Metal-Organic Frameworks for Fast Electrochemical Energy Storage: Mechanisms and Opportunities

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THE BIGGER PICTURE

Electrochemical energy storage devices are typically based on materials of inorganic nature which require high temperature synthesis and frequently feature scarce and/or toxic elements. Organic-based materials on the other hand can provide an attractive alternative, potentially yielding sustainable, safe, and cost-effective energy storage devices based on abundant elements (e.g. C, N, O, S, and H). However, attempts to incorporate organic and coordination compounds so far have led to sub-par cycling stability and charging rates due to insufficient structural and (electro)chemical stability, low electrical conductivity, and reduced performance at industrially relevant device scales.

In recent years metal—organic frameworks (MOFs) have gained attention as having the potential to rival or even supersede traditional energy storage materials. Functional properties such as electronic or ionic conductivity can be incorporated into these materials by judicious design of their constituent inorganic and organic building blocks. However, full realization of the potential of MOFs for electrochemical energy storage requires joint expertise from distinct fields. In particular, bridges must be formed between electrochemists and synthetic and material chemists to establish the unified approach necessary to develop MOF-based energy storage devices exhibiting competitive performance.

INTRODUCTION

Energy storage devices having high energy density, high power capability, and resilience are needed to meet the needs of the fast-growing energy sector.¹ Current energy storage devices rely on inorganic materials² synthesized at high temperatures² and from elements that are challenged by toxicity (e.g. Pb) and/or projected shortages of stable supply (e.g. Li, Co).³ In this context, systems that do not rely on Li as the charge carrier ion, such as those based on Na, and utilize organic-based materials as electrodes provide a conceptually attractive alternative.⁴,⁵ These systems can be made fully from abundant elements such as Na, C, N, and O and can be potentially produced by sustainable methods.⁴,⁵,⁶ Practical implementation of systems that possess such compositions remains challenging for several reasons: 1) transition to lithium-free energy systems requires the design of materials that are *capable of reversibly accommodating larger* cations;⁵-7 2) insufficient *chemical*³ and *electrochemical stability*⁵ of organic-based materials leads to unsatisfactory cycle life⁶,⁰; and 3) an additional set of requirements is imposed on material properties and electrode and device architectures for fast electrochemical energy storage (EES) applications (e.g., high power).¹0 In addition to a high density of redox active sites (and hence, high capacities), these sites should be readily accessible to both ions and electrons to achieve fast charging kinetics.⁵,¹¹¹ Hence, the material should have sufficient electronic conductivity and ion accessibility (Figure 1).²,⁰

Recently, metal–organic frameworks (MOFs) have emerged as a distinct class of materials for electrochemical energy storage. ¹² MOFs are constructed *via* coordination bonding between nodes (metal ions or ionic metal clusters) and ligands (organic molecules) that results in the formation of crystalline networks. The coordination geometry between metal node and ligand define the MOF's network topology and potential porosity. ^{13,14} The chemical identity of the nodes and ligands, along with MOF topology, play a deterministic role in the ultimate functionality of the material. Many MOFs possess accessible porosity and high specific surface areas; ¹⁵ consequently, their applications in gas storage, ¹⁶ fluid separations, ¹⁷ and catalysis ¹⁸ have been extensively studied. MOFs designed for these applications typically feature metal ions bound with redoxinactive ligands with *via* hard ¹⁹ coordinating motifs such as carboxylate ²⁰ or imidazolate, ²¹ yielding robust, porous frameworks that however lack the electrical conductivity and redox activity necessary for fast electrochemical energy storage. ²²

MOFs for fast EES must fulfill the set of stringent requirements outlined above to be competitive with (or superior to) state-of-the-art systems. Meeting these requirements necessitates an altogether different and distinct approach to MOF design. For example, learning from design principles of conductive coordination

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polymers, electrical conductivity can be introduced.²² An emerging class of electrically conductive MOFs shows a marked improvement in performance over their non-conductive counter parts and demonstrates substantial promise for high-rate energy storage.^{14,23} Along with electrical conductivity, density and accessibility of the redox active sites and MOF stability under operating conditions are equally critical to consider at the design stage.

This review aims to bridge the knowledge gap between electrochemists and scientists working in the field of metal–organic frameworks. We introduce basic concepts of energy storage devices and electrochemical characterization techniques. We then discuss design principles of MOFs in the context of what parameters are important to consider for their use in fast EES devices. Given the nascence of this field, many questions regarding energy storage mechanisms remain unaddressed, such as the effect of redox-active centers and porosity (e.g. size, shape, and pore volume) on device performance. Further, we identify opportunities for the rational design of MOFs for energy storage applications.

FARADAIC AND CAPACITIVE CHARGE STORAGE FOR FAST CHARGING

Knowledge of distinct charge storage mechanisms and understanding their advantages and drawbacks is critical to enable the design of next generation energy storage materials.²⁴ Fundamental differences in the operation principles exist between faradaic and non-faradaic charge storage mechanisms (**Figure 1**). As the name suggests, the former involves redox reactions that follows Faraday's law (in bulk or at the surface of the material) while the latter is a redox-free physical mechanism termed electrical double layer (EDL) capacitance.

The EDL mechanism relies on charge separation in the electrical double layer upon polarization that is achieved through the reversible electrosorption of ions from the electrolyte to the outer and inner (porosity) surfaces of the material (**Figure 1**). Therefore, to maximize EDL capacitance, electrode materials with high specific surface area (SSA) and good ionic accessibility are used. For example, highly porous carbons with SSA >1500 m²/g demonstrate the highest EDL capacitances²⁴ and are used as active materials in commercial supercapacitors (also known as electrochemical capacitors and ultracapacitors).²⁴ As the EDL capacitance does not involve redox processes, this mechanism is inherently fast.²⁴ As a result, devices that use EDL capacitance to store charge (e.g. supercapacitors) benefit from fast charging and are characterized by high power operation and excellent cycling stability (>100′000 cycles).²⁴ Electrical double layer capacitors (EDLCs) are characterized by the rectangular shapes of their cyclic voltammograms (CVs) and linear galvanostatic charge discharge profiles (**Figure 1**).

