Degradation Mechanisms and Mitigation Strategies of Nickel-Rich NMC-Based Lithium-Ion Batteries

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Abstract
The demand for lithium-ion batteries (LIBs) with high mass-specific capacities, high rate capabilities and long-term cyclabilities is driving the research and development of LIBs with nickel-rich NMC \((\text{LiNi}_x\text{Mn}_y\text{Co}_{1-x-y}\text{O}_2, x \geq 0.5)\) cathodes and graphite \((\text{Li}_x\text{C}_6)\) anodes. Based on this, this review will summarize recently reported and widely recognized studies of the degradation mechanisms of Ni-rich NMC cathodes and graphite anodes. And with a broad collection of proposed mechanisms on both atomic and micrometer scales, this review can supplement previous degradation studies of Ni-rich NMC batteries. In addition, this review will categorize advanced mitigation strategies for both electrodes based on different modifications in which Ni-rich NMC cathode improvement strategies involve dopants, gradient layers, surface coatings, carbon matrices and advanced synthesis methods, whereas graphite anode improvement strategies involve surface coatings, charge/discharge protocols and electrolyte volume estimations. Electrolyte components that can facilitate the stabilization of anodic solid electrolyte interfaces are also reviewed, and trade-offs between modification techniques as well as controversies are discussed for a deeper understanding of the mitigation strategies of Ni-rich NMC/graphite LIBs. Furthermore, this review will present various physical and electrochemical diagnostic tools that are vital in the elucidation of degradation mechanisms during operation to supplement future degradation studies. Finally, this review will summarize current research focuses and propose future research directions.

Keywords Lithium-ion battery · Ni-rich NMC · Degradation · Mitigation · Graphite · Diagnostic tools

1 Introduction
Due to significant advantages such as high energy densities, high galvanic potentials, wide temperature ranges, no memory effects and long service lifespans, lithium-ion batteries (LIBs) have been widely employed in various applications, including in the fields of communication, aviation and transportation. Overall, the chemistries of LIBs are constantly evolving, and in terms of cathode materials, LiNi\(_x\)Mn\(_y\)Co\(_{1-x-y}\)O\(_2\) (NMC) layered oxides have received increasing attention due to their enhanced specific capacity and thermal stability [1]. As compared with \(\text{LiMO}_2\) (M = Ni, Mn, or Co) cathodes, NMC-based cathodes also possess the combined merits of three transition metals in which nickel can offer high specific capacities, whereas Co and Mn can provide layered structures and enhanced structural integrity. In addition, NMC-based cathodes can outperform lithium iron phosphate (LFP) cathodes in many areas, particularly in terms of operational voltage in which LFP-based LIBs can only output voltages below 3.4 V [2] and suffer from high rates of self-discharge. Moreover, materials such as Li\(_2\)MnO\(_3\) and lithium titanate do not possess great industrial potential as cathodes due to short lifespans and high costs. Here, NMC-based cathode materials are much cheaper to produce (~ USD 23 per kg for NMC111) as suggested by researchers using a co-precipitation method [3], and the increase in NMC-based material prices from January 2017 to March 2018 (~ 43% for NMC532 and NMC622, and ~ 27% for NMC811) [4] indicates that more research is
being directed to the field of NMC-based cathode materials due to their commercial potential.

Following the initial commercial success of LiNi0.5Mn0.3Co0.2O2 (NMC333 or NMC111), NMC-based LIBs have become the mainstream with gradual improvements in NMC technology through the steady increase in the nickel content in each generation of cathode materials. And because Ni can cycle between Ni2+/3+ and Ni3+/4+ redox couples without large variations in voltage during the charge/discharge process, layer-structured Ni-rich NMC (LiNiMnCox−yO2, x ≥ 0.5) cathodes [e.g., LiNi0.5Mn0.3Co0.2O2 (NMC532), LiNi0.6Mn0.2Co0.2O2 (NMC622) and the most recent LiNi0.8Mn0.1Co0.1O2 (NMC811)] have been developed. Here, the main advantage of Ni-rich materials is their high discharge capacity (200–220 mAh g−1), which represents a large increase in energy density (~800 Wh kg−1) as compared with conventional LiCoO2 (~570 Wh kg−1) and LiMn2O4 spinel (~440 Wh kg−1) materials [5]. In addition, increasing the Ni content is the most effective method to enhance capacity in the current state-of-the-art LIBs without requiring advanced and novel battery chemistries. As a result, researchers are reviewing recent progress achieved in the development of Ni-rich cathode materials in an attempt to provide important clues for the future design of Ni-rich cathodes [6].

However, although increasing Ni percentages can allow for enhanced Ni-rich NMC-based battery performances as compared with other NMC combinations in terms of capacity, capacity fading becomes severely pronounced. To address this, significant efforts have been made to elucidate the mechanisms of capacity fading and associated growth of impedance [7, 8], as well as to find remedies [9–12]. And in recent decades, numerous studies and reviews have been published worldwide [13–15] in which various testing protocols [16], research methodologies, models [17–19] and physical characterizations [9, 20] have been used to interpret the behavior of NMC-based cathodes with or without modifications. Here, studies based on microscopy [21], crystallography [22], mass spectroscopy [23], gas measurements [24] and calorimetry [8] have revealed that parasitic reactions [7], cation mixing (leading to restructured surface regions) [25], active material dissolution [26] and oxygen release [23, 25] are primary factors responsible for cathode degradation at the atomic scale. In addition, scanning electron microscopy (SEM) [13] together with computational studies [27] has revealed that intergranular cracking can weaken connections between primary particles. Despite these findings, the majority of recently published studies are insufficient and only partially cover the processes through which degradation occurs. For example, Liu et al. [28] reviewed surface layers and surface chemistries during Ni-rich NMC-based LIB operations and presented cationic doping and metal oxide coating as recent research progress, whereas Myung et al. [29] in their review only provided a brief developmental history of NMC and NCA-based LIBs with a focus on the categorization of gradient layers and core–shell structures as remedies. A more recent and detailed review conducted by Hou et al. [30] covered more mitigation topics varying from ion substitution and electrolyte additives to gradient layers; however, to the best of our knowledge, no review has presented a comprehensive collection of mitigation methods or has categorized them based on characteristics. Similarly, controversies in mitigation strategies and trade-offs need to be considered, such as the selection of dopants or coating materials, all of which have yet to be discussed. Based on this, these vacancies in the literature will be covered in this review to assist in the further investigation of Ni-rich NMC-based LIBs.

Apart from the cathode, graphite—the anode material that has received the most attention in LIBs due to its high mass-specific capacity and economic advantages [26, 31, 32]—can function collaboratively with NMC-based cathodes to provide promising performances in coin cells [20, 33], pouch cells [9, 34], pouch bags [11] and T-cells [35]. Based on this, this review will also provide a complete picture of the aging of graphite anodes that can affect both electrodes in battery cells based on degradation mechanisms and effective mitigation strategies. Furthermore, the solid electrolyte interface (SEI) is a film that forms on graphite anode surfaces during initial charging [36–39] and thickens in subsequent cycles, which can protect and impede anodes and is vital in the discussion of the aging behaviors of graphite anodes. Therefore, detailed mechanisms of anode SEIs are also reviewed to supplement previously published reviews.

To further investigate the degradation mechanisms of LIBs, multiple physical methods have been used in both electrochemical tests and physical characterizations [e.g., electrochemical impedance spectroscopy (EIS) to characterize the resistance of electron transfer and surface films]. Conventional and modern physical characterization methods have also been utilized to reveal surface morphology, element compositions, crystal lattice dimensions and reaction processes, and are also important to the interpretation of the degradation of Ni-rich NMC/graphite LIBs. Based on this, these diagnostic tools will also be presented in this review.

To fully present the evolution of electrode performances in Ni-rich NMC/graphite LIBs, degradation mechanisms based on various experiments and recent mitigation strategies in which cell conditions were evaluated separately or as a whole are presented in this review. And overall, this review will provide a summary of the recent advances in Ni-rich NMC-based batteries and can supplement pioneering reviews published previously.
2 Degradation Mechanisms

Ni-rich NMC-based materials are some of the most promising cathode candidates for next-generation LIBs due to high capacities and large voltage windows. Despite impressive progress in the development of advanced materials and fabrication processes, the development of high-capacity electrodes with long-term stability and prolonged cycle lifespans remains an important challenge. To address this, the understanding of underlying degradation mechanisms presented in this review can serve as a starting point to learn and improve the performance of Ni-rich NMC-based cathodes. In addition, the information included in this review can provide valuable insights into the creation of new devices with improved efficiency and productivity. And for the sake of completeness in the understanding of degradation mechanisms in LIB full cells, parasitic reactions on NMC-based cathode surfaces, the aging of graphite anodes as well as SEI formation and aging are also briefly presented in this section.

