A novel 2D metal oxalatophosphite K₂Mn(II)(H₂O)₂C₂O₄(HPO₃)₂ (KMnCP) was hydrothermally synthesized and characterized. The hexagonal morphology and crystal growth faces of KMnCP were predicted by Bravais–Friedel–Donnay–Harker (BFDH) theory. Single crystal X-ray diffraction analysis revealed that the compound displays a unique double layered structure constructed from Mn(H₂O)(HPO₃) single layers linked by oxalate ligands, where the potassium levels interrupt the 3D connectivity through the organic bridges. Moreover, in order to classify this archetype a crystallochemical revision of the metal oxalatophoshites with anionic frameworks has been carried out. Several Secondary Building Units (SBUs) formed by metal-phosphate substrates and different roles of de oxalate bridges have been observed, what has led to propose a new structural classification for this kind of materials with common features between the classic inorganic metal phosphates and coordination polymers. Finally, the thermal, spectroscopic and magnetic properties together with the electronic structure of the studied compound are discussed.

**K₂Mn(II)₂(H₂O)₂C₂O₄(HPO₃)₂: A New 2D Manganese(II) Oxalatophosphite with Double-Layered Honeycomb Sheets Stabilized by Potassium Ions**

Joseba Orive, Ramesh Sivasamy, Roberto Fernández de Luis, Edgar Mosquera and Maria I. Arriortu

**Introduction**

The design of new sophisticated hybrid materials containing inorganic and organic linkers has experienced a clear interest reflected mainly in the development of a huge number of multifunctional metal–organic frameworks (MOFs) with applications in several fields. The reticular chemistry approach applied during the last decade to synthesize MOFs can be translated to obtaining other inorganic-organic compounds containing a mixture of carboxylate based linkers and inorganic anions, such as phosphate, vanadate or arsenate groups, just to mention some of them. One of these classes of hybrid frameworks is based on oxalate and phosphate groups in which the metal centers are coordinated by both types of ligands, giving rise to extended structures containing common features with the classic inorganic metal phosphates and coordination polymers. As in inorganic-organic classic metal phosphates, many of these called transition metal oxalatophosphates have been prepared using organic templates giving rise to crystalline materials with channels similar to those found in zeolites. Inorganic and organic templates act as structural directing agents, located at the pores of the structure, and balancing the negative charge of the inorganic skeleton.

Unlike classical 4-coordinated PO₄ units, HPO₃ characteristics such as a reduced charge and a fewer number of P-O binding sites can be exploited to design different connectivity patterns. In this sense, many successful examples of organically templated oxalatophosphites including different metals such as indium, cobalt, manganese, iron, gallium and lanthanides have been reported during the last decade.

In contrast to the organically templated zeotypes, examples of oxalate-phosphates and phosphites with alkali metal charge compensating cations as inorganic templating agents encapsulated into the metal organic-phosphate frameworks are scarcer. Due to their similitudes with the coordination polymers, M. Nagarathinam et al. defined these inorganic-organic materials as metal organic–phosphate open frameworks (MOPOFs). Such hybrid materials were tested as cathode materials for lithium ion batteries (LIBs), demonstrating to have great feasibility of exchanging K⁺ with Li⁺ ions and showing good capacity of reversible lithium insertion/extraction. Despite the insertion of the organic
ligand in the hybrid materials decreases the specific capacity of the cathodes, on the other hand it opens new frontiers to explore the lithium ion reversible exchange and migration within more flexible designs of inorganic-organic two-dimensional materials. The phosphate replacement by phosphite units gave rise to obtaining a new family of redox active vanadyl oxalatophosphites with $A_2[(VO)2(HPO3)2(C2O4)]$ ($A = Li, Na$ and $K$) formula.12b This approach has the advantage of getting similar archetypes to that found in the previous mentioned oxalatophosphates together with the increase of the theoretical specific capacity of the compounds due to decrease in the molecular weight.

Such achievements encourage the search of new hybrid materials by varying the alkali and transition metal, organic ligands, or polyanion units which could lead to better and rentable cathode materials with new architectures.

As part of our pursuit of new host lattices with redox-active metal centers which are able to encapsulate mobile ions, here we disclose an experimental and theoretical study of the first manganese (II) oxalatophosphate with an alkali metal charge compensating cation which presents a not very usual layered topology. Moreover, a new structural classification for metal oxalatophosphites has been formulated taking into account the dimensionality of the inorganic metal phosphite skeleton and the role of the oxalate in the structures.

**Experimental section**

**Synthesis and characterization**

$K_2Mn^+$$_2(H_2O)$$2C_2O_4(HPO_3)_2$ (KmncP) was prepared using mild hydrothermal conditions under autogenous pressure. A mixture of $H_3PO_3$ (7.5 mmol), $MnCl_2·4H_2O$ (0.75 mmol) and $C_6H_2O_4·2H_2O$ (1.5 mmol), was dissolved in distilled water (25 ml) and stirred in air. Then, around 5.4 ml of 2M KOH solution was added dropwise to increased up the pH of the resulting solution to 5.0. The reaction mixture was sealed in a 50 ml PTFE-lined stainless steel pressure vessel (fill factor 65%). After 3 days at 120 °C, large hexagonal plates and prismatic crystals of KmnCP, suitable for single crystal X-ray diffraction, were recovered and washed with water and acetone. Some unidentified acicular crystals were removed by decantation from the final product. Fine white powder as pure phase of KmnCP, for further characterization, was achieved by decreasing the heat treatment temperature to 100 °C.