Electrode materials in batteries on the other hand rely on redox reactions to store charge (i.e. the operation mechanism is faradaic) that are accompanied by ion insertion/intercalation and phase transformations.^{2,11,24} Redox-based charge storage is characterized by large specific capacities and an order of magnitude higher energy densities compared to supercapacitors. However, due to unavoidable phase transformations the operation of battery materials is limited by solid-state diffusion²⁵ during charge/discharge cycling.^{2,11,24} Furthermore, insufficient ionic and electronic conductivities of common battery electrode materials make them inherently slower than supercapacitors and impose limitations on the specific power achievable by batteries.^{2,26} CVs of battery electrode materials are characterized by well-separated redox peaks, while galvanostatic charge discharge profiles feature plateaus at the corresponding redox potentials. Performance trade-offs between supercapacitors and batteries can be visualized using Ragone plots that use specific energy and power as their corresponding x-and y-axes.²⁴

Pseudocapacitors are a sub-class of electrochemical capacitors that rely on fast redox reactions for charge storage. Their electrochemical features resemble those of EDLCs (**Figure 1**), however, unlike EDLCs, the charge storage mechanism in pseudocapacitors is faradaic in nature, enabling pseudocapacitors to store 5-10 times more charge compared to EDLCs. Depending on whether redox processes occur at the material surface or within the bulk, pseudocapacitance can be classified as surface redox or intercalation, respectively. ^{10,24} Both types are characterized by sloping galvanostatic charge-discharge profiles (**Figure 1**). Pseudocapacitors can potentially bridge the gap between batteries and EDLCs with energy density approaching that of batteries and power densities reaching those of EDLCs. It is important to emphasize however a key difference between batteries and pseudocapacitors as significant confusion exists in the literature: while both utilize faradaic mechanisms, pseudocapacitors are characterized by fast charging kinetics not limited by diffusion and do not undergo phase transformation during charge/discharge cycling. ¹¹ Moreover, only a select few materials show true pseudocapacitive behavior as they have to combine good electronic and ionic conductivity to display fast charging kinetics. ^{10,27,28} Known examples of pseudocapacitive systems illustrate that ion transport to a redox

active site can be assisted by the presence of (i) structural solvent²⁹, e.g. hydrated RuO_2^{30} , 2D transition carbonitrides (MXenes)^{11,31} or (ii) low-barrier ion diffusion channels, such as in T-Nb₂O₅.³²

The design of fast energy storage devices (that rely on the outlined mechanisms to store charge) requires an understanding and optimization of the many interdependent factors that ultimately define the final performance of the device. In the next section we introduce these factors.

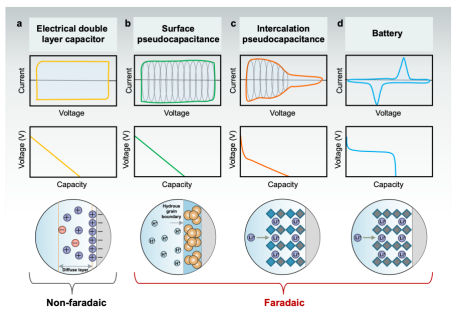


Figure 1. Faradaic and capacitive charge storage. Characteristic cyclic voltammetry and galvanostatic profiles and^{28,33} schematic diagrams depicting characteristics of each charge storage mechanism: (a) electrical double layer capacitor, (b) surface pseudocapacitance, (c) intercalation pseudocapacitance, (d) battery.

BASIC CONCEPTS OF ENERGY STORAGE DEVICES

Factors affecting performance of the electrochemical energy storage device

Researchers in academia and industry alike target concurrent improvements in the specific energy, power, and durability of electrochemical EES devices, all while lowering their overall costs. ^{34,35,} All of these device characteristics are typically presented as normalized by total device weight and volume. When comparing the performance and commercial viability of fast EES devices, it is important to take all these metrics into consideration. ³⁴ EES devices are constructed from the following key components: (i) Positive and negative electrodes that store charge. These electrodes are made of a mixture of active material, conductive additives (e.g. carbon black) that improve the electronic conductivity of the electrode, and a small amount of polymeric binder (typically PTFE or PVDF) that maintains the structural integrity of the electrode, (ii) electrolyte whose function is to conduct ions, (iii) a separator that prevents electrical contact and therefore short circuiting between the positive and negative electrodes, (iv) current collectors that ensure electrical contact to the active material, and (v) device casing. ^{2,36,37} All these components contribute to overall device weight and volume and therefore the respective device metrics (outlined below). ³⁷ Therefore, to achieve maximized performance all device components and their relative ratios require special optimization. ³⁷ In Figure 2 we depict the key aspects considered for device optimization at the electrolyte, electrode, and active materials levels, highlighting the interdependence of these factors on overall device performance.

Cost. While topic of cost optimization is outside the scope of this review, in **Figure 2** we list important factors that are known to affect the final device cost³⁸: while element scarcity represents a foundational cost-determining parameter, supply chain efficiency and scalability of material production are equally important as they can directly affect the cost of the device.³⁷. Another parameter that is increasingly taken into consideration is related to the recyclability. An exponential increase in EES device usage must be followed by an exponential increase in the number of end-of-life EES devices that must be disposed or recycled in sustainable, cost-effective ways.^{38,39}

Durability. The number of cycles over which a fast EES device can be operated before losing 20% of its original performance (the cycle life) is one of the main metrics used to characterize the device durability. ⁴⁰ Cycle life depends on the charge-discharge conditions (current density, cut-off voltages, temperature) as these

can directly affect coulombic efficiency by prompting irreversible reactions associated with electrode and electrolyte decomposition (*vide infra*).^{2,40} Other industry-relevant durability characteristics are self-discharge (characterized by the capacity loss with use)^{24,41} and calendar life (measurement of capacity retention during a fixed period of time)^{42,43}, safety tests such as hot box test (thermal behavior monitoring under heating condition at 180 °C)^{44,45} and nail penetration test⁴⁶ (to simulate internal short-circuit) are beyond the scope of this review.

Energy and fast charging capability (power). The specific energy of the device, W, is directly proportional to specific cell capacity, Q, (charge stored per total device volume or mass) and cell voltage V: W=QV for a battery and $W=\frac{1}{2}QV$ for a supercapacitor.^{47,48} Improving energy performance involves maximization of 1) the capacity of the positive and negative electrodes, 2) improving the volume and mass fractions of active charge storage material (e.g. higher electrode thicknesses/loadings), while reducing the fraction of inactive compounds (electrolyte, current collectors, casing)^{40,49} and 3) increasing the cell voltage.⁴⁰ The specific power of the device refers to its ability to deliver energy per unit of time and is proportional to energy density as a function of the charging rate (current density).²⁴ Specific power is affected by ion and electron transport within the electrodes and cell (vide infra).^{10,24}

Capacity. For redox-based charge storage, electrode capacity primarily depends on the charge storage capability of the active material, i.e. the density of the redox active sites per unit mass. 11,50 It can be estimated theoretically using the following equation: $Q_{theoretical}(mAh\ g^{-1}) = \frac{n\times F}{3.6\ W_F}$, where n is the number of electrons that material gains or losses during charging per formula weight (W_F) of the active material and F is the Faraday constant (96'485 C mol⁻¹). 2,10 The theoretical capacity can serve as a first benchmark to estimate whether a material might be attractive for energy storage applications. However, experimental values of specific capacity rarely reach theoretical values, in particular at high charging rates, due to the inherent difficulty of establishing excellent ion and electron accessibility at each redox active site and/or due to the instability of the material after electrochemical cycling. Therefore, materials for fast EES must have sufficiently high electronic conductivity. Furthermore, since most redox reactions require (de)insertion of ions to maintain charge balance, high ionic conductivities as also desirable in addition to a high density of redox active sites.

