**Operando Synchrotron X-ray Diffraction in Calcium Batteries: Insights into the Redox Activity of 1D Ca$_3$CoMO$_6$ (M = Co and Mn)**

A.P. Black, D. Monti, C. Frontera, D. S. Tchitchekova, R. G. Houdeville, F. Fauth, and M.R. Palacin

**ABSTRACT:** 1D Ca$_3$Co$_{2-z}$M$_z$O$_6$ (M = Co, z = 0; M = Mn, z = 1, and M = Fe, z = 0.4) were prepared and tested electrochemically. While the iron-containing phase was not found to be active, the iron- and manganese-containing phases were found to be potentially interesting as positive electrode materials for calcium metal-based high-energy battery technologies and were investigated by operando synchrotron X-ray diffraction. Results indicate that electrochemically driven calcium deintercalation from the crystal structure (ca. 0.7 mol per formula unit) takes place upon oxidation in both cases. The oxidized phases have incommensurate modulated crystal structures with the space group R 3m(001)0s and a = 9.127(1) Å, c$_1$ = 2.4226(3) Å and c$_2$ = 4.185(3) Å, and γ = 0.579 (M = Co) and a = 9.217(1) Å, c$_1$ = 4.9076(4) Å and c$_2$ = 4.3387(4) Å, and γ = 1.139 (M = Mn), which exhibit differences due to the presence of manganese and Mn/Co ordering. The degree of calcium re-intercalation within the structure was found to be extremely limited, if any. Complementary experiments carried out in lithium cells did not show any reversibility either, thus pointing at intrinsic structural/migration constraints in the oxidized phase rather than slow kinetics of high desolvation energies associated with divalent ion charge carriers.

**INTRODUCTION**

The quest for high-energy-density battery technologies has revived research on metal anode-based concepts, not only for the case of lithium, for which dendrite growth has to date been a technical showstopper, but also for other more abundant electrropositive metals such as sodium, potassium, magnesium, calcium, or aluminum. Among these, the first two represent a more straightforward option, as the know-how gained in the study of lithium-based chemistries can be most valuable. In contrast, multivalent concepts come with intrinsic significant differences, which make their research more complex in nature with previous lithium-based know-how being less useful. Yet, theoretical energy density coupled to an expected reduced cost makes multivalent chemistries appealing.

Despite the standard reduction potential for Ca$^{2+}$/Ca being ~2.87 versus normal hydrogen electrode, research on calcium batteries has been relatively limited until 2015. Moreover, in the absence of standards, even the development of reliable experimental protocols has required a significant effort. These are especially critical in the quest for new positive electrode materials, as the operation of calcium as the negative electrode can be plagued with a number of issues which can affect the results of positive electrode testing. For instance, electrolytes enabling calcium plating and stripping are limited and not optimized. Hence, electrochemical response per se is not enough to assess the redox activity and the use of additional advanced materials characterization techniques is compulsory. In addition to that, the length scale that can be probed with such tools needs to be considered as, for instance, X-ray photoelectron spectroscopy or other surface-sensitive techniques are not trustworthy to assess bulk behavior. Finally, sample modification during handling/measurement may also lead to biased results. In this sense, operando diffraction is the most appealing. Besides being nondestructive, it enables capturing metastable intermediates and ascertaining the influence of operation conditions on the redox mechanism.