Another possible mobile ions, such as lithium and sodium were tried to encapsulate in the system, but $A_2[(H_2O)2Mn_2(HPO_3)_2]_n$ ($A = Li, Na$) inorganic phosphites crystallize into hexagonal prisms with dimensions 0.269 x 0.083 x 0.073 mm.

The manganese, phosphorous and potassium contents were confirmed by inductively coupled plasma quadrupole mass spectrometry (ICP-Q-MS) analysis, performed with a Thermo Scientific XSERIES 2 spectrometer. The amount of C was calculated by elemental analysis. $K_2Mn_2(H_2O)$$2C_2O_4(HPO_3)_2$ (KmnCP). Calc: K, 16.56; Mn, 23.27; P, 13.12; C, 5.09; H, 1.28. Found: K, 13.5(4); Mn, 23.4(5); P, 12.7(5); C, 4.5(4); H, 1.6(2).

**Single crystal X-ray diffraction study**

A hexagonal prism with dimensions 0.269 x 0.083 x 0.073 mm of the studied compound was selected under a polarizing microscope and glued on a glass fiber. Intensity data were collected at 100 K on an AGILENT SUPERNOVA single source diffractometer with Mo Kα radiation using a CCD (Eos) detector. Data frames were processed (unit cell determination, intensity data integration, correction for Lorentz and polarization effects,17 and analytical absorption correction18 taking into account the size and shape of the crystal) using the corresponding diffractometer software package.19 The structure was solved by direct methods, SHELXS 97 computer program,20 in the hexagonal space group P6$_3$/m, and then refined by the full matrix least-squares procedure based on $F^2$, using the SHELXL 97 computer program21 belonging to the WINGX software package.22 TWINROTMAT option of PLATON program23 suggested the existence of a merohedric twin by 180° rotation around the [120] reciprocal lattice direction, in agreement with the twin’s law expressed by the [-100, 110, 00-1] matrix. After the refinement, the masses of the two components of the twin were found to have percentages of 58.9(2) and 41.1(2).

This scheme permitted us to find the positions of the manganese and phosphorous atoms, and all the other non-hydrogen atoms (K, O and C) were placed from subsequent Fourier-difference map calculations. Next, the hydrogen atoms related to the phosphite units were located. Finally, the hydrogen atoms related to the coordinated water molecule were first located and placed in geometrically ideal positions (O–H: 0.82(1) Å; H–H: 1.35(2) Å) and refined using the riding model. The atomic displacement parameters (ADPs) for all atoms except hydrogens were described using an anisotropic model.

Details of crystal data measurement and reduction, structure solution and refinement of KmnCP are reported in Table 1. The selected bond distances and angles are reported in ESI, Table S1. Structure drawings were made using the ATOMS 6.2.24

**Powder X-ray diffraction**

X-ray powder diffraction measurement for qualitative phase analysis using the pattern matching routine of the FULLPROF program25 for KmnCP was recorded at room temperature using a PHILIPS X’PERT PRO automatic diffractometer equipped with Cu Kα radiation source ($\lambda = 1.5418$ Å). The power generator was set to 40 kV and 40 mA and the pattern was recorded in 20 steps of 0.026° in the range of 5°–70°. The systematic 20 shift; peak shape (pseudo-Voigt), U, V, and W half-width parameters for the profile function, the previously obtained unit cell parameters, asymmetry parameters, and the background were refined. Starting from the space group and the lattice constants of the compound previously studied by single crystal X-ray diffraction, we reached to good agreement between the
experimental and the calculated diffractograms (Fig. S2). Besides, the inset of Fig. S2 shows the great similarity between the experimental powder pattern and the one simulated from the single crystal X-ray diffraction solution, confirming the phase purity of the as-prepared sample.

### Table 1 Crystallographic data and structure refinement parameters for KMnCP obtained by single crystal X-ray diffraction

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tr>
<td>Molecular weight (gmol⁻¹)</td>
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<tr>
<td>Space group</td>
<td>P6₃/m</td>
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<tr>
<td>a, c (Å)</td>
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<td>V (Å³), Z</td>
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<tr>
<td>ρ(Mo Kα), ρ(Mo Kβ) (gcm⁻³)</td>
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<td>Crystal size (mm)</td>
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<td>F(000)</td>
<td>1392</td>
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<tr>
<td>Diffractometer / Temperature (K)</td>
<td>AGILENT SUPERNOVA (omega scan mode) / 100(2)</td>
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<tr>
<td>μ (mm⁻¹), T/min./max.</td>
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<td>Radiation (Å)</td>
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<tr>
<td>Limiting indices h, k, l</td>
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<tr>
<td>Theta range (*)</td>
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<tr>
<td>Completeness (%)</td>
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</tr>
<tr>
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</tr>
<tr>
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<td>Parameters / Restrictions</td>
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<tr>
<td>R [&gt;2σ(int)]</td>
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</tr>
<tr>
<td>R [all data]</td>
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<tr>
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</tr>
<tr>
<td>Max. and Min. e. density (eÅ⁻³)</td>
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</tr>
</tbody>
</table>

