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On the strange case of divalent ions intercalation in V₂O₅

R. Verrelli^{a,1}, A.P. Black^{a,1}, C. Pattanathummasid^{a,b}, D.S. Tchitchekova^a, A. Ponrouch^a, J. Oró-Solé^a, C. Frontera^a, F. Bardé^c, P. Rozier^b, M.R. Palacín^{a,*}

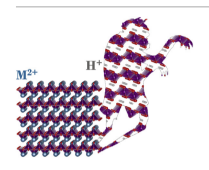
HIGHLIGHTS

- Controversial reports exist in the literature regarding Ca/Mg intercalation in V₂O₅.
- Crystal structure elucidation of intercalated phases has remained elusive to date
- Electrochemical and ex-situ XRD experiments evidence formation of protonated phases.
- Degradation of V₂O₅ has also been observed under some conditions.
- Oxidation of AV₂O₅ (A = Ca, Mg) prepared by solid state reaction was not feasible.

ARTICLE INFO

$$\label{eq:continuous} \begin{split} &\textit{Keywords:} \\ &V_2O_5 \\ &\textit{Mg batteries} \\ &\textit{Ca batteries} \\ &\textit{Multivalent ion intercalation} \end{split}$$

GRAPHICAL ABSTRACT



ABSTRACT

Vanadium pentoxide has been investigated for multivalent ion battery technologies but the structural characterization of inserted phases is poor, and conflicting reports exist in the literature. This study presents a critical overview of controversial aspects related to Mg and Ca insertion in α -V₂O₅ under diverse conditions by combined electrochemical and ex-situ XRD experiments. Galvanostatic tests are carried out in dry and wet alkyl carbonate-based electrolytes at RT and 100 °C. The formation of protonated phases with negligible divalent ion content if any is evidenced by Rietveld refinements of the XRD data, unambiguously dismissing the presence of AV₂O₅ (A: Mg, Ca) as electrochemical reduction products. Furthermore, thermal instability of V₂O₅ at 100 °C in alkyl carbonate solvents is demonstrated by XRD and TEM analysis and the formation of an orthorhombic phase with increased a parameter, most likely due to degradation favored by both water and temperature, is observed for both Mg and Ca. In order to assess the feasibility of the reverse reaction, fully intercalated AV₂O₅ (A = Ca, Mg) phases were also prepared by solid state reaction and oxidation attempted both electrochemically and chemically without evidence of any significant amount of Mg²⁺ or Ca²⁺ extraction, further corroborating the sluggish diffusion kinetics of divalent cations in α -V₂O₅.

1. Introduction

Development of new battery technologies involving multivalent

metal anodes (e.g. Mg, Ca, Al) has recently re-attracted the interest of the scientific community, as these rely on abundant, inexpensive and nontoxic metals and hold promise of very high energy densities [1].

E-mail address: rosa.palacin@icmab.es (M.R. Palacín).

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^a Institut de Ciència de Materials de Barcelona (ICMAB-CSIC), Campus UAB, E-08193, Bellaterra, Catalonia, Spain

^b Université Paul Sabatier, CIRIMAT, UMR CNRS 5085, 118 Route de Narbonne, 31062, Toulouse Cedex, France

^c Toyota Motor Europe, Research & Development 2, Advanced Material Research, Technical Centre, Hoge Wei 33 B, B-1930, Zaventem, Belgium

^{*} Corresponding author.

¹ Authors equally contributed.

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Amongst them, Mg batteries have deserved the most attention, with proof of concept being given as early as 2000 using Chevrel phases as positive electrode material [2], despite commercialization being still elusive due to diverse fundamental bottlenecks [3,4]. On the other hand, the recent reports on the feasibility of Ca plating and stripping [5,6] opened the quest for materials reversibly inserting and extracting Ca²⁺ that could be used as positive electrode to achieve proof of concept of a full Ca based battery. Besides the exploration of new materials [7–9], revisiting traditional layered intercalation hosts appear as a very useful tool to gain further insight into the fundamentals of divalent ion intercalation [10]. Indeed, even if the know how gained in the development of the Li-ion battery technology is precious to accelerate advances in alternative electrochemical energy storage systems, it cannot be directly translated to divalent chemistries, as many of the aspects taken for granted (reliability of reference electrodes, absence of side reactions etc) do not hold when moving to Mg or Ca based chemistries [11] and the use of a wide spectrum of characterization tools is compulsory [12-16] to avoid biased interpretation of results. The case of V₂O₅ is especially intriguing. Despite the expected low multivalent ion mobility in oxide hosts, it seems at first sight well established that it does intercalate both Ca2+ and Mg2+ ions. Nonetheless, the interpretation of some observed trends deserves further attention and some relevant controversies emerge from the available literature [16]. On one hand the addition of water to magnesium electrolytes as "shielding" agent was suggested to promote intercalation of Mg2+ ions but alternative effects of water or hydronium ions have not been unambiguously interpreted [17,18]. On the other hand, elucidation of the intercalated A_xV₂O₅ crystal structure remains elusive, despite some changes in the XRD patterns of pristine orthorhombic V₂O₅ observed after intercalation, which differ from one study to another (see next section).

Energy barriers for multivalent (MV) ion migration and the V₂O₅-MV ion phase diagram have been recently computed [19-21], which should contribute to clarify the situation. According to the first principles calculations performed by Sai Gautam et al. [20], the intercalation of Mg^{2+} in α - V_2O_5 under equilibrium conditions should evolve through a two-phase reaction, with nucleation of δ-Mg_xV₂O₅ (0 < x < 1) from demagnesiated α -V₂O₅, the absence of experimental evidence being attributed to hindered kinetics. Instead, the formation of the metastable ε -phase (with ordering at x = 0.5 resulting in a voltage drop on the electrochemical profile) is more consistent with experiments, since it does not require structural rearrangement of the host structure. Alternative studies [22] predict a phase transformation from α to δ polymorph while discussing a possible competition with Mg²⁺ ion diffusion (with estimated Mg hopping barriers of 1.26 and 0.81 eV for the α and δ phase, respectively). The α and $\delta\text{-}V_2O_5$ polymorphs differ in the stacking of the layers along the a direction, which results in a different coordination environment for the intercalation site (see Fig. 1) . Since Mg2+ migration is predicted to be faster and energetically more favorable in the δ polymorph, the deintercalation of magnesium from δ -MgV₂O₅, which can be directly synthesized by solid

state reaction [23], is suggested as a viable strategy to foster the Mg²⁺ mobility and achieve good electrochemical performance [20].

In the case of Ca^{2+} ions, the available DFT studies [21,24] predict migration barriers of about 1.7–1.9 eV in α -V₂O₅ with an average voltage of about 3.2 V vs Ca^{2+} /Ca. In contrast, the migration barrier in the δ -V₂O₅ phase is calculated to be much lower (0.2 eV). However, this polymorph is not known: CaV_2O_5 prepared by solid state reaction is not isostructural with δ -MgV₂O₅, but rather exhibits the α structure, which enables a more suitable coordination for the larger Ca^{2+} ion [25].

Aiming to move further steps towards the understanding of the critical aspects related to MV-ion intercalation into V_2O_5 , we have investigated the electrochemical behavior of orthorhombic α - V_2O_5 both in Mg and Ca cells, in dry and wet alkyl carbonate-based electrolytes, with a critical focus on the insertion-driven structural changes of the active phase as observed by ex-situ X-ray diffraction.

Approaching the subject from the opposite side, AV_2O_5 phases (A: Mg, Ca) were prepared by solid state synthesis and their oxidation was attempted chemically and electrochemically. This approach, to the best of our knowledge here originally explored, addresses the open questions on the possible structural transformation of fully intercalated AV_2O_5 phases into V_2O_5 (and vice versa), to date not yet thoroughly clarified. The results achieved in this study underpin the importance of a rigorous methodology combining electrochemical and alternative characterization techniques to unambiguously assess MV-ion intercalation.

