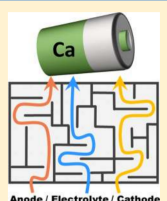
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Achievements, Challenges, and Prospects of Calcium Batteries

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ABSTRACT: This Review flows from past attempts to develop a (rechargeable) battery technology based on Ca via crucial breakthroughs to arrive at a comprehensive discussion of the current challenges at hand. The realization of a rechargeable Ca battery technology primarily requires identification and development of suitable electrodes and electrolytes, which is why we here cover the progress starting from the fundamental electrode/electrolyte requirements, concepts, materials, and compositions employed and finally a critical analysis of the state-of-theart, allowing us to conclude with the particular roadblocks still existing. As for crucial breakthroughs, reversible plating and stripping of calcium at the metal-anode interface was achieved only recently and for very specific electrolyte formulations. Therefore, while much of the current research aims at finding suitable cathodes to achieve proof-of-concept for a full Ca battery, the spectrum of electrolytes researched is also expanded. Compatibility of cell components is essential, and to ensure this, proper characterization is needed, which requires design of a multitude of reliable experimental setups and sometimes methodology development beyond that of other next generation battery technologies. Finally, we conclude with



Anode / Electrolyte / Cathode

recommendations for future strategies to make best use of the current advances in materials science combined with computational design, electrochemistry, and battery engineering, all to propel the Ca battery technology to reality and ultimately reach its full potential for energy storage.

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1. INTRODUCTION

Efficient and readily available energy storage is essential to help solving some of the grand challenges our modern society is facing today: air pollution, oil-dependency, and climate change. The magnitude of the challenge is huge and involves many interrelated aspects including energy independence, environmental sustainability, economics-and not the least technological development. Electrochemical energy storage is considered to be a key technology, and the prevailing lithium-ion battery (LIB) technology has made possible our society of portable electronic devices. It also offers a near-term solution for more sustainable transport with less environmental impact and stationary energy storage for renewable energies, such as solar and wind power. While the cost of LIBs has been reduced by 8% at the pack level annually during the past decade,¹ this technology is now reaching its fundamental limits in terms of

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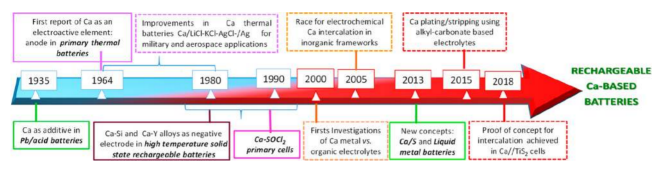


Figure 1. Timeline of Ca use in battery technologies.

energy density. In addition, while the quest for even higher energy densities has for long been the main driving force behind progress in battery technology, LIBs contain both Ni and Co, and the risks of limited supply and/or significantly increased prices cannot be ignored and prompt the development of more long-term sustainable battery chemistries.^{2,3} Furthermore, a transition from metal ion- to metal-anode-based battery chemistries is appealing, as it can substantially enhance the figures of merit.⁴⁻⁸ If also more abundant and nontoxic metals are used, the overall battery technology would also benefit from reduced cost and lower environmental impact. Moreover, the concept becomes particularly attractive if multivalent ions are used as charge carriers, as this in principle enables the capacities to be doubled as compared to using monovalent-ion charge carriers such as Li⁺ or Na⁺. Alternatively, one can consider that reaction of only a half of the ions would be needed to achieve the same capacity.^{8,9} Combining all of the above: Calcium is the fifth most abundant element in the Earth's crust, it is nontoxic, and its standard reduction potential is -2.87 vs NHE, which combined with its density of 1.54 g/cm³ and charge capacity of 1.34 Ah/g, result in a theoretical energy density of 2.06 Ah/cm³. In comparison, the graphite of today's LIBs is at a mere 0.97 Ah/ cm³—hence a 2-fold increase in this measure is achievable. In addition, the Ca²⁺ ion should be more mobile in liquid electrolytes because of its less polarizing character (charge/ radius ratio) than both Mg²⁺ and Al³⁺ ions, which are employed in battery concepts much more heavily researched. To enable the reader to more easily understand the (limited) development of Ca batteries until now, we here first provide a historical perspective before we turn to the (rapid) progress during the last 4 years.

This Review presents a comprehensive examination of the requirements and current state-of-the-art of electrolytes and electrodes for rechargeable Ca batteries. This section contains a minor historical recollection, section 2 covers the exploration of active materials and electrolytes, while section 3 deals with the (un)reliability of experimental setups used and the methodology development needed, before we assess full-cell layouts and finally present some concluding remarks.

1.1. Historical Perspective (-2015): A Myriad of Concepts

The first application of calcium at all in batteries was in 1935 but then as an additive for alloy strengthening the lead grids for Pb– acid battery cells.^{10–12} The first report of calcium as the electroactive element appeared much later, in 1964, and is related to primary thermal batteries, a technology mostly used in military and aerospace applications (Figure 1).¹³ These Ca batteries contained electrolytes with high melting points (such as mixtures of metal chlorides), kept at ambient temperature during storage periods (which may be extremely long) to avoid self-discharge and thermally activated to deliver power. The Ca/ KCl-LiCl-AgCl-K₂CrO₄/Ag cells were reported to discharge at 450 °C for 11 min, despite issues related to the formation of a Li–Ca alloy at the anode. This type of cell was used in military applications since World War II.¹⁴ Some attempts to modify the surface of the calcium metal anodes with acetic acid to yield CaCO₃ were reported,¹⁵ despite the chemistry being shown to be extremely complex involving also corrosion with formation of both Ca_2CrO_4Cl and $KCaCl_3$.¹⁶ Efforts to replace the chlorides by nitrates, with lower melting points, as electrolytes were also reported.¹⁷ A maximum exchange of one electron per mole of calcium was observed with the formation of a passivation film most likely consisting of CaO, hindering any fast kinetics. The addition of halides to the electrolyte was suggested as a way to break the passivation film and improve the kinetics. Other hightemperature concepts investigated in these early days involved the use of a Ca–Si alloy anode coupled to a Ca²⁺ conducting β'' alumina solid electrolyte for rechargeable cells operating at 580 $^{\circ}C_{1}^{18}$ a Ca-Y alloy anode coupled to a fluoride conducting solid electrolyte,¹⁹ and even a Ca-O₂ cell-also using a Ca-Si alloy anode complement with a zirconia-based electrolyte.²⁰ Despite a few charge-discharge cycles being reported for the above systems, the redox mechanism was not unambiguously ascertained, and no further studies were published.

Staniewicz was the first to report on the electrochemistry of Ca-SOCl₂ as an alternative to the Li-SOCl₂ primary cells, again a battery technology mainly used for military applications, and presented the impossibility of calcium plating upon cell reversal as a safety advantage.²¹ Further studies by Peled et al. attributed this feature to the formation of a passivation layer consisting mainly of $CaCl_2$ impermeable to the Ca^{2+} ions.²²⁻²⁵ A similar technology with a somewhat higher operation cell potential was also developed,^{26,27} but corrosion of the Ca metal electrodes was found to be an issue; the Ca-SOCl₂ cells lost ca. 10% of their capacity after only 2 weeks of storage.²⁸ The strategy to avoid corrosion and self-discharge was to use different additives such as Ca(AlCl₄)₂ formed by reacting Ca with $AlCl_{3}^{21}$ Sr(AlCl₄)₂, or Ba(AlCl₄)₂, changing the composition of the passivation layer.²⁹ Furthermore, corrosion of the stainless steel can material was also observed and attributed to reactivity with calcium metal, in analogy with lithium metal systems.³⁰ As the calcium-based concepts above did not provide enough advantages to replace the Li-SOCl₂ technology, they never reached the market,³¹ but this may also be related to the fact that these batteries are only used in niche applications (military or very low-temperature environments).

The issues related to electrochemical calcium deposition discovered in the studies above effectively prevented any further investigations of secondary, rechargeable, battery technologies based on calcium metal as the negative electrode. Yet, basic

studies related to the behavior of calcium metal anodes using organic electrolytes similar to those used at the time of emerging LIB technology were made by Aurbach et al.³² Their studies included solvents such as acetonitrile (ACN), tetrahydrofuran (THF), γ -butyrolactone (gBL), and propylene carbonate (PC) as well as $Ca(ClO_4)_2$ and $Ca(BF_4)_2$ salts together with noncalcium containing salts. The conclusion of this pioneering work was that the passivation layer formed on calcium metal does not enable transport of calcium ions, in full agreement with the results for the SOCl₂ battery technology, with the major consequence being the impossibility to develop any secondary Ca batteries using these electrolytes, as even if calcium stripping would be feasible, plating upon charge would not. By the year 2000, when the LIB technology was already well-established in the market,³³ the idea to use metal anodes coupled to multivalent-ion charge carriers re-emerged as a way to further boost the energy density. Indeed, proof-of-concept was achieved for Mg battery cells with metal anodes,³⁴ using magnesium organohaloaluminate salts in THF or glymes as electrolytes, which, despite a somewhat narrow electrochemical stability window of ca. 3 V, enabled reversible Mg plating and stripping. Yet, no similar electrolytes were available for Ca⁸ and hence, the study of Ca²⁺ intercalation and the quest for cathodes was not straightforward, the only possibility being the assembly of cells using alternative anodes, such as activated carbon.³

The basics of calcium-ion intercalation in transition-metal compounds were studied in the 1960s-1970s as part of early intercalation chemistry research.³⁶ This involved crystal structures exhibiting a van der Waals gap, such as metal sulfides WS_2 , TaS_2 , TiS_2 , and some metal oxides, e.g., MoO_3 , and V_2O_5 , in which intercalation of a wide range of neutral or charged species was possible, the latter concomitant to reduction of the transition metal of the host. Despite most studies being devoted to intercalation of lithium and other alkali ions, attempts to chemically intercalate other ions such as Ca²⁺ for comparative purposes were common, either from aqueous solutions of the corresponding hydroxides or from the metal dissolved in liquid ammonia.^{37–39} Due to the advent of the LIB technology in the 1970s, which concentrated most research efforts to Li⁺, research on multivalent systems fell into oblivion, but the field started to re-emerge in the early 2000s, with a few reports dealing with the feasibility of calcium intercalation in inorganic compounds. These included electrochemical studies of hexacyanoferrates using an aqueous electrolyte^{40,41} and combining electrochemistry and other characterization techniques (mostly powder X-ray diffraction, or XRD hereafter) with V₂O₅ as the host using $Ca(ClO_4)_2$ in either ACN or PC as electrolytes.^{42–44}

In the following decade and in line with the growing awareness about the imperative evolution toward electrification of transportation and energy storage needs for renewable energy, the efforts in the area were reinforced. This resulted in investigations of new concepts such as Ca/S cells,⁴⁵ which showed some electrochemical response but lacked reversibility, and more exotic alternatives such as liquid metal batteries.⁴⁶ The latter concept consists of two liquid metal electrodes separated by a molten salt electrolyte that self-segregate into three liquid layers based on immiscibility and density differences. While the solubility in molten salts and strong reducing power of Ca preclude its use as metal electrode, calcium containing alloys melting at moderate temperatures (around 600 °C) seem to exhibit promising behavior using molten halide electrolytes⁴⁷ given their thermodynamic redox potential, 50,51 but much engineering is needed for this technology-development of corrosion resistant cell components, effective seals, and thermal management, and so on—before any proper practical estimates can be made.

Reinvestigations of hexacyanoferrates were also made using both aqueous⁴¹ and organic⁵² electrolytes resulting in a very reversible electrochemical response and with enhanced capacity obtained by water addition to the organic electrolytes.⁵³ Despite some further studies,^{54,55} indicating minor, if any, modifications of the host structural framework, the redox mechanism and the possible role of water molecules, either present in the as prepared hexacyanoferrates or in the electrolyte, have not yet been fully elucidated. Interest in intercalation cathodes, such as those used in the LIB technology and using the vast know-how created in that field, also re-emerged, but the bottlenecks related to the differences between Li⁺ and Ca²⁺ as charge carriers were soon clearly realized and hence the need for specific materials design.^{56,57}

1.2. The Current Era (2016–): Realizing Metal-Anode- and Organic-Electrolyte-Based Cells

By the end of 2015, the feasibility of reversible calcium metal plating and stripping was assessed at moderate temperatures using conventional alkyl-carbonate organic-solvent-based electrolytes.⁵⁸ Despite issues associated with these electrolytes, this opened for a more extensive electrode materials screening. With Ca metal anodes being the most attractive choice, research was focused on electrolyte formulations enabling better Coulombic efficiency and, if possible, making operation at room temperature feasible, and at the same time, these were used for unraveling suitable cathode materials. Yet, alternative anodes have also been researched; the first calcium graphite intercalation compound (CaC₆) was achieved chemically at high temperature in 2005⁵⁹ and attempted electrochemically somewhat later.⁶⁰ An interest in Ca alloys has been present since 2011,⁶¹ but still, most efforts are directed toward cells using calcium metal anodes and intercalation cathodes.⁶

Most Ca battery cells are analogous with LIBs on the cathode side; during the discharge, the charge carrier ion (Ca^{2+}) migrates from the anode to the cathode through the electrolyte, while the electrons flow across an external circuit (Figure 2). These

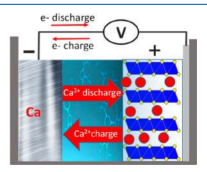


Figure 2. Schematic of a Ca battery using a Ca metal anode and an intercalation cathode.

processes are reversed upon charging. Except for a few more exotic concepts, with $air^{63,64}$ or sulfur cathodes,⁴⁵ the redox mechanism at the cathode involves insertion/deinsertion of Ca^{2+}

$$x \operatorname{Ca}^{2+} + 2x e^{-} + \operatorname{Ca}_{y} \operatorname{Host} \Leftrightarrow \operatorname{Ca}_{x+y} \operatorname{Host}$$
(1)

where Host represents the cathode active material wherein the Ca^{2+} ions are intercalated. For Ca batteries, the main challenge is

