Dalton Transactions



PAPER View Article Online



Cite this: *Dalton Trans.*, 2025, **54**, 10558

$Eu_3Ta_2ON_5$: an n = 2 Ruddlesden-Popper ferromagnetic oxynitride with long-range ordered anion vacancies

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The new compound $Eu_3Ta_2ON_5$ with an n=2 Ruddlesden–Popper structure has been prepared by solid state reaction between EuN, Eu_2O_3 and Eu_3N_5 at 1200 °C under EuN_2 . It crystallizes in the EuN_3 large group with unit cell parameters EuN_3 and EuN_3 at 1200 °C under EuN_4 . It crystallizes in the EuN_3 and EuN_4 large group with unit cell parameters EuN_3 and EuN_4 and EuN_4 and EuN_4 and EuN_3 and has anion vacancies in central A-site layers of the perovskite blocks, where tantalum is in a square pyramidal coordination polyhedron with composition EuN_4 and europium is 8-coordinated. The perovskite blocks intergrow with EuN_4 layers of the Ruddlesden–Popper structure. Electron Energy Loss spectrum imaging shows a mixed EuN_4 alence with preferred occupation of EuN_4 in the 8-coordinated sites. Magnetic susceptibility measurements are consistent with the formal composition EuN_4 and indicate ferromagnetic order of EuN_4 ions below EuN_4 ions below EuN_4 in the 8-coordinated containing non-magnetic EuN_4 ions EuN_4 ions below EuN_4 ions below EuN_4 in the 8-coordinated containing non-magnetic EuN_4 ions EuN_4 ions below EuN_4 in the 8-coordinated sites. Which is attributed to the covalency-reinforced EuN_4 and EuN_4 Ruddlesden–Popper compounds (EuN_4 in the 8-coordinated by solid sites.

Received 9th May 2025, Accepted 9th June 2025 DOI: 10.1039/d5dt01093d

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Introduction

Simple perovskite oxynitrides $ABO_{3-x}N_x$ -where A is an alkaline earth metal or rare earth metal and B is a transition metalhave been widely investigated because of their electronic and photocatalytic properties.1 There are numerous examples of compounds with different combinations of cations and anion stoichiometries, and some paradigmatic materials are nontoxic pigments La_{1-x}Ca_xTaO_{2-x}N_{1+x}, BaTaO₂N and SrTaO₂N with high dielectric permittivity,3 EuNbO2N that shows colossal magnetoresistance at low temperature, 4 and LaTiO₂N that is a photocatalyst for water splitting.⁵ Compounds with more complex structures are less common, and include magnetic double or triple perovskites such as La₂MnTaO₅N,⁶ $Ba_2MnWO_{4.42}N_{1.58}$ or $Eu_3Ta_3O_{3.66}N_{5.34}$, and polar hexagonal perovskite BaWON2.9 Perovskite related Ruddlesden-Popper oxynitrides $AX(ABX_3)_n$ (X = O,N), formed by intergrowth of n perovskite blocks with a rock salt AX layer, have been reported as a few examples restricted to aluminum, tantalum and niobium at the B sites. Existing n = 1 members for alkaline

earth cations at A sites are Sr_2TaO_3N , $Ba_2TaO_3N^{10}$ and Sr_2NbO_3N . Lanthanide derivatives were initially reported by Marchand for aluminum, R_2AlO_3N (R = La, Sm, Nd), and more recently the tantalum compounds $R_2Ta(O,N)_4$ (R = La, Nd, Sm, Eu) have been discovered. To the best of our knowledge the only reported n=2 compounds are $Sr_3Nb_2O_5N_2^{-11}$ and $Eu_3Ta_2O_3N_4^{-14}$. The strontium niobium compound was prepared using a solid state reaction at room pressure under NH₃ at 1050 °C, and crystallizes in the aristotype space group for Ruddlesden–Popper phases, I4/mmm. The europium tantalum phase was obtained in the form of single crystals by ammonothermal synthesis at 797 °C and 170 MPa. It was described as a europium mixed valence compound of detailed composition $Eu^{3+}_2Eu^{2+}Ta_2O_3N_4$, with a distorted symmetry $P4_2/mnm$ and cell parameters a=5.7278(1), c=19.8149(5) Å.

Reported perovskite oxynitrides of europium show ferromagnetic order of $\mathrm{Eu^{2^+}}~S=7/2$ spins at low temperatures. $\mathrm{EuNbO_2N}$, $\mathrm{EuTaO_2N^4}$ and $\mathrm{EuWO_{1-x}N_2^{15}}$ show Curie temperatures up to 12 K, and Ruddlesden–Popper n=1 $\mathrm{Eu_2TaO_{2.8}N_{1.2}}$ orders below 8 K. 13 Additionally the Nb and W perovskites show colossal magnetoresistances at low temperatures, originating from the coupling between the carrier spins of the transition metal and the localized Eu spins.

