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ARTICLE

Structure and Magnetic Properties of the AB(HCO₂)₃ (A = Rb⁺ or Cs⁺, B = Mn²⁺, Co²⁺ or Ni²⁺) Frameworks: Probing the Effect of Size on the Phase Evolution of the Ternary Formates Received 00th January 20xx. Accepted 00th January 20xx

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This work reports the synthesis and structures of six new AB(HCO₂)₃ (A = Rb⁺ or Cs⁺ and B = Mn²⁺, Co²⁺ or Ni²⁺) framework containing the largest monoatomic cations on the A-site. RbMn(HCO₂)₃ is found to adopt a distorted perovskite framew(with a $4^{12} \cdot 6^3$ topology and a mixture of syn-anti and anti-anti ligands, while the remaining compounds adopt a chiral hexagonal structure with a $4^9 \cdot 6^6$ topology. The structures of these frameworks clarifies the effect of ionic size on 1formation of the five known architectures adopted by the AB(HCO₂)₃ frameworks, which have attracted attention as a new class of potential multiferroics, and in particular the chiral hexagonal structure within this. This also highlights the role of molecular A-site cations in stabilising the 49.66 topology for frameworks where such cations are too large or small to support this structure on the basis of size alone, possibly due to hydrogen bonding. The magnetic properties of ti RbB(HCO₂)₃ and CsMn(HCO₂)₃ frameworks are also reported with the Rb⁺ compounds featuring weak ferromagnetic behaviour and the latter being purely antiferromagnetic. In conjuction with a comparison of the other isostructura' $AB(HCO_2)_3$ frameworks we find that compounds adopting the $4^9 \cdot 6^6$ topology have much higher magnetic ordering temperatures than those with the RbMn(HCO₂)₃ structure, highlighting the importance of understanding the structure property relationships of the ternary formates.

1. Introduction

Coordination frameworks, including dense Metal-Organic Frameworks (MOFs), have attracted significant attention in recent years for their ability to exhibit unique variations on the functional properties traditionally associated with metal oxides including magnetic order, multiferroicity, semi-conductivity and ionic conductivity. 1,2-4 This is due to the varied architectures they adopt because of the structure directing effect of the ligand, even while smaller ligands bring metal centres closer together enabling electronic communication between them. Magnetoelectrical properties include hydrogen-bond driven relaxor ferroelectric order,5,6 guestdependent magnetism^{2,7} and new routes to multiferroic materials.^{3,6,8} Arguably the most intriguing of these compounds are the AB(HCO₂)₃ frameworks, where A is a monovalent alkali metal, ammonium, imidazolium or a protonated alkylamine and B is typically a divalent first row transition metal. 4,6,9,10 Molecular A-site cations in these

The AB(HCO₂)₃ frameworks with three dimensional covalent connectivity all feature octahedral B-site cations linked in a corner-sharing fashion through the formate ligand with the A-site cation typically sitting in pockets found in the structure. Beyond that they can vary significantly with five different structure-types known, which can largely be classified by their topologies and the coordination modes the forma e ligands bind in (see Fig. 1).11 Those compound accommodating the largest A-site cations, usually moderately sized alkylamines, such as $(CH_3)_2NH_2$, or imidazolium, adopt a perovskite-like structure with a 4¹² •6³ topology. ^{11,13} Herein this structure is referred to as Perovskite I, consistent with Gómez-Aguirre et al. 11, and has all ligands binding the B-s e metals in an anti-anti fashion. Two other 4¹² •6³ topologi have been discovered for the AB(HCO₂)₃ frameworks. The first so-called Perovskite II, is the thermodynamically preferred phase where the A-site cation is K⁺ and the B-site cation is Mn²⁺, Co²⁺ and Cd²⁺;^{10,14,15} each B-site cation is connected to six others via a mixture of syn-anti and anti-anti ligands. In this structure every second channel down the c-axis is occupied by A-site cations while the formate hydrogen atoms fill the other

frameworks tend to exhibit hydrogen-bond driven order disorder transitions, near to or below room temperature, often leading to the emergence of ferroelectric order. 5,6,11 They typically exhibit weak ferromagnetic order, arising from spin canting and, while the coupling of electric and magnetic order are very weak, the combination of these two facets in a framework leads to a new route to potential multiferroics. 3,4,12

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Perovskite III is adopted by $NH_4Cd(HCO_2)_3$, ¹¹ which is polar at ambient temperature, and a less common polymorph of $NH_4Mn(HCO_2)_3$. ⁸ This structure closely resembles Perovksite II except all ligands connect M^{2+} in a *syn-anti* arrangement.