Nonetheless, the cell geometry must be compatible with the experimental setup while at the same time being electrochemically efficient, which can be problematic in the case of new technologies due to the lack of standard cell components. The use of synchrotron radiation improves data collection statistics and angular resolution. To the best of our knowledge, synchrotron operando diffraction in calcium cells has only been reported for graphite electrodes at low potentials using activated carbon (AC) as the counter electrode (CE) and Ca(TFSI)$_2$ in tetraglyme as the electrolyte and was helpful in assessing staging. Therefore, in situ and operando advanced materials characterization techniques and tools need to be considered as, for instance, X-ray photoelectron spectroscopy or other surface-sensitive techniques are not trustworthy to assess bulk behavior. Finally, sample modification during handling/measurement may also lead to biased results. In this sense, operando diffraction is the most appealing. Besides being nondestructive, it enables capturing metastable intermediates and ascertaining the influence of operation conditions on the redox mechanism. Nonetheless, the cell geometry must be compatible with the experimental setup while at the same time being electrochemically efficient, which can be problematic in the case of new technologies due to the lack of standard cell components. The use of synchrotron radiation improves data collection statistics and angular resolution. To the best of our knowledge, synchrotron operando diffraction in calcium cells has only been reported for graphite electrodes at low potentials using activated carbon (AC) as the counter electrode (CE) and Ca(TFSI)$_2$ in tetraglyme as the electrolyte and was helpful in assessing staging.
Looking for positive electrode materials for calcium-based batteries operating at high potential, intercalation compounds appear as an appealing option. Indeed, topotactic insertion of a guest species into a host is possible for several metal ions. The host compounds should exhibit an open framework of interconnected sites through which the intercalated ion can diffuse and an electronic band structure able to reversibly accept/donate electrons. When looking for compounds able to operate as positive electrode materials, hosts containing transition metals are the most suitable, which is the reason why these are used in the ubiquitous Li-ion batteries. Within this scenario, we turned our attention toward Ca$_3$Co$_2$O$_6$. This compound is an end member of the general A$_{1+x}$(AyB$_{1-y}$)$_x$O$_3$ ($0 \leq y \leq 1/2$) family of compounds related to the 2H ($y = 0$) hexagonal perovskite with $y = 1/2$, A = Ca, and A' = B = Co. Its crystal structure exhibits columns of face-sharing A'O$_6$ trigonal prisms and B'O$_6$ octahedra (in a ratio that is given by $y$) running along the c direction. This framework is similar to that of TiS$_3$, where the columns consist only of trigonal prisms, and for which lithium intercalation ability has been known for long, albeit at too low potential to present a practical interest. In the case of Ca$_3$Co$_2$O$_6$, high redox operation potential can be expected related to the oxidation of Co(III) to Co(IV) concomitant to electrochemical calcium deinsertion from the structure. Moreover, a high theoretical capacity (160 mA h/g) can be calculated considering the extraction of 1 mol of Ca$^{2+}$ per formula unit to yield capacity (160 mA h/g) can be calculated considering the extraction of 1 mol of Ca$^{2+}$ per formula unit to yield capacity.21

With the aim of fully assessing the reaction mechanism of Ca$_3$Co$_2$O$_6$ at room temperature and getting further insights into the reversibility and/or formation of alternative phases, we performed operando synchrotron diffraction studies using a modified coin-cell setup and an alternative electrochemical configuration enabling operation at room temperature while avoiding the use of calcium metal CE.22 The influence of crystal chemistry within the activity and reversibility of the system was further explored by broadening the study to other isostructural phases containing alternative transition metals which could act as redox centers. Indeed, these exhibit somewhat different cell parameters and hence different intercolumn spacings, which may have an impact on the barriers for the migration of calcium ions. The only suitable phases already reported contain iron or manganese and were previously investigated for the interest in their magnetic properties. Among these, the maximum amount of iron in the structure is achieved for Ca$_3$Co$_2$Fe$_x$O$_{6+x}$ in which Fe(III) occupies trigonal prismatic sites28 and has a similar unit-cell volume ($741.6 \text{ Å}^3$) compared to Ca$_3$Co$_2$O$_6$ ($740.9 \text{ Å}^3$). For the case of manganese, Ca$_3$CoMnO$_6$ has been reported,27 in which Mn(IV) occupies the octahedral sites and exhibits a larger cell volume ($764.1 \text{ Å}^3$), resulting in a larger intercolumn space within the structure.