2. Overview of a controversial background

Orthorhombic α - V_2O_5 is built up by layers of alternating edge- and corner-sharing VO₅ square pyramids linked together through Van der Waals interactions (S.G. Pmmn). It holds potential for multielectron intercalation reactions with the concomitant reduction of V5+ to V4+ and beyond, as demonstrated for Li+ and Na+ [26-28]. Computed voltages are about 2.3 V vs. Mg^{2+}/Mg and 3.2 V vs. Ca^{2+}/Ca for Mg^{2+} and Ca2+ intercalation [21], respectively, with a specific capacity of 294.7 mAh g^{-1} related to the formation of AV_2O_5 (A = Mg, Ca). The first experimental reports on the electrochemical intercalation of Mg²⁺ into V₂O₅ date back to the late '80s, with the pioneering work of Pereira Ramos [29] and Novak [30], achieving reversible Mg insertion in V₂O₅ in Mg(ClO₄)₂ based electrolytes, either in dimethylsulfone at 150 °C or acetonitrile at room temperature (RT). Further attempts to electrochemically intercalate Mg^{2+} in V_2O_5 have been carried out under very diverse experimental conditions (regarding active material morphology, electrolyte composition, temperature, cell set-up and electrochemical protocols). Despite the similarity between Li⁺ and Mg²⁺ ionic radius (76 and 72 pm, respectively), Mg²⁺ intercalation is hindered by its high polarizing character, which is expected to translate in slow migration kinetics and high (de)solvation energy barriers. In contrast with the vast knowledge available about the Li+ intercalation mechanism, little is known in the case of Mg²⁺ insertion in this host,

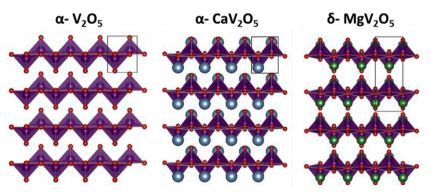


Fig. 1. Crystal structure of the pristine α -V2O5, α -CaV₂O₅ polymorphs on the b-c plane and δ -MgV₂O₅ polymorph on the a-b plane.

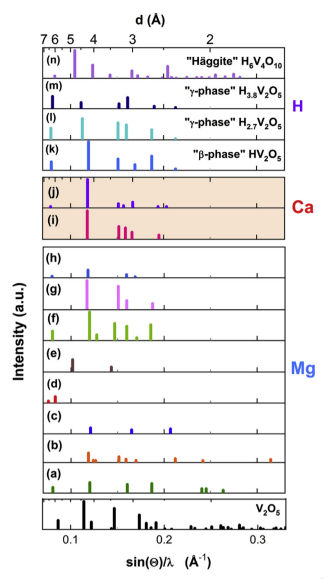


Fig. 2. XRD peaks reported in the literature for the phases formed upon Mg $^{2+}$ (a–h) and Ca $^{2+}$ (i–j) electrochemical intercalation in $\alpha\text{-V}_2O_5$ at RT (Ref. [17,30,33,38,39,43,48]); (k–n) Characteristic XRD reflections of chemically hydrogenated V $_2O_5$ phases (Ref. [44–47]). The pattern below corresponds to $\alpha\text{-V}_2O_5$.

despite a significant number of publications devoted to the topic. The elucidation of the nature and crystal structure of the phase formed is still elusive and the case of ${\rm Ca}^{2^+}$ is even less explored. An overview of the available literature data is provided below with a focus on the crystal chemistry and available diffraction data.

While interpretation of results converge to the existence of topotactic insertion of very low amounts of ${\rm Mg}^{2+}$ in $\alpha\text{-V}_2{\rm O}_5$ without major structural modifications (diffraction peaks being only slightly shifted (or broadened)) [31–38], progressive amorphization with intercalation has also been reported in some cases [17,33]. In contrast, a slight increase of the a cell parameter has been reported after a redox process assumed to result in a significant amount of calcium intercalation in Cainserted $\alpha\text{-V}_2{\rm O}_5$, but no structural model is given for the phase formed [39].

Fig. 2 (panels (a-j)) graphically summarizes the diffraction data available in the literature for presumably intercalated phases, formed upon electrochemical reduction of $\alpha\text{-}V_2O_5$. The peak intensities are rescaled between 0 and 1 (a.u.). The experimental conditions employed in each study are specified in Table 1. The actual interpretation and

assignment of the observed diffraction patterns, which can be significantly different from one study to another, remains elusive in most of the cases. The Mg content within the inserted phases reported in literature has been tentatively assessed by Inductively Coupled Plasma (ICP), Atomic absorption (AAS) and Energy Dispersive (EDAX) spectroscopy or simply deduced from capacity, thus not being completely reliable. Both types of methods may yield to an overestimation of the intercalated magnesium amount: in the case of chemical methods, due to the possible existence of magnesium containing surface layers resilient to washing of the particles [40] and, in the case of electrochemical methods, due to side reactions involving cell components other than the electrode active material (mostly the electrolyte). The formation of δ-MgV₂O₅ and α-CaV₂O₅ isostructural to the phases prepared by solid state reaction is generally not observed, or attribution is based on a single low intensity peak [17]. On the other hand, formation of the metastable ε-Mg_{0.5}V₂O₅ phase has only been detected locally by aberration corrected-STEM imaging [41], despite a much lower intercalation level being experimentally probed by combined atomic resolution STEM, EELS, electron diffraction and EDX techniques.

Novak and De Silvestro [42] and later other authors [17,36,43], speculated that the addition of water to the electrolyte results in enhanced electrochemical response of V2O5 towards magnesium intercalation basing on the higher current observed in cyclic voltammetry experiments and the higher capacity recorded. This hypothesis is related to the ability of H₂O molecules to strongly coordinate to magnesium ions and hence shield their high polarizing character, thus allowing faster diffusion kinetics within the host material [16]. However, the possibility of proton (co-)intercalation has never been fully dismissed and recent solid state NMR studies on electrochemically reduced (magnesiated) orthorhombic $\alpha\text{-}V_2O_5$ electrodes demonstrated that proton is indeed intercalated in a wet electrolyte (1 M Mg(TFSI)2 in diglyme, with 2600 ppm of H₂O) [17]. These results evidence the need to consider the possible formation of hydrogenated H_vV₂O₅ and/or coinserted Mg_vH_vV₂O₅ phases for the interpretation of the XRD patterns of electrochemically reduced V₂O₅. Indeed, structural studies on the chemical hydrogenation of V₂O₅ [44] revealed that proton is initially topotactically inserted in V_2O_5 to form α -H_X V_2O_5 (x < 0.4) phase with cell parameters only slightly different from the ones of pristine V₂O₅. At higher H⁺ concentrations (i.e. 0.4 < x < 3.2 and 3.2 < x < 3.9), two novel orthorhombic phases are progressively formed (β and γ, respectively), with a quasi-amorphous compound (y) obtained for x $(H^+) > 3.8$ [44]. The characteristic reflections of the hydrogenated β and γ H_XV₂O₅ phases are displayed in Fig. 2 (Panels k and l-m, respectively). This figure also exhibits the diffraction pattern of a monoclinic, "Häggite" phase with composition H₆V₄O₁₀ (Fig. 2- Panel n), a protonated oxide reported in 1958 by Evans and Mrose [45,46] and further observed by Srivastava et al. [47] upon chemical hydrogenation of Pt-activated V2O5 at 150 °C.