Table 1. Ele	ctrochemical Te:	sts of Ca An	Table 1. Electrochemical Tests of Ca Anodes in Two- or Three-Electrode Cells	ctrode Cells					
negative/work- ing electrode	counter electrode	reference electrode	electrolyte	cutoff voltage (cyclic voltam- metry)	passivation	coulombic efficiency	number of cycles	water content (ppm)	comments
Ca^{21}	SOCl ₂ /C	Ca	Ca(AlCl ₄) ₂ in SOCl ₂		$CaCl_2$	<5%	0		Ca stripping at ca. 100 mV vs Ca pseudo-RE.
Ca or Au ³²	Ca	Ca	0.05 to 1 M Ca(ClO ₄) ₂ or Ca(BF ₄) ₂ in THF, PC, γ BL, or ACN	–1.8 V vs Ca pseudo-RE	CaCl ₂ in ClO ₄ -based electrolyte, Ca(OH) ₂ in THF	%0	1	<30	No plating observed. Shift of the Ca metal RE: 750 mV.
Ca ⁷⁴	Ca	Ca	0.45 M Ca(BF ₄) ₂ in EC:PC (100 $^{\circ}$ C)	–1.4 V vs Ca metal	borate, carbonate, CaF_2	85%	30	<40	Plating observed above 75 $^\circ\mathrm{C}.$
Au^{73}	Ca	Ca	$1.5 \text{ M Ca}(BH_4)_2 \text{ in THF}$		CaH_2	95%	50	4>	
Pt^{79}	Ca	Ca	0.25 M CaBhfip in DME	-0.35 V vs Ca metal	CaF_2	80%	22		$5~{\rm CV}$ cycles at 80 ${\rm mV}\cdot{\rm s}^{-1}$ conditioning procedure.
Au ⁷⁸	Ca	Ca	0.5 M CaBhfip + 0.1 M Bu ₄ NCl in DME	–1.0 V vs Ca metal	CaF_2	%06	50		Bu4NCI appeared mandatory for long-term cycling.
CaSi ₂ ⁵⁸	Ca	Ca	0.45 M Ca(BF ₄) ₂ in EC:PC (100 $^{\circ}$ C)	0.8 V vs Ca metal			1	<30	Dealloying evidenced by XRD, alloy formation at very high overpotential.
Sn ^{81,82} fully re- duced: Ca ₇ Sn ₆ (?)	graphite and acti- vated carbon	,	0.8 M Ca(PF ₆) ₂ in EC:PC:DMC:EMC (2:2:3:3 in vol)	5 V for full cells	carbonate and CaF ₂	ca. 80%	350 and 1000		Stability issues of $Ca(PF_6)_2$ and cell balancing? Long cycling only at high C-rate (low Coulombic efficiency at low C- rate).
GeO ₂ 98	graphite rod	pyres-glass sealed Ag/AgCl	CaCl ₃ —NaCl (1:1) melt at 600 °C	-2.5 V vs Ag/ AgCl			1		Absence of Na alloy was ascertained by means of EDX and XRD.
MCMB ⁸⁹	expanded graphite	1	0.8 M Ca(PF ₆) ₂ in EC:DMC:EMC (5:3:2 vol)	5.2 V for full cells		ca. 80%	300		Stability issues of $Ca(PF_6)_2$ and cell balancing? Long cycling only at high C-rate (low Coulombic efficiency at low C- rate).
PNDIE ⁹⁷	copper hexacyano- Ag/AgCl ferrate	Ag/AgCl	2.5 M $Ca(NO_3)_2$ (aq) (pH = 5.1)	-0.9 V vs Ag/ AgCl		99% (at C/ 2)	2000	ı	Role of proton studied by cycling at pH 3, 5, 7, and 9.
PTCDA ⁹⁶	activated carbon	,	Ca(NO ₃) ₂ (aq) sat.				10	ı	Amorphization upon cycling. HER?

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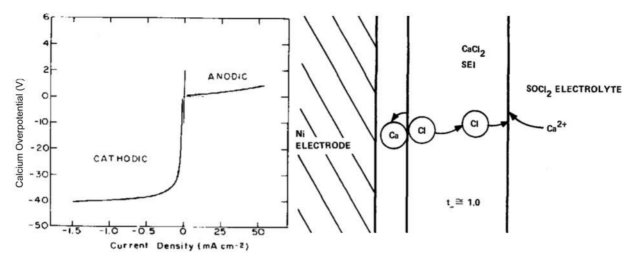


Figure 3. Left: I-V characteristics of a Ca²⁺/Ca solid electrolyte interphase (SEI) covered electrode in a SOCl₂-based electrolyte (showing "diodelike" behavior). Reproduced from ref 25 with permission. Copyright [1988] Elsevier. Right: Model for calcium deposition through an SEI of CaCl₂. The ionic current is carried by the anion as a result of the low, close to 0, transport number of Ca²⁺. Reproduced with permission from ref 21. Copyright [1980] The Electrochemical Society.

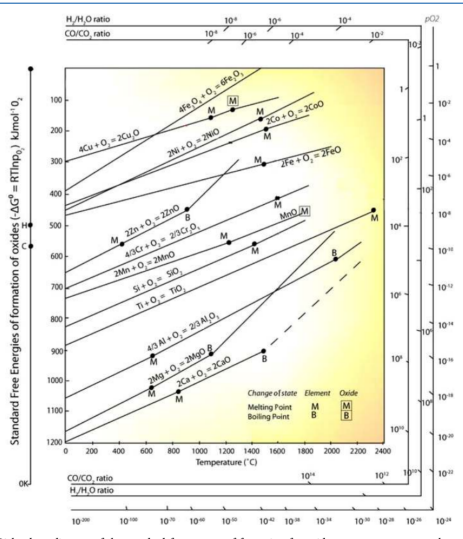


Figure 4. Ellingham–Richardson diagram of the standard free energy of formation for oxides vs temperature, together with the corresponding equilibrium O_2 partial pressure. Reproduced with permission from ref 72. Copyright [2018], University of Cambridge.

the larger size and charge of the Ca²⁺ ion as compared to the Li⁺ ion, and to less extent to the Na⁺ ion, for which many suitable

hosts exist. To date, the cathodes proposed have been very few and are treated in detail in section 2.3 below.

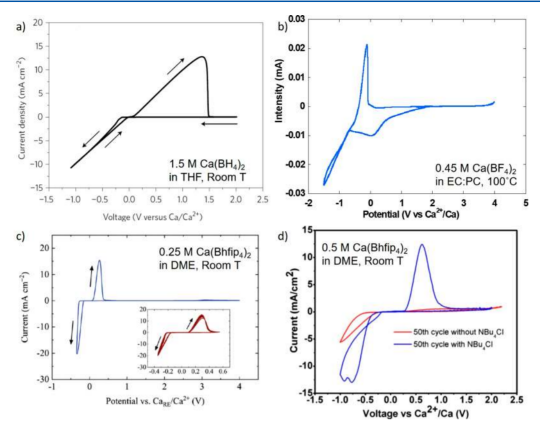


Figure 5. Cyclic voltammograms of calcium plating/stripping in (a) 1.5 M Ca $(BH_4)_2$ in THF (room temperature, 25 mV/s, working, reference, and counter electrodes Au, Ca, and Pt, respectively), reproduced from ref 73 with permission, copyright [2018] Nature Publishing Group, in (b) 0.45 M Ca $(BF_4)_2$ in EC:PC (100 °C, 0.5 mV/s, stainless steel working and Ca counter and reference electrodes), reproduced from ref 74 with permission, copyright [2016] Nature Publishing Group, in (c) 0.25 M Ca $[B(hfp)_4]_2$ in DME (room temperature, 80 mV/s, Pt working and Ca counter and reference electrodes), reproduced from ref 79 with permission, copyright [2019] The Royal Society of Chemistry, and in (d) 0.5 M Ca $[B(hfp)_4]_2$ in DME with or without 0.1 M NBu₄Cl (room temperature, 25 mV/s, Au working and Ca counter and reference electrodes), reproduced from ref 78 with permission, copyright [2019] American Chemical Society.

2. ANODES, ELECTROLYTES, AND CATHODES

The choice of active materials together with the electrolyte sets the theoretical limits in voltage and energy density but also affects the kinetics (power), life-length, safety, and so on. A special role is played by the *interfaces* and in many cases also the *interphases* formed between the electrolyte and the electrodes. While also current collectors and casing materials and so on affect the final product, these are covered in section 3 dealing with full-cell design, rather than in separate subsections here.

2.1. Anodes

Several Ca battery anodes have been electrochemically tested, either in three- or two-electrode cell configurations, this is to say, with or without a reference electrode distinct to the counter electrode (Table 1). While the description of cell configurations is exposed in section 3, this section focuses in the anode side, and for simplicity, the contents are organized into two subsections; section 2.1.1 covers Ca metal anodes with the related interfaces, and section 2.1.2 covers alloys, intercalation materials, and other alternatives.

2.1.1. Ca Metal Anodes and Interfaces. The use of Ca metal anodes requires reversible stripping (at discharge) and plating (at charge). Several unsuccessful attempts to electro-deposit divalent cations have demonstrated the large difficulties associated with trying to accomplish this.⁶⁵ Based on the studies of Auborn et al. and Behl et al. on electrolytes of inorganic Li salts, e.g., LiBCl₄, LiAlCl₄, and so on, in phosphorus oxychloride

 $(POCl_3)$ and thionyl chloride $(SOCl_2)$ vs Li metal anodes,^{66,67} Peled et al.⁶⁸ and Staniewicz²¹ reported on the electrochemical behavior of the M–SOCl₂ (M = Mg or Ca) electrolytes vs the corresponding metal anodes, with a cell reaction suggested to be analogous with the Li systems

$$2M + 2SOCl_2 \Leftrightarrow 2MCl_2 + SO_2 + S \tag{2}$$

In terms of specific energy density, the Ca-SOCl₂ cell reaction is ca. 300 Wh/kg (for sake of comparison, the Li system is ca. 700 Wh/kg). Although this technology was considered for the development of primary cells, efforts to electroplate Ca from SOCl₂-based electrolytes were made but failed. Indeed, while a high current density was recorded upon anodic polarization of a Ca metal electrode with relatively low overpotential (efficient stripping), a very high overpotential was recorded upon cathodic polarization (Figure 3), and solvent reduction led to SO_2 gas being formed. The Coulombic efficiency for the Ca plating reaction was estimated to be a mere 5%.²⁵ The low reversibility of the Ca plating/stripping reactions has in general mainly been attributed to the poor diffusion of divalent cations through the passivation layer formed (mostly CaCl₂) as well as its mixed ionic conductor nature, which precludes Ca plating (Figure 3). In contrast, Mg electrodeposition is possible, and the obtained deposit less prone to corrosion; a Ca deposit is completely consumed in less than 7 min after plating, whereas Mg electroplated on Ni appears to be stable for at least 2 h.^{23,68}

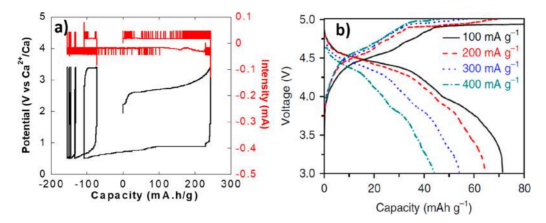


Figure 6. (a) Capacity vs potential profile from Potentiostatic Intermittent Titration Technique (PITT) of a CaSi₂ electrode at 100 °C. Reproduced from ref 58 with permission from Elsevier, copyright [2016]. (b) Charge–discharge performance of a Sn-expanded graphite Ca-ion cell at different current densities ranging from 100 to 400 mA g^{-1} . Reproduced from ref 82 with permission from the Nature Publishing Group, copyright [2018].