Electrode loading. Effective ion transport and electronic conductivity are equally important at the electrode level. For practical cells it is critical to demonstrate the scalability of active material performance at industry-relevant loadings of >10 mg/cm².⁵¹ Doing so necessitates inclusion of conductive additives (e.g., carbon black) for active materials possessing lower conductivity.⁶ While generally having larger content of conductive additives improves power characteristics when normalized just per active material weight, when total electrode weight is considered, overall electrode capacity can drop.^{11,51} Effective ion transport within the electrode is closely related to effective diffusivity that is a product of ion diffusivity within electrolyte and tortuosity: Deff=Dt.⁴⁹ Tuning the tortuosity of the electrode will have significant impact on ionic transport.

Voltage. The difference between the positive and negative electrode potentials defines the cell voltage and increasing the cell voltage constitutes one of the ways to improve the energy density of the device.²⁴ For redox-based energy storage, selection of positive and negative electrode materials that yield large differences between their reversible redox potentials is one of the ways to increase overall energy density.⁵² To ensure high charging efficiencies, irreversible reactions at the electrodes (e.g., leaching of cations or ligands, structure collapse, electrolyte decomposition) that lead to loss of capacity with cycling and should be avoided.^{6,10} In order to find out experimentally the *reversible* redox potential window for the active material, cycling voltammetry or galvanostatic cycling can be used in three-electrode configuration (that allows to separately track electrochemical behavior of the material of interest).^{53,54} Selection of positive and negative cut-off potentials can strongly affect the coulombic efficiency (defined as discharge capacity over charge capacity × 100) and cycle life (defined as the plot of discharge capacity with cycle number).^{11,25,41} Therefore, tests with variable cut-off voltages should be conducted to identify the optimal potential range in which the active material experiences the least degradation.⁴¹

Electrolyte. Electrolyte selection affects cell performance on the multiple levels. (i) To maximize energy density, the amount of electrolyte must be minimized while still ensuring proper electrode material wetting for optimal ion accessibility to active sites. 40 High ionic conductivity is necessary for fast charging (ensuring high rate performance and thus power). 55 (ii) The voltage window in which the electrolyte is stable (as defined by the electrolyte's oxidation and reduction decomposition potentials) constitutes another critical parameter. 56 This voltage window is affected by solvent selection and electrolyte concentration. 56 Electrolytes based on organic solvents (e.g. acetonitrile or organic carbonates) and ionic liquids offer higher stability

windows (> 3 V) compared to water, that has a thermodynamic stability window of 1.23 V.⁵⁷ To ensure high charging efficiency, the electrode redox potentials should fall within the electrolyte stability window. Otherwise, charge would be spent on electrolyte decomposition.¹¹ (iii) The electrolyte flash point is an important safety characteristic, and its low values, as in the case of organic electrolytes, can lead to safety concerns and limit device operation temperatures.⁵⁵ (iv) Electrolyte composition can also affect the chemical and electrochemical stabilities of the active material.⁵⁵ Careful selection of the electrolyte can help to suppress degradation and thus dramatically affect the cycle life of the device.⁵⁶ It is important therefore to first assess the performance of the material in a range of electrolytes.⁵⁶

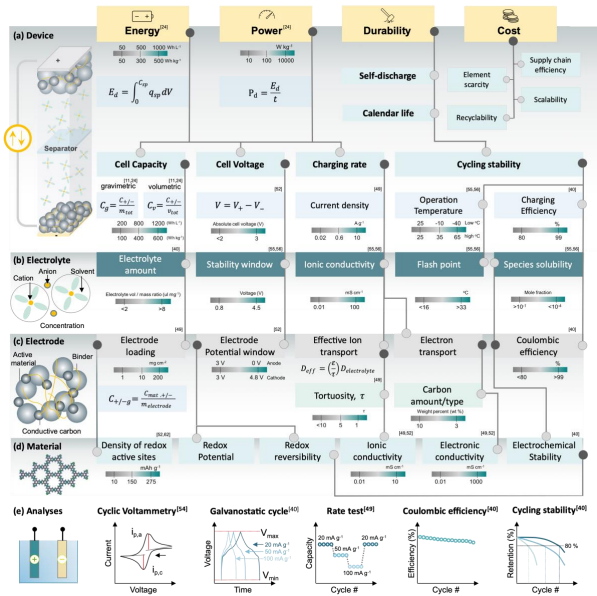


Figure 2. A guideline depicting the interconnected nature of how key characteristics of energy storage devices (a) are affected by electrolyte (b), electrode (c) and active material (d) selection as well as typical electrochemical characterizations used to assess them (e). Connecting line scheme: filled dark grey symbols indicate that metrics are dependent on the parameter marked by the light grey symbol. Bar gradients provide reference values, where teal regions correspond to the most desirable range, while grey indicates less desirable values.

DESIGN OF MOFS FOR FAST ELECTROCHEMICAL ENERGY STORAGE

As was emphasized above, sufficient electronic and ionic conductivity, a high density of redox active sites, and electrochemical stability are all necessary characteristics of next generation materials for fast EES. MOFs can potentially fulfil all these criteria. Below, we outline how each of these desirable characteristics can

be expressed in MOFs. The *a priori* design of MOFs for fast EES is challenging, and successful candidates have thus far been based on a narrow subset of possible conductive MOF candidates (*vide infra*). However, the parameter space for new MOF design is essentially limitless (**Figure 3a**). We provide our personal perspective on key design rules for new MOFs for fast EES, with the aim to increase both the diversity of MOFs studied to this end, and to improve their performance to an extent where they might achieve commercial relevance.

Redox Characteristics. For pseudocapacitive behavior, maximizing the density of redox-active sites is important for high device performance. Achieving this aim requires careful consideration of several factors that are discussed below.

To a first approximation, extending ligand length without changing the ligand's symmetry or coordination motifs will lead to MOFs having larger pores, but otherwise identical topology. ⁵⁸ The relevance of this paradigm has been confirmed for 2D conductive MOFs. ⁵⁹ Larger pores result in higher accessible surface area (see below) and therefore can enable increased contributions from double layer capacitance to charge storage. Furthermore, larger pores can provide improved ion accessibility to the redox active sites critical for fast charging. Yet, despite the noted advantages of larger pores and higher accessible surface area (see below), such characteristics come at the cost of inherently reducing the volumetric density of redox-active sites in the MOF. Moreover, where ligand extension does not concomitantly include additional ligand-based redox states, the gravimetric density of redox-active sites is also reduced. Hence, smaller ligands generally lead to greater capacity as long as redox-active sites remain accessible.