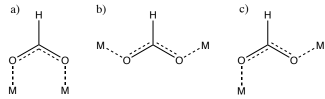


Fig. 1: Possible different coordination modes of the formate anion: a) syn-syn, b) antianti and c) syn-anti.

Two structures with other topologies have currently been observed in the AB(HCO₂)₃ family. Those containing smaller Na⁺ A-site cations adopt a chiral cubic structure with a $3^3 \cdot 5^6$ topology and all linkers adopting a *syn-anti* arrangement. 10,16,17 Chiral hexagonal or pseudo-hexagonal phases with a $4^9 \cdot 6^6$ topology are also known to form where the A-site cation is a small molecule such as NH₄ $^+$, 6,18 HONH₃ $^+$, 1,9 or NH₂NH₃ $^+$ (the later only when B is Co²⁺ and Mg²⁺) 5 . In this architecture all linkers adopt an *anti-anti* configuration. The NH₄ $^+$ members are known to adopt a polar structure at low temperature with ferroelectric or relaxor-like ferroelectric properties 6 while the properties of the NH₂NH₃ $^+$ compounds are also fascinating as, unusually, they undergo a transition from an antiferroelectric phase to polar phases as a result of partial cation disordering. 5

The relative sizes of the cations plays a critical role in determining which of these five structures are adopted by the AB(HCO₂)₃ frameworks. Kielslich et al.²⁰ have recently defined a tolerance factor for hybrid phases extending the classic concept of tolerance factors, traditionally used to judge whether the relative sizes of cations and anions in an ABX3 ionic solid will lead to a perovskite phase, to those containing Their building blocks. formularism $\alpha = (r_{A,eff} + r_{x,eff})/\sqrt{2}(r_B + 0.5h_{x,eff})$, where r refers to the effective molecular or ionic radius of the A-site, B-site and X(anionic)-site while $h_{x,eff}$ refers to the height of the anions in the direction between B-site cations. According to this almost all the AB(HCO₂)₃ compounds with an α of between 0.80 and 1 adopt the Perovskite I phase as expected from their ionic counterparts. 13,20 Gómez-Aguirre et al. 11 have recently extended the use of the tolerance factor to rationalize the order in which other AB(HCO₂)₃ phases appear. They find that Perovskite II is stable for an intermediate tolerance factor range, approximately $0.64 < \alpha < 0.69$, and that $NH_4Cd(HCO_2)_3$, the only compound for which Perovoskite III has been isolated as a pure phase, has an α of 0.63. The inclusion of Na⁺ decreases the tolerance factor further, to below 0.60 leading to the adoption of the chiral cubic phase. They were, however, unable to explain the adoption of the 49.6 chiral hexagonal structure in this scheme, which they described as spanning a range of $0.65 < \alpha < 0.68$ citing the $NH_4B(HCO_2)_3$ series, which overlaps with the Perovskite II series range. 11 Indeed the situation is even more complex than this as chiral hexagonal structures containing HONH3⁺ and NH2NH3⁺ cations have

tolerance factors as high as 0.85, well into the range of the Perovskite I structure. 13,19,21 Clearly other factors beyond ionic radii will influence the structures adopted by some of these materials e.g. the specific shape of the molecular guest and the precise hydrogen bonds it forms with frameworks may favour some structure types over others. It is also likely that polymorphism is present in some of these cases, found for $NH_4Mn(HCO_2)_3^{8,18}$ and $KCo(HCO_2)_3^{14}$ where in the later the kinetic product adopts a chiral hexagonal phase. Given the versatility and ease of use of the tolerance factor, however, it is clearly valuable for giving an approximate indication of what structure is likely to form in the fascinating AB(HCO₂)₃ family particularly in the cases where size effects are primarily responsible for the phases adopted. In an attempt to do this and in particular clarify the position of the 49.6 chiral hexagonal structure in this hierarchy, we have synthesised t $AB(HCO_2)_3$ frameworks where $A = Rb^+$ or Cs^+ and $B = Mn^{2+}$, Cc^{2+} or Ni²⁺. These compounds contain the largest possible monoatomic A-site cations, which leads to higher tolerance factors in the range of 0.66 to 0.73. Here we report the structure of these materials, clarifying the effect of size on the sequence of phases adopted by the AB(HCO₂)₃ family, along with their magnetic properties and thermal stability.