While the Ca metal anode is by far the most promising cell design, the electrochemistry of Ca metal is rather complex, including the role of the very stable calcium oxide. An Ellingham diagram⁶⁹ plots the standard free energy of formation of oxides with temperature, and it usually also provides the partial pressure of oxygen at equilibrium (Ellingham-Richarson diagram, as shown in Figure 4). The standard free energy is more negative for the more stable oxide; hence, CaO is clearly more stable than MgO and even Al₂O₃. The high stability of CaO is certainly a severe problem to overcome for those aiming at employing $Ca-O_2$ as the cathode. Since the entropy variation for the oxide formation is negative, the free energy becomes less negative as temperature increases. Therefore, it is not surprising that the first reports on Ca metal anodes in the early 1960s (section 1) were on cells operated at high temperatures (>450 °C).^{13-15,17} The electrolyte chemistry was very similar to Ca metal production; molten salts, typically eutectic mixtures of CaCl₂ and CaF₂, were employed and involved electrowinning and electrorefining processes performed at >850 °C.⁷⁰ Later on, the concept of liquid metal electrodes also used higher temperatures, typically operating at >500 °C.⁷¹

Ca batteries are very much dependent on the properties and performance of the electrolyte/electrode interfaces. In an early study by Aurbach et al.,³² the process of calcium plating at the anode from an organic electrolyte was explored using $Ca(BF_4)_2$ and $Ca(ClO_4)_2$ as the electrolyte salts and PC, gBL, THF, and ACN as the solvents. The plating was found by infrared spectroscopy to be limited by the electrolyte decomposing to form a passivation layer on top of the Ca metal, hence akin to the solid electrolyte interphase (SEI) formed on graphite in LIBs. In contrast to LIBs and why the term SEI should not be used here. among all the different electrolytes studied, none of them allowed for reversible plating/stripping of calcium, as the passivation films, mainly composed of CaCO₃ and Ca(OH)₂, did not conduct Ca²⁺ ions. Although these observations severely stalled the development of Ca metal-anode-based batteries, two recent studies have indeed reported on successful calcium plating and stripping using organic electrolytes in the presence of passivation layers: 1.5 M Ca(BH₄)₂ in THF⁷³ and 0.45 M $Ca(BF_4)_2$ in EC:PC.⁷⁴ The former electrolyte enables reversible cell performance at room temperature, with an electrochemical stability window of about 3 V, while the latter offers over 4 V but demands an elevated temperature (100 °C) (Figure 5). For $Ca(BH_4)_2$ in THF, the passivation layer was found to be entirely

composed of CaH₂ formed by the reaction of Ca metal with the solvent. Unfortunately, the CaH₂ passivation layer was found to continuously grow upon cycling and is highly reactive. Recently, Ta et al. have performed a systematic fundamental study on the Ca plating mechanism using this electrolyte and a Au or Pt rotating disc electrode setup, suggesting a chemical–electrochemical deposition mechanism, with the kinetics of the chemical step (involving hydrogen adsorption) being substrate-dependent and faster with Pt than with Au.⁷⁵

For the $Ca(BF_4)_2$ in EC:PC electrolyte-based cells, the situation is more complex; several solvent decomposition products were identified including CaF₂ and carboxylates but not which one(s) that enabled Ca migration.⁷⁴ Young et al. investigated possible reduction pathways for EC and $Ca(ClO_4)_2$ in EC (thus not the same anion as in the experiment) by means of DFT and AIMD simulations and concluded that CaCO₃, CaO, and $Ca(OH)_2$ should be the major inorganic components of the SEI.⁷⁶ Later on, a detailed study on the passivation layers formed from $Ca(BF_4)_2$ or $Ca(TFSI)_2$ in EC:PC electrolytes allowed calcium carbonates, fluorides, and borates to be identified—with the latter only being observed for BF₄⁻-based electrolytes, the electrolyte in which Ca plating and stripping is feasible, i.e., no plating is observed for the electrolyte containing TFSI.⁷⁷ Further experiments used an electrode prepassivation step by a BF₄⁻-based electrolyte. The prepassivation procedure consisted in polarizing a stainless steel current collector at low potential in order to produce a borate containing passivation layer. Subsequently, after a thorough rinsing procedure, the electrode was transferred to a cell with a TFSI containing electrolyte.⁷⁷ Thereby could not only Ca plating be observed, using the previously inactive Ca(TFSI)₂-based electrolyte, but also an enhanced electrochemical response-ca. 4 times higher current densities. This strongly suggests that the nature of the passivation layer is key to enable Ca plating and that the composition of the electrolyte plays a major role in the overall plating kinetics. Both the cation mobility in the electrolyte and interfacial phenomena such as desolvation are thus interconnected properties of utmost importance for practical Ca batteries.

In 2019, two independent studies demonstrated Ca plating and stripping at room temperature using Ca salts based on fluorinated alkoxyborate anions and dimethoxy ethane (DME) as solvent. Coulombic efficiencies between 80 and 90% were reported together with anodic stabilities above 4 V vs Ca^{2+}/Ca and the presence of CaF₂ as an electrolyte decomposition product.^{78,79} These studies represent a significant step forward in the development of a room temperature Ca metal-anode-based battery and should be benchmarked against cathode materials. Nonetheless, the need for a conditioning protocol (five cyclic voltammetry cycles at 80 mV·s⁻¹)⁷⁸ or addition of 0.1 M Bu₄NCl⁷⁹ in order to achieve the high Ca plating/stripping Coulombic efficiency or to improve cyclability also calls for additional work on surface passivation and in order to identify the electroactive species. The possibility/risk of Ca dendrite growth is also mentioned, as characteristic structures were observed growing through the separator, and a short circuit was recorded at a current density of 1 mA.cm^{-2.78}

2.1.2. Alloys, Intercalation Materials, and Other Alternatives. Alloys are considered a viable alternative to Ca metal anodes, as they often have both high specific capacity and low potential.^{61,80} As expected from the divalent nature of Ca and Mg cations, a similar volume expansion can be calculated for the formation of A_M (with A = Ca or Mg and M = Sn, Si, etc.) as for the formation of Li_{2x}M, and a similar amount of charge is stored.⁶¹ Si- and Sn-based alloys investigated electrochemically showed decalciation of CaSi2 to be possible using 0.45 M $Ca(BF_4)_2$ in EC:PC electrolyte at 100 °C, leading to an average particle diameter size decrease from 7.3 to 2.6 μ m.⁵⁸ A large voltage hysteresis was, however, observed upon reduction, and amorphization prevented any confirmation of alloy reformation. Tang's group investigated a Sn anode using a $0.8 \text{ M Ca}(\text{PF}_6)_2$ in EC:PC:DMC:EMC (2:2:3:3 by vol.) electrolyte and a graphite cathode (employing anion insertion).^{81,82} Since only twoelectrode cells were assembled, the capacity vs voltage profile of Sn is not really unambiguously determined, but yet, a much smaller voltage hysteresis was obtained as compared with the Si anode (Figure 6). Even though 350 cycles were reported, the low Coulombic efficiency (ca. 80%) suggests a significant amount of parasitic reactions, most probably involving electrolyte decomposition. In addition, the $Ca(PF_6)_2$ salt employed is known to be quite unstable.⁸³

A systematic screening and high-throughput density functional theory (DFT) investigation of a large number, in total 115, of Ca_xM alloys was very recently made by Wolverton's group.⁸⁰ The most promising alloy candidates were selected considering calciation voltage, volume expansion, and specific capacity. These criteria resulted in a wide range of metals being considered worthy of further investigation; M = Sn, Si, Sb, Ge, Al, Pb, Cu, Cd, CdCu₂, Ga, Bi, In, Tl, Hg, Ag, Au, Pt, and Pd. Pt and Au attract special interest, as these metals sometimes are used as substrates/working electrodes (Table 1), emphasizing that care must be taken to ensure that alloy formation does not bias the interpretation. Ca alloys were also considered in thermal batteries as a way to overcome the high melting point of Ca (842 °C). The operation temperature of such cells could be lower by using Ca alloys such as Ca-Bi,⁴⁸ Ca-Sb,⁸⁴ Ca-Ge,⁸⁵ and Ca-Mg.

Not only simple metal alloys are advocated for; twodimensional GeP₃ was recently investigated by DFT as a potential anode for several post-Li batteries,⁸⁶ Ca²⁺ was found to preferentially react with GeP₃ through a conversion reaction resulting in a theoretical specific capacity of 658 mAh/g.

Graphite is by far the most representative and studied insertion anode material for the LIB technology. While fellow alkali earth metals such as barium (Ba) and strontium (Sr) have been found to intercalate in graphite rather easily,⁸⁷ Ca intercalation using a similar procedure (contact of graphite

with the metal vapor) only leads to superficial intercalation.⁸⁸ So far, only dimethyl sulfoxide (DMSO)-solvated Ca has been intercalated readily into graphite.⁵⁹ Successful preparation of CaC₆ was, however, claimed by Emery et al. after immersing graphite in a molten lithium–calcium alloy at 350 °C for 10 days.⁵⁹ The authors suggested that only Ca is intercalated and mentioned that the CaC₆ crystal structure is unique (rhombohedral and $R\overline{3}m$ space group) as compared to other MC₆ compounds having hexagonal symmetry. Wu et al. assembled a dual-carbon electrode cell using a *meso*-carbon microbead (MCMB) anode and an expanded graphite cathode.⁸⁹ Upon cycling, reversible shifts of the (002) graphite diffraction peak were observed as well as a broadening, showing loss in graphitization degree (Figure 7). Unfortunately, since all

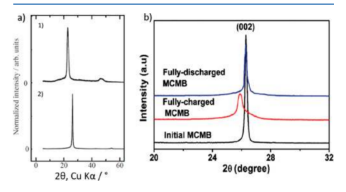


Figure 7. (a) XRD patterns of (1) Ca-intercalated BC₈ and (2) original BC₈ films. Reproduced from ref 90 with permission from The Chemical Society of Japan. Copyright [2018]. (b) XRD patterns of an initial, fully charged, and fully discharged MCMB anode in a Ca dual-carbon electrode battery (expanded graphite used as cathode) during the initial cycle. Reproduced from ref 89 with permission from Wiley-VCH. Copyright [2018].

tests were performed using two-electrode cells, no conclusions can be drawn with respect to the intrinsic performance and operating voltage of the MCMB anode. Ishikawa et al. proposed an interesting strategy for Ca insertion by first preparing BC₈,⁹⁰ which allowed for easier Ca intercalation using the standard method, something rationalized by the relationship between the ionization potential of Ca and the electron affinity of BC8. Borophene, the boron analogue to graphene, and hydrogenated defective graphene, have both been investigated by DFT for possible application as Ca battery anode materials.^{91,92} Higher binding energies were obtained for Ca than for Mg, Li, and Na, and specific capacities of 800 and 591 mAh/g were obtained, individually. Datta et al. investigated the adsorption of Ca (and Na) on graphene with divacancies and Stone–Wales defects⁹³ and concluded that adsorption is not possible on pristine graphene but enhanced by increasing the density of defects, with a calculated capacity of 2900 mAh/g for Ca^{2+} adsorption. As for every other high-surface-area anode material, however, it is very unlikely that graphene can be used in practice, as the sidereactions taking place would be highly detrimental and result in an excessive cation inventory loss. In addition, the low-density graphene-based anodes will result in low volumetric energy density cells. MXenes have also been considered as anodes for Ca, K, Na, and Li batteries.^{94,95} By DFT calculations, the adsorption energies were found to decrease as the coverage increased, and a larger effective ionic radius was found to increase the interaction between the alkali and alkali earth atoms, penalizing the capacity; 320 mAh/g for Ca^{2+} . However,

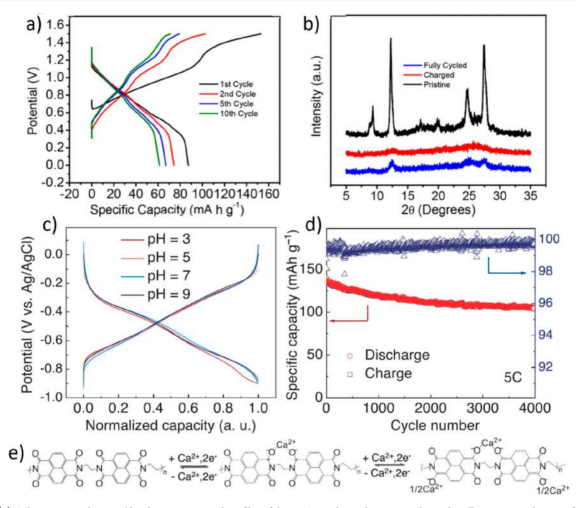


Figure 8. (a) Galvanostatic calciation/decalciation potential profiles of the PTCDA electrode in a two-electrode cell at a current density of 20 mA g^{-1} , and (b) ex situ XRD patterns of the PTCDA corresponding to the pristine (black), calciated (red), and decalciated (blue) electrode. Reproduced from ref 96 with permission from the American Chemical Society, copyright [2017]. (c) Charge–discharge profile of PNDIE electrodes in 2.5 M Ca(NO₃)₂ aqueous electrolytes with different pH. (d) Capacity stability and Coulombic efficiency of PNDIE at 5C rate (925 mA g^{-1}). (e) Possible reversible electrochemical redox mechanism of PNDIE. Reproduced from ref 97 with permission from Wiley-VCH, copyright [2017].

MXenes have the common problem of all high-surface-area anode materials: poor first cycle Coulombic efficiency, i.e., a large irreversible loss of capacity.

Organic compounds have also been tested as Ca battery anodes using aqueous electrolytes (Table 1). The aromatic molecular solid 3,4,9,10-perylenetetracarboxylic dianhydride (PTCDA) was tested in a three-electrode cell using activated carbon as the cathode.⁹⁶ Despite some signatures of reversible electrochemical activity, XRD studies indicated rapid deterioration/amorphization upon cycling (Figure 8a,b), which was attributed to the large size of Ca²⁺, as this behavior was not observed for Mg²⁺. For a polyimide (PNDIE) anode and a copper hexacyanoferrate cathode cell, a stable discharge capacity of 40 mAh/g (at C/10 rate) was demonstrated for 1000 cycles⁹⁷ This cycling was, however, performed at a 5C rate, thus potentially masking parasitic reactions/active material dissolution issues. Moreover, for most tests using aqueous electrolytes, the issue of reactivity of protons as well as the role(s) of the oxygen reduction and/or the hydrogen evolution reactions, ORR and HER, respectively, is rarely mentioned, even though the voltage window commonly explored is >1.4 V.

2.2. Electrolytes

Regardless of battery design and concept, the role of the electrolyte is primarily to efficiently separate the two electrodes, which is in practice solved by some kind of membrane/separator layout, and to efficiently transport the ionic charge carrier(s) of interest between the electrodes.⁹⁹ Due to the (very) hard acid– base nature of both the Mg²⁺ and Al³⁺ cations, the corresponding multivalent-battery technologies have very special electrolyte compositions to enable desolvation of the cations at the electrode/electrolyte interfaces-especially because the solvents are very cumbersome to create and handle.⁸ In contrast, Ca battery electrolytes are much more similar to both LIB and SIB electrolytes, and simple salt-in-solvent concepts using standard battery solvents from these technologies seem to provide the needed overall bulk electrolytes properties-likely beause Ca²⁺ is a much softer cation. The remaining open questions are mainly related to if these suffice to provide ample Ca²⁺ transport and exchange at the electrodes.

