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Cite this: *Energy Environ. Sci.*, 2018, 11, 2696

Review of electrical energy storage technologies, materials and systems: challenges and prospects for large-scale grid storage

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Increased interest in electrical energy storage is in large part driven by the explosive growth in intermittent renewable sources such as wind and solar as well as the global drive towards decarbonizing the energy economy. However, the existing electrical grid systems in place globally are not equipped to handle mass scale integration of intermittent energy sources without serious disruptions to the grid. It is generally agreed that more than 20% penetration from intermittent renewables can greatly destabilize the grid system. Certainly, large-scale electrical energy storage systems may alleviate many of the inherent inefficiencies and deficiencies in the grid system, and help improve grid reliability, facilitate full integration of intermittent renewable sources, and effectively manage power generation. Electrical energy storage offers two other important advantages. First, it decouples electricity generation from the load or electricity user, thus making it easier to regulate supply and demand. Second, it allows distributed storage opportunities for local grids, or microgrids, which greatly improve grid security, and hence, energy security. Currently, there is only 170 GW of installed storage capacity around the world, but more than 96% is provided by pumped-hydro, which is site-constrained and not available widely. Hence, a battery of technologies is needed to fully address the widely varying needs for large-scale electrical storage. The focus of this article is to provide a comprehensive review of a broad portfolio of electrical energy storage technologies, materials and systems, and present recent advances and progress as well as challenges yet to overcome. The article discusses the status and options for mechanical, thermal, electrochemical, and chemical storage. Where appropriate, it also provides tutorial level background information on fundamental principles for the interested non-expert. It is hoped that this article is of interest to the uninitiated as well as active scientists and engineers engaged in energy storage technologies, with particular focus on large-scale electrical energy storage.

Received 15th May 2018,
Accepted 27th July 2018

DOI: 10.1039/c8ee01419a

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Broader context

Electricity is a ubiquitous, indispensable and effective energy carrier. But nearly 2/3rd of it is produced from fossil fuels resulting in copious CO₂ emissions to the environment. Rapid decarbonization of global electricity infrastructure necessitates shifting from fossil fuels to renewable sources such as solar and wind. Due to their intermittency, however, this is difficult to realize without building sufficient storage capacities to assure grid reliability, quality, availability and security. Indeed, overbuilding solar and wind without adequate storage capacity has recently resulted in discarding electricity with negative pricing in China, Germany and California. Storage also helps reduce reliance on the central grid, while allowing distributed generation and storage for local microgrids. This article presents a comprehensive review of electrochemical, chemical, mechanical, and thermal energy storage systems, materials and technologies. It critically assesses their advantages and shortcomings, and discusses the challenges yet to be overcome. As storage requirements of the electric grid vary widely depending on specific applications, and often do not fully match the performance characteristics of any one particular storage system, the article highlights this mismatch and underscores the need to advance multitude of cost effective, efficient and scalable storage technologies compatible with grid storage requirements.

A. Background

Energy, and in particular electrical energy, is indispensable to maintain our standard of living. Globally, human activity has

consumed 575 Quads (or, 575×10^{15} Btu, or 606.7×10^{18} J) of energy in 2015, and the demand is expected to grow by 28% to 736 Quads (or, 736×10^{15} Btu, or 776.5×10^{18} J) in 2040.¹ As expected, most of this increase will be due to rapidly developing countries that fall outside of OECD (Organization for Economic Cooperation and Development) or called non-OECD, which badly need energy resources to drive their economic growth.

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Table 1 Global installed capacities of major power generation technologies with actual electric power produced in 2016 compared with expected production in 2040. Data compiled from ref. 1

| Power technology | Installed capacity (2016) | | Realized electricity generation (2016) | | Expected electricity generation (2040) | |
|------------------|---------------------------|------------|--|------------|--|------------|
| | Capacity (GW) | % of world | Generation (billion kW h) | % of world | Generation (billion kW h) | % of world |
| World total | 6637.8 | 100 | 23735.2 | 100 | 34049.0 | 100 |
| Renewables | 2112.4 | 31.82 | 5593.8 | 23.57 | 10702.3 | 31.43 |
| Hydro | 1081.5 | 16.29 | 3910.0 | 16.47 | 5677.9 | 16.68 |
| Solar | 278.3 | 4.19 | 275.6 | 1.16 | 1390.3 | 4.08 |
| Wind | 459.8 | 6.92 | 826.2 | 3.48 | 2524.5 | 7.41 |
| Geothermal | 13.8 | 0.21 | 77.1 | 0.32 | 353.4 | 1.04 |
| Other | 279.1 | 4.20 | 504.9 | 2.12 | 756.3 | 2.22 |
| Nuclear | 352.0 | 5.30 | 2510.1 | 10.58 | 3657.3 | 9.79 |
| Fossil fuels | 4173.4 | 62.88 | 15631.3 | 65.86 | 19689.4 | 57.83 |

More than 50% of this increase will be fueled by the needs of India and China.

Furthermore, detailed analyses and reports by reputable institutions such as the US Department of Energy-Energy Information Administration (DOE-EIA), International Energy Agency (IEA), the World Energy Council and others, all indicate that fossil fuels are expected to remain as the dominant primary energy resources for many long decades to come. Currently, coal's share in power generation in the US is over 35%, while it exceeds 70% in India and China. Globally, coal was responsible for 40% of electricity produced in 2016, and is expected to provide more than 30% in 2040.¹ Table 1 provides a comparative summary of the installed capacities for electricity generation with the actual and projected production quantities of major power generation technologies in 2016 and 2040, respectively.¹ In 2040, The International Energy Outlook 2017 report by the US-DOE's Energy Information Agency¹ expects fossil fuels to provide nearly 58% of the globe's electricity need, while renewables will collectively be responsible for more than 31%. These estimates agree in general with the earlier findings of the World Energy Outlook 2015 report by the International

Energy Agency that predicts 54% of the global electricity production will be by fossil fuels and 33% by renewables.²

Ongoing domination of fossil fuels in global electricity production (nearly 2/3rd in 2016) has significantly contributed to atmospheric CO₂ concentration to recently exceed 400 ppmv. Moreover, the amount of CO₂ emissions is expected to increase monotonically between 2015 and 2040, albeit at a somewhat reduced rate of 0.1% per year.¹ Fig. 1 illustrates the partitioning of CO₂ emissions among fossil fuel sources. It should be noted that burning of fossil fuels for power generation is the largest single source responsible for nearly 25% of the global greenhouse gas emissions.³ Fossil fuel-based electricity production, and in particular coal-fired power generation that has long been indispensable for reliable and cheap base-load power, faces significant social, political and environmental pressures to reduce carbon emissions. Increasing conversion efficiencies for fossil fuel power generation by developing advanced technologies such as fuel cells for less-carbon electricity holds the potential of dramatically reducing emissions and collecting CO₂ in capture ready form.^{4–6}

Another strategy that has already gained momentum and market interest in recent years involves shifting power generation from fossil fuels to renewable sources. Despite their impressive growth and deployment in the last decade, however, intermittent renewables still have modest capacities and collectively accounted for slightly more than only 5% of the global electricity production in 2016 (see Table 1). Moreover, the gap



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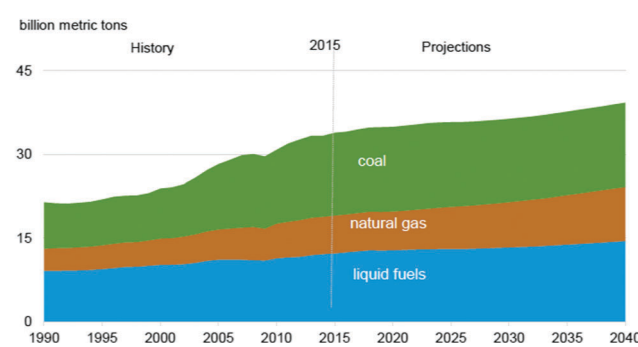


Fig. 1 Energy related global CO₂ emissions in billion metric tons by fuel type (1990–2040).¹

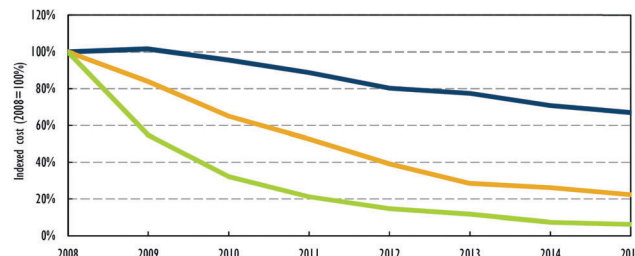


Fig. 2 Dramatic cost reduction in utility scale solar PV and wind power (and Light Emitting Diodes-LEDs) between 2008 and 2015.⁷ Color legend: blue; onshore wind, orange; solar PV-utility scale, green; LED.

between the installed capacities for intermittent sources such as solar and wind, and the actual amounts of electricity they generated clearly highlights the urgent need for utility-scale storage technologies.

An added advantage for most renewables is their modular nature, which provides opportunities for distributed power generation on-location that will help reduce the economic and environmental impact of power transmission and distribution from centralized power plants. Also, major advances in wind and solar technologies brought about major reductions in the cost of electricity (COE) over the last several years. Fig. 2 indicates 5-fold reduction in the cost of solar, while wind has decreased by more than 30% between 2008 and 2015, and recently they have started to compete favorably with coal-fired power in some countries. In fact, several news outlets have recently reported that solar and wind is now at a turning point, and the cost for wind at nearly \$50 per MW h now stands nearly one-half of the price of coal-fired power.⁸ Moreover, recent auctions of solar electricity contracts in India (\$64 per MW h) and Chile (\$29.10 per MW h) have demonstrated unprecedentedly competitive prices.¹⁰ Large-scale energy storage technologies hold the bottleneck in this respect and their rapid development and deployment is critically important in order to realize a clean and sustainable energy future.

A recent International Energy Agency (IEA) report “World Energy Outlook 2016” (WEO 2016) concludes that the pledges made at the 2015 Paris agreement by 195 signatory nations – minus the recent exit by US – only slows down the growth curve for CO₂ emissions but will fail to limit the total emissions to

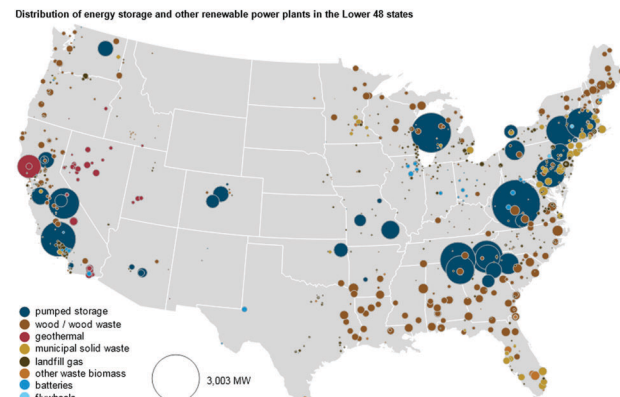


Fig. 4 Geographical distribution of US energy storage capacities by size (i.e., by diameter of circles scaled to 3003 MW as shown in inset) and location. Also included is power generation by renewable sources other than solar, wind, and hydro.¹²

450 ppmv CO₂ equivalents (so-called the 450 scenario).⁹ This is illustrated in Fig. 3. Indeed, the pledges by the signatory countries foresee 37% of power generation from renewables in 2040 (compared to 23% in 2016) and 150 million electric vehicles on the road by 2040 (compared to nearly 1.3 million today). WEO 2016 report also makes it clear that although the 2015 Paris agreement may indeed slow down the rise of greenhouse gas emissions, it falls far short of bending the emissions curve of Fig. 3 downward towards achieving carbon-neutrality (*i.e.*, adding no net carbon to the atmosphere) by 2100.

To limit the global temperature rise to within 2 °C above pre-industrial level by 2100, the WEO 2016 report posits that we need to achieve carbon-neutral energy by the end of this century, whereby renewables need to provide more than 60% of global power generation, and among other additional measures, 715 million electric vehicles need to be deployed on the road by 2040.

Achieving these ambitious targets pose major technological, political, social and financial challenges. As an example, even meeting the Paris Agreement pledges poses political challenges, as the recent departure by the US has demonstrated. Moreover, this report estimates that more than \$40 trillion in cumulative energy investments and an additional \$35 trillion for improvements in energy efficiency will be required to realize the 450 scenario.²

Despite such major hurdles and challenges towards achieving carbon neutrality, however, one thing is quite clear. Renewable energy needs to be a major component of the roadmap towards decarbonizing the energy sector. Indeed, among the energy sources, renewables is expected to grow the fastest with an annual rate of nearly 2.3%, or from 2112.4 GW of installed capacity in 2016, to 3733.8 GW in 2040, and are expected to produce 10702.3 billion kW h or nearly 31% of world's electricity by 2040.¹¹ Although this increase is very significant, it should be noted that much of this will still be provided by hydroelectric power that constitutes the lion's share of renewables, but it is restricted geographically to water-rich parts of the globe, as illustrated in Fig. 4 for energy storage and renewable power generation in US from sources other than wind, solar, and hydro.¹²

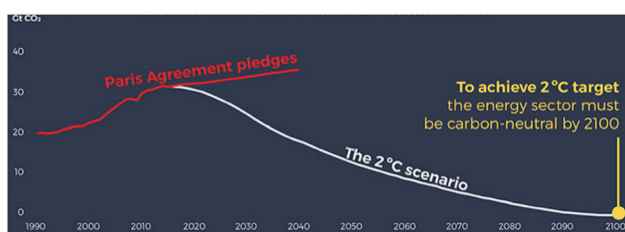


Fig. 3 Global CO₂ emissions under the Paris agreement pledges and the 450 scenario for decarbonization, where the latter assumes a 50% chance of limiting the total greenhouse gases to 450 ppmv CO₂ equiv., and the global temperature rise to less than 2 °C. For this, the energy sector needs to achieve carbon-neutrality by 2100.⁹

B. Introduction

Conventional power generation and distribution infrastructure currently employed globally demands a highly delicate, low error margin, and near-instantaneous balance between electricity supply and demand on the electrical grid system. To realize balance, thousands of turbines all over the world have to be brought online instantly or taken off in order to respond to the variability in electricity demand. Usually, most of these turbines stay idle majority of the time resulting in lost generation capacity. Also, the power generated during off-peak times by these plants is often wasted due to lack of proper storage capacity.

The basic premise of energy storage involves transforming one type of energy into another type that can, when needed, deliver back the stored energy in an efficient, cost-effective, and reliable manner. Similar to the wide diversity of power generation technologies, electrical energy storage technologies and systems show an equally rich diversity. Each offer a distinct set of advantages, but also disadvantages, challenges and shortcomings. As there is no single winning technology for power generation that can meet the wide range of requirements including environmental, cost, performance, availability, customer demand, portability, scalability, *etc.*, there is also no silver bullet to address our impending needs for electrical energy storage. Hence, a multi-prong portfolio strategy is best suited to develop a multitude of storage technologies and systems. Each offers its own strengths and weaknesses, as well as distinctly different technical basis governing its operating principle.

The global excitement, momentum and vast investments in intermittent renewables to grow them as one of our primary sources for electricity production is critical to pave the way towards decarbonizing our currently fossil fuel-dependent energy economy, and in particular, electricity production. But the intermittent nature of many renewables is a major impediment to reliable electricity generation, and for their mass deployment globally. The distributed nature of renewables provides great advantage to help reduce the economic and environmental impact of power transmission from centralized power plants, as is the case currently. Renewables are also important for poor and underdeveloped parts of the world, where large populations still lack electricity and other forms of energy even for their daily chores. In that regard, renewables can help jump-start economic growth and greatly improve quality of life in many of these societies.

However, rapid and utility-scale deployment of intermittent sources such as solar, wind, tide and wave is largely contingent on the availability of cost effective, efficient and scalable energy storage technologies. In fact, electricity storage or lack thereof may well be the “ball-and-chain” for intermittent renewables to break free and achieve the deployment levels foreseen by mid century.^{1,9} Indeed, Bloomberg Technology News reported on July 31, 2017 that Germany had to toss out 4% of wind energy in 2015 and China has discarded 17% of its renewable energy, while California had to give up 300 000 MW h of solar and wind power in the first half of 2017, all due to lack of sufficient electricity storage capacity.¹³ Without commensurate and

Table 2 Global installed energy storage capacities operational in 2016 by type of storage technology. New storage projects under construction are expected to add another 13 GW of storage capacity by 2018¹⁵

| Storage technology | Installed capacity (GW) | % Share |
|---------------------------------|-------------------------|---------|
| 2016 World total | 168.6 | 100 |
| Pumped-hydro | 162.2 | 96.2 |
| Electrochemical | 1.6 | 0.95 |
| Thermal | 3.2 | 1.9 |
| Mechanical | 1.6 | 0.95 |
| New projects under construction | | |
| 2016 | 2 | |
| 2017 | 5 | |
| 2018 | 6 | |

sufficient electrical storage capability, rapid buildup of solar and wind capacity can also be an undue financial burden. Case in point is California, which in the last decade has invested heavily in solar, not only was forced often to curtail solar power production due to power glut but also had to pay exorbitant prices, up to \$25 per MW h, to other states just to get them to take its excess power, *i.e.*, called ‘negative pricing’.¹⁴

Investments in energy storage technologies and projects in the last decade have improved the inventory of global installed capacities. Currently, world’s total installed storage capacity stands at about 169 GW (up from 132 GW in 2012), of which pumped-hydro storage provides more than 96% of the total capacity, with the remaining vested in mechanical, thermal and electrochemical storage technologies.¹⁵ Installed storage capacities and their breakdown are summarized in Table 2, where new capacity additions for a total of 13 GW to be completed by 2018 are also presented. A 2016 report predicts that the global demand for electricity storage capacity will increase to 11.89–15.72 TW h in 2030 from 4.67 TW h in 2017 in order to double the share of renewable energy in the global mix.¹⁶

For renewable energy to achieve its bold target of being a primary energy source by mid century, grid-compatible electricity storage technologies need to satisfy key requirements including, high capacity, rapid charge/discharge rates, high energy density, long cycle life, stable operation and performance, reliability, cost-effectiveness, and easy scale-up. For utility-scale storage of electricity, grid parity is also a critical consideration.

Achieving grid reliability and compatibility, managing peak demand, and integration of intermittent renewable energy sources into the grid all require utility-scale electrical energy storage that delivers electricity where it is needed and when it is needed. Electrical energy storage also decouples electricity generation from the load, or electricity user, thus making it easier to regulate electricity supply and demand. Distributed electrical storage also offers important opportunities to build local grids, or microgrids, which greatly improve grid security, and hence, energy security.

However, the requirements for energy storage are widely varying and diverse, and driven primarily by the type of application. No single technology meets all the requirements and the metrics for large-scale grid quality storage. Each technology

may offer attributes and merits for certain applications, but not for others. Indeed in some markets, electrochemical technologies such as Na-S and Li-ion batteries have become commercially viable even though they do not satisfy all the required metrics. Considering a suitable combination of different storage systems may alleviate such shortcomings towards properly meeting the specific requirements of a particular application.

Furthermore, not all storage technologies are at the same level of development or maturity. A recent report by US Department of Energy indicated a breakdown of the groups of storage technologies either deployed or under development towards large-scale deployment.¹⁷ Broadly, these technologies can be grouped under pumped-hydro, compressed air energy storage (CAES), a wide variety of batteries and other electrochemical systems, flywheels, and superconducting magnetic energy storage (SMES). The developmental status of storage technology groups is depicted in Fig. 5, which categorizes their maturity levels towards full commercialization and mass scale deployment.

In the US, pumped-hydro is the most dominant technology for grid storage, which constitutes nearly 95% of the grid storage capacity.¹⁷ A variety of other technologies including batteries, compressed air, flywheel, and thermal storage are being developed and some have already been utilized for storage operations, albeit at scales considerably smaller than the capacities that are actually needed. For example, 1.5 GW of global installed storage capacity for 2014¹⁸ depicted in Fig. 6 of different mechanical and electrochemical storage technologies makes up only a small fraction of more than 132 GW of global pumped-hydro storage capacity available in 2012. Despite major progress and effort, the cumulative share of electrochemical, thermal and mechanical storage capacities has remained below 4% of the total global storage capacity by 2016, where pumped-hydro provided the bulk capacity of nearly 96%. These numbers, however, still fall short of the combined storage needs for wind and solar (see Table 1).

While pumped-hydro is a mature and widely used technology for utility scale commercial storage where available, CAES, which relies on storing energy as compressed air in underground caverns and formations, is not as mature and fully developed

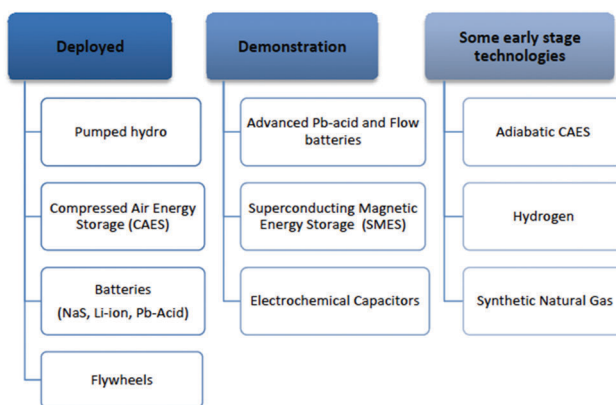


Fig. 5 Developmental maturity of major electricity storage technologies towards large-scale deployment.¹⁷

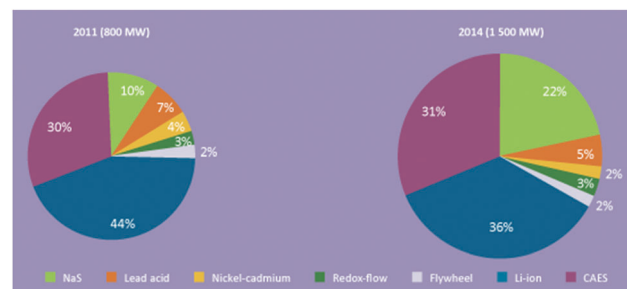


Fig. 6 Comparative breakdown of global installed capacities based on electrochemical and mechanical storage technologies between 2011 and 2014 (CAES: compressed air energy storage).¹⁸

as pumped-hydro. However, both have very specific geographical requirements that render their use site-specific.

It is expected that basic research focused on prospective technologies may expand and improve their merits and lower some of the technical barriers to better meet the required metrics for storage. This is identified as one of the strategies by US Department of Energy to drive down the cost, as investments in improving the technology and large-scale deployment will help lower costs.¹⁶ This is depicted in Fig. 7. However, cost alone does not guarantee success. Mass scale deployment of these grid-scale storage technologies also requires confidence buy-in from both the stakeholders and users such as utilities and customers as well as utility regulators for the safety, performance and reliability of these technologies.

A variety of electrical storage technologies and approaches can be organized into four general categories, namely, mechanical, chemical, electrochemical, and electrical. Many of these categories are reviewed in the next sections, and characteristic features for some of them are presented in Fig. 8, which shows the power ratings and typical discharge times of many of the electrochemical and mechanical electricity storage systems under development or in practical use.¹⁹ They exhibit a wide range of discharge characteristics and power ratings. The discharge times and system sizes may be used as helpful tools to select

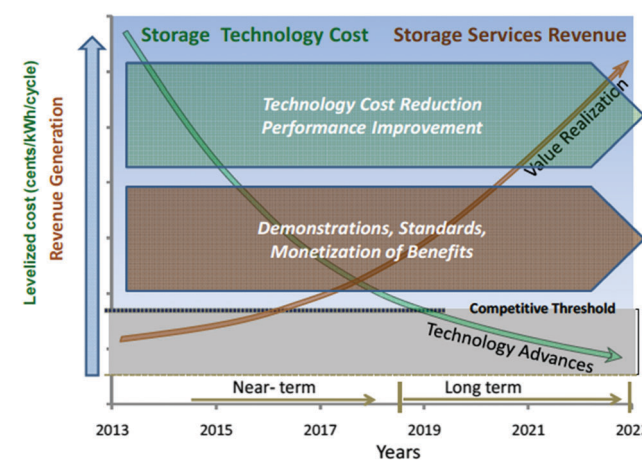


Fig. 7 US DOE's expected cost reduction timeline with the evolution and market penetration of storage technologies.¹⁷

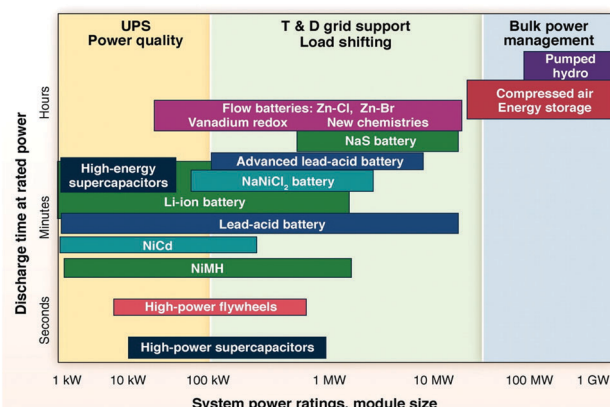


Fig. 8 Comparison of power ratings and discharge time scales of various electrical energy storage systems. From ref. 19, B. Dunn, H. Kamath and J. M. Tarascon, Electrical energy storage for the grid: a battery of choices, *Science*, 2011, **334**, 928–935. Reprinted with permission from AAAS.

suitable storage systems for a particular application. For example, grid management requires frequency regulation and load levelling, which usually demand short response times of the order of seconds that are suitable for flywheel and supercapacitor storage, while bulk power management may require longer response times of the order of hours where pumped-hydro and compressed air storage offer attractive opportunities.

However, the options for large-scale storage shown at the far right column in Fig. 8 are quite slim, and require augmentation of the storage portfolio by developing alternative technologies. Only pumped-hydro and CAES systems are commercially available for bulk storage at utility scale. The maturity levels of more advanced technologies, such as electrochemical storage, are still under active development and further require demonstration and deployment at utility scale. Moreover, the utility industry, which relies on low cost, scalability, and reliability, is typically averse to risk-taking. So they are understandably skeptical about many of these technologies due to cost, scale-up issues, and lack of track record on safe and reliable operation for prolonged use at industrial scale.

There have been excellent reviews of electrical energy storage technologies and materials in the literature. Some of the more recent ones are given in ref. 19 and 21–28. The interested reader is encouraged to seek these papers to gain full appreciation of the wide scientific footprint of energy storage and to expand the scope into more specialized areas. In large part, these review articles focus primarily on a single or select few storage technologies, such as electrochemical energy storage, or a sub-division, such as Li-ion batteries.

Instead, this article distinguishes itself by taking a broad brush and presenting an in-depth review of a wide spectrum of electrical energy storage technologies, materials and systems with an eye for large-scale electrical grid storage. The article provides a comprehensive discussion of the status of four major classes of electricity storage technologies including electrochemical, chemical, mechanical, and thermal storage systems covering a wide range of options from pumped-hydro and

flywheels, to hydrogen and ammonia, to supercapacitors, batteries, regenerative fuel cells, to flow batteries, and phase change materials storage. Where appropriate, the article also offers tutorial information for the interested reader to gain better insight and appreciation of the fundamental considerations and challenges in electrical energy storage. It is hoped that this review will be of interest and benefit to the uninitiated as well as active researchers and engineers engaged in electrical energy storage technologies.

C. Fundamentals of energy storage

The first law of thermodynamics (or, the law of conservation of energy) states that in a closed system, the total energy is fixed and energy can neither be created nor destroyed. It can only be converted from one form to another. This fundamental concept serves as the foundation of nearly all forms of energy conversion and storage. Accordingly, most storage technologies fall into four generalized categories, namely, mechanical energy storage, chemical energy storage, electrochemical energy storage, and electrical energy storage.

The maximum amount of electrical work that can be extracted from a storage system is given by,

$$G = H - TS \quad (1)$$

here, G is Gibbs free energy, H is enthalpy, T is temperature, and S is entropy. In other words, G represents the maximum energy available to do either mechanical or electrical work.

Mechanical energy storage, such as pumped-hydro, manifests itself by potential, E_{pot} , and/or kinetic, E_{kin} , energies that can be represented by,

$$E_{\text{pot}} = fd \quad (2)$$

$$E_{\text{kin}} = (1/2)mv^2 \quad (3)$$

here, f denotes force, d is distance, m is mass, and v denotes velocity. For a rotating body such as a flywheel or a wind turbine, the kinetic energy is described in a similar expression by,

$$E_{\text{kin}} = (1/2)I\omega^2 \quad (4)$$

where, I denotes the moment of inertia and ω is angular velocity of rotating system. In other words, stored energy increases with the system's inertia and the square of its angular velocity. The moment of inertia for a body of mass m , radius r , length l and density ρ can be given by

$$I = (1/2)mr^2, \text{ or } = (1/2)\rho l\pi r^4 \quad (5)$$

Accordingly, high-density materials with large radius help store more energy. For mechanical energy storage in flywheels, eqn (5) can also be expressed as,

$$E_{\text{kin}} = \sigma_m s / \rho \quad (6)$$

here, σ_m denotes maximum stress, s is a shape factor, and ρ is materials density. For modern flywheels made from reinforced high strength carbon fibers, it is possible to achieve storage capacities greater than 200 kJ per kilogram of flywheel mass.

Compressed air storage in salt caverns or underground aquifers relies on the gas law ($PV = nRT$), and the available work, w , is given by PdV integrated over the incremental volume change, dV .

Thermal storage relies on the amount of heat, q , one can store in a medium or material of defined volume, V , materials density, ρ , and specific heat, C_p , resulting a temperature rise of ΔT . This can be expressed by,

$$q = \rho C_p V \Delta T \quad (7)$$

In the case that the thermal energy storage medium or material undergoes a phase transformation during this process, the latent heat associated with this transformation should also be accounted for in the total energy stored.

Chemical storage relies on storing energy in the chemical bonds of fuels, which are inherently strong and thus provide high energy densities. However, one needs to use usually electrical or thermal energy to produce the fuels for chemical storage. The governing equation for the maximum work out of chemical storage is given by eqn (1) above.

Electrochemical storage systems are based on charge storage at the electrodes in the case of batteries, or at the electrochemical interface in the case of electrochemical capacitors or supercapacitors, or in the chemical bonds of fuels as in fuel cells. As the theoretical efficiency for electrochemical energy conversion is defined by the ratio of ($\Delta G/\Delta H$), electrochemical energy storage offers inherently high efficiencies.

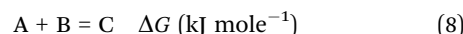
Finally, electrical and magnetic energy storage can also be realized. Examples include dielectric-based capacitors, magnetic materials, and superconducting magnets. However, these types of energy storage approaches lie outside the scope of this article, and are not covered here.

D. Electrochemical energy storage systems and materials

The underlying operating principle for electrochemical and photoelectrochemical devices involves three major process steps, namely, separation (or, ionization) of charge, transport of charged species, and recombining of charge. This basic principle governs the operation of a wide range of devices including batteries, fuel cells, supercapacitors, electrolytic capacitors, photoelectrochemical and photovoltaic devices. Their operating principles, however, involve different mechanisms where interfaces play a critical role. For example, while batteries store charge within the electrodes, in the case of fuel cells and flow batteries charge is stored in the fuel, which is fed externally on to the surface of the electrodes. Supercapacitors, on the other hand, store charge either in the electric double layer at the electrode/electrolyte interface, or as pseudocapacitance governed by surface redox reactions. Some of these electrochemical devices such as PEMFC, supercapacitors and batteries operate at room or ambient temperature, while others require elevated temperatures, *e.g.*, sodium–sulfur batteries or molten carbonate and solid oxide fuel cells. In other words, materials requirements

for each electrochemical storage system are different, and many have been reviewed in detail elsewhere.^{19,20,29–35} Furthermore, the fundamentals and operating principles of batteries, fuel cells and supercapacitors have been presented as a tutorial in a recent review article.³⁶

The fundamental concept behind electrochemical energy storage is the reciprocity between converting the chemical energy stored in the bonds of fuels into electrical energy, and expending electrical energy to synthesize chemicals or fuels by operating in the reverse direction. The driving force for this conversion is the Gibbs free energy change, ΔG , of the electrically neutral species at the electrodes participating in the chemical reaction,



This free energy change is the same if the reactants A and B were to undergo purely a chemical reaction as in eqn (8), or an electrochemical reaction where the reaction involves the transport of ions and electrons across the cell. As these species are electrically charged, the electrostatic energy transported across by a mole of such species is given by zEF , where F is Faraday's constant, z is the charge number of the transporting species, and E is the cell voltage. Under open circuit conditions, the cell voltage is related to the Gibbs free energy change by,

$$\Delta G = -zEF \quad (9)$$

In other words, the chemical potential difference between neutral species at the electrodes defines the underlying force driving an electrochemical cell *via* a chemical reaction among electrically neutral reactants, A and B, to form an electrically neutral product, C.

For electrochemical storage, the systems require basic components that have critically important roles to achieve this, namely, two specialized bodies of electrodes separated by an ionically conducting but electronically insulating “electrolyte”. The electrodes are preferably chosen from abundant and cost-effective materials that exhibit good electronic conductivity, good stability, and high catalytic activity. Some applications require that the electrode material also exhibit mixed ionic electronic conductivity, where both ionic and electronic defects are mobile.

The electrode where the reduction reaction takes place is called the cathode, and where the oxidation reaction takes place is called the anode. An electrochemical energy storage and conversion system, be it a battery, a fuel cell, or an electrochemical capacitor, can schematically be illustrated in Fig. 9, which shows storage of electrical energy in the form of chemical energy, and conversion of chemical energy for reaction (8) back into electrical energy that can be supplied to the grid to perform useful electrical work. Fig. 9(a) represents an electrolysis cell to produce fuel B using external power (*i.e.*, storage mode), while Fig. 9(b) depicts a fuel cell that employs fuel B and oxidant A to generate electricity (*i.e.*, power production mode). Electrolysis cells exclusively operate for the purpose of electricity storage in the form of chemical energy in fuels, while regenerative solid oxide fuel cells offer the opportunity to operate in a bi-directional

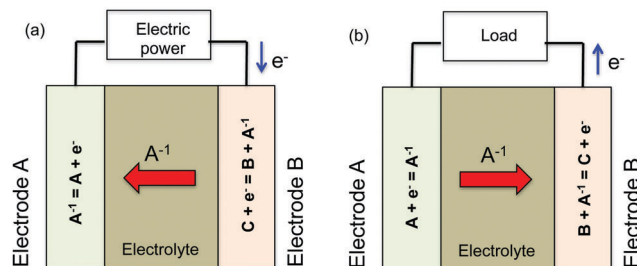


Fig. 9 Basic operating principle of electrochemical energy storage, illustrating (a) electrical energy to chemical energy conversion, and (b) chemical energy back to electrical conversion for reaction $A + B = C$.

manner (electricity to chemicals fuels, and chemical fuels back to electricity in the same fuel cell).

It should be noted that in the case of batteries, electrolytic capacitors and supercapacitors, specific energy and specific power are intimately coupled to each other by the battery chemistry and choice of battery materials. However, in fuel cells and flow batteries the energy density dictated by the choice of fuel is decoupled from power, which is dictated by the kinetic and transport properties of cell components. Such decoupling provides important advantages for many applications. Also important are the volumetric energy and power densities of electrical storage systems. Volume is especially important for transportation, space and mobile applications. Interestingly, there is an ongoing debate whether Li-ion batteries or fuel cells will eventually dominate mobile applications and in particular, which one will power the transportation sector.³⁷

D.1. Batteries

One of the most notable technological achievements of electrochemistry in recent years is the successful commercialization of Li-ion batteries that had a profound impact on the widespread use and deployment of portable electronic devices, and more recently, on transportation and stationary storage at the 100–200 kW h scale. In more general terms however, batteries, the first electrochemical device (*i.e.*, the Volta pile) built more than two centuries ago, have been commercially available for more than a century. They are efficient devices to store electrical energy, and hence used extensively for a wide range of applications including transportation, portable, mobile, as well as stationary back up power. Types of commercial battery systems available today or under development employ a diverse range of battery materials²³ and chemistries both organic³⁸ and inorganic^{21,33} in nature. Operationally, all battery systems can be grouped under primary (*i.e.*, non-rechargeable) and secondary (or, rechargeable) batteries. Primary batteries fall outside the scope of this review article, as these are in general not suitable for storage purposes.

Comparative operational characteristics of batteries are typically represented by Ragone diagrams. This is illustrated in Fig. 10, where specific power is plotted against specific energy for major rechargeable battery systems that compare various classes of batteries to those based on lithium chemistry.^{31,39–42} Superior properties of Li-based batteries to other alternative

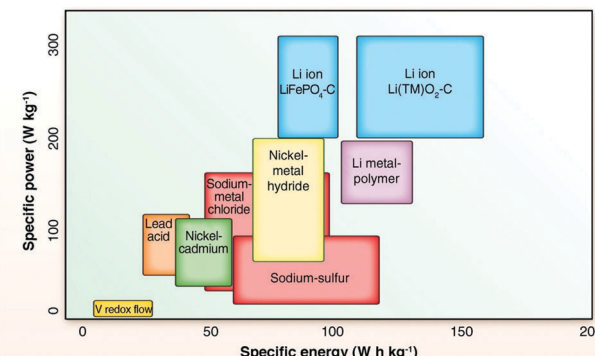


Fig. 10 Ragone diagram of specific energy versus specific power for major rechargeable battery systems. From ref. 19, B. Dunn, H. Kamath and J. M. Tarascon, Electrical energy storage for the grid: a battery of choices, *Science*, 2011, **334**, 928–935. Reprinted with permission from AAAS.

chemistries presented in Fig. 10 make it easy to understand the enormous academic and commercial interest in this system.

Batteries in general offer desirable solutions for cost-effective and compact storage, pollution-free operation with no moving parts, high overall efficiency, sufficient cycle life, considerable shelf and service life. They provide opportunities for distributed storage, as well as storage and frequency regulation from intermittent renewable sources. Several battery systems including redox flow batteries, alkali metal ion-based batteries, sodium sulfur batteries are under consideration and further development for utility scale storage purposes ultimately for the ‘smart grid’.^{19,32} Unfortunately, many of these systems are still too expensive for commercial deployment for large-scale storage applications, and require not only cost reductions but also major improvements on performance, reliability, and materials availability and stability.^{33,34} However, their growing market penetration is expected to lower battery prices.

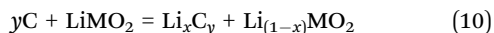
All batteries provide the convenience and portability of delivering stored chemical energy in the form of electrical energy at the point of use, with high conversion efficiency and without harmful emissions to the environment. The amount of stored electrical energy and the rate at which the energy can be extracted from a battery is indicated by its specific energy and specific power, respectively, and is a function of the cell voltage (V) and capacity (A h kg⁻¹) of a battery. These parameters are directly governed by the thermodynamic and kinetic constraints imposed by the battery’s chemistry and choice of materials for battery components. In other words, the amount of the electroactive species that can be stored in the electrode materials dictate the energy storage capacity of the battery, while the rates of interfacial charge transfer reactions and mass transport through the battery components govern its output power. It should be noted here that some of these concepts are in direct competition in terms of battery life and safety. As high storage capacity is related to cell potential, there has been a strong desire to develop electrode materials that yield a high cell potential. But this comes at the expense of battery safety, as high cell potentials are related to high activity that in turn facilitates reaction with the electrolyte and can lead to thermal runaway and fires.

This is particularly true in the case of Li-ion batteries employing organic electrolytes.

The driving force for improving the properties of batteries and battery materials is to achieve low cost, rechargeability, high voltage, safety, and fast charge/discharge capability. The desired battery system not only should exhibit high energy density (*i.e.*, specific energy) but also has high power density (*i.e.*, specific power). Compared to others, Li-Based storage systems clearly offer higher specific power and higher specific energy than others as indicated in Fig. 10.^{19,43,44} However, not all applications require high energy density simultaneously with high power density. This interplay may provide opportunities for other battery systems to offer suitable solutions for specific storage applications.

Among the systems shown in Fig. 10, Li-based battery systems have deservedly attracted the biggest commercial and research interest with excellent review articles reporting on the advances and challenges.^{21,31,39,43,45–48} Li-Based batteries currently dominate nearly 2/3rd of the global battery market due to their high energy density, portability, acceptable cycle life, and their compact size that provides flexibility in product design, especially for portable electronics and communication devices.