2. Experimental

The compounds synthesized in this work were obtained using a method similar to Eikeland et al. 10 using commercially available reagents. In a typical synthesis 10 mL of 0.73 M HCO₂H (reagent grade 95 %, Sigma-Aldrich) and 0.2 M of either RbOH•xH₂O or CsOH•H₂O (Sigma-Aldrich) in anhydrous methanol (Sigma-Aldrich 99.8 %) was placed at the bottom a glass tube. 10 mL of 0.025 M of MnCl₂•4H₂O (99+ % Across Organics), CoCl₂ (97 %, Sigma Aldrich) or NiCl₂•6H₂O (99.3 % Alfa Aesar) in anhydrous methanol was carefully layered on top. The tube was then sealed and kept undisturbed for up to two weeks, until crystals formed. This led to single crystals of RbMn(HCO₂)₃, CsMn(HCO₂)₃, RbNi(HCO₂)₃, CsNi(HCO₂)₃ and RbCo(HCO₂)₃ suitable for structure determination, the first three of these in pure form. A pure sample of RbCo(HCO₂)₃ was made using two 10 mL methanol solutions of 1.46 M HCO₂H and 0.6 M RbOH•xH₂O, and 0.05 M CoCl₂. Single crystals of CsCo(HCO₂)₃ suitable for structural determination were obtained by decreasing the bottom layer concentrations to 0.365 M HCO₂H and 0.1 M CsOH•H₂O.

Single crystal structure determinations were carried using two diffractometers, a Nonius Kappa CCD equipped with Mc K α radiation, generated by a conventional generator operating at 60 kV and 30 mA, and a dual source Oxford Diffracti n Supernova equipped with Mo K α and Cu K α micro-focus sources, operating at 0.8 kV and 50 mA, with multilayerectocusing optics and a Atlas CCD detector (see Table 1 for crystallographic details). Samples were cooled to 150 K using a Oxford Cryosystem cryostream with the samples held on a MiTeGen micro loop. Data were integrated and empirical or, in the case of CsCo(HCO₂)₃, analytical, absorption corrections

Journal Name

performed using the DENZO/SCALEPACK suite²² and the CrysAlisPro software suite version 171.36.32,²³ for the Nonius and Oxford Diffraction instruments, respectively. Structures were solved using direct or Patterson methods in SHELXS-2014²⁴ or charge-flipping in olex2.solve²⁵ and least-squares refinements were carried out using SHELXL-2014,²⁶ via the Olex2 graphical user interface.²⁷ The displacement parameters of non-hydrogen atoms were refined anisotropically and

hydrogen atoms were typically located geometrically using the AFIX commands in SHELXL-2014, with their displacement parameters constrained to 1.2 times that of the carbon on which they rode. The crystal of $CsMn(HCO_2)_3$ used was a merohedral racemic twin with two equal components within the precision of the refinement.

Table 1: Crystallographic Data for structures **1-6** determined by single crystal X-ray diffraction. Structure **3** was determined using Cu K α while Mo K α radiation was used in all other cases.

Compound	1	2	3	4	5	6
Formula	RbMnC3H3O6	RbCoC3H3O6	RbNiC3H3O6	CsMnC3H3O6	CsCoC3H3O6	CsNiC3H3O6
Formula Weight	275.46	279.45	279.23	322.90	326.89	326.67
T (K)	150(2)	150(2)	150(2)	150(2)	150(2)	150(2)
Crystal System	Monoclinic	Hexagonal	Hexagonal	Hexagonal	Hexagonal	Hexagonal
Space Group	C2/c (15)	P6₃22 (182)	P6 ₃ 22 (182)	P6 ₃ 22 (182)	P6 ₃ 22 (182)	P6 ₃ 22 (182)
a (Å)	11.2786(3)	7.2350(3)	7.2095(3)	7.58190(10)	7.5112(3)	7.4866(7)
b (Å)	9.1088(3)	7.2350(3)	7.2095(3)	7.58190(10)	7.5112(3)	7.4866(7)
c (Å)	7.0254(3)	8.2926(4)	8.1492(4)	8.3786(2)	8.0670(4)	7.8898(8)
α (°)	90	90	90	90	90	90
β (°)	96.6260(10)	90	90	90	90	90
γ (°)	90	120	120	120	120	120
V (ų)	716.93(4)	375.92(4)	366.83(4)	417.117(15)	394.15(4)	382.97(8)
Z	4	2	2	2	2	2
$ ho_{\sf calc}$ (g cm $^{ extstyle -3}$)	2.552	2.469	2.528	2.571	2.754	2.833
μ (cm ⁻¹)	8.565	8.694	11.807	5.865	6.709	7.197
Refl.	7071/822	3066/336	2630/263	9467/225	3248/363	3112/352
Meas./Unique	$[R_{\rm int} = 0.043]$	$[R_{\rm int} = 0.029]$	$[R_{\rm int} = 0.026]$	$[R_{int} = 0.046]$	$[R_{\rm int} = 0.028]$	$[R_{\rm int} = 0.048]$
Parameters Refined	55	19	19	21	19	19
R1, wR2 ^a (all)	0.026, 0.053	0.017, 0.033	0.014, 0.032	0.020, 0.049	0.019, 0.038	0.030, 0.069
R1, wR2 ^a (obs)	0.023, 0.053	0.015, 0.032	0.013, 0.032	0.019, 0.049	0.017, 0.037	0.027, 0.068
χ^2	1.09	1.12	1.18	1.36	1.163	1.09