Here we report the new n = 2 Ruddlesden-Popper oxynitride $\mathrm{Eu^{2+}}_2\mathrm{Eu^{3+}}\mathrm{Ta_2ON_5}$ that has been prepared by a solid-state reaction between EuN, $\mathrm{Eu_2O_3}$ and $\mathrm{Ta_3N_5}$ at room pressure, under

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Dalton Transactions Paper

N2 at 1200 °C, and shows a different crystal structure and richer Eu²⁺ contents than early reported Eu³⁺₂Eu²⁺Ta₂O₃N₄. 14 The differences in anion composition and crystal chemistry between these two compounds are due to the drastic changes in the synthesis conditions, as Eu³⁺₂Eu²⁺Ta₂O₃N₄ was obtained at much lower temperature and under high pressure conditions, starting with Ta, Eu, NaN3 and NaOH.14 Rietveld refinement of synchrotron X-ray diffraction data and integrated differential phase contrast imaging indicate that tantalum is in square pyramidal coordination analogously to the nitrides $R_3B_2N_6$ (R = La, Ce, Pr; B = Nb, Ta), ¹⁶ resulting from long-range ordered anion vacancies in the central europium layer of the perovskite blocks. The new compound represents the first example of a stoichiometric anion deficient oxynitride perovskite. Magnetization measurements show that Eu²⁺ ions, that are concentrated at the NaCl-type layers, are ferromagnetically ordered below 20 K. The Curie temperature is significantly larger than reported for Eu²⁺ perovskite-related oxides, where T_C has been found to be lower than 10 K. It will be argued that the ordering of magnetic Eu²⁺ and non-magnetic Eu³⁺ ions and the reinforced Eu-N/O-Eu superexchange interactions account for this remarkable observation.

Experimental

Synthetic procedures and chemical characterization

Black Eu₃Ta₂ON₅ samples of 100 mg were prepared by solid state reaction between Eu₂O₃ (Sigma-Aldrich 99.9%), EuN (Materion, 99.9%) and Ta₃N₅ at 1200 °C under N₂ gas (Air Liquide, 99.9999%) using one single treatment of 3 hours at this temperature. Ta₃N₅ was prepared by reaction of Ta₂O₅ (Sigma-Aldrich 99.99%) with NH₃ (Carburos Metálicos 99.9%) at 850 °C during several cycles of 15 hours with intermediate regrinding, using a flow rate of 600 cm³ min⁻¹. Eu₂O₃ was treated at 900 °C under vacuum of 10⁻³ torr during 12 hours for dehydration. The reactants were weighed, mixed and pelletized in a glove box under recirculating Ar. The pellets were placed in molybdenum crucibles covered by zirconium foil. Another crucible with zirconium was placed near the sample in the reaction tube to scavenge H₂O and O₂ impurities from the gas. The reaction tube was evacuated to 10⁻³ torr and purged three times with N₂ before heating at 300 °C h⁻¹ up to the maximum temperature. The obtained samples were stable in air.

Nitrogen analyses were performed in a Thermo Fisher Scientific instrument heating the samples up to $1060~^{\circ}\text{C}$ under O_2 , using MgO, WO₃ and Sn as additives and atropine as reference standard.

Structural characterization

Laboratory X-ray powder diffraction patterns were obtained on capillary samples of diameter 0.3 mm using a Bruker D8 Advance A25 diffractometer in a Debye Scherrer configuration with Mo K α_1 radiation (λ = 0.7093 Å). High resolution synchrotron X-ray powder diffraction (SXRPD) data were collected at room temperature on the same capillaries in the angular range

 $2.0^{\circ} \le 2\theta \le 60^{\circ}$ at the MSPD beamline of the ALBA Synchrotron (Cerdanyola del Vallès, Spain), ¹⁷ using multi analyser detector and 30 keV energy that resulted in a wavelength of 0.41376 Å determined by refining SRM640d NIST Si standard. Rietveld analysis was performed using the program Fullprof. ¹⁸ Background refinement was performed by linear interpolation and data were corrected from absorption.

Electron diffraction patterns were taken in a JEOL 1210 transmission electron microscope operating at 120 kV using a double tilt side entry specimen holder with tilting angles $\pm 60^{\circ}/\pm 30^{\circ}$. The samples were deposited in powder form on a copper grid coated by a Lacey carbon film with a continuous layer of ultrathin carbon film.

Atomic resolution images were obtained at the Joint Electron Microscopy Center at ALBA (Cerdanyola del Vallès, Spain) on a double-corrected Thermo Fisher Spectra 300 (S)TEM microscope operated at 200 kV. High-angle annular dark field (HAADF) images were acquired using a semi-convergence angle of 19.5 mrad. Integrated differential phase contrast scanning transmission electron microscopy (iDPC-STEM) method has also been used for imaging of all elements using the segmented Panther detector from Thermo Fisher. This imaging mode detects the deflection of the beam produced by the atomic electrostatic fields thus allowing to image simultaneously heavy and light atoms. 19 Electron Energy Loss Spectroscopy (EELS) spectra were collected in a continuum spectrometer equipped with a fast direct electron detection camera K3 from Gatan, using a collection angle of around 40 mrad, a probe current of around 90p A and a dwell time of 50 ms. A principal component analysis filter was used after acquisition to minimize the random noise of the EELS spectrum images. To improve the energy resolution, the beam was monochromated using an excitation of 0.6, a spot size of 14 and a 1µm diameter C1 condenser aperture. This leads to a zero-loss peak full width at half maximum of around 0.4 eV. EDX spectra were acquired using a four quadrant Super-X windowless silicon drift detector system and beam currents of ≈100-250 pA. The crystal composition was quantified using the Thermo Fisher Scientific Velox software, applying the Cliff-Lorimer approach with Brown-Powell ionization cross-section models.