D.1.1. Li-ion batteries (LIB). As the energy density of a battery is directly related to its working potential, or, voltage, while its power is proportional to the square of its voltage, the search for better systems and materials has often been driven by high voltage considerations. In this regard, battery systems based on Li are attractive because Li is the most electropositive element (Li/Li⁺ redox potential = −3.04 V vs. SHE) in the periodic table and the lightest metal (6.94 g mol^{−1}), thus offering a high energy density. Indeed, the theoretical energy density of Li-ion batteries is about 380 W h kg^{−1}, with commercially available Li-ion based rechargeable batteries providing 150–210 W h kg^{−1}.⁴⁴ The power densities vary in the range 500–2000 W kg^{−1}, with efficiencies around 90%. As the Li⁺ ion also has a small ionic radius (90 pm), it allows fast diffusion rates through the electrode material, offering fast charge–discharge rates. Because of these practically attractive properties, Li-based battery systems have gained strong attention in research and development as well as commercial activity. Indeed, Sony Corporation was the first to introduce the Li-ion batteries based on intercalation chemistry into the market in 1991. These commercial Li-ion batteries typically contain a graphite anode (the negative electrode), a lithium metal oxide (LiMO₂, where M is Co and/or Ni) cathode (positive electrode), and a Li⁺ ion conducting electrolyte consisting of an organic solvent, typically ethylene carbonate, dimethyl carbonate, and their mixture, in which a soluble lithium salt such as lithium hexafluorophosphate (LiPF₆) is dissolved. The corresponding cell reaction provides a cell potential of around 3.7 V and is given by,



here, the typical values of x and y are 0.5 and 6, respectively. The reaction involves the insertion (*i.e.*, lithiation) and extraction (*i.e.*, delithiation) of lithium ions between the two electrodes,

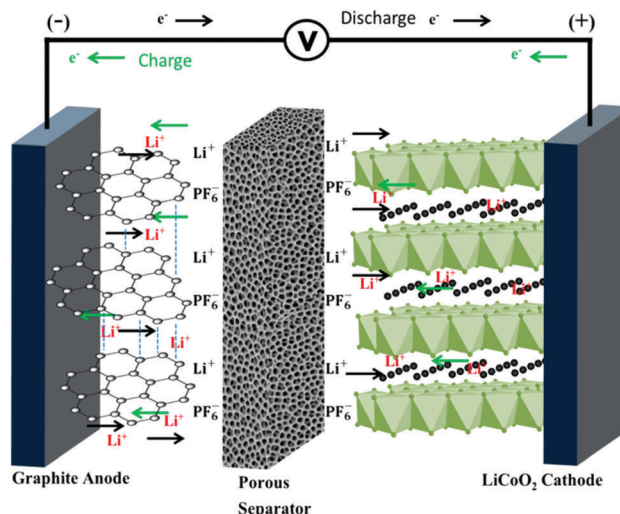


Fig. 11 Schematic illustrating the operating principle of the first commercial Li-ion battery, where the Li⁺ is intercalated (and, deintercalated) in the layered structures of the graphite negative electrode and the LiCoO₂ positive electrode.²¹ Reprinted with permission from (K. M. Abraham, Prospects and limits of energy storage in batteries, *J. Phys. Chem. Lett.*, 2015, **6**, 830–844). Copyright (2018) American Chemical Society.

while the electrons flow through the external circuit performing useful electrical work.

Both the C₆Li anode and the LiCoO₂ cathode materials have layered crystal structures so involve the intercalation (or, lithiation) and deintercalation (delithiation) of Li⁺ ions in between the structural layers of the anode and cathode materials. Battery operating principle and the charge and discharge cycles *via* transport through the electrolyte are depicted in Fig. 11. During the charging process, lithium is extracted from the positive electrode, whose working potential is higher than 2 V vs. Li⁺/Li, and intercalated into the graphite anode, whose working potential is lower than 3 V vs. Li⁺/Li. As Li is shuttled between the two electrodes, these rechargeable cells are sometimes referred to as “rocking-chair” batteries.⁴⁹

There have been excellent reviews of Li-based batteries and systems discussing the chemistry and electrochemistry, materials properties as well as the search for better electrode materials, performance and safety issues, and economics for large-scale storage applications.^{19,39,47,48,50–53} Most Li-batteries that are commercially available employ nonaqueous electrolytes,⁵⁴ although other types of electrolytes including ionic liquids,^{55,56} polymers,^{57,58} ceramics,⁵⁹ and even aqueous electrolytes^{60,61} have also been investigated. Much progress has been achieved in Li-battery technology in the last decades. The capacity of the first commercial Li-ion batteries, *i.e.*, the 18650 cell, was about 900 mA h, while the same size Li-batteries today deliver a much higher practical capacity of 2600 mA h.

D.1.1.1. Negative electrode materials. Although most commercial Li-ion batteries employ a graphite anode, which has a layered structure, the candidate materials under development for the negative electrode also include elemental Li, Li-alloys, or a nanostructured host material such Si-nanowire.^{61,62} Each offers its own set of merits and challenges.

Elemental lithium is the ideal desired anode material as Li is the most electropositive element in the periodic table and has an exceptionally high specific capacity of 3860 mA h g^{-1} . Unfortunately, elemental Li is quite reactive and the chemical stability of the battery electrolyte against Li poses serious concerns regarding safety, reliability and durability.⁶⁴ Also, the exothermic reactions between the Li anode and the battery electrolytes may lead to overheating and “thermal runaway”. More importantly, elemental Li anodes tend to grow dendrites during charge–discharge cycling due to the nonuniformity of the Li deposition process that tend to occur at preferential sites. The dendrites eventually penetrate through the electrolyte towards the cathode and may cause electrical shorting and other serious hazard problems, even explosion.

Several approaches have been pursued to reduce the driving force for chemical reactivity and dendrite growth. One is based on the thermodynamic concept of lowering the activity of Li, and hence decreasing its propensity to react and also to grow dendrites, by alloying it with other metals including aluminum,⁶⁵ antimony,⁶⁶ bismuth,⁶⁷ and especially those that alloy with lithium at low potentials such silicon,^{68,69} and tin.^{70,71} Interestingly, the first rechargeable Li-battery was developed in 1970’s by Exxon and employed a LiAl alloy negative electrode and a TiS_2 cathode.⁷²

Recent attention on other remedial strategies such as alloying or conversion chemistries has provided promising results. For example, Si-based alloy anodes provide a theoretical charging capacity around 4000 mA h g^{-1} , and volumetric energy densities in the range $700\text{--}972 \text{ W h L}^{-1}$, which are 1.5–1.8 times higher than for commercial cells, have already been demonstrated.⁷³ These and other electrode chemistries are reviewed elsewhere.⁴³ Although Si exhibits one of the highest capacities among others, its molar volume change during cycling can be as high as 400% depending on the microstructure, which of course can lead to rapid cell degradation and loss of capacity and performance.

It was reported that employing Si nanowires,^{69,74} and nano-size Sn anodes⁷¹ improved mechanical durability and cycle life. Indeed, to mitigate problems associated with dendrites and improve cell performance, micro- and nanostructured negative electrodes have been widely explored and reviewed elsewhere.^{75–79} However, a key challenge with nanostructured electrodes is to maintain mechanical integrity and electronic connectivity. A recent study employed electronically conducting polymer (PFFOMB) as a binder to both stabilize the Si nanoparticles mechanically and to serve as a current collector, which boost up the performance at 1C rate to 2050 mA h g^{-1} for Si and 1360 mA h g^{-1} of the electrode.⁸⁰

Lithium alloy anodes are essentially mixed ionic-electronic conductors of both electrons and Li^+ ions. This is a desirable property for reversible electrodes. Many of these alloys also offer fast ionic transport for Li^+ chemical diffusion in the alloy reaching values up to $2 \times 10^{-3} \text{ cm}^2 \text{ s}^{-1}$ as reported earlier for Li_3Sb at 360°C .⁶⁷ Alloying especially with tin or silicon also helps increase the anode capacity greatly. In fact, the theoretical specific capacities for Sn and Si are 994 mA h g^{-1} and

4200 mA h g^{-1} , respectively. The latter figure is about an order of magnitude higher than that for graphite (372 mA h g^{-1}). Lithium forms several alloy phases with Si and the highest Li content is reported for the $\text{Li}_{22}\text{Si}_5$ phase.^{63,68} In other words, 4.4 Li^+ ions can be inserted for every Si atom. This is a substantial increase in anode capacity when compared to the C_6Li stoichiometry for graphite-host anodes, which allow only 0.167 Li^+ ions for every C atom. However, this advantage comes at the cost of significantly increased volume expansion ($\sim 400\%$ for Si) during Li^+ insertion or lithiation of the Sn and Si anodes, which upon cycling eventually leads to mechanical breakdown and fracturing or pulverization of the anode structure. Using nanostructured anodes such as nanotubes instead of bulk Sn or Si greatly mitigates this problem and allows significant increase in charge–discharge cycle life.^{69,77} Besides nanostructuring, studies that pursue the holy grail of using elemental Li as the anode material have also employed interface engineering. For example, coating the Li anode with a monolayer of interconnected carbon nanospheres helped avoid dendrite formation up to a practically significant current density of 1 mA cm^{-2} and maintained 99% coulombic efficiency for more than 150 cycles.⁸¹ Also, a thin oxide layer on the Si nanostructure is shown to help improve the mechanical integrity of the anode.⁸²

Although much attention is given to achieving high gravimetric capacity and long cycle life, for most battery applications, volumetric capacity is equally important, and poses one of the barriers for practical implementation of Si-based anode hosts for Li-ion batteries. Intentionally designed void spacing allowed among the nanostructured Si anode architectural features to accommodate for the large volume change during cycling adversely impacts the volumetric capacity. Depositing SiC-free multilayer (2–10) graphene on the Si anode surface resulted in a 2D layered architecture and provided volumetric energy density of 972 W h L^{-1} in the first cycle and 700 W h L^{-1} at the 200th cycle.⁷³ These numbers represent 1.8 to 1.5 times higher values than the volumetric energy densities of currently available commercial Li-ion batteries.

Another strategy commonly practiced is to reduce the propensity for lithium dendrite growth by intercalating Li^+ ions into a suitable host structure, such as in between the graphitic sheets of a typically disordered “soft” carbon anode, as depicted in Fig. 11. Carbon exhibits a slightly less negative potential than Li (*i.e.*, nominally $100 \text{ mV vs. Li/Li}^+$). It also has a volumetric capacity of about 800 mA h ml^{-1} and hence takes up $\sim 50\%$ of the volume of commercial Li-batteries.⁷⁷ As lithium in the carbon host structure is always in ionic form, rather than in metallic form, this provides some advantage against dendrite formation and related safety and electrolyte stability issues. However, the proximity of the carbon potential to that of Li does not entirely eliminate the possibility of dendrite formation and Li plating during charging. Chemical and physical properties of carbons and carbon nanostructures as negative electrode hosts for Li-ion batteries have been reviewed elsewhere.^{78,79,83}

D.1.1.2. Positive electrode materials. Most of the strategies that aim to minimize or overcome the stability and dendrite

growth problems give rise to an increase in the potential of the negative electrode with respect to Li and correspondingly cause a reduction in the net cell voltage. In order to compensate for this voltage loss, cathode materials that exhibit high half-cell potentials and high capacities for insertion or intercalation have been the subject of intense research. Accordingly, more recent developments focused on the positive electrode, which is considered as the critical component of the battery that impacts its performance the most.^{43,45-48}

Many oxide systems have been investigated for prospective cathode materials over the last four decades, but most of them can be grouped into three different crystal structures, namely layered, spinel and olivine structures. The layered and spinel structure cathodes have the general formulas LiMO_2 ($\text{M} = \text{Co, Ni, Mn}$) and LiM_2O_4 ($\text{M} = \text{Mn, Ni}$), respectively. These materials are good conductors for Li^+ ions, and also have sufficient electronic conductivity due to the electronic transitions between the oxidation states of the multi-valent transition metal cations present in the structure. Furthermore, their electronic conductivities can be improved by appropriate doping.⁸⁴ For example, partial substitution of Mg^{2+} on the Co^{3+} -site that is compensated by the creation of electron holes in LiCoO_2 and improve the room temperature electronic conductivity by nearly two orders of magnitude to 0.5 S cm^{-1} .⁸⁵ Similarly, the conductivity of the electronically insulating ($<10^{-13} \text{ S cm}^{-1}$) spinel $\text{Li}_4\text{Ti}_5\text{O}_{12}$ can be improved many orders of magnitude to $10^{-2} \text{ S cm}^{-1}$ by Mg^{2+} doping on the Li^+ site that is compensated by the reduction of Ti^{4+} to Ti^{3+} .⁸⁶ Accordingly, such advances enabled the use of these structures suitable for cathode materials.

Indeed, the first commercial Li-ion battery employed the layered structure oxide LiCoO_2 as the positive electrode.⁴³ However, LiCoO_2 has an inherently low capacity of about 130 mA h g^{-1} . It is also expensive and poses safety problems due to the toxicity of cobalt, which makes up only 20 ppm of the Earth's crust. Better safety, higher capacity, more abundant and lower cost cathode materials are needed for consumer electronics and energy storage applications.

In this regard, olivine structure LiMPO_4 where M is a transition metal such as Fe offers lower cost, long cycle life, and safety.^{46,47} From a safety point of view, the strong P-O covalent bond in the structure prevents the release of oxygen that can react with the electrolyte and initiate a thermal runaway. The strong P-O bond also reduces the covalency of the Fe-O bond and improves Li intercalation as well as cell voltage (3.5 V vs. Li). Moreover, Fe is cheap and non-toxic, and offers a gravimetric capacity of 170 mA h g^{-1} . There has been a flurry of studies on olivine phosphates since the early results of electrochemical activity with a capacity of $100\text{--}110 \text{ mA h g}^{-1}$ reported for LiFePO_4 .⁸⁷ However, the olivine structure LiFePO_4 is electronically insulating ($10^{-10}\text{--}10^{-9} \text{ S cm}^{-1}$), and the Li extraction is accompanied by Fe^{2+} transition to Fe^{3+} , which results in two separate phases, namely LiFePO_4 and FePO_4 , which are both poor electronic conductors. To overcome this difficulty, use of fine size LiFePO_4 particles or mixing it with an electronic conductor such as carbon has become necessary for cathode fabrication.

Indeed, the rates of Li insertion and extraction reactions were improved in subsequent studies by incorporating carbon-based conductive additives⁸⁸ and redox mediators.⁸⁹ Remarkably, it was shown that a small (0.5%) of Nb^{5+} doping ($\text{Li}_{1-x}\text{M}_x^{z+}\text{FePO}_4$) helps increase the electronic conductivity by eight orders of magnitude.⁹⁰ Introducing disorder in the Li-sublattice by nanosizing and nanostructuring,²³ also improves reaction rates and battery performance. A recent study concluded that the desired size of battery-grade LiFePO_4 particles should be $<200 \text{ nm}$.⁴⁶ Indeed, nanosized LiFePO_4 based Li-ion batteries have recently gained practical interest and commercialized originally by the US company A123 for portable applications.

The effect of doping LiMPO_4 on the M and/or P sites as well as the properties of other prospective materials for positive electrodes including dichalcogenides, manganese oxide spinels, and silicates are extensively reviewed elsewhere.^{45,47,48,91,92} The cell potentials and storage capacities of various positive electrode materials of different crystal structures versus the graphite anode are presented in Fig. 12, where the voltage stability regime of the 1 M LiPF_6 /ethylene carbonate electrolyte is also indicated. For many of these materials, however, the useable energy density of the Li-ion battery is much smaller than the theoretically expected value. For example, in the case of electric vehicles the practically achievable energy density is approximately 25% of theoretical expected value for the active cathode material.⁹³ Unrealized differences between the theoretical storage capabilities and the actually achieved values are presented in Table 3 for various commercially available Li-ion cells. This gap is partly due to the dead weight of the inactive battery components. But part of this unused capacity is related to the structural properties of the cathode and the structural transformations the cathode undergoes during the lithiation and delithiation reactions

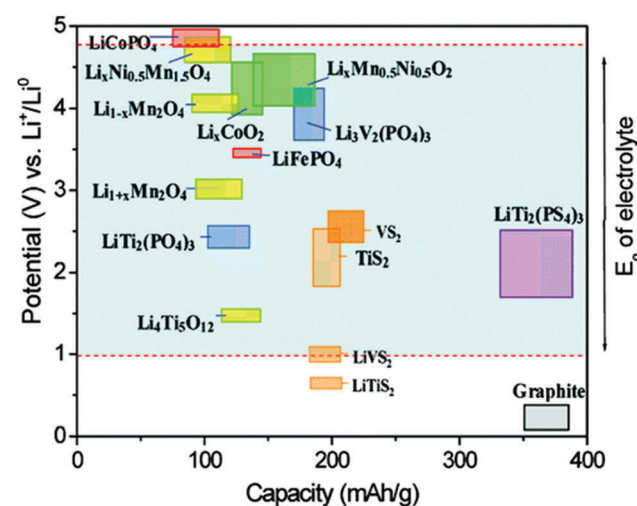
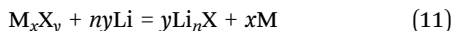


Fig. 12 Comparative relation between capacity and cell potential of Li-ion batteries with various positive electrode materials (cathodes) versus the graphite anode with respect to the stability window (red dotted line) of the electrolyte (1 M LiPF_6 dissolved in 1:1 mixture of ethylene carbonate: diethylene carbonate).⁴⁸ Reprinted with permission from (J. B. Goodenough and Y. Kim, Challenges for rechargeable Li batteries, *Chem. Mater.*, 2010, **22**, 587–603). Copyright (2018) American Chemical Society.

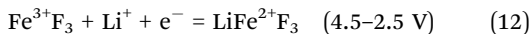
during battery operation.⁹⁴ Such low utilization of the inherent capacities depicted in the table highlights the need to greatly improve the reversibility of the lithiation and delithiation reactions at the electrodes.

For large-scale energy storage, or power applications such as electric vehicles much higher capacities and energy densities, with lower cost and better safety than offered by today's Li-ion batteries are desired. Many believe that about $\sim 300 \text{ mA h g}^{-1}$ may approach the upper limit of the capacity for intercalation chemistry⁵⁰ due to constraints posed by the crystal structure and the limited transitions allowed in the oxidation state of the transition metal in these layered structures. A more recent review article agrees with this general assessment and estimates the ultimate limit for intercalation chemistry for Li-batteries may be 0.5 kW h kg^{-1} and 1.5 W h L^{-1} for full cell configurations.⁴⁶ Therefore, intercalation chemistry most likely may not provide the roadmap to achieve much higher energy densities and capacities at lower cost with improved safety.

The search for improving the specific capacity of Li-ion batteries by utilizing all the accessible oxidation states of the electrode materials has lead to the concept of using conversion reactions, represented generally by the reaction,



here, M is a 3-d transition metal such as Co, Fe, Mn, Cu, and X is N, O, S, F. Indeed, nano-sized transition metal oxide based positive electrodes for Li-ion batteries gave electrochemical capacities up to 700 mA h g^{-1} with good cycling capability retaining nearly 100% up to 100 cycles.⁹⁵ Similarly, transition metal fluorides, and in particular FeF_3 , which offer high operating voltages due their highly ionic character, and large theoretical capacities (e.g., 712 mA h g^{-1} for FeF_3) gained much attention for positive electrodes.^{96,97} However, metal fluorides are electronically insulating materials, thus unsuitable for use as electrodes. But carbon composites of FeF_3 overcame this problem and gave encouraging results, where reversible capacity as high as 600 mA h g^{-1} of the composite was reported.^{96,98} The reactions at the positive electrode were proposed^{96,98} to proceed by the two-step sequence,



The theoretical capacity for reaction (13), where Li is inserted into the FeF_3 structure in the potential regime of 4.5–2.5 V is 237 mA h g^{-1} , while the capacity for reaction (13) in the 2.5–1.5 V potential region is 400 mA h g^{-1} . It should be noted

that the electrochemical lithiation of FeF_3 proceeds first with an intercalation step and multiple phase transitions, then followed by conversion process.⁹⁷ Binary metal fluorides FeF_2 and CuF_2 were also investigated for Li-battery positive electrode materials, and concluded that the lithiation of MF_2 occurs via a three-phase conversion process, where FeF_2 displays high reversibility for lithium via a network of Fe embedded in the insulating FeF_2 matrix, while CuF_2 shows lack of such reversibility.⁹⁹

D.1.1.3. Solid electrolyte interphase (SEI). The organic carbonate-based solvent used in the nonaqueous electrolyte helps the formation of a thin solid electrolyte interface (SEI) on the anode. The SEI layer first reported nearly four decades ago, forms on the anode even after the initial charging of the fresh battery.¹⁰⁰ This layer contains both organic and inorganic components that are blocking to the transport of both Li^+ ions and electrons. In the case of a graphite-based anode with ethylene carbonate/ LiPF_6 nonaqueous electrolyte, the SEI layer may contain grains of Li_2O , LiCO_3 and LiF . As Li_2O , Li_2CO_3 , and LiF are insulators for both Li ions and electrons, they block the transport of these species, impeding the anode reaction. Indeed, probing the electrolyte/electrode interface of ethylene carbonate/metal oxide system by high resolution X-ray reflectivity, a recent study reported interfacial layering of the Li-salt and electrolyte solvent molecules on the electrode surface, which increases the interfacial transport impedance.¹⁰¹

There is limited understanding of SEI formation at the cathode. Unlike the graphitic carbon anode held together by van der Waals forces that allow co-intercalation of the electrolyte solvent molecules, the ionically bonding character of the cathode crystal structure does not allow this to happen. Also, the potentials for most cathode materials with the exception of the emerging “5 V class” of cathodes such as $\text{LiNi}_{0.5}\text{Mn}_{1.5}\text{O}_4$ (4.6 V vs. Li/Li^+) and LiCoPO_4 (4.8 V vs. Li/Li^+) generally lie within the stability limit of the nonaqueous electrolytes avoiding the possibility of electrolyte decomposition.⁵⁴ Although the SEI at the anode may initially exhibit higher impedance, the cathode interphase over prolonged cycling builds up faster and becomes the more dominant resistive component of the two SEIs.

Despite its disadvantages, the SEI layer also offers important advantages. For example, the SEI layer that forms on the bulk Li anode, in the case of Li-metal battery, provides a protective passivation layer on the Li surface that prevents violent reaction of Li with the electrolyte. In other words, the SEI layer at the anode makes it possible to operate the Li-ion battery at potentials beyond the thermodynamic stability regime of the

Table 3 Theoretical and actual storage capabilities of Li-ion batteries for various positive electrode materials.⁴⁶ Theoretical values include only the active components of the battery

| Cathode chemistry | Battery size | Theor., W h l^{-1} | Actual, W h l^{-1} | % Realized | Theor., W h kg^{-1} | Actual, W h kg^{-1} | % Realized |
|--------------------------------|--------------|-----------------------------|-----------------------------|------------|------------------------------|------------------------------|------------|
| LiFePO_4 | 54 208 | 1980 | 292 | 14.8 | 587 | 156 | 26.6 |
| LiFePO_4 | 16 650 | 1980 | 223 | 11.3 | 587 | 113 | 19.3 |
| LiMn_2O_4 | 26 700 | 2060 | 296 | 14.4 | 500 | 109 | 21.8 |
| LiCoO_4 | 18 650 | 2950 | 570 | 19.3 | 1000 | 250 | 25.0 |
| Sr-LiMO ₄ panasonic | — | 2950 | 919 | 31.2 | 1000 | 252 | 25.2 |

organic electrolyte. The SEI layer also helps improve the kinetic stability of the cell by stabilizing the Li insertion and delithiation reactions during subsequent charge–discharge cycles, and provides a reversible capacity of 370 mA h g⁻¹ for the graphite anode. However, it should be noted that the nature of mobile species in the SEI layer is of critical importance. If the SEI layer exhibits mixed ionic electronic conduction properties where both Li ions and electrons are mobile, then the layer will continue to grow, whereas if it is ionically conducting but electronically insulating in nature, then it will stop growing after forming a chemically protective barrier on the electrode/electrolyte interface to prevent further reaction with the electrolyte.

Also, the SEI layer, undergoes constant structural changes and reformation during repeated charge–discharge cycles, which leads to loss of Li to the constant repair of the SEI layer. Moreover, repeated volume changes at the anode, even in the case of bulk lithium anode as lithium is added or inserted, and extracted from the electrode lead to mechanical failure, particle fracturing and separation, and concerns about battery safety. The details of the properties, mechanisms of formation, and the structure of SEI layers have been reviewed elsewhere.^{54,102}

D.1.1.4. Electrolytes for Li-batteries. Other approaches to address the battery safety, stability, and dendrite problems focused on modification of the battery electrolyte or search for improvements and/or new alternatives. A wide variety of electrolytes employed in Li-based battery research and development have recently been reviewed.^{103,104} The common electrolyte employed in Li-ion batteries is a non-aqueous electrolyte composed of lithium hexafluorophosphate (LiPF₆) dissolved in an organic solvent, typically ethylene carbonate or dimethyl carbonate. But organic-based electrolytes pose stability problems in the presence of high Li activities, *i.e.*, high working potentials. This was a common failure of high voltage batteries with Li-metal anodes, which vigorously react with the organic components of the electrolyte irrespective of whether the electrolyte is a liquid or polymer-based. Similar reactions can also occur at the positive electrode, as the organic component usually is not stable in the presence of high lithium activities. These exothermic reactions cause local heating in the battery that may eventually lead to thermal runaway. To avoid this problem, it is important to avoid operation at the potential where lithium at the electrode reacts with the organic component. Repeated cycling of the battery exacerbates this problem, possibly due to morphological changes that increase reaction surface area. A related problem is due to oxygen evolution or build up to a flammable mixture with the highly reducing organic component in the electrolyte and causing battery fires. This is generally related to the positive electrode material that can exhibit a high oxygen activity *versus* Li,¹⁰⁵ and facilitates oxygen evolution at high potentials that can lead to battery fires. Recently, several notable incidents caught the public's attention, ranging from aircraft battery fires to smartphones catching fire.

Efforts to modify liquid non-aqueous electrolytes by incorporating appropriate additives as inhibitors against dendrite growth as well as flame retardant chemicals for battery safety

have provided encouraging results.^{106,107} Even trace amount of water added to the nonaqueous electrolyte is reported to help eliminate dendrite formation.¹⁰⁸ Similarly, the use of Li⁺-ion conducting polymeric membranes,^{39,109,110} polymer-ceramic composite electrolytes¹¹¹ or impervious Li⁺-ion conducting ceramic electrolytes^{112–115} provides a physical barrier layer that impedes dendrite growth between the two electrodes. Alternatively, ionic liquids,^{56,116,117} which are essentially room temperature molten salts, have been considered as electrolyte solvents for their high stability with a wide electrochemical window typically in excess of 4 V,¹¹⁸ as well as for their inherent safety due to inflammability and low vapor pressure.⁵⁵ However, they are expensive, and have limited solubility for Li-salts, high viscosity, and generally moderate ionic conductivity at ambient temperatures. Usually, ionic liquids are also susceptible to moisture, which complicates processing and battery fabrication.

Alternatively, aqueous electrolytes offer high conductivity and are highly desirable for safety and cost considerations for large-scale energy storage,¹¹⁹ but water has a narrow stability window of only 1.23 V. However, it is well known that several aqueous battery systems such as lead-acid (~2.0 V) can clearly operate outside this potential regime. This is facilitated by the formation of electronically blocking but ionically conducting interlayers on the electrodes. In the case of Pb-acid battery, for example, the PbSO₄ interlayer covers both electrodes. Hence, modifying the electrolyte to promote the formation of such layers opens up opportunities for battery safety and cost. Indeed, a recent study reported a 4.0 V aqueous Li-ion battery where a fluorinated additive that is immiscible with the aqueous electrolyte provides a hydrophobic layer that minimizes the competing reaction of water reduction at the anode, and allows reversible cycling of both the graphite and Li-metal anodes.⁶¹

Primarily for reasons of safety, stability, battery performance, and cost of manufacturing especially in thin film¹²⁰ or flexible forms,^{121,122} there has long been a search for high conductivity ceramic and polymeric electrolytes for solid state Li-batteries.¹¹³ Many of the polymeric electrolytes are based on polyether or polyethylene oxide (POE) providing the framework into which an appropriate Li salt is dissolved. Similarly, inorganic electrolytes based on various oxides, silicates, glasses and other structures have also been investigated and reviewed.^{103,104,114} A more recent review provided a large compilation of comparative ionic conductivity data for a wide variety of Li-ion conducting polymer and ceramic electrolytes for Li-batteries.^{103,112,123} Fig. 13 presents a subset of Li-ion conducting solid polymer- and ceramic-based electrolytes from among a large collection of compositions. It is clear that most of these electrolytes are limited by their relatively poor room temperature conductivities typically in the range of 10⁻²–10⁻⁶ S cm⁻¹. This is especially true for most polymer and ceramic electrolytes whose conductivities are typically two or three orders of magnitude lower than for liquid organic electrolytes.

There is growing interest towards an all solid-state battery architecture that employs inorganic solid electrolytes, which are expected to exhibit sufficient chemical stability against high voltage cathodes, and potentially mitigate or eliminate

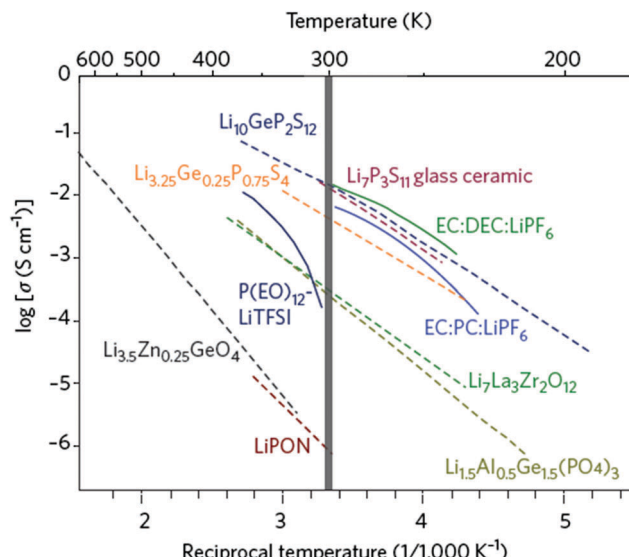


Fig. 13 Comparative Li^+ ion conductivities of ceramic, polymeric, and liquid electrolytes for Li-ion batteries.¹²³ Reprinted by permission from Nature/Springer, J. Janek and W. G. Zeier, A solid future for battery development, *Nat. Energy*, 2016, **1**, 16141. Copyright (2018).

safety concerns.¹²³ Indeed, a lithium-metal anode cell employing a polymer electrolyte of lithium (bisfluorosulfonyl)imide in polyether is recently reported as a promising candidate for an all solid-state lithium-metal battery.¹²⁴ Thermodynamically, however, most solid electrolytes with few exceptions including garnets lack sufficient stability against metallic Li. This necessitates the build up of SEI protective layers at the electrode/electrolyte interfaces. Nevertheless, the presence of a Li-ion conducting inorganic impervious electrolyte membrane would eliminate the dendrite problem, and associated safety concerns of electrical shorting and thermal runaway. On the other hand, Li-batteries with organic electrolytes typically exhibit nearly twice the storage capacities ($\sim 0.2 \text{ mA h cm}^{-2}$) and discharge rates than batteries with solid electrolytes ($\sim 0.1 \text{ mA h cm}^{-2}$) due to kinetic and mass transport limitations.¹²⁵

Recent advances on sulphide-based and garnet-based solid electrolytes, have shown significant improvements in ionic conductivities. For example, $\text{Li}_{10}\text{GeP}_2\text{S}_{12}$ was shown to exhibit an ionic conductivity of 0.012 S cm^{-1} at room temperature,¹²⁶ which is only an order of magnitude less than for organic electrolytes. More recently, an even higher ionic conductivity of 0.025 S cm^{-1} was reported for $\text{Li}_{0.54}\text{Si}_{1.74}\text{P}_{1.44}\text{S}_{11.7}\text{Cl}_{0.3}\text{S}_{12}$ at room temperature.¹²⁷

Garnet-based ceramics with cubic structure and the nominal composition of $\text{Li}_7\text{La}_3\text{Zr}_2\text{O}_{12}$ (LLZO) also exhibit high Li-ion conductivity and excellent stability against lithium metal, but stabilizing the cubic structure at room temperature is challenging. Room temperature conductivity of the tetragonal garnet LLZO is typically $\sim 10^{-6} \text{ S cm}^{-1}$, nearly two orders of magnitude lower than the cubic phase. It was reported that generation of Li vacancies by extrinsic doping with aliovalent cations disrupts ordering of the Li-sublattice, and this may present opportunities to stabilize the cubic structure and improve room

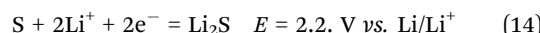
temperature Li-ion conductivity.¹²⁸ Incorporating ceramics into polymer electrolytes also improves ionic conductivity, expands the stability limit of the polymer electrolyte and also mitigates the dendrite problem.¹²⁹ For example, adding 15 wt% of $\text{Li}_{70.33}\text{La}_{0.557}\text{TiO}_3$ nanowires to polyacrylonitrile- LiClO_4 electrolyte has substantially improved the room temperature Li-ion conductivity to 0.024 S cm^{-1} .¹³⁰ However, typically low transference numbers for Li ions ($t_{\text{Li}} = 0.38$) in composite electrolytes need significant improvement to attain higher efficiency.

The prospects of identifying highly conductive and stable electrolyte compositions in the near future require further investigation and development. However, it was recently suggested that the challenge to materials design for Li-batteries is not necessarily to find the stable materials, but instead to search for materials whose instabilities are self-limiting.⁹⁴

D.1.2. Beyond Li-ion batteries. Although Li-ion batteries get most of the attention, their theoretical specific energy for the $\text{LiCoO}_2/\text{graphite}$ and $\text{LiFePO}_4/\text{graphite}$ systems is limited by the insertion chemistry to around 400 W h kg^{-1} .^{42,90} As a case in point, a battery power pack that can provide a cell-level specific energy of $350\text{--}400 \text{ W h kg}^{-1}$ would be required for electric vehicles to have a driving range of 300 miles on a single charge. Unfortunately, Li-ion batteries currently available today can only provide a specific energy nearly half of this value. Also, the desired life of the battery should last more than 1000 charge-discharge cycles at a rate greater than 2C capable of providing power greater than 600 W kg^{-1} .¹³¹

D.1.2.1. Li-Sulfur batteries. The demand for high performance batteries for transportation as well as stationary applications requires electrode materials that offer much higher specific energies, which can be achieved by increasing the cell potential and the storage capacity of the materials used for the anode and the cathode. But the stability windows of the battery electrolytes limit the cathode operating voltage to below 4.3 V.⁴⁸ Furthermore, with the capacity limited by insertion chemistries, Li-sulfur system offers many advantages, but also challenges.^{132,133} Fig. 14 presents a schematic diagram for the Li-S battery system illustrating the charge and discharge processes.

The use of sulfur for the positive electrode in Li-S batteries offers low cost, low weight, high rates, low toxicity, environmental friendliness, and high theoretical capacity (2600 W h kg^{-1} , *versus* $\sim 600 \text{ W h kg}^{-1}$ for current Li-ion batteries), which is due to the high specific capacity of sulfur (1675 mA h g^{-1} , *versus* $130\text{--}200 \text{ mA h g}^{-1}$ for current Li-ion battery cathode materials). The net cathode reaction during discharge is based on 2-electron transfer by,



Although the cell potential of 2.2 V is about 60% of the 3.7 V for conventional Li-ion batteries, this is more than made up by the high theoretical energy densities of 2600 W h kg^{-1} and 2200 W h L^{-1} ,⁴³ which are 4–5 times higher than for the graphite/ LiCoO_2 system. Potentially, Li-S batteries may be able to provide a driving range of nearly 300 miles.¹³¹ But the Li-S battery system also poses several important challenges.¹³⁴

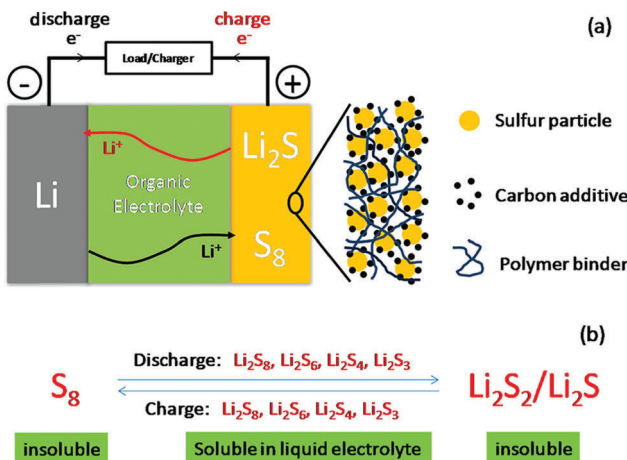


Fig. 14 Schematic depiction of in the Li–S battery architecture illustrating charge and discharge processes.¹³³ Reprinted with permission from (A. Manthiram, Y. Fu, S.-H. Chung and Y.-S. Su, Challenges and prospects of lithium–sulfur batteries, *Acc. Chem. Res.*, 2013, **46**, 1125–1134). Copyright (2018) American Chemical Society.

One is the poor electronic conductivities of sulfur (10^{-17} S cm $^{-1}$) and its discharge products of various polysulfides Li_2S_x ($x = 1\text{--}8$) deposited on the electrodes. These passivation layers increase the cell impedance and therefore require addition of an electronic conductor such as carbon or metals. Naturally, the addition of the conducting phase lowers the cell capacity, energy density, round-trip efficiency and cycle life. Another challenge is posed by the loss of sulfur from the cathode into the electrolyte due to the high solubility of the polysulfide ions (S_x^{2-}) formed on reduction of S_8 or oxidation of Li_2S . The dissolved polysulfide ions diffuse to the negative electrode and react with Li in a parasitic reaction to form lower polysulfides, which then diffuse back to the positive electrode and are re-oxidized. As expected, this internal shuttling mechanism so-called “polysulfide shuttle” lowers the Coulombic efficiency of the Li–S battery as well as the poor recyclability of the battery to achieve long cycle life. This is one of the key issues that can limit the commercial viability and prospects of the Li–S battery technology. In essence, the high order polysulfides dissolved in the electrolyte migrate toward the anode and react with the Li-metal anode and are reduced to the lower order sulfides, which migrate back to the cathode to form higher order polysulfides. Such continuous shuttling of polysulfides between the two electrodes not only leads to permanent capacity fading due to loss of sulfur, but also constitutes the underlying cause of self-discharge. This self-discharge occurs even at the resting state of the Li–S battery due to the continuous dissolution of polysulfides, which gradually lower the open circuit potential and discharge capacity of the battery.

Furthermore, the multiple conversion reactions between the polysulfide species bounded by the end members S and Li_2S are accompanied by large volume changes (up to 76%), which can lead to mechanical failure or disintegration of the electrodes, and eventually to capacity loss upon cycling. Furthermore, dendrite formation and stability issues posed by the Li-metal

anode and discussed in detail in the previous section are also valid for the Li–S system.

Much attention has been given to mitigate or overcome these technical challenges. Various strategies have been pursued to address and improve Coulombic efficiency, discharge capacity, cyclability, and self-discharge in Li–S batteries. Many of these strategies including carbon-composite or nanostructured electrodes,¹³⁵ trapping interlayers, selective coatings or solid state electrolyte layers on the Li-metal anode, and modification of electrolytes by additives have been investigated and recently reviewed.^{132,133,136} Much progress has been made. For example, nanostructured cathodes where sulfur is encapsulated inside hollow carbon nanofibers improved the cell capacity to 730 mA h g $^{-1}$ measured at C/5 rate after 150 charge–discharge cycles.¹³⁵ Similarly, sulfur-graphene oxide nanocomposite cathode modified by cetyltrimethyl ammonium bromide provided a high specific capacity of 800 mA h g $^{-1}$, a long cycle (1500) life with a remarkably low decay rate of 0.039% per cycle, and allowed discharge rates as high as 6C (1C = 1.675 A g-sulfur $^{-1}$) and charging rates as high as 3C.¹³¹ Furthermore, the addition of LiNO_3 to the electrolyte facilitates passivation of the Li-anode and prevents parasitic reactions with the polysulfides. Several studies employing LiNO_3 additives reported Coulombic efficiencies of 96–99% even after 1500 cycles.^{131,135} These advances provide encouragement for the future prospects of Li–S batteries as a potential candidate for efficient and cost effective storage system. However, for the Li–S system to compete potentially for the transportation sector, a recent techno-economic study considering a materials-to-systems approach predicted that high electrode loadings > 8 mA h cm $^{-2}$ are necessary to achieve > 300 W h L $^{-1}$ and $< \$125$ per kW h.¹³⁷ Currently demonstrated loading levels (~ 2.5 mA h cm $^{-2}$) and energy densities (~ 100 W h L $^{-1}$) indicate the major challenges yet to be overcome for the Li–S system to vie for transportation applications.

D.1.2.2. Non-Li-based batteries. Power modules based on Li-ion batteries have been the primary impetus for the explosive growth of portable electronic devices as well as the re-advent of electric vehicles in the last decade, albeit after a hiatus of more than one century. However, long-term sustainability of energy solutions requires a careful examination of the feasibility, cost, and availability of resources. This is especially important if Li-based batteries are to be considered for utility scale energy storage systems for the electric grid.

The relative abundance of lithium in the earth’s crust is only 20 mg kg $^{-1}$ (or, ppm). Furthermore, known lithium reserves are mostly concentrated in South Africa, Chile and Australia, which create source dependency, supply reliability and price volatility for Li-based battery technologies. In fact, large fluctuations observed in the past in the price of lithium also pose economic concerns for reliability and availability when projected into the future even only for the electrification needs of the transportation sector. Recycling lithium from Li-based batteries in a cost effective manner remains an unsolved technology challenge. Moreover, the use of lithium for large-scale stationary energy storage may diminish the known lithium resources.

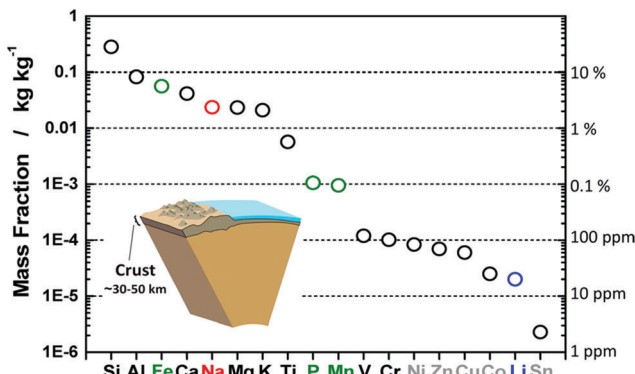


Fig. 15 Relative abundance of metals in the earth's crust.¹³⁸ Reprinted with permission from (N. Yabuuchi, K. Kubota, M. Dahbi and S. Komaba, *Chem. Rev.*, 2014, **114**, 11636–11682). Copyright (2018) American Chemical Society.