### EXPERIMENTAL SECTION

**Synthesis.** Ca$_3$Co$_2$O$_6$ was prepared by the Pechini method and 30 mL of a 1 M solution of Ca(NO$_3$)$_2$ (99% tetrahydrate, Sigma-Aldrich) and 20 mL of a 1 M solution of Co(NO$_3$)$_2$ (98%, hexahydrate, Sigma-Aldrich) in water were mixed with 10.5 g of citric acid (C$_6$H$_8$O$_7$, 99%, Sigma-Aldrich) and put together in a glass vial to which 6.25 g of ethylene glycol (C$_2$H$_6$O$_2$, 99%, Sigma-Aldrich) was added (corresponding to 3:2:5:10 molar ratios for Ca/Co/C$_6$H$_8$O$_7$/C$_2$H$_6$O$_2$). The solution was subsequently heated at 80 °C for 4 h and thereafter stirred overnight at room temperature to promote water evaporation, which induced enhanced viscosity of the mixture and the evolution of nitrous oxide gas. Once the gel formed dried, it was placed in an alumina crucible and heated in air first at 600 °C for 6 h and then at 950 °C for 20 h and subsequently cooled to room temperature. The synthesis of Ca$_3$Co$_2$Fe$_x$O$_6$ was attempted by both the Pechini method and a solid-state reaction. In the first case, 30 mL of a 1 M solution of Ca(NO$_3$)$_2$ (99% tetrahydrate), 10 mL of a 1 M solution of Co(NO$_3$)$_2$ (98%, hexahydrate), and 10 mL of a solution of Fe(NO$_3$)$_3$ (98%, hexahydrate, Sigma-Aldrich) in water were mixed with 10.5 g of citric acid (C$_6$H$_8$O$_7$, 99%, Sigma-Aldrich) and put together in a glass vial to which 6.25 g of ethylene glycol (C$_2$H$_6$O$_2$, 99%, Sigma-Aldrich) was added (corresponding to 3:1:1:5:10 molar ratios for Ca/Co/Fe/C$_6$H$_8$O$_7$/C$_2$H$_6$O$_2$). Thermal treatments analogous to those enabling the synthesis of Ca$_3$Co$_2$O$_6$ resulted in a mixture of Ca$_3$Fe$_x$O$_6$, Ca$_3$CoO$_6$, and/or Ca$_3$Co$_2$Fe$_x$O$_6$, and CaO despite a previous report suggesting the indexation of an analogous diffraction pattern with a single triclinic unit cell with no structural model provided.29 For the solid-state route, a stoichiometric mixture of CaCO$_3$, Fe(NO$_3$)$_2$, and Fe$_2$O$_3$, was successively heated at 850, 950, and 1000 °C for 24 h in each step, with intermediate regrinding, and single-phase Ca$_3$Fe$_{2-x}$Co$_x$O$_6$ was achieved for $x \leq 0.4$. Ca$_3$CoMn$_x$O$_6$ was obtained both using a solid-state route from CaCO$_3$, Co$_3$O$_4$, and MnO$_2$ after successive treatments in air at 850 °C for 20 h and 1200 °C for 24 h with intermediate regrinding and the Pechini method (from 3:1:3:5:10 molar ratio of Ca/Co/Mn/C$_6$H$_8$O$_7$/C$_2$H$_6$O$_2$ using Ca(NO$_3$)$_2$, Co(NO$_3$)$_2$, Mn(NO$_3$)$_2$, citric acid, and ethylene glycol with thermal treatments under synthetic air at 600 °C for 6 h and 1000 °C for 20 h).

**Characterization.** Scanning electron microscopy micrographs were acquired using an FEI Quanta 200 FEG microscope under high vacuum operating at 20 kV. X-ray powder diffraction patterns were acquired in a Bruker D8 Advance A25 diffractometer in a Debye–
Scherer configuration equipped with a Mo Kα radiation source ($\lambda = 0.7093$ Å) and a Johansson monochromator. Synchrotron X-ray diffraction (SXRD) patterns were collected on the position-sensitive detector MYTHEN and $\lambda = 0.6200$ Å. In both cases, the samples, either as-prepared or recovered after running electrochemical experiments and dismantling cells inside an Ar-filled glovebox, were embedded in a 0.5 mm diameter borosilicate glass capillary and spun during data collection to ensure proper powder averaging.

**Electrochemical Experiments.** Tape-casted electrodes were prepared from the mixture of as-synthesized powder samples with carbon black (Super P, Timcal, Switzerland) to enhance the electronic conductivity and polyvinylidene fluoride (Arkema) as the binder in a weight ratio of 80:10:10. The mixture was further dispersed in N-methyl-2-pyrrolidone (Aldrich, ≥99.9%), ground in a ball mill, casted on an 18 μm thick aluminum foil (Goodfellow, 99%) with a blade gap of 300 μm, and vacuum-dried at 80 °C for 24 h. Punched ca. 1 cm$^2$ electrodes were tested in three-electrode Swagelok cells in different configurations (loading ca. 2 mg/cm$^2$). Galvanostatic cyclic with potential limitation (GCPL) tests were carried out at C/100 (1 mol of electrodes exchanged in 100 h) at room temperature with sodium counter and reference electrodes and 1 M NaF6 (Strem Chemicals, 99%, EC/PC/DMC (0.45:0.45:0.1) as the electrolyte (all solvents, Aldrich, anhydrous, 99.0%) in EC/PC/DMC (0.45:0.45:0.1) as the electrolyte (all solvents, Aldrich, anhydrous, 99.0%) as the electrolyte to provide inputs for comparison with alternative setups. Ca metal was also used as counter and pseudoreference electrodes, with 0.3 M Ca(BF4)$_2$ dissolved in a 1:1 mixture of EC (Aldrich, anhydrous, 99.0%) and propylene carbonate (PC, Aldrich anhydrous, 99.0%) as the electrolyte, prepared from Ca(BF4)$_2$·H2O salt (Alfa Aesar). Prior to electrolyte preparation, Ca(BF4)$_2$ was dried under vacuum in a glovebox, were then further dried inside an Ar-filled glovebox with less than 1 ppm of O2 and H2O by heating at 90 °C overnight to achieve a water content lower than 80 ppm, as determined by Karl Fischer analysis. Cells were tested in the galvanostatic mode at C/200 (defined as the reaction of 1 mol of Ca$^{2+}$ in 200 h and equivalent to 1 mol of electrons exchanged in 100 h) on a Bio-Logic VMP3 potentiostat and were tested in the galvanostatic mode at C/200 (defined as the reaction of 1 mol of Ca$^{2+}$ in 200 h and equivalent to 1 mol of electrons exchanged in 100 h) on a Bio-Logic VMP3 potentiostat.