The redox characteristics of both metal and ligand are also important for maximizing the density of redox-active sites within the MOF. Both the ligand and metal (or metal cluster) should ideally be redox active and contribute to charge storage to maximize energy density. This aim can be achieved by redox matching, where both transition metal and ligand are redox-active in the same potential range.⁶⁰ While redox matching has been cited as enabling high conductivity in conductive MOFs⁶¹, direct evidence for matching ligand and metal redox potentials is still lacking. In fact, several authors have shown that pseudocapacitance in several 2D MOFs used for fast EES is ligand-centered.⁶²⁻⁶⁴ The discovery of redox-matched motifs can be aided by the study of small molecule coordination complexes having metal-ligand binding motifs analogous to those used to construct MOFs. Electrochemical studies of such complexes are amenable to solution-based spectroscopy methods to assess oxidation state changes in metal and ligand.^{23,61,65}. For example, a model tri-nickel hexaoxytriphenylene (HOTP) complex, used as an analogue for the conductive MOF Ni₉(HOTP)₄, exhibits welldefined redox waves during electrochemical cycling.65 A combination of CV and electron paramagnetic resonance (EPR) spectroscopy was used to show that redox is ligand-centered, with the oxidation state of Ni²⁺ remaining unchanged. In situ X-ray absorption spectroscopy can be used to determine whether changes in metal oxidation state occur at potentials identical to those at which ligand oxidation state changes are detected (discussed below). It is important to keep in mind that changes in the metal oxidation state can lead to changes in coordination geometry and therefore structural changes. Hence, structural characterization (e.g., via PXRD) of MOFs before and after cycling, and in each accessible redox state is necessary to determine whether structural changes do in fact occur upon cycling and if they do, whether or not they are reversible

As discussed earlier, the energy density of a device is not only proportional to electrode capacity but also to cell voltage. Hence, the redox potential(s) of a MOF must be considered during its design. To maximize cell voltage, the redox potentials of the MOF-based anode material should be minimized and those of the cathode material maximized (as discussed above). Since MOFs consist of inorganic and organic components, the potential ranges of both the ligand and metal redox activity strongly depend on their respective identities. The redox potentials for ligands and transition metals, determined computationally (by DFT)^{6,11} or experimentally (by CV)^{23,66}, can serve as starting points for the expected range of redox activity of the parent MOF. It is important to be cognizant of the fact that MOF assembly from these components can change these redox potentials. For example, even if the transition metal does not directly participate in charge storage, it can still influence the potentials at which ligand redox occurs in the MOF.²³ DFT calculations can be useful to pre-screen the relative effects of metal-ligand coordination on redox potentials that might ultimately be observed in the related MOFs. Finally, the redox potential of the MOF must lie within the electrochemical stability window of the electrolyte to achieve stable cycling.

Electronic conductivity. MOFs used for fast EES must have sufficient electronic conductivity. MOFs were initially designed with metal-ligand motifs that are unlikely to support conduction (e.g., cyano- 67 or carboxylate 68 -metal links), but in 2009, the first MOF with substantial guest-independent conductivity (6 × 10 S cm $^{-1}$ at 300 K) was reported. 69 Since then, a diverse set of 2D and 3D MOFs has been discovered with conductivities up to and even exceeding 10 S cm $^{-1}$. 22 Several design strategies have been developed to produce conductive MOFs; these have been divided into through-bond, extended conjugation, and through-

space approaches (**Figure 3b**, each described below).²² Through-bond conjugated MOFs possess unbroken chains consisting of metal cations and ligand heteroatoms (typically N or S) that serve as conduction pathways. The carbonaceous ligand cores are assumed to not participate in electrical conduction. It is important to note that favorable energetic overlap between metal and heteroatom orbitals is necessary to enable effective charge delocalization along such chains. Through bond MOFs featuring metal cations with adjacent oxidation states (e.g. Fe^{2+}/Fe^{3+}) constitute a special case where an additional boost to conductivity can be delivered. For example, $Fe_2(1,4\text{-benzenedipyrazolate})_3^{70}$ has a conductivity of 10^{-2} S cm⁻¹;⁷¹ allowing partial oxidation of the Fe^{2+} cations to Fe^{3+} in air increases the conductivity by two orders of magnitude.⁷²

Extended conjugation refers to the case where charge delocalization is established between conjugated ligands and metal nodes. Similar to MOFs with through-bond conduction, good geometric and energetic orbital overlap between metal ions and non-innocent (redox-active) ligands are essential to achieving satisfactory conductivity.²² Examples include MOFs that showed promising performance for fast EES based on conjugated ligands having multiple *ortho*-dithiol, -diamine, or -diol groups bound to transition metal ions (noting that the metal does not necessarily contribute significantly to charge storage – *vide infra*). However, there is no fundamental reason why other MOFs with conductivity originating from through-bond interactions or extended conjugation could not also be deployed for fast EES. For example, an iron-based triazolate MOF showed substantial conductivity and mixed iron redox states,⁷³ but studies using this MOF for fast EES have not yet been reported. While the pores of this specific MOF are relatively small (ca. 4.5 Å), the use of extended triazole ligands could allow electrolyte ions access to the interior of the MOF without compromising conductivity. The synthesis and structures of numerous other 3D conductive MOFs that could potentially be promising for fast EES has been detailed²²,however, their electrochemical performance has not been described.

MOFs with through-space conductivity possess conjugated organic ligands that are not covalently bound to one another but interact electronically via through-space π -orbital overlap. The iconic series of M_2 (ttftb) MOFs (ttftb = tetrathiafulvalene-tetrabenzoate) is exemplary of this mode of conduction, where conductivity originates from π - π interactions between stacked ligands. To date, however, conductivities in through-space conductive MOFs have been modest, typically not exceeding 10^{-2} S cm⁻¹. It is possible that introducing donor-acceptor motifs akin to those found in charge-transfer salts may produce through-space conductive MOFs with substantially higher conductivity, but this strategy currently poses significant synthetic challenges. Finally, through-space conductivity in MOFs can be introduced by guest molecules such as tcnq (tetracyanoquinodimethane).

It is our current perspective that MOFs based on through-bond and extended conjugation conduction will remain the best performers for fast EES due to their reproducibly high conductivities (achievable without limiting porosity), but improvements in through-space conductive MOF design²² may also lead to promising candidates for fast EES. Guest infiltration to enhance MOF redox characteristics and/or conductivity inherently limits ion access to pores, and therefore in our view is unlikely to compete with current state-of-the-art materials. Finally, it is critical to remember that theoretical capacity estimates (described above) should serve as a guide for the selection of promising motifs that introduce electrical conductivity in a MOF.