By contrast, other electropositive metals such as Na, K, Ca, and Mg are quite abundant in the earth's crust as indicated in Fig. 15, and they are also fairly uniformly distributed geographically. In fact, Na is the fifth most abundant element. Especially oceans provide a vast and almost infinite resource for these elements. So battery systems based on Na, K, Ca, and Mg chemistries can provide cost-effective alternatives to existing Li-based batteries. Also, Na is the second lightest element next to Li in the periodic table. The reduction potentials and metal-anode-based theoretical capacities of alkali metal and alkaline earth metals are provided in Fig. 16. Although the theoretical capacity for lithium is clearly superior to Na and K, the abundance and availability of these metals and others including Ca and Mg make them attractive candidates for alternative battery systems as discussed in recent reviews.^{138–144}

As the active materials make up nearly 50% of the cost of most batteries, the wide availability and low cost of sodium at ~\$150 per ton compared to ~\$15 000 per ton for lithium presents great economic incentives for sustainability. Hence,

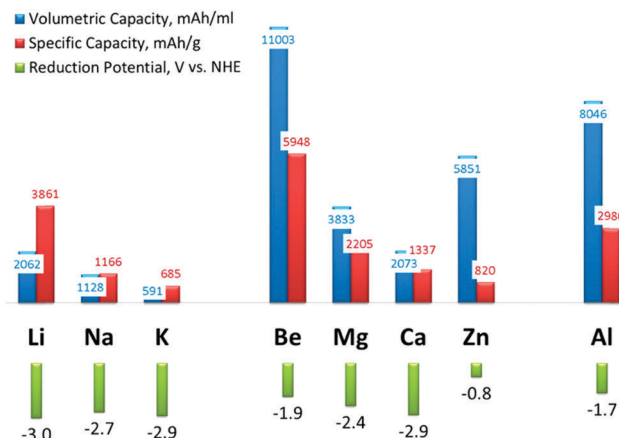


Fig. 16 Comparison of specific capacities and reduction potentials of prospective metals for battery anodes.¹³⁹ Reprinted with permission from (J. Muldoon, C. B. Bucur, T. Gregory, Quest for nonaqueous multivalent secondary batteries: magnesium and beyond, *Chem. Rev.*, 2014, **114**, 11683–11720). Copyright (2018) American Chemical Society.

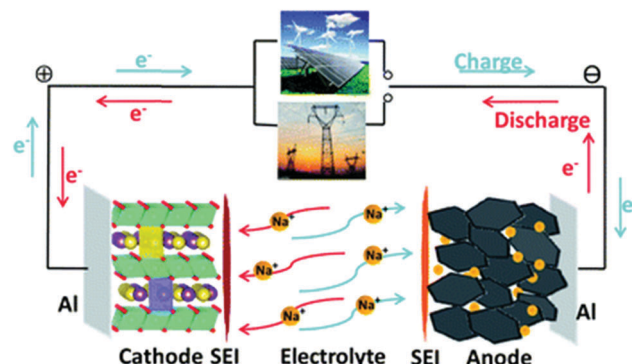


Fig. 17 Schematic depicting the operating principle of the Na-ion battery. Reproduced from ref. 142 with permission from The Royal Society of Chemistry.

sodium-based batteries attracted much attention. Na-Ion battery architecture, chemistry and materials share many similarities with the Li-ion battery system.¹⁴⁵ Fig. 17 illustrates the schematic depiction of the Na-ion battery. Because of such similarities in cell materials and their challenges, detailed discussion of Na-ion battery materials is not provided here. Instead, only salient features, recent progress and challenges are discussed briefly. For detailed information, the interested reader is referred to recent reviews on various aspects of Na-ion batteries.^{138,140,142,143,146}

Interest in the Na-ion system as a cheap and viable alternative to Li-based systems goes back to 1970's, almost concurrent with the intense research and development efforts in Li-ion batteries. Despite the similarities in most of their chemical and physical properties, however, progress has been slow for Na-ion batteries. One stumbling block (or barrier) has been the unsuitability of soft carbon-based anode materials such as graphite, which is commonly employed also in Li-ion batteries. Unfortunately soft carbon anodes exhibit low or no electrochemical activity when used in Na-ion batteries. Heating graphite in Na vapor seemed to be the only way to achieve intercalation. A major breakthrough came in 2000 when Na-ion cells employing hard carbon at the negative electrode were reported to provide a reversible capacity of 300 mA h g^{-1} .¹⁴⁷ The hard carbon anodes were prepared by the pyrolysis of glucose at 1000–1150 °C in an inert atmosphere to form randomly stacked carbon layers made of small aromatic fragments that resemble a “house of cards”, in between which the Na (and also Li) ions can be inserted. This high capacity, comparable to the anode capacities for the Li-ion batteries, generated great impetus and interest in Na-based storage systems research.

Na exhibits a slightly less reducing potential of -2.7 V compared to the reduction potential of -3.04 V for Li, and also lower volumetric and gravimetric capacities due to the heavier Na atom (see Fig. 16 above). A typical charge-discharge curve for a Na-NaCoO₂ cell is shown in Fig. 18, which also includes a similar curve for the Li-LiCoO₂ couple for comparison.¹⁴⁰ Both cathode structures are analogs of each other and consist of layers of CoO₂ slabs in between which Na and Li ions reside as shown in the inset. The figure also compares the operating cell

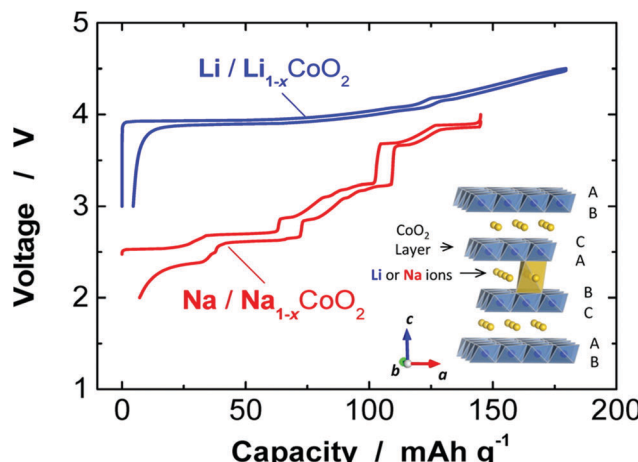


Fig. 18 Comparative charge-discharge curves for Li/Li_{1-x}CoO₂ and Na/Na_{1-x}CoO₂ cells. Inset depicts Li or Na insertion into the layered crystal structure of the cathode.¹⁴⁰ Republished with permission of The Electrochemical Society, from [K. Kubota and S. Komaba, Review – practical issues and future perspective for Na-ion batteries, *J. Electrochem. Soc.*, 2015, **162**, A2538–A2550]; permission conveyed through Copyright Clearance Center, Inc. (2018).

potential for NaCoO₂, which is >1 V lower than for the LiCoO₂ cell. Accordingly, the Na-ion battery has a lower energy density than the Li-ion battery, marked by the stepwise potential changes for NaCoO₂.

As expected, many of the technical challenges facing Na-ion batteries are similar to those faced by Li-ion batteries, including better materials for electrodes with higher capacities, insufficient cycle life, capacity fading upon cycling, electrolyte stability at high voltages, *etc.* There has been recent progress in some of these areas, especially in seeking earth abundant, low-cost, and environmentally friendly electrode host materials for sustainability. For example, inspired by the early report¹⁴⁸ of high theoretical capacity (560 mA h g⁻¹) for dilithium rhodizonate (Li₂C₆O₆), which suffered from poor cycle life due to dissolution and exfoliation, its sodium analog disodium rhodizonate (Na₂C₆O₆) was recently investigated as an organic cathode material and reported achieving a four-electron redox reaction with a reversible capacity of 484 mA h g⁻¹, energy density of 726 W h kg⁻¹ of cathode material, and an energy efficiency above 87%.¹⁴⁹ The key to the observed long cycle life of this material was to understand and kinetically suppress the electrochemically limiting irreversible phase transformation between α -Na₂C₆O₆ and γ -Na₂C₆O₆ during cycling by controlling particle size and morphology as well as using electrolyte additives. Also, this organic-based cathode material is derived from biomass, particularly from the corn extract myo-inositol, and hence, provides a sustainable and environmentally friendly synthesis route to produce the cathode material.^{31,148}

Similarly, a new class of non-rare-earth containing Fe-based cathode material with a composition Na₂Fe₂(SO₄)₃ was recently reported to provide an exceptionally high potential of 3.8 V *versus* Na/Na⁺ with an experimentally measured capacity of 102 mA h g⁻¹, which was highly reversible over 30 cycles.¹⁵⁰

The measured capacity corresponds to about 85% of the theoretical capacity of 120 mA h g⁻¹ based on the one electron Fe³⁺/Fe²⁺ redox transition of the cathode material. Similarly, an all-solid-state Na-ion battery employing a Na-metal anode coated with a thin polymeric interlayer to suppress dendrite formation, a solid Na-ion conducting solid electrolyte of the composition Na₃Zr₂Si₂PO₁₂ (*i.e.*, NASICON) and ionic conductivity of $\sim 10^{-3}$ S cm⁻¹ at temperatures ~ 65 °C, and a NaTi₂(PO₄) cathode provided 102 mA h g⁻¹ after 70 cycles at 0.2C at 65 °C with a Coulombic efficiency of 99.7%.¹⁵¹

As the ionic size of Na (radius = 97 pm), compared to Li (radius = 68 pm), is too large to facilitate mobility in a closed packed framework of oxide ions, it needs a host structure with sufficiently large interstitial sites. A recent study reported the synthesis of a low-cost Na-Mn-hexacyanoferrate cathode composition where lowering of the activation barrier for Na-ion diffusion was achieved *via* weakening the Na-oxygen bonding by replacing the oxide ions by (CN)⁻¹ ions, whereby providing a reversible capacity of >120 mA h g⁻¹ and a high potential of 3.4 V.¹⁵² Similarly, a family of Prussian blue structure cathode materials were reported for aqueous alkali-ion battery systems, including nanosized Cu hexacyanoferrate that showed no capacity fading after 1000 deep-discharge cycles and provided 95% round-trip efficiency when cycled at 5C¹¹⁹ and Ni hexacyanoferrate that showed complete reversibility with zero capacity loss after 5000 cycles for sodium and 1000 cycles for potassium.¹⁵³

Prussian blue (ferric hexacyanoferrate, or Fe₄[Fe(CN)₆]₃·xH₂O, or C₁₈Fe₇N₁₈) is a low cost, effective cathode host material for the insertion and extraction of potassium over thousands of cycles, and potassium is also cheap and abundant, which makes it an ideal anode material for K-ion batteries. A schematic depiction of the K-ion battery structure with Prussian blue cathode and carbon anode is shown in Fig. 19. An early demonstration of K-ion battery employed a Prussian blue-based cathode material and a potassium anode in a nonaqueous electrolyte consisting of 1 M KBF₄

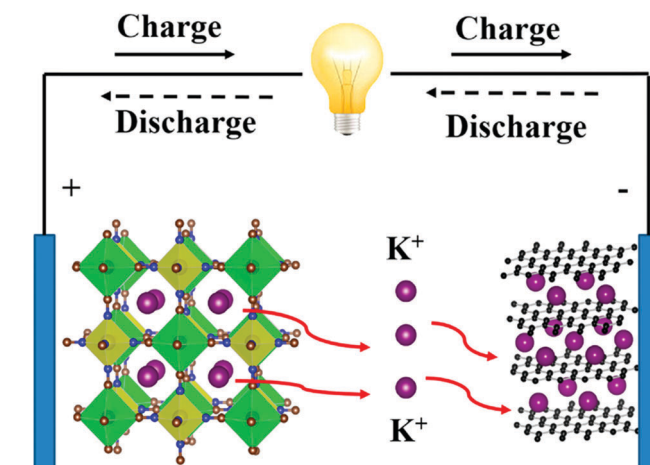


Fig. 19 Schematic illustration of a K-ion cell with Prussian blue structure cathode and carbon-based anode.¹⁴¹ Reprinted with permission from (A. Eftekhari, Z. Jian and X. Ji, Potassium secondary batteries, *ACS Appl. Mater. Interfaces*, 2016, **9**, 4404–4419). Copyright (2018) American Chemical Society.

dissolved in a 3:7 ratio ethylene carbonate/ethylmethyl carbonate mixture, and showed a cathode capacity of 80 mA h g⁻¹ and excellent cyclability for over 500 reversible cycles.¹⁵⁴ With the expected carbon anode capacity of 262 mA h g⁻¹ at C/10 rate and potential of 0.3 vs. K⁺/K, a complete K-ion cell can deliver an energy density of ~200 W h kg⁻¹ based on the mass of the electrodes, and a full battery may be able to provide ~110 W h kg⁻¹.¹⁴¹ Another advantage of K-ion batteries is the highly negative standard potential of -2.936 V vs. the standard hydrogen electrode (SHE) for the K⁺/K redox couple. This value compares quite well with the potential of -3.04 V vs. SHE for the Li⁺/Li couple for Li-ion batteries, while it is still more negative than the Na⁺/Na couple at -2.714 V vs. SHE, providing close to ~200 mV higher cell voltage for K-ion batteries than for Na-ion. Furthermore, transport of the "soft" K⁺ ion is much faster than for the strongly solvated Li⁺ ions in electrolytes, because of the smaller Stoke diameter (*i.e.*, solvation sheath) for potassium. Again, due to many similarities in materials aspects with Li-based battery systems that were discussed previously (see Section D.1.1.: Li-Ion batteries), the interested reader is referred to recent review articles for more information on K-based batteries.^{141,145,155}

Unlike the single-electron transfer reactions in alkali metal-based battery systems, battery chemistries based on charge transfer reactions involving multivalent cations provide a great incentive to design battery materials and systems that accomplish multi-electron transfer and higher storage capacities. The theoretical capacities are presented in Fig. 16. In that regard, magnesium, calcium, and even aluminum offer obvious advantages for progress beyond Li-batteries, albeit with their own challenges and shortcomings. Replacing Li⁺ with multivalent Mg²⁺, Ca²⁺, Al³⁺ cations increases the total charge transferred per ion, which naturally increases storage capacity. For example, the theoretical volumetric capacity of 3833 mA h ml⁻¹ for magnesium is nearly twice that for lithium metal (2062 mA h ml⁻¹), and it has a slightly less negative reduction potential of -2.4 V vs. SHE compared to lithium (see Fig. 16). Similarly, Al offers a theoretical volumetric capacity of nearly four times (*i.e.*, 8040 mA h ml⁻¹) that of lithium and a reduction potential of only -1.67 V vs. SHE, while Ca has a theoretical capacity of 2073 mA h g⁻¹ and a reduction potential of -2.87 V vs. SHE.¹³⁹ Moreover, Mg, Ca, and Al, are cheap and among the most abundant elements on the planet (see Fig. 15). They also do not show a high propensity for dendrite formation, unlike the case for metallic Li anodes. Also, these metals do not pose major safety issues and are easier to work with during processing. They do not need to be processed in glove boxes under tightly controlled and monitored inert and dry atmospheres.

Despite the great potential of these metals to be suitable battery anodes, however, progress in these systems has been slow due to multiple challenges. Major advances in the discovery of superior electrode and electrolyte materials, stability, reversible cyclability and long cycle life are needed in order to capitalize on the attractively high theoretical capacities offered by the chemistries of multivalent cations. Many of these challenges and materials issues for Mg, Ca, and Al battery systems have recently been reviewed.^{139,144,156–158}

There are critical barriers to designing a rechargeable battery based on these cations. For example, finding suitable high capacity cathode structures that can reversibly intercalate these multivalent cations at high potentials remains a major challenge. During charging of the battery, these metals tend to form a passive, electronically insulating oxide layer, which also blocks cation transport. Electrolytes with sufficient stability have yet to be identified that can facilitate the formation of conductive passivation layers on the electrodes at high potentials. Intercalation of these cations into host cathode structures has been difficult and slow due to low mobility, which presents a barrier for further development. The high oxidation state of these cations slows them during mass transport due to stronger Coulombic interactions with the host electrode structures. Also, their reduction potentials are lower than that for Li, and hence provide a lower operating cell voltage for the battery.

Nevertheless, some progress has been made in some of these fronts. For example, fast Mg²⁺ ion conductivity of 0.01–0.1 mS cm⁻¹ at 298 K in the MgSc₂Se₄ spinel was recently reported with a low migration barrier of only 370 mV.¹⁵⁶ It should be noted that previous reports of comparably high Mg²⁺ mobility in spinel and Chevrel phases were obtained only at temperatures (400–800 °C), while this recent study constitutes the first report of high Mg²⁺ mobility in a closed packed chalcogenide structure at room temperature. Similarly, one of the first demonstrations of reversible Ca²⁺ intercalation was recently reported for a Prussian blue structure manganese hexacyanoferrate (MFCN) cathode in a nonaqueous electrolyte giving a capacity of 80 mA h g⁻¹, which retained 50% of its value after 35 cycles.¹⁴⁴

Aluminum is cheap and one of the most abundant metals on Earth (see Fig. 15). As a trivalent ion, aluminum-based batteries provide a three-electron redox reaction that results in a theoretical capacity of 2.98 A h g⁻¹.¹⁵⁷ This is slightly less than the theoretical capacity of 3.86 A h g⁻¹ for lithium, while its volumetric capacity of 8.04 A h ml⁻¹ is much superior to 2.06 A h ml⁻¹ for lithium, 3.83 A h ml⁻¹ for magnesium, and 5.85 A h ml⁻¹ for zinc (see Fig. 16). However, early attempts to develop a rechargeable Al-based battery has faced numerous challenges including low cell discharge voltage, short cycle life (<100 cycles) in part due to disintegration of the cathode materials, and high capacity fading rate (26–85%). The first functional Al-ion rechargeable battery utilizing a Al metal anode, V₂O₅ cathode and an ionic liquid electrolyte containing AlCl₃ provided 305 mA h g⁻¹ in the first cycle and 273 mA h g⁻¹ after 20 cycles with a stable Al³⁺ insertion plateau at 0.55 V.¹⁵⁸ Recent progress using a Al metal anode, a graphitic foam cathode and ionic liquid electrolyte demonstrated electrochemical dissolution and deposition of aluminum at the anode, fast intercalation and de-intercalation of chloroaluminate ions in the graphite foam cathode that resulted in rapid (~1 min) charging times at high current densities of ~4000 mA g⁻¹, and exhibited a stable discharge plateau at ~2.0 V, a low specific capacity of 70 mA h g⁻¹, high Coulombic efficiency of 98%, and long cycle life >7500 without decay.¹⁵⁹ Despite the remarkable progress, this Al-ion/graphite system was able to provide an

energy density of only $\sim 40 \text{ W h kg}^{-1}$, which is comparable to Pb-acid or Ni-metal hydride batteries. On the other hand, however, it offered a high power density of $\sim 3000 \text{ W kg}^{-1}$ that is comparable to supercapacitors.

D.1.2.3. Na-S batteries. The active materials of this battery system, namely, sodium and sulfur, are cheap, abundant, and readily available. Moreover the Na-S system has a cell potential of 2.1 V and high theoretical energy densities of 790 W h kg^{-1} and 1180 W h L^{-1} ,⁴⁴ high energy efficiency, and great cyclability typically in the 1000's of cycles. These attractive properties have attracted great interest in the Na-S system in the last several decades. They have been commercialized in Japan since 2002 for stationary storage in MW scale systems, and generally used in load-leveling and peak-shaving applications by the utilities. There are hundreds of Na-S battery installations around the world, most of them in Japan, that provide in excess of 300 MW of discharge capacity for energy storage.

The technical challenges for Li- and Na-sulfur battery systems posed by the high solubility of alkali metal polysulfide species in the electrolyte that are the root cause of multiple problems including passivation layers building on the electrodes, polysulfide shuttling through the electrolyte, and capacity fading as discussed in previously (see Section D.1.2.1.: Li-sulfur batteries) can be largely overcome by replacing the liquid electrolyte with a solid impervious membrane that selectively transports only alkali ions across and provides a physical barrier that separates the alkali metal anode from the sulfur cathode. This has been the underlying reason and the rationale for the development of the Na-S battery, which employs a Na-ion conducting solid electrolyte between the molten sodium anode and the sulfur cathode.

Indeed, the discovery of fast sodium ion transport in a common refractory Na β -alumina ($\text{Na}_2\text{O}\cdot 0.11\text{Al}_2\text{O}_3$, or $\text{NaAl}_{11}\text{O}_{17}$) near room temperature in early 1960's made this possible and generated great interest in Na-ion based electrochemical storage systems. At 300°C , the ionic conductivity of Na β -alumina approaches the room temperature conductivity of $10^{-1} \text{ S cm}^{-1}$, comparable to common aqueous electrolytes. Another member of this family is the Na β'' -alumina, which exhibits an ionic conductivity of 1.3 S cm^{-1} at 500°C .¹⁶⁰ These highly conductive electrolyte materials opened up opportunities to explore sodium chemistries for power generation and storage applications, including Na-S and Na-metal chloride batteries.¹⁶¹

Schematic depiction of the Na-S battery tubular architecture and its operating principle is shown in Fig. 20. At the typical operating temperature of $300\text{--}350^\circ\text{C}$, both the Na anode and the sulfur cathode are in their molten state. Generally, a tubular sodium β'' -alumina impervious electrolyte membrane houses the molten Na (negative electrode), and separates it from the molten sulfur (positive electrode). Due to its low conductivity, the sulfur electrode is mixed with carbon to improve the electronic conductivity of the positive electrode.

Elemental sulfur has a high theoretical specific capacity of 1675 mA h g^{-1} based on the 2-electron half-cell reaction at the cathode. During discharge, Na is oxidized to Na^+ and is

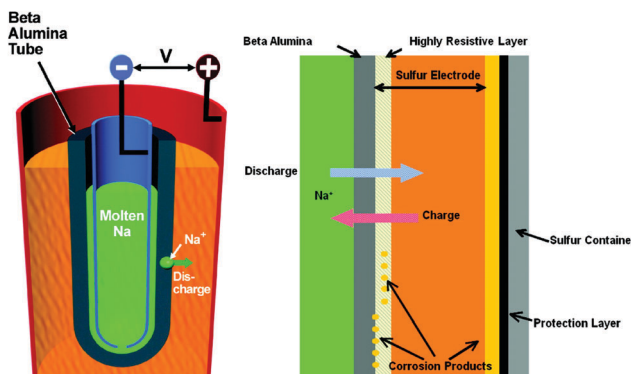


Fig. 20 Schematic cell structure (left) and operating principle (right) of the Na-S battery.³² Reprinted with permission from (Z. Yang, J. Zhang, M. C. W. Kintner-Meyer, X. Lu, D. Choi, J. P. Lemmon and J. Liu, *Chem. Rev.*, 2011, **111**, 3577–3613). Copyright (2018) American Chemical Society.

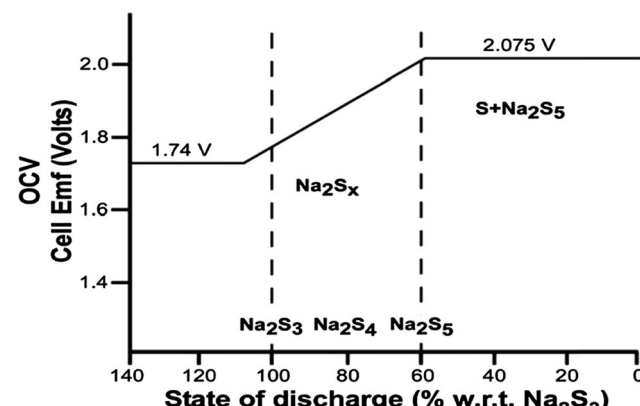


Fig. 21 Voltage regimes and phase boundaries of polysulfides in the Na-S system. Reproduced from ref. 162 with permission from The Royal Society of Chemistry.

transported across the electrolyte to react with sulfur and form sodium polysulfides (Na_2S_x). The phase boundaries of these polysulfides and the voltage regimes are presented in Fig. 21. At the open circuit potential of 2.075 V, there exists a two-phase region as Na_2S_5 is insoluble in sulfur, so forms a two-phase liquid mixture at the operating temperature. As the battery discharges, sulfur combines with sodium to form a single-phase region marked by Na_2S_4 in the figure. If unchecked, the level of molten Na drops due to consumption, so process control is important. At 1.74 V, Na_2S_4 is reduced to Na_2S_3 . For deeper discharge, the undesirable solid Na_2S_2 may form and increase the cell impedance. During charging, the reactions are reversed, but the re-formation of sulfur must be managed. Otherwise, it can deposit on the electrolyte and form an insulating layer, which increases cell impedance and lowers performance. The round-trip efficiency of Na-S batteries is around 80% with an estimated life of ~ 15 years for >4500 cycles at 90% depth of charge.¹⁶²

The Na^+ released at the positive electrode is transported back through the Na^+ ion conducting ceramic electrolyte into the negative electrode, where it picks up an electron from the

external circuit and deposits as metallic sodium. In order to facilitate sufficient rates (*i.e.*, current) of ion transport through the solid electrolyte, the battery operates typically at 350 °C. The heat produced during discharge/charge cycles is sufficient to maintain the battery temperature. Usually, no external heat input is needed.

Recent review articles provide more details into the operational, performance, and materials aspects of Na⁺-based batteries and their technical challenges.^{142,161,163}

Although Na-S battery systems are commercially available and offer attractive advantages including inexpensive, abundant, nontoxic, and environmentally benign materials, while offering low maintenance, cycling flexibility and long cycle life, high storage capacity and efficiencies,²⁴ they also face technical challenges still remain to overcome. One critical problem is related to corrosion and vapor loss. High operating temperatures compared to other storage systems reviewed in earlier sections, as well as the highly reactive nature of the molten electrodes pose hot corrosion problems, related in particular, to the highly corrosive nature of the S and Na₂S₅ species at these temperatures. This places expensive demands on materials selection for the container and other cell components such as hermetic seals. High operating temperatures required for achieving sufficient ion transport through the ceramic electrolyte is also dictated by the difficulty in achieving proper wetting of the solid electrolyte by molten sodium, which is one of the most challenging issues for Na-S batteries.

The solid electrolyte, namely β- (or β"-) alumina, is a critical and expensive component of the Na-S battery. Although ionic transport of Na⁺ through the electrolyte occurs anisotropically along the 2-dimensional conduction planes separated by close-packed Al-O blocks, the random positioning of the fine grains of powders that make up the finished polycrystalline electrolyte tubes mitigates the anisotropy problem greatly. Ionic conductivities of the tubular polycrystalline electrolyte are only a factor of 3–5 smaller than that for the single crystal electrolyte. However, mechanical fragility and process yields for the tubular β-alumina fabrication are major concerns and cost factors for mass production of these electrolytes.

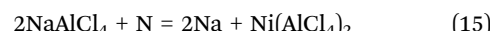
Reducing the operating temperature of the Na-S battery is important for improving its safety, cost, and durability by slowing down degradation mechanisms. A recent study attempted to reduce the operating temperature to 150 °C by employing a molten Na separated by a 600 μm thick β"-alumina electrolyte in series with a liquid catholyte of 4:1 molar ratio of S:Na₂S₄ dissolved in tetraglyme solution reported an initial discharge capacity of 473 mA h g⁻¹ of which 70% was retained after 60 cycles, and cell voltages around 2 V.¹⁶⁴ Similarly, alloying Na with 6.7 mol% of the low surface tension metal Cs markedly improves wetting of the β"-alumina solid electrolyte in the molten alloy and at the same time, helps bring down the operating temperature to 150–175 °C with a capacity fading rate of <3% over 100 cycles.¹⁶⁵

Others aimed at reducing the Na-S battery operation to room temperature by employing a sulfur–carbon composite cathode in nonaqueous electrolyte.¹⁶⁶ One critical challenge

to room temperature operation is the instability of the electrolyte to form electrochemically inactive alkali metal polysulfides. Porous carbon framework doped highly with nitrogen and sulfur was suggested to help trap the sodium polysulfides and inhibit reaction with the electrolyte, and provide a long cycle of over 10 000 cycles with less than 20% capacity fading (3% after 800 cycles) at a high current density of 4.6 A g⁻¹.¹⁶⁷ Similarly, a more recent study that employed a Na metal anode, a metal–organic framework (MOF)-derived microporous carbon–sulfur composite cathode in a liquid carbonate electrolyte containing 1 M NaClO₄ and an ionic liquid (1-methyl-3-propylimidazolium-chlorate) reported a reversible storage capacity of 860 mA h g⁻¹ at 0.1C rate (1C = 1675 mA g⁻¹) and 600 mA h g⁻¹ at 0.5C with a capacity fade rate of only 0.31% over 100 cycles at room temperature.¹⁶⁸

A different family ionically conducting solid electrolyte dubbed NASICON (for sodium-super-ionic-conductor) with the nominal composition Na_{1+x}Zr₃P_{3-x}Si_xO₁₂ (0 < x < 3) was introduced in the 1970's.¹⁶⁹ Unlike 2D transport of Na⁺ in the layer structured β-alumina, which poses challenges in processing and ohmic losses due to misalignment of grains in the electrolyte membrane in the actual battery, transport of Na ions in the NASICON structure is 3-dimensional. This of course offers many advantages, and also, isotropic conductivity of 6 × 10⁻⁴ S cm⁻¹ at room temperature,¹⁷⁰ while it increases to 0.2 S cm⁻¹ at 300 °C with an activation energy of 0.29 eV,¹⁶⁹ which is only 0.1 eV larger than that for β"-alumina. Despite the attractively high ionic conductivity for Na⁺, NASICON is known to be chemically unstable in the presence of molten sodium and degrade rapidly to eventual fracture.¹⁷¹ It is also unstable in water, and therefore not suitable for aqueous systems that require long service life.

Another Na-based storage system utilizing the β-alumina family of electrolytes is based on the Na–NiCl₂ chemistry (also called the ZEBRA cell) that also operates at moderately high temperatures of 270–300 °C. The ZEBRA battery has theoretical and practical energy densities of 788 and 120 W h kg⁻¹, respectively, theoretical capacity of 305 A h kg⁻¹, high cell voltage (2.58 V), maintenance free operation, zero self-discharge, and long cycle life up to 5000 cycles at 80% depth of discharge.¹⁶² The positive electrode structure contains interconnecting porous Ni particles dispersed in the semisolid composite of the electrochemically active NiCl₂ and a molten secondary electrolyte NaAlCl₄ that conducts Na-ions. During discharge, sodium is oxidized to Na⁺ and transported through the β-alumina electrolyte and then across the NaAlCl₄ electrolyte to the cathode, where it reacts with NiCl₂ to form NaCl and metallic Ni, giving a cell potential of 2.58 V. During charging, the reverse process takes place, where the Ni is oxidized and reacts with NaCl to form NiCl₂. Aside from conducting Na⁺ ions to the cathode during discharge, the other important function of NaAlCl₄ is to overcome overcharging the battery, as it serves as a source for Na by the reversible reaction,



ZEBRA batteries offer several advantages over Na-S, including higher cell voltage, safer operation, tolerance to overcharging,

and ease of assembly of the battery in the discharged state. The positive electrode primarily consists of solids, reducing corrosion and associated problems. It also eliminates the need to handle metallic Na for the negative electrode, as all Na is introduced in the form of salts using AlCl_3 and NaCl . However, capacity fading due to the solubility of NiCl_2 in the NaAlCl_4 and coarsening of the Ni particles pose operational challenges.¹⁶²

D.1.2.4. Metal-air batteries. The operating principle of metal-air systems in essence can be considered between that of traditional batteries and fuel cells. As in many battery systems, the fuel is a bulk metal serving as the negative electrode, while the reduction reaction at the positive electrode is accomplished by the oxygen supplied through a porous electrode from the ambient air, as in fuel cells. Elimination of the bulky positive electrode of the battery and replacing it with an air electrode is key to achieving exceptionally high theoretical energy densities in metal-air systems. Obviously, such advantage that potentially offers practical prospects especially in the transportation sector has generated interest in these systems that goes back several decades.¹⁷²

Although the name “metal-air” battery is used commonly to refer to these systems as also adopted in this article, this should be considered merely as an operational label. Strictly speaking, they are metal–oxygen batteries, as they utilize oxygen from the ambient air. Although this is indeed a great advantage as air is free, it also presents operational challenges that may vary for these systems depending on the nature of the electrolyte and the metal anode they employ. Regardless, however, almost all metal–oxygen batteries require some degree of pre-treatment or removal of air impurities primarily CO_2 and moisture, but also SO_2 , N_2 and possibly others depending on the metal–air system from entering the battery proper, and adversely interact with the electrolyte or battery components *via* parasitic reactions. For example, CO_2 is known to readily absorb in alkaline electrolyte solutions and precipitate in the porous cathode as solid carbonates, while moisture is detrimental to nonaqueous and aprotic electrolytes. In many cases, pure oxygen has preferably been employed. This is especially true for Li-air batteries, where air impurities are particularly problematic. Others employed polymeric membranes to avoid problems associated with air impurities.¹⁷³ Regardless, however, the need for such pre-treatment or clean-up processes or for oxygen separation increases the complexity of the energy storage

process and adds weight to the battery that collectively reduces both the system efficiency and the energy density.

Portable electronics, transportation and other types of mobile storage applications demand high gravimetric and volumetric energy densities, low cost, long cycle life, and safe and stable operation. These requirements are difficult to meet using currently available commercial battery chemistries. For example, commercial Li-ion batteries provide close 150 to 200 W h kg^{-1} , while the power system for an electrical vehicle with a 500 mile range would require nearly 10 times increase in the energy density of the battery,¹⁷⁴ which is most likely an unrealistic expectation. In search for improved electrochemical systems, metal-air batteries offer many advantages particularly with respect to the amount of energy they can store. Many of the characteristic features and operational properties of metal–air battery systems are summarized in Table 4 for comparison and reviewed elsewhere.^{175,176} By definition, alkaline electrolyte is an aqueous solution of a base such as NaOH . Unlike protic solvents such as water and many alcohols including methanol that contain labile protons bonded to oxygen or nitrogen, which can then be donated *via* hydrogen bonding, aprotic solvents such as propylene carbonate lack acidic hydrogen (*i.e.*, O–H bonds) and hence cannot make hydrogen bonding to donate their protons. Similarly, saline electrolyte in the table refers to an aqueous solution of a soluble salt such as NaCl .

While the cell architecture of the commercially available rechargeable batteries is closed, metal-air batteries have open architecture, where oxygen from air is accessed through the porous cathode. They have lower cost and considerably higher energy densities than other battery systems. They utilize the conversion chemistry and the high oxidation potential of the pure metal electrode with respect to oxygen. They also eliminate the need for the massive positive electrode that holds the electrochemically active species as in other battery systems, and replace it with a porous air electrode.

The Zn–air primary system is the only commercialized and widely used battery, most for hearing aids. Al–Air and Mg–air batteries are employed in the military for underwater propulsion. The other metal–air battery systems are still under development for commercial applications. Several of these battery systems, namely, Zn–air,^{175,176} Al–air,^{157,177} and Mg–air,¹³⁹ can be recharged mechanically, or refueled, by simply replacing the discharged anode and the spent electrolyte with fresh ones. This also allows the air electrode to be operational during

Table 4 Comparison of characteristic features and properties of metal–air battery systems. Adapted from ref. 175

| Battery systems | Li-Air | K-Air | Na-Air | Mg-Air | Zn-Air | Fe-Air | Al-Air |
|--|------------------------|------------------------|------------------------|---------|----------|----------|---------------------|
| Cost of metal ^a (\$ per kg) | 68 | 20 | 1.7 | 2.75 | 1.85 | 0.4 | 1.75 |
| Theoretical voltage (V) | 2.96 | 2.48 | 2.27 | 3.09 | 1.65 | 1.28 | 2.71 |
| Practical voltage (V) | 2.6 | 2.4 | 2.2 | 1.2–1.4 | 1.0–1.2 | 1.0 | 1.1–1.4 |
| Theoretical energy density ^b (W h kg^{-1}) | 3458 | 935 | 1106 | 2840 | 1086 | 763 | 2796 |
| Practical energy density (W h kg^{-1}) | Not known ^c | Not known ^c | Not known ^c | 400–700 | 350–500 | 60–80 | 300–500 |
| Electrolyte | Aprotic | Aprotic | Aprotic | Saline | Alkaline | Alkaline | Alkaline, or saline |
| Primary (P), or rechargeable (R) | R | R | R | P | R | R | P |
| Year invented | 1996 | 2013 | 2012 | 1966 | 1878 | 1968 | 1962 |

^a 2014 prices. ^b Includes oxygen. ^c Literature values are normalized to mass of catalyst.

discharge only as the cathode for oxygen reduction. Despite intense research efforts, however, achieving electrical recharging in a reversible, efficient and cost effective manner has been challenging, and presents the major obstacle for the commercialization of these high-energy density systems. Among the metal–air systems, however, Li–air and Zn–air batteries attracted the most interest partly because of their exceptionally high energy density and other advantages.¹⁷⁶

Cell type are labeled according to the anode material, while cell configurations can be grouped with respect to the nature of their electrolyte, namely, aqueous and non-aqueous (containing aprotic solvent). As expected, aprotic (no hydrogen bonding) electrolytes, usually employed for alkali metal–air batteries, are susceptible to moisture contamination leading to cell and performance degradation, while Fe-, Zn-, and Al-air batteries use aqueous electrolytes. Recent advances in Li-air battery materials and architecture also allow the use of aqueous electrolytes.^{174,178} Many of the metal–air battery systems and their cell components, electrode reactions and cell performances are reviewed elsewhere.^{174–176,178–180}

Despite differences in their chemistries, there are many technical challenges shared by metal–air batteries regarding the anode and cathode materials and processes. Typically, poor utilization of the anode metal and slow kinetics at the air electrode are the most common problems. The anodes typically undergo corrosion in the electrolyte, which leads to self-discharge and short shelf life. This also monotonically decreases the battery's Coulombic efficiency. The non-conductive oxide and/or hydroxide layer built after the first discharge cycle due to severe passivation of the anode metal surface impedes electrolyte access to the liquid–solid–gas interface at the anode and also blocks the active anode material to participate in the oxidation/dissolution reaction, and hence, slows down the anode reaction kinetics. To mitigate these problems, one strategy is to isolate one of the active battery components, *i.e.*, either the anode or the electrolyte, from the cell until the battery is ready for activation and use. Regardless, however, this does not solve the reversibility requirement for achieving rechargeability of the battery, as the blocking layer impedes plating of the metal on the anode during the charging cycle.

Likewise, the air electrode is arguably the most complicated and expensive component of metal–air systems. It suffers from intrinsically sluggish oxygen kinetics, high polarization losses, and poor reversibility of the oxygen electrochemistry.¹⁸¹ The requirements for developing an effective air electrode also depend on the chemistry of the metal anode as well as the class or type of the electrolyte employed for the battery. These considerations are recently reviewed in detail elsewhere.¹⁸⁰ At the air electrode, oxygen from air accepts electrons from the metal-ion during discharge and releases them during charging. Finding a bifunctional facile catalyst that can perform both the oxygen reduction reaction (ORR) and oxygen evolution reaction (OER) reversibly and fast at or near room temperature presents many challenges. The obvious candidates, *i.e.*, precious metal catalysts are prohibitively expensive and hence, not suitable. To cut costs and preserve performance, one study synthesized

PtAu nanoparticles (6.8 nm) supported on carbon and reported bifunctional catalytic activity for both the ORR and OER with a high round trip efficiency of 77% for the Li–air system.¹⁸² So there is a need for developing non-precious metal electrocatalysts for the air electrode. In that regard, earth abundant transition-metal oxides offer great incentives. For example, manganese oxides (MnO_x) have been shown to be effective catalyst for oxygen reduction in alkaline air electrodes, and MnOOH was identified as the bifunctional catalytic species.¹⁸³ Manganese oxide is also shown to be an effective bifunctional electrocatalyst for the oxygen evolution reaction.^{184,185} Likewise, various doped transition-metal oxide compositions of the perovskite structure have also been investigated. Guided by the importance of catalyst d-band center as an effective descriptor for the overvoltage behavior for oxygen reduction as suggested earlier,¹⁸⁶ a total of 15 A-site and B-site doped-perovskite compositions were explored and the results concluded that increasing the covalency between the B-site transition metal 3d and oxygen 2p orbitals helps reduce the overvoltage for the oxygen reduction reaction on these catalysts.¹⁸⁷

Aside from the importance of oxygen kinetics, the electrode design and structure must also consider avoiding the detrimental effects of CO_2 exposure in the case of aqueous electrolytes and moisture in the case of nonaqueous, aprotic electrolytes, which can lead to electrolyte degradation, while at the same time, provide a means for selectively introducing oxygen to the electrocatalyst surface of the air electrode.

D.1.2.4.1. Li-Air batteries. Arguably, the Li–air battery enjoyed the most interest and most intense efforts among the other metal–air systems, mostly due to its potential for large-scale energy storage applications. The oxidation of 1 kg of Li releases nearly 11 700 W h of energy, almost on par with the energy storage capacity of gasoline. On the system level, the Li–air system offers up to 10 times the theoretical energy density of the Li-ion battery. The bulky positive electrode for the intercalation or conversion reaction that is employed in Li-ion batteries is completely eliminated in the Li–air system, and is replaced by a porous oxygen-breathing cathode. The concept of the air electrode is not new, however, and has long been employed in various types of fuel cells and also in Zn–air batteries.