Magnetic measurements

Magnetic susceptibility measurements were performed on a Quantum Design SQUID magnetometer at temperatures between 2 and 300 K and magnetic fields of 25 Oe and 10 kOe. Magnetization-field loops were measured between -70 kOe and +70 kOe at temperatures below 50 K.

Results and discussion

Synthesis and structural study

The synthesis of the new anion deficient n = 2 Ruddlesden–Popper oxynitride takes place through the formal reaction:

$$2 Eu_2O_3 + 5 EuN + 2 Ta_3N_5 \rightarrow 3 Eu_3Ta_2ON_5 + 3/2 O_2$$

Paper **Dalton Transactions**

that involves an N/O ratio of 2.5 in the mixture of reactants. Under the synthesis conditions at 1200 °C in N2, 2/3 of europium is reduced from +3 to the +2 oxidation state, with concomitant oxidation and loss of one O²⁻ anion per formula. The initial N/O ratio was tuned by changing the proportion of Eu₂O₃ and EuN in the mixture, until the optimal ratio $Eu_2O_3 : EuN : Ta_3N_5 = 0.45 : 8.1 : 2$ with N/O = 13.4 produced a sample free of impurities according to synchrotron X-ray powder diffraction data (Fig. 1).

Syntheses starting with N/O ratios below this value led biphasic samples with different proportions of the n =1 member Eu₂TaO_{2,37}N_{0.63}. This phase is prepared at the same temperature but has a lower N/O stoichiometric ratio compared to Eu₃Ta₂ON₅, ¹³ hence the increase of N/O favours the formation of the n = 2 member. In the biphasic samples, Eu₂TaO_{2,37}N_{0,63} was detected either in the X-ray diffraction patterns or as small intergrowth domains in the scanning transmission electron microscopy (STEM) images.

Electron diffraction patterns of Eu₃Ta₂ON₅ (Fig. 2) were indexed in a cell with $a \approx b \approx 3.97$ and $c \approx 20.54$ Å in the I4/ mmm aristotype space group, with reflection conditions: hkl, h + k + l = 2n; hk0, h + k = 2n; 0kl, k + l = 2n; hhl, l = 2n; 00l, l = 2n2n; h00, h = 2n. For some crystals, very weak additional reflections were observed in one of the (110) planes, that could be indexed in a $\sqrt{2a_0} \times \sqrt{2a_0} \times c_0$ superstructure with the orthorhombic space group Amaa (no. 66), where a_0 and c_0 are the parameters of the I4/mmm cell.

High-resolution HAADF-STEM imaging in combination with EDX and iDPC allowed mapping of the position of all atomic sublattices in real space. Fig. 3a shows a high-resolution HAADF-STEM image of a Eu₃Ta₂ON₅ crystal with the characteristic n = 2 Ruddlesden-Popper structure. EDX maps (Fig. 3b) showed no Eu/Ta intermixing, giving a ratio Eu/Ta of 2.8(2)/2; the nitrogen content of 7.56% determined by combustion analysis indicated 4.88 atoms per formula, close to the formal stoichiometry Eu₃Ta₂ON₅ where the oxygen content has

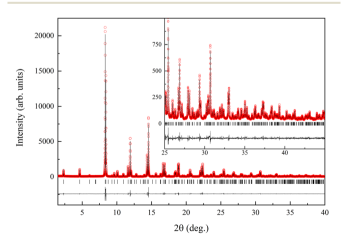


Fig. 1 Rietveld fit to synchrotron X-ray powder diffraction data (λ = 0.41376 Å) of Eu₃Ta₂ON₅, performed in the I4/mmm space group with a = 3.98240(1), c = 20.42020(6) Å. The inset shows the high 2θ region

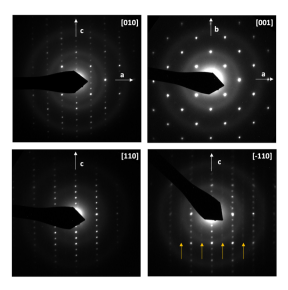


Fig. 2 Electron diffraction patterns of Eu₃Ta₂ON₅ along [010], [001] and [110] axes. Orange arrows indicate superstructure reflections observed in some crystals, that can be indexed in a $\sqrt{2a_0} \times \sqrt{2a_0} \times c_0$ unit cell.

been adjusted considering 6 anions. EDX spectra also indicated the presence of oxygen and nitrogen in the sample, but their quantitative analysis was not possible because of the large error observed for these light elements.