Ionic Conductivity. Ionic conductivity is a critical parameter for both pseudocapacitive and EDLC charge storage. To take full advantage charge storage sites throughout the volume of the MOF (not just the MOF particle surface), ionic conductivity within the MOF must be sufficient (Figure 3c). To date, the strategies used to render MOFs ionically conducting are focused on achieving high proton conductivity; the conduction of other ions more relevant for energy storage (e.g., Li⁺, Na⁺, Mg²⁺, etc.) is less studied. To achieve internal ionic conductivity, MOF linkers are typically decorated with acidic functional groups (e.g., carboxylic, sulfonic, or phosphonic acids)⁷⁷ – these functional groups can be added pre-synthetically (where they are part of the linker used to construct the MOF) or post-synthetically (where the MOF linkers are functionalized after the MOF is formed). While the former case can be more straightforward and does not suffer from sub-quantitative functionalization yield, the proclivity of acidic functional groups to coordinate directly to metal cations can lead to interference with MOF formation, rendering pre-synthetic functionalization challenging is some cases.78 Conduction of other cations can be achieved by replacing acidic protons with larger cations (e.g., Li+, Na⁺, Mg²⁺, etc.).⁷⁷ For MOFs with open metal sites (native to the MOF or introduced via defects, ⁷⁹ e.g., as in the accessible Cr cations in MIL-10180), ion conduction can be introduced by functionalization of these sites.81 Ionic conductivity in MOFs can emerge after infiltration with liquid electrolyte. Finally, for MOFs with sufficiently large pores, both electrolyte ions and solvent are expected to populate these pores, in which case ionic conductivity originates solely from electrolyte confined in the MOF pores.82

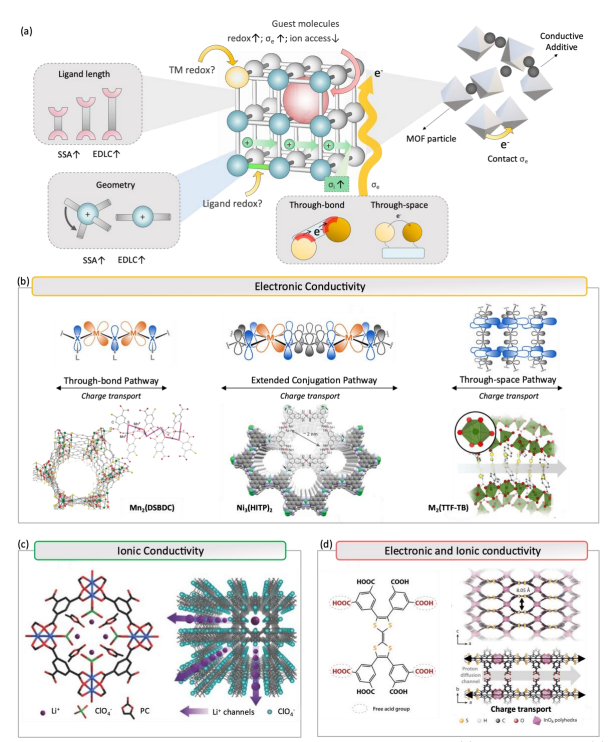


Figure 3. Relationships between electrochemical properties and MOF structure. (a) Schematic diagram. (b) Examples of high electrical conductivity MOFs with extended 2D conjugation. $^{22,87-89}$ (c) An example of a Li⁺ conductive membrane using HKUST-1 with Li⁺ and ClO_4^- bound to the open metal sites. 90 (d) An example of a Li⁺ conductive MOF with a high Li⁺ transference number. 91 (e) Tetrathiafulvalene octacarboxylic acid forming 2D sheets which have both electrical and ionic conductivity. 77

For MOFs with smaller pores, the correlation between a MOF's ionic conductivity and its pore size is not well established due to the difficulty of decoupling the effects of pore size and the distance between ion hopping sites. In a recent report, the proton conductivity was greater for a smaller-pore MOF compared to a larger-pore analogue.⁸² While this report of smaller pores yielding superior H⁺ conductivity is consistent with previous observations for Li⁺⁸³, the opposite trend was observed for Mg²⁺.⁸⁴ Furthermore, the effect of

dimensionality on ionic conduction has to our knowledge lacked systematic study, but using MOFs having 3D pore networks could also improve ion transport compared to MOFs having 1D channel pores of the same size.

Electronic conductivity is also a critical factor for a MOF's utility for fast EES, but few MOFs with simultaneous ionic and electronic conductivity have been reported.^{85,86} A possible strategy to obtain such dual-conducting MOFs would be to incorporate moieties known to support ion conduction (described above) into electronically conductive MOFs. However, it is critical to consider the negative effects of introducing redox inactive moieties on theoretical and experimental capacity (**Figure 3d**).

Particle-level considerations. The size and crystallinity of MOF particles can strongly affect the electrochemical response of MOF-based electrodes. For ionically and electronically conducting MOFs higher crystallinity is desirable. Larger MOF crystalline domains will result in higher intraparticle electronic conductivity. However, if larger crystalline domains have insufficient ionic conductivity, charge storage will be limited by solid-state ion diffusion. Thus, for MOFs with poor ionic conductivity smaller particle sizes can enable faster charging via increased accessibility to redox active sites by electrolyte ions. Meanwhile, for MOFs with high ionic (greater than that of the ionic conductivity of the electrolyte) and electronic conductivity (higher than that of conductive additives, e.g., carbon black), electrodes with smaller particles will display lower conductivity due to the increased role of interparticle transport, negatively affecting the overall charging rates.

IMPLICATIONS OF SYNTHESIS ON MOF PROPERTIES

The formation of extended coordination networks (MOFs) from ligand and transition metal occurs via their reversible ligand-metal coordination. Upon dissolution, both ligand and transition metal cation are solvated by solvent molecules and/or counterion. Therefore, MOF formation process in solution is strongly affected by solvent identity (and especially whether solvent(s) are protic or aprotic, or basic or acidic), reagent selection (i.e. counterion of the transition metal salt or ligand protonation) and concentration, the presence (or absence) of assisting agents such as modulators and co-solvents, and impurities (e.g., residual water). Each of these parameters as well as reaction temperature can additionally tune the rate of network formation and hence MOF crystallinity and particle size. If the rate of MOF formation is too high, the quality/crystallinity of the product can suffer.92 Hence, numerous strategies that enable the control of crystallization have been developed: (1) Ligands are typically introduced in protonated form; their deprotonation, required to realize the ligand-metal coordination that drives MOF assembly, is enabled by introduction of base. Examples include N,N-dimethylformamide (DMF) that serves both as a solvent and a base,93 organic amines (e.g., triethylamine)94, and inorganic base (e.g., aq. NH₃).95 The strength of the base and its rate of introduction can both impact the product characteristics: slower rates tend to yield higher-quality (e.g., more crystalline) MOFs. 96,97 (2) Various modulators can be used to suppress MOF nucleation and slow crystallite growth. Modulators are small molecules (e.g., ethylenediamine⁹⁸) or anionic species (e.g., benzoate⁹⁹) that compete with the ligand for transition metal cation coordination, thereby reducing the rate at which these cations can participate in MOF formation. Modulators have previously proven to be effective to enable the growth of single-crystals instead of micro- or nanocrystalline powders of network materials. 100,101 (3) Higher temperatures (ca. 60-120 °C) can assist in MOF crystallization, however it is essential to understand that increasing the reaction temperature can change the thermodynamically favorable phase, potentially yielding undesirable products. 102