The similarities between the diffraction patterns observed after attempts of electrochemical Mg and Ca intercalation and those of the different phases formed by chemical hydrogenation of V2O5 raise a question on the actual contribution of protons to the overall intercalation process, which can only be carefully assessed through additional characterization techniques. Despite promising steps in this direction [17,18], detailed structural characterization of the phases formed is only starting. Interesting results have been reported by Hong et al. [43] in the course of the present work, reporting the formation of a $Mg_{0.17}H_xV_2O_5$ (0.66 < x < 1.16) phase upon reduction of submicrometric α-V₂O₅ sample in beaker-type (flooded) cell at room temperature, using $0.5 \,\mathrm{M}\,\mathrm{Mg}(\mathrm{ClO_4})_2 + 2 \,\mathrm{M}\,\mathrm{H_2O}$ in ACN (acetonitrile) electrolyte (Panel g of Fig. 2). The structural model of the inserted phase is resolved through ab-initio methods, by combined synchrotron XRD data and DFT calculations. The formation of another phase is observed upon further reduction, which resulted to be difficult to characterize (especially considering that the intensity of the peaks attributed to this phase increases upon reoxidation to about 0.2 V vs Ag⁺/

Table 1 Experimental conditions of the electrochemical intercalation tests recalled in Fig. 2 and assignments made to explain the additional diffraction peaks. ACN: Acetonitrile, G2: Diglyme, PC: Propylene Carbonate. Commercial V_2O_5 exhibits micrometric particle size.*:Phase observed after full reduction $(1.8 \le \Delta x \le 2 \text{ mol e}^-)$ and following oxidation $(2 \le \Delta x \le 1 \text{ mol e}^-)$;#: phase observed for $0.5 \le \Delta x \le 2 \text{ mol e}^-$; $^{\circ}$: after reduction (2.5 cycles) and $^{\circ}$: after reduction (2.5 cycles).

	W.E.	Electrolyte	C.E	Assigned Phase	Ref.
j	Commercial V ₂ O ₅	0.5 M Ca(ClO ₄) ₂ , ACN	CaCo ₂ O ₄	Ca _X V ₂ O ₅	[48]
i	Commercial V ₂ O ₅	1 M Ca(ClO ₄) ₂ , ACN	Ca	$Ca_{0.2}V_2O_5$	[39]
h	Commercial V ₂ O ₅	0.5 M Mg(TFSI) ₂ , PC	Coated Mg	Unknown	[38]
g	Submicrometric V ₂ O ₅	$0.5 \mathrm{MMg(ClO_4)_2},\mathrm{ACN}+2\mathrm{M}\mathrm{H_2O}$	Activated C	Unknown*	[43]
f	Submicrometric V ₂ O ₅	$0.5 \mathrm{MMg(ClO_4)_2},\mathrm{ACN}+2\mathrm{M}\mathrm{H_2O}$	Activated C	$Mg_{0.17} H_x V_2 O_5^{\#} (0.66 < x < 1.16)$	[43]
e	Commercial V ₂ O ₅	1.0 M Mg(TFSI) ₂ /G2 (15 ppm H ₂ O)	Mg	MgV_2O_5	[17]
d	Commercial V ₂ O ₅	1.0 M Mg(TFSI) ₂ /G2 (15 ppm H ₂ O)	Mg	Unknown	[17]
c	V ₂ O ₅ thin film	0.1 M Mg(TFSI)2, ACN (32 ppm H2O)	Activated C	Unknown	[33]
b	Commercial V ₂ O ₅	$1 \text{ M Mg(ClO}_4)_2$, ACN + $1 \text{ M H}_2\text{O}$	Mg	$Mg_xV_2O_5$	[30]
a	Commercial V ₂ O ₅	$1 \text{ M Mg}(\text{ClO}_4)_2, \text{ ACN } + 1 \text{ M H}_2\text{O}$	Mg	$Mg_xV_2O_5^{\alpha}$	[30]

Ag). However, pristine, orthorhombic α -V₂O₅ was recovered upon subsequent oxidation. Surprisingly, authors state that the formation of Mg_{0.17}H_xV₂O₅ (together with an additional unknown phase) does also take place when the reduction is carried out in dry (2 ppm H₂O), 0.5 M Mg(ClO₄)₂ ACN electrolyte, without the formation of any other phase upon further reduction. At this point the need for a careful evaluation of the actual water content in the electrolytes is worth recalling, since conventional drying methods may not be adequate for some of the compositions used [43,49]. Moreover, alternative side reactions involving other proton sources (e.g. electrolyte breakdown reactions) should also be considered.

3. Experimental

3.1. Materials and synthesis

Commercial V_2O_5 (Sigma Aldrich, 99.6%), with micrometric particle size, was used as active material. MgV_2O_5 was synthesized either by i) solid state reaction between MgO and VO_2 [23] or by ii) comproportionation of MgV_2O_4 and MgV_2O_6 . The conditions for these reactions are detailed in the following paragraphs:

- i) A stoichiometric powder mixture of pre-heated (500 °C, 2 h) MgO and VO $_2$ was annealed for 6 h at 900 °C. VO $_2$ was prepared by comproportionation of V $_2$ O $_5$ and V $_2$ O $_3$. Stoichiometric amounts of V $_2$ O $_3$ and V $_2$ O $_5$ powders were mixed and ground together and further annealed at 650 °C for 10 h under vacuum. Platinum sheets rolled in an Al $_2$ O $_3$ crucible were employed as sample holders. The V $_2$ O $_3$ precursor was prepared by reduction of V $_2$ O $_5$ under pure H $_2$ gas flow (100 ml/min) generated by water electrolysis. Two consecutive 2 h annealing steps at 500 and 700 °C were applied and the sample was further naturally cooled under H $_2$ gas flow for about 10 h
- ii) Stoichiometric amounts of MgV_2O_6 and MgV_2O_4 were ground together and annealed at 770 °C for 12 h under vacuum. The MgV_2O_6 precursor was prepared by mixing V_2O_5 with a Mg^{2+} source such as MgO, $Mg(OH)_2$, or $4MgCO_3 \cdot Mg(OH)_2 \cdot 5H_2O$ (independently from the Mg source employed, the same product was formed). The mixture of precursors, placed in Pt sheets rolled in Al_2O_3 crucible, was calcinated for 15 h at 650 °C in air. MgV_2O_4 was prepared by reduction of MgV_2O_6 under H_2 gas at 700 °C for 4 h.

Preparation of $Mg_xV_2O_5$ solid solutions (x = 0.7 and 0.9) was attempted by solid state reaction of MgV_2O_5 and V_2O_5 powders (mixed in x: (1-x) molar ratio) or, else, by reacting MgO, VO_2 and V_2O_5 in x: 2x: (1-x) molar ratio. In both cases, the mixed precursors were annealed at 900 °C for 6 h, under vacuum. All these resulted in mixtures of mostly MgV_2O_5 and VO_2 and thus, will not be discussed further. Alternative efforts to achieve $Mg_xV_2O_5$ (x \leq 1) by chemical reduction of V_2O_5 with MgI_2 (with V_2O_5 : MgI_2 molar ratios of 1:0.5 and 1:1 employed) also

failed. CaV_2O_5 was prepared by a comproportionation reaction from CaV_2O_4 and CaV_2O_6 . Repeated annealing treatments of the mixed powder precursors were performed under vacuum at 650 °C (two 10 h steps), 750 °C (12 h) and 800 °C (two 12 h steps). Pt sheets rolled in Al_2O_3 crucibles were used as sample holders, except for the last annealing step, in which Mo crucible was employed. CaV_2O_6 was prepared by solid state reaction between CaCO_3 and V_2O_5 (650 °C, 12 h, air atmosphere) while CaV_2O_4 was obtained by reduction of CaV_2O_6 under H_2 gas at 700 °C for 4 h. Attempts to chemically oxidize MgV_2O_5 and CaV_2O_5 powders were made by reaction with NO_2BF_4 (100% mol excess) at 80 °C in ACN for 6 and 24 h under uninterrupted Ar bubbling. Once the reactions were completed, the samples were washed with dry ACN, vacuum filtered, dried and collected for XRD.

3.2. Structural, morphological and compositional characterization

X-Ray powder diffraction patterns were acquired in a Bruker D8 Advance A25 diffractometer in a Debye Scherrer configuration equipped with Mo K α 1 radiation source (λ = 0.7093 Å) and Johansson monochromator.

High resolution synchrotron X-ray diffraction (SXRD) patterns were collected at MSPD beamline (ALBA synchrotron Light Source, Cerdanyola del Vallès, Spain) using Mythen detector and $\lambda=0.6199~\textrm{Å}.$ 0.5 mm diameter borosilicate glass capillaries were used as sample holders and rotated during data collection.

Scanning Electron Microscopy (SEM) analyses were performed with a FEI Quanta 200 FEG microscope.