Powder electrodes were prepared by simply mixing active materials with carbon black (Super P, Timcal, Switzerland) in a weight ratio of 75:25, respectively. Operando electrochemical tests were performed in two types of in situ cells: drilled 2032 coin cells with a 4–5 mm glass window and Leriche-type cells. A$^2$ An aluminum foil (3 μm, Goodfellow) was placed on the positive electrode side to protect the window and ensure good electrical contact. One glass fiber filter disk (Whatman, GE Healthcare, 420 μm thick) was used as a separator. Electrochemical experiments were carried out in a galvanostatic mode at room temperature and C/50 (defined as the reaction of 1 mol of Ca$^{2+}$ in 50 h) using the same electrolyte as the solid-state reaction, in which some low-intensity impurity peaks are clearly visible in the pattern corresponding to 3% CaO (as deduced from the refinement). Since previous neutron powder diffraction studies have shown that both Mn and Fe replace Co in the octahedral position, we have presumed this ordering of transition metals to take place. All patterns have been successfully refined using the Rietveld method. As expected, the Pechini method delivered more homogeneous and smaller (<5 μm) particles than solid-state reactions.

**RESULTS AND DISCUSSION**

**Synthesis and Electrochemical Characterization.** Pure Ca$_3$Co$_{1.6}$Fe$_{0.4}$O$_6$, Ca$_3$CoMnO$_6$ and Ca$_3$Co$\_x$Fe$_x$O$_6$ were obtained using the conditions reported in the Experimental Section. Figure 1 depicts the corresponding XRD patterns and representative scanning electron micrographs. As expected, the Pechini method delivered more homogeneous and smaller (<5 μm) particles than solid-state reactions.

Rietveld refinements indicate that the prepared compounds are free from impurities (except for Ca$_3$CoMnO$_6$ prepared by the solid-state reaction, in which some low-intensity impurity peaks are clearly visible in the pattern corresponding to 3% CaO as deduced from the refinement). Since previous neutron powder diffraction studies have shown that both Mn and Fe replace Co in the octahedral position, we have presumed this ordering of transition metals to take place. All patterns have been successfully refined using the Rietveld method. As expected, the Pechini method delivered more homogeneous and smaller (<5 μm) particles than solid-state reactions.

Operando experiments have shown that both Mn and Fe replace Co in the octahedral position, we have presumed this ordering of transition metals to take place. All patterns have been successfully refined using the Rietveld method. As expected, the Pechini method delivered more homogeneous and smaller (<5 μm) particles than solid-state reactions.
The feasibility of electrochemical oxidation on iron- and manganese-containing compounds was tested at room temperature in cells using sodium counter and reference electrodes and sodium-based electrolytes, in conditions which had previously enabled the oxidation of Ca$_3$Co$_2$O$_6$ concomitant to Ca$^{2+}$ extraction. Figure 2A depicts the potential versus capacity profile of Ca$_3$CoMnO$_6$ composite electrodes upon oxidation in sodium cells. The potential increases rapidly to 3.9 V versus Na$^+$/Na and then a sloping region is observed, with a capacity of 267 mA h/g being achieved at 4.4 V. This behavior is similar to the one of Ca$_3$Co$_2$O$_6$ with most likely some electrochemical capacity being related to electrolyte decomposition at high potential. Analogous experiments attempting to oxidize Ca$_3$CoMnO$_6$ were carried out in calcium cells using AC as the CE and a silver wire as the pseudoreference electrode. Figure 2B depicts the characteristic potential versus capacity profile during a single oxidation/reduction galvanostatic cycle using 0.3 M Ca(BF$_4$)$_2$ in EC/PC as the electrolyte (54 ppm H$_2$O) at the C/50 rate. The oxidation evolves through a pseudoplateau centered around 1.5 V versus AC with an associated capacity of 100 mA h/g. Further oxidation involves a linear increase in potential up to 2.1 V (160 mA h/g). Upon reduction, the potential decreases rapidly down to −1.2 V versus AC and a sloping region is further observed down to −2 V versus AC (corresponding to a capacity of 120 mA h/g). Figure 2C depicts the XRDE patterns of the (a–f) samples shown in Figure 2A oxidized to different extents in sodium cells and an additional pattern (g) oxidized in a calcium cell (200 mA h/g) at room temperature in 0.3 M Ca(BF$_4$)$_2$ in EC/PC using Ca metal as counter and reference electrodes. Upon oxidation, patterns gradually evolve with peaks at Q = 1.37, 2.23, and 2.37 Å$^{-1}$ of the pristine phase decreasing in intensity with their position remaining unchanged and new (strong) peaks appearing at 2.95 and 1.36 Å$^{-1}$, together with some tiny peaks in the region of Q ~ 2 Å$^{-1}$. Thus, regardless of the CE and electrolyte used, the changes in the diffraction pattern observed upon oxidation are the same and hence attributed to electrochemical calcium extraction from the crystal structure to form Ca$_3$MnCoO$_6$, similar to what had been observed for Ca$_3$CoO$_6$.