 a w = $1/[\sigma^{2}(F_{o}^{2}) + (aP)^{2} + bP]$ and $P = (\max(F_{o}^{2}, 0) + 2F_{c}^{2})/3$; R1 = $\sum ||F_{o}| - |F_{c}|| / \sum |F_{o}|$ and $wR2 = \sqrt{[(F_{o}^{2} - F_{c}^{2})^{2}/\sum w(F_{o}^{2})^{2}]}$

Phase purity of the samples created during this study were assessed using powder X-ray diffraction at ambient temperatures. This was carried out using a PANalytical Empyrean diffractometer, operating in Bragg-Brentano geometry, with monochromated Cu K α_1 radiation and a PIXcel 1D detector. Diffraction patterns of RbMn(HCO₂)₃, CsMn(HCO₂)₃, RbNi(HCO₂)₃, and RbCo(HCO₂)₃, held on a glass plate, were fitted using the Le Bail method²⁸ in the program Rietica²⁹ and shown in Fig S1-4, confirming their purity. Diffraction patterns of CsNi(HCO₂)₃ and CsCo(HCO₂)₃ always indicated the presence of an impurity phase, which was unknown and whose structure it was not possible to identify.

Magnetic properties of samples determined to be pure phase by diffraction were characterized by Quantum Design magnetometers, either a MPMS 5 or MPMS XL instrument, equipped with 5 T superconducting magnets. Samples were enclosed in gelatin capsules mounted inside a pierced straw with a uniform diamagnetic background. Thermal stabilities were studied using a Perkin-Elmer TGA 7 Thermogravimetric Analyzer, with samples held in a platinum pan and heated at 10 °C min⁻¹. Infrared spectra were measured between

4000 cm⁻¹ and 500 cm⁻¹ using a Bruker Tensor-27 ATP spectrometer. These spectra have only four features present at about 2850 cm⁻¹ (C-H stretch), 1570 cm⁻¹ (C=O stretch), 1370 cm⁻¹ (C-O stretch) and 770-790 cm⁻¹ (O-C-O symmetric deformation).³⁰ The sharp O-C-O and C-O bands in Rb(HCO₂): are split in two, indicating inclusion of two distinct formates.

3. Results and Discussion

3.1 Crystal Structures

Of the six compounds reported in this paper one of them. RbMn(HCO₂)₃, adopted the Perovskite II structure in $C2^{\prime}c$ monoclinic symmetry, with the remainder adopting $4^{9} \cdot i^{6}$ chiral hexagonal structures in $P6_{3}22$ symmetry. The asymmetric unit of RbMn(HCO₂)₃ contains one Rb⁺ cation, one Mn²⁺ cation and one and a half formate anions, with the Rb⁺ Mn²⁺ and C atom on the second formate anion sitting or special positions with half the occupancy of the general position (see Fig. 2). The framework can be described as a heavily distorted ReO₃ structure with half the channels parallel