2.1 Ni-Rich NMC-Based Cathodes

The degradation of Ni-rich NMC LIBs involves different types of capacity loss, including initial capacity loss (ICL), sudden capacity loss (SCL) and gradual capacity loss (GCL), all of which are accompanied by impedance formation and aging are also briefly presented in this section.

2.1.1 Surface Degradation During Cell Operation

**Lattice Expansion/Contraction** NMC oxide possesses an $R\bar{3}m$ structure (rhombohedral symmetry) with a Li-layer on the $3a$ site, an NMC layer on the $3b$ site and an oxygen layer on the $6c$ site [7] and can usually be indicated in the splitting of (110) and (108) peaks and of (006) and (102) peaks. For example, the expansion and contraction of the $c$-axis (14.21 Å) of NMC811 lattices can be observed by using in situ XRD measurements in the cell voltage windows of 3.0–4.0 V and 4.0–4.4 V, in which (003) peaks can decrease from 18.96° (3.0 V) and subsequently rise to 19.09° (4.4 V) [7, 10, 43]. This can also be observed for the movement of (104), (015) and (108) peaks. Here, researchers suggest that the expansion of the $c$-axis is due to the repulsive force generated from MO$_6$ slabs, which are positively charged in a highly delithiated state, whereas the loss of electrons can shrink the radius of transition metal ions to reduce the $a$-axis [44]. In addition, the (110) peak can shift to higher angles in experiments, suggesting that the $a$-axis can contract during charging due to the smaller radius of transition metal ions at elevated valences [45]. Researchers have also reported that the $a$-axis of NMC811 lattices remains constant at potentials exceeding 4.3 V [10] and that a steep drop of the $c$-axis occurs at potentials exceeding 4.2 V in which the drop becomes steeper as the Ni content increases [7, 20, 43]. Here, researchers proposed that this drop is due to the fact that the repulsion between O$^{2-}$ layers decreases with more covalent M–O bonding at higher delithiated states [46, 47] (Fig. 1a, b). Furthermore, studies have also confirmed two pathways (Fig. 1c2, c3) involved in the delithiation of Ni-rich NMC-based cathodes [40, 48], including oxygen dumbbell hopping and tetrahedral site hopping, in which at the beginning of charging, Li can exit through oxygen dumbbell hopping until a certain degree of delithiation is reached and continue through tetrahedral site hopping due to increased energy barriers as caused by Li–O bonds during delithiation.

**Surface Reconstruction to Fm3m Rocksalt** Surface reconstruction of the particles of NMC-based cathodes can initiate if delithiation exceeds 70% as shown in NMC111 [49]. In addition, increasing the Ni content can lead to increased portions of Ni$^{3+}$ (usually Ni ions with higher valences) in the total amount of Ni. Furthermore, elemental Ni partially located in the transition metal layer as Ni$^{2+}$ (0.69 Å) possesses a similar radius to Li$^+$ (0.76 Å) [50], and Li$^+$ and Ni$^{2+}$ can exchange positions in a delithiated state through Ni$^{2+}$ diffusion to the octahedral sites of Li$^+$ (energetically favorable) through a neighboring tetrahedral site (Fig. 1c) [42, 51]. Here, the degree of Ni$^{2+}$ migration can be evaluated by calculating the $c/a$ ratio of the NMC lattice because this migration reduces the $c/a$ ratio. And based on XRD
measurements, ratios of $I_{003}$ to $I_{104}$ can decrease to less than 1.2 after significant cation mixing and can therefore serve as an indicator of the degree of cation mixing [52]. Moreover, researchers reported that cation disorder can lower spacing between atomic layers in the lattice of the NMC-based cathodes, hinder Li$^+$ movement and reduce the amount of active Ni and Li [53]. In the case of higher Ni percentage cathodes, Mn$^{4+}$ can partially be replaced by Ni, which increases the valence of Ni and therefore lowers the possibility of Ni atoms migrating to 3a sites [8]. However, because the total amount of Ni also increases, Ni$^{2+}$ migration still occurs to a larger extent. To minimize Ni$^{2+}$ migration, researchers reported that mitigation methods can be used to modify cathodes [54]. Furthermore, a disordered spinel structure ($Fd_{3}m$) due to non-ideal cation mixing [55] can form if LIBs are charged to high voltages (e.g., 4.8 V) [21, 56–60] and consists of a Li$_2$M$_2$O$_4$ spinel structure and a M$_3$O$_4$ spinel structure if Co migration into tetrahedral sites occurs [42] (Fig. 2a). Xiong et al. [61] and Eom et al. [62] also reported that spinel phases can be formed on charged cathodes stored at 90 °C for a week. Moreover, researchers found that migrated Mn and Co possess high valences and smaller radii for movement and more vacancies in the Li-layer in a highly delithiated state [63]. Researchers also reported that long-term cycling to 4.2 V (NMC 622) [64] can result in phase changes to an $Fm\bar{3}m$ structure consisting of Ni$^{2+}$, Mn$^{2+}$ and Co$^{2+}$ with lowered valence states as compared with a spinel structure [63, 65] and ion-insulating [56, 63, 66] (a cubic rocksalt phase) (Fig. 2b). Here, the conversion from a layered structure to a cubic rocksalt phase (usually confirmed by HRTEM/EELS [9]) can release O$_2$ into the cell, which can react with electrolytes to generate CO$_2$ and result in increased electrode interfacial resistances due to kinetic barriers in the Li$^+$ insertion/desertion process [50].

To address this, researchers reported that NMC811 cathodes pretreated with ramping to 4.5 V limited the growth of the rocksalt layer in subsequent cycles and can serve as a pillar structure to stabilize the lattice and isolate the surface of NMC particles [67]. Furthermore, researchers reported that the oxidation states of Mn and Co decrease in the surface layer/region of NMC811 and NMC442 after cycling [9, 63].

**Phase Transformations** Two hexagonal phases of NMC811 can coexist on the surface of particles if freshly prepared in which the evolution of one phase is irreversible and vanishes after initial charging (with lattice constants remaining at the initial value), whereas the second phase is reversible (except for a small difference between charging and discharging above 4.2 V). Here, the fading of the (110) peak at 65.12° can track the conversion of the region in which the two hexagonal phases coexist and are converted to a single phase [7]. In addition, the broadening of the (003) peak during cycling as recorded by using in situ XRD measurements is of great interest in which researchers [25] suggested that this phenomenon was due to the creation of a second phase in the potential range of 3.8–4.0 V (H2, evolved from the monoclinic phase in contrast to the phase H1 following the stoichiometric composition) that possesses less Li. Researchers also reported that this process occurred earlier with higher Ni portions (Fig. 2c) and that at higher voltages, the conversion from H2 to H3 occurs with impedance growth (volume contraction) only for NMC811 (or NMCs

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**Fig. 1** The c-axis (a1), the a-axis (a2) and potential (a3) of an NMC 811 cell as a function of specific capacity. The c-axis (b1) and the a-axis (b2) as a function of cell potential during the second cycle and dQ/dV of the second cycle as a function of cell potential (b3) (Reprinted with permission from Ref. [7]. Copyright © The Author(s) 2015. Published by ECS). **c1** Lattice of an NMC layered structure. **c2** Tetrahedral site pathway and **c3** oxygen dumbbell pathway for Li-ion diffusion in an NMC layered structure (Reprinted with permission from Ref. [48]. Copyright © 2015 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim)
with Ni portions larger than 80%) [25]. Researchers have also found that the phase transition from H2 to H3 can lead to (003) peaks shifting to higher degrees [68] and that Li-reordering (Li diffusing into Ni4+-rich sites) can further convert H3 to H3-2 [69] in which the fast shrinkage of the c-axis at potentials above 4.15 V is a clear indication of this phase transition. Moreover, the generated microregion of NiO2 in the intralayer of the NMC811 lattice as a result of the phase transition from H2 to H3 can lead to the irreversible transformation of the NMC811 structure [45] and this phase transition becomes severer as Ni portions further increase to extreme numbers as indicated by rising peaks of the differential capacity curve centered at 4.15 V [68]. Here, the fading of this peak leads to capacity fade and is referred to as the “deleterious effect.” Furthermore, phase changes to a second hexagonal phase for other NMC-based cathodes have yet to be clarified. (Fig. 2d illustrates crystal lattice evolution during delithiation.)