Despite the apparent simplicity of solvothermal synthesis, the morphology, crystal structure, and even chemical composition can be extremely sensitive to synthesis conditions. For example, Dincă group showed that apart from parameters discussed above experimental variables such as exposure to air vs. inert environment and even the volume of the reaction flask influence the aspect ratio of the formed MOF.¹⁰³ Despite the importance of experimental parameters on the outcome of synthesis, there are few systematic studies exploring this subject.¹⁰⁴ As such, several precautions should be taken when attempting to reproduce a MOF synthesis from the literature or synthesize a new MOF. Changes in reaction conditions (e.g., solvent, temperature, time) can result in significant changes in network topology.¹⁰⁵ Moreover, defects can play an important role on MOF porosity, pore size, and access to potentially redox-active metal nodes.¹⁰⁶ Defects can be introduced (or avoided) by using modulators – typically HCl or capping ligands of coordination chemistry similar to that of the ligands used to form the MOF itself.^{106,107} Handling and treatment of MOFs post-synthesis is also important. For example, pore (and structure) collapse upon guest removal¹⁰⁸ or even solvent exchange¹⁰⁹ can occur. Hence, caution should be applied when drying MOFs prior to or after electrode fabrication, as this process may lead to structure changes such as intralayer displacement or structural collapse¹¹⁰; these changes may be detected by PXRD and gas sorption analysis.

Finally, when synthesis of new MOFs is pursued, several principles can be applied. Transition metal coordination and ligand geometries can dictate MOF structure. To this end, existing MOFs can be useful to inform how components with similar geometries might self-assemble. However, as discussed above, practical realization of such *a priori* prediction of MOF structure can be challenged by the complex solution chemistry that drives MOF phase selection (e.g., counter ions and solvent molecules can incorporate into the MOF structure, leading to the formation of unexpected MOF phases). If no relevant references are available, the following experimental parameters can serve as reasonable starting points: DMF as a solvent, temperatures of 25 and 100 °C (temperatures commonly used for successful MOF synthesis), the presence or absence of water (which may influence the crystallinity of the resulting MOF)⁹⁶, and the inclusion (or absence) of common modulators such as sodium acetate. ¹¹¹

CHARACTERIZATION OF MOFS

MOFs are crystalline materials, and diffraction methods therefore form the basis for their characterization (Figure 4). New MOFs for fast EES generally possess strong metal-ligand coordination bonds of low kinetic reversibility, making it especially difficult to obtain crystals of size and quality suitable for single crystal X-ray diffraction (Figure 4a). As such, structure solution from high-resolution, synchrotron-based X-ray diffraction of powder samples is often a necessary starting point to elucidate their structures. Structure solution from powder diffraction is rarely straightforward and available methods generally require at least a moderate level of expertise in both the powder diffraction theory and appropriate software (the interested reader is referred to a recent review on the subject and references therein).¹¹² TEM imaging and electron diffraction enables analysis of the smaller crystallites as well as can provide information about secondary or amorphous phases.¹¹³ However, due to the sensitive nature of MOF samples it is essential to ensure that sample doesn't undergo beam damage during data acquisition. To do so low-dose TEM approach is applied. 114 In cases where the structure of the MOF is known, powder X-ray diffraction is used to confirm the identity of synthesis products. This is generally a far more straightforward process, whereby standard laboratory X-ray diffractometers are sufficient for data collection, and data analysis typically involves a simple visual comparison of peak positions and intensities. Rietveld refinement provides a more quantitative and reliable approach to confirm phase identity, but is rarely carried out in practice. For example, Rietveld refinement can be used to quantify potential differences in stacking modes and intralayer stacking distances for 2D MOFs. 115,116 The pair distribution function obtained from diffuse scattering data can be used to describes the atom-atom distances of the sample and therefore capture the structure of any amorphous phase present.¹¹⁷ Amorphous regions typically lack the high porosity of the intended crystalline phase¹¹⁸ and complicate interpretation of electrochemical data (for example, determining to what extent internal redox active sites are electrochemically accessible). Finally, we stress that the crystallinity of the MOF should be assessed both before and after operation in an electrochemical device: MOFs may be sensitive to exposure to the conditions needed to operate fast EES devices, and observed device performance may be related to the MOF degradation products rather than to the MOF itself. Even post-synthesis treatments such as guest exchange, guest removal, and temperature changes can influence the MOF's structure¹¹⁸, so analytical data before and after such changes is necessary to ensure that the structure of the synthesized MOF is retained when used as an electrode material.

The crystallographic model of a MOF belies the complexity that is possible with experimental MOF samples. Synthesized MOFs can contain a wide variety of irregularities including surface defects, missing linkers or nodes, occluded guests (solvent, counterions, metal salts, etc.), stacking faults, non-uniform metal oxidation states and coordination environments, and differences in particle morphology. All these irregularities can dramatically impact the characteristics of the experimental sample. For example, accessible internal surface area and pore size can both be increased¹¹⁹ or decreased¹²⁰ by structural defects, and occluded guests and stacking faults usually prevent access by ions (with exceptions^{63,121}). We emphasize here that *the basic characterization of MOFs solely by X-ray diffraction is entirely insufficient to reveal these irregularities, and that these irregularities are fundamental to the performance of MOFs in fast EES devices*. We provide here several examples that describe the analytical techniques used to identify the important characteristics of experimentally obtained samples (**Figure 4b**).

We begin with gas sorption, which is typically used to measure the specific surface area¹²² and pore size distribution. The specific surface area derived from gas sorption measurements can under- or overestimate the electrochemically active surface area since ion intercalation kinetics are a complex function of pore characteristics (size, shape, chemistry), potential for intercalation⁶³, and solvent. Furthermore, gas sorption is best-suited for the characterization of micro- and mesopores, doing a poor job of capturing

macroporosity and >100 nm-scale surface roughness that can impact electrochemical behavior.¹²³ Pore size distributions are obtained from density functional theory (DFT) fits that are designed for traditional sorbents such as activated carbons and zeolites. Hence, pore size distributions derived from applying these models to experimental gas sorption data can potentially be quite inaccurate and provide little insight into whether the MOF's internal pores play a significant role in electrochemical behavior. A combination of gas sorption and electrochemically active surface area measurements (see below) is recommended to provide a more complete picture of the role porosity plays on the electrochemical behavior of the MOF.