to the c-axis occupied by Rb⁺ cations, alternating along the aand b-axis with channels occupied by hydrogen atoms from the ligand (see Fig. 3). The octahedral Mn2+ are linked to six neighbouring octahedra through four equatorial syn-anti formate linkers and two axial anti-anti ligands along the a-axis, forming sheets of skewed squares. The presence of ligands with different coordination modes is consistent with splitting observed in the infrared spectra. The bond valence of the Mn²⁺ cations are 2.08, consistent with a divalent cation.³¹ The Mn-Mn intra-sheet distances are 5.7515(2) Å, significantly shorter than the 6.290(3) Å inter-sheet distance, although the likely superexchange pathways for magnetic interactions are both about 6.85 Å. The Rb[†] cation is bonded to eight oxygen atoms with a bond valence of 1.21.31 Down the directions in which octahedra are connected via formate bridges they can be viewed as being tilted out-of-phase with each other in a way resembling an a a c Glazer tilt pattern. 32 The same tilt system is observed for the K-compounds 10,14 and the C2/c symmetry oxides with this tilt system is maintained despite the three atom formate bridge which skews the structure somewhat.

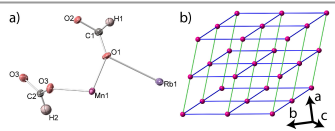


Fig. 2: a) The asymmetric unit of RbMn(HCO₂)₃ shown with 50 % probability ellipsoids. The Mn^{2+} , Rb^+ , C, O and H atoms are pink, purple, black, red and pale pink. An additional oxygen atom, as required to complete the second formate ligand in the structure, is shown. b) The topology of the anionic $Mn(HCO_2)_3$ framework with axial and equatorial formate linkers shown in light green and dark blue.

The other five frameworks adopt the same chiral hexagonal structure reported for the NH₄B(HCO₂)₃ frameworks at room temperature (see Fig. 4 and 5 for depictions of CsMn(HCO₂)₃).¹⁸ The asymmetric units of these compounds contain one alkali metal, one transition metal cation and half a formate ligand, most of the atoms in the structure are on special positions — the metals are on sites with a sixth of the occupancy of the general position and the H and C atoms on sites with half of the general occupancy. The octahedral transition metals in these materials have bond valencies of 2.06 for CsMn(HCO₂)₃ and between 1.96 and 1.98 for all remaining compounds, confirming divalent cations.³¹ The chirality of this architecture comes from helical chains of these octahedra down the c-axis, linked by syn-anti formate linkers forming hexagonal channels; the A-site cations occupy these in highly distorted pseudo-octahedral sites, with bond valencies between 0.93-0.95.31 The framework features puckered hexagonal sheets of transition metal cations linked by intersheet zigzag linkages, neighbouring transition metals in a sheet are significantly closer than between sheets (c.f. separations of 6.0589(1) Å to 8.3789(2) Å for CsMn(HCO₂)₃). This is consistent with the reported antiferromagnetic structures of the NH₄B(HCO₂)₃ compounds which reveals antiferromagnetic order of nearest neighbours in a sheet.⁸ It was observed for RbCo(HCO₂)₃, RbNi(HCO₂)₃ and CsMn(HCO₂)₃ that the *a*-lattice parameter increases on heating while the *c*-lattice parameter shrinks, which is similar to the anisotropic negative thermal expansion in the NH₄B(HCO₂)₃ compounds and is likely due to a similar wine-rack mechanism.⁸ As was also the case for RbMn(HCO₂)₃ a combination of single crystal and powder X-ray diffraction did not suggest any phase transitions occur in these materials between 150-300 K. This is not surprising as the phase transitions found in related structures are caused by hydrogen bonding between the molecular A-site cation and the framework, which alkali metals are unable to support.

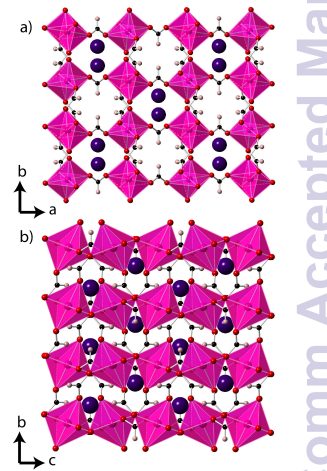


Fig. 3: The structure of RbMn(HCO₂)₃ showing a) the ab plane and b) the bc plane. Colours are the same as Fig 2.