Table 5 summarizes the key reactions at the negative electrode, and compares the energy densities and cell potentials of Li-based battery systems. It is clear that the Li–S and Li–air cells offer significantly higher theoretical energy densities than the Li-ion chemistry. The actual cell-level energy densities depend strongly on the air cathode porosity and electrolyte. For a Li–air cell operating at 2.9–3.1 V with a cathode of 70% porosity, the maximum theoretical cell capacities are estimated to be 435 mA h g^{-1} and 509 mA h ml^{-1} (or, 1300 W h kg^{-1} and 1520 W h L^{-1}) for alkaline electrolyte, and 378 mA h g^{-1} and 452 mA h ml^{-1} (1400 W h kg^{-1} and 1680 W h L^{-1}) for acidic electrolyte, while the capacities for the same cathode in nonaqueous electrolyte are nearly two times higher at 940 mA h g^{-1} and 950 mA h ml^{-1}

Table 5 Theoretical energy densities of Li-air batteries compared to other Li-based storage systems⁴³

| Battery system and net cell reaction | Cell voltage (V) | Theoretical specific energy (W h kg ⁻¹) | Theoretical energy density (W h L ⁻¹) |
|--|------------------|---|---|
| Li–O ₂ battery (nonaqueous) 2Li + O ₂ = Li ₂ O ₂ | 3.0 | 3505 | 3436 ^a |
| Li–O ₂ battery (aqueous) 2Li + $\frac{1}{2}$ O ₂ + H ₂ O = 2LiOH | 3.2 | 3582 | 2234 ^b |
| Li-Ion battery $\frac{1}{2}$ C ₆ Li + Li _{0.5} CoO ₂ = 3C + LiCoO ₂ | 3.8 | 387 | 1015 |
| Li–S battery 2Li + S = Li ₂ S | 2.2 | 2567 | 2199 ^c |

^a Based on the sum of initial volume of Li and the final volume of Li₂O₂ after discharge. ^b Based on the sum of initial volume of (Li + H₂O) consumed and the final volume of LiOH after discharge. ^c Based on the sum of initial volume of Li and the final volume of Li₂S after discharge.

(or, 2790 W h kg⁻¹ and 2800 W h L⁻¹).¹⁸⁸ This is due to the fact that excess electrolyte is needed to counter the consumption of the aqueous electrolyte during cell operation. However, a practical Li-air battery employing nonaqueous electrolyte has not been developed yet.¹⁸⁹ So, it is not clear if the estimated capacities reported above for Li-air batteries with nonaqueous electrolytes may or may not be realistic. Nevertheless, the potential for achieving exceptionally high energy density batteries offered by the Li–oxygen chemistry is the underlying reason for the intense interest in this system.

Although the concept of Li-air goes back nearly two decades, the first rechargeable Li-air battery was reported in 2006 using a Li metal anode, MnO₂/C cathode, and 1 M LiPF₆ dissolved in propylene carbonate nonaqueous electrolyte, and showed that Li₂O₂ is key to reversibility of the battery, and forms on discharge and decomposes back to Li and O₂ during charging.¹⁹⁰ Since then, a variety of Li-air battery designs and materials have been reported, and the progress has been reviewed elsewhere.^{174,176,189,191–196} The cell designs that are currently being pursued are illustrated in Fig. 22 and can be grouped under four general classes of electrolytes, namely, aprotic liquids, aqueous electrolytes, solid electrolytes, and the hybrid, or mixed, electrolyte design concept where the cathode is immersed in an aqueous electrolyte while the anode resides in an aprotic electrolyte. Examples of aprotic electrolytes used in these studies include organic carbonates such as ethylene carbonate, propylene carbonate, and dimethyl carbonate, ethers such as tetrahydrofuran (THF) and dioxolane, and esters such as γ -butyrolactone, where soluble Li-salts such as LiAsF₆, LiPF₆, LiSO₃CF₃, or LiN(SO₂CF₃)₂ are dissolved.

Both in the aprotic electrolyte and hybrid designs, where the Li-metal anode resides in the aprotic electrolyte compartment, the carbonates employed in the electrolyte reacts with the lithium to form a solid electrolyte interphase (SEI) layer that protects the electrolyte from reacting further with the Li anode. It should be noted, however, that organic carbonate-based nonaqueous electrolytes are later shown to decompose irreversibly at the cathode on discharge into several parasitic products. The microstructural aspects of the SEI layer are discussed earlier (see Section D.1.1.3.: Solid electrolyte interphase). Alternatively, a

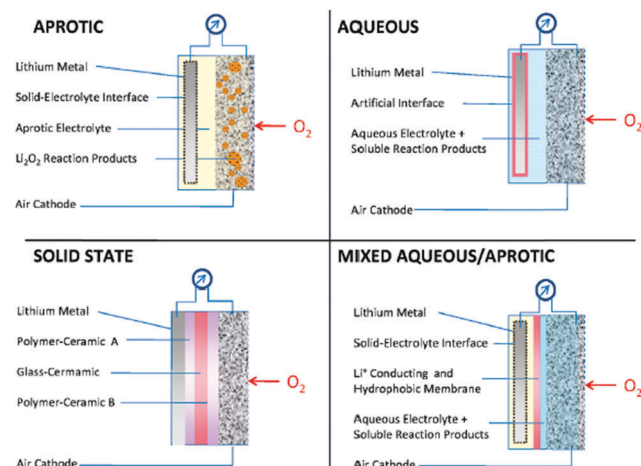


Fig. 22 Li-Air battery architectures using four different nonaqueous, aqueous, solid state and mixed (or, hybrid) electrolytes. Dashed boundaries indicate the spontaneously formed SEI layers, while artificial SEI layers are indicated by solid lines.¹⁷⁴ Reprinted with permission from (G. Girishkumar, B. McCloskey, A. C. Luntz, S. Swanson and W. Wilcke, Lithium–air battery: promise and challenges, *J. Phys. Chem. Lett.*, 2010, **1**, 2193–2203). Copyright (2018) American Chemical Society.

selectively Li⁺-ion conducting but electronically insulating solid state electrolyte membrane either of ceramic or polymer in nature may be coated over the Li anode and serves as the artificial SEI. The natural and artificial SEI concepts are illustrated in Fig. 23. A similar SEI layer is also formed naturally on the positive electrode, demonstrating the importance of understanding and control of the electrode/electrolyte interface.

In the aprotic electrolyte cells, the reaction at the anode during discharge involves oxidation of metallic Li to Li⁺ to form Li₂O₂ at the cathode, and also possibly Li₂O, yielding a cell potential of 2.96 V, although the practical discharge potential is typically around 2.7 V, while that for charging is over 4 V as depicted in Fig. 24. The large difference between the charge and discharge potentials leads to low round-trip efficiencies around 65%. As Li₂O₂, which is electronically and ionically insulating, builds up and clogs up the pores of the porous positive electrode, it increases polarization losses and is believed to be the controlling factor in governing the end point of discharge. The formation of Li₂O₂ as well as the reduction of oxygen at the cathode requires the presence of a catalyst for activating these reactions. Even though several transition metal oxides including Fe₂O₃, CuO, Co₃O₄ and, in particular MnO₂, are found to be effective catalysts for this purpose¹⁹⁷ discharge overpotentials of 0.3–0.4 V below the theoretical value as well as total cell overpotentials around 1.5 V are typically observed in practical cells between charge and discharge curves (see Fig. 24). The status and advances in cathode materials and processes are extensively reviewed elsewhere.¹⁹⁸

The blocking of catalytic sites by the solid Li₂O₂ is a major problem not only on discharge but during the charging cycle as well, leading to nucleating the reaction products and high polarization losses. Moving to aqueous electrolytes would provide partial solution to the blocking problem at

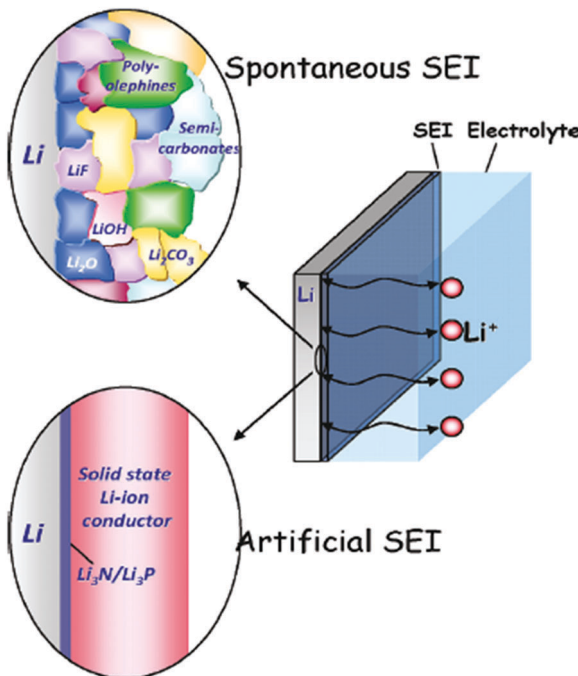


Fig. 23 Schematic depiction of the spontaneous and artificial SEI layer on the Li anode of the Li-air battery.¹⁷⁴ Reprinted with permission from (G. Girishkumar, B. McCloskey, A. C. Luntz, S. Swanson and W. Wilcke, Lithium–air battery: promise and challenges, *J. Phys. Chem. Lett.*, 2010, **1**, 2193–2203). Copyright (2018) American Chemical Society.

the cathode by producing the soluble LiOH species, but aqueous cells exhibit significantly lower cell voltage and reduced capacity. Indeed, a protective Li-ion conducting solid electrolyte such as LISICON (for “lithium super ion conductor”, e.g., $\text{Li}_{1+x+y}\text{Al}_x\text{Ti}_{2-x}\text{Si}_y\text{P}_{3-y}\text{O}_{12}$) layer coated on the Li anode is needed to physically separate the water-reactive Li metal anode from the aqueous electrolyte. This innovation opened up attractive opportunities towards realizing aqueous Li-air batteries.^{199,200} Similarly, a garnet-type of Li-ion conducting ceramic $\text{Li}_{6.4}\text{La}_3\text{Zr}_{1.4}\text{Ta}_{0.6}\text{O}_{12}$ (LLZTO) was employed for a Li-air battery, where it was reported to provide a specific capacity of $20\,300\, \text{mA h g-C}^{-1}$ at $80\,^\circ\text{C}$ and at $20\, \mu\text{A cm}^{-2}$.²⁰¹ The all solid-state Li-air battery concept^{115,123} based on ceramic electrolytes eliminates some of the issues related to other battery designs such as safety and fire hazards related to flammable organic based electrolytes, overcharging, and electrical shorting by dendrite formation, and offers compact battery design, but brings along its own technical challenges including insufficient ionic conductivity and narrow stability window of the electrolyte against Li, interfacial charge transfer resistance, mechanical integrity due to volume changes in the electrodes, and poor cycling life. By contrast, use of solid polymer electrolytes provide flexibility to accommodate for volume change in the battery electrodes, but they have narrow windows for electrochemical stability and poor ionic conductivities that usually allow anion transport also. Their polymer matrix is also sensitive to the presence peroxide and superoxide species and deteriorates in the battery environment.²⁰² Extensive reviews of published

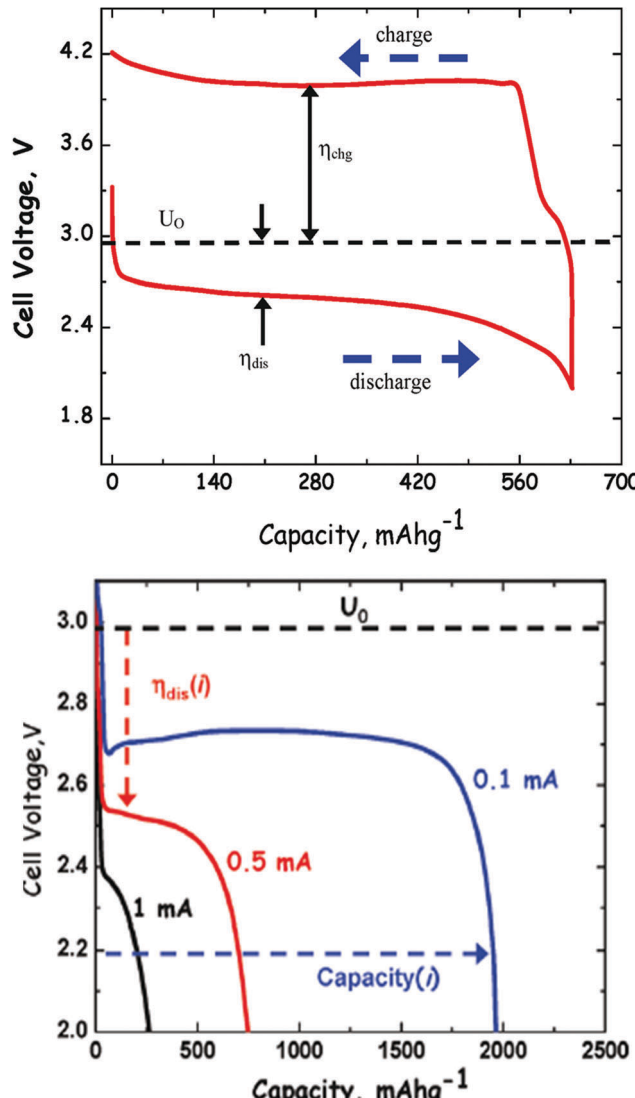


Fig. 24 Capacity and cell charge–discharge overvoltages for a rechargeable Li-air battery with active electrode areas of $1.2\, \text{cm}^2$ in aprotic electrolyte having a theoretical OCV of $2.96\, \text{V}$ shown by the dashed line. (left) Single discharge–charge cycle at a current density of $0.1\, \text{mA cm}^{-2}$ with dashed arrows indicating the charge and discharge directions. (right) Capacity loss as a function of cell current during discharge (per gram of carbon in the cathode).¹⁷⁴ Reprinted with permission from (G. Girishkumar, B. McCloskey, A. C. Luntz, S. Swanson and W. Wilcke, Lithium–air battery: promise and challenges, *J. Phys. Chem. Lett.*, 2010, **1**, 2193–2203). Copyright (2018) American Chemical Society.

work exist on the variety of liquid electrolytes as well as Li-ion conducting ceramic and polymer electrolytes.^{114,115,195,202}

Although there has been immense interest and activity in Li-air battery research and development, there still remains much work before these systems can offer their inherent potential for commercial applications. Of utmost importance is the lack of mechanistic details of the electrode reactions and battery processes, which are highly impacted by the sensitivity of the $\text{Li}-\text{O}_2$ electrochemistry to the nature of the battery electrolyte.^{196,203–206} This makes it difficult to compare and rationalize experimental findings on a common platform.

In fact, a recent review felt the need to caution discretion in interpreting claims of cycle life and efficiency, and battery performance.¹⁸⁹ Moreover, the results are complicated and affected by parasitic reactions both due to the electrolyte solvent and the impurities in the ambient air that may occur in parallel to the Li–O₂ electrochemistry.¹⁹⁶ For example, the parasitic decomposition reaction of the electrolyte solvent dimethoxyethane is identified as the dominant reaction that is catalyzed by carbon, Pt and MnO_x especially above 3.5 V, while the onset of OER starts slightly above OCV indicating that the evolution reaction did not necessitate electrocatalytic promotion.²⁰⁵ It is generally assumed that the parasitic reactions in the electrolyte involve the highly reactive intermediates LiO₂, or O₂[−] such that most strategies for down-selecting a suitable electrolyte have been based on the electrolyte's stability against these species. Recently, singlet oxygen (¹O₂) was detected in the electrolyte above 3.5 V.²⁰⁶ This highly reactive species forms on the cathode upon Li₂CO₃ oxidation in aprotic electrolytes,²⁰⁷ and has been shown to account for parasitic reaction products of electrolyte decomposition.²⁰⁸ Superoxide and peroxide are found to play a role in the formation of the singlet oxygen, which is also promoted by the presence of traces of water in the electrolyte. These important findings must be taken into consideration in the design and interpretation of Li-air battery studies to avoid reaching erroneous conclusions regarding the oxidation of Li₂O₂. If indeed the singlet oxygen holds the key to the reversibility of the Li–O₂ electrochemistry, this will require a shift from electrolyte-focused strategies to finding suitable electrolyte additives that can effectively trap or quench singlet oxygen formation.²⁰⁹

Significant gains have been made in the last few years regarding the mechanistic understanding of electrode processes.¹⁹⁴ The fate of Li₂O₂ during oxygen reduction upon discharge has always been a critical issue. It was shown that whether Li₂O₂ deposits on the cathode as a blocking film or remains in solution as particulates inside the electrolyte is controlled primarily by the solubility of the single electron reaction intermediate LiO₂, which governs the mechanism of cell discharge.²¹⁰ As the solubility of cations in aprotic solvents depends on the solvation of the cation by the solvent molecules defined largely by the Gutmann donor number (DN),²¹¹ aprotic solvents with high donor number exhibit stronger solvation. Thus, for electrolytes with high donor number that strongly solvate Li⁺, the oxygen reduction reaction to form Li₂O₂ occurs inside the electrolyte, while for electrolytes with low donor numbers, Li₂O₂ forms an electronically blocking film on the cathode, passivating the electrode surface and giving rise to low rates, high polarization losses and poor performance. On the other hand, aprotic solvents with low donor numbers are generally more stable, and hence desirable. This conflict is mitigated by adding a soluble redox mediator to the electrolyte that alters the pathway for the oxygen reduction reaction and allowed the use of a low donor number aprotic solvent such as ether.²¹²

Despite the potential of achieving extremely high energy densities and improved specific power,²¹³ most Li-air cell designs still suffer from poor specific capacity and power due to their slow kinetics and high polarization losses that result in

low current densities. During charging, the large overpotential is reported to be associated with deposition of the solid discharge products of electrolyte decomposition at the oxygen evolving surface or the Li₂O₂/electrolyte interface.²¹⁴ They also have stability and materials issues that require further optimization for cost and performance. For example, H₂O contamination in the aprotic electrolyte cells and CO₂ contamination in aqueous cells can lead to parasitic reactions with the Li metal anode or Li₂O₂. Interestingly, addition of trace amounts (500–400 ppm) of water to ethereal electrolytes helped increase discharge capacity by inducing a solution mechanism for the growth of Li₂O₂, while also enhancing parasitic reactions as expected.²¹⁵ Li-Air batteries also exhibit high charge–discharge overvoltages that result in modest round trip efficiencies around 65%.¹⁷⁴ Moreover, these cells generally are not 100% reversible, as the amount of oxygen consumed during discharge is considerably more than the oxygen evolved during charging.²⁰⁹ On the high side, 95% retention of capacity after 100 cycles was reported for Au–C nanoparticle cathode with dimethyl sulfoxide electrolyte, providing a capacity of 3000 mA h g^{−1} (or 300 mA h g^{−1} for carbon–gold combined).²¹³ However, most studies report much lower values.

So at the present state of art, they are not particularly suitable for transportation applications for electric vehicles yet, but can be considered for low load stationary storage applications. However, further understanding and advances in battery electrode materials, catalysts and SEI materials can improve kinetics by, for example, nanostructuring, doping, interface engineering, and other strategies. Similarly, exploring ionic liquid electrolytes,⁵⁵ which offer high ionic conductivity and a wide thermodynamic stability regime up to 5 V, offers exciting possibilities to improve cell stability and kinetics. A recent study using an ionic liquid electrolyte reported a highly reversible charge–discharge cycle, low overvoltage for the charging reaction, and an energy efficiency of 83% for the Li–oxygen battery.²¹⁶ Similar advances were made using non-aqueous electrolytes with high stability. Using tetra(ethylene)-glycol dimethyl ether–lithium triflate (TEGDME–LiCF₃SO₃) electrolyte in a Li–oxygen cell, exceptionally high capacities of up to 5000 mA h g^{−1} and rates of 3 A g^{−1} were reported without the use of a catalyst.²¹⁷ At the average discharge voltage of 2.7 V, this capacity value corresponds to a exceptionally high theoretical energy density of 13 500 W h kg^{−1}.

Although much progress has been made, the jury is out on the commercial prospects of the Li-air battery. The technical challenges are far from being resolved completely. The short cycle life of these systems is still a major problem. Optimistically, even a 500 W h kg^{−1} Li-air battery is only 2–3 times better than the commercial Li-ion battery technology. A recent review article discusses the prospects of the Li-air battery in light of the technical challenges and suggests pursuing instead a holistic rather than an issue or problem-based approach as had been the case so far, to address and resolve these problems.¹⁹³

D.1.2.4.2. Zn–Air batteries. Primary Zn–air batteries have been commercially available for more than half a century, and

this relatively mature technology has widely been employed in hearing aids, medical devices, telecommunications and other applications. The rechargeable Zn-air battery system, however, is a more recent development, and is considered as one of the contenders for electric vehicle applications provided that the technical challenges related to the Zn-anode and air catalyst can be resolved effectively. Recent reviews provide extensive discussions on the details of this battery system.^{175,176,218,219}

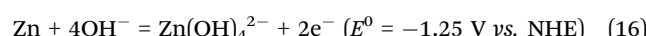
Zinc is a non-toxic, earth-abundant, and relatively cheap material, and is relatively stable to corrosion in aqueous and alkaline solutions. Zn-air batteries are light weight and low cost at <\$10 per kW (> 10 times cheaper than Li-ion batteries), and have high theoretical energy density of 1086 W h kg⁻¹, which is more than 5 times higher than currently available Li-ion batteries on the market. Zn-Air system provides the highest energy density among all primary batteries, and as a rechargeable battery offers a competitive, safe and cost-effective alternative to Li-ion batteries. Recently, flexible Zn-air battery architectures have been reported that may provide innovative and attractive opportunities as portable power for mobile and wearable electronics.^{220,221}

Typical cell design of the Zn-air battery is schematically illustrated in Fig. 25. It consists of three major components, a Zn metal negative electrode, the air cathode and an electrochemically inactive but ionically permeable membrane separator to transport the hydroxyl (*i.e.*, OH⁻) ions. The Zn anode is generally made of fine powdered Zn metal bound by the aid of binders and other additives to make a porous body. Such Zn foam anode structures can provide high energy densities in the range of 300–500 W h kg⁻¹. Other micro- or nanostructured Zn anodes have also been developed in many different forms including nanospheres, nanofibers, flakes, rods, or sheets in

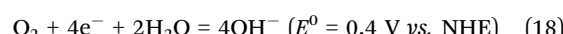
order to increase surface area and battery performance.²²² Indeed, these attempts resulted in increasing the capacity by 40%, energy by 50% and Zn utilization by 30%.¹⁷² The air-breathing electrode, especially for rechargeable Zn-air batteries, is arguably the most critical component of the battery as it greatly impacts battery performance and reversibility. The air electrode for the Zn-air battery typically contains a porous carbon matrix with a gas diffusion layer to provide access for oxygen, and a catalyst deposited on the carbon matrix to promote the oxygen reduction (during discharge) for the primary battery, or a bifunctional catalyst that also promotes the oxygen evolution (during charging) reaction for rechargeability. Finally, the aqueous electrolyte is typically KOH, although other alkaline solutions such as NaOH and LiOH are also employed.

The net electrode reactions during cell discharge can be summarized in the following sequence.

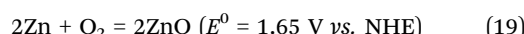
Negative electrode:



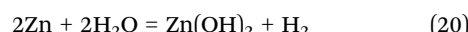
Positive electrode:



Overall cell reaction:



Parasitic corrosion reaction at the negative electrode:



During discharge, oxygen from air is reduced to hydroxyl ions by reaction (18) and zinc is oxidized in accordance with reaction (16) to the soluble zincate ions (*i.e.*, Zn(OH)₄²⁻) until supersaturation in the alkaline electrolyte is reached, after which the zincate ions decompose to form the insoluble product ZnO *via* reaction (17). Zincate ions can also cross over through the semipermeable separator membrane to the positive electrode compartment. This migration decreases the capacity of the Zn-air battery. At the same time, the parasitic reaction (20) between Zn and water takes place on the anode to give off hydrogen gas. This corrosion reaction leads to self-discharge and gradual loss of Zn metal resulting in decreased utilization, efficiency and shelf life. Attempts to minimize the impact of self-discharge by the parasitic reaction (20) and slow down corrosion include alloying Zn with other metals such as Bi, Sn, Pb, Cd, In, Mg, Ni, and Al to stabilize it,²²³ adding surfactants²²⁴ or organic-based corrosion inhibitors to the electrolyte,²²⁵ and using surface modification techniques such as coating the Zn surface with conductive polymers (PMMA, polypyrrole, polyaniline) or inorganic films such as lithium boron oxide (*i.e.*, Li₂O-2B₂O₃) and Al₂O₃,²²⁶ all of which are aimed to modify and improve the electrochemical behavior of the negative electrode and suppress hydrogen evolution.

The alkaline electrolyte employed in Zn-air batteries is susceptible to CO₂ contamination from the air supply, resulting

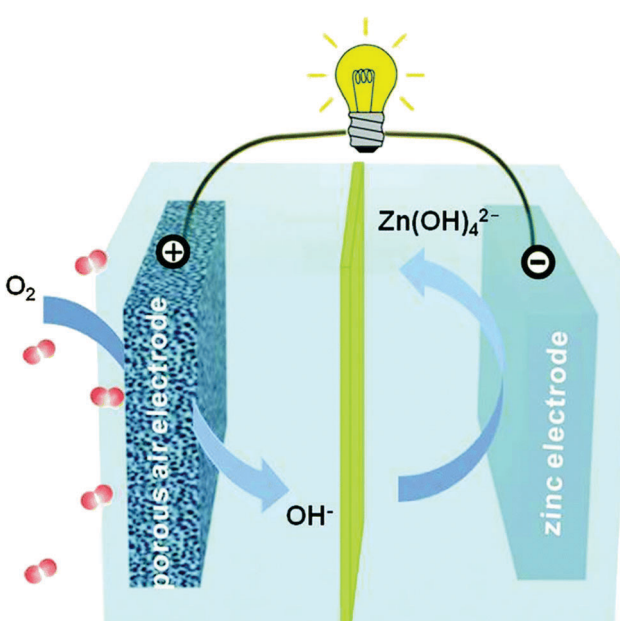


Fig. 25 Schematic cell design of the Zn-air battery, indicating the porous air cathode, Zn anode, and the separator. Reproduced from ref. 175 with permission from The Royal Society of Chemistry.

in the formation of insoluble carbonates at the air electrode and parasitic consumption of the electrolyte, which in turn lowers the electrolyte conductivity and clogs up the pores of the gas diffusion layer on the cathode. Naturally, carbonate formation adversely impacts the performance and service life of the battery. Generally, aqueous NaOH or KOH solutions are employed for Zn-air battery electrolytes. Typically, 7 M (or, 30 wt%) KOH solution is the electrolyte of choice due to its higher ionic conductivity, higher oxygen diffusion coefficient and lower viscosity, but suffers from higher solubility for CO_2 than NaOH solutions. Recent interest in aprotic electrolytes, especially highly conductive and stable ionic liquids, may provide new opportunities to suppress these parasitic reactions. It should be noted, however, that the air electrode structure has been specifically optimized for aqueous electrolytes, and hence, switching to aprotic electrolytes may require a design change.

The theoretical OCV for the Zn-air battery is 1.65 V vs. NHE, as given by the overall cell reaction (19). In practice, however, working cell potentials are typically less than 1.2 V due to activation, ohmic, and concentration polarization losses. In particular, the activation loss at the cathode for ORR largely dominates the overall cell losses, as is also the case for PEM and other types of fuel cells. The ORR reaction occurs at the triple phase boundaries (tpb) where the electrolyte, catalyst impregnated carbon current collector and the gas phase (*i.e.*, air) meet. Wettability of the cathode structure is as important as the access of the gas phase to the tpb sites. In attempts to improve the activation for ORR, both unifunctional (*e.g.*, Pt, Ag, $\gamma\text{-MnO}_2$, doped-perovskites) and bifunctional (*e.g.*, $\text{MnO}_2\text{-Co}_3\text{O}_4$, NiCo_2O_4 , La_2NiO_4) catalysts have been reported.¹⁷⁵

As discussed in some detail (under Section D.1.4.: Metal-air batteries) above, the air electrode is the most critical component of the Zn-air battery, as reviewed recently elsewhere.²¹⁹ Specifically, bifunctionality poses a major challenge to achieving rechargeability of the Zn-air battery. It requires facile catalysis of the ORR during discharge, and fast promotion of the OER during charging. For example, Pt is known to be the best ORR catalyst, but even then, it exhibits significant overpotential due to surface coverage by strongly adsorbed O and OH species, which are found to be stable.¹⁸⁶ Pt also shows poor activity as an OER catalyst due to the coverage of its surface with an oxide layer at the charging potentials in alkaline media. Moreover, Pt is cost-prohibitive to employ widely as a bifunctional electrocatalyst in practical systems. Hence, there has been an intense search for earth-abundant, inexpensive, and high activity bifunctional catalysts for air-breathing electrodes for many electrochemical devices and applications, including all types of metal-air batteries.

Among many electrocatalyst compositions reported in the literature, nanostructured MnO_2 , an inexpensive, nonprecious and abundant material, shows promise and its activity was shown to approach to that of precious metal-based catalysts.¹⁸⁴ Fig. 26 compares the ORR and OER performance of various precious metal electrocatalysts with manganese oxide. As a bifunctional electrocatalyst for ORR and OER, MnO_x clearly performs favorably with precious metal electrocatalysts Pt, Ir and Ru. Other transition metal electrocatalysts have been

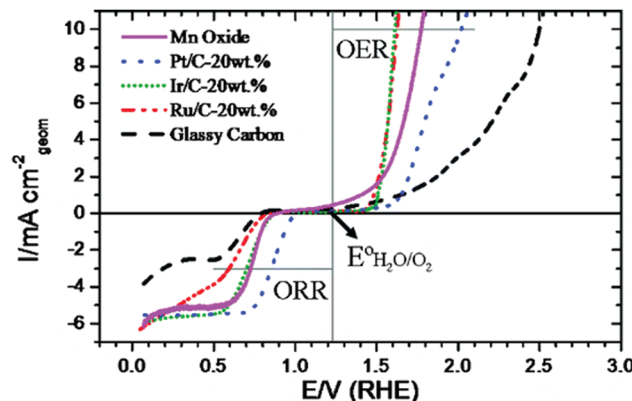


Fig. 26 Comparison of the ORR and OER activities of nanostructured MnO_x with the best precious metal catalysts.¹⁸⁴ Reprinted with permission from (Y. Gorlin and T. F. Jaramillo, A bifunctional nonprecious metal catalyst for oxygen reduction and water oxidation, *J. Am. Chem. Soc.*, 2010, **132**, 13612–13614). Copyright (2018) American Chemical Society.

investigated also. A carbon-free, mesoporous Ni-Fe nitride (Ni_3FeN) bifunctional electrocatalyst was reported to show excellent activity with low overpotential of 0.355 V measured at 10 mA cm^{-2} for OER, and a more positive half-wave potential ($E_{1/2}$) of 0.78 V (vs. RHE) for ORR compared to RuO_2 (0.54 V), but smaller than Pt (0.84 V).²²⁷ Moreover, the Ni_3FeN electrocatalyst showed excellent cycling stability for 310 cycles over 100 hours, with a low voltage-gap of 0.7 V between discharge-charge cycles, and only 3.9% decrease in the round-trip efficiency after 90 cycles. Similarly, 3D carbon- and binder-free air electrodes made of Co_3O_4 nanotubes nominally 300 nm in outside diameter (50 nm inside) and 15 μm in average length were deposited on a stainless steel (SS) mesh current collector and exhibited stable charge and discharge potentials of 2.0 V and 0.98 V, respectively, that remained unchanged after 100 cycles, and retained 94% of the discharge potential and 97% of the charging potential after 600 hours.²²⁸ A similar study employed carbon nanotube-supported CoO and carbon-free NiFe nanoplates as electrocatalytic air-electrodes and reported a high discharge power density of 265 mW cm^{-2} and energy density $>700 \text{ Wh kg}^{-1}$ for the former composition under primary battery mode, while the combination of the two electrocatalysts in the rechargeable battery mode gave a small charge-discharge polarization of only 0.7 V and good cycling stability during 4 to 20 h long charge-discharge cycles at $20\text{--}50 \text{ mA cm}^{-2}$.²²⁹ Doped-perovskites are potential candidates but so far, redox stability of these bifunctional catalysts during repeated cycling has not been satisfactory. More work is needed to develop nonprecious metal-based bifunctional ORR-OER catalysts, not only for achieving the full potential of Zn-air batteries, but also for most fuel cell systems.

Indeed for the Zn-air battery, the overpotential at the Zn anode is relatively small, while the cathode overpotential for the ORR reaction to form hydroxyl ions *via* reaction (16) makes up the major portion of the cell polarization losses. During charging, an external potential significantly larger than the OCV needs to be applied to reverse the cell reactions. Typical voltages

required for the charging process are 2 V or higher. Because of such losses during charging and discharging, typical round trip efficiencies of Zn-air batteries are less than 60%.¹⁷⁵

Designing an electrically rechargeable Zn-air battery pose many challenges. For long and stable cycle life of Zn-air batteries between charge–discharge cycles, it is imperative to avoid irreversible and morphological changes in all battery components. It is well known, however, the Zn metal in the negative electrode undergoes shape changes and dendritic growth during the dissolution/electrodeposition cycles while discharging and charging, respectively. This is shown in Fig. 27, which illustrates the growth of dendrites during electrodeposition of Zn from a ZnCl_2 solution.

Dendrite growth is driven by the inherent instability of the solid/liquid interface during electrodeposition. When the current density is sufficiently high to cause depletion of the solute in the electrolyte near the electrode surface, the local gradient of the solute's chemical potential near the surface becomes positive and drives any microscopic surface feature to grow at a much faster rate than the solid/liquid interface. The roughened surface eventually leads to the formation of dendrites and/or filaments.

The poor cyclability of the Zn-air battery is largely due to the highly soluble zincate ion that is formed *via* reaction (16) during discharge, and not returning to the same surface site to electrodeposit Zn metal during the charging process. Needless to say, this process is very difficult to control. It causes shape change, and eventually leads to the formation of dendrites. As expected, it degrades cell performance and, in extreme

cases, may even lead to electrical shorting of the battery as the dendrites cross over to the positive electrode.

One of the practical solutions to get around the electrical rechargeability problem for Zn-air batteries is mechanical charging. This approach may particularly be attractive for automotive or other transportation applications. Mechanical charging employs the primary Zn-air battery concept and simply involves replacing the Zn-anode assembly mechanically. This simplifies or eliminates many of the technical and materials challenges posed by electrical rechargeability, including using simple unifunctional ORR catalysts, avoiding dendrite and other shape change problems at the negative Zn electrode, and improve battery efficiency. The spent Zn anodes can be recycled in centralized regeneration facilities back to Zn anodes. For transportation applications, this provides attractive opportunities, but require both the distribution of Zn-fuel anodes and collection of spent anodes for regeneration. The vast network of gasoline service stations in place already may be appropriate to serve as Zn anode replacement stations. However, a collection system for spent Zn, and centralized regeneration facilities to recover and regenerate the Zn anodes are required.

D.1.3. Flow batteries. Flow batteries offer many advantages for electrical energy storage including ease of scalability, fast response, high power density that can respond to rapid changes in the load, high round-trip efficiency, and long service life for repeated charge–discharge cycles. Arguably, however, the major advantage of redox flow batteries is their ability to decouple rated power from energy storage capacity, unlike batteries where the two are intimately linked. This advantage provides a considerable latitude and flexibility for designing flow battery systems with respect to the specific requirements of particular applications. In flow batteries, while the power output is controlled by the size of the cell stack, the size of the storage tanks that hold the electrolyte solutions for the redox couple defines its energy capacity. These attributes make flow batteries attractive for grid scale stationary energy storage applications for electricity generation and transmission industries. Although there are a large number of flow battery configurations and chemistries, flow batteries can be grouped in two general types of chemistries from a mechanistic point of view, namely, redox flow batteries (RFB) and hybrid flow batteries (HFB). Prominent members of each group are discussed in the next two sections.

Typical schematic design and operating principle of RFBs is given in Fig. 28, which also illustrates the operating principle of the battery with respect to the generalized redox couples A and B dissolved in appropriate aqueous or organic liquid electrolytes. The electrolytes for the redox couples A and B need not be the same, and in general are different. The anode and cathode compartments (or, anolyte and catholyte, respectively) of the cell is partitioned by an ionically-conducting membrane or a separator that prevents the two electrolytes from mixing with each other, while providing selective transport for the charge carrying ions. The electrochemical process involves pumping of the stored electrolyte from the tank to the flow-through electrodes in the cell stack. During discharge, the chemical energy of the reversible redox couple is converted into

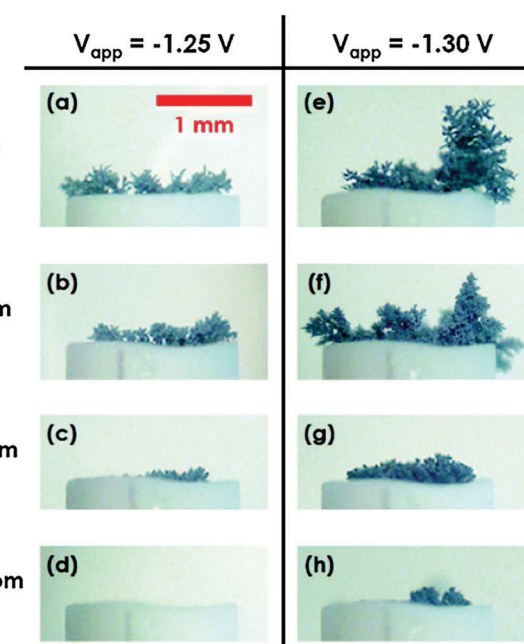


Fig. 27 Optical images of dendritic growth of Zn during electrodeposition at two different potentials (vs. Ag/AgCl) on a Zn wire from a 0.1 M ZnCl_2 electrolyte solution with various concentrations of the additive polyethylene glycol (PEG). Reproduced from ref. 175 with permission from The Royal Society of Chemistry.

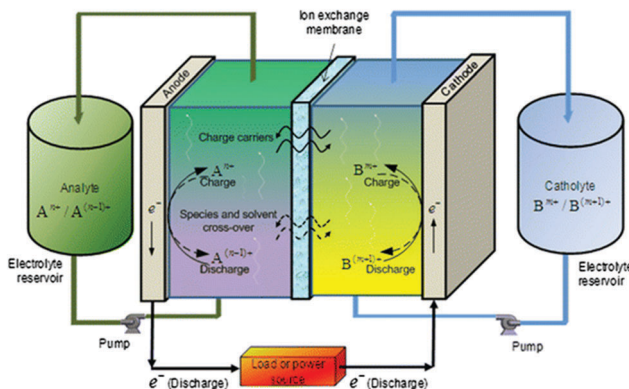


Fig. 28 Basic cell design and operating principle of a redox flow battery based on the generalized redox couples A and B.²³⁰ Republished with permission of The Electrochemical Society, from [M. L. Perry and A. Z. Weber, Advanced redox-flow batteries: a perspective, *J. Electrochem. Soc.*, 2016, **163**, A5064–A5067]; permission conveyed through Copyright Clearance Center, Inc. (2018).

electricity, while during charging the reaction is reversed and the redox couple is regenerated from an external power source such as off-peak power from the grid, or preferably from a renewable source. For this reason, RFBs operate like rechargeable batteries in many ways. As the redox couple can be considered as the fuel, however, RFBs also resemble very much the operation of a fuel cell, and hence, are sometimes referred to as “regenerative fuel cells”.

There has been a wide range of organic and inorganic redox chemistries that were explored and reported in the literature since the advent of RFBs in the early 1970's. Most of this work has been reviewed recently.^{230–235} More recently, a special focus issue of the Journal of The Electrochemical Society was entirely dedicated to most recent progress in RFBs.²³⁶ Among the multitude of redox couples that were studied or developed for RFBs, the standard potentials for some of the more common ones are provided in Fig. 29 that also indicates the electrolyte

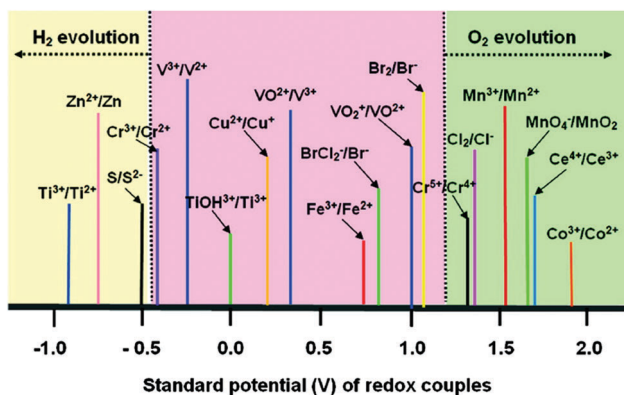


Fig. 29 Standard potentials vs. SHE of selective redox couples in aqueous electrolytes, also indicating the boundaries for H_2 and O_2 evolution on carbon electrodes.²³¹ Reprinted from W. Wang, Q. Luo, B. Li, X. Wei, L. Li and Z. Yang, Recent progress in redox flow battery research and development, *Adv. Funct. Mater.*, 2013, **23**, 970–986, with permission from John Wiley and Sons.

stability boundaries, including the overpotentials for hydrogen and oxygen evolution on carbon electrodes. It is clear that the choices for the appropriate redox couples within this window are not unlimited. Also important is the consideration for the solubility of the redox couples in the electrolytes.

Among the various chemistries, the Fe–Cr redox chemistry with an OCV of 1.18 V was the first truly RFB system that was developed for the space program by NASA in the 1970's. Since then, a diverse range of organic-²³⁷ and inorganic-based^{230–234,236,238} and even organic-inorganic-based^{235,239} RFB chemistries have been reported. The redox cycles, electrochemically active species, and cell potentials of some of the RFB systems are illustrated in Fig. 30. The working potentials of RFBs employing non-aqueous electrolytes are nearly twice higher than those with aqueous electrolytes. Among the multitude of inorganic redox chemistries, the all-vanadium-based RFB system has arguably received the most attention. There has also been renewed interest in Zn–Br₂ RFBs, although other interesting chemistries have also been investigated. Recently, an exceptionally high power density was reported for a quinone–bromine flow battery that exhibited 1 W cm⁻² at peak power,²³⁹ which is more than 75% higher than the best power reported for the vanadium-based RFBs.