Selected area electron diffraction (SAED) studies were performed using a JEOL 1210 transmission electron microscope operating at 120 kV and equipped with a Gathan sample holder. The specimens for electron microscopy were prepared dispersing the powders in dimethyl carbonate and depositing a droplet of this suspension on a carbon coated film supported on a copper grid.

3.3. Electrochemical tests

Slurries were prepared by dispersing powder mixtures of active material (80% wt), Super P carbon (10% wt) and polyvinylidene difluoride (PVdF, Kynar) binder (10% wt) in N-Methyl Pyrrolidinone solvent (NMP, Sigma-Aldrich). The dispersion was ball milled for 1 h at 500 Hz with a RETSCH PM100 miller before being doctor-blade casted on Al foil (Goodfellow, 99%). Once casted, the tapes were dried at 120 °C under vacuum and cut into discs of 11 mm diameter, with average active material load of about 2 mg. These electrodes were pressed at 8 Tons before being used.

The electrochemical tests were performed in three electrode Swagelok cells assembled in Ar-filled dry box, employing Al and stainless steel current collectors for the working and reference(RE)/counter(CE) electrodes, respectively, and Whatman borosilicate glass fiber separator soaked in 0.6 ml of electrolyte solution. Mg and Ca metal

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discs were used as both counter and reference electrodes.

Dry (< 25 ppm H₂O) and wet (> 15000 ppm H₂O) solutions of 0.3 M Magnesium bis trifluoromethane sulphonyl imide (Mg(TFSI)₂, Solvionic, 99,5%) salt in 1:1 v mixture of ethylene carbonate (EC) and propylene carbonate (PC) (Solvionic, 99.9%) and 0.3 M Magnesium perchlorate (Mg(ClO₄)₂, Alfa Aesar) in EC:PC (1:1 v) were used as electrolytes for the electrochemical tests in Mg cells. In addition to those, wet (> 15000 ppm H₂O) 0.3 M Mg(ClO₄)₂ in ACN was also used for comparative purposes. The electrolyte solutions used in Ca experiments were 0.45 M Ca(BF₄)₂, 0.3 M Ca(ClO₄)₂ and Ca(TFSI)₂ in EC:PC (1:1 v) and 0.3 M Ca(ClO₄)₂ in ACN with H₂O content < 25 ppm in all cases. For the sake of comparison, tests were also made with the same electrolytes exhibiting > 5000 ppm of water. The water contents of the electrolytes were measured by Karl-Fisher titration.

As Mg plating cannot be achieved in Mg(TFSI) $_2$, EC:PC electrolyte at 100 °C, the electrochemical oxidation of the MgV $_2$ O $_5$ phase was carried out in Li cells with Li metal as counter and reference electrodes at RT with 1 M LiPF $_6$ in EC:DMC (1:1 v) (LP30, Solvionic) or at 100 °C using an electrolyte compatible with high temperature studies: 0.1 M Lithium Bis Oxalate Borate (LiBOB, Sigma Aldrich) in EC:PC (1:1 v).

The electrochemical tests were performed on Bio-Logic VMP3 and MPG2 potentiostat/galvanostats, with either potentiodynamic cycling with galvanostatic acceleration (PCGA) technique, applying potential steps of 5 mV and current threshold of C/100 (equivalent to intercalation of one mol of divalent ions in 100 h) in order to ensure full reactivity at each step, or by Galvanostatic Cycling with Potential Limitation (GCPL) at C/100 rate. Cells were left for 5 h at the open circuit potential (OCP) prior to measurements to enable temperature equilibration.

Once tested, the cells were disassembled in Ar-filled dry box and the electrodes were washed with Dimethyl Carbonate solvent (DMC, Sigma-Aldrich, \geq 99%) and further collected in borosilicate glass capillaries for ex-situ XRD measurements, as mentioned above.

4. Results

4.1. Electrochemical tests

Aiming to clarify the viability of α - V_2O_5 as a reference cathode material for divalent ion battery technologies, galvanostatic cycling (GCPL) measurements were performed on V_2O_5 electrodes in Mg and Ca cells using dry alkyl carbonate-based electrolyte solutions. As no electrochemical activity was observed neither at RT nor at 55 °C, moderate temperatures of 100 °C were employed to enhance the diffusion kinetics of Mg^{2+} and Ca^{2+} (note that the efficiency of calcium plating/stripping in $CaBF_4$ in EC:PC was found to be higher at this temperature than at 50 °C) [5].

The characteristic potential vs. composition profile of V_2O_5 electrodes subjected to GCPL tests at $100\,^{\circ}\text{C}$ and C/100 rate in Mg cells employing both $0.3\,\text{M}\,\text{Mg}(\text{ClO}_4)_2$ and Mg(TFSI) $_2$ in EC:PC electrolytes is displayed in Fig. 3(A). The ex-situ XRD diffraction patterns of the electrodes recovered at different stages of reduction and re-oxidation are displayed in Fig. 3(B). Analogous experiments were carried out in Ca cells using $0.3\,\text{M}\,\text{Ca}(\text{TFSI})_2,\,0.3\,\text{M}\,\text{Ca}(\text{ClO}_4)_2$ and $0.45\,\text{M}\,\text{Ca}(\text{BF}_4)_2$ in EC:PC (1:1 v) and 1 M Ca(ClO $_4$) $_2$ in ACN as electrolytes (Fig. 3(C)) and the corresponding ex-situ XRD patterns are presented in Fig. 3(D).

In Mg cells, the characteristic potential vs. capacity profile for V_2O_5 evolves through a continuous sloping trend upon reduction and oxidation (Fig. 3(A)). The capacity observed is much higher than the theoretical value for full reduction of V^{+5} to V^{+4} (i.e. 294.7 mAh g $^{-1}$), which may reasonably be related to electrolyte decomposition side reactions, expected to be enhanced in the testing conditions employed (high temperature and very low rate). The corresponding ex-situ XRD patterns at different redox stages displayed in Fig. 3(B) clearly reveal the appearance of extra peaks upon magnesiation which disappear upon oxidation up to the recovery of pristine V_2O_5 . The capacity upon

reoxidation is slightly higher than theoretical, which may indicate some thermally enhanced electrolyte oxidation even at this moderate potential. At early reduction stages in Mg(ClO₄)₂, EC:PC electrolyte (capacity close to 140 mAh g⁻¹), extra peaks were observed at d values of 3.1, 3.3, 4.2 and 6.2 Å, despite $\rm V_2O_5$ being still the major phase. These peak positions are consistent with some of the patterns reported in the literature for presumably Mg²⁺ or Ca²⁺ intercalated V₂O₅, which seem also closely related to protonated H_xV₂O₅ phases prepared chemically (Fig. 2, panels f-j and panels l-m, respectively). This set of reflections was tentatively indexed with an orthorhombic Pmmn symmetry (see Fig. 4(A)) and refined using the structural model reported by Yoshikawa et al. [44] The cell parameters obtained were a = 12.393(3). b = 3.4070(1) and c = 4.2675(1) Å (Table 2). Despite the impossibility to refine the hydrogen occupation from XRD data and the low signal to noise ratio of the pattern corresponding to the reduced sample, refinements considering a possible partial occupation of magnesium in the proton positions led to larger χ^2 values likely indicating absence of Mg²⁺ in the phase. Upon further reduction (total capacity close to 500 mAh g^{-1}), additional extra peaks formed at d = 2.45, 3.07, 4.11, 4.84 Å, while the reflections previously observed at 3.1, 3.3, 4.2 and $6.2\,\text{Å}$ and those corresponding to V_2O_5 exhibit very low intensity if any. These peaks are consistent with the formation of a protonated Häggitetype phase $(H_xV_4O_{10})$, with C2/m symmetry (see its crystal structure in Fig. S2 in S.I.). The cell parameters obtained from Rietveld refinement (Fig. 4(B)) are a = 12.437(5) Å, b = 3.043(1) Å, c = 4.926(3) Å and $\beta = 98.48(3)$ Å. Note that refinement attempts placing Mg²⁺ in the proton position led to negative occupation values. Nonetheless, given the low crystallinity of the sample, the possible presence of Mg²⁺ in very low amount could not be conclusively dismissed. Similar results were obtained in Mg(TFSI)2, EC:PC electrolyte, for which the ex-situ XRD patterns at different redox stages are shown in Fig. 3(B) (red patterns). In this case, the Häggite-type phase is the only product observed upon full reduction (500 mAh/g). Similar XRD patterns were also obtained after GCPL tests on cells with magnesiated active carbon as counter electrode and for both Mg(ClO₄)₂ and Mg(TFSI)₂ based electrolytes, allowing to detect the formation of the Häggite-type phase again after a second reduction.