### Physicochemical characterization techniques

Thermogravimetric analysis was performed under synthetic air on a Netzsch STA 449 F3 Jupiter simultaneous TG-DSC thermomicrobalance. An alumina crucible containing around 15 mg of the sample was heated in air at a rate of 5 °C/min from room temperature to 800 °C. Thermodiffractometric experiment was carried out in air with a Bruker D8 Advance diffractometer (Cu Kα radiation) equipped with, a variable-temperature stage (HTK2000), a Pt sample heater and a Vantec high-speed one dimensional detector with six degrees of angular aperture. The powder patterns were recorded in the 5 ≤ 2θ ≤ 38° range (step size = 0.033° and time per step = 0.5 s) at intervals of 15 °C, increasing the temperature at 10 °C/min from room temperature to 810 °C. The infrared (IR) spectrum (KBr pellet) was obtained with a JASCO FT/IR-6100 spectrophotometer in the 400-4000 cm⁻¹ range, using the KBr pellet technique. Raman spectroscopy was measured on a powdered sample using a Renishaw RM-1000 spectrophotometer equipped with a thermoelectrically cooled charge-coupled device (CCD) detector. A radiation of 785 nm wavelength laser was used to collect Raman spectrum (operating at maximum power, 1 mW power on sample surface). The laser was focused onto the samples using a Leica microscope with a 50× objective lens. Diffuse reflectance spectrum was measured at room temperature using a Varian Cary 5000 spectrophotometer in the 200–2500 nm range. Magnetic measurements on the powdered sample was performed in the temperature range 5.0-250 K, at 0.1 and 0.01 T using a Quantum Design MPMS-7T SQUID magnetometer.

### Computational methods

The peripheral morphology and possible crystal growth faces of KMnCP crystal have been predicted by Bravais-Friedel-Donnay-Harker (BFDH) method. BFDH is strictly based on the symmetry of the crystal lattice to generate an ordered list of possible growing faces. The relative growth rate of (hk) (G_i) is inversely proportional to the interplanar distance d_(hk) (G_i=1/d_(hk)). The larger the d_(hk), the more pronounced the morphological importance of the face. The electronic structure calculations were executed using DMOl3 code of Accelrys package. The calculation is based on density functional theory (DFT) with the plane-wave pseudopotential method. The Perdew-Burke-Ernzerhof (PBE) of generalized gradient approximation (GGA) was used as the exchange-correlation function. A plane wave energy cut-off was set to 300 eV (ultra-soft pseudopotential) and Brillouin-zone integrations were performed using Monkhorst-Pack k-points of 2 × 2 × 1. The criterion of convergence for the residual forces is set to be less than 0.01 eV/Å and the change of total energy less than 5 × 10⁻⁶ eV.

### Results and discussion

#### Structure description

The asymmetric unit of KMnCP contains 13 non-hydrogen atoms (C, K, Mn, O and P), 3 hydrogen atoms related to the phosphite units and 2 ones associated to a coordinated water molecule (Fig. S3). Mn(1), K(1) and the six oxygen atoms crystallographically independent occupy general positions. The three phosphorous atoms are situated in special positions 4e (P1) and 4f (P(2) and P(3)) with 3... site symmetry, meanwhile C(1) and C(2) carbon atoms occupy 6h special positions over m. site.

The structure of KMnP is built of two 2D Mn(H₂O)₂(P₂O₇)₄ single layers linked by oxalate ligands giving rise to 2D anionic double sheets of [Mn(H₂O)₂(P₂O₇)(C₂O₄)]⁻ formula placed parallel to the ab-plane (Fig. 1a). The distorted MnO₆(C₂O₄) octahedra are corner-shared with three different (HPO₃)²⁻ units forming a honeycomb hexagonal pattern (Fig. 1b). The coordination sphere of the manganese cations is completed by terminal aqua ligand and an oxalate unit (Fig. 1c). The double anionic sheets are stacked along the c-direction by K⁺ ions giving rise to a 3D structure (Fig. 1a).

In the present structure, it is noteworthy that the oxalate anion as tetradentate ligand bonds to two manganese atoms in bidentate-chelating mode, thus forming the peculiar double anionic sheets (Fig. 1c). The shortest metal-oxygen distances are those which are linked to phosphite units in the ab plane ranging from 2.125(1) Å to 2.161(2) Å. M-O bond lengths increasing up to 2.227(2) Å and 2.237(2) Å are observed for the linkages shared with oxalate units. The longest manganese-
oxygen distance (2.245(1) Å) is ascribed to the terminal aqua ligand. All the metal-ligand distances are in good agreement with 2+ oxidation state for the manganese atom, based on the bond valence sums (BVS) calculations.26 [Mn(1)O₅(H₂O)] c/s angles vary from 73.41(5)º to 99.55(7)º, while trans angles values are in the 163.85(6)-175.92(6)º range. The S(Oh) value, calculated by Continuous Symmetry Measure,27 is 0.889, near the value for the octahedral ideal symmetry (S(Oh) = 0.00).