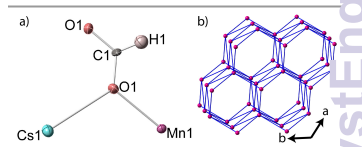


Fig. 4: a) The asymmetric unit of $CsMn(HCO_2)_3$ shown with 50 % probability ellipsoids. The Cs^+ atoms are represented in blue and all other colours are the same as Fig. 2. Ar additional O^{2^-} , required to complete the formate in the structure, is shown. b) The topology of the anionic $Mn(HCO_2)_3$ framework with formate linkers shown in dark blu

Journal Name

ARTICLE

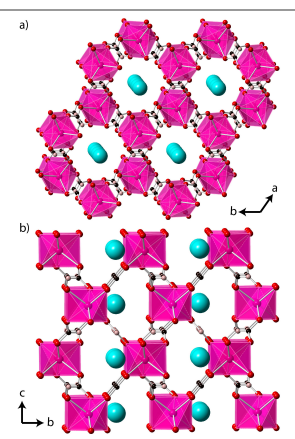


Fig. 5: The structure of CsMn(HCO_2)₃ showing a) the ab plane and b) the bc plane. The colours are the same as Fig. 4.

Tolerance factors for these six new phases were calculated using the method of Kielslich et al. ²⁰ For RbMn(HCO₂)₃ α = 0.66 and the other compounds have 0.68 < α < 0.73. This suggests that, when only monoatomic cations are present in an AB(HCO₂)₃ structure, the tolerance factor cut-off between the Perovskite II structure and the $4^9 {ullet} 6^6$ chiral hexagonal phase is close to 0.66-0.68. This boundary is consistent with the observation of both phases for $KCo(HCO_2)_3$ whose $\alpha = 0.65$. The Perovskite II structure of KCo(HCO₂)₃ is denser that the 49-66 chiral hexagonal phase, 14 which typically makes this structure more thermodynamically stable, as well established for other framework compounds.³³ If, however, the A-site cation is too large it appears that it becomes preferable for an AB(HCO₂)₃ framework to adopt the chiral hexagonal structure due to its larger channels (as a rough guide using a 1.1 Å probe in Mercury CSD 2.0³⁴ on the two KCo(HCO₂)₃ structures, from which K⁺ has been removed, indicates that the channels in which the A-site cation sit take up 12.3 % and 2.5 % of the volume of the chiral hexagonal and Perovskite II phases, respectively). We checked for any indication of phase transformations in the new compounds reported in this work but did not notice any indication of this over a month.

This work clarifies the effect of ionic size on the evolution of the phases adopted by the larger $AB(HCO_2)_3$ family, with the $4^9 \cdot 6^6$ phase appearing to be formed for tolerance factors intermediate of the Perovskite I and Perovskite II phases (see Fig. 6). Based on our results we predict that the $KB(HCO_2)_3$ (B =

 Fe^{2+} , Ni^{2+} and Zn^{2+}) frameworks, which have not been reported to date, would adopt the Perovskite II structure while $RbB(HCO_2)_3$ ($B=Fe^{2+}$ and Zn^{2+}) are likely to adopt the $4^9 \cdot 6^6$ phase. We propose that the tolerance factor range of the $4^9 \cdot 6^6$ phase is extended to lower values, for the $NH_4M(HCO_2)_3$ frameworks, and to higher values, for those including $HONH_3$ and NH_2NH_3 cations, because of the presence of hydroger bonding between these molecular A-site cations and the framework, which the alkali metals cannot replicate. 5,18,19 This is likely because the directionality of hydrogen bonds means the number and strength of these bonds will depend on the precise shape of both the molecular cation and the architecture into which they are incorporated.

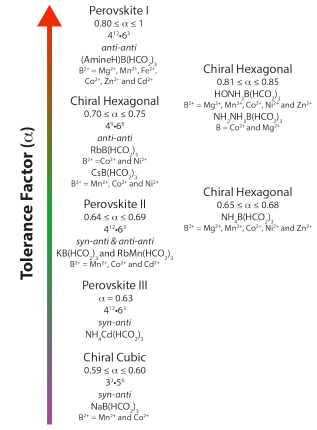


Fig. 6: The evolution of the structures of the known three dimensional $AB(HCO_2)_{\perp}$ frameworks with the tolerance factor, with an emphasis of the role of the molecular cations on the stability of the $4^9 \cdot 6^6$ at lower and higher tolerance factors than obsermed for monoatomic A-site cations.

