2.9, 28.5
hkl range	-20:20;-33:34;-20:22
Reflexes measured, independent	41 875, 4134
R _{int}	0.029
R_1, WR_2, S	0.0290, 0.0742, 1.03
$\Delta \rho_{\rm max}, \Delta \rho_{\rm min} [e {\rm \AA}^{-3}]$	-0.26, 0.43
CCDC number	1541606

Aldrich molecular sieves. Tetrakis(4-bromophenyl)methane was prepared according to a previously known procedure.⁶⁶

Optimized Pd-catalyzed synthesis of tetrakis[4-(diethoxyphosphoryl)phenyl]methane, (L)

Tetrakis(4-bromophenyl)methane (2.50 g, 3.92 mmol) and 40 cm³ dioxane were placed in a two-necked round-bottom flask and heated to 85 °C. Nitrogen was purged *via* a reaction system for *ca.* 15 minutes. After purging, anhydrous $Pd(OAc)_2$ (9.7 mg, 0.043 mmol, 1.1 mol%) and dppf (49 mg, 0.088 mmol, 2.2 mol%) were added and stirred at 85 °C until a dark brown coloration of the palladium–phosphine complex appeared. Next, diethyl phosphite (3.03 cm³, 23.52 mmol, 6 eq.), triethylamine (3.28 cm³, 23.52 mmol, 6 eq.), and anhydrous potassium acetate (0.300 g, 3.05 mmol, 0.78 eq.) were added in single portions. Heating continued for 15 hours at that temperature and under a nitrogen atmosphere.

The mixture was cooled down to room temperature and a white precipitate of triethylammonium hydrobromide was filtered and discarded. After evaporation of the filtrate under reduced pressure, a slightly viscous solid was purified using column chromatography (silica gel 60 Å (63–200 µm)), a 10:1 v/v mixture of ethyl acetate/ethanol ($R_{\rm f} = 0.04$, silica gel on PET foils with fluorescent indicator 254 nm, UV lamp for the visualization). Yield: 2.85 g (84%) mp. (ethyl acetate) 177.1–183.8 °C.

Alternatively, the compound can be purified by recrystallization from an ethyl acetate–cyclohexane mixture,³⁵ but at the expense of lower yield (around 70%). Note that palladiumcatalyzed phosphonylation was previously achieved using the unusual $Pd(PPh_3)_4$ -triisopropyl phosphite system, at $220 \ ^{C}$.^{17,67}

HRMS (ESI): calc. for $C_{41}H_{57}O_{12}P_4$ (M + H⁺): 865.2800; found: 865.2831 (*m*/*z*). ¹**H NMR** (400 MHz, CDCl₃) δ 7.69 (dd, ³*J*_{P-H} = 12.9 Hz, ³*J*_{H-H} = 8.4 Hz, 2H), 7.28 (dd, ³*J*_{H-H} = 8.4 Hz, ⁵*J*_{P-H} = 3.8 Hz, 2H), 4.21–4.05 (m, 4H), 1.32 (t, ³*J*_{H-H} = 7.1 Hz, 6H) ppm. ³¹P{¹H} **NMR** (162 MHz, CDCl₃) δ 18.7 (s) ppm. ¹³C{¹H} **NMR** (151 MHz, CDCl₃) δ 149.31, 131.53 (d, ²*J*_{P-C} = 10.3 Hz), 130.85 (d, ³*J*_{P-C} = 15.1 Hz), 127.00 (d, ¹*J*_{P-C} = 190.9 Hz), 65.61 (s), 62.43 (d, ²*J*_{P-C} = 5.6 Hz), 16.44 (d, ³*J*_{P-C} = 6.3 Hz) ppm. NMR spectra are presented in Fig. S6–S8, ESI.†

Synthesis of $2Co(H_2O)_4^{2+} \cdot 2CoCl_4^{2-} \cdot L$, (2-Cl). Ligand L (23.0 mg, 0.0265 mmol) and anhydrous $CoCl_2$ (20.7 mg, 0.159 mmol, 6 eq.) were dissolved in 96% ethanol (1 cm³) in a 10 mL glass vial. Upon addition of chloroform (0.5 cm³), the solution changed the color from pink to dark blue. Then on the top of the vial a screw-cap was placed and lightly closed, to allow vapor diffusion. Next, the vial was placed in a 250 mL chamber filled with diethyl ether (40 mL). After two weeks, dark blue parallel-piped crystals appeared. Yield with respect to the ligand: 20.4 mg (50%).

Anal. calcd for $C_{41}H_{72}Cl_8Co_4O_{20}P_4$: C, 32.22; H, 4.74. Found: C, 32.44; H, 4.59 [%]. MIR and FIR spectra are presented in Fig. S9 and S10, ESI,† respectively. IR tentative assignments are given in Table S3, ESI.† Thermogravimetric characterization and discussion are presented in the ESI (Fig. S11†).