**Surface Stabilizer** In general, NMC-based cathodes with more Mn content tend to possess better cycling stability [13, 70], whereas NMC-based cathodes with more Ni content are believed to be incapable of reaching high cutoff voltages due to the lack of Mn4+ as a structure stabilizer [65]. Apart from Ni2+, structural instability also applies to Mn3+, although with the high Ni content, Mn migration is not considered to be significant. Here, due to low octahedral site stability energy, NMC-based cathodes possess a tendency to form spinel-structured LiMn2O4 on surfaces [71] and Mn4+ possesses a similar tendency to migrate to the Li-layer [33, 71] and decrease capacity. Nevertheless, diffused Mn4+ on particle surfaces can stabilize structures during long-term cycling [50].

### 2.1.2 Impurities and Parasitic Reactions

**Surface Impurities** To maintain a layered structure, excessive Li is used for Ni-rich NMC-based cathodes. However, side reactions of Li with water vapor and CO2 in air can lead to LiOH and Li2CO3 in which exposure to air can result in a reconstructed surface layer 3 nm thick (evidenced by STEM) that thickens if cycled [9]. A surface layer with similar impacts can also form as cathodes come into contact with electrolytes [65]. Here, researchers suggest that this reconstructed layer can be tuned through different synthesis methods of the cathode powder [63] and that in the case of NMC111, surface impurities can also grow on top of cathode particles as resistive films over the same time period for uncycled Ni-rich NMC-based LIBs [35] that contain mainly insulating hydroxides and carbonates as revealed through Raman spectroscopy (Fig. 3a). Here, the circumstances in which Li2CO3 (Eq. 1) or NiCO3 is favored remain unclear [35, 72, 73]. Researchers also reported that high humidity can lead to more hydroxide (e.g., LiOH) on NMC622 [72] and that surface impurities can affect cell impedances in subsequent cycles through reactions with electrolytes to form a Li+ diffusion inhibiting layer.

\[
\text{Li(Ni, Mn, Co)O}_2 + \frac{x}{2}\text{CO}_2 + \frac{x}{4}\text{O}_2 \\
\rightarrow \text{Li}_{1-x}(\text{Ni, Mn, Co})_2\text{O}_2 + \frac{x}{2}\text{Li}_2\text{CO}_3
\]

**NMC Dissolution** The dissolution of NMC materials can also result in capacity attenuation [26, 74] because it can decrease Li+ insertion sites and is triggered by the products of electrolyte decomposition such as HF from the reaction of PF6− with H2O (Eqs. 2, 3) [31]. In addition, higher voltages can accelerate the dissolution of NMC111 due to the release of more acidic components as a result of rapid electrolyte decomposition [75]. Despite these findings, effects on Ni-rich NMC have yet to be reported. The dissolution of NMC materials can also cause a concurrent issue involving the production of resistive MF, as a side product on the surface of NMC particles. Moreover, the dissolution of transition metals can pose a threat to anodes through electrodeposition [32, 76], catalysis of solvent reduction [77] and formation of inorganic layers in SEIs [78], all of which can impede Li+ intercalation and reduce capacity [79]. Here, Mn4+ disproportion into Mn2+ and Mn3+ was proposed to explain Mn dissolution [26, 75] as previously reported in Li–Mn–O spinel structures [80, 81] and various electrolytes have been utilized to form protective surface films containing carbon on electrodes to isolate surfaces from electrolytes [33].

\[
\text{LiPF}_6 \rightarrow \text{LiF} + \text{PF}_5
\]

(2)

\[
\text{LiPF}_6 + 2e^- + 2\text{Li}^+ \rightarrow \text{LiF} + \text{Li}_4\text{PF}_6
\]

(3)

**Self-redox Reaction** At highly delithiated states, the valence of transition metals increases, and because the low-spin Co3+/4+ band overlaps with the 2p band of O2− (Fig. 3b), electron transfer from O2− to Co3+/4+ can occur and reduce oxidized transition metal ions, thus releasing O2 into battery cells [82]. Oxygen release based on this also occurs at the first charging period in which O2− reduces transition metals oxidized to 4+ (similar effects on Ni4+ also exist and dominate oxygen release in Ni-rich NMC-based cathodes) [14, 42].

**Parasitic Reaction: Effects of Ni4+** Highly delithiated (highly charged) states tend to generate large amounts of Ni4+ that can react with electrolytes. (No reaction schemes have been proposed.) This side reaction can significantly thicken cathode–electrolyte interfaces (CEIs) and reduce the number
Parasitic Reaction: CO₂ generation Large Ni⁴⁺ portions at higher voltages can cause increased CO₂ evolution at increased upper cutoff voltages [85] in which cathodes with higher Ni content suffer more from parasitic reactions with Ni⁴⁺ that are a source of emitted CO₂. In addition, because highly reactive oxygen can be produced through O₂⁻ from self-redox reactions, surface reconstructions and highly delithiated cathodes (ca. >80%) [86], CO and CO₂ can also be traced back to the reaction between O₂ and alkyl carbonate electrolytes (Eq. 4) [24, 25]. Alternatively, sweeping to higher voltages can also cause the electrochemical decomposition of electrolytes to produce CO₂ (Eq. 5, Fig. 3c) [23, 24, 87]. Furthermore, Gasteiger et al. [88] have experimentally shown three other mechanisms that may contribute to CO₂ emission, including solvent hydrolysis (if trace amounts of OH⁻ exist), electrolyte impurity oxidation and HF reacting with lithium carbonate as a surface impurity. Moreover, the decomposition of lithium carbonate has also proven to be a source of CO₂ [89, 90]. Researchers have also suggested that...
NMC-based cathodes with Ni-rich surfaces behave differently than other cathodes with different portions of Ni [7]. For example, the degradation of NMC811 cells was studied by using different electrolyte additives including VC and PES211 to suppress side reactions and it was revealed that NMC442/graphite cells and NMC111/graphite cells showed better performances with PES211, whereas NMC811/graphite cells aged slower with VC (with slower capacity fading and impedance growth) [79]. Here, PES was also reported to be a viable additive for the suppression of gas evolution [91] and impedance growth during cycling to higher voltages (above 4.3 V), and rocksalt surface layers formed with the use of VC but not with PES211, concluding that electrolyte additives can significantly affect the rate of parasitic reactions and that parasitic reactions were the main reason for capacity fading [7, 92].

\[
\text{EC} + 2\text{O}_2 \rightarrow 2\text{CO}_2 + \text{CO} + 2\text{H}_2\text{O} \quad (4)
\]

\[
\text{EC} \rightarrow \text{CO/CO}_2 + \text{R-H}^+ + e^- \quad (5)
\]

### 2.1.3 Other Issues

#### Initial Capacity Loss

Initial capacity loss is strongly linked to the loss of available Li and the growth of impedance in anodes due to SEI formation. Here, researchers have reported that NMC-based cathodes cannot return to their fully lithiated states after initial cycling [26] in which separate mechanisms have been proposed, including parasitic reactions consuming Li\(^+\) [93] and the sluggishness of Li\(^+\) diffusion into the few vacancies of the Li-layer [94] (circumstantially proved by Gasteiger et al. [26]). In addition, researchers have also suggested that initial capacity loss is related to the C-rate because it is kinetically influenced. Furthermore, partial losses of available cathodic Li occur to compensate for losses of anodic Li\(^+\) to SEIs [26], and other losses of Li in the cycling period can be attributed to parasitic reactions including the immobilization of Li in the SEI and enlarged polarization, which become significant at higher cutoff potentials [26, 95].
**Cracking of Secondary Particles** Reduced performances due to the cracking of spherical secondary particles have been extensively reported and are of significant concern [96]. Here, the expansion and contraction of the c-axis during repeated cycling at above 4.2 V can lead to microstrains on particles in cathodes, leading to the generation of microcracks in the core of primary particles with some initially generated cracks being able to close in subsequent cycles [68]. These cracks can cause poor connection between particles and micropores and even the cracking of secondary particles [13], all of which contribute to degradation at the microscale. To address this, Lim et al. [27] have used first-principle calculations to study these cracks generated by gaps between primary particles and found that they were a result of anisotropic and average contractions. In addition, researchers reported that these cracked particles possessed enlarged surface areas and therefore increased the possibility of parasitic reactions [13]. Moreover, these microcracks were also found to partially originate from the gas evolution of NMC particles [13] (Fig. 3d), and recent studies on coated NMC (76 14 10) suggested that the infiltration of liquid electrolytes into the gap between primary particles can also cause cracking [97]. Alternatively, various NMCs have also been found with no cracking by using SEM [7, 20]. Here, Dahn et al. [20] suggested that the shell of core–shell-structured NMC cathodes (a Mn-rich shell and a Ni-rich core) was resistant to cracking due to the absence of compressive stress on the core and tensile stress on the shell. Based on all of this, numerous mitigation methods including carbon matrices and gradient layers have been proposed to resolve cracking at the microscale (further details in Sect. 4.1).