Attempts to oxidize Ca$_3$Co$_{1.6}$Fe$_{0.4}$O$_6$ were unsuccessful, as no changes were observed in the XRDE patterns despite a plateau being observed at a high potential (3.9 V vs Ca$^{2+}$/Ca) with a capacity close to 200 mA h/g, which was thus attributed to electrolyte decomposition. In this case, one must conclude that any electrochemical activity for this compound, if existing, takes place beyond the electrochemical stability window of the electrolyte used.

**Operando Diffraction Studies.** *Operando* synchrotron diffraction experiments involving materials for calcium batteries are challenging. Beyond the general requirement of compatibility of all the components that are integrated in the *in situ* cells in terms of corrosion stability, the sluggish diffusion rates of the calcium ion into the crystal host material impose a limitation in terms of C-rate and hence unavoidably long experiments. Finally, the lack of electrolytes enabling calcium plating/stripping at room temperature and a fast C-rate which are also stable at high potentials (>4 V vs Ca$^{2+}$/Ca) constitutes a showstopper for these materials.

In the present study, and in order to enhance the efficiency of the full experiment, a setup enabling the alternative sequential measurement of four cells was used (see Figure 3) inspired in ref 24.

*Operando* experiments aiming at getting further insights into the redox activity of 1D Ca$_3$CoMnO$_6$ (M = Co and Mn) were carried out using 0.3 M Ca(BF$_4$)$_2$ in EC/PC at the electrolyte (54 ppm H$_2$O) and consisted of a single oxidation/reduction cycle at C/50 within an electrochemical potential window between 2.5 and −2.5 V versus AC. Since the main objective was to assess the reversibility of the process, if any, the switch between oxidation and reduction steps was made manually once there was clear evidence of significant calcium extraction.

![Figure 2](https://doi.org/10.1021/acs.energyfuels.1c01343)
Since C/50 is a rather slow rate, in order to optimize the beamtime, some cells were cycled \textit{ex situ} and placed under the beam to be monitored only when they reached a certain state of charge. This also enabled us to assess the reproducibility of the redox behavior both under and outside the beam.

In the case of \( \text{Ca}_3\text{Co}_2\text{O}_6 \), different cells were monitored \textit{operando} starting at different states of charge. Cell A was launched to cover the initial oxidation up to a specific capacity of 52 mA h/g. Cell B was initially oxidized out of the beam (160 mA h/g, equivalent to 1 mol of \( \text{Ca}^{2+} \) extracted if this was the only redox process taking place, see the upper abscissa axis in Figure 4a) and placed under the beam for further oxidation up to 213 mA h/g, followed by a reduction step. The electrochemical curve evolves through a pseudoplateau centered at about 1.5 V versus AC upon oxidation, and a more sloping trend upon reduction, centered at \(-1.2\) V versus AC. Selected \textit{in situ} SXRD patterns for those cells displayed in Figure 4b–f are in full agreement with the results from previous experiments using the same electrolyte but carried out at 100 °C and using calcium metal CE.\textsuperscript{21} The (300) and (100) Bragg reflections, at \( Q=2.396 \) and 1.383 Å\(^{-1}\), respectively, progressively decrease in intensity, whereas new peaks grow at \( Q=2.386 \) and 1.377 Å\(^{-1}\). The latter are ascribed to \( \text{Ca}_3\text{Co}_2\text{O}_6 \) with \( x \sim 0.7 \), which exhibits an incommensurate modulated crystal structure [space group \( R\ 3m(00\gamma)0\alpha \), \( a=9.127(1) \) Å, \( c_1=2.4226(3) \) Å and \( c_2=4.1857(3) \) Å, and \( \gamma=0.579(2) \)]. The \( \text{Ca} \) content in this phase is straightforwardly related to the propagation vector, determined by the position of the tiny peak appearing at \( Q \sim 2 \) Å\(^{-1}\) among others.\textsuperscript{21} The patterns depicted in Figure 4b–f correspond to pristine \( \text{Ca}_3\text{Co}_2\text{O}_6 \) intermediate, and final stage of oxidation as well as the intermediate and final stage of reduction. From the evolution of patterns represented in Figure 4e, a progressive intensity increase for the (100)\(^{\prime}\) reflection at \( Q=1.377 \) Å\(^{-1}\) as a function of oxidation time can be inferred. The patterns observed at the end of oxidation are consistent with the electrodes containing a mixture of both oxidized \( \text{Ca}_3\text{Co}_2\text{O}_6 \) and pristine \( \text{Ca}_3\text{Co}_2\text{O}_6 \) phases. No significant evolution of the peaks, neither in position nor in intensity, is seen when the polarity of the cell is reversed. Indeed, Figure 4f depicts the patterns taken upon reduction and no changes are evidenced in the region of (300)\(^{\prime}\) (\( Q=2.386 \) Å\(^{-1}\)) throughout the process (the last pattern corresponds to a capacity of 325 mA h/g). Such results are fully consistent with those derived from diffraction patterns taken \textit{ex situ} after electrochemical experiments were carried out at 100 °C and using Ca metal as the CE and indicate that the electrochemical extraction of calcium from the crystal structure is irreversible in the experimental conditions used and the capacity recorded is only related to side reactions involving the electrolyte.