Spectrally-resolved third harmonic generation

For the wide-wavelength range THG measurements of 2-Cl powdered samples and KDP with the same particle size (125–177 µm), which was used as a reference material, were placed between microscope glass slides and excited by tunable fs laser pulses from a laser system consisting of a Quantronix Integra-C regenerative amplifier operating as an 800 nm pump and a Quantronix-Palitra-FS BIBO crystal-based optical parametric amplifier. This system delivers wavelength tunable pulses of ~130 fs length and was operated at a repetition rate of 1 kHz. The incident beam was directed onto the plane of the sample at about 45 degrees while the detector was placed in front of the sample. Laser beam intensity was attenuated by using a Glan laser polarizer and spectra of the THG emission were recorded by using an Ocean Optics 2000 fiber coupled CCD spectrograph for excitation wavelengths in the range from 1125 to 1750 nm. The geometry of the experimental setup and the intensity of the laser beam were exactly the same for the whole experiment and all samples at a given wavelength. Unwanted excitation wavelengths that are usually present in the output of an OPA were removed using color glass filters. After setting the output wavelength (1125-1750 nm, approximately 25 nm step), the intensity of the beam was attenuated, when needed, by using polarizing optics, in order to avoid sample decomposition and white light continuum generated by femtosecond laser pulses. A different integrating time was used for coordination polymers (typically 5000-30 000 ms) and for the KDP sample (500-5000 ms) in order to ensure a reliable S/N ratio. The intensity of the signal was normalized to the same time of collection. CAUTION. Work with a high-power laser brings danger to the eyes, especially in the spectral range in which the beam is invisible. Adequate eye protection should be used during measurements.

Other methods

Powder X-ray diffraction (PXRD) patterns of the coordination polymers were obtained on a PANanalytical X'Pert diffractometer equipped with a Cu-K_{α} radiation source ($\lambda = 1.54182$ Å). Elemental analysis was performed using a CE Instruments CHNS 1110 elemental analyzer. Magnetic susceptibility down to 1.7 K at a field of 30 mT was measured with a Quantum Design SQUID magnetometer. Thermogravimetric analysis (TGA) and differential thermal analysis (DTA) were conducted using a Setaram SETSYS 16/18 thermoanalyzer in the range 20–800 °C at a heating rate of 5 °C min⁻¹ under a flowing 80% nitrogen-20% oxygen atmosphere. Diffuse reflectance spectra (DRS) were obtained on a Cary 500 spectrophotometer in the range 200-2000 nm using BaCO₃ as a reference. The absorption spectra were calculated from the Kubelka-Munk function: $F(R) = (1 - R)^2/2R = K/S$, where R is the diffuse reflectance, K is the molar absorption coefficient, and S is the scattering coefficient. Mid-infrared (MIR) and far-infrared (FIR) spectra were obtained on a VERTEX 70 V FT-IR spectrometer (Bruker Optik GmbH, Ettlingen, Germany) in attenuated total reflection (ATR) measurement mode at room temperature. ¹H, and

 $^{31}\mathrm{P}\{^{1}\mathrm{H}\}$ NMR spectra were recorded on a Jeol JNM-ECZ 400S Research FT NMR spectrometer (JEOL Ltd, Tokyo, Japan) operating at 400 and 162 MHz, respectively. The $^{13}\mathrm{C}\{^{1}\mathrm{H}\}$ spectrum was recorded with the use of a Bruker Avance 600 spectrometer operating at 151 MHz.

Conclusions

The structural, topological and spectrally-resolved THG properties of the centrosymmetric $O_{\rm h}$, $T_{\rm d}$ -cobalt(II) tetraphenylmethane-based coordination polymer, belonging to the rare family of diester-phosphonates, have been characterized. Based on the reactivity of ligand **L** towards cobalt(II) halides as well as on the detailed analysis of coordination and supramolecular interactions, we have shown that O-H…Cl HB bonds are fundamental interactions sustaining the structure of the **2-Cl** network.

By measurements of THG in a broad wavelength range (1125–1750 nm), we determined that the maximum of the THG response is obtained when approximately 1300 nm pumping wavelength radiation is applied. SR-THG was combined with the analysis of the self-absorption effects of pumping and of third-harmonic radiation; based on these results we suggested that the main influence on the THG response is the ability of the material to absorb the third harmonic rather than the extent of self-absorption of the pumping radiation. This demonstrates the importance of further investigations of SR-THG (also SR-SHG) of CPs in which resonant and nonresonant processes may compete.

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