**The Effects of Temperature** Studies have shown that cell degradation can accelerate at elevated temperatures above 30 °C and that at low C-rate cycling, cell degradation is mainly affected by the time in which cells are placed under high temperatures and not by cycling [98]. Long-term operation of LIBs can produce large amounts of heat. And although this released heat may even be useful in cold weather conditions, heat emissions are a major issue of Ni-rich NMC-based LIBs due to thermal instability [33] in which thermal runaway can occur if NMC-based cathodes react with LiPF6 from the electrolyte and limit the commercial and practical use of these battery packs. To improve thermal stability, coatings such as SiO2 [62], TiO2 [99] and Li2ZrO3 [100] have been chosen to prevent contact between cathodes and electrolytes, and an ALD-coated NMC532 with Al2O3 has even been reported [101] to resolve the issue of uneven coatings and prevent the failure of cathode separation from electrolytes.

### 2.2 Graphite Anode

Similar to the cracking of NMC secondary particles, cracking can also occur in graphite anodes due to Li intercalation/deintercalation, which can expand and contract the distance between graphite layers [102]. In addition, recent research indicated that unit cell expansion (13.2% in total) during Li intercalation is neither continuous nor linear, but staged (Fig. 4a–e) [103]. Here, Li intercalation from C6 (Li-free graphite) to LiC24 (Stage 2L) showed no in-plane ordering of intercalated Li, whereas from 2L to LiC12 (Stage 2), the ordering of Li appeared and the lattice can be assigned to the space group P63/mmm. Furthermore, in the transition from Stage 2 to LiC6 (Stage 1), total volume change suddenly increases as the amount of x exceeds 0.6 in LiC6 (where Stage 1 appears as Li presents in every interlayer). These results showed that optimal composition was between 20 and 80% charged in which Stage 2 stably exists or extends into.

Furthermore, “dead lithium,” which is Li-ions electrochemically plated onto the surface of graphite anodes, is the principal reason for capacity fading, especially at final stages [104]. This phenomenon occurs on graphite surfaces that possess limited mass transport, slow diffusion within the graphite or low charge transfer speeds [105] and is strongly promoted by low temperatures/large polarizations (including high electrolyte resistances induced by high temperatures) during lithiation as well as the properties of the graphite electrode (e.g., morphology [102], loading). And as Li+ intercalation is replaced by Li plating, issues such as dendrite growth become more pronounced, leading to increased impedance and more importantly, potential safety hazards due to internal shorting. To avoid this, the limits of charging currents and the state of charge need to be established to maintain polarization in optimal ranges [106]. However, large polarizations also need to be adopted to minimize charging durations to meet industrial application demands. Here, researchers reported that the reduction of graphite thicknesses and improvements in porosity can decrease charge transfer impedances [105, 107].

### 2.3 SEI

Electrolytes used in LIBs contain two main components including Li salt and organic solvents. Here, the use of additives as film-forming agents can allow for stable SEIs and, in general, the concentration of Li salt is maintained at 0.8–1.3 M to optimize electrolyte viscosity and ion conductivity. As compared with cathodes, the potential of LIB anodes is low and can trigger electrolyte reduction (SEI formation), which can cause Li loss during initial charging (Fig. 4f) [31]. However, because SEIs are electronically insulating, further reductions of electrolytes do not occur
in which formed SEIs are usually 10–50 nm thick and are conductive to Li$^+$ ions. Researchers have extensively studied the constituents of passivating SEIs on graphite anodes and have reported that lithium ethylene dicarbonate (LEDc, Eq. 6, Fig. 5a–d) [108] was the main component of SEIs formed with ethylene carbonate (EC) solvent [109] apart from electronically insulating LiF and gaseous products (H$_2$ and C$_2$H$_4$). Propylene carbonate (PC) as a solvent has also been well investigated due to its thermal stability and wide voltage windows. Furthermore, DFT calculations have shown that stable SEI films can be produced by oligomers of SEI film components (SFCs) [108], which are first created through solvent decomposition. LiF is also a product of almost all common Li salts such as LiPF$_6$ (the only commercially available salt), LiBF$_4$, LiTFSi, LiTSi, etc., in which researchers reported that the amount of LiF produced from LiBF$_4$ was large and would result in the formation of a grainy film instead of a smooth film.

$$2\text{EC} + 2\text{Li}^+ + 2\text{e}^- \rightarrow \text{Li}_2\text{EDC} + \text{C}_2\text{H}_4 \quad (6)$$

As aging continues during cycling, researchers believe that the inorganic portions of SEIs also increase as LiF (can be used as an indicator of SEI maturation) is produced through LiPF$_6$ reacting with lithium carbonates (Eqs. 7, 8) [37].

$$2\text{RCO}_3\text{Li} + \text{LiPF}_6 \rightarrow 2\text{LiF} + 2\text{RF} + \text{LiPO}_2\text{F}_2 + 2\text{CO}_2 \quad (7)$$

$$\text{RCO}_3\text{Li} + \text{LiPF}_6 \rightarrow 2\text{LiF} + \text{RF} + \text{POF}_3 + \text{CO}_2 \quad (8)$$

After SEI formation, the resistance of anodes/SEIs toward divalent transition metal ions such as Ni$^{2+}$, Co$^{2+}$, Cu$^{2+}$ and Mn$^{2+}$ (relatively more impactive) is significantly enhanced, preventing the reduction of EC solvent by these ions. However, formed SEIs remain targets to many reductive substances in which reductive decomposition can degrade SEIs (Li$_2$CO$_3$ and C$_2$H$_4$ are produced as a result) and expose thermodynamically unstable graphite anodes to reactive environments [38]. Here, if the exposure of graphite anodes occurs, a series of damage can occur on the graphite anode, including the intercalation of solvated Li$^+$, which leads to larger sizes and the potential risk of the breaking of the graphite layer. In addition, Mn$^{2+}$ has been reported to be able to penetrate SEIs and reside in the inner area between the SEI and the anode through the exchange of positions with Li$^+$ ions located in the SEI [110]. Furthermore, the recurrent generation of C$_2$H$_4$ suggests that Mn$^{2+}$ is a destructive substance to both solvents and graphite anodes in which accumulated Mn$^{2+}$ near graphite can function as a catalyst, suggesting that it is reduced at the surface of the anode but can reduce EC afterward, leading to thickened SEI films and the immobilization of active Li (Fig. 5e) [110]. In addition, the reduction of Mn$^{2+}$ can cause the gradual transfer of active Li$^+$ to SEIs or the conversion of active Li$^+$ into compounds deposited onto graphite anodes, both of which lower storage capacity [38]. Researchers have also proposed another mechanism of Li$^+$ that suggests that even monolayer Mn$^{2+}$ can impede the movement of Li$^+$ through the graphite and the SEI [111]. As for PF$_5$, it is produced from the decomposition of various electrolytes and can trigger the open-ring polymerization of EC, which further increases SEI decomposition along with the production of HF [112]. Moreover, the damage/repair process caused by gaseous PF$_5$ consumes active Li-ions until Li-ions are depleted in the system [113]. Researchers have reported, however, that this degradation can be mitigated through the addition of Lewis-basic additives. The presence of H$_2$O in electrolytes can also reduce the production of C$_2$H$_4$ (usually used as an indicator of EC reduction) and promote H$_2$ production. Here, the existence of SEIs can reduce the production of trace amounts of water that usually generate H$_2$ and CO$_2$ [114].

An automotive scale NMC pouch cell was also analyzed by Dahn group using ultra-high precision cycling technologies, and it was revealed that the performance of graphite anodes was decisive and that corresponding capacity fading (proportional to $i^{1/2}$) was related to SEI thickness and was inversely proportional to the rate of SEI growth, suggesting that the gradual lowering of SEI thickening rates can lead to stable SEIs [115, 116]. Here, alternative relationships proposed for SEI growth need to take stability as a factor into consideration [117].

### 2.4 Summary

Overall, the degradation of Ni-rich NMC-based cathodes originates from the insertion/desertion of Li$^+$ during charge/discharge in which the expansion and contraction of crystalline lattices (primary particles) gradually generates tension, leading to the cracking of secondary particles (Fig. 6). Electrolyte infiltration as well as oxygen release can further deteriorate this issue. On an atomic scale, Ni$^{4+}$ can accumulate during charging because delithiation can elevate the valence state of Ni together with high-valence Mn and Co ions. And due to smaller radii, easier diffusion can lead to surface reconstruction (from rhombohedral to rocksalt), which can further inhibit Li$^+$ insertion/desertion. Moreover, increased voltage levels of high-valence ions can trigger side reactions such as the oxidation/decomposition of electrolytes and the further dissolution of NMC materials. Electrochemical potentials of electron-depleted Co and Ni can also fall below the fermi level of O$_2$ to transition metal ions and the release of O$_2$ into systems, which can also cause electrolyte decomposition. Throughout the operation of Ni-rich NMC-based LIBs, four types of unwanted substances exist on the surface of cathodic particles, including other phases, surface impurities (carbonates and hydroxide), rocksalt structures and surface films. To
sum up, aging effects on both micrometer and atomic scales can result in two common outcomes in which the first is the chemical (through chemical reactions) and mechanical (through the breakage of crystallites) damage/loss of active materials and the second is the production of detrimental substances on electrode particle surfaces, including surface films (both SEIs and CEIs), surface impurities and surface layers. And although many differences in the performance of various Ni-rich NMC-based cathodes have been discovered, these remain unexplained. Therefore, future studies should focus on the interpretation of different aging behaviors in NMCs with or without mitigation methods.