Figure 5a displays the characteristic potential versus capacity profiles for \( \text{Ca}_3\text{CoMnO}_6 \) electrodes during galvanostatic testing at C/50 in the course of the \textit{operando} experiment. The red curve corresponds to the oxidation step, which evolves through a pseudoplateau around 1.3 V versus AC [equivalent to 4.5 V vs \( \text{Ca}^{2+/3} / \text{Ca} \) (see the right axis in Figure 5a)\textsuperscript{,}f] up to a capacity of about 133 mA h/g that would correspond to a nominal extraction of 0.82 mol of \( \text{Ca}^{2+} \) from the crystal structure (top abscissa axis in Figure 5a). The reduction step is denoted in green for the section monitored under the beam (\textit{operando}) to a capacity of 112 mA h/g and in blue beyond that value, with a final diffraction pattern taken after extended reduction to a capacity of 325 mA h/g (black dot). Oxidation evolves through a pseudoplateau centered at about 1.3 V versus AC. The reduction process proceeds through a sloping region down to \(-3.8\) V versus AC and ca. 260 mA h/g, after which noisy behavior is observed, probably due to bubbling related to electrolyte decomposition.\textsuperscript{35} The SXRD patterns collected throughout the experiment are shown in Figure 5b. Zoom-in images (c–f) highlight the regions where the evolution of the phase composition is clearly observed, with red patterns corresponding to oxidation and green patterns to reduction. Upon oxidation of \( \text{Ca}_3\text{CoMnO}_6 \), a progressive decrease in the intensity of the peaks at \( Q=1.365 \) and 2.363 Å\(^{-1}\) occurs at the expense of new peaks appearing at \( Q=1.375 \) and 2.382 Å\(^{-1}\) [zoom-in images (d) and (e)] and also at \( Q=2.03, 2.09, 3.00, \) and 3.30 Å\(^{-1}\), which gradually grow upon oxidation. These are consistent with the formation of a new \( \text{Ca}_3\text{CoMnO}_6 \) phase upon oxidation, in agreement with the results of \textit{ex situ} experiments (Figure 2c).

The patterns corresponding to \( \text{Ca}_3\text{CoMnO}_6 \) bear great similarity with those corresponding to \( \text{Ca}_3\text{Co}_2\text{O}_6 \) (\( x \sim 0.7 \)). Peaks (100) and (300) progressively decrease, while reflections attributed to \( \text{Ca}_3\text{Co}_2\text{O}_6 \) appear at slightly lower \( Q \)-values (Figure 5c, D) than for \( \text{Ca}_3\text{Co}_2\text{O}_6 \). In addition to those, other low-intensity peaks appear, which are not consistent with the incommensurate structure of \( \text{Ca}_3\text{Co}_2\text{O}_6 \). This does not come as a surprise because despite pristine compounds being...
isostructural, the incommensurate structure of $\text{Ca}_3\text{Co}_2\text{O}_6$, with only one Co crystallographic position, is not compatible with the Mn/Co ordering present in $\text{Ca}_3\text{CoMnO}_6$. Indeed, in the incommensurate structure of $\text{Ca}_3\text{Co}_2\text{O}_6$, any transition metal position is suitable to switch (when varying the Ca content) from octahedral to prismatic, but it is unlikely that Mn would occupy a prismatic position. According to neutron powder diffraction (and our own refinements), Mn–O bond distances are close to 1.905 Å and consistent with Mn$^{4+}$, far from the bond distances