In order to probe the O and N sublattices, iDPC-STEM imaging mode was used, which is a direct phase imaging process that relies on atomic number. As the atomic number increases, the scattering intensity and signal brightness also rise. This imaging principle results in a linear relationship between the intensity in iDPC images and the atomic number. 19 In contrast, high-angle annular dark field (HAADF) images exhibit an approximate proportionality between intensity and the square of the atomic number. Consequently, iDPC-STEM offers greater information about lighter elements, which is highly valuable for investigating the O and N sublattices of Eu₃Ta₂ON₅. Notice that the iDPC contrast of the image of a crystallite oriented along [010] allows for the identification of all sublattices in Eu₃Ta₂ON₅. Remarkably, the planes signalled with an arrow show a much dimmer contrast due to an anion deficiency, consistent with a n = 2 Ruddlesden-Popper structure with 6 anions per formula and vacancies located between the two [TaX2] planes of the perovskite block, resulting in square pyramidal coordination of tantalum atoms.

Rietveld refinement of synchrotron X-ray powder diffraction data (Fig. 1) were performed in the I4/mmm space group starting with the structural model of La₃Ta₂N₆ that has two crystallographically independent anions at 8g and 4e sites, corresponding respectively to equatorial and axial positions of the TaX_5 pyramids. The refined cell parameters were a = 3.98240(1), c = 20.42024(6) Å, with agreement factors of $R_p = 6.90\%$, $R_{\rm wp} = 8.95\%$, $\chi^2 = 6.35$ and $R_{\rm Bragg} = 1.97\%$ (Table 1 and Fig. 4). Refinements were also performed in the Amaa (no. 66) space group using a unit cell with parameters $\sqrt{2a_0} \times \sqrt{2a_0} \times c_0$ and three positions for the anions, but the observed Bragg R factor

Dalton Transactions Paper

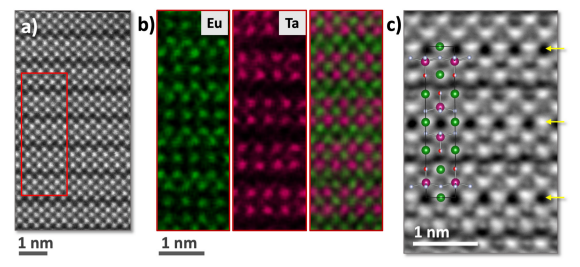


Fig. 3 (a) HAADF-STEM image of a $Eu_3Ta_2ON_5$ crystal along the [010] zone axis. The image shows an n = 2 Ruddlesden-Popper structure, with two perovskite blocks intergrowing with a rock salt-type layer. The inset shows the corresponding area for the elemental EDX maps. (b) From left to right, atomic resolution EDX maps of Eu (in green), Ta (in magenta) and a combined colour map. (c) iDPC image with the Eu₃Ta₂ON₅ structure overlay (Eu: green, Ta: magenta, O: red and N: light gray). Yellow arrows point to the planes in which the anion vacancies are located.

Table 1 Atomic coordinates in the I4/mmm space group, cation and anion occupancies and isotropic temperature factors for Eu₃Ta₂ON₅ from the refinement to synchrotron X-ray powder diffraction data at 300 K using $\lambda = 0.41376 \text{ Å}^{a}$

Atom	Wyckoff site	х	у	z	B^b (Å ²)	Occ. factor
Eu1	2b	0	0	0.5	0.44	1
Eu2	4e	0	0	0.31422(2)	0.65	1
Ta	4e	0	0	0.09922(3)	0.47	1
N1	8g	0	0.5	0.0738(3)	1.04(9)	1
O2/N2	4e	0	0	0.1910(3)	1.04	0.5/0.5

Bond	length	(Å)
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Ta-N1	$2.0595(6) \times 4$	Ta-O2,N2 1.865(6)
Eu1-N1	$2.493(4) \times 8$	
Eu2-N1	$3.036(5) \times 4$	Eu2-O2,N2 2.525(6)
Eu2-O2,N2	$2.8177(2) \times 4$	

^aO/N occupation factors were fixed considering the predicted distribution using Pauling's second crystal rule. Refined cell parameters were a = 3.98240(1), c = 20.42024(6) Å. Agreement factors: $R_p = 6.90\%$, $R_{wp} = 8.95\%$, $\chi^2 = 6.35$, $R_{Bragg} = 1.97\%$. ${}^bB_{eq}$ in Å² is calculated for Eu and Ta atoms from refined anisotropic temperature factors (×10⁴): Eu1 $\beta_{11} = \beta_{22} = 64(3)$ $\beta_{33} = 3.1(2)$; Eu2 $\beta_{11} = \beta_{22} = 120(4)$ $\beta_{33} = 2.5(2)$; Ta $\beta_{11} = 3.1(2)$ $\beta_{22} = 59(2) \beta_{33} = 3.9(1)$. The temperature factors were common for the two anion sites.

was significantly larger than for the I4/mmm model (5.06% vs. 1.97%).