As for graphite anodes, degradation primarily occurs due to the plating of Li dendrites and the cracking of layered graphitic structures. In addition, SEIs tend to thicken during cell operation as EC solvent is reduced electrochemically. Furthermore, transition metals, especially $\text{Mn}^{2+}$ from dissolved NMC, can cause significant capacity loss due to massive damage to SEIs. A comprehensive list of identified degradation mechanisms for Ni-rich NMC-based cathodes, graphite anodes, SEIs and CEIs is presented in Table 1. Here, the degradation mechanisms of SEIs are categorized based on anodes and the degradation mechanisms of CEIs.
are categorized based on cathodes because certain degradation mechanisms may involve both electrode materials and interfaces.

3 Diagnostic Tools

3.1 Electrochemical Techniques

Throughout the development of LIBs, a variety of electrochemical methods have been employed to test, analyze and diagnose LIBs. To provide a comprehensive overview, these methods are categorized into three groups, including charge/discharge tests, cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS).

3.1.1 Charge/Discharge Tests

Charge/discharge tests include basic electrochemical characterizations that can be used to determine the voltage at which a certain reaction (including Li⁺ insertion/desertion reactions that generate current in external circuits and side reactions that negatively impact cell performance) occurs and that can be used to evaluate the reversibility of cycling performances. Data from these charge/discharge tests can be analyzed by using many methods and can provide useful information on cell performance, reaction and reversibility, degradation and associated mechanisms.

**Voltage versus capacity** Charge/discharge curves are common and useful plots contained in almost every published study in the field of batteries. These plots are usually displayed in the format of voltage versus capacity (normalized by the mass of the active material) for a clear understanding of total capacity within a voltage range and capacity loss for certain cycles, including initial capacity loss (ICL), sudden capacity loss (SCL), gradual capacity loss (GCL) and voltage versus time. In some studies, the horizontal axis is changed to the content of Li in NMC compounds for easier comparison with physical characterization results in terms of lattice parameters/structures. Furthermore, the reversibility of cycling can be interpreted through these curves in which the presence of a reaction that can reconstruct the cathode at a certain voltage can be clearly predicted if the curve does not repeat itself in the next cycle after reaching that point [35]. Moreover, a small but important feature (a small peak) often found in the charge/discharge curve at the beginning of the initial charging can indicate the existence of impurities (most probably nickel hydroxide and nickel carbonate) during storage (a typical example showing the reaction of surface impurities of NMC811 after long-term storage during cycling is illustrated in Fig. 7a) [35]. Finally, the polarization of battery cells can be estimated by comparing the voltage levels of different cycles at fixed capacities.

**Differential Capacity Curve** Differential capacity curves are another kind of plots derived from charge/discharge curves [7].
These curves are also referred to as the incremental capacity and can be obtained by plotting $\frac{dQ}{dV}$ versus voltage. (For similar reasons mentioned above, the vertical axis is converted to $\frac{dx}{dV}$ in some cases, as shown in Fig. 7b.) The main use of these curves is to cross-check features already shown in charge/discharge curves at certain voltages but can also provide other information. Here, the most important feature revealed based on differential capacity curves for Ni-rich NMC-based cathodes is the phase transition in which three peaks can be identified in both charging and discharging processes, indicating phase transition from H1 (hexagonal) to M (monoclinic), and subsequently to H2 and H3 (see Sect. 2.1.1) as Li is extracted from the cathode [45]. In addition, differential capacity curves can be plotted with in situ X-ray diffraction results to correlate phases to corresponding lattice parameters [119]. Moreover, the loss of available Li between graphite layers or in the Li-rich layers of cathodes can be revealed through the fading of these peaks. And by integrating peak areas and calculating the ratio of these peak areas to the total charge/discharge capacities for each cycle, trends in the pathways/steps of Li transfer can be implied (e.g., more dominating, remaining the same or fading). Furthermore, the position of peak potential is indicative of cell impedance in which positive shifting during cycling implies an increasing cell impedance. Similarly, a differential voltage plot $\frac{dV}{dQ} \text{ vs. } Q$ can be used to show the fluctuation of voltage during charge/discharge and is also linked to phase transitions [98].

**Maximum Capacity** To determine maximum capacity and associated fading, electromotive force (EMF) data [120] can be derived from the discharge curve of the same cell at different C-rates. Here, comparisons of EMF curves with charge/discharge curves can reveal the impact of C-rates on cell degradation. In addition, discharge capacities at each time slot during discharge can be plotted versus corresponding C-rates and a trendline fitting of all data points taken at the same voltage can be drawn and extrapolate to the $Q$-axis, thus revealing the relationship between maximum capacity $Q_{\text{max}}$ and voltage. Moreover, maximum capacity fading plots based on EMF curves can be used to determine capacity loss (irreversible capacity loss $\Delta Q_{\text{ir}}$) at different C-rates. Differential voltage curves can also be applied to EMF, and $\frac{dV}{dQ}_\text{EMF}$ can be regarded as a nondestructive method in which peak shifting is indicative of cell degradation and can reveal anode material decay and voltage slippage effects [121].

**Capacity Fading** Capacity fading curves can provide more macroscopic views of capacity loss during long-term cycling and can be derived from capacity fading plots in which cell capacity at different conditions is displayed. And because electrode materials and aging conditions vary significantly in difference battery systems, the speed of capacity fading can also vary significantly. Therefore, the degree of cell aging needs to be normalized in which the vertical axis of capacity fading curves is sometimes converted to represent normalized capacity. A modified version has also been recently presented [16] to investigate the effects of the amount of Li$^+$ ions entering/exitng Li-layers in cathodes rather than focusing on the number of alterations. Here, an accumulative discharge capacity fading plot can be drawn and used to compare capacity fading in different currents, which can provide insights into the possible relationships between the amount of Li$^+$ transferred and structure deformations.
Based on capacity fading plots, the number of electrons consumed by side reactions (both at the cathode and at the anode) can be measured by subtracting the discharge capacity by the corresponding charge capacity. Here, a charge efficiency (CE) plot focusing on the degree of side reactions is commonly used in which the charge efficiency of an anode can be calculated by \( \frac{Q_{\text{charge,anode}}}{Q_{\text{charge,anode}}} \), because it describes the percentage of electrons charged to the anode that can be used to release \( \text{Li}^+ \) (remaining electrons consumed by side reactions during charging), whereas the number of electrons that can be discharged from the cathode can be calculated by \( \frac{Q_{\text{charge,cathode}}}{Q_{\text{discharge,cathode}}} \). And in general, the two percentages should be around 100% (CE\text{anode} is a bit less, Table 1