3.2 Magnetic Properties

Magnetic property measurements were carried out on purephase, by diffraction, RbB(HCO₂)₃ and CsMn(HCO₂)₃ sample 5. The 100 Oe zero-field cooled (ZFC) susceptibil v measurements of the Rb⁺ containing phases exhibit a maximum in their magnetic susceptibility at 4, 9 and 27 K, for Mn²⁺, Co²⁺ and Ni²⁺ (see Fig. 7 for RbCo(HCO₂)₃ and Fig. S5 and S6 for other Rb⁺ frameworks); this is consistent with the onset of antiferromagnetic order. Below the ordering temperatures however, the 100 Oe field cooled (FC) and ZFC measurements diverge indicating a ferromagnetic component is present. This

is suppressed rapidly with increasing applied fields so the ferromagnetic signal is essentially absent in fields over 1 kOe but clear, albeit subtle, hysteresis is observed in magnetization measurements of all three compounds, confirming the presence of a weak ferromagnetic component (see Fig. 8 for RbCo(HCO₂)₃ and Fig. S7 and S8 for other Rb⁺ compounds). The magnetization measurements are not saturated at 50 kOe, consistent with predominantly antiferromagnetic behaviour.

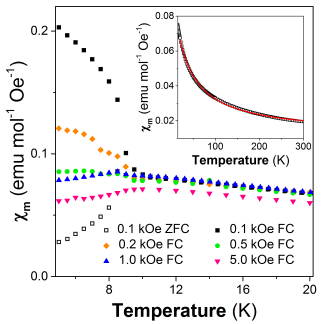


Fig. 7: Plot of variable temperature magnetic susceptibility measurements of RbCo(HCO₂)₃ in various magnetic fields with the insert showing the fit of the Curie-Weiss fit to the 100 Oe ZFC susceptibility measurement with a temperature independent component.

Weak ferromagnetic behaviour has previously been observed in the (NH₄)B(HCO₂)₃ and KB(HCO₂)₃ with similar ordering temperatures. 6,10,14,18 This was attributed to spin-canting, specifically due to Dzyaloshinsky-Moriya interactions in the 49 •6⁶ compounds, which lack an inversion centre, 6,18 and it is possible that symmetry lowering associated with magnetic ordering may also enable this in the Perovskite II structure. The magnetic susceptibility of RbNi(HCO₂)₃ increases further below 5 K, exhibiting a paramagnetic tail most likely caused by the presence of a trace impurity in the sample not detected by X-ray diffraction (see Fig S6). Above the Néel temperatures the behaviour of the Mn and Ni compounds are well fitted by Curie-Weiss behaviour with effective magnetic moments of 5.99 and 3.35 $\mu_{\text{B}}\text{,}$ consistent with the expected values for octahedral Mn²⁺ and Ni²⁺, and Curie-Weiss temperatures of -6.0 K and -62 K (see Fig. S5 and S6). The magnetic susceptibility of RbCo(HCO₂)₃ could only be successfully fitted by the addition of an temperature independent term to the Curie-Weiss equation, yielding an effective magnetic moment of 4.61 μ_B , a Curie-Weiss temperature of -29 K and a temperature independent term of 120 × 10⁻⁴ emu mol⁻¹ Oe⁻¹ (see Fig. 7 insert). While the effective magnetic moment of RbCo(HCO₂)₃ is well within the range expected for octahedral Co^{2+35} it is somewhat lower than that reported for the isostructural K and NH₄ phases, 5.5-5.6 μ_B . ^{14,18}

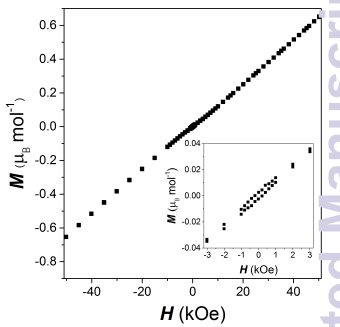


Fig. 8: Isothermal magnetisation measurements of RbCo(HCO₂)₃ at 5 K with the insert highlighting the subtle hysteresis at low applied magnetic fields.

The 100 Oe magnetic susceptibility measurements CsMn(HCO₂)₃ reveal an antiferromagnetic cusp without deviation of the FC and ZFC measurements, consistent with purely antiferromagnetic behaviour below 9 K (see Fig. 9) Consistent with this a magnetization measurement at 5 K exhibits linear behaviour that does not saturate at the maximum field examined, 50 kOe. The magnetic susceptibility measurements in the paramagnetic phase can be well modelled by Curie-Weiss behaviour yielding a Curie-Weiss temperature of -13 K and an effective magnetic moment of 6.14 μ_B , close to the expected spin-only value of 5.92 μ_B for high-spin Mn²⁺ (see Fig. S9). The higher magnetic ordering temperature of this compound compared to the RbMn(HCO₂) phase indicates that the magnetic interactions in the chiral hexagonal phases are significantly stronger than in the Perovskite II phase. This is consistent with a comparison of the other known compounds with these structures, which shows that interchanging between the NH₄⁺, K⁺, Rb⁺ and Cs⁺ cation does not significantly change the magnetic ordering temperature provided the structure and the transition meta. remains the same; changing from the chiral hexagonal structure to the Perovskite II structure, however systematically weakens the magnetic interactions in the e materials regardless of the transition metal present. 6,10,14 This emphasises the importance of understanding the structure-property relationship in the complex AB(HCO₂)₃ family of compounds in the search for new functiona magnetic materials.