However, it is important to note that unlike batteries that store energy in its electrodes, whose mass defines its energy capacity while its kinetic and transport properties in turn govern the battery's power output, RFBs store energy in the

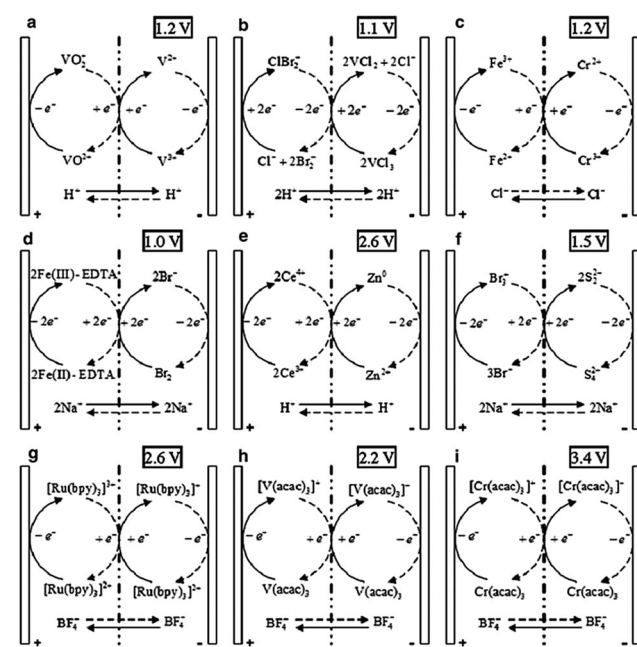


Fig. 30 Redox chemistries, transport pathways and working potentials of select RFBs: (a) all vanadium, (b) vanadium/bromine, (c) iron/chromium, (d) Fe-EDTA/bromine, (e) zinc/cerium, (f) bromine/polysulfide, (g) non-aqueous ruthenium/bipyridine, (h) non-aqueous vanadium/acetlyacetonate, (i) non-aqueous chromium/acetlyacetonate.²³² Reprinted by permission from Springer Nature, A. Z. Weber, M. M. Mench, J. P. Meyers, P. N. Ross, J. T. Gostick and Q. Liu, Redox flow batteries: a review, *J. Appl. Electrochem.*, 2011, **41**, 1137–1164. Copyright (2018).

form of chemical energy of the soluble redox couple held in storage tanks that can easily be scaled up to demand. As the tank sizes are necessarily large, the volumetric energy densities for RFBs are usually less than 25 W h L⁻¹.²³¹

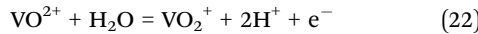
Again in contrast to batteries whose electrodes usually undergo morphological, compositional, structural and molar volume changes, and even sometimes phase transformations during each discharge–charge cycle, which place a heavy toll on the mechanical integrity, stability and useful lifetime of the electrodes, RFB electrodes merely provide an electrochemically active surface for the redox reactions without the burden of physical or chemical changes. This simplifies the operation, and provides high stability, long cycle and service life, low cost, and scalability for RFBs. Using cell performance criteria as well as cell components and materials costs, a techno-economic study of various electrochemical storage technologies suggested that it may be possible to achieve system level energy storage costs in the range \$106–148 per kW h for non-specific aqueous RFBs and \$114–156 per kW h for non-specific nonaqueous RFBs with the right combination of cell and system attributes.^{227,249} For comparison, this study also suggested the future cost of storage for V²⁺/V³⁺ RFBs (or, VRB) at \$118–207 per kW h and Zn/Br₂ RFB at \$103–160 per kW h, which compare favorably with \$193–254 per kW h estimated in a similar manner for Li-ion batteries. It should be noted however, that by comparison, the cost target by the US Department of Energy (DOE) is \$100 per kW h,²⁴¹ while the system level cost estimates for current VRBs are reported to be around \$500–600 per kW h.²⁴² Obviously, there remains much catching up to meet these cost and performance targets for RFB storage.

D.1.3.1. Vanadium redox flow battery. The vanadium redox flow battery (VRB) is arguably the most extensively studied RFB and is a promising technology that was pioneered in the early 1980's. Materials issues and technological challenges for VRBs have been reviewed extensively elsewhere.^{242–245} The all-vanadium redox battery was originally proposed as an attempt to minimize or overcome the chemical cross talk across the separator of the different redox species employed in earlier RFB chemistries. By exploiting vanadium's four oxidation states, the same element (*i.e.*, vanadium) can be employed in both the anolyte and the catholyte, which provides major improvements in reducing cross contamination and extending battery life. The anolyte and the catholyte employed in the VRB consist of soluble V²⁺/V³⁺ and V⁴⁺/V⁵⁺ (in the form of VO²⁺/VO₂⁺), respectively. Both electrolytes are made of aqueous 5 M H₂SO₄ solution separated by an ionically conducting membrane. The half-cell reactions are given by,

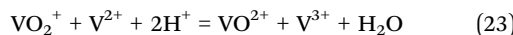
Negative electrode:



Positive electrode:



Overall cell reaction:



The overall cell reaction (23) proceeds to the right during discharge, and to the left during charging. The standard cell potential corresponding to the overall cell reaction is 1.26 V at room temperature, but the OCV for VRB can be as high as 1.6 V when the anolyte and catholyte contain of ~2 M V²⁺ and V⁵⁺, respectively. The four different oxidation states can easily be identified by the color of the solution, such that the color of the V²⁺, V³⁺, V⁴⁺, and V⁵⁺ states correspond to violet, green, blue and light yellow, respectively. For these vanadium electrolytes, an energy density of 25 W h L⁻¹ was reported.²³³ However, the VRB performance is severely impacted by the limited solubility (~2 mol l⁻¹) and stability of vanadium ions in sulfuric acid solutions. This in turn constrains the operating temperature regime between 10 and 40 °C, and requires active heat management, which lowers efficiency. The low fuel utilization due to solubility limitation also impacts performance adversely. When VRB is over-charged, hydrogen and oxygen gas evolution may be observed at the negative and positive electrodes, respectively. But such gas evolution can impact the pH, increase cell resistance by disruption of the liquid electrolyte flow, and oxidation of the carbon at the positive electrode.

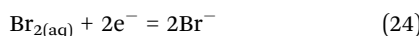
Moreover, the highly corrosive nature of V⁵⁺ makes it necessary to employ more expensive membranes or separators, and containers as well as cell components. Accordingly, the volumetric energy density remains modestly low and does not exceed 25–35 W h L⁻¹, while the round trip efficiencies are around 60–65%. Similarly, the current densities are usually in the 50–80 mA cm⁻² corresponding to a power density below 100 mW cm⁻²,²³⁴ which is much lower than provided typically by PEM fuel cells. High cost and price fluctuations of vanadium and its salts makes it difficult to reliably do economic forecasting, which poses challenges for commercial development of VRBs. For these reasons, a multitude of different redox chemistries based on vanadium and others has been reported in aqueous as well as in non-aqueous electrolytes, but many pose challenges.^{231,233,243,244,246} For example, the cell membrane (typically Nafion) makes up more than 40% of the total cost of the VRB stack.²⁴² Advanced hydrocarbon-based cost-effective membranes that display adequate stability and selectivity may have a major impact on stack costs. Vanadium is expensive so another major cost item is the vanadium-based electrolytes. More cost-effective redox chemistries, including organic redox couples, may help improve both VRB economics as well as increasing the cell operating voltage and hence, the energy density. On the other hand, developing nonaqueous electrolytes with sufficient solubility and stability for the redox couple is challenging. Redox chemistries that can undergo multiple-electron transfer reactions and yield high discharge potentials are highly desirable for advancing RFB technology. Achieving high utilization and effective use of large electrode areas in large-scale storage applications, chemical stability, low cell impedance, and long term recycling capability will significantly improve overall performance.

Commercially, VRB units in the kW to MW scale have been developed for nearly two decades for local grid load-levelling and other storage applications in countries like Japan, Australia,

and South Africa. Although they have fast cycling properties, the capital cost of \$3000–3310 per kW for VRBs are still high, which limits their use for specialized applications and niche markets.¹⁹ A recent study estimated the energy storage cost for a 2 kW (30 kW h) all-vanadium RFB unit to be ~\$100 per kW h, corresponding to \$0.10 per kW h for the cost of stored electricity.²⁴⁵ It should be cautioned, however, this estimate is far too optimistic than more recent estimates of \$500–600 per kW h for current VRB systems,²⁴² and the projected estimates of \$118–207 per kW h for the future cost of storage for VRB.²⁴⁰

D.1.3.2. Zn–Br₂ flow battery. The Zn–Br₂ flow battery is actually a hybrid flow battery (HFB), different from the true definition of a redox flow battery system. As in the case of the Zn–air battery, the Zn–Br₂ flow battery also employs Zn metal as the negative electrode, where Zn is dissolved and electro-deposited during discharge and charge cycles, while the Br₂ gas soluble in the electrolyte is at the positive electrode, where Br[–]/Br₂ conversion takes place. This is schematically illustrated in Fig. 31. The half-cell reactions are given below, and provide an open circuit potential of 1.85 V.

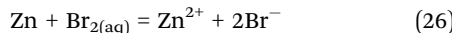
Positive electrode:



Negative electrode:



Overall cell reaction:



Although the Zn–Br₂ battery concept was originally proposed more than century ago, research on this battery system was revived by commercial interest in the 1970's and 1980's. More recently, there has been renewed interest propelled in large part by the urgent need for developing large-scale energy storage systems.

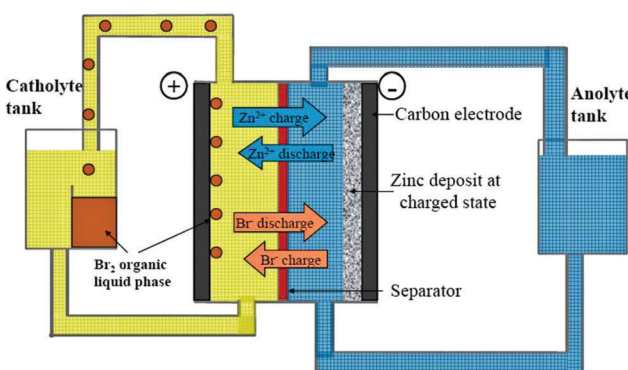


Fig. 31 Schematic flow diagram of the Zn–Br₂ flow battery.²³³ Republished with permission of The Electrochemical Society, from [M. Skylas-Kazacos, M. H. Chakrabarti, S. A. Hajimolana, F. S. Mjalli and M. Saleem, Progress in flow battery research and development, *J. Electrochem. Soc.*, 2011, **158**, R55–R79]; permission conveyed through Copyright Clearance Center, Inc. (2018).

Generally, Zn–Br₂ flow batteries offer higher energy densities (65–75 W h kg^{–1}) than traditional redox flow batteries.²³³ This is in part due to the use of Zn metal anode that eliminates the need for large volumes of anolyte, otherwise required in RFBs. Also, the OCV for Zn–Br₂ flow batteries is relatively higher than those for RFBs. They have storage efficiencies 70% or higher, with high Coulombic and voltage efficiencies of 90% and 85%, respectively.²³³ However, Br₂ gas is highly corrosive and hazardous, and leakage of this gas presents public health and environmental problems. It also has the propensity to cross over the membrane separator due to the unsatisfactory selectivity of the membrane for Br[–] ions that lowers the working cell potential, contributes to polarization losses, and reduces reactant utilization.

Repetitious dissolution and electrodeposition of Zn on the anode during discharge and charge cycles leads to irreversible shape or morphological changes, surface roughening, and dendrite formation (see Fig. 27). The latter can eventually lead to electrical shorting. At the same time, a parasitic reaction (20) between Zn and the aqueous electrolyte takes place to give off hydrogen gas. This corrosion reaction leads to gradual loss of Zn metal resulting in decreased cycle life and in particular, zinc utilization, which can be as low as 40%.²⁴⁷

Furthermore, the slow reduction reaction (24) kinetics of bromine gives rise to polarization losses at the positive electrode, leading to lower voltage efficiency and useful cell potential. High surface area carbon is usually employed to mitigate this loss, but due to gradual consumption of carbon by oxidation, it is difficult to preserve this high surface area over long service life.

Due to these challenging problems, the highly negative reduction potential of the Zn²⁺/Zn couple makes this half-cell reaction less attractive for flow batteries. The performance of Zn–Br₂ flow batteries does not compare favorably with the VRB system, which generally offers better performance. A new single-chamber, membrane-free design concept that eliminates many of the expensive components in the traditional Zn–Br₂ HFB system is reported to operate at >60% energy efficiency for over 1000 cycles, and achieved >90% Coulombic efficiency with a volumetric energy density of 9 W h L^{–1} at a projected cost of <\$100 per kW h.²⁴⁸ The latter figure meets the US DOE cost target for electrical storage systems.²⁴¹ Nevertheless, Zn–Br₂ flow batteries are generally limited to small scale storage applications up to 500 kW h.²³³ However, the economics of Zn–Br₂ batteries are more favorable than VRBs. The average capital cost of \$3000–3310 kW^{–1} for VRBs is significantly more expensive than the capital cost of \$1650–2015 per kW for Zn–Br₂ batteries, which competes favorably well with other electric storage and even generation systems.¹⁹

D.2. Supercapacitors

Supercapacitors are in essence electrochemical capacitors that lie between dielectric capacitors and batteries, and store large amounts of charge. Similar to batteries, their design architecture involves two electrodes separated by an electrolyte and a separator to keep the electrodes from shorting. They offer

exceptionally high capacitances ranging from several 100s of farads per gram up to 1700 F g^{-1} ²⁴⁹ depending on the storage mechanism, choice and microstructure of the electrode materials, and the electrolyte. Although they store charge orders of magnitude more than dielectric capacitors, their relatively low energy density limits their practical use as stand-alone storage systems. However, they can be useful for many high-power but low energy applications. Recent review articles extensively discuss the fundamental aspects of this technology, and report progress in advanced materials for electrodes and electrolytes.^{249–255}

Power densities of supercapacitors are generally around $10\,000 \text{ W kg}^{-1}$, which is 2–3 orders of magnitude higher than batteries, but on the other hand, they have much lower energy densities, typically $<5 \text{ W h kg}^{-1}$.^{252,256} Most electrochemical storage systems exhibit low power densities, partly due to the long diffusion distances and slow mobility of charge carrying ionic species either in the electrolyte and/or in the electrode material as they rely on bulk diffusion for mass transport. In supercapacitors, however, diffusion lengths are significantly shorter (of the order of nanometers) due to the nature of charge storage mechanism occurring at interfaces or surfaces. Hence, they usually have small time constants, or short response times, that allow them to store and release charge at fast rates. Accordingly, they are suitable for fast power drain applications as they can be discharged and charged in a manner of seconds to minutes, as opposed to batteries that typically take several hours to recharge.

Another important feature of supercapacitors is their exceptionally long cycle life, up to several 100 000 cycles.^{250,255} Typically batteries store charge by insertion or chemical reactions at the electrodes, which result in induced strain due to structural or volume changes that limit the battery's cycle life. In supercapacitors, however, charge is stored at interfaces or surfaces and does not involve changes in the microstructure or volume of the electrodes during discharge–charge cycles. Without such strain on the electrodes, these devices have long service and cycle life. These attractive attributes make supercapacitors highly efficient (85–98%) electrical energy storage devices that fill the gap between batteries and conventional capacitors.

Operationally, supercapacitors behave differently from batteries. The cell voltage of an ideal battery remains constant during the charging and discharging operations, while the ratio of the discharge-to-charge potential is a measure of battery's efficiency. In supercapacitors, however, the cell voltage linearly decreases during discharge, and increases linearly during recharging. This is illustrated in Fig. 32. As the voltage of the supercapacitor during charge/discharge cycles varies linearly, it is quite easy to assess the state of charge or discharge of the device as indicated by the dashed line in Fig. 32, while the flat cell voltage makes this assessment significantly more difficult in the case for batteries. Moreover, the energy of a battery is proportional to its cell voltage and defined by the area under the flat discharge potential, while the energy of a supercapacitor is proportional to the square of the cell voltage and

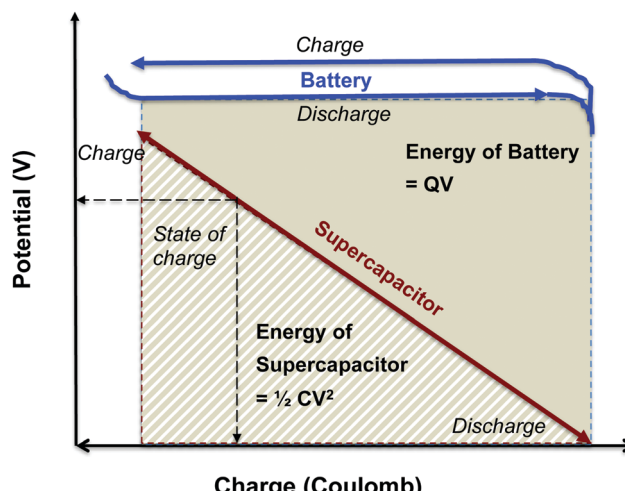


Fig. 32 Schematic comparing the characteristic performance of ideal batteries with supercapacitors. Batteries ideally charge and discharge at constant potential plateaus providing energy at fixed voltage, while the potential of supercapacitors decrease with discharge, which makes it easy to assess the state of charge by simply measuring its voltage, as indicated by the dotted arrows. The energy for the battery is defined by the rectangular shaded area under the discharge plateau, while the striped triangular area defines the energy of the supercapacitor.

defined by the triangular area in Fig. 32, which makes it clear why batteries pack more energy. Recently, an interesting question was raised regarding where batteries end and supercapacitors begin especially for nanostructured devices, where the distinction may sometimes be blurred between the discharge behavior of a nanostructured Li-ion battery from an EDL capacitor,²⁵⁷ and suggested cyclic (*i.e.*, sweep) voltammetry to determine characteristic charge–discharge behavior and device identity.

High power-low energy density property of supercapacitors can complement the low power-high energy density characteristics of batteries. In fact, batteries and supercapacitors are considered as power trains for electric vehicles, where useful life and cost are the most critical factors for implementation.^{257,258} In addition, suitable combinations of these systems may open up new design opportunities to build battery–supercapacitor hybrid storage systems for high-power – high-energy applications.

There are generally two major classes of supercapacitors, those that rely on charge storage at the electrochemical double layer and those that rely on surface redox reactions, or pseudocapacitance. In both cases, capacitance is proportional to the active surface area. Hence, synthesis routes for producing ultrafine particle size, and nanostructuring methods to fabricate 0D, 1D, 2D, and 3D active materials as well as composite structures have been explored for supercapacitor electrodes.^{249,251,254}

Electrochemical double layer (EDL) supercapacitors (also called, ultracapacitors²⁵⁶) have been commercially available for more than three decades, first introduced in Japan by Nippon Electric Corporation in early 1980's, and later by others like Panasonic. However, supercapacitors are expensive systems with an estimated cost of \$10 000–20 000 per kW h,^{24,255} which is more than one order of magnitude more expensive

than Li-ion batteries. Although cost is a major impediment for large-scale industrial deployment of this technology, supercapacitors are employed in select military applications or in niche markets such as power supplies for Airbus A380 aircraft emergency doors.

Supercapacitors that are based on the principle of electrochemical double layer capacitance (also called electrostatic supercapacitors) usually involve high surface area activated or nanostructured carbon electrodes connected to metal current collectors. The charge is stored electrostatically in the electrochemical double layer by adsorbed ions at the carbon/electrolyte interface, and the resulting capacitance is usually in the 5–25 $\mu\text{F cm}^{-2}$ range.²⁵⁹ This type of charge storage does not involve a chemical reaction, and involves only atomic scale distances. Hence, double layer supercapacitors can be charged and discharged in a matter of seconds and have very long cycle lives up to 500 000 cycles with 100% depth of discharge, but suffer from self-discharge with rates up to 14% loss of capacity per month, which limits shelf life.²⁴ Naturally, the higher the surface area, higher is the total cell capacitance. In this regard, work has been focused on nanostructured carbons with increased surface area such as nanospheres, nanofibers and nanorods, nanotubes and mesoporous carbons, foams and aerogels that can offer up to 3000 $\text{m}^2 \text{g}^{-1}$ of active surface area. Electronically conducting 2D materials such as graphene and graphene-oxide offer high surface area (up to 2675 $\text{m}^2 \text{g}^{-1}$), high electronic conductivity of $\sim 10^5 \text{ S cm}^{-1}$ and high thermal conductivity that make them ideal electrode materials for electrochemical energy storage applications.²⁶⁰ Indeed, these graphene-based 2D materials provide high specific capacitance up to 528 F g^{-1} (at 0.3 A g^{-1}) for EDL superconductors, and a high energy density of 12.5 W h kg^{-1} and 110 kW kg^{-1} power density in graphene– MnO_2 composite pseudocapacitive asymmetric supercapacitor cells.²⁴⁸

A typical activated carbon-based supercapacitor has a smooth and near-rectangular shaped cyclic voltammetry curve, reminiscent of conventional capacitors. The capacitance, C , of the double layer is adequately defined by Helmholtz as,

$$C = \varepsilon_r \varepsilon_0 A / d \quad (27)$$

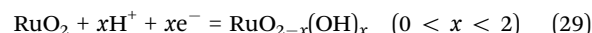
here, A is surface area, d is the thickness of the double layer (or, charge separation distance), ε_r and ε_0 are the dielectric constants of the electrolyte and of free space, respectively. The energy storage capacity varies with the square of cell potential, V , and is given by,

$$E = \frac{1}{2} C V^2 \quad (28)$$

The second category of supercapacitors operates on the principle of pseudocapacitance, which is a Faradaic charge storage mechanism that relies on fast and highly reversible surface or near-surface redox reactions. Many electronically conducting transition metal oxides such as RuO_2 , NiO , MnO_2 , Fe_2O_3 as well as electronically conducting polymers such as polypyrrole and polyaniline exhibit pseudocapacitive behavior and have been developed for supercapacitor applications.^{23,249,252}

Capacitances of transition metal oxides are significantly higher than for carbon-based EDL supercapacitors, as they offer multi-electron transfer during the redox reaction. Among these, hydrated RuO_2 is the most studied, because it has three oxidation states that are accessible within 1.2 V, and offers an energy storage capacity of 240–440 W h kg^{-1} , or a theoretical capacitance of 1200–2200 F g^{-1} ²⁵⁴ although others reported much smaller capacitance values ranging from 600 F g^{-1} ²⁵² to 750 F g^{-1} ,³³ and 900–1300 F g^{-1} ,²⁴⁹ highlighting the importance of the thermal history, synthesis method and processing conditions, particle size, and surface area of RuO_2 on its pseudocapacitive behavior.

The charge storage reaction for RuO_2 in acidic electrolytes involves proton insertion (and de-insertion) and can be expressed by,



Key to the storage mechanism is fast and reversible electron transfer step concurrent with electrosorption of protons on the RuO_2 surface.²⁶² The continuous change of x between the limits 0 and 2 during proton insertion and extraction processes occurring within the 1.2 V window, provides a capacitive storage mechanism operating within the few nanometers of the surface.

Due to the prohibitively high cost of RuO_2 , however, there have been efforts to develop less expensive and abundant materials for surface redox active components for supercapacitor. Among cheaper materials, earth abundant transition metal oxides and other materials ranging from nitrides, carbides, and oxynitrides to polymers have also been reported,^{249–252} including composite electrodes made by mixing high surface area carbon- or graphene-based materials with redox-active pseudocapacitive materials.^{254,261} Inspired by battery design, supercapacitor devices with asymmetric electrode architecture that marry a carbon-based EDL electrode with a redox-based electrode have been reported. A wide range of materials properties and active components, fundamental mechanisms as well as device architectures and performance of such devices have been reviewed elsewhere.^{23,249,251,252,255,263}

In a similar manner reminiscent of Li-ion batteries, the concept of lithium intercalation chemistry has recently gained interest for pseudocapacitive behavior. These hybrid devices with asymmetric cell design typically employ a high surface area activated or nanostructured carbon as the positive electrode and nanostructured transition metal oxide Li^+ intercalation negative electrode materials such as $\text{Li}_4\text{Ti}_5\text{O}_{12}$, MnFe_2O_4 , LiMn_2O_4 ,^{249,264,265} and mesoporous orthorhombic Nb_2O_5 .²⁶⁶ This type of asymmetric device architecture exhibits attributes derived both from batteries and supercapacitors and is capable of storing 5–10 times more energy than the all-carbon based EDL type supercapacitors. Furthermore, as the nanostructured negative electrodes experience no strain during discharge/charge cycling, these hybrid asymmetric supercapacitor devices exhibits not only fast rate capability, *e.g.*, charging in 1 minute at 60C rate for Nb_2O_5 ,²⁶⁶ but also excellent cycling capability for long service life. Indeed, the $\text{C}/\text{Li}_4\text{Ti}_5\text{O}_{12}$ hybrid supercapacitor architecture has maintained more than

95% of its charge after more than 4000 cycles and performed at nearly the same level as a carbon-based EDL supercapacitor.³⁰

As energy stored in supercapacitors is proportional to square of the potential, electrolytes that offer a wide potential stability window are desirable. In that respect, organic electrolytes and ionic liquids provide a significant advantage. They usually increase the cell potential to exceed 3.5 V, and also significantly widen the typical operating temperature window from -30 to 80 °C range to -50 to 100 °C in the case of ionic liquid electrolytes.²⁵⁵ Although most of the early double layer supercapacitors were based on alkaline or acidic aqueous electrolytes, and operated around 1 V, more recent ones employ nonaqueous electrolytes. However, this comes at a price. For example, the typical capacitance of carbon-based materials is $100\text{--}300$ F g⁻¹ in aqueous electrolytes, while this value drops to <150 F g⁻¹ for organic electrolytes,²⁵¹ and $20\text{--}70$ F g⁻¹ for ionic liquids.²³ There is usually a trade off between operating potential of the supercapacitor and the ionic conductivity of its electrolyte. Organic electrolytes and ionic liquids generally have ionic conductivities that are at least one order of magnitude lower than that for aqueous electrolytes, which results in high equivalent series resistance (ESR) indicative of the large ohmic losses that limit the power and energy densities.

With advances in cost reduction and materials improvements, the exceptional ability of supercapacitors for high power at fast charge-discharge rates is expected to open up wide range of applications that may include short duration peak power boost on the grid, peak backup for uninterrupted power sources (UPS), or for storing energy during regenerative braking in transportation vehicles. They can also be combined with batteries to complement the strengths of the two storage systems for improved electrical storage. Such hybrid systems offer the collective advantages of the two systems, increasing cycle life especially when used with Li-ion batteries. However, the costs of both Li-ion batteries ($>\$270$ per kW h) and supercapacitors ($\$10\,000\text{--}20\,000$ per kW h) are currently too high for large-scale deployment for utility grid storage applications.

D.3. Regenerative fuel cells

Fuel cells are essentially electrochemical engines that provide inherently efficient conversion of the chemical energy of fuels directly into electrical energy. They are environmentally friendly and operate under isothermal conditions without generating noise.

There are many types of fuel cells operating at different temperature regimes, and employing different types of liquid or solid electrolytes. They employ different sets of cell components including electrolytes or separators, electrocatalyst and/or electrodes, current collectors, and seals. Fig. 33 illustrates some of the more prominent types of fuel cells and indicates the corresponding net electrode reactions, ionic species transported across the electrolytes, and their operating temperature regimes. The fuel cell types in the figure are arranged from low to high with respect to their typical operating temperatures. Low temperature fuel cells including aqueous alkaline (AFC) and polymer exchange membrane fuel cell (PEMFC) generally employ hydrogen as fuel. Bulk of the global hydrogen

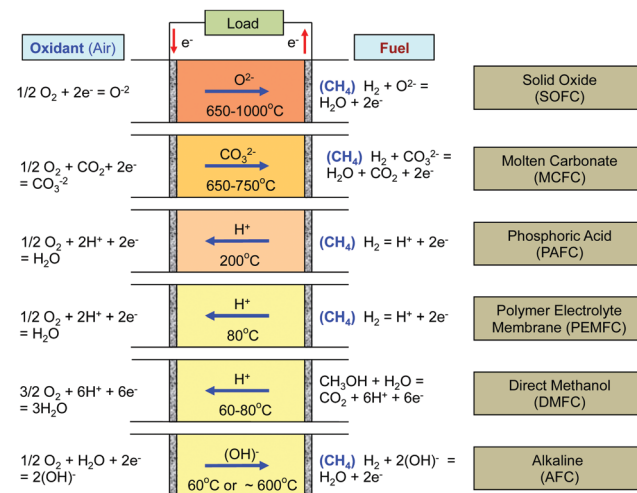


Fig. 33 Schematic depiction of the operating details such as net electrode reactions, transporting ions, and temperature regimes of major types of fuel cells.⁶ Reprinted with permission from (T. M. Gür, Comprehensive Review of Methane Conversion in Solid Oxide Fuel Cells: Prospects for Efficient Electricity Generation from Natural Gas, *Prog. Energy Combust. Sci.*, 2016, **54**, 1–64). Copyright (2018) American Chemical Society.

production is based on steam reforming of methane, which yields a hydrogen stream with small quantities of CO even after extensive separation and purification. Unfortunately, the Pt catalyst at the PEMFC anode is susceptible to CO poisoning leading to degradation of cell performance even at trace levels around several ppm. Also, only precious metals such as Pt are the only catalysts that have sufficient activity, but add considerable cost to PEMFCs.

Due to its high operating temperature, by contrast, solid oxide fuel cells (SOFC) employ non-precious metal and/or earth-abundant oxide electrocatalysts and offer other important advantages^{29,267} including fuel flexibility.²⁶⁸ Indeed, SOFC configurations can successfully utilize and convert gaseous^{269–272} and liquid hydrocarbons,^{273–275} as well as solid fuels⁵ including carbon,^{276–278} biomass,²⁷⁹ and coal²⁸⁰ into electricity. Furthermore, CO, which is an undesirable impurity for PEMFCs is in fact a viable fuel for power generation in SOFCs as recently demonstrated in a 1 kW planar stack.²⁸¹ However, not all fuel cell systems depicted in Fig. 33 can be employed in the electrolysis mode with relative ease or without major cell modification. Only aqueous alkaline (AFC), aqueous polymer exchange membrane (PEMFC), and solid oxide fuel cells (SOFC) have been investigated for regenerative operation for storage purposes.^{282,283}

There is lack of unanimous agreement in the literature on a common terminology to describe the fuel cell systems that can switch their operating mode from electrolyzer to fuel cell and back in the same cell, without modifications. The term “reversible fuel cells” seems to be used more than others especially in the SOFC literature, but the term “regenerative” has also been employed. Yet others especially in the PEMFC community use the terminology in quite a different context such that the regenerative fuel cell refers to a pair of discrete

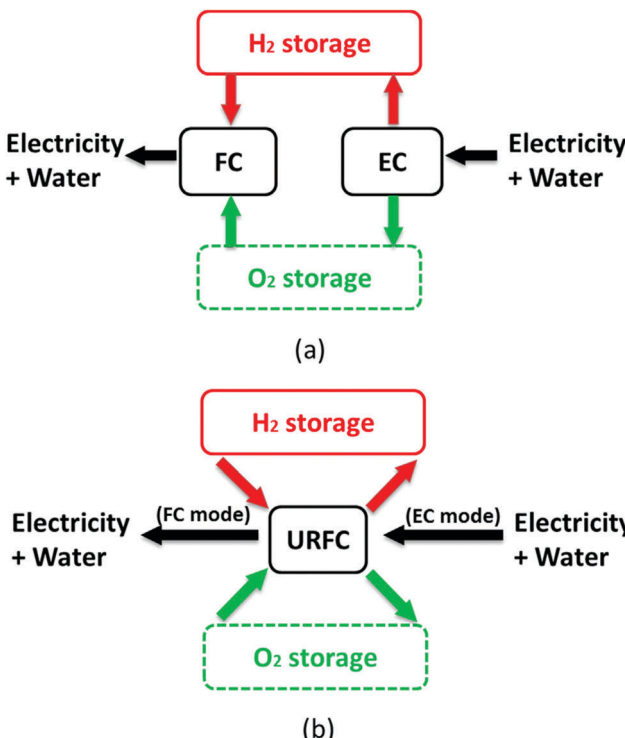


Fig. 34 Electrochemical energy storage systems (a) electrolyzer-fuel cell combination, and (b) regenerative fuel cell (also called, unitized regenerative fuel cell (URFC) in the PEMFC community).²⁹¹ Reprinted from Y. Wang, D. Y. C. Leung, J. Xuan and H. Wang, A review of unitized regenerative fuel cell technologies, part A: unitized regenerative proton exchange membrane fuel cells, *Renewable Sustainable Energy Rev.*, 2016, **65**, 961–977, with permission from Elsevier.

electrolyzer and fuel cell units, where the fuel, *e.g.*, H_2 , and the oxidant O_2 are first produced in the electrolyzer, stored in tanks, and then recombined in a fuel cell to generate electricity when needed, while calling the single cell configuration that can switch without modification between electrolyzer and fuel cell modes, a “unitized” regenerative fuel cell.^{283–285} The latter two concepts are schematically illustrated in Fig. 34. However, labelling separate electrolyzer-fuel cell configurations as “regenerative” may be misleading. More appropriately, this is merely a coupled electrolyzer-fuel cell configuration, and in the view of this author, is unfitting to describe the intended functionality of a bimodal electrolyzer/fuel cell operation. As for the terminology “reversible fuel cells”, reversibility implies well-defined connotations in both kinetics and thermodynamics, and hence, it too may be unsuitable to describe this system. Hence, the term “regenerative fuel cells” is preferred in this article and is used in the context of operating the very same cell sequentially in electrolyzer or fuel cell modes without modifications, as indicated in Fig. 34(b).

Low temperature aqueous fuel cell configurations such as alkaline and PEMFC have been successfully transformed into electrolyzer configurations for distributed production of hydrogen *via* splitting of water, their kinetic reversibility for regenerative operation mode is poor due to large activation overvoltage and scarcity of cost-effective dual action catalysts

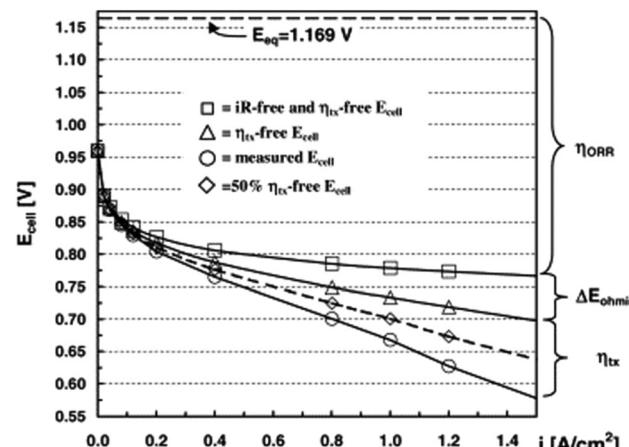


Fig. 35 Electrochemical performance of a H_2/O_2 PEMFC at $80\text{ }^\circ\text{C}$ with 50wt% Pt/C ($0.4/0.4 \text{ mgPt cm}^{-2}$) catalyst-loading and stoichiometric flows of H_2 and O_2 . The figure shows the partitioning of individual voltage losses η_{tx} for mass transport, and η_{ORR} for the oxygen reduction reaction, which makes up more than half of the total voltage losses of the cell.²⁸⁶ Reprinted from H. A. Gasteiger, S. S. Kocha, B. Sompalli and F. T. Wagner, Activity benchmarks and requirements for Pt, Pt-alloy, and non-Pt oxygen reduction catalysts for PEMFCs, *Appl. Catal., B*, 2005, **56**, 9035, with permission from Elsevier.

for the oxygen and hydrogen electrodes. In the acidic environment of the PEMFC, the catalytic electrodes have to perform the difficult tasks of both the water oxidation and oxygen reduction reactions (ORR) at the oxygen electrode, and hydrogen reduction and hydrogen evolution reactions (HER) at the hydrogen electrode. To activate these reactions selectively and at sufficiently fast rates, highly selective specialized bifunctional catalysts are needed. Only precious metals and their oxides such as Pt, RuO_2 , PtIrO_2 , PtRuO_2 have shown satisfactory rates, but still exhibit appreciable overvoltages for these reactions that result in considerable irreversibility. For example, Pt is the preferred electrocatalyst for ORR even though the relatively sluggish ORR kinetics on Pt accounts for nearly 50% of the total cell voltage loss in PEMFCs.²⁸⁶ This is shown in Fig. 35. Furthermore, Pt also lacks sufficient activity for OER.²⁸⁷ The oxides of Ir and Ru exhibit the lowest overvoltages for OER among other electrocatalysts, but they are not suitable for ORR. Similarly for the hydrogen electrode, Pt is the preferred catalysts and economically not suitable for large-scale applications. Nevertheless, no cheaper alternatives have been found yet to compete with the performance of precious metal-based catalysts for regenerative PEMFC electrodes. The prohibitively expensive cost of these catalysts presents a barrier for large-scale energy storage applications.

In alkaline fuel cells, the bifunctional catalyst need to effectively drive the ORR and OER at the oxygen electrode, and hydrogen oxidation (HOR) and water reduction (*i.e.*, hydrogen evolution (HER)) reactions at the hydrogen electrode. Unlike PEMFC, the alkaline environment of AFCs offer catalyst alternatives other than Pt, which is nearly two orders of magnitude less active as hydrogen electrocatalyst in alkaline environments.^{288,289} Various metal hydride

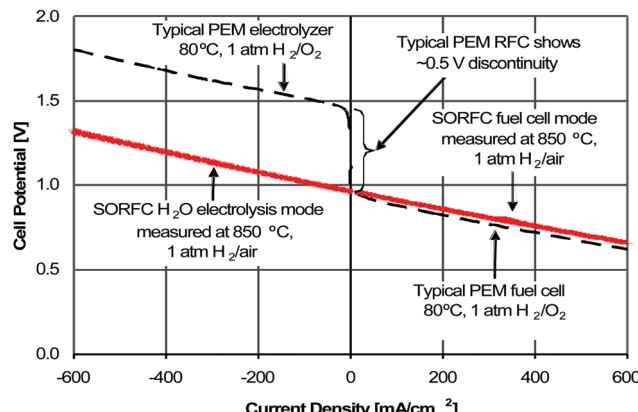


Fig. 36 Comparative performance of regenerative SOFC versus PEMFC, where the latter exhibits a large activation loss around OCV.²⁹² Republished with permission of The Electrochemical Society, from [K. R. Sridhar, J. McElroy, F. Miltitsky, V. Venkataraman and M. C. Williams, Applications and markets for solid oxide regenerative fuel cells, *Proc. – Electrochem. Soc.*, 2005, 295–305]; permission conveyed through Copyright Clearance Center, Inc. (2018).

compositions have been investigated for hydrogen electrodes in alkaline fuel cells, where they serve not only as the catalyst to promote HOR and HER, but also function as medium for hydrogen storage.²⁹⁰ For the oxygen electrode, various transition metal oxides including perovskites, spinels and MnO_x have been reported. The detailed reviews of electrode materials and cell performance of low temperature regenerative cells are provided elsewhere.^{282,283,285,291}

The importance of operating temperature is highlighted in Fig. 36, which compares the nearly reversible SOFC-based regenerative operation at 850 °C with the large irreversibility of PEM-based regenerative fuel cell operation at 80 °C, where the predominant activation loss around the open circuit voltage (OCV) for the PEM fuel cell/electrolyzer operation is evident with a discontinuous drop of nearly 0.5 V. In the case for solid oxide based electrolyzer (SOEC), however, the polarization behavior of the solid oxide-based regenerative cell is continuous and smooth across the OCV indicating sufficiently high kinetic reversibility for steam reduction and hydrogen oxidation reactions at elevated temperatures.^{292,293}

Accordingly, solid oxide fuel cells (SOFC) are offer attractive opportunities for regenerative operation, where off-peak electricity can be employed to store the excess energy for electrolytic production of useful fuels, *e.g.*, hydrogen, which can then be oxidized in the fuel cell mode to generate electricity when demand increases. Many of the regenerative SOFC systems and related materials have been reviewed recently.^{294–296} It should be noted that kinetic reversibility has been limited to the H_2O-H_2 couple only. Compared to liquid fuels such as ethanol ($21\,200 \text{ MJ m}^{-3}$) and gasoline ($34\,560 \text{ MJ m}^{-3}$),²⁹⁷ hydrogen has quite a low volumetric energy density of 11 MJ m^{-3} . Hence, it is highly desirable to achieve reversible cell operation cycling between a liquid hydrocarbon fuel and its oxidation products CO_2 and H_2O in the same regenerative SOFC. But this is yet to be demonstrated. Thermodynamic constraints imposed

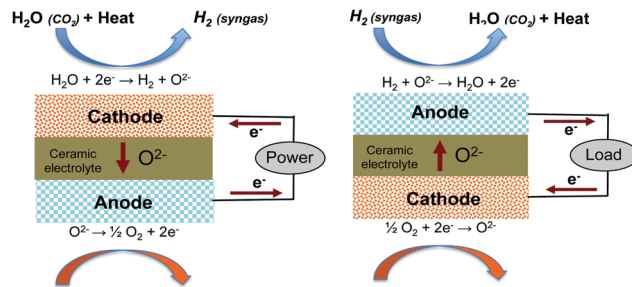


Fig. 37 Schematic depiction of the general concept of a regenerative, or, "reversible" solid oxide fuel cell operating on the H_2/H_2O couple.

by the high operating temperatures of regenerative SOFCs, and the technical challenges posed by kinetics and selectivity issues have hampered progress to achieve electrosynthesis of high value added liquid fuels.