| No. | Condition | Degradation mechanism | Electrode | References |
|-----|-----------|-----------------------|-----------|------------|
| 1   | N/A       | Initial SEI formation consuming active Li | Anode     | [108]      |
| 2   | N/A       | Li diffusion into the NMC layer | Cathode   | [26]       |
| 3   | N/A       | Sluggish Li diffusion back into the cathode | Cathode   | [94]       |
| 4   | High voltage, high temperature | Sudden capacity loss caused by surface film formation and thickening under harsh conditions | Anode/cathode | [31]       |
| 5   | High SOC  | Ni\textsuperscript{2+} migrating to vacant Li\textsuperscript{+} sites (cation mixing), reducing the number of Li sites and blocking Li diffusion pathways | Cathode   | [51]       |
| 6   | N/A       | Structural changes from rhombohedral (layered) to spinel and later to rocksalt (NiO, nickel carbonate and minor quantities of hydroxides) as a result of massive cation mixing | Cathode   | [64]       |
| 7   | Long cycling | Mn\textsuperscript{3+} forms a spinel structure from a layered (rhombohedral) structure because of low octahedral site stability energy (based on studies of Mn-rich NMC cathodes) | Cathode   | [71]       |
| 8   | High SOC  | Ni\textsuperscript{2+} oxidizing electrolytes and thickening CEI (self-discharge) | Cathode   | [83]       |
| 9   | Storage in air | Surface impurities formed on particle surfaces due to long-term storage in air | Cathode   | [35]       |
| 10  | N/A       | NMC dissolution in HF (less active sites and higher charge transfer resistances) | Cathode   | [74]       |
| 11  | Extreme high temperature | Thermally unstable NMC reacting with LiPF\textsubscript{6} and causing thermal runaways (fire and explosion) | Cathode   | [33]       |
| 12  | N/A       | Release of reactive oxygen due to surface reconstruction causing chemical oxidation of electrolytes to produce CO and CO\textsubscript{2} | Cathode   | [25]       |
| 13  | High voltage | CO\textsubscript{2} produced from the electrochemical oxidation of electrolytes | Cathode   | [87]       |
| 14  | High voltage (above 4.6 V) | Oxygen reducing metal ions due to the hybridization of electrons and orbitals (self-redox reaction) | Cathode   | [82]       |
| 15  | Low voltage | Disproportion of Mn\textsuperscript{3+} (based on studies of Mn-rich NMC cathodes) | Cathode   | [75]       |
| 16  | N/A       | Carbonate layers reacting with electrolyte | Cathode   | [35]       |
| 17  | Long cycling, high DOD | Secondary particles cracking along grain boundaries and enlarging surface areas for more side reactions to occur | Cathode   | [13]       |
| 18  | N/A       | Loss of active Li metal to SEI as immobilized during cell operation | Anode     | [110]      |
| 19  | High voltage | SEI layer decomposition facilitated by the precipitation of transition metals into the SEI (huge capacity loss) | Anode     | [38]       |
| 20  | N/A       | Volume expansion and contraction of graphite during operation cracking the graphitic layer | Anode     | [102]      |
| 21  | N/A       | Metal ions from the cathode inserting into the anode and blocking active sites | Anode     | [111]      |
| 22  | N/A       | Mn\textsuperscript{2+} reacting at the surface of the anode to generate Li\textsubscript{2}CO\textsubscript{3}, which can crack graphite surfaces | Anode     | [38]       |
| 23  | N/A       | Solvents getting into the holes of graphite | Anode     | [31]       |
| 24  | N/A       | Solvated Li\textsuperscript{+} cracking graphite during Li intercalation | Anode     | [110]      |
| 25  | N/A       | Li dendrite plating (may trigger short circuiting if growth becomes too large) | Anode/cathode | [104]      |
| 26  | N/A       | Continuous thickening of surface films during cell operation | Anode/cathode | [37]       |

Note: N/A in the condition column indicates a common degradation mechanism occurring in many conditions; Only one representative reference is provided for each degradation mechanism because many of them have been reported extensively in many studies.
Fig. 7  

a Charge/discharge curves of the 1st, the 2nd and the 3rd cycles of NMC811-Li (Reprinted with permission from Ref. [35]. Copyright © The Author(s) 2018. Published by ECS).

b The c-lattice constant and differential capacity as a function of cell voltage for NMC811 (LiNi0.8Mn0.1Co0.1O2) (3–4.4 V) (Reprinted with permission from Ref. [20]. Copyright © 2016, American Chemical Society).

c Cyclic voltammetry of the first three cycles of an NMC 811/Li coin cell (Reprinted with permission from Ref. [33]. Copyright © 2018, Elsevier B.V. All rights reserved.).

d The Nyquist plot of NMC 811 (The inset: the equivalent circuit model used for data fitting). e The Nyquist plot of SiO2-coated NMC 811 (Reprinted with permission from Ref. [123]. Copyright © 2015, Elsevier B.V. All rights reserved.)
but $CE_{\text{cathode}}$ is a bit more); however, the existence of kinetic barriers may cause enlarged differences.

To study impedance growth during cycling, average polarization plots combined with discharge voltage decay plots can be obtained and can reveal capacity retention behaviors by measuring $\Delta V_{\text{ave}}$ increases, which can indicate impedance growth through cell degradation as well as potential kinetic barriers of Li$^+$ insertion/desertion [9].

3.1.2 Cyclic Voltammetry

As a supplementary method, cyclic voltammetry (CV) can be used to determine the reversibility of charge/discharge in electrodes by measuring the difference in charge/discharge peak potentials, which is related to major delithiation/lithiation processes (Fig. 7c) [33]. As such, CV can assist in the investigation of transition metal oxidation and reduction in which the plot of the initial CV can differ from subsequent cycles due to the formation of SEIs during the first delithiation. In addition, upper cutoff voltages can also be determined by using CV [122].

3.1.3 Electrochemical Impedance Spectroscopy

Electrochemical impedance spectroscopy (EIS) is another common method to investigate cell impedance evolution and degradation mechanisms and involves the application of AC sinusoidal waves (voltage or current) to target cells. Here, AC sinusoidal waves are applied at an amplitude with different frequencies because semicircles in resulting Nyquist plots can overlap with each other if represented processes occur in the same frequency range. In addition, half-cell EIS can be carried out with either 2-electrode symmetric cells (e.g., Swagelok 2-electrode symmetric configuration cells) or 3-electrode cells by installing a reference electrode in which the resulting EIS curve often involves a vertical short tail, two depressed semicircles (only if a half cell is tested) and a long tail at a certain degree to the horizontal axis from high to low frequencies. Here, the short tail usually appears at extremely high frequencies, indicating the inductance of cables, which is rarely seen in test results. Solution resistances can also be measured at the intercept of the short tail with the real ($Z$) axis and therefore can be represented by a resistor. As for the first semicircle at high frequencies (usually a small one), this is indicative of the resistance of the CEI, and as the CEI continuously reacts (thickening and decomposing), the impedance cannot be taken as merely resistive. The second semicircle can be attributed to charge transfer at the cathode. (Double-layer capacitance is generated because of voltage decay across the interface.) In typical EIS curves, these semicircles can appear as depressed due to surface roughness (unevenness), which can cause varied capacitance from point to point. To address this, a resistor in parallel with a constant phase element (CPE) can be used to model each process. As for the long tail attached to the two semicircles in the EIS curve, this is related to mass-transfer resistance and therefore can be represented by using a Warburg element. During aging, both semicircles commonly increase (Fig. 7d, e), indicating large cell polarizations and thickened CEIs; however, with remedial efforts, the increase in both resistances (linked to both semicircles) can usually be reduced. For example, researchers have reported that impedance growth of surface films and charge transfer can be significantly reduced through a SiO$_2$-layer coating of NCM811 (Fig. 7e) [123].

3.2 Physical Techniques

3.2.1 Crystal Lattice and Particle Morphology

**X-ray Diffraction** In situ and ex situ XRD methods are the most frequently used physical characterization methods to determine lattice parameters and crystal structures. Here, in situ XRD can be used to measure the evolution of lattice parameters (both $a$-axis and $c$-axis) of NMC-based cathodes during cycling [43]. For example, the use of in situ XRD can evidence the shrinking of the $a$-axis to a constant value and the expansion and rapid shrinking of the $c$-axis during delithiation in Ni-rich NMC cathodes through the shifting of (003) and (006) peaks to a lower angle together with the shifting of the (101) peak to a higher angle (Fig. 8a).

In addition, in situ XRD can allow for the comparison of the lattice shrinkage percentage of the $a$-axis to that of the $c$-axis, leading to the conclusion that lattice shrinkage is anisotropic and that lattice parameters are linked to corresponding phases (H1, M, H2, H3). In situ XRD can also be used to test the thermal stability of NMC cathodes and show temperature-resolved (TR) results [42] similar to voltage-resolved results. For example, TR-XRD results of heated NMC-based cathodes can be used to examine thermal stability in which researchers found that the structures of all NMC cathodes tested experienced surface reconstruction from rhombohedral to spinel (first the LiMn$_2$O$_4$ type and later the Mn$_3$O$_4$ type) and subsequently to rocksalt structure (Fig. 8b). In another study, in situ XRD plots were combined with gas chromatography–mass spectroscopy (GC–MS) data to reveal the relationship between the Li content and evolved gas from NMC-based cathodes [90].

Ex situ XRD with Rietveld refinement is easier to conduct than in situ XRD but conveys less information because it can only illustrate the crystal structure of NMC-based cathodes. For ex situ XRD data, more peaks can be seen if a bulk phase of impurities or coating materials is formed on the
surface of cathode materials. Here, the ratio of $I_{003}/I_{104}$ in the ex situ XRD data indicates the degree of cation mixing and can be used to determine the ameliorative performance of a mitigation method in which ratios lower than 1.2 represent significant cation mixing. Moreover, the peak splitting of (110)/(108) and (006)/(102) represents a well-layered hexagonal structure (Fig. 8c) [45] and the evolution of the full width at half maximum (FWHM) of peaks represents change in crystallinity. Surface species can also cause peak shifting [e.g., (003) of NMC622] if stored [72], or generate new peaks, and therefore, ex situ XRD can also be used to cross-check the existence of impurities.