While Ca battery electrolytes basically must only fulfill the condition of fast and large enough Ca^{2+} -ion transport, through the bulk and at the interfaces, to not limit the power performance of the cell—and in some cases make it at all cyclable—there is the additional issue of enough (electro-)

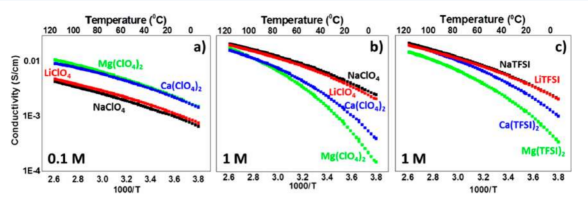


Figure 9. Arrhenius plots (between -10 and 120 °C) for various Li/Na/Ca and Mg salts in EC_{0.5}:PC_{0.5} with (a) 0.1 M and (b,c) 1 M salt concentrations. Reproduced from ref 101 with permission from The Electrochemical Society, copyright [2017].

chemical stability. This is due to the electrochemical potential of Ca being so low and close to Li. From the perspective of electrolyte and anode compatibility, basically the cathodic stability of the salt anion(s) and the solvent(s) used, two main approaches have been employed:

- (i) electrolytes that form a passivation layer on Ca metal anodes by limited decomposition of salt and/or solvents, i.e., are metastable. The main issue is then Ca²⁺ mobility through the passivation layer;
- (ii) electrolytes that do not form any passivation layer—thus being intrinsically stable, primarily suited for nonmetallic anodes, rendering cells with lower energy density.

From the electrolyte vs cathode perspective, both electrolyte designs above hitherto often have rather limited electrochemical stability windows (ESWs), in principle hindering the use of really high-voltage cathodes—but at the same time no practical Ca cathodes (see section 2.3) exist to date.

A major limitation common to all Ca battery electrolytes is the limited number of Ca salts commercially available. Most work has so far employed one or more of these five simple Ca salts: calcium perchlorate, $Ca(ClO_4)_2$, calcium tetrafluoroborate, $Ca(BF_4)_2$, calcium bis(trifluoromethanesulfonyl)imide, Ca-(TFSI)₂, calcium nitrate, $Ca(NO_3)_2$, and calcium borohydride, $Ca(BH_4)_2$. As the ClO_4^- anion is laden with safety issues, the latter four salts are strongly recommended for any practical studies. For aqueous electrolytes, the salt of choice has distinctively been $Ca(NO_3)_2$. Up to now, there has been no systematic investigation on the impact of the nature of the salt in divalent-cation aqueous systems; the choice of nitrate salt most probably is rooted in historical reasons.

In 2015, however, Lipson et al.¹⁰⁰ reported on the synthesis of calcium hexafluorophosphate, $Ca(PF_6)_2$, salt, and furthermore, in 2017, Keyzer et al.⁸³ showed a direct anhydrous synthesis route for the same salt-something they previously developed for the Mg analogue. The latter study is particular significant, as it makes quite an effort targeting the chemical stability of the PF_6^- anion—noting that the chemically softer Ca²⁺ (as compared to Mg²⁺) would have a weaker solvent interaction and thereby accelerate the anion decomposition by stronger cation-anion interaction, i.e., ion-pairing. However, the latter is also weakened by the cation being softer, and to us, it is not clear what interaction is affected the most: cation-solvent or cationanion, calling for computational and spectroscopic local interaction studies. The synthesis is also significant given that the PF_6^- anion is part of the standard LIB electrolyte design, stressing further possibilities to create Ca battery electrolytes

using LIB know-how but with the difference that the anion must comply with cycling vs Ca metal anodes.

Apart from water as the solvent for aqueous electrolytes, more or less standard LIB solvents such as THF, ACN, gBL, PC, DMC, DEC, EMC, DMF, dimethoxyethane (DME), and mixtures of EC and PC have all been applied as matrices for the above Ca salts—even if not all possible combinations. The crucial feature of the metastable electrolytes according to (i) above is that the degradation products arise electrochemically by reduction, often due both to solvent and salt anion, creating a covering and stable film onto the anode, i.e., an SEI. These electrolytes and the truly stable ones are discussed further in section 2.2.1.

In addition to the standard salt-in-solvent design, there are designs relying on ionic liquids (ILs) as solvents and also a few solid-state electrolytes (SSEs), ceramics, and polymers, which we briefly touch upon in section 2.2.2.

Below, the studies centering on Ca conducting electrolytes so far are summarized briefly, but many general Ca battery studies simply use one or two electrolytes without any motivation and from this extrapolate/generalize. We would therefore like to stress that for further development, there is a need for work with the electrolyte in focus to enable:

- (i) a better understanding of the reactions at the interfaces to tailor the SEI (if any),
- (ii) a large mapping of salt(s), solvent(s), concentration(s), and additives and how these alter densities, viscosities, conductivities, ESWs, Ca²⁺ transport, etc.,
- (iii) a use of more LIB and SIB electrolyte know-how as well as from Mg and Al batteries.

2.2.1. Liquid Electrolytes. Nonaqueous organic-solventbased liquid electrolytes have by far been the prevailing choice for Ca batteries, even if aqueous electrolytes have been shown to improve the kinetics of Ca intercalation at the electrolyte/ electrode interface. Systematic electrolyte studies are hitherto very scarce. We note the early "screening" study by Aurbach et al.³² investigating the electrochemical behavior of a Ca metal anode in different organic electrolytes composed of ACN, THF, gBL, and PC and the salts $Ca(ClO_4)_2$ and $Ca(BF_4)_2$ and the similar study by Hayashi et al.⁴² using $Ca(ClO_4)_2$ as the salt and PC, DMC, DEC, EMC, gBL, DMF, and ACN as solvents. In contrast, most often a single or at most two different electrolytes have been employed to study an anode or cathode in half- or fullcell designs. The three most prominent examples are the studies on reversible Ca plating and stripping using a Ca metal anode by Ponrouch et al.⁷⁴ using different Ca salts in EC:PC, by Wang et al.⁷³ employing Ca(BH₄)₂ in THF, and by the groups of Nazar⁷⁸ and Fichtner⁷⁹ using Ca salt based on fluorinated alkoxyborates anions in DME. These studies are covered extensively in section 2.1 with respect to the functionality for the anode and interface creation. We note here, however, that these research groups used different salt concentrations—with 0.25 M Ca[B(hfip)₄]₂, 0.45 M Ca(BF₄)₂, 0.5 M Ca[B(hfip)₄]₂, or 1.5 M Ca(BH₄)₂—a far from small design difference and significant for the electrolyte properties. Another note is that these electrolytes have a fair concentration of oxygen atoms by virtue of the carbonate solvents, and combined with the stability of the CaO (Figure 4), this might have profound effects on the cyclability of Ca-metal-based cells (see section section 2.1, Figure 5).

In a recent study, the Ca salt type and concentration was systematically varied in a matrix of EC:PC.¹⁰¹ The target was truly electrolyte oriented to connect various fundamental physicochemical electrolyte properties such as ion conductivity, viscosity, and the Ca²⁺ cation solvation by solvent and anions, with the practical electrochemical performance. It also compared Ca²⁺ vs Li⁺, Na⁺, and Mg²⁺ in order to strengthen the fundamental understanding of the importance of underlying interactions for the phenomena and correlations observed. It was deduced that for low salt concentrations (0.1 M), when almost total ion-ion dissociation can be expected, the divalent systems are more conductive than the corresponding monovalent-cation-based systems, while for higher concentrations (1.0 M), both the viscosity and the ion-pairing increases relatively more for the divalent systems, and consequently, the ion conductivity drops (Figure 9). At elevated temperatures, such as those used to enable the reversible stripping/plating in the former studies, however, these differences become less obvious (Figure 9b,c). Also, the nature of the anions mattersemploying the TFSI salt renders more conductive electrolytes for the 1.0 M system (Figure 9b,c)—primarily by less extensive ion-pairing.

The ionic conductivities were all above the target value often given for LIB electrolytes (1 mS/cm): ca. 3.5 mS/cm even at room temperature for the 0.1 M systems and slightly lower, ca. 2.0–2.8 mS/cm for the 1.0 M systems. The differences correlate with the viscosity of ca. 3 cP increasing to a vast 70 cP, which again can be mitigated by raising the temperature (Figure 10).

At a molecular level, the differences, especially with respect to the ion-pairing, can be analyzed by vibrational spectroscopy, showing that the cation solvation numbers are lowered from 6.7 to 5.3 from 0.1 to 1.0 M Ca(TFSI)₂ concentration but also that Ca^{2+} has a larger first solvation shell than all other cations (Li/Na/K/Mg).³² This should affect the mobility in the bulk as well as the (de)solvation dynamics at the electrolyte/electrode interfaces.

A note of caution is also needed with respect to the discussion of ion conductivities above; a 0.1 M $M(TFSI)_2$ electrolyte has nominally 50% more ions than a 0.1 M MTFSI electrolyte. Hence, analysis of (cation) transference numbers and not only the total ion conductivities is crucial.

Comparing the above systems, the EC/PC-based electrolytes exhibit a 4 V ESW and are thus in principle suited for medium—high-voltage cathode materials, while the THF-based electrolyte is more limited (<3 V).

Turning to water, real aqueous electrolytes as well as water present in conventional aprotic electrolytes have been shown to stimulate formation of a calcium hydroxide film passivating a Ca metal anode.³² Hexacyanoferrates were used as cathodes in the

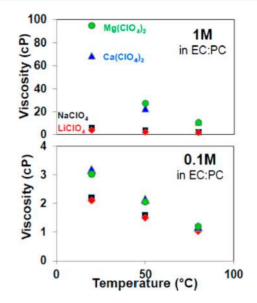


Figure 10. Viscosities for $Ca(ClO_4)_2$ in EC:PC electrolytes for the two salt concentrations (0.1 and 1.0 M) as a function of temperature. Reproduced from ref 32 with permission from the The Electrochemical Society, copyright [1991].

early 2000s,⁴⁰ and more recently, this system has been reinvestigated,⁴¹ also using nonaqueous electrolytes,⁵² showing both a very reversible electrochemical response and enhanced capacity from the water addition. As an example, ca. 17% of water in a Ca(ClO₄)₂ in ACN electrolyte led to enhanced redox activity, attributed to solvation effects.⁵³ The role of water in the electrolytes for the intercalation is not fully elucidated—some studies point to minor changes in the host.⁵⁵ We caution for other events such as proton intercalation,¹⁰² electrolyte decomposition, and/or current collector corrosion, which all can lead to misleading conclusions/assumptions with respect to calcium intercalation.

Protic electrolytes have also been employed for $Ca-O_2$ batteries, but to reduce the reactivity of Ca with water, a solvent mixture of 2:1 methanol:water without any Ca salt inside was employed—as its role is more to transport O_2 to the anode¹⁰³ than to transport any Ca²⁺.

A bit more fundamental aqueous electrolyte properties were partly investigated by Wang et al.⁴¹ using 1 M Ca(NO₃)₂ as salt. The target was fast migration of Ca²⁺ in electrodes, but Lee et al. showed more elaborately that highly concentrated aqueous electrolytes, also known as water-in-salt electrolytes (WISEs),¹⁰⁴ are also possible for Ca²⁺.¹⁰⁵ The lowered hydration number, by increasing the Ca(NO₃)₂ concentration and thereby a greater amount of anions coordinating the cation,^{105,106} diminished the activation barrier for intercalation, leading to improved cell cycling performance.

For all the above aqueous electrolytes, the issue of CaO formation, in contrast to the carbonate-based nonaqueous electrolytes, is a nonissue, as cells are not targeting Ca metal anodes.

2.2.2. Other Concepts. As well as there are few modern studies of alternative Ca battery concepts, the same is true for the electrolytes employed (disregarding the molten salt electrolytes covered extensively in section 1.1). Yet, application of solid-state electrolytes (SSEs)⁸⁴ and ionic-liquid (IL)-based electrolytes⁶³ has been given some minor attention.

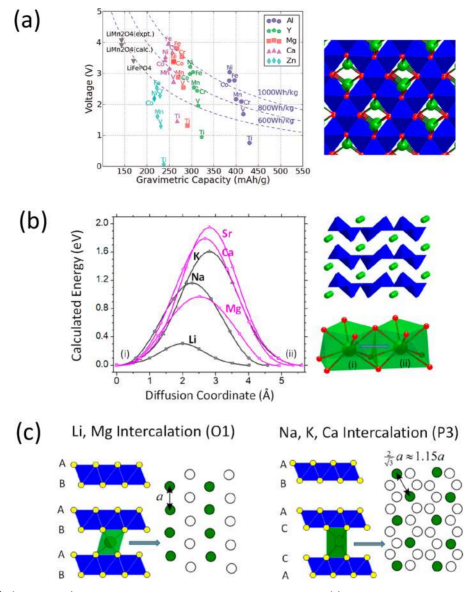


Figure 11. Some Ca^{2+} (comparative) intercalation electrode properties calculated by DFT: (a) intercalation voltage vs specific capacity in spinel- $Ca_{T}[M_{2}]_{0}O_{4}$, reproduced with permission from ref 57, copyright [2015] The Royal Society of Chemistry. The right panel shows the spinel structure. Color code: O in red, TM in blue, and Ca in green. (b) Diffusion barriers in $A_{x}V_{2}O_{5}$ for A = alkali or alkali earth cation, adapted with permission from ref 123, copyright [2014] American Chemical Society. The right panel shows the structure of $V_{2}O_{5}$ with potential sites for Ca insertion and the two sites (i) and (ii) involved in the Ca hop. Color code: O in red, TM in blue, and Ca in green. (c) The most stable $A_{0.5}$ TiS₂ structures for A = Li, Mg, Na, K, and Ca. Color code: S in yellow, TM in blue, and Ca in green. Reproduced with permission from ref 128, copyright [2016] American Chemical Society.