The general concept of regenerative SOFCs has been around for several decades²⁹⁸ and is schematically illustrated in Fig. 37, which shows how the symmetrical SOFC can both store electrical energy in the form of hydrogen fuel by splitting steam, and then re-utilize the hydrogen to generate electricity by oxidizing it back to steam. The details of this high temperature steam electrolysis process have been extensively studied and reported recently, using oxide-ion conducting SOFC configurations,^{294,295,299–302} and also using protonically conducting SOFC configurations.^{303–305}

As expected, this scheme demands stringent requirements for the catalytic electrodes, which must serve both the dual function of anode and cathode depending on whether it is the storage or generation cycle. For example, the electrode in the storage cycle serves a cathode to reduce steam into hydrogen, while in the generation cycle it serves as the anode to oxidize hydrogen back to steam. As expected, this requires that the electrode material possess high catalytic activity for both steam reduction and hydrogen oxidation reactions. Moreover, it has to exhibit exceptional redox stability over a wide range from reducing to oxidizing environments at these elevated temperatures. So there have been intense efforts to achieve high redox stability in catalytic electrodes for regenerative SOFCs, such as the composition $La_{0.75}Sr_{0.25}Cr_{0.5}Mn_{0.5}O_3$ (LSCM).^{293,306} Indeed, kinetically highly reversible fuel cell behavior that remains stable over several cycles and 120 hours has been reported for (LSCM).²⁹³ Other perovskite-based electrodes for reversible operation have been explored for regenerative SOFCs, and reviewed elsewhere.^{295,307} There have also been attempts to both lower the operating temperature of these systems.^{308–310} The performance of a reduced temperature regenerative SOFC cell shown in Fig. 38 features a doped-perovskite ($La_{0.9}Sr_{0.1}Ga_{0.8}Mg_{0.2}O_3$ (LSGM)) electrolyte with Gd-doped ceria (GDC) modified catalytic electrodes, and exhibits a smooth and continuous transition between the power generation (fuel cell) and electrolysis (storage) modes across the OCV and a low value of area specific resistance (ASR) of 0.18 ohm cm^2 at 650°C .³⁰⁷

Various storage systems based on regenerative SOFCs have been proposed.^{296,299,307} These storage schemes involve H_2O ,

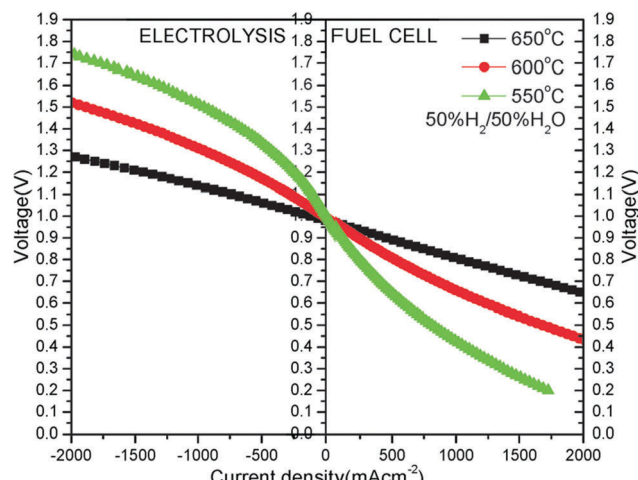


Fig. 38 Performance of a regenerative SOFC operating between 550 and 650 °C. Reproduced from ref. 307 with permission from The Royal Society of Chemistry.

CH_4 , or CO_2 chemistries, and explore combining a regenerative SOFC with solar, wind or other renewable sources. One of these storage concepts involves integration of a 250 MW SOFC stack with underground storage of CO_2 and CH_4 having a total capacity equivalent to 500 GW h of storage. The modeling study of this system concept predicted 70% round-trip system efficiency and electricity storage costs of \$0.03 per kW h.³⁰⁷

A similar concept was proposed for a 100 kW regenerative SOFC with 800 kW h of storage capacity in the form of fuels.²⁹⁶ Assuming 600 °C and 1 atm pressure, 0.2 A cm^{-2} current density, a H/C ratio of 9.9, and a fuel utilization of 65%, this modeling study predicted round-trip conversion efficiency of 74% and storage density of 90 kW h m^{-3} (or, 324 MJ m^{-3}), resulting in a capital cost of \$2.33–3.17 per kW h for storage. The encouraging results may warrant exploring further into similar storage schemes that incorporate regenerative fuel cells with intermittent renewable energy sources.

However, despite their immense potential for large-scale energy storage, major progress in regenerative SOFCs has been slow due to degradation issues largely related to materials instability, kinetic reversibility, and microstructural and architectural integrity of the cells.³¹¹ Delamination is the most common problem observed at the oxygen electrode/electrolyte interface at high current densities due to build up of high internal pressures during oxygen discharge. Also, oxygen evolution at this interface may often lead to chemical changes to form secondary phases increasing ohmic losses. Hence interface engineering and control is important for stable cell operation, and several different approaches to improve stability and cell performance have been suggested including impregnation, exsolution, and impurity control.³¹² Interestingly, it was recently reported that microstructural degradation at the oxygen electrode, which is dominant during electrolysis in particular, could be eliminated by reversibly cycling regenerative SOFCs between fuel cell and electrolysis modes.³⁰¹ Reversible cycling seems to slow down microstructural changes and

mitigate the build up of high internal pressures at the interface and restore sintering. However, detailed mechanisms of degradation (and recovery) processes in these systems are not well understood. Accordingly, the mechanical integrity, chemical stability and microstructure of regenerative SOFC interfaces are important considerations that require further research for development.

E. Chemical energy storage

Our addiction to fossil fuels is no accident, because energy storage in chemical fuels offers the highest energy density, most convenient, cost-effective, scalable, flexible, and highly portable way to store and transport large amounts of energy in the chemical bonds of fuels. The comparative numbers for the gravimetric and volumetric energy densities of fuels provided in Table 6 clearly demonstrate the superiority of chemical storage over electrochemical storage. In this regard, converting excess electrical output or renewable electricity into chemical fuels, which can then be converted efficiently back to electricity naturally holds much merit. However, many fuels including hydrogen and ammonia do not exist in nature in large quantities, so need to be synthesized. This is of course a thermodynamically uphill process and is quite energy intensive.

The starting chemicals to synthesize most fuels are typically hydrogen- and carbon-containing species, and more commonly H_2 (H_2O) and CO. There are multiple synthesis routes to produce high energy density chemical fuels. Examples include electro-reduction of CO_2 to fuels, photoelectrochemical reduction of CO_2 to fuels, solar thermochemical reduction of CO_2 to fuels, and partial oxidation of methane to syngas or coal gasification to syngas, which can then be converted to higher hydrocarbon fuels via Fischer-Tropsch (F-T) process. Some of these routes e.g., gasification to F-T, are mature technologies that have been in use commercially for decades, while others like photoelectrochemical and electrochemical reduction of CO_2 to fuels are under development. Regardless, however, all these process schemes involve carbon-containing species that contribute to global carbon emissions, and fail to provide carbon-free solutions for the electrical energy enterprise. Accordingly, carbon-containing synthesis schemes are only briefly covered in this article. Also, photocatalytic splitting of water to produce hydrogen is not reviewed here.

Chemical storage topics covered in this section focuses on converting excess or off-peak electrical energy preferably from renewable resources such as solar and wind into environmentally benign, carbon-free, high energy density transportable fuels via electrosynthesis. In this regard, electrochemical production of hydrogen and ammonia are presented and reviewed below. It is important to note that chemical energy storage in hydrogen and ammonia also offers environmentally benign solutions towards decarbonization of the global energy systems. The proponents of hydrogen see this energy carrier on which to build a carbon-free energy economy,^{313,314} while opponents of hydrogen-based economy have long argued that the heavy

energy and carbon emission penalties paid during hydrogen production, the difficulties in handling and transporting hydrogen, and lack of earth-abundant cost-effective materials that can provide >6.5 wt% H₂ for long term storage are major barriers for practical adoption and implementation at a wider scale. Ammonia as an environmentally benign energy carrier has recently gained interest as a viable option for chemical storage, but electrosynthesis of this chemical faces similar obstacles to be overcome.

E.1. Chemical storage in hydrogen

Exceptionally high gravimetric energy density of hydrogen makes it an ideal candidate for chemical storage and a carbon-free energy carrier. As hydrogen does not occur naturally, it is produced largely from natural gas. More than 75% of the industrial hydrogen production in the world is *via* steam reforming of methane, which is a highly efficient (60–85%) and mature technology but is cost effective only when it is produced centrally and at large scale. This process is also highly energy intensive and consumes nearly 2% of the world's energy output. Most of the industrially produced hydrogen is utilized on-site (*i.e.*, captive hydrogen) for chemical, catalytic and petrochemical processes. A small fraction is supplied to the market (*i.e.*, merchant hydrogen) for other industrial processes or uses.

Hydrogen is an environmentally benign energy carrier with the highest gravimetric energy density of 142 MJ (HHV) kg⁻¹ among the common liquid fuels (see Table 6), but it has also one of the lowest volumetric energy densities of about 10 MJ l⁻¹ in its gaseous form at atmospheric pressure and room temperature. The latter property poses challenges in wider utilization of hydrogen for portable or mobile power generation applications, including transportation. Although there have been intense research activity in the last two decades to develop effective storage medium or materials, cost effective solutions for extended use capacity has been elusive. Among the options, mechanical storage *via* cryogenic liquefaction as well as compression are mature technologies but both have practical challenges including safety, cost, and energy penalties.³¹⁵ For

example, a pressurized hydrogen tank offers a system energy density of less than 2 MJ l⁻¹ at 350 bar and 3.4 MJ l⁻¹ at 700 bar. Compared to other fuels listed in Table 6, the energy density of pressurized tank storage is quite limited.

Storing hydrogen in solids *via* chemical or physical routes has many advantages, but also often have low gravimetric capacities that add extra weight to the mobile application. Many also have either undesirable thermodynamics that limit reversibility of storing and releasing hydrogen and/or slow kinetics that lead to long charge and/or discharge times. Many storage materials or methods have been under development but none has reached commercial maturity.

Many materials can store hydrogen by absorption or hydride formation (*e.g.*, Pd and its alloys), adsorption (*e.g.*, carbon nanostructures), or chemical reactions (*e.g.*, MgH₂, NaBH₄, alanates). An in-depth review of hydrogen storage materials and technologies fall outside the scope of the present article. The interested reader is referred to detailed reviews published in the literature.^{316–318} Gravimetric capacities of various materials for hydrogen storage are provided in Fig. 39 for comparison purposes.

Storage and transportation of hydrogen is costly and inefficient. That is why distributed generation may be critically important for an effective carbon-free energy economy for the future. In this regard, water electrolysis driven by renewable electricity presents attractive opportunities for local production of hydrogen.

Currently, low temperature electrolysis of water is the only commercial technology available for distributed generation of hydrogen. Thermodynamically, water splitting is an uphill process and hence, is highly energy intensive. It needs to overcome the thermodynamic barrier as presented in Fig. 40, which indicates the thermal ($T\Delta S$), electrical (ΔG), and total energy (ΔH) demands to water (and steam) electrolysis as a function of temperature. For this, an external bias that opposes and is greater than the theoretical open circuit voltage (1.23 V at 298 K) must be applied to the electrolysis cell in order to break the O-H bonds in water, permitting the formation of the H-H bond. A large fraction of the electricity used in conventional water electrolyzers is consumed merely to overcome this

Table 6 Gravimetric and volumetric energy densities of liquid fuels compared with select storage systems. Most values are from ref. 297

| Fuel | Gravimetric energy density (W h kg ⁻¹) | Volumetric energy density (W h l ⁻¹) |
|-------------------------------|--|--|
| Gasoline | 12 330 | 9.060 |
| Diesel | 12 700 | 10 700 |
| Propane | 12 870 | 7490 |
| Butane | 12 700 | 7190 |
| Ethanol | 7490 | 5890 |
| Methanol | 5620 | 4470 |
| Liq. hydrogen | 33 570 | 2200 |
| Liq. ammonia | 5170 | 3750 |
| Batteries | | |
| Li-Ion battery ^a | 150–210 | 450 |
| Ni-Metal hydride ^a | 70 | 220 |
| Zn-Air battery ^a | 300 | 240 |

^a Practical values.

Weight % H₂

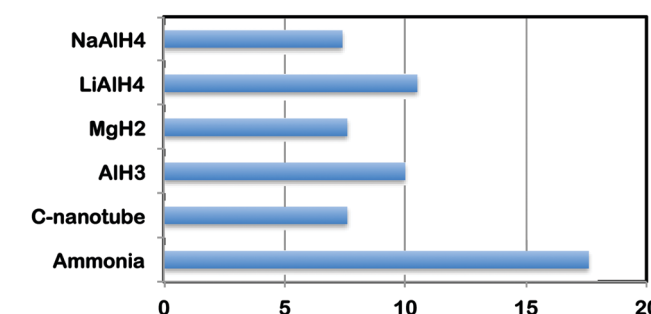


Fig. 39 Gravimetric storage capacities of various materials given in weight percent hydrogen.

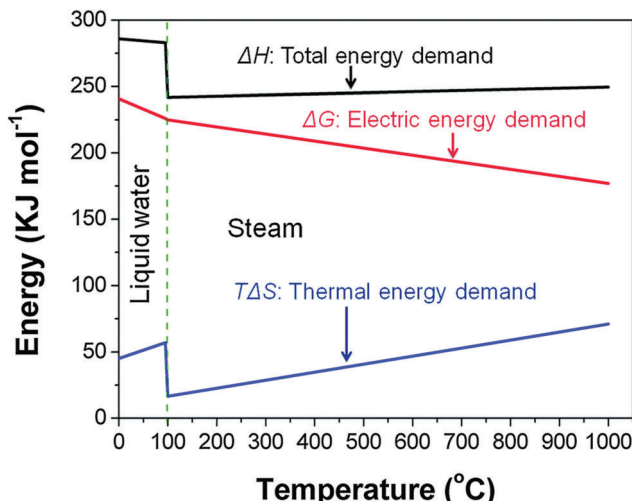


Fig. 40 Energy demands for water electrolysis as a function of temperature. Reproduced from ref. 303 with permission from The Royal Society of Chemistry.

potential barrier, which reduces efficiency. Dissociation of water to hydrogen is also an endothermic reaction with $\Delta H = +286 \text{ kJ mol}^{-1}$ at room temperature, which corresponds to the thermoneutral potential of 1.48 V. Clearly, this is greater than the theoretical cell potential of 1.23 V. So if no external heat source is employed, the applied potential required to split water has to be greater than 1.48 V. In practice the externally imposed potential is much higher, up to 2.0 V or above, depending on the type of water electrolyzer and the electrolyte used.

By Faraday's law, the rate of hydrogen production is linearly related to the amount of current supplied to the electrolytic cell, as electrolysis generally occurs under nearly Faradaic conditions with close to 100% coulombic efficiency. Hence, the cost of electricity is an important consideration in assessing the economics of electrolytic hydrogen production. The strong dependence of the production cost for electrolytic hydrogen on the cost of electricity is shown in Fig. 41. The high sensitivity to the cost of electricity renders electrolysis a costly process for distributed hydrogen production. An earlier study concluded that the electricity costs must be below the \$0.045–0.055 per kWh

range in order to produce hydrogen using current commercial electrolyzers at a cost of \$3 per kg of H₂ to stay competitive with gasoline.³¹⁹ Although the cost of electricity cost is a major contributor to electrolytic hydrogen production, the origin of electricity for electrolysis is another important consideration. Most of the electricity from the grid is derived from fossil fuels, and hence yield greenhouse gas emissions. So hydrogen production by electrolysis is best suited for renewable power such as wind and solar in order to avoid upstream carbon emissions.

The appeal of hydrogen as a viable and sustainable energy carrier drives commercial interest in high temperature electrolyzers and several development efforts are being pursued, but the technology is far from being ready for commercialization. Besides the technical challenges, the economics of hydrogen production in SOECs is very likely to be considerably more expensive than the average cost target for wind-based low temperature water electrolysis of \$3.10 per kg of hydrogen for central and \$3.70 per kg for distributed production recently set forth in 2015 by the US Department of Energy (DOE)³²⁰ based on the 2011 NREL analysis.³²¹ It should be noted that the cost estimates for high temperature SOECs may be necessarily inaccurate as the cost models are largely based on laboratory scale results. In fact, US DOE Office of EERE possibly refrained from providing a cost target for hydrogen production in SOECs³²² most likely for the reason that the reported cost estimates may be somewhat premature.

E.1.1. Hydrogen production by water or steam electrolysis.

Combining water or steam electrolysis with renewable energy provides an integrated distributed storage system whereby hydrogen produced can be either utilized for end use in a fuel cell or for chemical energy storage. Unlike steam electrolysis still under development, hydrogen produced *via* the electrolysis of water is a mature technology and has been available commercially for many decades. Water electrolyzers serve relatively a small market, where access to merchant hydrogen is limited by geography or other reasons. The operational parameters, cell performance and materials, and economics of water electrolyzers for hydrogen production have been reviewed recently.^{319,323–326} These commercial electrolyzers typically operate at 70–80 °C with an imposed potential of 1.75–2.0 V and have electrical conversion efficiencies in the range of 60–70%. But unless this electricity for electrolysis is derived from renewable sources, the primary efficiencies of commercial electrolyzers drop to around 20–25% when electricity from conventional power is used. Also, the cost of hydrogen production using electrolyzers cannot compete with the large-scale centralized production of hydrogen *via* steam reforming of natural gas, which provides more than 75% of the world's industrial hydrogen production. Capital costs for commercial water electrolyzers reported in the literature range from >\$3000 per kW for small (<50 kW) alkaline electrolyzers to \$400–700 per kW for larger plants with life times from 5 to 10 years.³²⁶ A more recent review estimates the capital costs for commercial alkaline electrolyzers in the range of \$1200–1500 per kW and polymer electrolyte-based electrolyzers in the \$1300–2000 per kW range.³²⁷ Driven in

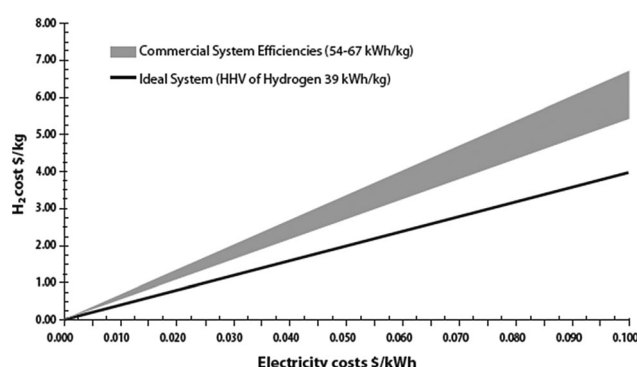


Fig. 41 Cost of hydrogen via electrolysis varies linearly with cost of electricity, according to a 2004 NREL study.³¹⁹

Table 7 Typical structure and operational characteristics of the four major classes of electrolyzers for hydrogen production named after their electrolytes, where PEM stands for polymer exchange membrane, and SOEC denotes solid oxide electrolysis cell. (YSZ: yttria-stabilized zirconia, BYZ: yttria-doped barium zirconate)

| | Alkaline | Acid | PEM | SOEC |
|-----------------------|--|--|---|---|
| Electrolyte | NaOH | H ₂ SO ₄ | Polymer (Nafion™) | Ceramic (YSZ, BYZ) |
| Carrier ion | KOH OH ⁻ | H ₃ PO ₄ H ⁺ | H ⁺ | O ²⁻ (YSZ) H ⁺ (BYZ) |
| Electrodes | Ni | Pt/C, IrO ₂ | Pt/C, IrO ₂ | Ni, ceramic |
| Operating temperature | 80 °C | 150 °C | 80 °C | 600–900 °C |
| Strengths | Cheap materials Commercial technology | High activity electrodes | High activity electrodes Commercial technology | Cheap materials Lower barrier to split H ₂ O Higher efficiency |
| Short-comings | High barrier to split H ₂ O Lower efficiency | High barrier to split H ₂ O Expensive electrodes Cell durability Developing technology | High barrier to split H ₂ O Expensive electrodes Hydrogen crossover Cell durability | High operating temperature Materials stability Fabrication costs Developing technology |

large part by cost concerns, water electrolyzers have been generally employed for niche markets, for specialized applications in remote locations or confined spaces, or where abundant hydroelectric power or other renewables are readily available.

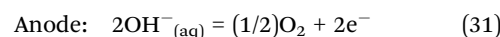
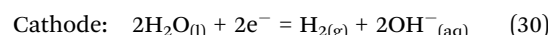
Electrolysis cells are usually named after their electrolytes. Typical materials, operating temperatures, advantages and disadvantages of the four major classes of water or steam electrolysis technologies are summarized in Table 7. The simplicity and modular design of electrolytic cells provide many advantages including ease in scaling up simply by adding units, and suitability for distributed chemical storage, which is particularly attractive for intermittent renewable sources that usually generate electricity at remote locations far from the populated areas or points-of-use. Electricity generated from intermittent renewable energy sources makes water (or, steam) electrolysis an attractive way to store high quality energy in a clean and environmentally benign fuel such as hydrogen. Indeed, hydrogen storage has gained renewed interest for distributed energy storage driven by the recent and rapid expansion of solar and wind energy.³²⁸

Commercial electrolyzers for water electrolysis are similar in design to fuel cells, and compose of an anode and a cathode immersed in an electrolyte and separated by an inert separator, diaphragm, or an ionically conducting polymer exchange membrane (PEM) such as Nafion®. Most commercial electrolyzers operate at temperatures <80 °C and generally employ alkaline electrolytes such as NaOH and KOH solutions, which allows the use of inexpensive catalysts such as Ni in contrast to electrolysis in acidic electrolytes that would necessitate the use of precious metal catalysts such as Pt. However, they are slightly less efficient (60–70%) when compared to efficiencies of PEM-based acidic electrolyzers in the 65–82% range.³²⁵ PEM-based acid electrolyzers can sustain high current densities and high production rates, high voltage efficiencies, and also higher purity of product gases H₂ and O₂ due to minimal crossover across the solid polymer electrolyte membrane. Unlike alkaline electrolytes that provide better stability and longer service life, however, acidic electrolytes also cause severe corrosion problems that require more expensive materials for cell components. On the other hand, alkaline electrolyte-based electrolyzers suffer from low current densities leading to low

hydrogen production rates, large ohmic losses up to 25% of total cell loss across the electrolyte and the separator, low ionic conductivity of the electrolyte, crossover of product gases across the porous separator or diaphragm that yields impure products, and unwanted transport of hydrogen gas into the anode chamber, which can accumulate beyond the explosion limit of >4% H₂ in O₂ to cause safety concerns. Despite these disadvantages, commercial alkaline electrolyzers seem to be the preferred product compared to acid electrolyzers, due mostly to the cost of the precious metal catalysts used at the electrodes, cell stability and service life concerns.

In order to improve upon the low current densities of alkaline electrolyzers, high temperature alkaline electrolyzers that typically operate at temperatures 150–200 °C have gained interest.²⁹⁹ As expected, these cells need to operate under pressure in order to minimize evaporation losses and accommodate the high operating temperatures of the aqueous electrolyte. High temperature operation lowers cell voltage from the typical >2.0 V to <1.5 V, which improves efficiency, and reduces ohmic losses due to improved electrolyte conductivity. High pressure operation on the other hand, reduces size of gas bubbles evolving at the gas diffusion electrodes that result in less disruptions to current stability and distribution, but at the same time may cause bubbles to block the micropores of the electrodes and render those sites ineffective and inactive. Hence, the operating cell pressure should be kept lower than for steam condensation, but high enough to avoid evaporation losses from the electrolyte. In an alkaline electrolyzer using aqueous KOH immobilized in mesoporous SrTiO₃ and operating at 250 °C under 42 bar pressure, high current densities of 1000 mA cm⁻² at 1.5 V and 2000 mA cm⁻² at 1.75 V were achieved using Ni foam gas diffusion electrodes.³³⁰

Conceptual cell design and half-cell reactions for water electrolysis are shown in Fig. 42. The thermodynamic potential corresponding to the half-cell reactions is 1.23 V, but cell polarization losses require significantly higher voltages around 1.8–2.2 V to be applied externally to drive the water splitting reaction. The half-cell reactions in alkaline electrolyte are,



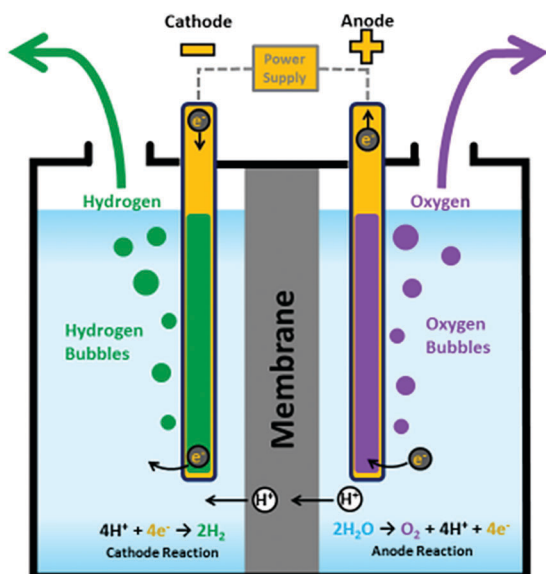
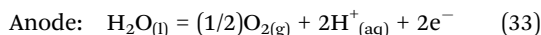
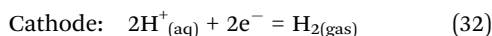


Fig. 42 Hydrogen production via water electrolysis.³²⁹

while in acidic electrolytes, the corresponding reactions are,



Regardless of the type of electrolysis cell, splitting of water requires highly active electrocatalytic electrodes for the hydrogen evolution reactions (HER) (30) and (32), and the oxygen evolution reactions (OER) (31) and (33). HER is a 2-electron proton-coupled reaction while OER involves 4 electrons. As expected, the activation barrier or overpotential for OER is significantly higher than for HER. Lowering the barriers require the use of effective catalysts. High efficiency, low cost and long durability or lifetime considerations are important criteria for the desirable catalyst. Platinum is an excellent catalyst for both HER and OER but is prohibitively expensive, but still serves as the benchmark for HER when comparing the performance of other electrocatalysts. For OER, IrO_2 and RuO_2 are the best electrocatalysts with the lowest overpotentials in both acidic and alkaline electrolytes compared to others. They are also excellent electronic conductors and can serve as their current collectors as well. Under high anodic potentials, however, both RuO_2 and to a lesser extent IrO_2 suffer from stability problems in both electrolytes because they oxidize to RuO_4 and IrO_3 , respectively, and form soluble products that dissolve in the electrolytes. So besides cost, these precious metal catalysts have durability or lifetime problems.

There has been an intense search for efficient and cheap catalysts for HER and OER that are made of non-precious metals, as well as the oxides, sulfides, phosphides and carbides of earth abundant materials.^{331,332} Many strategies for catalyst development have been employed including advanced processing techniques such as nanostructuring to fabricate metastable structures, core-shell structures, and engineered surfaces and interfaces,³³³ atomic layer deposition for atomic

scale control of nanostructure, doping and morphology of the catalyst,^{334–336} as well as theoretical simulations and computational tools coupled to experimentation for rational design.^{332,337} One of the key parameters that can guide catalyst design efforts is the adsorption free energy of the electroactive species on the catalyst surface. In accordance with the Sabatier principle, the surface binding energy of the species should be moderate for a good catalyst, and neither too weak (then adsorption becomes rate limiting) nor too strong (then desorption becomes rate limiting). When the free energy of adsorption of the electroactive species is plotted against a performance parameter such as overpotential or exchange current density on various electrolytic surfaces, the resulting “volcano” plots present a descriptive tool to observe the trends and select the electrocatalyst candidates exhibiting the best performance criteria, *e.g.*, the lowest overpotential, or the highest exchange current density. Insights for the selection process of several catalyst systems for various electrochemical reactions have been presented recently.³³⁷ For example, guided by the results of an earlier theoretical study,³³⁸ nanoparticles of earth-abundant MoS_2 prepared in a double gyroid morphology with 3-dimensional mesoporous microstructure in order to maximize the edge-site density have shown great promise for HER with overpotentials only 150–200 mV away from Pt.³³⁹ Similarly, nanostructured MnO_2 has shown great promise as an earth abundant effective catalyst for OER.¹⁸⁴ The OER activity of MnO_2 is already presented in Fig. 26, which compares MnO_2 quite favorably with the precious metal OER catalysts Ir, Pt and Ru. A detailed review of many classes of prospective OER catalyst materials is provided elsewhere.³³¹

It is also possible to achieve water splitting using steam in high temperature solid oxide electrolysis cells (SOEC) operating at 600–800 °C and employing a ceramic membranes that exhibit selective ion transport for either O^{2-} such as yttria-stabilized zirconia (YSZ) and gadolinia doped ceria (GDC), or for H^{+} as in proton-conducting perovskites such as yttria-doped barium zirconate (BYZ) and yttria-doped barium cerate (BYC), as also indicated in Table 7. In the case of steam electrolysis using an oxide-ion-conducting ceramic electrolytic cell, steam is fed on the cathode side and air is in the anode compartment. By the use of an external power, oxygen is abstracted from steam at the cathode and transported across the solid electrolyte and discharged at the anode as molecular oxygen into the air compartment. In the meantime, hydrogen is produced at the cathode. In the case of steam electrolysis using a proton-conducting ceramic electrolyte, steam is fed on the anode side and the hydrogen evolves at the cathode. But the steam decomposition reaction is thermodynamically uphill and requires an external power source and the associated cost of electricity.

Operating the SOEC at elevated temperatures lowers the electrical energy requirement (see Fig. 40). It also increases the kinetic and transport rates significantly. Accordingly, non-precious metal or oxide catalytic electrodes can be employed at high temperature SOECs. Furthermore, a thermodynamic modeling study showed that the difference between the energy and exergy efficiencies for SOECs operating at elevated

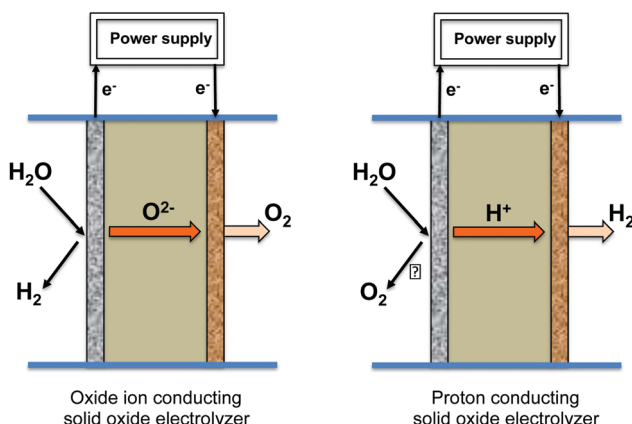


Fig. 43 Schematic cell configurations for steam electrolysis and directions of ionic transport in solid oxide electrolyzers featuring (a) oxide ion conducting solid electrolyte, and (b) proton conducting solid electrolyte.

temperatures is quite small, unlike electrolyzers running at low temperatures, due to the fact that at high temperatures, the energy quality is high and also the thermal energy input constitutes a relatively small portion of the total energy requirement.³⁴⁰

Steam electrolysis at elevated temperatures can be achieved either by using an oxide-ion conducting solid oxide electrolyte^{299,300} such as yttria-stabilized zirconia, or by a proton-conducting solid oxide electrolyte^{300,303,341} such as yttria-doped barium zirconate (Y₂BaZrO₅, or BYZ) or cerate (Y₂BaCeO₅, or BYC). These electrolytes are made dense and impervious to block chemical shorting, and allow only mobile ionic species O²⁻ and H⁺ to transport selectively through the crystal lattice of the ceramic membrane. Steam electrolysis schemes for the two types of ceramic electrolytes are illustrated in Fig. 43, which also indicate the direction of ionic transport through the solid oxide electrolytes.

The impervious ceramic electrolyte membrane physically separates the anode and cathode compartments and eliminates chemical shorting or mixing of the product gases. This results in high purity hydrogen production. The electrolyte transport mechanisms are respectively, lattice diffusion *via* oxygen vacancies in the oxide-ion conducting fluorite structure solid oxide electrolytes, and Grotthuss mechanism for proton transport in the perovskite-type protonically conducting solid oxide electrolytes. As the activation energy for proton transport in perovskites is considerably smaller than that for oxide ion transport in fluorites, protonic SOECs can be operated a lower temperatures (<600 °C) than the oxide ion-based SOECs, which typically operate at 700–1000 °C. The major elements of cell voltage loss in SOECs are ohmic losses due to finite ionic conductivity of the electrolyte and activation losses at the electrodes due to slow kinetics and transport. To overcome these difficulties, progress has been made at the laboratory scale in advanced catalytic electrodes and cell architectures, which have been reviewed in detail elsewhere.^{299,300,303}

E.1.2. Spontaneous hydrogen and electricity co-generation. As emphasized in Fig. 41, it is desirable to minimize or preferably avoid the electrical costs for electrolysis. Indeed, the

nascent steam-carbon fuel cell (SCFC) concept proposed earlier³⁴² aims to accomplish hydrogen production from steam without the need for external power, and in fact, spontaneously generates electricity and hydrogen at the same time. Essentially, the SCFC concept represents the steam gasification reaction of carbon carried out electrochemically. This is accomplished by replacing the more expensive electrical energy requirement for electrolysis (see Fig. 41) with much cheaper chemical energy, which provides a sufficiently low oxygen activity to reverse the sign of the activation barrier and drive the steam decomposition reaction downhill. As the equilibrium (chemical) potential for oxygen in the carbon–oxygen system is significantly lower than that for the H₂–H₂O system, the uphill barrier disappears and the gradient becomes downhill, allowing the generation of carbon-free hydrogen and electricity simultaneously. This concept is illustrated in Fig. 44, and shows the uphill barrier of about –0.9 V for steam electrolysis at elevated temperatures of 800–1000 °C, which reverses its sign by the presence of solid carbon at the anode and becoming instead a downhill potential gradient with a driving force of about +0.5 V that allows the cell to operate in a fuel cell mode, while co-producing hydrogen and electricity spontaneously.^{343,344} The driving force is governed by the prevalent H₂/H₂O and CO/CO₂ ratios at the cathode and anode compartments, respectively. The SCFC concept utilizes a typical solid oxide fuel cell design and materials, where the only difference is that air is replaced with carbon fuel at the anode. The impervious solid oxide electrolyte membrane serving as a physical barrier between the anode and cathode gas environments allows the production of carbon-free hydrogen, which is critically useful for PEMFC applications. Moreover at high fuel utilization, the anode product gas is highly concentrated in CO₂ with only a small amount unreacted CO in the balance, and hence, nearly ready for capture.

The SCFC concept accomplishes the steam gasification reaction of carbon inside a fuel cell with discrete anode and cathode reaction gas streams, producing power and carbon-free hydrogen in the process. Fig. 45 presents the experimental results for simultaneous and spontaneous co-generation of hydrogen and electrical energy in a button-type SCFC employing a yttria-stabilized zirconia (YSZ) solid electrolyte at 850 °C with activated carbon at the anode and steam at the cathode, which provided a downhill driving force of 0.56 V open circuit potential that enabled achieving steam electrolysis in a fuel cell mode.

A similar concept, this time using reducing gases such as methane and CO at the anode, was also proposed.³⁴⁶ As expected, the placement of these gaseous fuels at the anode greatly reduces the energy barrier. However, the open circuit potentials reported in this study were less than 90 mV, not sufficiently large enough to spontaneously drive the steam electrolysis reaction. Hence, electricity from an external power, albeit less than otherwise needed for conventional steam electrolysis cells, is still required to drive the electrolysis reaction and produce hydrogen using this configuration.

E.2. Chemical storage in ammonia

Ammonia is an effective energy carrier with a high-energy density. For comparison, liquid ammonia provides 13.7 MJ l⁻¹

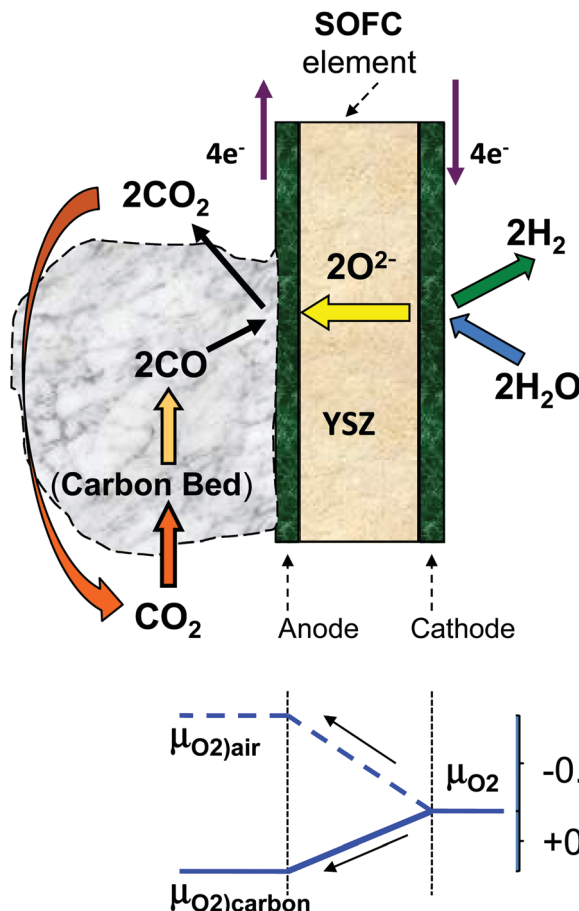


Fig. 44 Schematic of the steam–carbon fuel cell concept, indicating its operating principle, and the chemical potential gradients for steam electrolysis.⁵ While steam electrolysis in SOEC is uphill by about -0.9 V at operating temperatures above $800\text{ }^\circ\text{C}$, replacing air with carbon at the anode compartment lowers the oxygen activity such that the oxygen chemical potential gradient reverses sign and provides SCFC a downhill potential of about $+0.5\text{ V}$. Reprinted with permission from (T. M. Gür, Critical Review of Carbon Conversion “Carbon Fuel Cells”, *Chem. Rev.*, 2013, **113**, 6179–6206). Copyright (2018) American Chemical Society.

(or 3800 W h L^{-1}) versus 8 MJ L^{-1} (or, 2200 W h L^{-1}) for liquid hydrogen. It is an environmentally benign, carbon-free liquid fuel that when combusted in a turbine or converted in a fuel cell back to electricity, its reaction products are only nitrogen and water vapor. Hence, ammonia is ideally suited for environmentally benign chemical storage. Although 79% of the earth’s atmosphere is made up by nitrogen, it is difficult to utilize this abundant resource to make ammonia or other nitrogen chemicals, as molecular N_2 is difficult to activate due to the unusual strength (940 kJ mol^{-1}) of the N–N triple bond. Actually, this is a kinetic hindrance because thermodynamically the reaction between H_2 and N_2 to make ammonia is favorable and spontaneous below $175\text{ }^\circ\text{C}$ and at 1 bar pressure.

Fixation of nitrogen to make fertilizers was made possible by the Haber–Bosch process invented in 1904 that catalytically achieved reduction of nitrogen to produce ammonia at practical production rates in the presence of a Fe-based catalyst using high temperatures ($\sim 500\text{ }^\circ\text{C}$) and high pressures (150–300 bar).