**Microscopy and Electron Diffraction** Scanning electron microscopy (SEM) and transmission electron microscopy (TEM) can be used to reveal microscopic structures of NMC-based cathodes and assess structural damage [124]. In addition, diameters of primary and secondary particles can be measured from obtained images together with the thickness of any surface coating layers [100]. Here, aggregates of primary particles, i.e., secondary particles, are usually observed in electron microscopy images possessing elongated or even needle-like shapes before degradation, whereas microcracks on secondary particles can usually be seen after repeated cycling, which can cause the breakage of particles. As a result, the spherical shape of secondary particles is usually distorted, and subfigures of SEM images obtained from samples aged under different conditions are usually displayed together for comparison. Another indispensable microscopic method is high-resolution transmission electron microscopy (HRTEM), which can reveal microregions of $Fm\overline{3}m$ NiO and spinel structures and allow for the distance between atomic layers to be clearly marked (Fig. 9a shows a surface reconstruction region on a NMC-based cathode and an electron diffraction pattern of selected spots) [56, 70]. A drawback of HRTEM, however, is that it is unable to distinguish between phases with similar structures and lattice parameters [50].

Electron diffraction is a useful tool to characterize phases and microregions in NMC lattice crystals and includes selected area electron diffraction (SAED) and nano-beam electron diffraction (NBED, in which a condenser lens and an upper objective lens are used to achieve nano-beams of electrons). Here, Kang et al. [56] have used electron diffraction to measure the space group of NMC-based cathodes with the zone axis at the bulk and surface regions of a NMC particle. In addition, electron diffraction is usually conducted separately on different regions of particles such as surface regions, transitional regions and bulk regions. Here, the existence of secondary phases can be confirmed based on obtained patterns in which spinel structures can lead to an additional spot (Fig. 9a2–a5), whereas $Fm\overline{3}m$ space groups will show two strong diffraction spots that indicate the formation of rocksalt phases with high symmetry. Phase conversion can also be tracked and compared by using HAADF-STEM results [9].

High-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) measurements can provide array images of transition metal atoms located from the bulk to the surface of NMC particles and spectra of electron energy loss spectroscopy (EELS). Here, orderly arrays of atoms can clearly display layered structures of the crystal lattices of NMC (Fig. 9b) [63, 124, 125]. Here, the degree of surface reconstruction can be determined based on changes in atom patterns and a clear boundary between reconstructed and unreconstructed phases can be traced. In addition, region thicknesses can also be measured based on HAADF-STEM images.

### 3.2.2 Chemical Analysis

**Elemental Analysis** Elemental analysis is necessary for the physical characterization of NMC-based LIBs and the composition of NMC-based cathodes as well as the investigation of impurities and remedial coatings/dopants. In addition, the semiquantitative properties of various elemental analyses are useful in the quantification of relative increases/decreases of certain elements during operation or storage. Here, energy-dispersive spectroscopy (EDS) mapping analysis can illustrate elemental distributions (Li, transition metals, C, etc.) semiquantitatively on cathode particle surfaces. For example, the dissolution of these surfaces through attack with HF can be evidenced by the increase in F element signals and the loss of transition metals.

Alternatively, electron-probe X-ray microanalysis (EPMA) accompanied with SEM can allow for the examination of transition metal element concentrations within different regions of NMC particles with a spatial resolution of ~0.3 μm and can often be used to determine the concentration gradient layer of modified-NMC particles [126]. Researchers have also reported that the average composition of Ni, Mn and Co can be measured using atomic absorption spectroscopy (AAS) [126].

X-ray photoelectron spectroscopy (XPS) is another elemental analysis method that can detect the chemical and electronic states of transition metals and oxygen in NMC cathodes as well as elemental carbon from current collectors and binders [73]. Here, the 2p orbitals of Ni, Mn and Co and the 1s orbital of O can be captured to reveal oxidation states by checking the peak positions of $2p_{1/2}$ and $2p_{3/2}$ (Fig. 9c) [127]. For example, the binding energies of Ni $2p_{3/2}$ in NiO and in $Ni_2O_3$ were found to be located at 854.0 and 856.0 eV, respectively [50]. In addition, the shift in oxidation states can be clearly shown by changes in peak positions,
and changes in the amount of Ni can be deduced from the increase in peak area and intensity. Moreover, the use of cycle-dependent XPS can allow for the change of valence states to be attributed to a certain range of cycles, whereas a decrease in XPS peaks can indicate the loss of elements or the coverage of elements with surface impurities. Furthermore, XPS can enable the identification of new substances on cathode surfaces that evolve during storage or operation [35]. As for organic elements including C (1s), O (1s), F (1s), N (1s) and S (2p) that are contained in the components of SEI layers, XPS can also be used to track the existence of these elements in different molecules versus voltage [128].

To validate XPS data, X-ray absorption spectroscopy (XAS) can be used as a supplementary technique to monitor transition metal valence changes during charge/discharge and include extended X-ray absorption fine structure (EXAFS) and X-ray absorption near edge structure (XANES) [129]. For example, stronger Ni^{2+} signals over Ni^{4+} in NMC532 at 4.5 V reveal that the redox reaction of Ni^{4+} with electrolytes can occur, which reduces Ni^{4+} to Ni^{2+} [130]. And due to the viability of XAS in the examination of transition metal concentrations and oxidation states, XAS has also been widely utilized for anodic elemental analysis (e.g., transition metals deposited on graphite anodes) under various cycling conditions [113], leading to the discovery that the massive dissolution of Mn can occur if NMC111/graphite cells are charged above 4.6 V [76].

Electron energy loss spectroscopy (EELS) is another elemental analysis method that can investigate elemental energy loss and possesses three modes, including auger electron yield (AEY), total electron yield (TEY) and total fluorescence yield (TFY), each of which provides information from different depths (1–2 nm, 2–5 nm and 50 nm, respectively). Due to smaller probing depths, EELS signals reflect information on the top layer of cathodes [63], and the shift in energy loss peaks from time to time or from the bulk to the surface can indicate shifts in oxidation states of elements, in which lower energy losses translate to lower oxidation states of transition metals [9]. In general, each EELS curve consists of two peaks, including a pre-edge peak and a primary peak. Here, the pre-edge peak of oxygen is thought to be caused by the transition of electrons from the 1s to the unoccupied 2p state, which can hybridize with the 3d orbitals of transitional metals [66]. Alternatively, the primary

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Fig. 8 a In situ XRD patterns of an NMC 811 cathode over a voltage range from 3.330 to 4.298 V during the first charge (Reprinted with permission from Ref. [43]. Copyright © 2015 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim). b Contour plots of TR-XRD patterns at the selected 2θ range for charged b1 NMC433, b2 NMC532, b3 NMC622 and b4 NMC811 (Reprinted with permission from Ref. [42]. Copyright © 2014, American Chemical Society). c Ex situ XRD patterns of NMC-811 and Li_{2}MnO_{3}-coated (10% and 30%) NMC-811 (Reprinted with permission from Ref. [45]. Copyright © 2016, American Chemical Society)
peak originates from the transition of electrons from the 1s to the 2p state of O or to the 4sp state of transition metals. Moreover, area-integrated EELS data can be compared to check the homogeneity of NMC particles [63].

Electron paramagnetic resonance (EPR) spectroscopy can also be applied to NMC materials such as NMC622 to investigate the effects of doping with Al and Fe on cation mixing in which the change of the Ni\(^{2+}\) line indicates the redistribution of Ni within the structure. Hubert Gasteiger’s group have also demonstrated Li plating onto graphite anodes quantitatively and in real time by using operando EPR [106] quantified the amount of Li-ions not available for future cycling. Despite these uses, EPR cannot provide more detailed structural information, which requires other physical characterization methods [131].