The potential vs. capacity profile for experiments carried out in Ca cells at C/100 rate, depicted in Fig. 3(C), revealed to be significantly dependent on the electrolyte used: for Ca(TFSI)2 and Ca(ClO4)2 in EC:PC 1:1v the profile follows a sloping trend down to specific capacity values between 180 and 250 mAhg⁻¹ while for Ca(ClO₄)₂ in ACN at RT, a plateau at 1.7 V vs. Ca²⁺/Ca is observed up to specific capacities of 300 mAh g^{-1} followed by a sloping region up to 430 mAh g^{-1} at 0 V vs. Ca²⁺/Ca in agreement with previous reports [39]. In the case of Ca $(BF_4)_2$ in EC:PC a plateau is reached at 1.2 V vs. Ca^{2+}/Ca which extends to a specific capacity of 550 mAh g^{-1} . It is worth mentioning that the OCP observed in Ca(BF₄)₂ electrolyte is approximatively 0.7 V higher than those observed for the other three electrolytes, which might be related to the formation of different passivation layers on the Ca RE. The ex-situ XRD patterns of the V2O5 electrodes taken at different reduction stages are displayed in Fig. 3(D). No additional reflections are observed at early stages of reduction (> 1.2 V vs. Ca²⁺/Ca, < 100 mAh g⁻¹) in any of the electrolytes tested and only a very minor change in lattice parameters of V₂O₅, if any, is detected (see Fig. S1 in S.I.), which points at ascribing the observed capacity to side reactions most likely involving the electrolyte. Upon further reduction additional peaks are observed at 3.1, 3.3, 4.2 and 6.2 Å (see Fig. 3(D) patterns (b), (d), (h) and (i)]. These reflections, appearing at the same d values as those observed for Mg cells (see Fig. 3(B)), seem compatible with the patterns of electrochemically reduced and chemically protonated V2O5 phases reported in literature (Fig. 2, panels f-j and 2, panels l-m, respectively). Only in the case of Ca(ClO₄)₂ in EC:PC 1:1 v the appearance of other peaks, the most intense at d = 4.85 Å is observed upon full reduction (see Fig. 3(D) pattern (d)), which seems to correspond to the Häggite phase (H_xV₄O₁₀) observed in the Mg experiments previously discussed R. Verrelli et al.

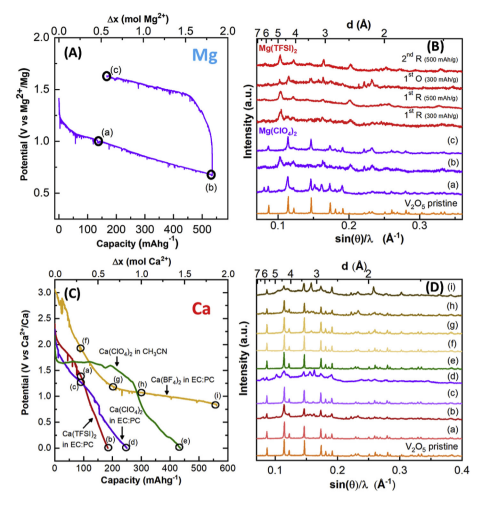


Fig. 3. (A) Characteristic GCPL voltage profile versus specific capacity (bottom axis) and versus moles of inserted ion (Δx) (top axis), considering that all electrochemical response is related to divalent ion intercalation, of a Mg/0.3 M Mg(ClO₄)₂, EC:PC (1:1 v)/V₂O₅ cell (C/100 rate, 100 °C) and corresponding ex-situ XRD patterns (B). The XRD patterns of V₂O₅ upon cycling in Mg(TFSI)₂ electrolyte (C/100 rate, 100 °C) are also shown in (B); (C) Characteristic GCPL voltage profile (C/100 rate) versus specific capacity (bottom axes) and versus hypothetical number of moles of inserted ion (Δx) (top axes) of Ca/0.3 M Ca(TFSI)2, EC:PC (1:1 v)/V2O5 cell, 100 °C (red line), Ca/0.3 M Ca(ClO₄)₂, EC:PC (1:1 v)/V₂O₅ cell, 100 °C (purple line), Ca/1 M Ca(ClO₄)₂ EC:PC (1:1 v)/V₂O₅ cell, 100 °C, Ca/0.3 M Ca(ClO₄)₂, ACN/ V₂O₅ cell, RT (green line), Ca/0.45 M Ca(BF₄)₂, EC:PC (1:1 v)/V2O5 cell, 100°C (yellow line) and corresponding ex-situ XRD patterns (D) The pattern of pristine V₂O₅ is also given for comparison. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

(see Fig. 3(B)). After significant reduction in $Ca(BF_4)_2$ electrolyte (Fig. 3(D) pattern (i)) two peaks at d values of 1.93 and 3.16 Å are indicative of the formation of CaF_2 . Thus, we speculate that the long plateau observed in the GCPL curve is related to the electrolyte decomposition. Finally, despite the capacity observed upon reduction at RT using $Ca(ClO_4)_2$ in ACN as electrolyte, no changes in the XRD pattern can be detected (see Fig. 3(E) pattern (e)) and hence we are tempted to again ascribe it to side reactions involving the electrolyte.

The effect of water impurities in the electrolyte on the electrochemical response of the V_2O_5 electrodes in Mg and Ca cells was investigated by performing galvanostatic cycling at C/100 rate and both RT and 100 °C in wet electrolytes (water content between 4000 ppm and 18000 ppm) (Fig. S3) and carrying out ex-situ XRD experiments (Fig. 5(A) and (B)). Independently from the temperature, enhanced specific capacities are recorded in the water containing electrolytes as compared to the dry ones (See Fig. S3 (A) and (B) in the S.I.). For instance, in Mg cells (Mg(ClO₄)₂,EC:PC; 100 °C) about 500 mAh g $^{-1}$ were recorded when V_2O_5 was reduced down to 0.7 V vs Mg $^{2+}$ /Mg in dry electrolyte (Fig. 3A), while cycling in wet electrolyte led to a capacity of almost 600 mAh g $^{-1}$ with a cutoff voltage above 1 V vs Mg $^{2+}$ /Mg (Fig. S3A) reasonably including contributions from uncontrolled side reactions (possibly including water reduction).

The lattice parameters of V_2O_5 remained virtually unchanged after the reduction tests performed in both Ca and Mg cells. The appearance of additional reflections at d values of 3.2, 4.15 and 6.4 Å was observed upon both presumed magnesiation and calciation, independently from the electrolyte and temperature. These values are close to those observed after galvanostatic reduction tests in dry Mg and Ca electrolytes previously discussed and may be tentatively assigned to an

orthorhombic phase related to γ -H_xV₂O₅. It is important to highlight that this orthorhombic V₂O₅ related secondary phase, characterized by a larger a cell parameter, appears in a meaningful fraction at room temperature in presence of water (Fig. 5(A) patterns (a), (b) and (d)) and its formation seems to be favored by both water and temperature.

Furthermore, the main peak of the Häggite phase $(H_xV_4O_{10})$ at d value of 4.8 Å was also observed in the samples reduced in Mg(ClO₄)₂, EC:PC + H₂O at 100 °C and in wet Ca(TFSI)₂ and Ca (ClO₄)₂EC:PC + H₂O electrolytes at RT (Fig. 5(A)-pattern (c) and Fig. 5(B) patterns (b) and (c), respectively). In the case of wet Ca(BF₄)₂ electrolyte, peaks corresponding to V₂O₅ were no longer observable and the pattern presented only the peaks at d = 3.2, 4.15 and 6.4 Å. The position and intensities of these peaks seen again compatible with a phase related to γ -H_xV₂O₅ with an a parameter further expanded to approximatively 12.86 Å inducing the reflections (110) and (400) to merge (see Fig. S9 in S.I.).