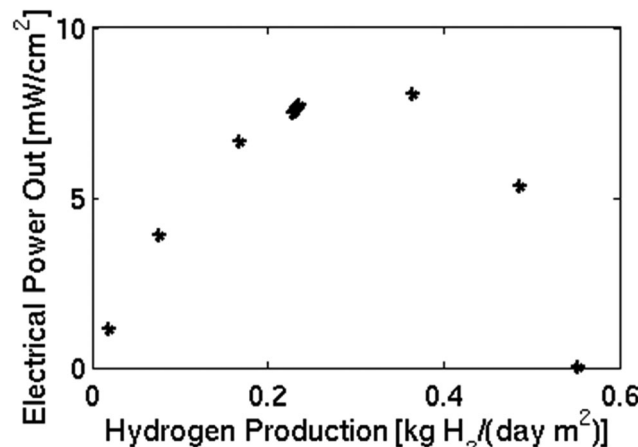


Fig. 45 Experimentally measured electrical power output and co-production of hydrogen in SCFC operating at $850\text{ }^\circ\text{C}$ illustrating the tuneable nature of SCFC output between the two products. The SCFC button cell used in this experiment consisted of 300 micrometer thick YSZ disk with Pt electrodes, and steam at the cathode and activated carbon placed on the anode surface.³⁴⁵

High pressures are needed to shift the synthesis reaction equilibrium to the right due to the Le Chatelier principle, and high temperatures are required to achieve practically high reaction rates, despite the fact that the synthesis reaction between nitrogen and hydrogen is actually exothermic in nature. Accordingly, the Haber–Bosch is an energy intensive process and consumes nearly 2% of the world’s energy production. As the hydrogen for ammonia synthesis is derived from natural gas, it also consumes 3–5% of the global natural gas supply while emitting large quantities of CO_2 . Hence for both energy and environmental reasons, it is desirable to synthesize ammonia near ambient pressures and at lower temperatures, as ammonia tends to decompose above $175\text{ }^\circ\text{C}$ into N_2 and H_2 , and thermodynamically is not stable at high temperatures. Progress made in new catalytic routes for ammonia synthesis at low temperatures and ambient pressures has recently been reviewed elsewhere.³⁴⁷

E.2.1. Electrochemical synthesis of ammonia. Unlike the case for the reaction between H_2 and N_2 , however, thermodynamics of ammonia synthesis using H_2O as the reactant is unfavorable. But this can be overcome, in principle, by applying externally a sufficiently large potential to overcome the thermodynamic barrier and drive the synthesis reaction towards ammonia formation. Indeed, using water as the hydrogen source and nitrogen from air, previous work has demonstrated ammonia synthesis in electrochemical reactors running at atmospheric pressure. The operating temperature of these reactors varied from near room temperature using liquid³⁴⁸ or polymer electrolytes³⁴⁹ such as Nafion[®],³⁵⁰ to temperatures up to $570\text{ }^\circ\text{C}$ using proton-conducting solid oxide electrolytes such ytterbium-doped strontium cerate (Yb:SrCeO_3).^{351,352} Most of the electrochemical studies for ammonia synthesis on precious metal and oxide catalysts utilizing both solid-state and liquid electrolytes have recently been reviewed elsewhere.³⁵³

Electrochemical synthesis of ammonia from N_2/H_2 and N_2/steam mixtures at production rates of $10^{-10}\text{ mol cm}^{-2}\text{ s}^{-1}$

and 10^{-11} mol cm $^{-2}$ s $^{-1}$, respectively, was demonstrated under atmospheric pressure at 220 °C on carbon paper cathodes containing Ru and Ag–Pd alloy catalysts in a solid-state cell that featured a protonically conducting $\text{CsH}_2\text{PO}_4/\text{SiP}_2\text{O}_7$ composite electrolyte.³⁵⁴ Ammonia synthesis at atmospheric pressure and at 450–700 °C was demonstrated at an electrochemical production rate of 2.9×10^{-9} mol cm $^{-2}$ s $^{-1}$ at 550 °C in a solid-state dual chamber reactor featuring a protonically conducting $\text{BaZr}_{0.7}\text{Ce}_{0.2}\text{Y}_{0.1}\text{O}_{2.9}$ (BZCY) ceramic electrolyte and Ni/BZCY cermet catalytic cathode,³⁵⁵ where it was suggested that the protons stored in the Ni/BZCY in the form of highly reactive hydride reacts with adsorbed nitrogen to promote the formation of ammonia.³⁵⁶ Ammonia synthesis by electrolysis of steam and N₂ on nanoscale Fe₂O₃ catalyst particles suspended in a 0.5NaOH/0.5KOH molten electrolyte at 200 °C was reported to achieve 35% Coulombic efficiency.³⁵⁷ A similar study employing Fe₃O₄ nanoscale catalysts in a NaOH/KOH molten electrolyte has recently demonstrated ammonia synthesis from N₂ and H₂ at a rate of 6.53×10^{-10} mol cm $^{-2}$ s $^{-1}$ at 2 mA cm $^{-2}$ at 210 °C with a Faradaic efficiency of 9.3%.³⁵⁸ More recently, ammonia was synthesized at a high rate of 5.3×10^{-8} mol cm $^{-2}$ s $^{-1}$ with a Faradaic efficiency of 10.2% at room temperature in an electrochemical cell featuring a metal–organic framework-derived nitrogen-doped highly disordered carbon electrocatalytic cathode and 0.1 M KOH alkaline electrolyte.³⁵⁹

In all studies, however, production rates for ammonia have generally been low (typically 10^{-9} – 10^{-10} mol cm $^{-2}$ s $^{-1}$) due to low kinetic rates resulting from large polarization losses at the cathode. Also, with few exceptions, the Faradaic efficiencies for ammonia were fairly low below 1% largely due to the competition between the hydrogen evolution and ammonia synthesis reactions at the cathode, as the protons do not only react with nitrogen to form ammonia, but may also receive electrons from the cathode to form H₂ gas. Such competition highlights the importance of selective electrocatalysis with high activity for nitrogen reduction towards ammonia synthesis, but with low activity (*i.e.*, high overvoltage) for the hydrogen evolution reaction at the cathode.

Indeed, theoretical studies also suggest that electrosynthesis of ammonia is limited by the competitive processes of nitrogen reduction reaction (NRR) and hydrogen evolution reaction (HER) on the electrocatalytic sites at the cathode, where the HER usually has a smaller overpotential than for NRR.³⁶⁰ In search of new catalysts, density functional theory (DFT) investigation of the energetics of electro-reduction of nitrogen on $\gamma\text{-Mo}_2\text{N}$ electrocatalyst indicated that among other Mo₂N surfaces, the (111) surface shows the highest reactivity and requires the least cell potential (−0.7 V) to activate the N–N bond of the adsorbed N₂ for nitrogen reduction.³⁶¹ Similar theoretical studies of electrochemical ammonia synthesis reported a thermodynamic limiting-potential of −0.71 V for nitrogen reduction on Ru electrocatalysts, but also indicated the competitive evolution of hydrogen that limits the efficiency of conversion to ammonia.³⁶⁰ More recently, DFT studies of nitrogen electro-reduction to ammonia and the hydrogen evolution reaction on cubic MoC surfaces suggested that the

MoC(111) surface is capable of both adsorbing nitrogen at low overpotentials (−0.3 V vs. SHE) and suppress hydrogen evolution,³⁶² while a lower potential of −0.53 V for nitrogen activation on a MoS₂ nanosheet was predicted for the single Mo atom on the sheet identified as the active site.³⁶³

It is generally accepted that the associative mechanisms among surface intermediates favor the undesirable HER over the desired NRR on the cathode surface, resulting in low Faradic efficiencies reported often for the electrochemical synthesis of ammonia. A detailed density functional theory (DFT) investigation of the energetics of nitrogen electro-reduction to ammonia on more than one dozen precious and transition metal catalysts identified that the potential-limiting step for nitrogen activation on most of these catalyst surfaces is the reductive adsorption of *N₂ to form *N₂H adsorbate requiring a theoretical overpotential of at least −0.5 V.³⁶⁴ The study also proposed strategies such as stabilizing the adsorbed *N₂H while destabilizing the *NH₂ adsorbate by designing new catalyst that can bind *N₂H more strongly on the surface, but without affecting or strengthening the binding energy of *NH₂. As the binding energies of these two species require individual optimization for selective electro-reduction of nitrogen, designing bifunctional catalysts that can bind the two species on different surface sites may be effective pathway to address this difficult problem of nitrogen activation. In this regard, designing catalysts for experimental verification should be guided by theoretical simulations and finding to identify opportunities and gain better insight into the mechanistic aspects of surface processes.

E.3. Other chemical options for storage

Compounds of hydrogen and/or nitrogen offer chemical storage options that release carbon-free environmentally benign products when oxidized for electricity generation. Among such energy carriers, and excepting the rocket fuel hydrazine (N₂H₄) which is toxic and dangerously unstable, energy storage in hydrogen peroxide has recently gained interest as a potential energy storage chemical.^{365–367} Hydrogen peroxide is produced commercially and mostly *via* the hydroquinone process. Alternatively, power from renewable resources can also be employed for cathodic reduction of oxygen in air on an electrocatalyst (typically Co-based) to produce hydrogen peroxide.^{367,368} Hydrogen peroxide can be converted directly into electricity in a fuel cell.^{366,369} Alternatively, electrolysis of hydrogen peroxide can produce H₂ and O₂, which can be electrochemically converted back into electricity in a fuel cell. Hydrogen peroxide stores 5.88 wt% H₂ and 94.12 wt% O₂, so it can be used as an oxidant or a fuel in fuel cell.³⁷⁰ However, its theoretical energy density is only 1.72 MJ kg $^{-1}$ of a 50 wt% aqueous H₂O₂ solution (or, 478 W h kg $^{-1}$) with respect to its decomposition products of water and oxygen. This is not an attractively high energy density for large scale mobile storage applications, and is several orders of magnitude lower than the energy density of 142 MJ kg $^{-1}$ of gaseous hydrogen, and 33 570 MJ kg $^{-1}$ for liquid hydrogen (see Table 6). Also, hydrogen peroxide is corrosive and reactive with metal surfaces, so it must be stored and transported only in

plastic tanks and containers. It also is unstable against alkaline medium. In its dilute form, hydrogen peroxide solutions are safely employed in household cleaners and disinfectants. However, it is not stable above 52 wt% in water and this may place additional constraints on hydrogen peroxide as a practical energy carrier.

At the carbon end of the chemical storage spectrum, there is obvious incentive to recycle CO₂ emissions for reutilization by converting to chemical fuels. However, the most obvious source, which is the atmosphere, is too dilute in CO₂ (~400 ppmv), so finding pure sources for CO₂ may pose challenges. Nevertheless, multiple avenues for CO₂-to-fuels conversion have been explored, including low temperature^{371–373} and high temperature electrochemical reduction of CO₂ to hydrocarbon fuels,^{374,375} or to CO,³⁷⁶ or a mixture of CO + H₂ (*i.e.*, syngas).³⁷⁷ Reduction products CO and/or syngas can further be converted catalytically into fuels *via* the Fischer-Tropsch synthesis process.³⁷⁸ Power from renewable sources can be employed to accomplish CO₂-to-fuels conversion without adding new CO₂ emissions to the environment. Alternatively, solar energy can be employed directly for CO₂ conversion using photoelectrochemical reduction,³⁷⁹ and solar fuel production *via* photocatalytic conversion^{380,381} or solar-thermal splitting.³⁸²

Conversion of CO₂ to fuels in electrochemical, photoelectrochemical or photocatalytic cells can offer incentives for carbon-neutral re-utilization and repurposing of carbon-emissions.³⁸³ However, conversion of CO₂ to fuels is challenging, as it requires both the reduction of CO₂ and the oxidation of water. Reduction potential of CO₂ (−1.06 to −1.40 V *vs.* SHE) is not vastly different from the H₂O splitting reaction (−1.23 V *vs.* SHE), but CO₂ reduction over various metal catalysts usually exhibits high overpotentials due to multiple electron transfer steps as well as the poisoning and deactivation of the catalytic surface by the reaction products. Moreover, the reduction reaction over most catalytic surfaces yields a wide product distribution that may include hydrocarbons and oxygenates as well as CO and H₂, making product separation challenging.³⁸⁴ Hence, selectivity is an important consideration.

Search for earth abundant, high activity, efficient and selective catalysts for the multi-electron reduction reaction have been elusive. After the earlier reports that Cu is an active electrocatalyst for CO₂ reduction,³⁸⁵ many transition metal catalysts have been investigated. Recent progress has shown that nanostructured catalysts^{371,386,387} and reticular processing of metal-organic framework catalysts with atomic level chemical and structural control may offer promising results.³⁸⁸ However, reduction of CO₂ in low temperature aqueous cells is greatly limited by its low solubility (typically 0.03 mol l^{−1}), so pressurization is required to improve solubility. Reduction in non-aqueous electrolytes such as propylene carbonate improves CO₂ solubility. Low product yields and catalyst selectivity need significant improvement too, where the formation of formic acid is often favoured as the primary product of the CO₂ reduction reaction.

High temperature (*i.e.*, 600–800 °C) electrolytic reduction of CO₂ to synthetic methane first reported several decades ago^{374,389}

overcomes some of these difficulties in converting CO₂ to useful fuels, but pose other challenges including product stability and high reduction overpotentials that require search for non-noble metal electrocatalyst with low overpotentials for CO₂ reduction. Often, carbon deposition is observed during reduction on transition metal catalytic anodes including Ni, a commonly employed anode catalyst in high temperature solid oxide electrolyzers (SOE). Moreover, most hydrocarbons are unstable above 300–400 °C and undergo thermal cracking. Hence, high temperature electrolysis of CO₂ poses limitations on the viability of this approach, so lowering the electrolysis temperature is desirable. Many of these difficulties can be minimized by adding steam to SOE for the concurrent reduction of CO₂ and H₂O into hydrocarbon fuels.³⁹⁰ On the other hand, if employed for the production of CO and/or syngas only, high SOE operating temperatures should not pose an issue for product stability.³⁹¹

F. Mechanical energy storage

In mechanical storage systems, inexpensive electrical energy is converted into mechanical energy during off-peak times. Mechanically stored energy is converted back into electricity when the supply of electricity falls short of demand. It should be noted that the stored mechanical energy can easily be converted to heat also, but this is outside the scope of this article. Similarly, mechanical energy storage systems and materials such as springs, pendulums, piezoelectrics that do not lend the potential to store utility scale electrical energy are left outside the scope of this review. Accordingly, only pumped-hydro, flywheels and compressed air energy storage systems are reviewed here.

F.1. Flywheels

Mechanical energy storage in flywheels generally offers high reliability, rapid charge/discharge rates, high power density and storage efficiency, low maintenance costs, long cycle life, and environmentally friendly clean operation.^{392–395} Flywheels offer effective and efficient storage for applications where fast charge/discharge rates, long cycle life (up to hundreds of thousands), and medium to high power (kW to MW) are needed. For these attractive attributes, flywheels are increasingly finding applications in the areas of electric power storage, telecommunications, automotive, marine and aeronautics. Especially for storing energy from variable speed wind generators, they can improve the quality of power delivered to the grid.³⁹⁶

The amount of kinetic energy that can be stored in a flywheel is related linearly to the mass of the flywheel and the square of its angular velocity (see eqn (4)–(6)). Hence, its energy density is largely dependent on the mass and type of the material used for the manufacturing of the flywheel. Moreover, monitoring the angular velocity of the flywheel allows a simple but accurate way of determining its state-of-charge.

The flywheel can be designed to operate either at moderately low speeds (up to 6000 rpm) or at high speeds reaching up to 50 000 rpm. Low speed flywheels are usually constructed of

Table 8 Comparative properties of composite materials considered for flywheel rotors³⁹⁵

| Material | Density (kg m ⁻³) | Tensile strength (MPa) | Max. energy density | | Cost (\$ per kg) |
|----------------------|-------------------------------|------------------------|------------------------|--------------------------|------------------|
| | | | (MJ kg ⁻¹) | (kW h kg ⁻¹) | |
| 4340 stainless steel | 7700 | 1520 | 0.19 | 0.05 | 1 |
| Composites | | | | | |
| E-glass | 2000 | 100 | 0.05 | 0.014 | 11.0 |
| S2-glass | 1920 | 1470 | 0.76 | 0.21 | 24.6 |
| Carbon T1000 | 1520 | 1950 | 1.28 | 0.35 | 101.8 |
| Carbon AS4C | 1510 | 1650 | 1.1 | 0.30 | 31.3 |

steel, and because of their weight can achieve typical energy densities of around 5 W h kg⁻¹ (or 18 kJ kg⁻¹). Since achieving high angular velocities requires the use of high strength materials, carbon fiber reinforced composite materials are typically being used in most high-speed flywheel construction. Rotors made from such lightweight high strength materials greatly improve the energy density of high-speed flywheels up to 100 W h kg⁻¹ (or, 360 kJ kg⁻¹). A further advantage of such lightweight flywheels is that they can come up to speed in a matter of minutes rather than hours, and can provide 1 kW kg⁻¹ of power. It is also worth noting, however, that nanomaterials such as carbon nanotubes exhibit greater tensile strength and Young's modulus than carbon fibers, and hence offer opportunities to further improve the energy density of flywheels. Table 8 illustrates some of the pertinent properties of composite materials explored for flywheel rotors,³⁹⁵ where storage capacities in excess of 1 MJ kg⁻¹ may be achievable.

The general design of a flywheel shown in Fig. 46 consists of a massive rotating drum, *i.e.*, the spinning wheel in the figure, an electromotor generator, and a magnetic suspension bracket, or magnetic bearing that supports the massive rotating drum. In more advanced flywheel designs, superconducting bearings are explored to reduce frictional losses.³⁹⁷ These components are housed in a physical enclosure generally maintained in low-pressure helium gas or vacuum to minimize frictional and self-discharge losses. The interface with the power system

includes a motor/generator, and a variable speed power converter and controller. The motor/generator usually consists of a permanent magnet integrated to a rotor serving as a synchronous generator.

When cheap electricity is available during off-peak times, the electromotor drives the flywheel to high angular velocities where the electrical energy is stored as kinetic energy in the flywheel. When there is demand for electricity, the kinetic energy stored in the flywheel is converted to electricity in the generator as the flywheel slows down. This process is generally efficient, with round-trip storage efficiencies as high as 90%. They also have good cycling capability that range from 10^4 to 10^6 cycles. By virtue of converting mechanical to electrical and back, flywheel can essentially be considered as a mechanical battery.

The typical shortcomings of flywheels include low energy density suitable only for short-term storage, inability to directly produce high output voltage (typically > 36 kV),³⁹⁵ self-discharge during idling, and high capital cost (\$5000 per kW h).³⁹⁴ Storing large amounts of electricity from the power grid requires high capacity, and hence massive size flywheels. However, frictional losses, which scales with size, reduces rotational speed and are the primary cause of self-discharge that lower the flywheel efficiency. For example, the frictional losses in a 200 ton flywheel are estimated to be about 200 kW, which lowers the efficiency of the flywheel to 78% after 5 hours of operation, and down to 45% after 24 hours.²² In others, self-discharge rates as high as 20% per hour have been reported.²⁴ Such dramatic impact of frictional forces, even in vacuum environment, renders flywheels unsuitable for long-duration storage. However, this handicap may not pose a major barrier for short periods. In the case of integration with intermittent renewable sources, whose electricity output fluctuates by the hour, flywheels offer attractive advantages due to their fast response times. This is illustrated in Fig. 47, where a flywheel coupled to a wind turbine successfully smoothens the fluctuating and intermittent nature of power generation by peak shaving.³⁹⁴ Even with conventional power generation, where there is a mismatch between the electricity demand (or, load) and the electricity supply, flywheels can rapidly supply stored energy back to the grid and also provide power conditioning.

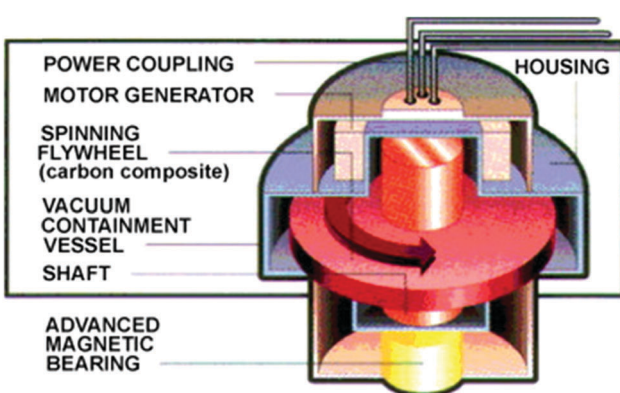


Fig. 46 Schematic cross-section illustrating the basic design and operation of a flywheel energy storage system.²⁴ Reprinted from I. Hadjipaschalos, A. Poulikkas and V. Efthimiou, Overview of current and future energy storage technologies for electric power applications, *Renewable Sustainable Energy Rev.*, 2009, **13**, 1513–1522, with permission from Elsevier.

F.2. Compressed air energy storage (CAES)

The idea of storing energy in the form of compressed air pumped into large underground cavities goes back many decades,³⁹⁸

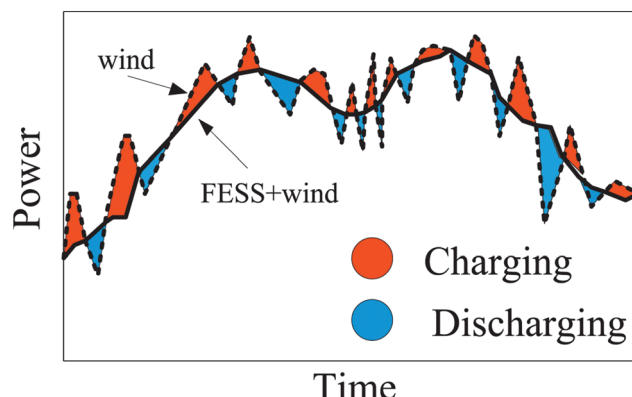


Fig. 47 Power conditioning and peak shaving by the use of flywheel energy storage system (FESS) coupled to a wind turbine.³⁹⁴ Reprinted from A. A. K. Arani, H. Karami, G. B. Gharehpetian and M. S. A. Hejazi, Review of flywheel energy storage systems structures and applications in power systems and microgrids, *Renewable Sustainable Energy Rev.*, 2017, **69**, 9–18, with permission from Elsevier.

but early progress has been somewhat slow in large part due to the lack of need or interest for grid storage until recently, where rapid growth and deployment of intermittent renewable sources has generated renewed interest in compressed air energy storage (CAES).^{399,400}

Unlike flywheels, the typical discharge times for compressed air energy storage (CAES) are long and measured in hours up to days. Their start-up times are also relatively longer than flywheels, measured in minutes.³⁹⁹ Furthermore, CAES requires ready availability and proximity to large underground cavities, aquifers, and caverns that exhibit impervious rock formations to ensure gas tight containment of compressed air. Hence, its availability and feasibility is necessarily restricted by geological constraints. This is one of the major shortcomings of this technology. On the other hand, it eliminates large and costly installations for building a storage facility. Where geological conditions are favorable, CAES offers an attractive and cost-effective technological option to store off-peak power from the grid in the form of compressed air that is kept underground reservoirs during low demand. When make-up power is needed, the high-pressure compressed air is converted back to electricity with an estimated efficiency up to 70%.²² This process is schematically depicted in Fig. 48.

The typical charging cycle of CAES starts with compressing air to high pressures in several stages by electrically drive compressors during low electricity demand, usually with cooling in between successive compressions before injecting into the underground storage cavity, which is maintained at air pressures up to 100 bar. Temperature of the injected air is typically kept at or below 40 °C to avoid thermal stress in the underground cavity. During high electricity demand, i.e., discharge cycle, compressed air from the underground cavity is preheated in a recuperator or heat exchanger, and mixed with fuel in a combustor. The hot combustion gases are expanded in a turbine to generate electricity.

There are a variety of CAES processes being considered or developed for different applications, but all of them can be

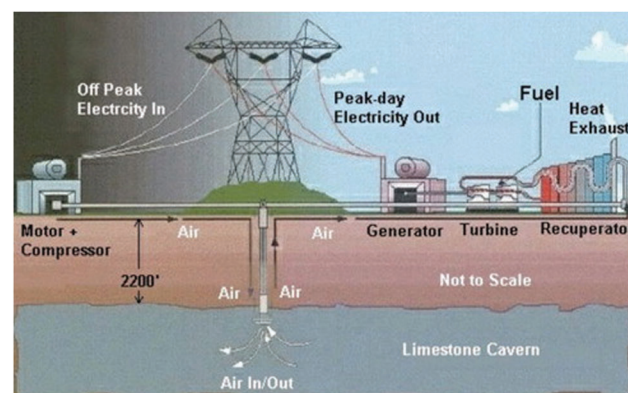


Fig. 48 Schematic illustration of compressed air energy storage.²² Reprinted from H. Ibrahim, A. Ilinca and J. Perron, Energy storage systems – characteristics and comparisons, *Renewable Sustainable Energy Rev.*, 2008, **12**, 1221–1259, with permission from Elsevier.

grouped under three categories differentiated by how the heat is managed during charging and discharging cycles, namely, adiabatic CAES, diabatic CAES and isothermal CAES. In the adiabatic CAES process, the heat of compression during the charging cycle is fully captured into a thermal energy storage system, and re-used to preheat the compressed air prior to expansion during the discharge cycle. In the diabatic CAES process, by contrast, the heat of compression is lost to the surrounding atmosphere during the charging cycle by cooling the compressed air, which then necessitates an external heat source to reheat the compressed air during the discharge cycle. Unlike in diabatic and adiabatic CAES processes, heat of compression needs to be minimized or eliminated completely during isothermal CAES to keep the air temperature near ambient at all times. Consequently, performance parameters such cycle efficiencies, energy densities, and start-up times widely vary among the three process groups.

There are a number of installations of CAES systems around the world. The first CAES system, namely the Huntorf facility, was installed near Bremen, Germany in 1978. It had a charging time of 8 hours, discharge time of 2 hours, and maximum charging and discharging power of 60 MW and 290 MW, respectively, with a power plant efficiency of 42%.⁴⁰⁰ In 1991, another CAES system built over a 700-meter deep $2.55 \times 10^6 \text{ m}^3$ cavern in McIntosh, Alabama, USA, and operated at 54% efficiency but with significantly longer charge/discharge times of 40/26 hours, respectively, had provided 110 MW of power for a duration of 226 hours, and has been in use for 1770 hours per year.²² In the last decade, several CAES projects with storage capacities in the 175–315 MW range have been initiated in the US³⁹⁹

It is estimated that 0.7–0.8 kW h of off-peak electricity is required to compress the air into the underground formations in order to supply 1 kW h of peak-demand electricity back into the grid.²² As more than 2/3rd of the available power in a power plant is consumed in compressing the combustion air for the gas turbines, it makes sense to separate this process in time and use the off-peak power to compress the air, and use it to convert to electricity when peak demand increases.

Due largely to their geological requirements and constraints, CAES can be used only at large scales. In fact among all other storage technologies and systems, CAES is currently the only technology other than pumped-hydro that has the ability to provide large-scale energy storage capacity, typically, less than 300 MW. CAES also offers the longest storage duration of the order of 20–40 years without significant losses, and long cycle life reaching more than 13 000 charge/discharge cycles. However, the volumetric energy density for CAES is low, typically 2–12 kW h m⁻³ (or, 7.2–43.2 MJ m⁻³). By comparison, the typical volumetric energy densities of liquid fuels are nearly 3 orders of magnitude larger than these values. A recent economic analysis of the Danish electric system that produces a significant portion (42% in 2015) of its electricity from offshore wind has concluded that investments in CAES carry risks and do not offer more cost-effective options than other storage alternatives.⁴⁰¹ Thermodynamic analysis and modeling of CAES systems using first and second law (*i.e.*, exergy) efficiencies predicted more optimistic results regarding the viability of CAES for large scale storage with efficiencies as high as 86% when combined with thermal storage, but these studies still caution on the technical challenges that still remain to overcome by this technology.^{399,400,402}

F.3. Pumped hydro storage

Hydropower constitutes the major segment among the renewable energy sources with a global share of more than 16% of the global electricity production in 2016, and is a mature commercially proven technology. It has been used at industrial scale since the turn of the 20th century. Globally, more than 25 countries depend heavily on hydropower to supply a large fraction of their electricity production.⁴⁰³ Fig. 49 indicates the geographical distribution for the global potential of hydropower with respect to the continents and highlights the fact that significant potential remains unexploited. Likewise, Table 9 provides information about the world top ten countries with the largest installed and generation capacities of hydropower.

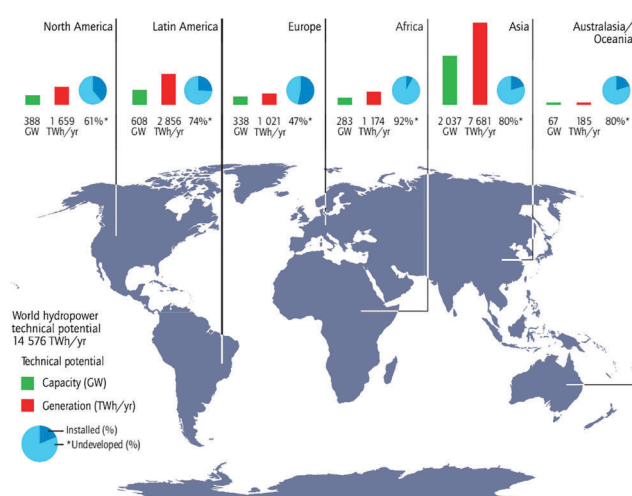


Fig. 49 Continental potential of hydropower generation, also indicating the percentage of undeveloped hydropower potential as of 2009.⁴⁰⁴

Table 9 Top-ten countries in the world with the largest installed and generation capacities of hydropower in 2010. Data compiled from ref. 404 and 405

| Country | Installed capacity (GW) | Hydro electricity production (TW h) | Share of electricity generation (%) |
|---------------|-------------------------|-------------------------------------|-------------------------------------|
| China | 210 | 694 | 14.8 |
| Brazil | 84 | 403 | 80.2 |
| Canada | 74 | 379 | 62.0 |
| United States | 79 | 328 | 7.6 |
| Russia | 50 | 165 | 15.7 |
| India | 38 | 132 | 13.1 |
| Norway | 30 | 122 | 95.3 |
| Japan | 28 | 85 | 7.8 |
| Venezuela | | 84 | 68 |
| Sweden | | 67 | 42.2 |
| World total | 936 | 3300 | 16 |

When and where geographical conditions are satisfied, hydropower plants can be employed for pumped hydro energy storage (PHES). Indeed, pumped hydro constitutes by far the largest segment of the electricity storage technologies globally. Extensive reviews of PHES technology addressing the various aspects of the technical challenges,⁴⁰⁶ technology trends,^{407,408} coupling to other renewable sources,^{406,409} and international PHES projects^{407,410} are provided elsewhere.

However, the availability and cost of this mature technology is site-dependent and only feasible where geographic conditions are favorable for its implementation. PHES provides more than 96% of the global installed storage capacity⁴¹¹ (see also Table 2). It provides cost effective solutions to load-leveling and peak-shaving in the grid and during peak demand, eliminates the need for generating make-up power by gas turbine plants using natural gas. Thus, pumped-hydro displaces the use of fossil fuels and thus helps reduce carbon emissions.

In principle, PHES employs the potential and kinetic energy of vast quantities of water stored between two water reservoirs with sufficiently large volumes and separated by a significant difference in altitude. Some of the pumped-hydro facilities employ abandoned mines or artificial formations as reservoirs, but most employ natural reservoirs such as lakes.

During off-peak times where the electricity demand is low, excess generation capacity is used to pump water from a lower reservoir to a higher one, such as a volcanic lake. When electricity demand is increased beyond generation capacity, water is released from the upper reservoir to turn a turbine to generate make-up electricity. The general layout of the key features of a PHES plant is schematically shown in Fig. 50. The turbine serves dual purposes, both to pump water to the upper reservoir and also to generate electricity when water is released from the upper into the lower reservoir.

PHES generally provides large storage capacities with power ratings in some places ranging up to 1000's of MW (or, 100's of GW h of energy). However, the storage capacity depends strongly on the separation height between the two reservoirs and the volume of water transferred between the two. For example, 1 ton of water released from 100 meters can

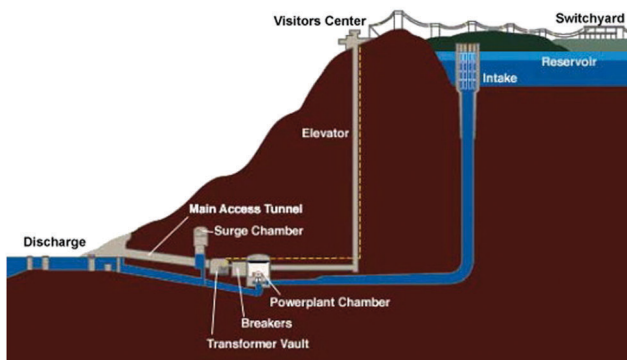


Fig. 50 A typical layout of pumped-hydro storage plant operating between two water reservoirs.⁴¹² Reprinted from I. Hadjipaschalidis, A. Poullikkas and V. Efthimiou, Overview of current and future energy storage technologies for electric power applications, *Renewable Sustainable Energy Rev.*, 2009, **13**, 1513–1522, with permission from Elsevier.

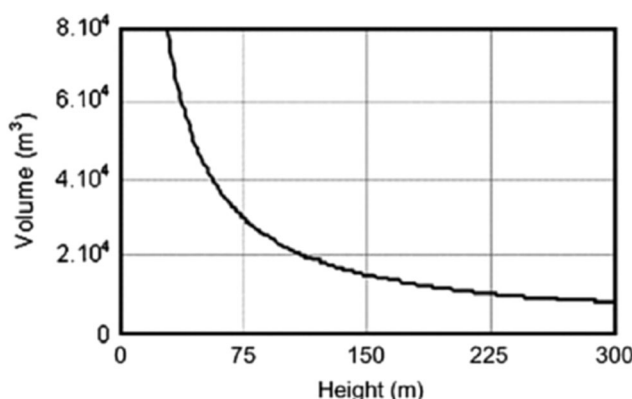


Fig. 51 Volume of water needed to store 6 MW h of electrical energy as a function of separation height between reservoirs.²² Reprinted from H. Ibrahim, A. Ilincă and J. Perron, Energy storage systems – characteristics and comparisons, *Renewable Sustainable Energy Rev.*, 2008, **12**, 1221–1259, with permission from Elsevier.

potentially generate 0.27 kW h of electricity. Fig. 51 illustrates the relation between the volume of water and the vertical separation height between the two reservoirs required to store about 6 MW h of electrical energy.²²

One of the major shortcomings of pumped-hydro is its strong reliance on both the availability and suitability of two large bodies of water located in reasonable physical proximity of each other and separated by a sufficient vertical distance. As Fig. 51 shows clearly, relatively low energy density of this technology requires both large volumes of water and large differences in altitude. The capital costs for plant construction may vary widely from 470 Euro per kW to 2170 Euro per kW (in 2009 Euros)⁴¹⁰ and strongly depends on geographic constraints. A more recent report estimated the average capital cost for a 500 MW green field hydropower generation plant in the USA to be about \$3500 (+35%) in 2012 with capacity factors ranging between 25 and 90% and a leveled cost of electricity between \$20 per MW h and 190 per MW h for large plant operations.⁴⁰³

Sometimes, conventional hydroelectric power generation plants can operate also for storage purposes provided that another large body of water at a lower, or higher, altitude is conveniently available nearby, which is more an exception than a norm. When available, such combined hydropower generation-storage plants provide the opportunity for peak-leveling and shaving in an environmentally friendly manner.

An unavoidable problem with pumped hydro storage is evaporation losses from the large surface areas of the two large bodies of water, especially the ones at high altitudes, where such losses can be significant. This leads to lost work from pumped water due to natural causes, and reduces the efficiency for storage.

Nevertheless, PHES offers high conversion efficiencies, where 70–85% of the energy used to pump water to the upper reservoir can be recovered during the generation cycle. Pumped-hydro storage offers response times that may range between minutes to hours, which allows it smooth out changes in load or demand conditions on the grid. This is critical for power management and frequency regulation on the network. By contrast, thermo-electric power requires relatively longer start up times, which can lead to frequency and voltage instabilities in the grid.

With significant cost reductions in wind and solar, there is growing interest in hybrid storage systems that explore various combinations of wind, solar and PHES technologies for consideration for large-scale energy storage as well as autonomous or isolated electric grids.^{407,408,413} Mostly, these studies employ system modeling to simulate widely varying conditions for optimum performance, cost and efficiency. Although the outcomes are impacted by site-specific constraints, it is suggested nevertheless that coupling PHES to intermittent renewable power sources help increase the penetration level of these renewable sources into the power mix.

As inland PHES is limited by geographical conditions, seawater presents a nearly unexplored potential for PHES. Currently, there is only one seawater-based PHES facility in the world operating since 1999 in Okinawa, Japan with a design a capacity of 30 MW and a head of 136 meters. There are few others in conceptual or design stage, including the Dead Sea Power project located on the Red Sea with a design capacity of 1500–2500 MW, and one proposed seawater PHES project in Ireland to provide 480 MW of storage capacity.⁴⁰⁶

G. Thermal energy storage

Central to electricity storage is the gap between generation and consumption, and thermal energy storage is a fundamentally sound approach that can provide attractive opportunities to achieve storage at large scale. Already, thermal energy storage is ubiquitous in our daily lives including hot water heaters and residential heating and cooling. But converting thermal energy back to electricity is generally not a highly efficient process. Moreover, the “energy quality”, which sometimes is also referred to as the quality of heat, may be a design consideration for many applications. For thermal storage, the temperature, T , at which the energy, or heat, is delivered, is an important factor.

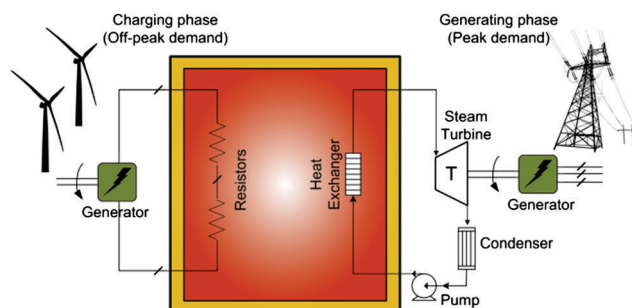


Fig. 52 Schematic process flow diagram for sensible heat thermal storage from wind power. Reprinted from ref. 414 (X. Luo, J. Wang, M. Dooner and J. Clarke, Overview of current development in electrical energy storage technologies and the application potential in power system operation, *Appl. Energy*, 2015, **137**, 511–536).

As Gibbs free energy, G , defined in eqn (1) represents the maximum amount of work that can be extracted from a system, the energy quality can be defined as the ratio of the Gibbs free energy to enthalpy by,

$$G/H = \Delta T/T = 1 - (T_0/T) \quad (34)$$

here, T_0 denotes the ambient temperature. It is clear from eqn (34) that higher the delivery temperature, T , higher will be the energy quality, or the quality of the heat delivered.

There have been renewed interest in thermal energy storage materials and technologies, and several excellent reviews are available in this topical area.^{415–419} Most thermal energy storage systems can be grouped under two general categories, namely sensible heat storage and latent heat storage. The latter employs the latent heat that often accompanies a phase change such as fusion or melting, or a structural transformation. Those that utilize the sensible heat given in eqn (7) rely on the specific heat and mass of a material. The schematic of thermal storage using sensible heat is illustrated in Fig. 52, where electricity from a wind farm is used to heat up a thermal mass of desirable heat capacity. The heated mass may be in liquid or solid form, which is then employed to make steam and generate electricity

when the demand is high. Many of these systems consider molten salts or their eutectic mixtures for thermal storage, which brings about potential problems related to hot corrosion, and thermal and chemical stability issues for the container materials in the highly corrosive environment of the molten salt. To mitigate such undesirable stability problem, the use of costly construction materials is often required. So cost and stability, and hence useful service life, are important considerations in the proper evaluation and selection of the appropriate thermal storage medium, materials and system design.

G.1. Sensible heat storage in molten salts and other materials

Thermal storage via sensible heat typically involves heating up a mass of a material in liquid or solid form and raising its temperature by ΔT without undergoing a phase change or structural transformation. The process of converting the stored thermal energy back to electricity typically involves using the sensible heat to make steam that drives a turbine, and finally the generator. Excess electricity during low demand, electricity from renewable sources, or thermal solar energy can all be employed as viable sources to heat up the storage material or medium. As the required mass or volume of the storage medium is typically large in order to store sufficient energy, the economics of this scheme favor relatively abundant and inexpensive materials with sufficiently high heat capacities. For that reason, solids are mostly preferred but molten salts are also being employed as well. Representative physical properties of select solids, liquids and molten salts are provided in Table 10.

Some of the materials listed in the table offer attractive properties for thermal energy storage applications but some have disadvantages or shortcomings such as low energy density, thermal conductivity or specific heat. Most are cheap and abundantly available, and several such as iron and water offer high volumetric energy densities around 4 MJ m^{-3} . Particularly, water possesses nearly two times larger specific heat than most other materials, and offers an energy density of 4.19 MJ m^{-3} , but it is only operable below 100°C . So the energy quality of water as a storage medium for large-scale storage

Table 10 Comparison of materials properties for thermal storage

| Materials | Melting point ($^\circ\text{C}$) | Thermal conductivity ($\text{W m}^{-1} \text{K}^{-1}$) | Density (kg m^{-3}) | Specific heat, C_p ($\text{J kg}^{-1} \text{K}^{-1}$) | Vol. energy density (kJ m^{-3}) |
|--|------------------------------------|--|--------------------------------|---|--|
| Solids | | | | | |
| Aluminum | 660 | 205 | 2700 | 920 | 2484 |
| Iron (cast) | 1150 | 58 | 7200 | 540 | 3890 |
| Glass | N/A | | 2710 | 837 | 2272 |
| Brick | > 1500 | 1.3 | 1970 | 921 | 1813 |
| Concrete | N/A | 1.0–1.8 | 2000 | 880 | 1760 |
| Granite | 1215–1260 | 1.7–4.0 | 2400 | 790 | 1896 |
| Liquids | | | | | |
| Water | 100 | 0.58 | 1000 | 4190 | 4190 |
| Molten sodium | 98 | | 750 | 1260 | 945 |
| 50 : 50 ethylene glycol–water mix | | | 1075 | 3480 | 3740 |
| Molten salts | | | | | |
| 50 wt% KNO_3 –40% NaNO_2 –7% NaNO_3 | 149 | | 1680 | 1560 | 2620 |
| LiNO_3 – NaNO_3 – KNO_3 | 117 | | 1710 | 1540 | 1056 |
| NaNO_3 – KNO_3 | 222 | | 1750 | 1530 | 756 |

applications is necessarily low. Environmental impact should also be taken into consideration. For example, underground energy storage using hot water may cause irreparable damage to the ground or the geological structure. However, water is a suitable energy storage medium for heating and cooling for residential buildings and workplaces where low quality heat is acceptable. Similarly, water–ethylene glycol mixtures are also limited by their operating window for storage, but are suitable for heating and cooling applications that do not require high quality heat.

On the other hand, most inexpensive solids offer a wide window of operating temperatures that can extend close to their melting points. Thus the quality of energy for these materials can be quite high, and makes them attractive for large-scale storage applications. Furthermore, their high densities nearly twice that for liquid and molten medium helps render the storage medium to be compact. Unlike water, other liquids, or molten salts, low vapor pressure of solids makes containment easy and inexpensive, while reducing materials loss to the surroundings. Indeed, solar energy can be stored up to 36 kJ kg^{-1} (or, 10^5 kJ m^{-3}) in rocks or concrete for space heating applications.⁴¹⁸

There is, however, another important consideration that has to do with how fast energy is transferred to or retrieved from the storage medium. In this respect, thermal conductivity is an important material parameter for system design purposes. As shown in Table 10, metals possess significantly higher thermal conductivity than many of the refractory solids, as well as liquids. For molten salts, thermal conductivity information is rather scarce, but for several fluoride-, carbonate- and nitrate-based molten salt eutectics, thermal conductivity values between 1.09 and $1.21 \text{ W m}^{-1} \text{ K}^{-1}$ have been reported.⁴²⁰ Clearly, nearly two orders of magnitude difference between the thermal conductivities of metals and refractory solids, liquids, or molten salts can be an important factor in the selection of the proper storage material and system design for a specific application.