The europium atoms in Eu₃Ta₂ON₅ show two well differentiated polyhedra with coordination numbers 8 and 9 for Eu1 at 2b and Eu2 at 4e sites respectively, and corresponding average bond distances of 2.493(4) and 2.871 Å. Considering the difference between both distances and the ionic radii of Eu^{3+} for CN = VIII (1.01 Å) and Eu^{2+} for CN = IX (1.30 Å), ²⁰ the oxynitride can be formally formulated Eu²⁺₂Eu³⁺Ta⁵⁺₂ON₅, a charge-ordered compound where Eu³⁺

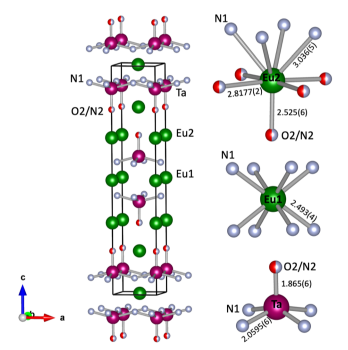


Fig. 4 Structural model in the I4/mmm space group and cation coordination polyhedra of Eu₃Ta₂ON₅ showing bond distances.

occupies the 8-coordinated sites and Eu²⁺ ions are in the 9-coordinated sites.

The distribution of O and N in the 8g and 4e sites could not be determined from X-ray diffraction because this technique does not provide enough contrast between the two anions. Neutron powder diffraction is the optimal technique to investigate the anion ordering in oxynitrides, but for Eu₃Ta₂ON₅ the large absorption cross section of europium

Paper

would make difficult to obtain accurate information from these data. Pauling's second crystal rule (PSCR) can be used to predict the anion distribution in mixed anion compounds, 21,22 from the calculation of the bond strength sums (b) with the equation:

$$b = \sum_i \frac{z_i}{\nu_i},$$

where z_i and ν_i are the electrical charge and coordination number of each cation bonded to a given anionic position.

According to PSCR, the electrical charge of each anion (q) tends to compensate the strength of the electrostatic valence bonds from the cations (b). Considering the ordering of Eu³⁺ at 2b sites and Eu²⁺ at 4e sites, the calculated bond strength sums are 2.85 and 1.94 for the anions at 8g and 4e positions respectively, predicting the preferred occupancy of N atoms at the equatorial positions of the $[TaX_5]$ pyramids. Accordingly, in the refinement we used fixed 100% occupancy of nitrogen on 8g sites and 50/50 N/O at 4e sites.

The observed equatorial Ta–N bond distance (2.0595(6) Å) is similar to those reported for La₃Ta₂N₆ and Ce₃Ta₂N₆ (2.053 (3) and 2.0465(19) Å respectively).¹⁶ In contrast, the Ta–O,N apical distance (1.865(6) Å) is slightly shorter than in the two nitrides (1.96(3) and 1.951(19) Å for La₃Ta₂N₆ and Ce₃Ta₂N₆ respectively). This difference is consistent with the mixed O/N occupancy at axial sites, according with the larger ionic radius of N^{3–} compared to O^{2–} (1.46 vs. 1.38 Å, both for CN = IV).²⁰

EELS measurements provided detailed atomic scale information of the oxidation state of both Eu2 and Eu1 sites by analysing the energy onset of the Eu M4,5 edge. Fig. 5(a-c) shows a HAADF-STEM image of a Eu₃Ta₂ON₅ crystal and two EEL spectra from the crystallographic sites Eu2 and Eu1. The fine structure of the two Eu M4,5 spectra is clearly different, although the source is not as straight forward. For Eu²⁺ species the onset is lower in energy than that of the Eu³⁺ species.²³ If both Eu²⁺ and Eu³⁺ are present, EEL spectra typically display two peaks with the stronger one belonging to the dominant oxidation state. The thickness of the sample may also play an important role, as it widens the electron beam, it lowers the spatial resolution and increases the contribution of nearest unit cells,²⁴ which may be the reason of a presence of a Eu²⁺ signal in the spectra acquired from the Eu1 site, or of Eu3+ in the spectra acquired for the Eu2 site. Nonetheless, these EELS measurements support the preferred occupation of Eu2+ and Eu³⁺ ions in Eu2 and Eu1 sites, respectively, of Eu₃Ta₂ON₅.

Magnetic properties

The temperature-dependent magnetic susceptibility $\chi(T)$ (black symbols) and $\chi^{-1}(T)$ (red symbols) of Eu₃Ta₂ON₅, recorded at H = 5k Oe, are shown in Fig. 6. It can be appreciated that $\chi^{-1}(T)$ displays an appreciable bending, suggesting an additional temperature-dependent contribution other than the Curie–Weiss behavior. As indicated by the solid lines thorough the data, the susceptibility per mole can be well fitted by the contributions from $n_{\text{Eu}^{2+}}$ moles of Eu²⁺ (S=7/2, J=7/2), modelled as a Curie–Weiss law, and a small temperature-

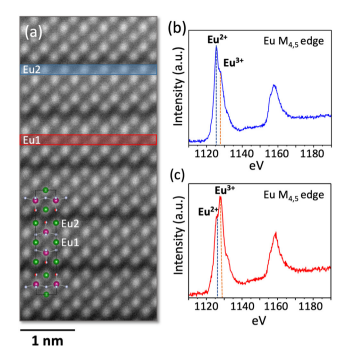


Fig. 5 (a) HAADF-STEM image of the $Eu_3Ta_2ON_5$ crystal along the [010] zone axis with the $Eu_3Ta_2ON_5$ structure overlay (Eu: green, Ta: magenta, O: red and N: light gray). (b) and (c) show the averaged EEL M4,5 spectra obtained from Eu2 and Eu1 sites of the $Eu_3Ta_2ON_5$ structure, marked in blue and red in (a), respectively.