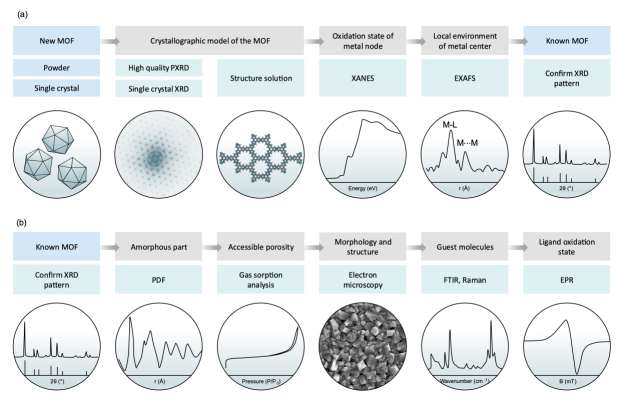


Figure 4. Characterization algorithm for new (a) and known (b) MOFs to assist with comprehensive characterization of their structure and properties.

Measurements with X-ray absorption near edge spectroscopy (XANES) and extended X-ray absorption fine structure (EXAFS) can be used to determine the metal oxidation states and coordination number, respectively¹²⁴, the latter potentially hinting at coordinated guests that are not part of the crystallographic model or stacking faults present between sheets in 2D MOFs. ¹²⁵ For example, EXAFS analysis of the conductive 2D MOF Ni₃(HITP)₂ was used to show, on the basis of Ni–Ni distances, that a slipped-parallel best characterizes interlayer stacking. ¹²⁶ The stacking mode is critical: staggered geometries can render potential pores inaccessible, preventing ion ingress and egress and thereby hindering electrochemical device performance. ¹¹³ Furthermore, these techniques can be used *in situ* during energy device cycling. For example, XANES was used to show that the metal oxidation state of Ni₃(hexaaminobenzene)₂ does *not* participate in redox processes during pseudocapacitive operation, indicating that changes in oxidation state are fully attributable to the non-innocent ligands within the material. ⁶² Once the structure of the MOF is known, EPR can be used to provide insights into the electronic states of the MOF. For non-innocent ligands, EPR can establish the presence of radicals (or lack thereof) and assist in the determination of their oxidation states. However, as it is very sensitive to the presence of unpaired electrons, it should be coupled with other techniques to definitively establish the electronic states present in the MOF. ¹²⁷

The morphology of synthesized MOFs is well-known to have a strong impact on the fabrication and behavior of MOF-based electrodes. Scanning electron microscopy (SEM) is generally the most straightforward way to determine MOF particle size and shape. The researcher must, however, obtain statistically relevant sample sizes and use image processing software to determine particle dimension and aspect ratio computationally to reduce user bias where possible. Care should also be taken when interpreting particle size data in that the observed particle size does not necessarily reflect the crystallite size.

The chemical purity of MOFs should be assessed by several analytical approaches. Since most MOFs are porous, the chemical contents of these compounds changes with the contents of the pores themselves. To

circumvent this complication, the MOF must be "activated" prior to chemical analysis: activation typically involves rinsing the MOF several times with polar and non-polar solvents, and then heating under reduced pressure for several hours to days. ¹⁰⁸ The MOF should then be subject to elemental (CHN, and other elements where applicable) analysis and inductively coupled plasma mass spectrometry (ICP-MS) or optical emission spectroscopy (ICP-OES) to determine the metal content. ¹²⁸ These techniques together show whether the expected chemical formula accurately describes the synthesized material. Nuclear magnetic resonance (NMR), infrared, and Raman spectroscopies can be used to detect occluded or trapped guests. Taking advantage of NMR for MOF chemical analysis typically requires digestion (dissociative dissolution) of the MOF in an appropriate deuterated solvent. The often modest solubility of the linkers used to form the MOF can pose a challenge. Hence, experimentation with digestion solutions to ensure complete dissolution of the MOF is often necessary. Good starting points for this experimentation can be strongly acidic solutions (e.g., trifluoroacetic acid, conc. HCl), strongly basic solutions (e.g., aqueous or DMF solutions of NaOH), and others. ¹²⁹

UNDERSTANDING OF CHARGE STORAGE MECHANISM AND DEGRADATION

It is essential to understand the charge storage and degradation mechanism of MOF electrodes. This knowledge can provide important guidelines towards improving the performance of existing systems as well as insight for the design of next-generation MOF-based high-performance electrodes. In **Figure 5** we depict a set of characterization guidelines that can assist in determining the dominant storage mechanisms. Below, we discuss the advantages and challenges associated with the use of each technique. As a first step, the contribution of the double-layer capacitance (F/g) can be estimated based on the accessible surface area⁶² using the following equation: $C_{dl} = C_A \cdot A$, where C_A is the areal capacitance (typically in the range between 5-20 μ F/cm², depending on the solvent and material; for an upper-bound estimate higher values of ~20 μ F/cm² can be used for conductive MOFs)⁶², and A is the specific surface area (m²/g) that can be obtained from gas sorption analysis.¹²² If the estimated C_{dl} value is much lower than the capacitance calculated from the electrochemical measurements it can serve as a clear indicator that other mechanisms (i.e. pseudocapacitance) are at play.^{11,32}

To gain insights into the contributions of the ligand and transition metal redox to charge storage, changes in their respective oxidation states have to be monitored during charge/discharge.⁶² For example, using in situ hard X-ray absorption spectroscopy (XAS) one can quantify the valence change of the transition metal by tracking changes in its edge energy. 130,131 Furthermore, analysis of the EXAFS region coupled with in situ XRD can provide important information about the reversibility of the changes in the transition metal coordination environment as a function of cycling and applied potential. 131,132 Insights into potential electrochemistry-induced degradation pathways can also be extracted from such measurements. 133 As discussed above, ligand redox can also significantly contribute to charge storage. However, quantitative assessment of its role in charge storage is not always easy. While in situ XAS can provide quantitative information regarding changes in the ligand's oxidation state, soft X-rays must be used to probe light atoms such as C, O, N and S.^{134,135} Due to the strong absorption of soft X-rays by most materials¹³⁰, the measurement set-up should be designed to minimize the loss of X-ray intensity due to absorption by the electrolyte or window material or even gas atmosphere.¹³⁰ Therefore, soft XAS has to be collected using specialized experimental configurations that feature electrochemical cells with SiN or SiC membrane windows at the working electrode that allow transmission of the soft X-rays^{130,136}. These configurations should also include gas-free paths for emitted X-rays to the detector. 130

Due to pronounced experimental challenges related to *in situ* XAS methodology, *ex situ* XAS (both hard and soft X-ray) and XPS measurements are used at different states of charge.^{62,137} However it is important to keep in mind the risks related to self-discharge during sample preparation that can result in underestimation of the metal/ligand redox contribution.¹³⁸ Therefore, in most cases other *in situ* techniques must be utilized in tandem to provide insights into ligand redox. As such, *in situ* Raman spectroscopy is another widely used method to analyze the chemical state of the organic ligand during charging.⁶² For instance, the evolution of the benzene-ring-related vibrations at different charge/discharge states were established by *in situ* Raman spectroscopy, revealing the reduction of the ligand within MOF structure.⁶² However, it is important to note that exact assignment of Raman peaks is not trivial and in many cases requires DFT calculations¹³⁹ or information from prior experiments with ligands that have been deuterated to assist with peak assignment.¹⁴⁰