4.2. Evolution of electrodes at OCP

Aiming to detect any possible evolution of the $\alpha\text{-}V_2O_5$ electrodes resulting from contact with the electrolyte that could bias the results of the electrochemical tests, different cells were assembled and kept at OCP (i.e. without any current applied) at different temperatures for various periods of time. The cells were then opened and the electrodes recovered and characterized. Fig. 6(A) and (B) display the corresponding ex-situ XRD patterns of V_2O_5 electrodes, in Mg and Ca cells respectively.

As noticeable in Fig. 6(A) (pattern (c), extra peaks at d values of 6.2, 4.9, 3.3 and 3.1 Å, were observed after $100\,h$ at OCP at $100\,^{\circ}C$ in

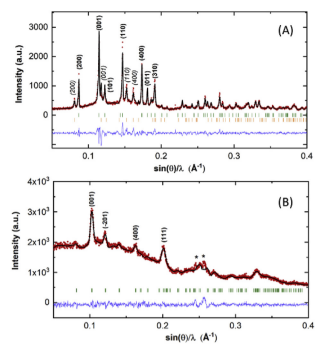


Fig. 4. Observed and calculated X-ray powder diffraction patterns of V_2O_5 tape electrode **(A)** partially reduced (140 mAh g $^{-1}$) in 0.3 M Mg(ClO₄)₂ EC:PC (1:1 v), **(B)** at full reduction (500 mAhg $^{-1}$) in 0.3 M Mg(TFSI)₂ EC:PC (1:1v). Tick marks indicate allowed reflections for α- V_2O_5 (Upper set in **(A)**), H_xV₂O₅ (lower set in **(A)**) and Häggite phase H₆V₄O₁₀ (green ticks in **(B)**). Reflections indicated with * are instrumental contribution due to direct beam stoppers. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

Table 2 Crystallographic data refined for V_2O_5 electrode after partial reduction (140 mAh g $^{-1}$) in 0.3 M Mg(ClO₄)₂ EC:PC (1:1 v) and full reduction (500 mAh g $^{-1}$) in 0.3 M Mg(TFSI)₂ EC:PC (1:1 v).

	α -V ₂ O ₅	γ -H _x V ₂ O ₅	$H_6V_4O_{10}$
Radiation System	X-ray Mo k _{α1} Orthorhombic	X-ray Mo k _{α1} Orthorhombic	X-ray Mo k _{α1} Monoclinic
Space group	Pmmn	Pmmn	C2/m
a (Å) b (Å)	11.540(2) 3.5697(4)	12.393(3) 3.4070(1)	12.437(5) 3.043(1)
c (Å)	4.3703(6)	4.2675(1)	4.926(3)
β (°)	90.0	90.0	98.48(3)

0.3 M Mg(TFSI)₂, EC:PC 1:1 v electrolyte. Such peaks were not observed when the tests were carried out at RT (a) and at 100 °C for shorter time (i.e. 24 h, pattern (b)). Furthermore, a progressive phase amorphization could be detected at 100 °C. In contrast, no evolution of the diffraction pattern was detected in 0.3 M Mg(ClO₄)₂, EC:PC 1:1 v electrolyte even after OCP for 100 h at 100 °C (d), which indicates possible effects of the electrolyte salt anion on the reactivity towards the electrolyte. The exsitu XRD patterns after 4 days OCP in Ca cells at RT in dry 0.3 M Ca (TFSI)₂, 0.3 M Ca(ClO₄)₂, 0.45 M Ca(BF₄)₂ in EC:PC 1:1 v or 0.3 M Ca (ClO₄)₂ in ACN electrolytes did not present any observable sign of degradation (Fig. S4 S.I). Moreover, no degradation was noticed even after 7 days in 0.3 M Ca(TFSI)₂ and 0.3 M Ca(ClO₄)₂ at 100 °C in EC:PC 1:1 v as shown in Fig. 6(B) (pattern (a) and (b)). The peaks at d values of 6.2, 4.9, 3.3 Å could be observed when 0.45 M Ca(BF₄)₂ in EC:PC was used, which points also to an effect of the electrolyte salt anion on the overall active phase/electrolyte side reactivity. Interestingly, the positions of these reflections are similar to those observed in early stages of reduction in several Mg²⁺ and Ca²⁺ dry electrolytes (see Fig. 3(B) and (D) (pattern (b) and (d), (h), respectively) and, in a greater fraction, in all electrodes reduced in wet electrolytes, independently of the electrolyte and temperature. As discussed earlier, these peaks are consistent with the patterns reported in the literature for presumably Mg²⁺ or Ca²⁺ intercalated V₂O₅ which seem closely related with those of chemically obtained H_xV₂O₅ phases (Fig. 2, panels f-j and panels l-m, respectively). Such findings seem again to indicate that the orthorhombic V2O5 related phase, characterized by a larger a cell parameter, is most likely related to a degradation of V2O5 electrochemically favored in wet electrolyte at high temperature but also spontaneously formed upon prolonged exposure to dry electrolyte.

4.3. V_2O_5 stability

With the aim to rationalize the results discussed above and to understand the effect of the different electrolyte components (salt and solvent) in the reactivity observed, further experiments were carried out by simply soaking pristine V_2O_5 powder at RT and $100\,^{\circ}\text{C}$ in diverse electrolytes including bare solvents. Changes in the suspension color, indicative of possible reduction of V^{+5} , were observed to gradually grow upon time at $100\,^{\circ}\text{C}$, even in pure alkylcarbonate solvents. The XRD patterns of such samples revealed the presence of a set of additional peaks, the most intense at $d=3.2\,\text{Å}$. These peaks are found to grow with time. After 6 weeks in EC:PC (1:1 v) at $100\,^{\circ}\text{C}$ the new phase with main reflections at d=3.2, 4.15 and $6.4\,\text{Å}$ is predominant and peaks corresponding to pristine V_2O_5 have very low intensity. (See Figs. S5 and S6 in the S.I.). These stability test has evidenced that this orthorhombic V_2O_5 related phase observed after $100\,\text{h}$ at OCP at $100\,^{\circ}\text{C}$ in $0.3\,\text{M}\,\text{Mg}(\text{TFSI})_2$ or $\text{Ca}(\text{BF}_4)_2$, EC:PC 1:1 v and in reduction in wet or dry

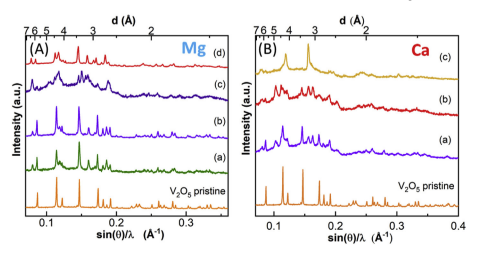
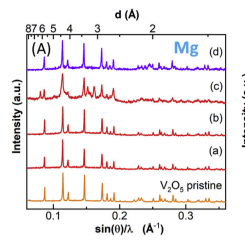


Fig. 5. (A) Ex-situ XRD patterns of the V_2O_5 electrode after galvanostatic reduction at C/100 rate in 0.5 M Mg(ClO₄)₂, ACN + 1 M H₂O at RT (500 mAh/g): (a) 0.3 M Mg(ClO₄)₂, EC:PC + 1 M H₂O at RT (200 mAh/g); (b) 100 °C (600 mAh/g); (c) in 0.3 M Mg(TFSI)₂, EC:PC + 1 M H₂O at RT (120 mAh/g) (d). (B) Ex-situ XRD patterns of V_2O_5 after reduction at 100 °C and C/100 rate in (a) 0,3 M Ca (ClO₄)₂, EC:PC + H₂O (280 mAh/g) (b) 0,3 M Ca (TFSI)₂, EC:PC: H₂O (410 mAh/g) and (c) 0,45 M Ca (BF₄)₂, EC:PC + H₂O (390 mAh/g).