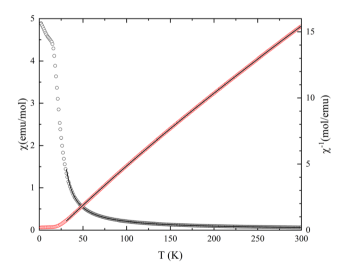


Fig. 6 Magnetic susceptibility $\chi(T)$ (left axis) and the reciprocal $\chi^{-1}(T)$, (right axis) recorded at 5 kOe. Solid lines are the results of the fit described in the text.

dependent Van Vleck contribution arising from $n_{\text{Eu}^{3+}}$ moles of Eu³⁺, in addition to a background contribution (χ_0). Notice that although Eu³⁺ ground state is nonmagnetic ($^7\text{F}_0$, S=3, L=3, J=0), thermal excitation to higher lying states (for instance the first one ($^7\text{F}_1$, S=3, L=3, J=1) is only at about 46 meV²⁵) shall produce an additional contribution to the magnetic susceptibility ($\chi_{\text{Eu}^{3+}}$), that will add to the Eu²⁺ contribution, and to

Dalton Transactions Paper

the diamagnetic contribution. Accordingly, the magnetic susceptibility per Eu ion can be expressed as:^{8,13}

$$\chi(T) = n_{\mathrm{Eu}^{2+}} \chi_{\mathrm{Eu}^{2+}}(T) + n_{\mathrm{Eu}^{3+}} \chi_{\mathrm{Eu}^{3+}}(T) + \chi_0,$$

where:

$$\chi_{\text{Eu}^{2+}}(T) = \frac{N_{\text{A}}\mu_{\text{B}}^2}{3k_{\text{B}}} \frac{g_J^2 J(J+1)}{(T-\theta)} (\text{with } J = \frac{7}{2} \text{ and } g_J = 2),$$

$$\chi_{\mathrm{Eu^{3+}}}(T) = rac{\sum\limits_{J=0}^{6}{(2J+1)\chi_{J}(T)\exp[-\lambda J(J+1)/k_{\mathrm{B}}T]}}{\sum\limits_{J=0}^{6}{(2J+1)\exp[-\lambda J(J+1)/k_{\mathrm{B}}T]}},$$

with:

$$\chi_{J}(T) = \frac{N_{\rm A}\mu_{\rm B}^{2}g_{J}^{2}J(J+1)}{3k_{\rm B}T} + \frac{2N_{\rm A}\mu_{\rm B}^{2}(g_{J}-1)(g_{J}-2)}{3\lambda},$$

and $g_I = 3/2$, except g_0 that equals 2 + L = 2 + S = 5.

Fitting to experimental inverse susceptibility was done by varying $n_{\rm Eu^{2+}}$, $n_{\rm Eu^{3+}}$, θ and χ_0 , while fixing $\lambda/k_{\rm B}=531.5$ K (equivalent to an energy splitting, $\lambda=46$ meV, between the nonmagnetic ground state of Eu³⁺ and its first excited state), 13,26 as well as the effective paramagnetic moment of Eu²⁺. From the fit, we obtain $n_{\rm Eu^{2+}}=1.98$, $n_{\rm Eu^{3+}}=1.02$, $\theta=20.2$ K and $\chi_0=4.6\times 10^{-3}$ emu mol⁻¹. These results are quite consistent with chemical analysis and crystal structure results, that suggest 2 Eu²⁺ and 1 Eu³⁺ per formula. The extrapolated Curie–Weiss temperature is $\theta>0$ (≈20 K), indicating the prevalence of ferromagnetic interactions.

In agreement with the susceptibility data of Fig. 6, the M(H) loops (Fig. 7) develop a ferromagnetic-like shape at $T \leq 20$ K. At 2 K the saturation magnetization is about $14\mu_{\rm B}$ per f. u., which is fully consistent with the presence of two ferromagnetically ordered Eu²⁺ ions per f.u in this oxynitride. The ferromagnetic ordering occurring at $T \approx 20$ K is evidenced in the zero-field-cooling and field-cooling susceptibility data recorded at low field (inset in Fig. 7), that show a perceptible hysteresis.

The most remarkable feature of these results is that the Curie temperature (\approx 20 K) is significantly larger than those reported for similar systems, such as Ruddlesden-Popper n=1 Eu₂TiO₄ and n=2 Eu₃Ti₂O₇, 25,28 where $T_{\rm C}$ is around 9–10 K. To rationalize the observed dramatic enhancement of $T_{\rm C}$ in Eu₃Ta₂ON₅, it is worth to compare these results with isostructural Eu₃Ti₂O₇, that for this purpose is described as a sequence of EuTiO₃ (perovskite-type) and Eu₂TiO₄ (K₂NiF₄-type, rock salt) blocks . We recall that, as argued above, the non-magnetic Eu³⁺ ions in Eu₃Ta₂ON₅ occupy Eu1 sites equivalent to those in EuTiO₃ structure and the magnetic Eu²⁺ ions occupy Eu2 sites as in Eu₂TiO₄. Therefore, the magnetic interactions in Eu₃Ta₂ON₅ are analogous to those for Eu₂TiO₄.