The water molecule coordinated to the manganese polyhedron establishes a triangular patterned intralayer H-bonding scheme (Fig. S4). The average bond length distances and angles O-H···O with the oxygen atoms O(4) and O(6) of the oxalate units are ~ 2.67 Å and in the 160-165º range respectively, suggesting a moderate to strong character of the interactions.28 All P atoms adopt pseudo-pyramidal coordination geometry linking to three Mn atoms via bridging oxygen atoms and to one H atom. The P-O distances are very regular ranging from 1.526(2) to 1.531(2) Å and the P-H distances vary from 1.30(4) to 1.33(4). The O-P-O and H-P-O angles (106.00(7)-112.71(6)º) are in the usual range for phosphite based compounds.29

The framework stoichiometry of [Mn(H₂O)(HPO₃)(C₂O₄)] creates a net charge of -2, which is balanced by two interlayer potassium ions. So, the potassium counter ions play a predominant role in the stabilization of the three-dimensional network by ionic bonding contacts involving the oxygen atoms of the phosphite moieties and the coordinated water molecule (Table S1, Fig. S5a). The nine-coordinated potassium cations are surrounded by nine neighboring [K₂O₉] polyhedra, sharing corners with six of them (K-K = 5.497(1)-5.724(1) Å), via one common edge O(1)-O(1) (K-K = 3.606(1) Å) and via two face-shared arrangements (3.640(1) Å). The volume occupied by the K⁺ ions was analyzed using the ToposPro 5.0 program30 by means of Voronoi–Dirichlet polyhedra (VDP)31, obtaining a value of 270.62 Å³, which supposes the 14.65% of the cell total volume (Fig. S5b).

From a topological point of view, the double-layered honeycomb sheets of KMnCP can be represented as a binodal (3,4)-connecting net with a point symbol of {6₁³}{6₁}, in which Mn(1) is considered as a 4-connecting node and the three crystallographically independent phosphorous atoms are 3-connecting nodes. After this simplification a known topology type 3,4L147 is obtained (Fig. 2). Such underlying net is not very frequent. In fact, it has only been reported in three compounds, such as in a Cd(II) coordination polymer32 as an interpenetrated topology and in two lanthanide MOFs33 as not interpenetrated frameworks (Fig. S6). As can be seen in the comparative view of Fig. 2, KMnCP displays hexagonal graphite-like honeycomb double-layers stacked in an ABAB sequence where the double-layers are related by a binary screw axis, in contrast with the lantanide MOFs that show a double-layered pattern made of irregular hexagons.
Crystal growth

The predicted equilibrium morphology of \textit{KCMnP} shown in Fig. 3, is a hexagonal prism with (001) basal face and (hk0) prismatic faces with truncated edges. Experimentally, two different crystal habits can be distinguished. The major fraction of crystals displays hexagonal plate habits. These plate crystals are accompanied by some hexagonal prism morphologies, whose (001) faces exhibit clearly smaller areas which is in good agreement with a preferential growth of the (hk0) faces along the c-axis direction. According to the Gibbs-Curie-Wulff theorem\textsuperscript{34}, the surface free energies play essential roles in the crystal growth processes. Since the main habit are hexagonal platelets, the (001) basal planes have the lowest surface energy and the slowest growth rate, while the six lateral planes grow faster with an equivalent rate.

Although length to width proportion of the crystals generally increases with crystallization temperature, the final size and shape of the crystals formed at a given temperature depend on many parameters that affect the values of the apparent activation energies, and thus the growth rates of different crystal faces. As a symmetry-based method, BFDH produce valuable information to limit the number of possible growing faces to be studied by the more expensive and demanding energy-based approaches.\textsuperscript{35} Nevertheless, some computational studies on quinoline derivatives establish that symmetry constraints of the more symmetric systems make BFDH predictions more realistic than in lower symmetry ones\textsuperscript{36}. Since \textit{KCMnP} crystallizes in a high symmetry space group (P6\textsubscript{3}/m), we assume that our morphology prediction based on BFDH laws is a realistic approximation in a good agreement with the as-grown crystals observed in Fig. 3.

J. Meng et al.\textsuperscript{37} studied the synergistic effect between vanadium oxide layer surface configurations and the K ions in terms of electrochemical stability. While the layer surface of KV\textsubscript{3}O\textsubscript{8} is composed of single- and tri-connected oxygen atoms, all of them on the layer surface of K\textsubscript{2-x}V\textsubscript{2}O\textsubscript{5} are single-connected, which can bond and provide strong interaction with K ions. So, more strongly bonded K ions in the interlayers act as pillars to maintain the structural integrity. In our case, the oxygen atoms that establish bonds with the potassium ions on the double-layered sheets surfaces are di-connected (O(1), O(2) and O(3)) to the manganese and phosphorous atoms and single-connected in the case of the coordinated water molecule. In terms of crystal growth dynamics potassium layers should be the most probable endings for basal faces, in such a way that the crystal growth is preferential in \textit{a} and \textit{b} axis (Fig. S7) as can be seen in the preferential crystallization as hexagonal plates.