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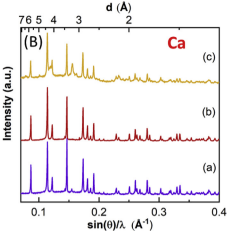


Fig. 6. Ex-situ XRD patterns of the V_2O_5 electrodes after OCP tests in Mg (A) and Ca (B) 2-electrode cells. (A) $0.3\,\mathrm{M}\,\mathrm{Mg}(\mathrm{TFSI})_2$, EC:PC 1:1 v (red plots): (a) $100\,\mathrm{h}$ at RT, (b) $24\,\mathrm{h}$ at $100\,^\circ\mathrm{C}$, (c) $100\,\mathrm{h}$ at $100\,^\circ\mathrm{C}$ and $0.3\,\mathrm{M}\,\mathrm{Mg}(\mathrm{ClO_4})_2$, (d) EC:PC 1:1 v (violet plot) after $100\,\mathrm{h}$ OCP at $100\,^\circ\mathrm{C}$. The diffraction pattern of pristine V_2O_5 phase is also given for comparison. (B) Ex-situ XRD patterns after 1 week OCV at $100\,^\circ\mathrm{C}$ in (a) $0.3\,\mathrm{M}\,\mathrm{Ca}(\mathrm{ClO_4})_2/\mathrm{EC:PC}$, (b) $0.3\,\mathrm{M}\,\mathrm{Ca}(\mathrm{TFSI})_2/\mathrm{EC:PC}$, (c) $0.45\,\mathrm{M}\,\mathrm{Ca}(\mathrm{BF_4})_2/\mathrm{EC:PC}$. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

electrolyte are also spontaneously formed after long exposition of V_2O_5 to pure dry EC:PC electrolyte at 100 °C and are thus presumably related to the same phase(s). However, no significant evolution could be detected in the XRD patterns of the powders soaked for 1 week in an electrolyte containing Mg(TFSI)_2 and Mg(ClO_4)_2 based electrolytes, which indicates that the degree of degradation, if any, was small or either resulted in amorphous side products.

Even if a thorough understanding of the possible mechanisms underneath the V2O5 reductive decompositions goes beyond the scope of the present study, additional experiments aiming at shedding some light on the transformations undergone by V_2O_5 at 100 °C in alkyl carbonate-based electrolytes were conducted. Since a previous TEM study of the V₂O₅ thermal decomposition process [50] revealed that VO_v and 7- and 24-fold superstructure phases related to oxygen loss and vacancy ordering could be locally observed in V₂O₅ samples annealed at 100 °C for 5 days, we decided to carry out electron diffraction to further characterize the phase resulting from V₂O₅ reactivity with the electrolyte. Fig. 7 displays a comparison of the representative SAED patterns along the [001] and [012] zone axes of the V2O5 powder sample in its pristine state ((A) and (C), respectively) and after 100 h at OCP at 100 °C ((B) and (D), respectively). As expected, reconstruction of the reciprocal space, obtained by tilting around the a* axis for pristine V₂O₅, led to the determination of the cell parameters (a = 11.53 Å, b = 3.59 Å and c = 4.36 Å) with the observed reflections being consistent with the orthorhombic space group Pmmn (No. 59) [51]. In contrast, an additional phase was detected in the sample stored for 100 h at OCP at 100 °C. The reconstruction of the reciprocal space from SAED patterns yielded an orthorhombic cell with cell parameters a = 12.39 Å b = 3.44 Å and c = 8.25 Å, which are somewhat related to those of pristine V_2O_5 (a = 11.53 Å, b = 3.59 Å, c = 4.36 Å), with a longer a parameter and doubling of c. Furthermore, weak additional reflections are observed in several planes, such as those marked with orange arrows in the [001] pattern of Fig. 7(B), which, according to Tilley et al. [50], can be interpreted as twins due to the presence of a VO_v phase.

Besides giving further proof of the reactivity of V_2O_5 at moderate temperature (100 °C) in contact with electrolyte components, these observations open the room for speculation about reaction mechanisms. Indeed, in addition to chemical redox reaction involving proton (coming from water impurities or electrolyte side reactions) or divalent cation intercalation, oxygen loss seems to also be plausible, but in this case no reversibility could be expected.

4.4. AV_2O_5 (A = Mg, Ca): synthesis and electrochemical reactivity

Given the impossibility to achieve significant multivalent intercalation into V_2O_5 , we approached the subject from the opposite side, by synthesizing fully intercalated AV_2O_5 (A: Mg, Ca) phases and investigating the viability of MV ion extraction both electrochemically and chemically.

Pure, orthorhombic δ -MgV₂O₅ phase (S.G. Cmcm) was obtained either by i) direct synthesis from MgO and VO₂ or by ii) comproportionation reaction of MgV₂O₆ and MgV₂O₄, with cell parameters of a = 3.6920(4) Å, b = 9.9737(1) Å and c = 11.0208(10) Å and a = 3.6908(2) Å, b = 9.9747(7) Å and c = 11.0229(6) Å for i) and ii), respectively. Both syntheses, described in the Experimental section, led to the formation of micrometric δ -MgV₂O₅, however with slight differences in the particle morphology (See Fig. S8 in S.I.).

Orthorhombic α -CaV $_2$ O $_5$ phase (S.G. Pmmn) with cell parameters a = 11.3310(9) Å, b = 3.605(3) Å and c = 4.9000 (4) Å was obtained by comproportionation reaction of CaV $_2$ O $_6$ and CaV $_2$ O $_4$ (See the Experimental Section for details). The concomitant formation of V $_2$ O $_3$ (S.G. R–3c) and Ca $_3$ V $_2$ O $_8$ (R3c) as impurity phases in amounts approaching, respectively, 7% and 2% in weight, could not be avoided by repeated annealing treatments (See Fig. S8 in S.I.)

Attempts to chemically deintercalate Mg²⁺ from MgV₂O₅ by reaction with NO₂BF₄ at 80 °C in ACN did not result in any change in the crystal structure (See Fig. 8(A)) and no meaningful variation of the cell parameters was inferred from Rietveld refinement. When reacted with NO₂BF₄ oxidant in ACN at 80 °C, the brown CaV₂O₅ powder samples got rapidly dissolved and formed blue colored suspensions indicating the presence of V⁺⁴. The XRD pattern of the powder after decantation of the obtained suspension revealed the formation of an amorphous phase together with broadened and noisy reflections tentatively ascribed to V-O phases, with triclinic V₄O₇ appearing as one of the possible compounds present. However, the breath of the peaks and low signal-tonoise ratio in the patterns hampered any unambiguous peak indexing. The electrochemical oxidation tests revealed that the AV₂O₅ (A: Mg, Ca) phases are inactive at RT while specific capacities approaching the theoretical value (i.e. $\Delta x = 1 \text{ mol A}^{2+}$) could be achieved at 100 °C, with voltage profiles displaying a plateau between 3.8 and 4.2 V vs $\mathrm{Li}^+/$ Li and between 3.4 and 3.55 V vs Ca^{2+}/Ca for the MgV₂O₅ and the CaV₂O₅ electrodes, respectively (See Fig. S9 (A) and (B) in S.I.) The exsitu XRD patterns of the oxidized samples (Fig. 8 (A) and (B)) revealed neither meaningful variation of the cell parameters nor phase amorphization with respect to the pristine phases. The extra reflections observed in the ex-situ XRD diffractograms of the electrochemically oxidized MgV2O5 phase may be ascribed to LixBvOz phases, reasonably due to electrolyte salt (LiBOB) decomposition.

5. Discussion

The results from this study evidence that V_2O_5 is electrochemically inactive towards MV ion intercalation at RT in dry electrolyte solutions.