Figure 4. Potential vs capacity profiles from GCPL experiments on $\text{Ca}_3\text{Co}_2\text{O}_6$/AC cells using Ca(BF$_4$)$_2$ in EC/PC as the electrolyte at RT and C/50 rate (a), corresponding SXRD patterns (b) and zoom-in images (c,d), selected patterns correspond to the first and last pattern of cell (A) in black and light red, last oxidation and last reduction of cell (B) in red and dark green, and last ex situ pattern cell (B) in blue. (e,f) Zoom-in images of reflections 100 and 300 of in situ SXRD patterns of cells (A,B), respectively.

Figure 5. Potential vs capacity profiles from GCPL experiments on $\text{Ca}_3\text{CoMnO}_6$/AC cells using Ca(BF$_4$)$_2$ in EC/PC as the electrolyte at RT and C/50 (a), corresponding SXRD patterns (b) and zoom-in images (c,d), selected patterns correspond to pristine and last pattern in oxidation in black and red, last reduction in dark green, and last ex situ pattern of cell (C) in blue. (e,f) Zoom-in images of reflections 100 and 300 of in situ SXRD patterns, respectively.
Figure 6. (a) Crystal structure model for Ca$_{2.28}$MnCoO$_6$ (Co is placed at the center of the dark blue polyhedra and Mn at the center of magenta octahedra, and oxygen atoms are represented by small red spheres and Ca by light blue, large spheres) showing the proposed incommensurate modulation. (b) Rietveld refinement of the crystal structure based on the diffraction pattern using the R3m(00γ)0s structure plus a remaining fraction of the pristine Ca$_3$CoMnO$_6$ phase and a small signal coming from the Al foil.
observed in prisms (>2.1 Å). Thus, an alternative incommensurate crystal structure model (Figure 6a) has been built, which is compatible with the Co/Mn order while keeping Mn in the octahedral environment and is detailed in the Supporting Information.

The Rietveld refinement of the last pattern collected upon oxidation using this model is plotted in Figure 6b (background has been subtracted for clarity). The refined values of the cell parameters are $a = 9.217(1)$ Å, $c_1 = 4.9076(4)$ Å, and $c_2 = 4.339(3)$ Å ($\gamma = 1.139$) for the incommensurate phase [48(2) wt%] and $a = 9.150(1)$ Å and $c = 10.596(1)$ Å for the pristine phase [52(2) wt%]. The calcium content can be deduced from the value of the propagation vector and is consistent with $\text{Ca}_2\text{CoMnO}_6$. At the end of oxidation, $\text{Ca}_{2.3}\text{CoMnO}_6$ represents about 48(2) wt% and is hence equivalent to 0.35 mol of $\text{Ca}^{2+}$ extracted per formula unit. This value is significantly smaller than the one estimated from the electrochemical capacity (0.82), which indicates that partial contribution to the current comes from a parasitic redox process, most likely related to electrolyte decomposition. The reasons for the extraction of a lower amount of calcium for this compound are not clear at this stage. It could be speculated that the ordering of Co and Mn in $\text{Ca}_3\text{CoMnO}_6$ makes calcium deinsertion entropically less favorable with respect to that of $\text{Ca}_3\text{CoO}_6$, as the position of pyramidal environments is restrained to Co sites. Moreover, the migration mechanism for calcium ions (see the Supporting Information) implies the simultaneous rotation of two oxygen triads versus one single rotation in the case of $\text{Ca}_2\text{CoO}_6$.

Rietveld refinements of the pattern taken at the end of reduction indicate that despite a slight growth of (100) and (300) peaks at $Q = 1.374$ and 2.381 Å$^{-1}$, the amount of $\text{Ca}_{2.2}\text{CoMnO}_6$ is still 42(3) wt%, which would point at only 0.04 mol of $\text{Ca}^{2+}$ ions being reinserted in the crystal structure upon reduction. Moreover, this result must be taken with caution as the precise region of the sample where the measurement is done could not be exactly the same after relocating the cell under the beam.

To further assess whether the limited reversibility observed, if any, is related to the sluggish diffusion of $\text{Ca}^{2+}$ ions within the oxidized phase or the strong $\text{Ca}^{2+}$ solvent interactions, causing high desolvation energies which may hinder calcium reinsertion, or related to electrolyte decomposition, resulting in the formation of some blocking films on the surface of the particles, an additional operando experiment was conducted in lithium cells. In this case, standard electrolytes known to be stable at high potentials are used. The charge compensation mechanism during the oxidation of $\text{Ca}_3\text{CoMnO}_6$ should also be related to the extraction of calcium ions from the crystal structure, while upon reduction, the lithium ions present in the electrolyte (much more mobile than calcium ions) might be inserted in the crystal structure.