Starting with the latter, the high ionic mobility, large ESWs, and great solubility power and safety trends—low or no vapor pressure—of ILs have attracted the attention of the battery community at large as electrolyte solvents. For Ca batteries, however, we know only of one single study, wherein Shiga et al.⁶³ investigated a nonaqueous Ca–O₂ battery design employing 0.1 M Ca(TFSI)₂ in the IL DEME TFSI and observed Ca plating/stripping at 60 °C. The CV indicates only a slightly reversible deposition of Ca, and the SEI formed was by Raman spectroscopy revealed to have a composition characteristic of TFSI anion decomposition.

Turning to SSEs, solid CaF_2 was used early vs alloys such as Ca-Bi and Ca-Sb.^{84,107} Much more recently, metal borohydride electrolytes were investigated by DFT calculations, but in a comparative study, the Ca^{2+} diffusion was predicted to be too slow for practical application.¹⁰⁸ Overall, very few papers have

reported on Ca²⁺ SSEs—despite the current "hype" on ASSBs, likely due to the difficulty to obtain materials with high-enough ionic mobility for this multivalent chemistry.

Solid polymer electrolytes (SPEs) containing Ca have been until now reported at the materials level but not with any real battery tests, including an early work almost 25 years ago using "home-made"(!) Ca(TFSI)₂ dissolved in a poly(ethylene oxide) matrix investigating basic properties such as cation coordination, ion conductivity, phase transformations, and so on.¹⁰⁹ We are aware that continued research on SPEs has been pursued in several laboratories since then but with no results in the open literature. A deviation from the SPE concept is gel polymer electrolytes (GPEs), and very recently, a Ca conducting GPE based on Ca(NO₃)₂ and a PEGDA cross-linked network was presented.¹¹⁰

2.3. Cathodes

As only a few Ca battery cathodes have been successfully tested electrochemically, this section follows a different layout than for the anode and electrolyte, and it also emphasizes the role of computational studies. Section 2.3.1 analyzes the relation between intercalation and crystal chemistry, which is the base for Ca-cathode identification/design. The following subsections discuss targeted cathode properties: the energy density, which is maximized by increasing both the electrochemical capacity and the operation potential (section 2.3.2), and the practical power rates which depend, among other factors, on the Ca mobility in the cathode (section 2.3.3). Finally, alternatives to intercalation materials are presented in (section 2.3.4).

2.3.1. Intercalation and Crystal Chemistry. Intercalation, i.e., reversible insertion into a host with minimal structural change, is possible for a variety of electrochemically relevant metal ions (35,36,41,111–113). The basic requirements are an open framework of interconnected sites wherein the intercalated ion can diffuse and an electronic band structure able to reversibly accept/donate electrons.¹¹⁴⁻¹¹⁸ The high redox potential needed to enable high cell voltages limits the suitable host materials to transition-metal (TM) containing compounds, just as for LIBs. The quest for cathode materials often relies on chemical intuition, and then, attempts are done to locate trends for a given host and different intercalated ions (44,56,57,101,112,119-128). Computational studies have been particularly important, since DFT permits the exploration of both existing and virtual materials, i.e., materials not (yet) made experimentally.^{129,130} DFT (comparative) studies of Ca²⁴ include intercalation voltages (57,121-123,131-133) (Figure 11a), migration barriers (56,57,120,123,133) (Figure 11b), and phase stability (Figure 11c).^{120,122,128,134} These studies uncover that trends are not straightforward. For Ca²⁺, its divalent charge promises similarities with Mg²⁺ intercalation chemistry but is also alike Na⁺ in size (ionic radii in metal oxides: $r(Ca^{2+})_{VI} = 1.0$ Å and $r(Na^+)_{VI} = 0.99$ Å¹³⁵), resulting in similar site preference. Is then Ca intercalation reminiscent of Mg and/or Na intercalation? A plausible answer is that each metal is unique, and accordingly, the suitable host materials may be substantially different from one metal ion to other. The relation of intercalation to crystal chemistry is essential to identify possible cathode host materials and can be further analyzed looking at the host materials used as today's commercial LIB cathodes: olivine-LiFePO₄, spinel-LiMn₂O₄, and layered-LiCoO₂ (115,117,136,137).

In spinel-LiMn₂O₄ (Figure 11a), the Mn_2O_4 framework of edge-sharing octahedra provides a three-dimensional network of interconnected tetrahedral sites where the Li ion is located and can diffuse.¹³⁸ DFT investigations of virtual spinels $Ca_{T}[M_{2}]_{O}O_{4}$ (Ca in tetrahedral coordination) predicted some promising cathode properties (Figure 11a),⁵⁷ but yet, the preference of Ca ions for larger sites makes the preparation of such compounds very unlikely. Indeed, the CaMn₂O₄ stable polymorph-0.9 eV more stable than the spinel-is the marokite with Ca in 8-fold coordination sites.^{134,139} While there are a few nitrides with Ca in tetrahedral coordination,¹⁴⁰ CaNiN¹⁴¹ and Ca₃N₂,¹⁴² the Inorganic Crystal Structural Database (ICSD) does not include any oxides with Ca in tetrahedral sites, and the ionic radii of tetrahedral Ca are not even listed for halides, chalcogenides,¹⁴³ or oxides.¹ Metastable structures with tetrahedral Ca could, however, be attempted by soft chemistry routes such as cation exchange, solvothermal, coprecipitation, or sol-gel processes, but even if $Ca_T[M_2]_OO_4$ spinels could be prepared, prolonged battery cycling might promote the thermodynamically stable structure, as previously observed for Mg spinels.^{144–147} Site preference is certainly a prime difference between the intercalation chemistry of the larger Ca²⁺, Na⁺, or K⁺ ions and the smaller Li⁺ or Mg²⁺ ions, which is also influenced by their charge.

Turning to the olivine LiFePO₄, its hexagonal close-packing of oxygen has two different octahedral sites, M1 and M2,¹⁴⁸ where only the former provides channels for Ca diffusion.¹⁴⁹¹⁵⁰ The M1 octahedron geometry is comparatively more distorted and smaller, whereas the M2 octahedron is more regular and larger. In LiFePO₄, the Li⁺ ions occupy the smaller M1 site, and the Fe²⁺ cations occupy the M2 site. In olivine-MgFeSiO₄ and MgMnSiO₄, some degree of cation mixing is observed,^{151,152} because of the similarity in the radii and equal charge of the two cations. In Ca olivines, however, as in the mineral kirschsteinite CaFeSiO₄,¹⁵³ the larger Ca²⁺ occupies the M2 site and is thus immobile.¹⁵⁴ The radii and the oxidation state of the intercalant and the TM ions combined regulate the cation distribution.¹⁵⁵ and thus the suitability of given the structural type to intercalate different metals.

The structure of O3-LiCoO₂ (α -NaFeO₂ structural type) can be viewed as "ordered rock salt", in which layers of octahedrally coordinated Li⁺ and Co³⁺ ions alternate within the cubic close packed oxygen array. The lithium ions can be reversibly removed from and reinserted within the triangular lattice of sites formed by the Li ions in a plane.^{156–158} Like Li, Na combines with trivalent TM to form ordered rock-salt type structures with potential application as cathode materials in Na batteries.^{159,160} In contrast, Ca²⁺ and TM ions preferably crystallize in structural types such as perovskites, postspinels, K₄CdCl₆, and so on, and due to cation distribution rules, few ordered rock-salt structures have been achieved: nanoclusters of CaMnO2 superstructures¹⁶¹ and layered-Ca_{0.47}CoO₂.¹⁶² Cabello et al. prepared the latter metastable phase (see ref 163 for phase stability in the Ca-Co-O system) by the Pechini method¹⁶⁴ and reported a reversible capacity of ca. $30-100 \text{ mAh/gCaCo}_2O_4$ for 30 cycles, but deeper structural characterization is required to determine the Ca and Co distribution and its effect on the (low) capacity and cycling stability.^{165–16}

Other crystal chemistries to analyze are the "traditional" insertion host materials, such as V_2O_5 , MoO₃, and TiS₂, ^{36,168,169} that were explored long ago and found to exhibit a rich intercalation chemistry for a variety of intercalants. These materials have been extensively studied for Ca—especially $V_2O_5,^{35,43,125,170}$ a material that intercalates Li, 171 Na, 172 and even K.¹⁷³ Under ambient conditions, V₂O₅ crystallizes in a layered structure consisting of VO5-square pyramids sharing edges and corners (Figure 11b). As layered structures favor intercalation reactions and the appealing high redox potential of the V^{5+}/V^{4+} couple, V_2O_5 was one of the first materials tested for Ca intercalation. Pioneering work by Amatucci et al.³⁵ reported a reversible capacity of 465 mAh/g for V_2O_5/PC nanocomposites,⁴⁴ further studied by Hayashi et al.^{42,43} However, DFT investigations suggest a more sluggish kinetics for Ca intercalation than for Li, Na, and even Mg^{120,123} (Figure 11b and section 2.3.3), and experiments show formation of protonated phases (section 3).^{125,170} Another classical material in intercalation chemistry, TiS_2 ,^{168,174,175} is among the very few compounds for which reversible Ca intercalation has been reported.^{124,176} Recent DFT investigations indicate that distinct site preference of the intercalant ions drives the intercalation reaction in TiS₂ across different intermediate A_xTiS₂ phases

(Figure 11c).¹²⁸ Thus, Mg and Li occupy the empty octahedral sites in the interlayer space of the initial O1 lattice (CdI₂-type structure), forming stage compounds as intercalation proceeds. In contrast, the large intercalants Na, K, and Ca occupy the prismatic sites in a P3 stacking (Figure 11c) at intermediate concentrations ($x \approx 0.5$), as this intercalant site topology minimizes the in-plane electrostatic repulsions.^{128,177,178}

Hence, attempting to accomplish Ca battery cathode design with a large amount of parallelism between Ca and better known intercalation chemistries (Li, Na, or even Mg) is not really possible, as the crystal chemical analogies are too intricate.

2.3.2. Energy Density: Voltages and Capacities. In an insertion reaction, the intercalated ions are incorporated in the crystalline structure of the host compound, and electrons are added to its band structure. In a simplified model,^{169,174,179} the intercalant metal ions are fully oxidized, donating electrons to the unoccupied levels of the band structure of the host compound, which arise from the antibonding d-states of the TM. Thus, the nature and the oxidation state of the transition-metal ion are the main determinants of the intercalation voltage, which depends also on the ionocovalent character of the cation–anion bond, primarily driven by the nature of the anion and in a lesser extent by the crystal structure.^{169,179,180} In short, the Ca intercalation voltage for a given TM redox couple should follow the trend known from Li-cathode intercalation materials: chalcogenides < oxides < poly-oxoanionic.^{136,179,181}

As there is so far very limited experimental data on Ca intercalation voltages available, DFT calculations have been extensively used to predict average values for several prospective materials (57,119,121,124,131–134,147,182). In a first step, the voltage is usually calculated between the fully intercalated and deintercalated compounds; this is x = 0, 1 in Ca_xTiS₂,¹⁸³ x = 0, 2 in Ca₂Fe₂O₅,¹⁸⁴ x = 0, 3 in Ca₃Cr₂(SiO₄)₃,¹⁵⁴ and so forth. While some oxidation states assumed might not be realistic—as the Mn⁷⁺ of fully deinserted Ca₃Mn₂O₇ and Ca₄Mn₂O₇¹³³—this gives a fast overview of voltage trends. Subsequently, the average intercalation voltages for any intermediate compositions can be calculated using structural models of ordered calcium/vacancy arrangements, as has been done for instance in the perovskites Ca_xMO₃¹³² or the Chevrel phase Ca_xMO₆S₈.

The DFT-predicted Ca2+ average intercalation voltages for oxides are similar to those of Li⁺ (Figure 11a), which is not surprising taking into account the metals' similar reduction potentials. For the Mn³⁺/Mn⁴⁺ couple, the predicted voltages are 3.8 V (marokite) and 3.1 V (postspinel and spinel),^{57,134} and for the Co^{3+}/Co^{4+} couple, the predicted voltages are 3.3 V $(Ca_{3}Co_{2}O_{6})$, 3.2 V $(Ca_{3}Co_{4}O_{9})$, and 3.8 V $(Ca_{2}Co_{2}O_{5})$.¹⁸⁵ As expected, the effect of the anion is critical to tune the intercalation potential, following the trends widely reported for Li materials.^{136,181,186} Covalent hosts display lower voltages than the more ionic oxides, as evidenced by the predictions for the series of Chevrel phases: Mo_6S_8 (2.1 V), Mo_6Se_8 (2 eV), and $Mo_6Te_8 (1.4 V)$.¹³¹ Å nice comparison is offered for oxo- vs thiospinels,⁵ ^{7,121} where the calculated voltages for the Mn³⁺/Mn⁴⁺ couple are 3 V for CaMn₂O₄ and 1.3 V for CaMn₂S₄. Similarly, the inductive effect raises the intercalation voltage in polyoxoanionic compounds: the Mn^{2+}/Mn^{4+} couple is at 3.6 V for pyroxene-CaMn $(SiO_3)_2$ and 3.8 V for kutnahorite-CaMn- $(CO_3)_2$.¹⁵⁴ Unfortunately, any comparison of these DFT predictions with experimental data is biased by issues related to the experimental setups (section 3); yet, some reported values are 3.5 V for $Ca_3Co_2O_6^{-187}$ (DFT: 3.36 V¹⁸⁵), 3.03 V for $V_2O_5^{-32}$ (DFT: 3.28 V¹²⁰), 1.5 V for TiS₂¹²⁴ (DFT: 1.7 V¹⁸³), and no more than 1.3 V for MoO_3^{188} (DFT: 2.23 V¹⁸⁸). The large discrepancy for the latter supports that side-reactions might be taking place, as discussed in more detail in section 3.