Fig. 8 illustrates the most relevant magnetic nearest-neighbor (nn) (J_{11} and J_{12}) and next-nearest-neighbor (nnn) magnetic interactions (J_{21}) in Eu₃TaON₅. Notice that we omit magnetic interactions involving Eu³⁺ as this ion is nonmagnetic. Here, we follow the nomenclature of Chien *et al.*²⁵ who

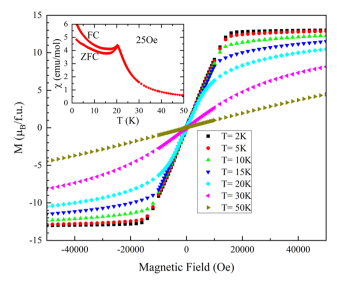


Fig. 7 Magnetization loops recorded at several temperatures (indicated) up to ± 5 T. Inset: zero field-cooled/field-cooled magnetization *vs.* temperature measured under 25 Oe applied magnetic field.

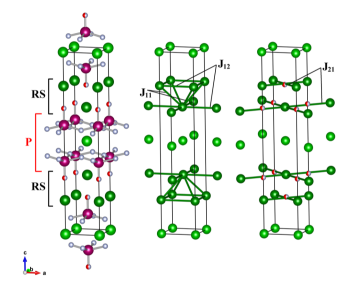


Fig. 8 Sketch of the structure and the most relevant nn $(J_{11}$ and $J_{12})$ and nnn (J_{21}) magnetic interactions in Eu₃Ta₂ON₅. Ta, N and O atoms are represented by magenta, light gray and red spheres respectively; Eu³⁺ and Eu²⁺ are shown as light and dark green spheres respectively. The two structural blocks for Eu are labelled as P, (perovskite) and RS (rock salt, K₂NiF₄-type blocks). For comparison with ref. 25, the structure has been shifted by (0.5,0.5,0) with respect to Fig. 4.

employed a mean field approach to describe the $T_{\rm C}$ of Eu₃Ti₂O₇ in terms of the nn and nnn magnetic interactions (eqn (19) in ref. 25). The model can be easily adapted to the present case by keeping only interactions involving Eu²⁺ ions; namely J_{11} , J_{12} and J_{21} . It follows that:

$$T_{\rm C} = \frac{16S(S+1)(J_{11} + J_{12} + J_{21})}{\kappa_{\rm B}}$$

Paper **Dalton Transactions**

where S (= 7/2) is the Eu²⁺ spin and κ_B is the Boltzmann constant. Structural data indicates that nn Eu-Eu distances in Eu_2TiO_4 (3.814 Å)²⁵ are only slightly shorter (1.3%) than in Eu₃Ta₂ON₅ (3.864 Å). Similarly, the next-nearest-neighbor nnn Eu-O-Eu distance in Eu₂TiO₄ (5.465 Å)²⁵ are also slightly shorter (1.8%) than nnn Eu-(N/O)-Eu distance in Eu₃Ta₂ON₅ (5.566 Å). Therefore, taking the nn interactions J_{11} and J_{12} as in Eu_2TiO_4 ($\approx 0.11 \text{ K}^{25}$) and using the expression above, we derive the nnn Eu-(N/O)-Eu interaction $J_{21} \approx 0.13$ K that is much (×4) larger than J_{21} in Eu₂TiO₄ (0.037 K).

We note that the 4f7 electrons of Eu2+ have well localized wave functions with tiny overlapping with neighboring Eu2+ ions. It follows that the direct 4f-4f interaction is negligible. Instead, as proposed long ago by Goodenough²⁹ and Kasuya,³⁰ intra-atomic 4f-5d intermixing offers a mechanism to propagate magnetic interactions. The 4f-5d intermixing is dictated by the relative position of the 4f and 5d orbitals, the latter being determined by the crystal field. 25,29,30 As the coordination polyhedra of Eu2+ in Eu3Ta2ON5 and Eu2TiO4 have rather similar bond lengths, it can be suspected the 4f-5d mixing in these compounds to be similar. As the nn distances in both compounds differ only by some 1.5%, it follows that the nn Eu-Eu interactions (J_{11}, J_{12}) should be also similar, justifying the assumption made above.