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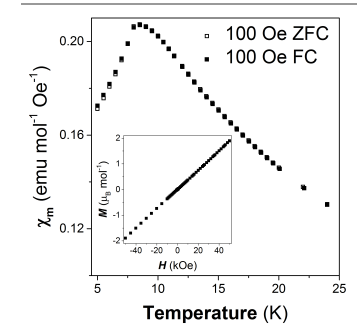


Fig. 9: Plot of variable temperature magnetic susceptibility measurements of CsMn(HCO₂)₃ in a 100 Oe applied field with an insert displaying its isothermal magnetisation measured at 5 K.

3.3 Thermal Stability

Thermogravimetric analysis carried out in air suggest that RbMn(HCO₂)₃, RbCo(HCO₂)₃, RbNi(HCO₂)₃ and CsMn(HCO₂)₃ begin to decompose at approximately 180, 210, 200 and 190 °C and is essentially complete by between 400 and 450 °C (see Fig S10-13). The bulk of the weight loss occurs in one step, as observed for other alkali metal formates, with weights at the end of this step consistent with the formation of alkali metal carbonates and metal oxides. 10,14,17 Only the decomposition of RbMn(HCO₂)₃ is delayed significantly when carried out in nitrogen, occurring at 270 °C, while all other compounds examined begin to decompose within 20 °C of the temperature at which they do in air. Duan et al. 14 have previously suggested that the decomposition pathway of KCo(HCO₂)₃ is different depending on whether it adopts the Perovskite II or chiral hexagonal polymorph and it is also possible that the differences in the structures of these materials may have some effect on their decomposition pathways in the absence of oxygen. The decomposition temperatures observed for these Rb⁺ and Cs⁺ compounds is somewhat lower than those observed for AB(HCO₂)₃ frameworks containing Na⁺ or K⁺, which are reported to decompose above 330 and 230 °C, respectively. 10,14,17 This suggests that the thermal stability of this family of alkali frameworks decreases significantly with increasing size of the A-site cation, likely because their decreased charge density leads to weaker interactions between them and the anionic framework.

4. Conclusions

This study reports the structure of six new AB(HCO₂) frameworks, where $A = Rb^{+}$ or Cs^{+} and $B = Mn^{2+}$, Co^{2+} or Ni^{2+} RbMn(HCO₂)₃ adopts a perovskite-like structure with $4^{12} \cdot 6^{3}$ topology with a mixture of syn-anti and anti-anti ligand coordination modes, while the other five compounds in this study all exhibit a chiral hexagonal structure with 49 • 65 topology. Adding this knowledge to the structure of knowr related compounds we have proposed a phase diagram rationalising the effect of ionic size on the formation of the various structures known to be formed by the AB(HCO₂)₁ frameworks, in particular clarifying the placement of the 49-6 chiral hexagonal frameworks within this. This also emphasises the likely importance of hydrogen bonding in stabilising the chiral hexagonal phases, which are commonly reported to ferroelectric, compared to perovskite-like topologies where a molecular A-site cation is present. The Rb⁺ containing compounds were all found to be weak ferromagnets, most likely due to spin canting, while CsMn(HCO₂)₃ appears to be ar antiferromagnet. The magnetic interactions in RbMn(HCO₂)₃ are much weaker than in the other compounds reported ir this study, confirming the trend that the magnetic interactions in the perovskite-like structure it adopts are much weaker than in the $4^9 \cdot 6^6$ phases.

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Notes and references

i We have (re)calculated the tolerance factors throughout this manuscript using the ionic radii of the high spin state, as determined by Shannon, ³⁶ found experimentally to be adopted by the paramagnetic B-site cations in these compounds. For consistency with Gómez-Aguirre et al., ¹¹ we have used the relevant eight coordinate ionic radii for the alkali metals. ³⁶

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