Overall hosts with intercalation voltages of 2-3 V are appealing for electrolytes with low to medium anodic stability. Yet, given that electrolytes with higher anodic stabilities are developed, ^{78,79} the use of high-voltage cathodes (and here many oxides and poly-oxoanionic materials are in the 3-4 V range) would enable high-energy density Ca battery cells, pending the capacities.

The specific capacity for any intercalation host is dictated by the amount of Ca²⁺ that can be cycled between the oxidized and reduced state and the overall formula weight, and the former depends on the available crystallographic sites for Ca ions and the operating redox couples. If one considers for instance the perovskite structure CaMO₃, a frequent structural type for Catransition-metal oxides, looking at the crystallographic sites, the maximum theoretical capacity corresponds to the exchange of 1 Ca ion per TM, but this would drive the TM to have a formal oxidation state of +6, which is clearly unrealistic for most cases (excepted Mo).¹³² Hence, theoretical capacities should be calculated considering reasonable TM oxidation states. The calcium cobalt oxides, Ca₃Co₂O₆, Ca₂Co₂O₅, and Ca₃Co₄O₉, is another set with the capacity being regulated by the redox couple (Co^{4+}/Co^{3+}) , despite a larger amount of available Ca sites.¹⁸⁵ It is thus most often the oxidation states of the TM ions and not the available crystallographic sites which determine the maximum specific capacity for Ca cathodes. In such cases, the specific capacity is similar to that achievable by Li⁺ intercalation. For TiS₂, DFT calculations predict a large voltage step at the stable intermediate $Ca_{0.5}TiS_2$ composition (Figure 11c),¹²⁸ with calculated average voltages of 1.7 V (Ti^{4+}/Ti^{3+}) and 0.9 V (Ti^{3+}/Ti^{3+}) Ti²⁺).¹⁸³ Hence, there would be a significant voltage drop if both redox couples were considered, and Ca insertion is likely limited to 0.5 mol Ca/TM and the Ti^{4+}/Ti^{3+} redox couple, delivering a similar specific capacity as 1 mol Li/TM. The consequence is that statements found claiming double specific capacity for Ca materials as compared to their Li analogues should be assessed carefully. Light TMs able to exchange two or more electrons are a priori: V^{3+}/V^{5+} , Mn^{2+}/Mn^{4+} , Cr^{3+}/Cr^{6+} , and Mo^{3+}/Mo^{6+} (not so light), and thus, their compounds should be the best candidates to surpass the capacities today achieved for Li cathodes.

For decades, the dream of the LIB research community has been to exchange two electrons per TM ion. As exposed by Goodenough, two redox couples can be accessed where the cation redox couples are "pinned" at the top of the O 2p bands, but to take advantage of this possibility, it must be realized in a framework structure that can accept more than one Li atom per transition-metal cation.^{115,180} The latter requirement restricts the candidate materials, and in the Li-ion battery technology prompted attention to the Li_2MSiO_4 silicates (M= Fe, Mn, Co in tetrahedral sites) which could in theory exchange two electrons for each TM involving the M^{3+}/M^{2+} and M^{4+}/M^{3+} couples.¹⁸⁹⁻¹⁹¹ For these cathodes, the structural collapse driven by the instability of the tetrahedral TM ion as its oxidation state varies is a pending issue.^{190,192} For Ca, this obstacle seems solvable as Ca²⁺ combines with octahedral Fe²⁺ and Mn²⁺ in major classes of minerals (pyroxene, garnets, double carbonates) and certain ternary oxides. Another great challenge in the LIB community is the effective utilization of the V^{5+}/V^{4+} , V^{4+}/V^{3+} , and even V^{3+}/V^{2+} redox couples,¹¹⁷ and this also

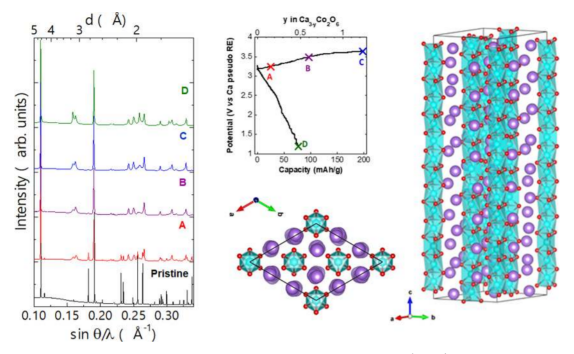


Figure 12. Potential vs capacity profiles from potentiodynamic cycling with galvanostatic acceleration (PCGA) experiments at 100 °C and C/200 rate in $Ca_3Co_2O_6//Ca$ cells, progressively oxidized (labeled A in red, B in purple, C in blue) or oxidized and reduced (D in green), and corresponding diffraction patterns (black corresponding to pristine $Ca_3Co_2O_6$ electrode). The crystal structure of the oxidized $Ca_{2.3}Co_2O_6$ phase (corresponding to sample C) showing an incommensurate modulation is also depicted. Adapted from ref 187, copyright [2018] The Royal Society of Chemistry.

represents a priority route to achieve high-capacity Ca intercalation.

Realization of theoretical into practical capacities requires that the crystal and electronic structures are stable within the compositional limits of the intercalation reaction. Severe structural rearrangements are a major concern, and in the limit, irreversible structural phase transformations are a threat for reversible specific capacity even if they occur at the microstructure level.¹⁹³ Unit cell volume variation upon calcium deinsertion/insertion is a first measure and can be inferred from preliminary characterization: 18% for $Ca_x TiS_2$, ¹²⁴ 1% for $Ca_x MoO_3$, ¹⁸⁸ 3% for $Ca_x V_2O_5$, ⁴³ and 3% for $Ca_x NaFePO_4F$. ¹⁹⁴ Small volume variations found experimentally might, however, also indicate limited Ca insertion or even proton insertion. The actual amount of intercalation is often unknown, as sidereactions related to electrolyte decomposition might significantly contribute to the observed specific capacity, especially for tests carried out at high temperature, leading to overestimation in the value of x. For TiS_2 the measured specific capacity is 500 mAh/g, corresponding to an $x \approx 0.9$ nominal, while phases with $x \approx 0.2$ and 0.5 are detected by differential absorption tomography at the Ca L2 edge and XRD, respectively.¹²

Since the TM oxidation state varies along the charge/ discharge of the battery, the host compound can become metastable with respect to other crystal structures at intermediate calcium contents. This is the case of Ca deinsertion from $Ca_3Co_2O_6$, for which DFT predicts a cell volume variation of 3% for 1 mol Ca extraction, which is in good agreement with experiments.^{154,187} Regardless of the small volume variation, a phase transformation occurs upon Ca deinsertion. $Ca_3Co_2O_6$ crystallizes in the K₄CdCl₆-type structure¹⁹⁵ where the Co–O atoms form chains of alternating face-sharing CoO₆ octahedra and CoO₆ trigonal prisms along the hexagonal *c*-axis (Figure 12), separated by eight coordinated calcium atoms that according with DFT results¹⁸⁵ can diffuse in the structure albeit with sluggish kinetics. It has been demonstrated¹⁸⁷ that ca. 0.7 Ca ions can be extracted from $Ca_3Co_2O_6$ at ca. 3.5 V (Figure 12) but with a hindered reinsertion process presumably due to a large Ca desolvation energy at the electrolyte/electrode interface-hence, not an intrinsic property of the cathode material itself. The authors also concluded that Ca extraction results in the formation of a modulated structure with all Co ions in octahedral coordination,¹⁹⁶ and parallel DFT investigations¹⁸⁵ confirmed that the driving force for this phase transformation is the trend of Co ions to adopt octahedral coordination according with crystal field stabilization energies. In addition to this change, DFT indicates that the Ca sublattice rearranges making it unlikely to insert Ca ions to regain the initial Ca₃Co₂O₆. Under this perspective, irreveribility is an intrinsic issue of the material itself. Thus, further investigations with improved electrolyte formulations are needed to properly assess the reversible Ca intercalation in $Ca_2Co_2O_6$.

2.3.3. Diffusion in Host Materials. While the abovediscussed capacity and voltage values determine the specific energy achievable, basically by thermodynamics, cathodes must also enable acceptable diffusion rates (kinetics) for the intercalated Ca^{2+} ions at any degree of insertion. Note that diffusivity may vary substantially with the host composition due to Coulombic repulsion between the diffusing Ca^{2+} ions and the TM cations in different oxidation states.^{149,197} At present, limited mobility of the Ca^{2+} ions is one of the major concerns in multivalent cathode design. Factors determining diffusion are rooted in crystal chemistry; the pathways should be wide enough for the ions and exhibit a favorable topology to diminish the electrostatic interactions between the diffusing Ca^{2+} cation and the lattice constituents.

A good ionic diffusion coefficient ($D \approx 10^{-12} \text{ cm}^2 \text{ s}^{-1}$ for a C/2 rate) is a prerequisite for battery electrode materials. There are no direct measurements of Ca diffusion coefficients reported, and estimates from electrochemistry are hampered by the poor

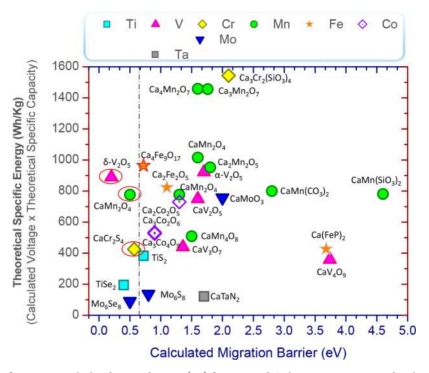


Figure 13. Theoretical specific energy vs calculated energy barriers (eV) for proposed Ca hosting TM compounds. The specific energy is estimated from the calculated average voltage for a particular redox couple and the corresponding theoretical specific capacity. The vertical line denotes the criterion for good cathode performance (<0.650 eV, see text). The red circles indicate virtual compounds. Data taken from 57, 120, 121, 131–134, 154, 182–185, 200, 201.

reliability of testing protocols and problems with the stabilities of electrolytes and interfaces (section 3). The migration energy barrier, i.e., the activation energy for an ionic hop, can be used as proxy,¹⁴⁹ and for a reasonable C-rate such as C/2, it should be <0.525 and <0.650 eV for micrometric and nanosized particles, respectively.^{9,56} These barriers can be extracted from DFT calculations using the nudged elastic band method (NEB),¹⁹⁸ as shown in Figure 11b for a single A ion diffusing in a V₂O₅ lattice; the energy landscape along the diffusion path has a maximum at the saddle point, in this case, a face of the shared triangular face.

In Figure 13, reported calculated energy barriers for Ca migration in various TM compounds are depicted, and notably, <0.650 eV has been predicted for some virtual spinel- $Ca_{T}[M_{2}]_{O}O_{4}$ with Ca occupying the tetrahedral sites.⁵⁷ Rong et al. provided useful design guidelines to enhance ionic mobility in multivalent cathodes and identified the local topology and site preference as key parameters.⁵⁶ Hence, a good mobility was predicted in spinel-Ca_T $[M_2]_0O_4$, since the Ca ion would diffuse from a nonpreferred site (tetrahedral) to a preferred site (octahedral). Similarly, a barrier as low as 0.2 eV is predicted for the virtual δ -CaV₂O₅, where Ca ions occupy a nonpreferred site (bicapped-tetrahedral).¹²⁰ However, while such a δ -AV₂O₅ structure is known for A = Mg and Li, in the stable form of CaV₂O₅, the Ca ions adopt an 8-fold coordination.¹⁹⁹ Interestingly, the calculated barriers for the latter are >1.5 eV, in agreement with the unsuccessful attempts to extract Ca ions from CaV₂O₅.¹²⁵ Combining the above trends leads to the conclusion that metastable (and in many cases virtual) compounds are a better research target for unravelling calcium conducting cathodes. Yet, the already mentioned difficulties in synthesis, possible instability with cycling, and in general a voltage penalty are downsides of metastable materials.

In contrast, minerals are stable naturally occurring compounds. The garnet, pyroxene, and dolomite mineral groups all possess suitable Ca migration pathways, and promising capacities and voltages have been predicted for garnet- $Ca_3Cr_3Si_3O_{12}$ (uvarovite), pyroxene- $CaMn(SiO_3)_2$, and double carbonates- $CaMn(CO_3)_2$ (kutnahorite).¹⁵⁴ Nevertheless, their calculated energy barriers are larger than for oxides (Figure 13), with a maximum of 4 eV for the pyroxene. A possible reason for this is the higher concentration of cations in these 3D polyoxoanionic structures, arising from the transition metals and the silicate/carbonate groups. Paradoxically, the garnet structure is well-known for fast Li-ion conductors.^{202,203} The hampered Ca mobility excludes not only these minerals but also the synthetic materials that exhibit these crystal structures. On the other hand, Lipson et al.¹⁹⁴ reported Ca intercalation into the layered structure of NaFePO₄F, boosting expectations of Ca intercalation in related 2D hosts.