In contrast, Fourier-transform infrared spectroscopy (FTIR) can offer direct information about the chemical state of the ligand at different states of charge. 141 For example, Cai et al. used triazine-based covalent

organic framework as a Li⁺ battery anode and performed *in situ* FTIR to investigate the redox mechanism of the electrode during Li⁺ (de)insertion. From the experiment, a stretching vibration peak corresponding to C=N functional group became stronger during lithium insertion and weaker during deinsertion. Strong -C-N-Li ionic bonds form during Li⁺ insertion process and the bonds successfully transformed back to C=N showing the reversibility of the redox process. However, such *in situ* FTIR studies must be performed using reflection or thin layer cell geometry to mitigate interfering spectroscopic signatures of the solvent/electrolyte. ^{141,142} For *ex situ* FTIR, the standard approach to preparing samples involving dilution in KBr pellets can easily disrupt the metastable charged state of MOF ligand (i.e., self-discharge) and therefore yield misleading results. Meanwhile, the use of pure MOF pellets can result in low-quality FTIR data.

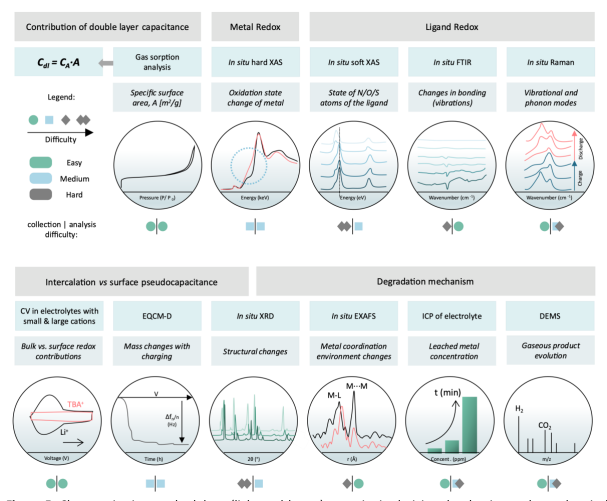


Figure 5. Characterization methodology (light teal boxes) to assist in deriving the dominant electrochemical charge storage and degradation mechanisms in MOFs (grey boxes).

Distinguishing the intercalation and surface pseudocapacitance mechanisms can be achieved by several methods. First and perhaps the most straightforward approach entails comparing MOF capacitance obtained in electrolytes containing alkylammonium cations larger than the MOF pores with that obtained in electrolytes with smaller cations such as Li⁺ or Na⁺.^{32,62} Capacitance values that do not differ significantly are indicative of surface, rather than intercalation, pseudocapacitance. As cation insertion can lead to changes in MOF lattice parameters, coupling such measurements with *in situ* XRD can yield further insights into structural changes (and their reversibility) that the MOF may experience during charging.¹⁴³ Another technique that can provide important mechanistic insight is electrochemical quartz crystal microbalance (EQCM) measurements that can track mass changes at the electrodes upon charging.¹⁴⁴ Quantitative information can be extracted about species entering the electrode material during charging by correlating electrode mass changes with the mass of the charge carrier ion. As such, EQCM can be used to reveal the charge carrier ion's solvation state upon entering the pores of the electrode.¹⁴⁵

Understanding MOF degradation pathways is the key to engineering MOF-based electrodes with long cycle life.²² The chemical instability of MOFs in certain environments (e.g., electrolytes) is typically caused by

competition between ligand, solvent, and/or anion for coordination to the metal center.¹⁴⁶ As of now, an understanding of the electrochemically-driven degradation of MOFs is lacking. However, presumably, much like for inorganic materials², the electrochemical instability of MOFs originates when the coordination between metal and ligand is compromised by metal or ligand redox processes ^{146,147}, leading to cumulative structural degradation of the MOF upon repetitive cycling. ^{147,148} The roles of solvent and salt selection can be critical to the electrochemical degradation rate, hence, optimal choice of solvent and salt can assist with stabilization of the structure. ^{149,150} *In situ* XAS and XRD can help to reveal changes in metal site coordination, irreversible structural changes of the electrode material, and formation of secondary phases. ^{135,151} MOF collapse, that also results in transition metal leaching from the electrode, can be tracked *via* time dependent analysis of transition metal concentration in electrolyte using inductively coupled plasma (ICP) analysis. ¹⁵² As previously noted, electrolyte decomposition can also result in poor cycling stability. Differential electrochemical mass spectrometry (DEMS) can be used to track the formation of gaseous products (e.g. H₂, CO, CO₂ and O₂) that are associated with electrolyte decomposition. ^{153,154}

OUTLOOK

In this review, we outlined MOF design principles for fast EES and highlighted the importance of gaining insights into MOF charge storage mechanisms. Significant progress has been made in recent years in the area of MOF-based EES devices, especially with the introduction of electrically conductive MOFs. However, we see numerous directions in need of concerted efforts from both MOF chemists and electrochemists to further advance this field and render MOF-based EES devices competitive with or better than traditional inorganic-based energy storage electrodes.

First, thus far, most of the promising examples of MOF-based energy storage are of MOF-based negative electrodes (anodes). Meanwhile, the field of MOF-based cathodes remains in its infancy primarily due to the lack of conductive MOFs that are reversibly redox-active at higher potentials. Hence, new MOFs are needed that simultaneously incorporate redox moieties active in the potential range > 3 V vs. Li/Li⁺ and can also deliver high capacities.

Second, to date, most conductive MOFs display ligand-based redox, while transition metal cations have not directly contributed to charge storage during EES device cycling. Hence, to yield further improvements in achievable pseudocapacitance, it is necessary to redox-match transition metals with their partner ligands and may require exploration of MOF architectures that have received little attention for fast EES to date (for example, 3D conductive MOFs). The study of small molecule coordination complexes (as discussed above) can assist in selecting metal-ligand motifs that support redox-matching in analogous MOFs. DFT simulations can further accelerate the discovery of promising motifs by rapid screening¹⁵⁵ for robust redox-matching.

Third, past work examining the behavior of carbide-derived carbons with uniform pore size has shown that double-layer capacitance is maximized when the pore size matches the size of the infiltrating organic ion. ¹⁵⁶ We foresee that MOFs can also benefit from enhanced EDL capacitance if MOF pore size is matched to the size of the solvated ions. This can be achieved by systematically changing the MOF's ligand length without changing the ligand geometry (see above). ⁵⁸

Finally, as cycle life is critical for fast EES, it is important to achieve MOF electrode cycling stability over at least several thousand cycles. To this end, the mechanisms by which MOFs electrochemically degrade, and the role of electrolyte on MOF degradation, must be better understood.

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