Structural classification

In order to better understand the metal-phosphite-oxalate structural chemistry, a crystallo-chemical revision of the crystal architectures constructed from metal phosphites containing HPO\textsubscript{3} groups and oxalate bridges have been carried out (22 archetypes found at August 10\textsuperscript{th}, 2017 in the Cambridge Structural Database (CSD)\textsuperscript{38}). The structures have been analyzed based on their metal phosphite framework, as well as by the role of the oxalate anion. Several common characteristics have been found:

i) All the metal-phosphite oxalate frameworks have negative charge needed to be compensated by a cationic inorganic or organic template.

ii) The metal-phosphite substructures can be described based on seven different Structural Building Units (SBUs) showed in Fig. 4. Each SBU has been denoted taking into account the number of metal polyhedra and phosphite groups.

iii) The most common SBU observed in more than the half of the crystal structures is the M\textsubscript{2}P\textsubscript{2}-cycle, the basic building block for several observed chains, layers and even some 3D inorganic frameworks.

iv) Depending on the connectivity between a single or multiple SBUs, 1D to 3D inorganic scaffolds have been found.

v) The oxalate group can act in a mono-bidentate mode, chelating the metal centers, or in a bis-bidentate fashion acting as a bridge between the metal centers of the inorganic framework.

vi) As a general rule, the connectivity of the inorganic substructures through the oxalate ligand gives rise to an increase of the crystal structure dimensionality. This is reflected in the high number of 2D and 3D structures in comparison with the 1D or 0D ones.
Table 2 summarizes the formula, SBU unit, oxalate role and a brief description of the crystal structure. In the last column of the table the compounds have been classified according to the dimensionality of the inorganic metal phosphite skeleton and the role of the oxalate in the structures. (follow Fig. 5 for structural classification and Fig. S8 for structural visualization of the analyzed compounds). A two letter based code has been assigned to each structure, MP\textsubscript{x}O\textsubscript{y}. MP stands for metal-phosphite, and O for oxalate. \(x\) defines the dimensionality of the inorganic metal phosphite net and \(y\) the dimensionality of the complete inorganic-organic net considering the connectivity of the metal centers through the oxalate bridges. Based on the proposed MP\textsubscript{x}O\textsubscript{y} code, nine crystal architectures depicted in Fig. 5 have been identified.

There is only an example of MP\textsubscript{0}O\textsubscript{0} discrete clusters, which can be also described as M\textsubscript{2}P\textsubscript{2} cycles with oxalate groups chelating the metal centers. The extension along one direction of the M\textsubscript{2}P\textsubscript{2} cycles found in the abovementioned zero-dimensional metal oxalatophosphite gives rise to ladder-like MP\textsubscript{1}O\textsubscript{1} chained structures.

The MP\textsubscript{0}O\textsubscript{2} archetype presents a honeycomb-like layer usually found in transition-metal oxalates. In the special case of the unique compound found within this group, the metal-oxalate net is partially interrupted by phosphite linkages. The interlinking of the ladder-like M-O-P chains through bis-bidentate oxalate ligands in-plane fashion form 2D anionic layers found in MP\textsubscript{1}O\textsubscript{2} compounds.

When the connectivity of the inorganic chains is given by oxalate bidentate bridges in- and out-of-plane fashions a three dimensional crystal structure can be formed, such as in the MP\textsubscript{1}O\textsubscript{3} crystal architectures.

Metal phosphite layers are commonly observed in inorganic-organic phosphites, but the role of the oxalate group determines the effective connectivity of the 2D sheets in 3D arrangements. For MP\textsubscript{2}O\textsubscript{2} the oxalate acts commonly as monobidentate units projected on both sides of the sheets, preventing the pillaring of the inorganic layers through covalent bridges. A special case of MP\textsubscript{2}O\textsubscript{2} architecture is the title compound, in which the oxalate units act as bis-bidentate ligand, but their connectivity is limited to two adjacent layers, giving rise to the doubled inorganic-organic substructure described above.

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**Fig. 4** (a) Polyhedral representation of the different SBUs found in metal oxalatophosphites. M and P represent the number of metal polyhedra and HPO\textsubscript{3} units respectively. (b) Distribution of the dimensionality and the various SBUs found in the classified compounds.

**Fig. 5** Schematic representation of the different structural archetypes observed in metal oxalatophosphites with anionic frameworks.
When the connectivity of the bis-bidentate oxalate ligands is extended through all the inorganic layers a MP2O3 crystal architecture is generated. The nature of the metal-phosphite inorganic layers found in MP2O2 and MP2O3 type structures is quite diverse, and can be formed by corner shared M2P2 cycles, M3P2 dimmers, M4P4 clusters or even corner linked M5P6 cages. When the inorganic framework is three dimensional, MP2O3, it is not possible a dimensionality increase through the oxalate groups, but they act as organic pillars connecting metal centers within the inorganic substructure.

Through this literature little review and subsequent classification we have shown the different roles that the oxalate anion adopts in the formation of different structural