The energy barriers >1.5 eV reported for perovskite-CaMoO₃, α -V₂O₅, and marokite-CaMn₂O₄ TM oxides (Figure 13) are in agreement with these materials being electrochemically inactive in Ca cells.^{125,132134} In these oxides, the divalent charge of Ca^{2+} combines with its size to produce these large energy barriers. As a comparison, across ions, the migration barriers in α -A_xV₂O₅ (Figure 11b) follow the order $Li^+ < Mg^{2+} < Na^+ < K^+ < Ca^{2+} <$ Sr^{2+123} The same trend is reported for the marokite-A_xMn₂O₄, reaching a value of 1.8 eV for Ca^{2+} .^{122,134} In both $\dot{V_2O_5}$ and marokite, the diffusing A ions occupy eight-coordinated sites, which share a triangular face to form a channel running along one crystallographic direction. Triangular faces of the highcoordination-number polyhedra in V₂O₅ and marokite are too small to enable large cations through. This plays against Ca²⁺ diffusing better than Mg²⁺ ions in many inorganic structures. The larger rectangular faces shared by trigonal prisms seem more favorable for conduction of large cations (Ca^{2+}, Na^{+}, K^{+}) , as inferred from the experimental diffusion coefficients for trigonal prismatically vs. octahedrally coordinated Na ions.^{36,17}

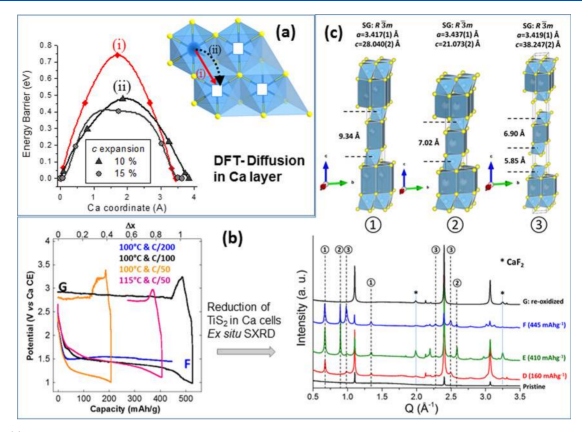


Figure 14. (a) Calculated energy barrier for Ca diffusion in TiS₂. The diffusing Ca ion jump from an occupied to a vacant octahedral site across either the S–S dumbbell (i) or the intermediate tetrahedral site (ii). (b) Curves from Ca//TiS₂ cells cycled at 100 °C and C/200 (blue), C/100 (black), and C/50 (orange) or at 115 °C and C/50 (pink). Note that the indicated potential is E_{WE} – E_{CE} . (c) SXRD collected at different stages of TiS₂ reduction in Ca cells and the associated refined structural model. For clarity, each pattern has been labeled with the corresponding capacity achieved upon reduction. Reprinted with permission from ref 124, copyright [2018] American Chemical Society.

Compared to monovalent cations, multivalent intercalants will suffer stronger electrostatic interactions with the surrounding cations (repulsive) and anions (attractive).^{56,197} In the perovskite structure CaMO₃ (M = Cr, Mo, Mn, Fe, Co, Ni), the high energy barrier for Ca migration (2 eV) arises from the facesharing between the transition-metal (TM) octahedra and the empty sites available for Ca diffusion.¹³² In the related brownmillerite structure, which has a crystal structure analogous to the perovskite but with ordered oxygen vacancies, the latter defines a possible pathway for Ca diffusion, and weaker electrostatic repulsion lowers the calculated energy barrier down to 1.3 eV for Ca₂Co₂O₃¹⁸⁵ and 1.0 eV for Ca₂Fe₂O₅,¹⁸⁴ but this is still not low enough to consider them as viable Ca cathodes.

Chalcogenide networks provide better screening of electrostatic interactions and consequently lower the migration barriers—those calculated for Mo_6S_8 (0.8 eV),¹⁸² Mo_6Se_8 (0.5 eV),¹⁸² TiS₂ (0.72 eV),¹²⁴ and TiSe₂ (0.4 eV)²⁰⁰ are so far the lowest reported for existing compounds. For reversible electrochemical Ca²⁺ intercalation in TiS₂ (Figure 14),¹²⁴ the first phase formed upon reduction is found to be the result of an ionsolvated intercalation mechanism, with solvent molecule(s) being cointercalated with the Ca²⁺ cation. Upon further reduction, new non-cointercalated calcium containing phases seem to form to the expense of unreacted TiS₂. In the noncointercalated phases, the Ca ions are likely in trigonal prismatic sites, in agreement with the above-discussed P3structure stabilization for large intercalants¹²⁸ and forestalling an enhanced ionic mobility respective to the initial octahedral coordination. From the combination of experimental and computational techniques, it was concluded that the cationsolvated intercalation mechanism improves Ca^{2+} diffusion, by both expanding the interlayer space (here the DFT barrier lowers to 0.5 eV with a more stable transition state denoted as (ii) in Figure 14a) and the solvent-screened cation charge. In practice, however, solvent cointercalation should be avoided to enable high cell energy densities.

2.3.4. Alternative Cathode Materials. The overall sluggish Ca diffusion in inorganic intercalation host materials is a major concern for enabling useful high-energy rechargeable Ca batteries with reasonable power performance. Alternative cathode concepts are therefore given attention, including organic electrodes offering mechanical flexibility, mild synthesis approaches, processability, and ample structural and chemical tuneability that can result in high specific capacities, up to 500 mAh/g for quinones, at attractive intercalation voltages.^{204–206} A circular battery economy is viable by preparing the materials from renewable sources.^{207,208} The limitations are primarily the large solubility of active material in electrolytes, capacity fading, and low rate performance. Even if many advances have been made within different classes of organic cathodes: carboxylates, organic radicals, quinones, imides, and so on, for Li and Na batteries,^{204–206,209} investigations in organic electrodes for Ca batteries are so far limited to anodes in aqueous batteries, as previously discussed in section 2.1.2.96,97

Also, sulfur and air cathodes have been suggested for Ca batteries—thus, Ca/S and Ca/air batteries, respectively.^{20,45,103,210} While such cathodes and battery concepts

Table 2. Summary o	of Electrochemic	al Test for C	Table 2. Summary of Electrochemical Test for Ca Cathodes in Either Three- or Two-Electrode Cell Configuration	ree- or Two-]	Electrode (Cell Co	nfiguration
positive electrode	negative electrode	reference electrode	electrolyte	V range	capacity (mAh/g)	cycles	comments
V ₂ O ₅ ^{35,44}	activated carbon	Ag	0.5 M Ca(ClO ₄) ₂ in PC	1 to -0.5	200/465	3/8	(xerogel/PC composite VOx/PC)
$V_2O_5^{42,43}$	Ca	Ag/AgNO ₃	1 M Ca(ClO ₄) in ACN	-0.5 to -1.5	400	0.5	XRD of "intercalated phase" given.
$V_{2}O_{5}^{125}$	Ca	Ca	0.3 M Ca(ClO ₄) ₂ in ACN	0 to 3	200-600	0.5	Capacity attributed to side-reactions involving the electrolyte.
			0.3 M Ca(BF4) ₂ in EC:PC				
			0.3 M Ca(TFSI) ₂ in EC:PC				
			0.3 M Ca(ClO ₄) ₂ in EC:PC				
1 70 1 70			(100 °C) 2 5 M G-(THET) :- HC BC	1 4 - 1 6	0.21	L.	
V2U5	activated carbon	Я	$0.5 \text{ M} \text{ Ca}(1 \text{ FS}1)_2 \text{ m} \text{ EC:} PC$	C.1- 01 I	150	s	water has an influence in overpotential. AKU given.
			water content from $H_2O/Ca = 0$ to 9.5				
$CaV_2O_5^{125}$	Ca	Ca	0.45 M Ca $(BF_4)_2$ in EC:PC (100 °C)	3.6 to 2.6	250	0.5	Capacity attributed to side-reactions involving the electrolyte, enhanced at high T. Attempts of chemical oxidation using NO $_{2}BF_{4}$ also unsuccessful.
$\rm NH_4V_4O_{10}^{227}$	Pt	Ag/AgNO ₃	$\begin{array}{c} 1 \ \mathrm{M} \ \mathrm{Ca}(\mathrm{ClO}_4)_2 \times \mathrm{H}_2 \mathrm{O} \ \mathrm{in} \\ \mathrm{ACN} \end{array}$	1 to -0.2	100	100	Changes in XRD reported, but mechanism not fully elucidated.
$Ca_{0.5}Co_2O_4^{164}$	V_2O_5	ı	0.5 M $Ca(CIO_4)_2$ in ACN	1 to 0	80	30	XPS/XRD to study mechanism but not fully elucidated.
$Ca_3Co_2O_6^{187}$	Ca	Ca	0.45 M Ca(BF ₄) ₂ in EC:PC 100 °C	3.5 to 1	150	0.5	Reversibility very limited, if any.
MoO ₃ ¹⁸⁸	Ca	Ca	$0.5 \text{ M Ca}(\text{TFSI})_2 \text{ in DME}$	2.2 to 0.7	80	12	XPS/XRD/Raman XRD similar to protonated phase.
MoO ₃ ¹⁸⁸	activated carbon		0.1 M $Ca(TFSI)_2$ in ACN	2.8 to 0.8	150	s	
$M_0O_3^{228}$	activated carbon	Ag/AgNO ₃	0.5 M Ca(TFSI) ₂ /AN	1 to -1	120	б	XPS and XRD, but mechanism not elucidated.
$K_{0,31}MnO_2.0.2SH_2O$ (birnessite) ²²⁹	activated carbon	Ag/AgCl	1 M Ca(NO ₃) ₂ 4H ₂ O in water	0.9 to -0.4	150	50	XRD, EDX, possibly proton participation.
${f Mg_{0.19}Na_{0.07}MnO_2^{\cdot}}\ 0.37H_2^{\cdot { m O}}\ ({ m todorokite})^{230}$	Ca and V ₂ O5 (two counterel)	Ag/AgNO ₃	0.2 M Ca(CF ₃ SO ₃) ₂ in ACN	1 to -2	100	10	XAS, but mechanism not fully elucidated.
${ m K_2BaFe(CN)_6}^{231}$	carbon paper	Ag/AgC1	1 M Ca(ClO ₄) ₂ in ACN	0 to 0.8	55.8	100	
$Na_{0.2}MnFe(CN)_6$	$Ca_x Sn$	Ca ²⁺ /Ca	$0.2 \text{ M Ca}(\text{PF})_6 \text{ in EC} + \text{PC}$	0 to 3.5	70	35	
KNiFe(CN)6 ⁵⁴	activated carbon	Ag^{+}/Ag	0.5 M $Ca(TFSI)_2$ in ACN	-1 to 1	60	12	
TiS_{2}^{124}	Ca	Ca	0.45 M Ca(BF ₄) in EC:PC 100 °C	3.5 to 1	48	2	Electrolyte solvent cointercalation, Ca insertion assessed by tomography.
TiS_{2}^{45}	Li	·	0.1 M Ca(CF ₃ SO ₃) ₂ in PC or PC:DMC	3 to 1.5	100	б	Minor changes in XRD, Li present as Li counter electrode.
Mo ₆ Se ₈ ²³²	CaHg ₁₁ –Ca	CaHg ₁₁ –Ca	0.5 M Ca(ClO ₄) ₂ in ACN 25–75 °C	2 to 0.4	3-10	100	
NaFePO ₄ F ¹⁹⁴	carbon	·	0.2 M Ca(PF ₆) ₂ in EC:PC $3:7$	-1.5 to 3	80	50	Changes in XRD reported, but mechanism not fully elucidated.
CaTaN ₂ ²⁰¹	Ca	Ca	0.45 M Ca(BF ₄) ₂ in EC:PC (100 $^{\circ}$ C)	4.5 to 0.5	60	1	Loss of crystallinity upon oxidation.
graphite ⁸²	Sn	1	0.8 M Ca(PF ₆) ₂ in EC:PC:DMC:EMC (2:2:3:3)	3 to 5	70	350	Graphite cathode operating through a redox mechanism based on anion intercalation. XRD consistent with the presence of very crystalline Ca_7Sn_6 after 300 cycles.
S infiltrated mesoporous carbon ⁴⁵	Ca	$Ag/AgCIO_4$	0.5 M Ca $(ClO_4)_2$ in ACN	0.75	600	0.5	Primary system. Nature of reduction products not fully elucidated.
O ₂ ⁶⁴		Ag^{+}/Ag	0.4 M Ca(ClO ₄) ₂ in DMSO				Superoxide and peroxide detected upon reduction by differential electrochemical mass spectroscopy.

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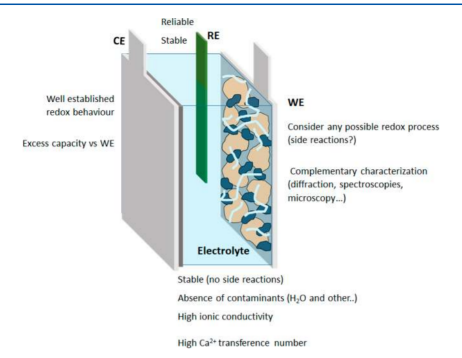


Figure 15. Scheme of a three-electrode Ca metal-anode-based cell setup with key properties to be developed/studied.

hold promise of very low cost and potentially would be an option for large scale energy storage, significant bottlenecks remain to be addressed-many of them covered in the research on Li/S and Li/air batteries.²¹¹ We note the "early" work on Ca/S resulting in rather high-capacity cells, but primary batteries due to an irreversible process upon reduction are still not wellunderstood.,⁴⁵ Nevertheless, the large potential, primarily in terms of cost and energy density, calls for further investigations.²¹⁰ The Ca/air concept is truly challenging and besides early papers discussing the possibility of operating a cell at high temperature,²⁰ there has been limited research efforts to develop systems operating at room temperature using a DMSO-based electrolyte suggested to allow formation of a Ca superoxide.⁶⁴ Some insoluble side-products were obtained but were not fully characterized, while the soluble species seem to reoxidize to oxygen with high Coulombic efficiency despite a high overpotential and with a reversibility decreasing as a function of water content in the electrolyte.⁶⁴ So far, no studies have been reported dealing with operation of an oxygen electrode using electrolytes enabling reversible calcium plating and stripping, despite that some efforts have been reported using ionic-liquidbased electrolytes.⁶³ Overall it is clear that there is a long way to go before any reliable proof-of-concept can be achieved and the true technological prospects are evaluated.¹⁰³

3. EXPERIMENTAL SETUPS, METHODOLOGY DEVELOPMENT, AND FULL-CELL ASSESSMENTS

While the previous sections have approached the Ca battery from a materials perspective, and in many cases, predicted and theoretical performance is quoted, Table 2 summarizes the practical achievements to date for different materials and a wide range of setups. However, before discussing the results in detail, the experimental protocols are to be covered as developing new battery chemistries is far from being trivial and the absence of any reliable standards makes the process tricky. Figure 15 highlights various complexities in the experimental setup of Ca cell studies and construction.