Needless to say, heat loss to the surrounding in the forms of radiation, convection and/or conduction is another important consideration in energy storage, because it directly impacts overall system efficiency. Such losses are dependent not only on the temperature difference between the storage medium and its surrounding, but also on other factors including exposed surface area, emissivity, and thermal conductivity. Hence, proper design, materials selection, and effective thermal insulation of the storage system are important considerations.

G.2. Latent heat storage in phase change materials

This approach utilizes the enthalpy change during phase transitions without a change in composition, *i.e.*, congruent phase transitions, where typically the first derivative of Gibbs free energy during the phase change is discontinuous. This type of energy storage may involve either the enthalpy of fusion, typically during melting, or the enthalpy change during a solid-state crystal structure transformation below the melting point of the material. As in the case of sensible heat, energy is stored in the material either by inducing melting or a structural phase

change, and retrieved back in the reverse process. Since most congruent phase changes occur at sufficiently elevated temperatures, the energy quality of energy storage using latent heat is typically high. Also, the entropies of solid materials are considerably smaller than those of liquids or melts. So the entropy change during melting can be considerable. Hence, the enthalpy change across the phase transformation is relatively larger for fusion rather than for solid-state phase transformations. On the other hand, congruent melting and solidification usually involves considerable molar volume change in the storage material, which makes containment and system design more complicated, while molar volume changes during solid-state phase transformations are typically much smaller. Table 11 lists a number of cheap and readily available materials that can be considered for thermal energy storage *via* phase change. However, it also illustrates that the amount of storable energy may be quite limited, as the energy densities for $0.3\text{--}0.5 \text{ MJ kg}^{-1}$ for many phase change materials are several orders of magnitude lower than those for chemical fuels such as hydrogen (142 MJ kg^{-1}), gasoline (44 MJ kg^{-1}), or methanol (20 MJ kg^{-1}).

Nevertheless, phase change materials can store 5–14 times more energy per unit volume than sensible storage materials such as water, most rocks and many other solids. Unlike sensible heat storage materials, they store and release heat at nearly constant temperature, which is an advantage. Most phase change materials can be grouped in two major categories, namely, organic and inorganic materials, and their eutectics. The inorganic group exhibits nearly twice the volumetric capacity of $250\text{--}400 \text{ kg dm}^{-3}$ for storage compared to $128\text{--}200 \text{ kg dm}^{-3}$ for organic phase change materials.⁴¹⁶

The most common organic phase change material is the paraffin group of hydrocarbons with melting points slightly higher than room temperature and moderate enthalpies of fusion. The latter increase with chain length. They possess many desirable properties including low vapor pressure, chemical stability in some cases up to 500°C , low toxicity and corrosivity, commercial availability at large quantities and at low cost, and negligible supercooling. On the hand, they undergo considerable volume change during solid–liquid phase transition and exhibit low thermal conductivities that greatly limit the rate at which heat is stored and released.

An important group among inorganic phase change materials is salt hydrates (see last three rows in Table 11), which break up at the melting temperature into water and an anhydrous salt (or, water and the lower hydrate). They offer attractive properties such as high volumetric latent heat of fusion, significant thermal conductivity nearly twice that of paraffins, small volume change during phase transition, and mostly non-corrosive and non-toxic and low cost. However, many inorganic salt hydrates melt incongruently, *i.e.*, the water released during the phase transition is not sufficient to dissolve all the solid salt phase(s). Another important disadvantage of salt hydrates is supercooling. As the rate of nucleation at the phase transition temperature is rather slow, the solution requires supercooling to lower temperatures to achieve a

Table 11 Latent heats of fusion of phase change materials. Compiled from ref. 416 and 421

| | Transition temperature (°C) | Enthalpy of fusion (MJ kg ⁻¹) |
|--|-----------------------------|---|
| Water | 0 | 0.335 |
| Paraffin | 20–60 | 0.14–0.28 |
| LiNO ₃ | 250 | 0.37 |
| NH ₄ NO ₃ | 170 | 0.12 |
| NaNO ₃ | 307 | 0.13 |
| NaOH | 318 | 0.15 |
| Ca(NO ₃) ₂ | 561 | 0.12 |
| LiCl | 614 | 0.31 |
| FeCl ₂ | 670 | 0.34 |
| MgCl ₂ | 708 | 0.45 |
| KCl | 776 | 0.34 |
| NaCl | 801 | 0.50 |
| LiNO ₃ ·2H ₂ O | 30 | 0.296 |
| FeCl ₃ ·6H ₂ O | 37 | 0.223 |
| Na ₂ PHO ₄ ·12H ₂ O | 40 | 0.279 |

reasonable nucleation. Hence, the energy stored in the material can only be released at a temperature lower than the fusion point. The undesirable effect of supercooling naturally renders inefficiency in the storage cycle.

Most studies concluded that low thermal conductivities of phase change materials pose a major limitation for their implementation for large scale energy storage applications. Therefore, recent attention has turned to designing microstructures and methodologies such as adding dispersions of metals, graphite and other high thermal conductivity materials, micro-tubes, fins, and encapsulation, all intended for enhancing the rate of heat transfer in these materials.³⁶⁰

H. Grid storage

The electrical grid depicted in Fig. 53 is a critical component of the vital connection between the point of use and power generation. In many parts of the world, the existing grid infrastructure is insufficient, old or obsolete. In the US, the Department of Energy estimates that 70% of the transmission lines and 70% of power transformers are 25 years or older, and 60% of the circuit breakers are more than 30 years old.¹⁷ The expected growth of 43% in the global electricity production (see Table 1) urgently requires major upgrading of the grid system to make the grid more secure, reliable, and responsive. Certainly, large-scale deployment of distributed storage capacity will help improve grid resilience and security, reduce line congestion and line loss, lower the need for generation

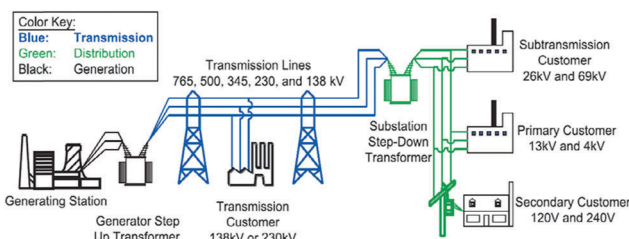


Fig. 53 Basic structure and lay out of the US electric grid system.⁴²²

during peak demand times, and also extend the capability of the transmission grid, for example, by positioning the storage system downstream on the load side. A White House 2013 report from the Executive Office of the President also concluded that energy storage is important for emergency preparedness during natural disasters and also in increasing grid resilience and robustness to weather outages.³⁶⁷

Rapid deployment of intermittent renewables such as wind and solar has highlighted the difficulties in fully integrating these renewable sources into the grid. A recent report by the US National Renewable Energy Laboratory (NREL) has suggested that more than 20% penetration of intermittent renewables such as solar into the electric grid can cause operational challenges that can adversely affect grid reliability.⁴²³ Although the scope of this study was rather limited to the grid systems in the islands of Oahu and Maui in Hawaii, it strongly underscores the importance of storage technologies to cope with dynamic loads. Assuming 20% penetration by renewable sources by 2020, a recent report by Pacific Northwest National Laboratory (PNNL)⁴²⁴ estimated that the US would need a total of 37.67 GW for intra-hour balancing capacity, and an energy storage capacity of 14.3 GW h that can provide power at rated capacity for 20 minutes. By comparison, the US has 24.6 GW of storage capacity (~2.3% of total electricity production capacity), including announced projects.¹⁷ Unfortunately, 95% of this storage capacity is by pumped-hydro, whose availability is site-specific and constrained by location, and only 1.2 GW of this capacity is provided collectively by CAES, flywheel, thermal, and battery storage. The situation is not much different elsewhere. For example, Europe and Japan have considerably higher fractions of grid storage than US, but it is largely dominated by pumped-hydro also. The global energy storage capacity has now grown to 168.6 GW (see Table 2), where pumped-hydro makes up more than 96% of the total capacity. It is clear that distributed storage can help remove or mitigate the logistic mismatch problem between the siting of storage availability and demand.

To appreciate the need for storage, one needs to look no further than intermittent sources such as solar and wind, whose power generation show large variability up to 80% during time scales of minutes to hours, as depicted in Fig. 54

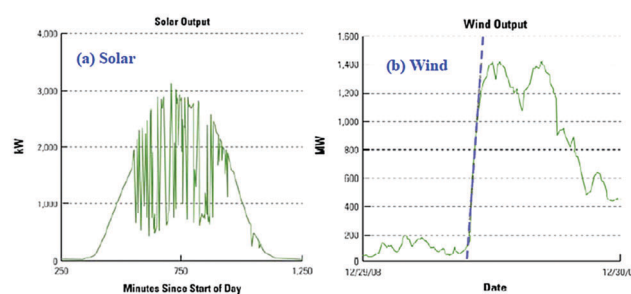


Fig. 54 Time-dependent intermittency of solar and wind power varying >80% within time scales of minutes to hours in a single day. The high rate of change of power (i.e., ramp rate) of the order of MW min⁻¹ is illustrated for wind power.²⁴¹

The ramp rates, especially for wind power which can be of the order of MW min^{-1} or higher, requires energy storage with characteristic time constants and capacities compatible with power production. US Department of Energy recommends²⁴¹ three time scales to accommodate variability of intermittent power sources; (a) seconds-to-minutes, with storage capacity of 5–7% of generated power for frequency and voltage support – for US this is about 20–50 GW of storage capacity for the domestic generation capacity of more than 1100 GW of power; (b) minutes-to-hours of capacity for up to one hour of storing 20% of power from renewable sources for smoothing intermittency – or, nearly 220 billion kW h of storage capacity in order to accommodate more than 1100 billion kW h of solar and wind power generated globally in 2016 (see Table 1); and (c) hours-to-days of power storage capacity for peak shaving and 20% integration of renewables into the grid, which requires a power capacity on the order of 200 GW and 1000 GW h for 20% integration of renewable energy. These estimates were based on the assumption that 200 GW generation from renewable resources would be required to meet 40% of daily load with 100% load shifting for 5 hours.²⁴¹

The need for cost-effective storage is highlighted in Fig. 55, which shows the 2010 costs of power and energy for various storage technologies, as well as the cost targets set by US DOE ARPA-E.²⁴¹ The figure also marks the required response times for 10 minute and 1 hour reserve storage capacity. Pumped-hydro and CAES with time scales from hours to days provide cost-effective storage where available, but rapid development and deployment of a portfolio of diverse storage technologies with the required characteristic response times and energy capacities are needed for broader siting of storage facilities, especially for intermittent renewable sources. Clearly, the figure indicates that major cost reductions are needed for electrochemical technologies to meet the cost target.

Sizing of a storage system for grid applications requires several operational parameters to be specified, including the power rating (in MW) to meet the target load, the energy rating

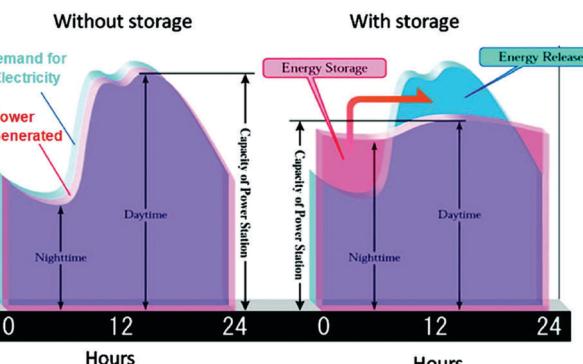


Fig. 56 Schematic illustration of balancing the demand and generation by load-leveling via storage.³² Reprinted with permission from (Z. Yang, J. Zhang, M. C. W. Kintner-Meyer, X. Lu, D. Choi, J. P. Lemmon and J. Liu, Electrochemical energy storage for green grid, *Chem. Rev.*, 2011, **111**, 3577–3613). Copyright (2018) American Chemical Society.

(in MW h) that the storage unit is required to deliver to the grid or absorb from it at any given cycle, and the characteristic response time. Large variations between on and off-peak electricity demand and generation can effectively be balanced and smoothed out using large-scale electrical storage to provide peak-leveling and shaving options, as shown schematically in Fig. 56. Real time regulation of the grid to control and respond to momentary fluctuations caused by the imbalance between load and generation is also a critically important need that can be achieved when combined with proper storage technology. Currently, grid regulation is in large part accomplished by placing generator units that are ready to go online or offline, or increase or decrease their power as needed. Besides lost generation capacity when off-line, such variable power and on/off fluctuations cause their wear and tear on the generation equipment and plants, especially for thermal base-load generators. Another consideration is frequency response, which is similar to regulation but responds to sudden fluctuations, usually in time periods of seconds. Fig. 57 illustrates the effects of using grid regulation and frequency response. Electrical storage not only provides time-shift duty, *i.e.*, buying electricity

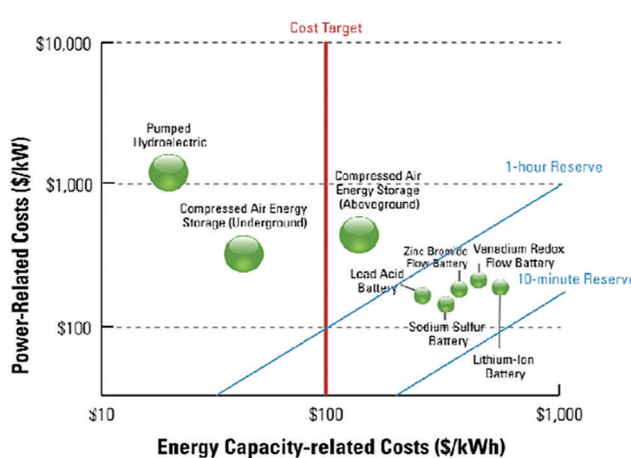


Fig. 55 Cost of energy storage for various technologies, showing cost targets and minimum required response times. Small circles denote response times of seconds, large circles indicate time scale in minutes.²⁴¹

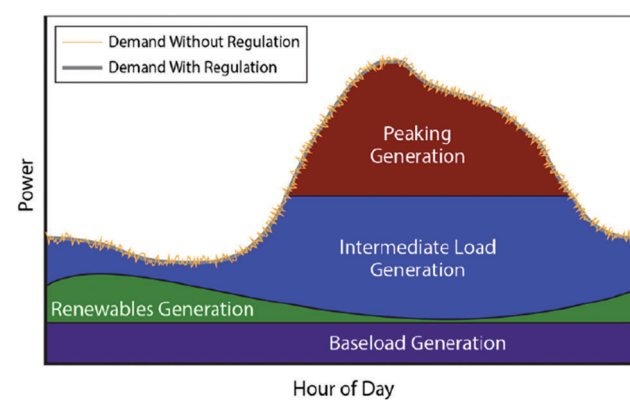


Fig. 57 Variation of the load on the grid with and without regulation and frequency response.¹⁷

off-peak times or when prices are low, but also prevents wasting of excess generation.

Load-following, depicted in Fig. 57, is characterized by power output that changes frequently within minutes in response to the changing balance between demand and electricity generation and to maintain the scheduled system frequency and establish the interchange with other generation areas. This is particularly important for intermittent and constantly varying renewable power sources such as wind and solar. Storage can alleviate some of these problems. However, large scale deployment of electrical energy storage technologies and systems for grid applications require that these widely diverse technologies need to overcome several barriers before they can be fully adopted and deployed by the utility industry. Namely, these storage technologies need to demonstrate the following¹⁷:

Cost competitiveness: This includes bringing down the cost not just of the storage component (*e.g.*, flywheel, battery, *etc.*), where most of the present focus and attention is concentrated, but also of the balance of plant that makes up the entire storage system, where the storage component makes up roughly 30–40% of the entire system cost. Based on published techno-economic analyses by others, an independent study suggested a target cost for energy storage at <\$100 per kW h (and power at <\$600 per kW).⁴²⁵ Also important is to quantify the value of storage in terms of the services provided to the grid, such that a single storage unit may capture several revenue streams and achieve economic viability. It is likely that the early deployment for storage systems will happen in high value applications. Hence, it is important to focus on key factors such as life-cycle cost, and performance parameters such as round-trip efficiency, energy density, cycle life, and capacity fading.

Validated performance and safety: Performance and safety evaluation of the storage system should be verified in accordance with the accepted industry codes and standards. For example, the true lifetime of the storage system and the time its takes to generate revenue is important for industry stakeholders and decision makers. Hence, validation of safety, reliability, and performance of the storage system is critically important for gaining the confidence of the user, investor and the insurer.

Equitable regulatory environment: Incentive to invest and install electrical energy storage requires a reliable revenue generation model or roadmap for storage operators. This is largely lacking due partly to inconsistent pricing and market plans, and also to uncertainty in the use-case economics, all of which deter investment. Hence, the value proposition for adopting storage systems for the grid relies on lowering the regulatory barriers.

Industry acceptance: This arguably poses the highest barrier among all. Due to uncertainties about how well these systems perform in practice and over time, system operators many of whom have no or only limited experience with them, are skeptical about adopting storage technologies. Equally important is the inability of today's utility planning, transmission and distribution tools to analyze energy storage as an integral part of their portfolio. Naturally, integration of storage options into industry's planning tools helps build confidence in storage

technologies. Acceptance and adoption of grid storage by the industry requires verified confidence that the storage system will deliver and perform as predicted and expected.

A recent US DOE workshop has identified several near- and long-term cost and performance targets for grid applications in order to aid and guide storage technology researchers and developers.¹⁷ In the near term, the workshop report recommended demonstration of AC energy storage systems involving redox flow batteries, sodium-based batteries, lead-carbon batteries, lithium-ion batteries and other technologies to meet the following electric grid performance and cost targets; system capital cost – under \$250 per kW h, levelized cost – under 20 ¢ per kW h per cycle, system efficiency – over 75%, and cycle life – more than 4000 cycles, with system capital cost targets under \$1750 per kW. For the long term, the workshop's recommendations included research and development for new technologies based on advanced materials and chemistries to meet the following AC energy storage system targets; system capital cost – under \$150 per kW h, levelized cost – under 10 ¢ per kW h per cycle, system efficiency – over 80%, cycle life – more than 5000 cycles, with system capital cost targets under \$1250 per kW. It also recommended system capital cost of <\$15 per kW h, system efficiency of 95%, cycle life of 10 000 cycles for concentrated solar power (CSP) storage systems.

These targets, especially for emerging storage technologies are ambitious to meet, and to gain commercial interest storage systems need to add value and be competitive with existing storage technologies. Also, to make progress, it is important to understand the inherent limitations and factors that govern the technical and economical viability of these storage technologies. As a case in point, the global incentive for electric vehicles is driven largely by the commercial availability of Li-ion batteries. But it is estimated that replacing the gasoline engines in transportation vehicles with 15 kW h Li-ion batteries would use up more than 30% of the world's lithium reserves.³¹ This may pose a serious limitation. The cost of storage is also a major concern. Although the cost, for example, of the Li-ion battery pack for a 240-mile driving range was estimated several years ago to be nearly \$30 000 at \$400–800 per kW h, this has steadily decreased to \$273 per kW h in 2016.⁴²⁶ However, this is still too expensive, compared to the cost target of <\$100 per kW h.^{241,425} Certainly, source, cost and performance limitations of storage systems need to be overcome. Basic research that can address the challenges and meet the required cost and performance targets for these and other applications will help move practical storage technologies forward.

I. Concluding remarks and summary

The 2003 blackout in the US northeast that immersed New York City and the surrounding states into darkness, and the 2011 power failure in southern California and Arizona are recent examples of how critical is the availability of cost-effective, reliable electrical power for our living standards, security, economy, and industry. Clearly, effective energy storage systems would

have helped mitigate such major disruptions. But more so, it is widely agreed that electrical energy storage will help catalyze rapid transformation of the global energy infrastructure towards clean energy technologies, achieve efficient load-leveling and regulation, accelerate mass penetration of renewables, realize distributed generation and local grids, and improve energy security.

A case in point is the explosive growth of wind and solar as well as electric vehicles, which in large part is enabled by storage. Deployment and integration of intermittent renewables into the grid without effective storage capabilities poses major challenges, and compromises the stability and integrity of the aging grid systems. Storage can serve as a buffer between electricity supply and demand, and provide flexibility of the grid to allow greater integration of variable renewable resources. Storage also allows power generation, when needed, at or near load centers, thereby augmenting the electricity infrastructure with distributed generation. Technology advances to date provided significant cost reductions. However, full adoption and deployment of storage technologies still requires that the users take advantage of multiple benefit streams.

Several extensive techno-economic analyses recently claimed that it is possible to realize low-cost grid reliability with 100% penetration of renewables,⁴²⁸ and further proposed an electrification roadmap to achieve 100% renewables (wind, water, solar) for 139 countries in the world.⁴²⁹ However, the methodology and assumptions employed in these analyses as well as their assessments and conclusions came under question and were challenged by others.⁴³⁰ It is generally agreed that transition towards a decarbonized energy economy will require the development of a multitude of clean electricity generation and storage technologies. Indeed, as much diversity as there is in power generation, a broad portfolio of electrical energy storage technologies and systems is needed to respond appropriately and cost-effectively to the wide spectrum of specific requirements of divergent electricity production systems. Each electrical storage technology offers its unique set of advantages and challenges, as well as operational characteristics such as discharge times that may be best suited for a particular application. This is depicted in Fig. 7 above, and also in Fig. 58 and Table 12. For example, flywheels and supercapacitors can provide high power with fast response times in a matter of seconds to respond to fluctuations in demand but have low energy densities, whereas metal-air and flow batteries offer high energy densities. In other words, the choice for the right system requires matching the system performance characteristics and cost, with the requirements of the desired application. Many of these electricity storage applications and their characteristic system and storage requirements are summarized in Table 12, where the terminology “arbitrage” refers to the trade practice in the energy business whereby electricity that was bought (and stored) at low price during low demand is sold back to the market during high-priced demand, “voltage support” refers to injection and absorption of reactive power to the system to maintain the desired voltage levels in the distribution and transmission systems, and “black start” refers to restarting the electrical supply system after catastrophic power

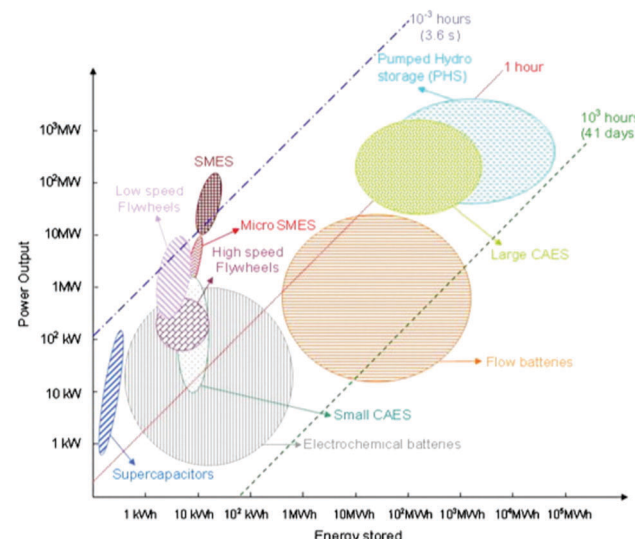


Fig. 58 Distribution of storage technologies according to their energy stored and power output, and typical discharge time windows.²² Reprinted from H. Ibrahim, A. Ilinca and J. Perron, Energy storage systems – characteristics and comparisons, *Renewable Sustainable Energy Rev.*, 2008, **12**, 1221–1259, with permission from Elsevier.

failure without pulling power from the grid. Likewise, “spinning reserve” refers to the response time of less than 15 minutes for the stored reserve electricity capacity to kick in in order to compensate for the loss of generation capacity elsewhere in the system, and “non-spinning” reserve relates to response times longer than 15 minutes.

For comparison and compatibility purposes, Table 13 presents typical performance characteristics of various energy storage systems and technologies, which show widely varying ranges of performance parameters. As also discussed in Section H.: Grid storage, the storage times and capacities vary widely depending on the application. Comparing Tables 12 and 13 makes it clear that no single storage system or technology can alone meet these requirements fully, *i.e.*, there is no silver bullet. This necessitates the development of multiple storage technologies to fully meet the varying needs of grid electricity storage.

Distributed storage offers many advantages, although not all storage systems reviewed in this article are suitable for this purpose. Pumped-hydro and CAES require specific geographic conditions and may not be a viable option available widely. On the other hand, systems such as batteries, regenerative fuel cells, flywheels, and chemical storage may be available regardless of location to couple into renewable energy sources or into the local grid. For example, instead of building a centralized power plant of 1 GW, placing 100 units of 10 kW battery storage capacity in utility service areas would help reduce the loads, but more importantly, eliminate expensive “spinning reserves”, *i.e.*, supplementary power generated typically by gas turbines that fire up to respond to daily spikes in electric demand. Moreover, such distributed storage also helps improve energy security greatly.

The infrastructure for power transmission and distribution in the US as well as in many other parts of the world is old and

Table 12 Characteristic performance requirements for electrical (e) and thermal (t) storage technology applications⁴²⁷

| Application | Output (electricity, thermal) | Size (MW) | Discharge duration | Cycles (typical) | Response time |
|---|-------------------------------------|---------------|-----------------------|----------------------|-----------------------|
| Seasonal storage | e, t | 500 to 2000 | Days to months | 1 to 5 per year | Day |
| Arbitrage | e | 100 to 2000 | 8 to 20 hours | 0.25 to 1 per day | >1 hour |
| Frequency regulation | e | 1 to 2000 | 1 to 15 minutes | 20 to 40 per day | 1 min |
| Load following | e, t | 1 to 2000 | 15 minutes to 1 day | 1 to 29 per day | <15 min |
| Voltage support | e | 1 to 40 | 1 second to 1 minute | 10 to 100 per day | Millisecond to second |
| Black start | e | 0.1 to 400 | 1 to 4 hours | <1 per year | <1 hour |
| Transmission & distribution (T&D) congestion relief | e, t | 10 to 500 | 2 to 4 hours | 0.14 to 1.25 per day | >1 hour |
| T&D infrastructure investment deferral | e, t | 1 to 500 | 2 to 5 hours | 0.75 to 1.25 per day | >1 hour |
| Demand shifting and peak reduction | e, t | 0.001 to 1 | Minutes to hours | 1 to 29 per day | <15 min |
| Off-grid | e, t | 0.001 to 0.01 | 3 to 5 hours | 0.75 to 1.5 per day | <1 hour |
| Variable supply resource integration | e, t | 1 to 400 | 1 minute to hours | 0.5 to 2 per day | <15 min |
| Waste heat utilization | t | 1 to 10 | 1 hour to 1 day | 1 to 20 per day | <10 min |
| Combined heat and power | t | 1 to 5 | Minutes to hours | 1 to 10 per day | <15 min |
| Spinning reserve | e | 10 to 2000 | 15 minutes to 2 hours | 0.5 to 2 per day | <15 min |
| Non-spinning reserve | e | 10 to 2000 | 15 minutes to 2 hours | 0.5 to 2 per day | <15 min |

Table 13 Typical performance characteristics of various energy storage systems and technologies⁴³¹

| | Power rating (MW) | Discharge time | Cycles or lifetime | Self-discharge (%) | Energy density (W h l ⁻¹) | Power density (W l ⁻¹) | Efficiency (%) | Response time |
|------------------------------|-------------------|----------------|--------------------|--------------------|---------------------------------------|------------------------------------|----------------|---------------|
| Pumped-hydro | 100–2500 | 4–16 h | 30–60 year | ~0 | 0.2–2 | 0.1–0.2 | 70–85 | 10 s-min |
| Compressed air | 100–1000 | 2–30 h | 20–40 year | ~0 | 2–6 | 0.2–0.6 | 40–70 | Min |
| Flywheel | 0.001–20 | s-min | 20 000–100 000 | 1.3–100 | 20–80 | 5000 | 70–95 | < s |
| Li-Ion battery | 0.05–100 | 1 min–8 h | 0.1–0.3 | 200–400 | 1300–10 000 | 85–95 | < s | |
| Lead-acid battery | 0.001–100 | 1 min–8 h | 6–40 year | 0.1–0.3 | 50–80 | 90–700 | 80–90 | < s |
| Na-S battery | 10–100 | 1 min–8 h | 2500–4500 | 0.05–20 | 150–300 | 120–160 | 70–90 | < s |
| Flow battery | 0.1–100 | Hours | 12 000–14 000 | 0.2 | 20–70 | 0.5–2 | 60–85 | < s |
| Supercon. magnet | 0.1–1 | ms-s | 100 000 | 10–15 | ~6 | ~2600 | 80–95 | < s |
| Super-capacitor | 0.01–1 | ms-min | 10 000–100 000 | 20–40 | 10–20 | 40 000–120 000 | 80–95 | < s |
| Hydrogen | 0.01–100 | Min–week | 5–30 year | 0–4 | 600 (200 bar) | 0.2–20 | 25–45 | s-min |
| Synthetic natural gas | 1–100 | Hour–week | 30 year | ~0 | 1800 (200 bar) | 0.2–2 | 25–50 | s-min |
| Molten salt (latent thermal) | 1–150 | Hours | 30 year | N/A | 70–210 | N/A | 80–90 | min |

even antiquated. For political, legal, environmental and social reasons, replacing or upgrading the existing grid system is challenging and expensive. Distributed storage may alter this paradigm, and shift power generation from centralized stations that require a vast network of expensive and lossy transmission and distribution systems, to having local control and energy security. Naturally, such distributed mode is easier to manage than a vast national grid, which requires a network of computers working in tandem to control the flow of electricity and are vulnerable to hacking and other malicious attacks.

As economics of storage is as important as performance, the corresponding cost of electricity storage of various electrical storage systems are compared with respect to their round-trip efficiencies and discharge times in Fig. 59. The data suggests that regenerative SOFCs with their high round-trip efficiency, long discharge times, and low storage cost can compete favorably with pumped-hydro and CAES technologies, albeit their maturity is far from being deployed on a commercial scale. Similarly, cost of electricity storage for various rechargeable battery systems are presented in Fig. 60. Although there is considerable uncertainty in the estimated cost figures, Na-S and Li-air batteries show favorable economics, while cost

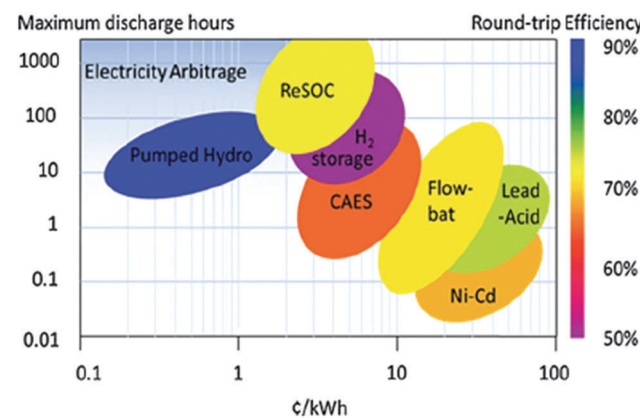


Fig. 59 Comparison of cost of electricity storage, round-trip efficiencies and discharge times of various storage systems.³² Reprinted with permission from (Z. Yang, J. Zhang, M. C. W. Kintner-Meyer, X. Lu, D. Choi, J. P. Lemmon and J. Liu, Electrochemical energy storage for green grid, *Chem. Rev.*, 2011, **111**, 3577–3613). Copyright (2018) American Chemical Society.

reductions in others, such as VRB and Li-ion batteries, are needed for wide scale implementation of these systems. Unfortunately, none of the battery systems singly satisfies the

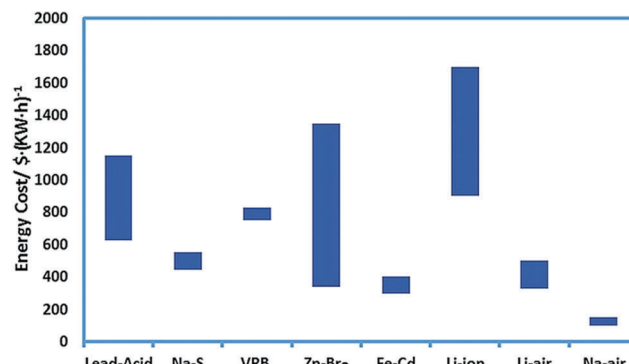


Fig. 60 Energy costs (\$ per kWh) of various battery systems.⁴³² Reprinted from W.-W. Yin and Z.-W. Fu, The potential of Na-air batteries, *ChemCatChem*, 2017, 9, 1545–1553, with permission from John Wiley and Sons.

full set of performance, size and economic requirements for effective electrical storage. Instead, each storage technology reviewed here offers characteristic properties and specific benefits as well as shortcomings and challenges. In other words, there is no silver bullet for electrical energy storage. One size does not fit all. Therefore, a multi-prong technology approach is desirable to adequately address the global needs for electrical energy storage in a comprehensive manner.

For example, there may be merit in considering smaller size batteries in the 5–10 kW range for energy storage for the grid. One prevailing view for megawatt-scale storage batteries is to place them on the high voltage (>10 kV) side of the utility transformer, which would then require multiple step-down transformers and other controls that would complicate the hook-up process. On the other hand, smaller battery units in the 10 kW range can be placed instead on the 100 V side of the transformer, which would render a simple hook up process and enable to achieve switching using inexpensive solid-state relays.

In assessing storage systems, source availability and access also need to be considered in the life cycle analysis. For example, Li-based batteries may be vulnerable to the cost and availability of lithium, which is largely mined and produced in select countries in the world. Other active chemicals, such as cobalt that are currently used in commercial Li-ion batteries pose also concerns regarding cost, scarcity, and availability. Other battery systems employ toxic elements in their electrodes such as lead, cadmium, and mercury, which are of great health and environmental concern. Precious metal catalysts used in other electrochemical storage systems such as low temperature aqueous electrolyzers are also expensive and source-dependent elements. So recycling battery components may supplement availability, and is definitely of practical, environmental and economic interest. For example, it takes 250 tons of mineral ore and 750 tons of brine to produce 1 ton of lithium, compared to only 28 tons of used Li-batteries.⁴³³ There have been recycling efforts to collect and re-process used batteries to recover active elements. However, for wide scale and cost-effective recovery of these spent resources, we need to develop more efficient, higher yield and inexpensive recycling processes, and also implement global environmental policies that mobilize a concerted recycling effort.

In a manner similar to cost of storage, another important consideration should be given to the amount of emissions produced during the life of storage systems. For example, based on life cycle analysis results it was estimated that to make a 1 kW h Li-ion battery, nearly 400 kW h of energy is consumed,⁴³³ resulting in 75 kg of CO₂ emissions during its production. Simple arithmetic suggests that the service life of the Li-ion battery needs to be >400 cycles to pay back the energy in terms of storage. By comparison, a coal-fired power plant emits only 1 kg of CO₂ for every 1 kW h of electrical energy it produces. Accordingly, these comparative examples highlight the importance of long service life for the storage system before its benefits can have a positive impact on the environment.

It should be noted that theoretical energy densities for electrochemical storage systems are usually calculated based on the weight and volume of the active materials, which constitute only a portion of the final packaged product. Hence, real energy densities are necessarily lower than those calculated theoretically. This is illustrated in Table 14, where average efficiencies for major systems are in range of tens of %. Only, Li-ion batteries seem to reach a relatively high energy utilization level between 42 and 58%. Surely, there is room for major improvements in active materials for electrochemical storage technologies that will help increase storage efficiencies of these systems. Better sealing and packaging processes and cost-effective materials will also help reduce this discrepancy between calculated and actual utilization capacities. So more work and innovation is needed to improve existing technologies and expand the storage options. For example, in most electrochemical storage systems such as batteries, only single-electron transfer reactions have been utilized. In today's Li-ion batteries, for example, less than only one Li⁺ ion is intercalated per transition metal redox center in the cathode material. Obviously, this limits the ultimate capacity of the battery. A recent article challenges this paradigm and suggests that for next generation batteries, it may be worth exploring multi-electron transfer chemistries, by either intercalating multivalent cations per redox site or intercalating multiple monovalent cations.⁴³⁴ Certainly, accomplishing this goal will greatly improve the practically realizable capacity for electrochemical storage. Similarly, development of earth-abundant materials with superior catalytic properties for regenerative fuel cells, electrolyzers for hydrogen production, electrochemical

Table 14 Theoretical and actual energy densities and actual utilization percentages for select electrochemical storage with various battery systems compared with the H₂/O₂ fuel cell⁴⁴

| System | Calculated energy density (W h kg ⁻¹) | Real energy density (W h kg ⁻¹) | Utilization (%) |
|--------------------------------|---|---|-----------------|
| Pb-Acid | 171 | 25–55 | 15–32 |
| Na-S | 792 | 80–150 | 10–19 |
| Ni-MH | 240 | 50–70 | 20–29 |
| Li-Ion | 360 | 150–210 | 42–58 |
| Li-S | 2654 | 250–350 | 9–13 |
| Li-O ₂ | 5217 | — | — |
| Zn-O ₂ | 1094 | 150–200 | 14–18 |
| H ₂ -O ₂ | 3525 | 120–800 | 3–23 |

Table 15 Major electrochemical storage projects that are announced, contracted or under construction by technology and country¹⁶

| Country | Electrochemical (unspecified) | Electrochemical capacitor | Lithium-ion battery | Flow battery | Vanadium redox flow battery | Lead–acid battery | Metal–air battery | Sodium-based battery | Total (kW) |
|-------------------|-------------------------------|---------------------------|---------------------|--------------|-----------------------------|-------------------|-------------------|----------------------|------------|
| United States | 500 398 | | 61 959 | 3030 | 20 250 | 21 500 | 14 250 | | 621 397 |
| Australia | 122 010 | | 9400 | | | | | | 131 410 |
| Germany | 30 000 | | 92 000 | 210 | | | | | 122 210 |
| India | 110 000 | | 125 | | | | | | 110 125 |
| Republic of Korea | | | 48 500 | | | | | | 48 500 |
| Canada | 12 150 | | 12 010 | 4000 | 5000 | | | | 33 160 |
| Egypt | | | 30 000 | | | | | | 30 000 |
| Italy | | 1920 | 20 000 | 1950 | | | 4000 | | 27 870 |
| Kazakhstan | | | | 25 000 | | | | | 25 000 |
| United Kingdom | 1000 | | 20 300 | 140 | | | | | 21 440 |
| Top 10 | 775 558 | 1920 | 294 304 | 34 330 | 25 250 | 21 500 | 14 250 | 4000 | 1 171 112 |
| World | 784 258 | 2920 | 333 404 | 34 965 | 25 250 | 21 500 | 5650 | 4800 | 1 212 747 |

reactors for ammonia synthesis, highly conductive, stable and selective membranes used in most electrochemical storage systems, and cost-effective EDL and pseudocapacitive electrode materials for supercapacitors will help expand the materials space, improve device performance, and bring down systems costs. In that regard, two additional tools, namely theoretical and computational simulations guiding experimentation and rational design of materials, as well as nanostructuring to control and enhance desirable properties will help shorten the pathway to innovation and discovery. Moreover, nanostructured materials may form highly active metastable structures and exhibit properties different from bulk, and offer attractive opportunities in overcoming some of the kinetic and mass transport limitations observed in energy storage systems.

Certainly, storage should be an essential component of the global energy systems of the future, especially if renewables were to take on 80% of the global need.¹⁶ The individual maturities, or technology readiness levels (TRL), of storage technologies vary widely as illustrated in Fig. 5. Similarly, the US DOE's view of the evolution, cost reduction and market penetration of storage technologies is expected to follow the trends depicted in Fig. 7, *i.e.*, there will be both short term and long term milestones and goals that need to be achieved.

In recent years, there has been an increasing drive to build more storage capacity. Some of the projects around the world that are based on electrochemical energy storage are summarized in Table 15 and categorized according to the storage system and country.¹⁶ It is clear that the installed capacities provided in Tables 2 and 14 are quite low compared to the storage needs for solar and wind (see Table 1) that may require durations between hours to days, and in some remote applications, even weeks.

The variability of the storage technologies, systems and materials make it hard to predict which technologies will excel, but successful commercialization of any one or more of the different systems will likely depend on the confluence of several important factors. Certainly, the technical challenges discussed and reviewed in this article must be overcome to achieve reliable, robust, long service and cycle life, efficient, and scalable technology options. Likewise, materials and fabrication costs must be reduced further for many of these systems and technologies to become truly competitive. Large investments both public and

private are required to achieve these goals. Market forces of supply and demand fueled by the rapid expansion and growth in renewables such as solar and wind, will also drive the need for large-scale storage. But this may still not be sufficient to achieve complete decarbonization of the electricity infrastructure, which we urgently need to make happen. Without globally accepted policies to implement an agreeable price on carbon emissions, storage may not receive the attention or the investment it deserves, and we may not achieve complete decarbonization of the global energy system, risking undesired consequences.

In summary, as much as we need a broad range of electric power technologies to meet our needs, we need an equally diverse portfolio of energy storage technologies with matching operational characteristics and properties. The urgent need for large-scale cost effective storage has become increasingly pronounced by the rapid growth and deployment of renewable power generation sources. It is difficult to predict which of the energy storage technologies will dominate the future, but most likely, a broad range of storage systems will be needed to accommodate the specific but diverse requirements of the equally diverse power generation technologies we employ today, and in the future. Meeting the requirements of such a rich diversity of individual system characteristics certainly poses technical challenges as discussed in this article, but it also provides a wide range of options and prospects for storage applications, especially when these technologies may be employed in synergistic combinations that complement each other's strengths and compensate for their deficiencies. It also offers opportunities for innovation and advancements in storage technologies, materials and processes, where the scientific community has an important role to play.

Conflicts of interest

The author has no conflicts to declare.

Acknowledgements

The author gratefully acknowledges partial support from the Stanford Energy 3.0 Program, and many helpful discussions with Prof. Friedrich B. Prinz of Stanford University.

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