| Table 2 Metal oxalatophosphites with anionic frameworks. Metal-phosphite inorganic SBUs together with the role of the oxalate to get the dimensionality of the final archetype |
|--------------------------------------------------|------------------|------------------|------------------|
| Compound | Metal-HPO3 SBU | Oxalate | Class. | Ref. |
| (C2N2H12)2[Mn(HPO3)2(C2O4)2]·4H2O | M3P2 cycles | 0-dimensional | Mono-bidentate terminal unit | MP2O3 | [5a] |
| (C2N2H12)2[Fe(HPO3)2(C2O4)2]·2H2O | M3P2 ladder-like chains formed by corner linked M3P2 cycles | 1-dimensional | Mono-bidentate terminal unit | MP2O3 | [9a,b] |
| (C2N2H12)2[Ga(HPO3)2(C2O4)2]·2H2O | M3P2 ladder-like chains formed by corner linked M3P2 cycles | 1-dimensional | Bridges metal centers of adjacent inorganic chains | MP2O3 | [9b] |
| (C2N2H12)2[Mn(HPO3)2(C2O4)2]·3H2O | M3P2 chains linked through M3P2 cycles | 1-dimensional | Chelates metal centers in both sides of the layers | MP2O3 | [5c] |
| K2[Mn(H2O)2(C2O4)(HPO3)2] | M3P3 hexagonal layers | 2-dimensional (double layers) | Bridges metal centers of two adjacent single layers | MP2O3 | This Work |
| (C2N2H12)2[In2(HPO3)2(C2O4)3] | M3P2 chains generated from corner linked M3P2 cycles interconnected by phosphite units | 2-dimensional (double layers) | Dual role bridging adjacent layers in a bilayer arrangement and acting as terminal units | MP2O3 | [5c] |
| (C2N2H12)2[Mn2(HPO3)2(C2O4)2]·4H2O | M2P2 chains | 3-dimensional | Each chain connects four adjacent ones through oxalate bridges | MP2O3 | [8a,b] |
| (C2N2H12)2[Mn(HPO3)(C2O4)] | M2P2 chains | Each chain connects four adjacent ones through oxalate bridges | MP2O3 | [7a] |
| (C2N2H12)2[Fe(HPO3)2(C2O4)2]·2H2O | M2P2 chains | 3-dimensional | Connects metal centers of adjacent layers | MP2O3 | [8a] |
| (C2N2H12)2[Ga(HPO3)2(C2O4)2]·2H2O | M2P2 clusters | 3-dimensional | Connects metal centers of adjacent layers | MP2O3 | [9b] |
| (C2N2H12)2[Ga(HPO3)2(C2O4)2]·2H2O | M2P2 clusters | 3-dimensional | Acts as an intralayer connector of metal centers | MP2O3 | [5d] |
| (C2N2H12)2[Mn(HPO3)(C2O4)] | M2P2 clusters | 3-dimensional | Acts as an intralayer connector of metal centers | MP2O3 | [8a] |
The thermodiffractometry in air of calculated weight loss. Calcination product are due also to an oxidation process extra peaks observed in the diffractogram of the TGA bond (Fig. S.11(b)). The P-O bonds vibrations of such phosphite (Fig. S.9(b)). The calculated weight loss due to decomposition with some other less intense peaks of an unidentified phase the decomposition. In parallel, the studied compound was observed weight loss was only 20.74%. This discrepancy can be calcined at 800ºC in argon atmosphere obtaining KMnPO 4 as pure phase (Fig. S.9(b)). So it is possible that the unidentified system. Terminal oxalates are found mainly in zero- and one-dimensional structures. The connection of chains trough bis-bidentate oxalate units in an in-plane fashion gives rise to two-dimensional layers. The stacking of the layers by oxalate moieties out-of-plane can complete the three-dimensional connectivity or generate a special structural feature as the double-layered two-dimensional structure found in the studied compound, KMnCP.

Thermal study

The thermal decomposition can be described as a three-step process. The initial weight loss (~8.2%) observed between 175 ºC and 300 ºC corresponds to the departure of two coordinated water molecules (7.63% calculated) per unit formula. From this point to 350 ºC an abrupt weight loss (~7.9%) associated with the thermal breakdown of some oxalate moieties is observed. Such process is marked by a sharp exothermic peak at 335 ºC on the DSC curve. The third step of weight loss (~4.2%), in the 350-500ºC temperature range, follows ascribed to the oxalate units release. Finally, up to 800ºC only 0.5% of mass is lost (see Fig. S.9(a)). The final decomposed product of KMnCP identified by powder XRD was KMnPO 4 (S.G. P-1 (2), a= 5.4813(5) Å, b= 8.6274(10) Å, c= 8.8865(13) Å, α= 87.728(10)º, β= 89.101(10)º, γ= 88.009(10)º) with some other less intense peaks of an unidentified phase (Fig. S.9(b)). The calculated weight loss due to decomposition of water molecules and oxalate unit is 26.27%, but the observed weight loss was only 20.74%. This discrepancy can be explained by considering the oxidation of P(III) to P(V) during the decomposition. In parallel, the studied compound was calcined at 800ºC in argon atmosphere obtaining KMnPO 4 as pure phase (Fig. S.9(b)). So it is possible that the unidentified extra peaks observed in the diffractogram of the TGA calcination product are due also to an oxidation process supporting the difference between the observed and the calculated weight loss.

The thermodiffactometry in air of KMnCP shows that the diffraction peaks of the phase remain unchanged until 225 ºC, then decreasing in intensity until disappearing at 285 ºC (Fig. S10). So the crystalline structure of the compound collapses due to the coordinated water molecule decomposition, as observed in the TGA study.