Starting with electrochemical setups, for technologies such as LIBs, new electrode materials are simply tested using a standard electrolyte and vice versa, typically in two-electrode cells with lithium metal as both the counter and reference electrode (a socalled half-cell). For less well-studied systems, the use of a separate reference electrode (through which no current flows) in a three-electrode cell setup is compulsory to be able to independently control and monitor the behavior of the working/counter electrode within the cell and assist in ascertaining the origin of observed features. The choice of the reference electrode is, however, not trivial.¹⁰¹ Silver wires have been often used as pseudo-reference electrodes when testing new cathode materials, but their potential is highly dependent on the anion present in the electrolyte. To address this issue, strategies such as calibration with a well-known standard as Fc⁺/ Fc or use of a reference electrode cell compartment in which the wire is in contact with a specific electrolyte have been used.²¹² While calcium metal would be the first obvious choice as a counter electrode for cathode development, the fact that plating/stripping is only viable under very specific conditions casts severe restrictions on its domain of application. If used in electrolytes not enabling calcium plating, reduction of the working electrode would be feasible (if calcium stripping takes place at the counter electrode), but reoxidation of the working electrode would result in unknown redox processes at the counter electrode, most likely related to electrolyte decomposition, which may affect the electrolyte speciation and the electrochemical response. Instead, activated carbon has often been used (Tables 1 and 2), operating through a capacitive mechanism, but electrode balancing is not trivial and rarely mentioned. Oversizing the capacitive counter electrode, which typically has low capacity, vs the electrode material to be investigated, most often operating through a faradaic redox mechanism and, hence, higher capacity, can be cumbersome in a practical cell. Despite a Sn-Ca alloy being reported as a counter electrode for testing $Na_{0.2}$ MnFe(CN)₆, its capacity was very low, <50 mAh/g, and the electrochemical alloying not fully assessed,

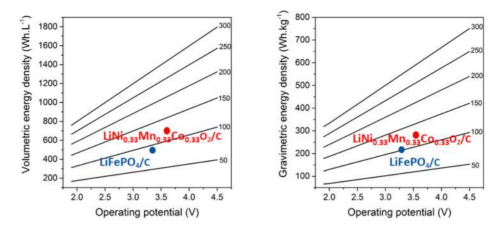


Figure 16. Volumetric and gravimetric energy densities for hypothetical calcium metal-based batteries. The straight lines are calculated energy densities as a function of operation potential and capacities (denoted on the right of each line) of the positive electrode material. All calculations were made using the model developed by Berg et al.²²⁶ Reproduced with permission from ref 225. Copyright 2019 Frontiers.

the electrolyte side-reactions were inferred.¹⁰⁰ Formation of crystalline Ca_7Sn_6 after 300 cycles when using Sn foils as anodes has been reported⁸² but also with electrolyte decomposition products being detected on the electrode surface.

Another factor affecting the electrochemical setup and results is that very few calcium salts are commercially available, and the most commonly used salt is $Ca(ClO_4)_2$. This salt is typically hydrated and also difficult to obtain in anhydrous form due to its explosive nature. This means that many results in the literature are likely obtained in the presence of a non-negligible amount of water in the electrolyte, seldom reported in the experimental sections until very recently, which may result in side-reactions such as H^+/H_3O^+ intercalation or water reduction/oxidation. This in turn can result in some observed "false" electrochemical capacity, as was reported for Mg^{2+} intercalation in V_2O_5 .¹⁰² While the $Ca(TFSI)_2$ and $Ca(CF_3SO_3)_2$ salts also are commercial but much less used, the synthesis of $Ca(PF_6)_2$ was reported only very recently^{83,100} but with purity issues due to a very high tendency of anion hydrolysis. $Ca(NO_3)_2$ is the salt most commonly used in aqueous electrolytes, and again, sidereactions related to H^+/H_3O^+ intercalation probably deserve further investigation, as these have been shown to be an issue for Zn²⁺-based aqueous chemistries,²¹³ and there are no major reasons to believe that the situation would be different for Ca²⁺.

Moving from pure electrochemical setups, the use of advanced materials characterization techniques is often warranted to properly assess and understand the Ca intercalation reactions, in particular to elucidate the real intercalation degree and associated structural changes in cathodes as well the nature and composition of the interfaces created between anodes and electrolytes.

To properly consider the length scale that can be probed, e.g., surface for XPS, bulk for XRD, and atomic for TEM (transmission electron microscopy), is also a must.^{214–218} Moreover, the precision and reliability of each technique also deserves attention as well as if ex situ, in situ, and/or operando setups are to be preferred. For instance, the amount of intercalated Ca ions cannot be assessed from the electrochemical capacity unless the absence of side-reactions can be confirmed, including issues such as simple corrosion of current collectors.²¹⁹ Chemical analysis techniques such as inductively coupled plasma (ICP), atomic absorption (AAS), and energy dispersive (EDS) spectroscopy can possibly be applied, but these methods may all overestimate the degree of Ca²⁺ insertion,

as there might be calcium containing electrolyte salt residues or surface layers resilient to the sample preparation of washing, as shown for Mg²⁺ intercalation studies.²²⁰ In the absence of reliable structural determination of the phases involved in the redox mechanism showing crystal sites and occupation for calcium ions or in the presence of mixed and/or amorphous phases, spectroscopic techniques specifically probing calcium are useful.²²¹ This is especially true if a 3D distribution can be achieved by tomography—a technique which recently enabled to unambiguously detect reversible calcium intercalation in TiS₂.¹²⁴

Also, the polymorphs used can be an issue and not always under control. V₂O₅ has attracted a lot of attention for Ca batteries and despite some studies dealing with samples prepared in the laboratory, including a xerogel composite (Table 2), most studies deal with the commercially available orthorhombic α -V₂O₅, for which a slight increase of the *a* cell parameter was reported in the very first studies of calcium cells.^{42,43} The redox process was then assumed to result in a significant amount of calcium intercalation, but no structural model was provided. The XRD diffraction pattern is, however, different from that of the isostructural α -CaV₂O₅ obtainable by solid-state synthesis^{222,223} and also from the δ -polymorph, for which lower migration barrier was predicted^{129,224} and which seems to be unstable with respect to the α -structure. Instead, there are notable similarities to the patterns observed for some $H_xV_2O_5$ and $H_xV_4O_{10}$ phases, a fact clearly deserving further attention.¹²⁵ Attempts to electrochemically oxidize α -CaV₂O₅ did not result in any change in the XRD pattern, in agreement with the high-energy migration barriers for Ca²⁺ predicted for this polymorph (discussed in section 2.3).^{129,224} Combining all of the above makes us propose that the electrochemical capacity observed likely is related to electrolyte decomposition and possibly proton insertion, at least partially-further stressing the need for combined electrochemical and advanced material analysis.

In line with the above, the electrochemical reduction of MoO_3 in calcium containing electrolytes has been shown to result in changes in the Mo oxidation state, and XRD indicates the formation of a new phase for which the Ca content could not be fully assessed.²²⁵ The authors comment that it exhibits the *Cmcm* space group, a space group previously reported for some H_xMoO_3 phases.^{164,188} Prussian blue analogues (PBAs) present even more complexity, as these materials are known to exhibit variable amounts of water in their crystal structures. This water is sometimes difficult to remove, and also, the versatility of PBA structural frameworks results in difficulties to extract meaningful data of occupation sites and factors from XRD. Yet, given the good rate performance reported to date for intercalation of single-valent ions, especially using aqueous electrolytes,²¹³ further exploration of the mechanisms taking place in divalent systems should be encouraged.

Last but not least, care should be exercised in generalizing results achieved for a specific electrolyte. Besides the possibility of side-reactions occurring due to insufficient (electro-)chemical stability, there are many other intrinsic issues related to the electrolyte composition, e.g., viscosity—affecting both ion transport and wetting of the electrodes, ion—ion and ion—solvent interactions, (de)solvation dynamics, and so on, which also may affect the electrode behavior. Recent reports demonstrate that Ca ions can be extracted from Ca₃Co₂O₆¹⁸⁷ and CaTaN₂.²⁰¹ but the insertion processes are elusive. Here the lack of reversibility could be attributed either to, e.g., a difficult Ca desolvation during discharge or to intrinsic cathode material properties/decomposition reactions.

Despite the numerous challenges remaining with respect to the development of functional Ca cells, the promising results of some chemistries (TiS_2 , $Ca_3Co_2O_6$) call for realistic estimates of (prospective) full-cell figures of merit. To quantify these at the cell level, the energy-cost model developed by Berg et al.²²⁶ was used.²²⁵ Using a scheme of possible operation potentials and specific capacities of some selected existing or virtual materials, the energy densities for a set of different cell configurations were estimated and benchmarked vs other rechargeable battery technologies, including LIBs (Figure 16). Specific LIB technologies are depicted with symbols, while the calcium concepts are represented by straight lines of energy densities as a function of operating voltage and specific capacity of the cathodes. The results indicate that the theoretical energy densities for calcium batteries could easily top the state-of-theart LIBs while most likely being cheaper. Overall, even cells with only moderate operating voltages of 2.1 or 2.5 V and cathode capacities of 250 or 200 mAh \cdot g⁻¹ would yield higher energy densities than the best state-of-the-art LIBs. Moreover, for combinations of 3.0 V/250 mAh·g⁻¹ or 3.5 V/200 mAh·g⁻¹, the volumetric energy densities would be >1000 Wh \cdot L⁻¹, hence higher than any of the (prospective) sulfur-cathode-based battery technologies. The cost-effectiveness for the battery configurations was also estimated, indicating that calcium batteries would be on par with state-of-the art graphite/NMC LIBs even for very high hypothetical calcium cathode material costs (>80 $\cdot kg^{-1}$).

4. CONCLUDING REMARKS

Reviewing the state-of-the-art of rechargeable Ca batteries, we find that most studies focus on developing inorganic intercalation host materials possessing TM ions combined with a variety of simple or complex anions (oxides, sulfides, hexacyanoferrates, etc.), electrolytes with high anodic stability able to reversibly plate/strip Ca metal efficiently, and the anode/ electrolyte interface. Very few, if any, full-cell studies exist, which both assures that the reported capacities truly originate in the redox reactions aimed for and at the same time covers all the aspects we today take for granted for the LIB technology: capacity fading at different C-rates and associated Coulombic efficiencies, calendar/shelf life vs cycle-life assessments, and so on. Indeed, very few studies are able to even name a target application due to the incomplete characteristics at hand which though is natural at this early stage of development. According to energy-cost models, however, the prospects for the Ca battery technology are encouraging, and this is further strengthened by the current technology advances—both on the materials and cell levels. Alternative directions such as Ca/O₂ and Ca/S batteries might be feasible, but must solve the same intrinsic bottlenecks associated with air and sulfur electrodes that are not yet mastered for the corresponding Li, Na, or Mg technologies and might in some aspects be even more difficult for Ca-based batteries, e.g., the very stable CaO potentially being formed.

The battery performance possible to attain certainly depends on the intrinsic material properties. Along this Review we have, however, underlined that the actual observation of electrochemical activity in laboratory cells requires using appropriate testing protocols and cell configurations-not the least when new chemistries are to be explored. The literature offers several examples of "failed" experiments dealing with materials not showing electrochemical activity. This may be due to a diversity of causes, such as bad inherent properties of the electrode materials, issues related to electrolyte degradation, or to difficult Ca desolvation at the interfaces, and therefore exemplifies the need for improved and a variety of experimental setups-not the least complementing the electrochemical by advanced materials analysis. As long as findings are rationalized and rigorously discussed, any result (positive or negative) represents a small step in the long and winding road to the design of new battery technologies, in general, and calcium-based systems, in particular. Finally, we would like to stress that since the current trend of the increasing need of batteries both in sheer amount but also in terms of diversification, devices, and applications, is expected to progress more rapidly in the not-so-distant future, the development of different and complementary battery technologies to the prevailing LIBs of today is the only longterm sustainable strategy. Herein, Ca batteries surely have a role to play.

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Notes

The authors declare no competing financial interest.

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Alexandre Ponrouch is currently a staff researcher at the Institut de Ciència de Materials de Barcelona (ICMAB-CSIC). He received his Ph.D. from the Institut National de la Recherche Scientifique (INRS-EMT, Canada) in 2010 working on electrodeposition of metals, alloys, and oxides for application in fuel cells and supercapacitors. He further joined ICMAB-CSIC to embrace research in battery electrolytes. His current research is mainly focused on developing new electrolytes for multivalent batteries following fundamental electrochemistry approaches.

Patrik Johansson is Full Professor in Physics at Chalmers University of Technology, Gothenburg, Sweden, which he joined in 1999. He has a Ph.D. in Inorganic Chemistry from Uppsala University, Sweden (1998), and he made a postdoctoral stay at Northwestern University. He is currently codirector of ALISTORE-ERI (CNRS FR 3104), Europe's largest industry—academia network within the field of modern batteries. His science combines understanding of new materials, mainly various electrolytes, at the molecular scale with next generation battery concept development and real battery performance—both via experimental and modeling efforts.

M. Rosa Palacín is a research Professor at the Institut de Ciència de Materials de Barcelona (ICMAB-CSIC) and actively involved in ALISTORE-ERI (which she codirected 2010–2017). After a Ph.D. in solid-state chemistry (UAB, 1995) and a postdoctoral stay at LRCS (Amiens, France), her research career at ICMAB-CSIC has been fully focused in rechargeable battery materials with specific emphasis set in the study of electrode materials. She worked on nickel-based and lithium-based technologies to more recently deviate to alternative chemistries such as sodium-ion and also multivalent systems.

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