Similarly, as the Eu-Eu nnn distances in both structures are also similar (within 1.8%), we conclude that the reinforced ferromagnetic Eu-Eu interaction results from the fact that the Eu-(N/O)-Eu hybridization is much enhanced. This can be explained by the higher hybridization (lower electronegativity of 2p(N) orbitals compared to 2p(O)) with 4f-5d orbitals of Eu. Enhanced ferromagnetic interactions induced by the large covalency of Mn-N bonds have also been observed in Ba₂MnWO_{4.42}N_{1.58} oxynitride.⁷

In short, the selective occupation of Eu³⁺ and Eu²⁺ in the two A sites of Eu₃Ta₂ON₅ precludes magnetic dilution, that would be present if both cations would occupy the same site, and the reinforced hybridization of Eu-(N/O)-Eu bonds strengthens the ferromagnetic superexchange. On the other hand, this selective occupation implies the absence of bridging magnetic ions in the perovskite block (see Fig. 8). It follows that ferromagnetic order takes place basically within the Eu²⁺containing bilayers of the rock-salt block. Therefore, a natural question arises: if the Eu²⁺ bilayers are magnetically decoupled, why do they appear similarly magnetically aligned in the magnetic measurements shown above? Tiny magnetic interactions between them may exist, either via direct dipolar coupling or via the presence of a small fraction of Eu²⁺ at the perovskite positions. However, none of those mechanisms, nor the combination of them, seem to be compatible with the observed large $T_{\rm C}$ (≈ 20 K). Alternatively, the internal field associated to the Eu^{2+} ordering, may induce mixing of J = 1and J = 0 in the ground state of Eu³⁺ (${}^{7}F_{J}$) mimicking the role of an external field which is responsible for the temperatureindependent paramagnetism of Eu3+, with the consequent induction of a magnetic moment in an otherwise non-magnetic ion. The induced moment in Eu³⁺ may be responsible for the interlayer ferromagnetic coupling, as observed in $Eu_{1-x}Y_xMnO_3$. Therefore, the possibility arises that the two RS blocks of Eu²⁺ in the Eu₃Ta₂ON₅ structure spontaneously order separately but are further aligned due to the magnetic applied field. Elucidating this possibility would only be possible by characterizing the magnetic order at zero field (e.g. by muon spectroscopy).

Conclusions

The new n = 2 Ruddlesden-Popper oxynitride $Eu_2^{2+}Eu^{3+}Ta_2ON_5$ has been prepared by a solid state reaction between EuN, Eu₂O₃ and Ta₃N₅ during 3 hours under N₂ at 1200 °C. It crystallizes in the I4/mmm space group, with cell parameters a = 3.98240(1)and c = 20.42024(6) Å. Contrast inverted STEM iDPC images show long range ordered anion vacancies in the Eu layers between the TaX2 planes of the perovskite blocks, which decrease the coordination number of the transition metal from 6 to 5 anions, analogously to reported rare earth nitrides R₃B₂N₆ (R = La, Ce, Pr; B = Nb, Ta). The anion vacancies generate two different environments for europium, with 8-coordinated Eu3+ located between the TaX₂ planes and 9-coordinated Eu²⁺ in rock-salt layers. The magnetization data of Eu₂²⁺Eu³⁺Ta₂ON₅ shows ferromagnetic order of Eu^{2+} S = 7/2 spins below $T_C =$ 20 K. The selective occupation of Eu³⁺ and Eu²⁺ in the two sites gives rise to Eu3+ magnetically inert perovskite blocks that alternate with rock-salt blocks of ferromagnetic Eu2+ bilayers, where the presence of nitrogen at the anion sites leads to stronger Eu-(N/O)-Eu superexchange interactions and a record ordering temperature. The nitride introduction in the n = 2 Ruddlesden-Popper structure allows electronic and chemical orders and subsequent tailoring of magnetic interactions. New magnetic properties may emerge by further modification of the N/O ratio that certainly will modify the balance between the two europium oxidation states.

Author contributions

J. O. S. and J. R. G.: investigation, formal analysis. B. M.: investigation. C. F. and J. G: investigation, formal analysis, methodology, writing. A. F. and J. F: conceptualization, supervision, writing.

Data availability

The data used in this publication are available from the author on reasonable request. The synchrotron X-ray powder diffraction data are available through the following link https://doi. org/10.20350/digitalCSIC/17261.

Conflicts of interest

There are no conflicts to declare.

Dalton Transactions Paper

Acknowledgements

This work was supported by grants PID2023-146263NB-I00, PID2023-152225NB-I00, Severo Ochoa MATRANS42 CEX2023by MCIN/AEI/10.13039/501100011033 funded (Ministerio de Ciencia e Innovación/Agencia Estatal de Investigación) and FEDER, EU, TED2021-129857B-I00 and PDC2023-145824-I00 funded by MCIN/AEI/10.13039/ 501100011033 and European Union Next Generation EU/PRTR, and grants 2021SGR00439 and 2021SGR00445 funded by the Generalitat de Catalunya. We thank ALBA synchrotron (experiment number AV-2024028275) for the provision of beam time; we also thank Dr François Fauth (ALBA) for assistance during data collection and Dr Bernat Bozzo (ICMAB-CSIC) for performing the magnetic measurements. JRG acknowledges AEI predoctoral fellowship PRE2018-085204. We are thankful for the assistance of ICMAB Scientific and Technological services of X-ray Diffraction, Low Temperature and Magnetism and Electron Microscopy. We acknowledge the Joint Electron Microscopy Center at ALBA (JEMCA) and funding from Grant IU16-014206 (METCAM-FIB) to ICN2 from the European Union through the European Regional Development Fund (ERDF), with the support of the Ministry of Research and Universities, Generalitat de Catalunya.

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