Infrared, Raman and UV-Vis spectroscopy

A vibrational spectroscopic study was carried out to advance in a better knowledge in the structure of KMnCP (Fig. S11 and Table S2). IR spectrum confirms the existence of the (HPO 4 ) 3- groups by the presence of three sharp bands situated at 2402 cm -1, 2344 cm -1 and 1018 cm -1 (IR and Raman) corresponding to the stretching vibrational mode and deformation of the P-H bond (Fig. S.11(b)). The P-O bonds vibrations of such phosphite moieties are found at frequencies below 1100 cm -1 (see Fig. S.11(a)). The comparison of the IR and Raman vibrational spectra (Fig. S11) shows that the differences in the intensities of the bands and the shifts in their positions are considerably more pronounced for the vibrational modes of the organic ligand, oxalate, than for the inorganic part represented by the phosphate units. For instance, the ν s(C-O) vibrations of the oxalate group are found at the same energies, 1714 (Raman: 1713 cm -1 and 1635 cm -1), in IR and Raman spectra, but the intensity in 1714 cm -1 is significantly stronger in the IR spectrum in comparison with Raman signal, which is very weak. Additionally, ν s(C-O) vibrations (IR: 1360 cm -1, 1318 cm -1, Raman: 1480 cm -1, 1440 cm -1) are shifted around 120 cm -1 to higher energies in Raman counterpart. And the same phenomenon is observed for the δ(OCO) vibrational mode, in this case with a lower shift of around 50 cm -1 (see Fig. S.11(a) and Table S2). Finally, The ν(O-H) stretching vibration of the coordinated water molecule appears as a very strong absorption centered at 3275 cm -1 (IR) (Fig. S.11(b)), which is consistent with the relatively strong character of the hydrogen bond inferred from the structural analysis, where the O-H···O distances are around 2.67 Å. And furthermore, the weak bands situated around 748 cm -1 and 723 cm -1 (IR) could be tentatively associated to the librational modes of the coordinated water in accordance with a previous study on a dihydrogenated iron(II) oxide where the 755/714 cm -1 doublet disappears after deuteration.40

The diffuse reflectance spectrum exhibits three weak spin-forbidden d-d bands, at approximately 403, 437 and 530 nm similar to those of other octahedrally coordinated Mn 3+ complexes which correspond to transitions from the 4A 2 (S) ground state to 4E(G), 4T 2 (G) and 4T 1 (G) terms (Fig. S12(a)). The Dq and Racah parameters for an octahedral d 7 high-spin system cannot be calculated because the frequencies of the transitions to 4T 1 (D) or 4E(D), needed for the calculation and usually found at higher energies, are strongly forbidden and do not appear in the spectrum.

Absorption (K/S) data were calculated from the following Kubelka-Munk function:42

\[ F(R) = \frac{(1 - R)^2}{2R} = \frac{K}{S} \]  

with R representing the reflectance, K the absorption, and S the scattering. In a K/S vs E (eV) plot (Fig. S12(b)), extrapolating the linear part of the rising curve to zero provides the onset of absorption. The value of energy band gap calculated is 3.08 eV.

Electronic structure

The electronic structure of KMnCP was studied systematically through the DFT calculation of the band structure and total density of states (TDOS) (Fig. S13). The distance from the top of the valence band (VB) and bottom of the conduction band (CB) determines the energy necessary for an electron to move from VB to CB which is known as band gap. The band structure of the compound reveals the direct band gap with the top of the VB and bottom of the CB at Γ point. The CB electrons are contributing more at the Fermi level than the VB electrons, which indicates that the crystal belongs to p-type semiconductors with the narrow band gap of 0.027 eV (0.73 eV). Further insight into the electronic structure we examined the partial density of states (PDOS) for all the elements: Mn, P, K, C, H, and O (Fig. S14). In PDOS, the
significant contributions of top of the VB are from Mn-3d and O-2p states which confirm the hybridization between p-d states of Mn-O atoms. Similarly the VB composed of P-2p and O-3p is in the range of -0.2 to -0.4 Ha and -0.8 to -0.7 Ha, K-2p and O-2p mixing is from -0.49 to -0.54 Ha, C-2p with O-2p from 0.4 to -0.5 Ha and H-1s and O-2p hybridization is found from 0.7 to -0.8 Ha. The spins in the bottom of the CB is mostly composed of O-2p states with a small contribution of C-2p states. Hence the Mn-3d states act as electron acceptors which show more contribution at VB region near at fermi level and the O-2p as electron donors. Hence, the optical activity of the crystal is originated by the electronic transition from Mn-3d to O-2p states.

Magnetic behaviour

The temperature dependence of the molar magnetic susceptibility ($\chi_m$) measurement was performed on powdered sample of KMnCP from 250 to 5 K, in the Zero Field Cooling (ZFC) mode at 1 kOe. The thermal evolution of $\chi_m$ follows the classical Curie–Weiss law above 13 K, with Weiss temperature $\theta_w = 20.6$ K and Curie constant $C_m = 4.80$ emu·K/Mn·mol·Oe. The effective paramagnetic moment ($\mu_{eff}$) value calculated for the paramagnetic region, $\mu_{eff} = 2.828(C_m)_{1/2}$, gives the value of 6.19 $\mu_B$/Mn$^{2+}$ ion, very close to the theoretical one ($\mu_{theo} = 5.91$ $\mu_B$) obtained from $\mu_{eff} = g[S(S = 1)]^{1/2}$, considering high-spin $S = 5/2$ Mn$^{2+}$ cations with no orbital contribution.

$\chi_m$ increases from 250 K as temperature decreases describing a smooth inflection point at low temperature. The negative $\theta_w$ value, together with the reduction of the effective magnetic moment ($\chi_{mT}$) observed when the temperature is decreasing (lower inset of Fig. S.15) suggests global antiferromagnetic interactions. ZFC and subsequently FC measurements were also carried out under 100 Oe at low temperatures (upper inset of Fig. S.15). The curves do not show a clear maximum but an inflection point at around 10K, defining the Neel temperature. Moreover, no irreversibility is observed, discarding the existence of any ferromagnetic interaction.