Figure 7a displays the characteristic potential versus capacity profiles of $\text{Ca}_3\text{CoMnO}_6$ electrodes during operando galvanostatic oxidation and reduction at C/50 with 1 M LiPF$_6$ in EC/DMC as the electrolyte and using AC as the CE. The potential versus capacity profiles upon oxidation in lithium and calcium cells are similar, while the reduction takes place at a slightly higher potential for the case of lithium, which may be related to the different ionic conductivities of the electrolytes inducing different cell polarizations. Representative operando diffraction patterns taken in lithium cells are depicted in Figure 7b. Upon the oxidation of $\text{Ca}_3\text{CoMnO}_6$, the evolution observed is fully consistent with that observed in calcium cells, reaching a stage in which both pristine and oxidized phases still coexist for a capacity of 160 mA h/g. Upon reduction, no changes in the patterns are evidenced, which forces us to conclude that the capacity observed is related to side reactions involving the electrolyte. These results point at the irreversibility of the redox process being related to intrinsic structural/migration constraints rather than to slow kinetics or high desolvation energies. Calcium ion rearrangement in the oxidized phase sublattice may hinder mobility in $\text{Ca}_3\text{CoMnO}_6$, as suggested for $\text{Ca}_3\text{Co}_2\text{O}_6$. Indeed, oxidation of $\text{Ca}_3\text{CoO}_6$ and $\text{Ca}_3\text{CoMnO}_6$ does not follow a topotactic mechanism and it is reasonable to assume that the phase transformation that gives rise to the formation of the incommensurate modulated structure induces changes in the Ca migration pathways, with the intercolumn space not making any significant difference.

### CONCLUSIONS

The operando synchrotron diffraction experiments indicate that the extraction of $\text{Ca}^{2+}$ from K$_2$CdCl$_4$-type $\text{Ca}_3\text{CoMO}_6$ (M = Co, Fe, or Mn) structures is achieved at room temperature for M = Co and Mn only. Different incommensurate modulated structures are formed upon oxidation for M = Co and M = Mn, as a result of Mn/Co ordering, with the propagation vector enabling the direct estimation of the calcium content, which is similar for both oxidized phases (ca. 2.3 mol per formula unit). The reversibility of the redox process was found to be extremely limited, if any. Further operando experiments performed in lithium cells also resulted in the electrochemical extraction of calcium from the crystal structure upon oxidation,
but intercalation of Li\(^+\) in Ca\(_{3-x}\)CoMnO\(_6\) upon reduction was not found to be feasible either. Thus, the main factor behind the absence of redox reversibility seems to be related to intrinsic structural/migration constraints rather than to slow kinetics or high desolvation energies commonly associated with divalent ion charge carriers.

Finally, although still not conventionally used in the emerging field of calcium batteries, we believe that the experimental protocols reported herein can be generally extended to the screening of new multivalent ion positive electrode materials and be a valuable tool to unambiguously assess reactivity and reversibility.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.energyfuels.1c01343.

Proposed description of the incommensurate structure of Ca\(_{3-x}\)CoMnO\(_6\) (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**
M.R. Palacín — Institut de Ciència de Materials de Barcelona, ICMAB-CSIC, 08193 Bellaterra, Spain; orcid.org/0000-0001-9465-3106

**Authors**
A.P. Black — Institut de Ciència de Materials de Barcelona, ICMAB-CSIC, 08193 Bellaterra, Spain
D. Monti — Institut de Ciència de Materials de Barcelona, ICMAB-CSIC, 08193 Bellaterra, Spain
C. Frontera — Institut de Ciència de Materials de Barcelona, ICMAB-CSIC, 08193 Bellaterra, Spain; orcid.org/0000-0001-7351-2005; Email: rosa.palacin@icmab.es

D. S. Tchitchekova — Institut de Ciència de Materials de Barcelona, ICMAB-CSIC, 08193 Bellaterra, Spain; orcid.org/0000-0003-0214-4115

R. G. Houdeville — CELLS—ALBA Synchrotron, 08290 Cerdanyola del Vallès, Spain
F. Fauth — CELLS—ALBA Synchrotron, 08290 Cerdanyola del Vallès, Spain; orcid.org/0000-0001-9465-3106

Complete contact information is available at: https://pubs.acs.org/doi/10.1021/acs.energyfuels.1c01343

**Notes**

The authors declare no competing financial interest.

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