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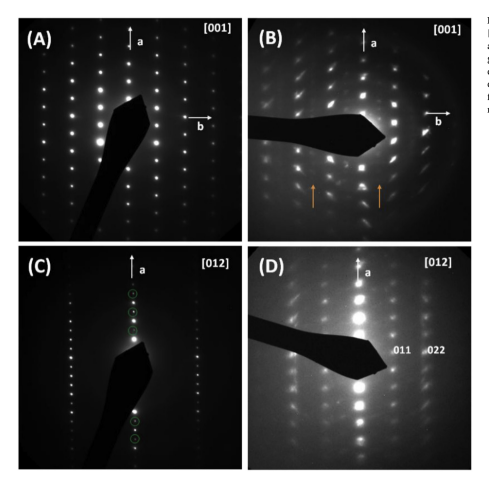


Fig. 7. Electron diffraction patterns corresponding to [100] and [012] zone axis for V_2O_5 pristine [(a), (c)] and after 100 h at the OCP at 100 °C [(b), (d)]. The green circles marked in (c) correspond to double diffraction and the orange arrows marked in (b) indicate possible twins. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

In contrast, relatively high values of specific capacity are obtained either by increasing the testing temperature to $100\,^{\circ}\text{C}$ or by increasing the water content in the electrolyte. Despite not being able to fully elucidate all the side reactions taking place and the phases involved, our findings unambiguously show that V_2O_5 is not stable in contact with the electrolytes (or even the electrolyte solvents alone) at $100\,^{\circ}\text{C}$ and that partial reduction of V^{+5} takes place. Hence, even if it is clear that applied current may modify the degradation mechanism, thermal instability has to be considered when assessing the results of electrochemical tests done above RT. The extent of the V_2O_5 reductive decomposition reaction is enhanced with time, but also depends on the electrolyte composition, being higher for magnesium than for calciumbased electrolytes, the lowest degradation being observed with

 $A(\text{ClO}_4)_2~(A=\text{Mg},\text{Ca})$ salts. Whether the charge compensation mechanism is related to oxygen loss (presumably irreversible), proton intercalation or both, is unclear at this stage and the nature of the phases formed is not unambiguously elucidated. Furthermore, the source of protons is also a matter of debate: since electrolytes with very low water contents were used in our study, we speculate that protons may alternatively come from complex decomposition reactions involving most likely electrolyte solvents, but further studies beyond the scope of the present paper should be carried out to assess that point. Yet, this is in agreement with previous work by Hong et al. [43] reporting the formation of similar phases upon electrochemical magnesiation of $V_2 O_5$ in both dry and wet electrolytes.

Despite a significant part of the electrochemical capacity observed

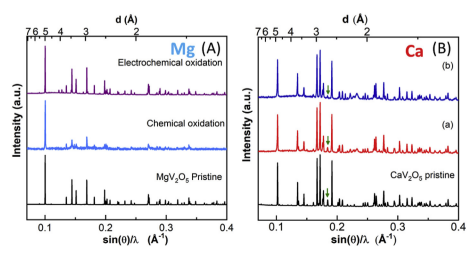


Fig. 8. Ex-situ XRD patterns of **(A)** MgV₂O₅ pristine electrode (black) and after chemical (light blue) and electrochemical (purple) oxidation; **(B)** CaV₂O₅ pristine electrode (black): **(a)** after 7 days OCP and **(b)** after electrochemical oxidation in Ca(BF₄)₂/ EC:PC at 100 °C. Green arrow indicates the main peak of V₂O₃ impurity. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

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in the experiments being related to side reactions involving the electrolyte (especially for Ca(BF₄)₂), changes in the diffraction pattern of V₂O₅ upon reduction at 100 °C are clearly observed, with variations being more evident in Mg than in Ca cells. In some of the electrolytes employed (such as the case of Mg(ClO₄)₂ in EC:PC), no evident signs of V₂O₅ thermal degradation could be detected by XRD (neither after OCP nor after soaking experiments), while sets of peaks due to the formation of additional phases were clearly observed along the reduction process. Besides the formation of phases analogous to those formed after long OCP periods (in amounts dependent on the employed electrolyte), the full reduction experiments led to the formation of a Häggite-type phase with a negligible divalent ion content, if any. In this respect, the formation of a fully inserted MgV₂O₅ or CaV₂O₅ phase was unambiguously dismissed by Rietveld refinement, this addressing a critical point in the controversial literature background related to the topic. The formation of a Häggite-type phase was observed both in Mg and Ca cells (on a minor extent in dry electrolytes than in wet ones). The reversibility of this reaction (despite some loss of crystallinity) does again raise the question of the competition between divalent and proton intercalation and points to the existence of alternative proton sources other than water impurities in the electrolyte.

Last but not least, no changes in the diffraction pattern were observed at RT unless water was added to the electrolyte. Despite the stability of V_2O_5 in the electrolyte at RT, its reduction in Mg cells with wet electrolytes led to the formation of a phase consistent with the one formed upon degradation at $100\,^{\circ}\text{C}$ (both at the OCP and upon reduction in Mg cells and in Ca cells only with $\text{Ca}(BF_4)_2$ based electrolyte). The diffraction pattern observed seems closely related to chemically prepared $\gamma\text{-H}_xV_2O_5$ phases, which do also bear strong similarities with most of the patterns reported in the literature upon presumable MV ion intercalation in dry and wet electrolytes at RT.

The results achieved in the course of this study indicate that, despite the similarity between the AV_2O_5 (A = Ca, Mg) and V_2O_5 crystal structures, the formation of a solid solution between α -V₂O₅ and α -CaV₂O₅ is not possible, neither electrochemically nor chemically. Moreover, attempts to intercalate magnesium in α -V₂O₅ do not lead to the formation of the closely related δ-MgV₂O₅. While for the case of the α polymorphs this can be related to higher migration barriers, in agreement with DFT results, the impossibility to extract Mg²⁺ from δ-MgV₂O₅ either chemically or electrochemically comes as a surprise, as moderate migration barriers were predicted in this polymorph [20]. Our most relevant and puzzling observations relate to the existence of several possible side reactions during the electrochemical reduction of α - V_2O_5 related to thermal instability or reactivity with the electrolyte components and water impurities. Unfortunately, none of the experiments performed enabled to assess a true significant amount of Ca²⁺ or Mg^{2+} intercalation in $\alpha\text{-}V_2O_5.$ While surprising at first sight, this is consistent with the controversial data available from previous literature, evidencing the impossibility to extract meaningful structural correlations from diffraction patterns with low signal-to-noise ratio or broad peaks.

6. Conclusions

The results obtained in this study contribute to raise the awareness about open questions related to MV ion intercalation into $\alpha\text{-}V_2O_5,$ generally regarded as a prototypical cathode host for MV batteries even without a clear understanding of the overall intercalation process, especially from a crystal chemistry perspective.

Our findings shed some light into the existing controversy and point at the unsuitability of V_2O_5 as cathode for Mg and Ca batteries. Indeed, no evidence of significant MV ion intercalation can be inferred from the refinements of the ex-situ XRD patterns of the electrodes collected after electrochemical reduction experiments carried out under various conditions. The formation of protonated Häggite-type phases $H_xV_4O_6$ (C2/m symmetry) rather than magnesiated or calciated V_2O_5 is observed in

both dry and wet electrolytes, with protons most likely coming from complex electrolyte decomposition requiring further studies to be fully understood.

Furthermore, a thorough analysis of the XRD pattern of V_2O_5 stored in the employed electrolytes (or even in the bare solvents) or in Mg and Ca cells kept at the OCP revealed the progressive appearance of a set of peaks closely related to those of chemically prepared γ -H_xV₂O₅. These patterns are also similar to most of those reported in the literature after presumable MV ion intercalation in dry and wet electrolytes at RT, its origin being not even tentatively elucidated in most of the cases. Our findings are consistent with this orthorhombic V₂O₅-like phase with expanded a parameter resulting from a reductive decomposition of V₂O₅ that requires further investigation.

The surprising results achieved in the course of this study underpin the importance of a rigorous methodology for a thorough and reliable investigation of the reduction process. Thermal and chemical stabilities need to be assessed, especially for tests done above RT, and characterization techniques complementary to electrochemistry are compulsory to demonstrate MV ion intercalation.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.jpowsour.2018.08.024.

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