Taking into account the crystal structure, the most probable magnetic exchange pathways due to the Mn···Mn distances and number of atoms involved in the paths are:

i) $J_1$ (Mn-O-P-O-Mn): within the manganese-phosphite single inorganic layers, the Mn(ll) cations are connected through phosphite pseudo-tetrahedra defining a hexagonal honeycomb Mn(ll) layer. Mn···Mn intralayer distances takes three different values, 5.484, 5.513 and 5.878 Å. In an ideal honeycomb layer, the Mn(ll) atoms would be spaced by the same distance, but the inorganic layers of the studied compound show a distortion in which the Mn(ll) cations are grouped in triangular clusters with the abovementioned Mn···Mn spacing (Fig. S16).

ii) $J_2$ (Mn-O-C-O-Mn): manganese cations of each individual inorganic layer are connected through the oxalate organic ligand to the adjacent one, defining Mn-Ox-Mn interlayer dimmers with Mn···Mn bond distances of 5.725 Å.

$J_1$ and $J_2$ involve very similar Mn···Mn distances, and the same number of atoms in the magnetic pathway, so most probably both of them are effective within the same range of temperatures. Magnetic exchange through potassium cations ($J_3$) is expected to be less effective at high temperatures, with Mn···Mn distances of 6.373 Å, but short enough to be involved in a magnetic ordered structure at low temperatures.

So, bearing the crystallographic information in mind we have tried to fit the magnetic data to a honeycomb hexagonal Heisenberg layer model, and to a manganese dimmers model taking into account an intermolecular $J'$ exchange. These models are near to the experimental data, but far from describing a reasonable model from the obtained $J$ values. For both attempts, $g$ values near 2 are obtained, in good agreement with the isotropic electronic nature of high-spin $S = 5/2$ Mn$^{2+}$ cations. The positive $J$ exchange value result from the honeycomb layer Heisenberg model fitting (Fig. S17) is opposite to the antiferromagnetic behavior observed in the decreasing $\chi_m$ curve (lower inset of Fig S.14).

When an intra- and inter- dimeric magnetic exchange is considered, $J = -0.97(1)$ K and $J' = -4.28(10)$ K values are obtained. In an ideal intra- and inter-dimmer exchange, $J$ would be at least one order of magnitude greater than $J'$, but in the specific case of KMnCP, the $J_1$ and $J_2$ exchange are too similar to be considered as intra- and inter-dimmer contributions. Considering only the intra-dimmer magnetic exchange in the fitting a value of $J = -1.63(2)$ K is obtained, and the fitting is not too far from the experimental data. It is clear that the $J$ values obtained from these dimer based models are not accurate, but the negative values obtained for the fittings are in good agreement with the antiferromagnetic behavior derived from the Weiss temperature.

There is a need to seek a more complex theoretical model or neutron diffraction experiments to obtain a correct magnetic model. In fact the Neel ordering suggested by the magnetic susceptibility is not trivial. Examples of the great variety and complexity of magnetic interactions and possible antiferromagnetic orderings in this type of hexagonal systems are the metal-oxalate layered coordination compounds, the inorganic metal halides or some transition metal oxides showing centered hexagonal honeycomb metal layers. In fact, the complexity of the magnetic exchange/ordering in the title compound could be even greater, giving the fact that the single honeycomb layers are connected also by the oxalate groups, and that the magnetic exchange through the potassium cations could become effective also at low temperatures.

Conclusions

In summary, we have reported the synthesis and characterization of a novel 2D manganese(II) phosphite with anionic framework containing oxalate bridges. BFDH analysis turned out to be a realistic approximation to compare the predicted morphology with the hexagonal plates and prisms morphologies experimentally observed.

The role of the oxalate generally determines the final dimensionality of the archetype. In that sense, the special bilayer arrangement observed in the structural study of KMnCP is consequence of the connection of two single sheets
through bis-bidentate oxalate units in an out-of-plane fashion. The three-dimensional connectivity observed in other compounds, is disrupted in this case by the interlayer spaces occupied by the potassium counter ions. The manganese octahedra join in-plane with 3-connecting HPO$_4$ groups and out-of-plane with other metal octahedron through the oxalate unit giving rise to the double-layered honeycomb sheets with not very frequent 3,4L147 topology, previously observed in a coordination polymer and two metal-organic frameworks.

Such special features have facilitated us to devise the structural chemistry of the metal oxalatophosphates with anionic frameworks. Several Secondary Building Units (SBUs) formed by metal-phosphate substructures and different roles of de oxalate bridges have been described promoting a new structural classification for the mentioned compounds. M$_2$P$_2$-cycle is the most common basic building block observed in more than the half of the crystal structures. The connectivity of the inorganic substructures through the oxalate ligand generally gives rise to an increase of the crystal structure dimensionality, which is obvious from the prevalence of 2D and 3D structures in comparison with mono-dimensional or supramolecular ones.

With a broader view, the crystal-chemical analysis here developed could be extrapolated for other tetrahedral (phosphate, arsenate, sulfate or vanadate) or pseudotetrahedral (selenite, tellurite or sulfite) metal hybrid phases containing short organic linkers.

Finally, KMnCP displays major antiferromagnetic interactions. Neutron diffraction experiments to obtain a correct magnetic model is needed since magnetic data fitting to honeycomb hexagonal Heisenberg layers model or to a manganese dimmers model are near to the experimental data but far from a real situation.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

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