Inductively coupled plasma–atomic emission spectroscopy (ICP-AES or ICP) is a cutting-edge technique that can provide relatively precise information on the amounts of Li [132], transition metals [133–135] and impurities in examined samples in which concentrations can be indicated by the intensity of characteristic waves emitted from excited atoms or ions. In addition, changes in the elemental composition of electrodes or powder samples due to operation or modification can be reflected in ICP-AES data and the results (in ppm) are often compared with theoretical values or results obtained by using other elemental analysis tools to observe differences. Researchers have also combined ICP with mass spectroscopy in studies. For example, ICP-MS was used to reveal significant increases in the Ni content on graphite anodes related to the dissolution of Ni from NMC 811 cathodes [136] and was also used to conduct a series of Li content measurements in sampled electrolytes, including sample preparation procedures [137], in which the

**Fig. 9** HRTEM images and FFTs after 50 cycles under 3.0–4.8 V conditions. **a1** Lattice image of the surface region where **a2-a5** correspond to the FFTs of regions 1–4, respectively. (11-1)c is the diffraction spot of the rocksalt phase of the metal monoxide (Reprinted with permission from Ref. [56]. Copyright © 2013 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim). **b1** Low-magnification STEM-HAADF image of one surface primary particle of the x=0.10 sample. **b2** Corresponding X-ray spectrum from the highlighted area in (a). **b3** High-resolution STEM-HAADF image showing the lattice structure from the highlighted area in (b1) (Reprinted with permission from Ref. [124]. Copyright © 2016, Elsevier B.V. All rights reserved.). **c** XPS data for lithiated FCG-Mn-F Li(Ni\(_{0.60}\)Co\(_{0.15}\)Mn\(_{0.25}\)) O\(_2\) : **c1** Ni 2p, **c2** Co 2p and **c3** Mn 2p (Reprinted with permission from Ref. [127]. Copyright © 2013, American Chemical Society)
results were pre-calibrated with known Li concentrations and proved the credibility of the ICP-MS method.

**Gas Measurement** Ex situ gas measurements have been adopted by many researchers in the study of LIBs in which the volume of evolved gas is measured as the cell is charged to certain voltage levels [9, 85]. Gas evolution under other conditions can also be measured, including the addition of electrolyte additives or under different compositions of Ni, Mn and Co. And if combined with mass spectroscopy, the content of emitted gas can be identified (usually including H$_2$, C$_2$H$_4$, CO and CO$_2$) and allows for volume or moles versus voltage to be plotted with various curves related to different gases [24]. Furthermore, online electrochemical mass spectrometry (OEMS) is a tool that can facilitate the detection of gas in real time and can quantify the amount of H$_2$, CO$_2$ and O$_2$ from different mass to charge ratios [23, 90] in which gas evolution (e.g., O$_2$) is related to cathode surface reconstruction (if aimed at cathodic processes) through comparisons of voltage profiles (or SOC) (Fig. 10a). Similarly, anodic processes with gas evolution can also be tracked by using OEMS. Researchers have also combined differential electrochemical mass spectrometry (DEMS) with differential electrochemical infrared spectroscopy (DEIRS) to identify and quantify gases emitted from NMC 532/graphite cells [84] and found that potential-resolved gas evolution partially originates from SEI formation. And if combined with thermogravimetry (TG), the mass loss of cells can be correlated to oxygen evolution from NMC-based cathodes heated to a certain temperature, showing that the majority of weight loss can be attributed to oxygen release as they both rapidly rise to around 175 °C [85].

**Impurities and Organics** Raman spectroscopy can be used to determine the existence of surface impurities that can cause capacity fading such as nickel carbonate, lithium carbonate and nickel hydroxide [35]. Because of this, Raman spectroscopy is a useful tool in the identification of impurities produced during long-term storage (Fig. 3a). Similarly, Fourier transformed infrared spectroscopy (FT-IR) can be used to identify impurities on cathode surfaces [72] or organic coatings [138]. For example, the O–H stretching mode from LiOH possesses a characteristic band between 3650 and 3200 cm$^{-1}$ in corresponding FT-IR spectra, whereas 1450 and 870 cm$^{-1}$ are regarded as characteristic peaks of the C=O mode originating from Li$_2$CO$_3$ [61]. Raman spectra of NMC-based cathodes can also be voltage dependent to monitor band shifts of A$_{1g}$ and E$_g$ of Ni, Mn and Co. Moreover, FT-IR and Raman spectroscopy are widely used in the characterization of anodic graphite and SEIs because they can identify substances forming SEI layers such as lithium alkyl carbonates and therefore can reveal reaction mechanisms involved in the SEI formation process and present differences in mechanisms if multiple electrolyte additives are used [139].

### 3.2.3 Thermal Stability

Differential scanning calorimetry (DSC) is a method that utilizes microcalorimeters to measure heat flow from cells (W g$^{-1}$) in which the thermal stability of NMC-based cathodes with/without electrolyte contact is recorded to reveal heat flow from the NMC-based cathode as it is heated to a certain temperature [33, 140]. For example, in two NMC-based cathodes and LiNi$_{0.7}$Mn$_{0.3}$O$_2$ with a concentration gradient layer, one major peak can be found for each DSC curve centering at a certain temperature in which a higher onset temperature of the DSC peak indicates better thermal stability (Fig. 10b) [126]. In addition, exothermic peaks can be integrated to calculate the amount of heat released, which can translate into the degree to which corresponding reactions occur [99]. Here, the application of common mitigation methods usually results in smaller exothermic peak areas and higher onset temperatures [100]. However, small endothermic peaks can also be seen for some cases, indicating the decomposition of Li salt in the electrolyte [101]. In addition, TG is often combined with DSC to record mass loss during the heating of NMC-based cathodes (Fig. 10c), including processes such as the addition of carbon matrixes to remedy cathode degradation [141] or the release of oxygen due to surface reconstruction [140], or through the loss of surface impurities and adsorbed species [72].

Accelerating rate calorimetry (ARC) is another method adopted by researchers to measure self-heating rates (SHRs) versus temperature [8, 142]. For example, in the comparison of the SHR of 4 types of Ni-rich NMC-based cathodes (Fig. 10d), the onset temperature of a rapidly increasing SHR marks the upper limit of the temperature range for the safe operation of a cell [7] in which cathodes with higher SHR onset temperatures are considered to be better LIB cathodes for industrial applications. Isothermal microcalorimetry (IMC) has also been used to examine heat flow induced by polarization, entropy and parasitic reactions versus the state of charge of NMC811 and has revealed that heat flow originating from parasitic reactions can increase sharply if the relative SOC rises from 0 to 1 at certain voltages (Fig. 10e) [7].

### 3.2.4 Advanced Physical Characterization

Many advanced physical characterization methods have also been used by researchers to study LIBs. For example, 3D transmission X-ray tomography mapping [92] can be used to determine the distribution of transition metals in NMC particles in which 3D mappings can allow for elemental analysis within cross-sections at different depths. Surface potential mappings of NMC-based cathodes can also be conducted by using Kelvin probe atomic force microscopy (AFM) [143] in which smaller surface potentials can allow for easier electron
The use of nano-secondary-ion mass spectrometry (nano-SIMS) analysis can provide good spatial resolution (less than 50 nm) for elemental mapping [144]. And with higher resolution, different phases can be identified by comparing the ratios of Mn to Ni along a direction across the particle. In situ neutron powder diffraction (NPD) can also be used to obtain more accurate results of lattice parameters as compared with in situ XRD [145], in which although the energy of neutrons is lower than X-rays, the penetration
depth is higher. In situ NPD can also be used to determine the d-spacing of anodic graphite during Li intercalation and therefore the amounts of LiC₆, LiC₁₂, and LiC₂₄ [44]. Despite these advantages, researchers have reported that hydrogen-containing species can attenuate neutrons. As for prompt gamma activation analysis (PGAA), this method can be used to quantify the amount of transition metals deposited onto graphite anodes by placing the sample in a neutron beam and collecting signals [26]. The use of PGAA allowed researchers to discover that cycling and high cutoff voltages can promote transition metal deposition onto graphite anodes.

4 Mitigation Strategies for Ni-Rich NMC-Based Cathodes and Graphite Anodes

4.1 Ni-Rich NMC-Based Cathodes

Ni-rich NMC-based cathodes suffer significantly more from capacity loss and impedance growth as compared with low Ni content NMC-based cathodes such as NMC111. This is because large amounts of Ni³⁺ on cathode surface regions can trigger side reactions and Ni²⁺ can cause cation mixing. In addition, excessive Li is usually added in the preparation of NMC-based cathodes and tends to accumulate on the surface of particles and react with electrolytes to convert to lithium carbonate and hydroxide. Furthermore, the continuous charging and discharging of these cathodes can cause the cracking of secondary particles.

To address these issues, researchers have proposed many strategies. One common method is the mixing of LMO with NMC [45, 118, 124], which can enhance capacity retention (not reviewed in this review because this method does not directly modify cathode particles) and another method involves the addition of electrolyte additives. Here, organic additives can be sacrificed at high voltages (4.3–4.5 V) because their HOMOs are higher than the HOMOs of organic solvents to preferentially lose electrons and be oxidized to prevent the oxidative decomposition of solvents [146]. In addition, the electropolymerization of electrolyte additives can also provide electrons to counteract the decomposition of solvents [147]. Moreover, the use of electrolyte additives can allow for the formation of more even CEIs on cathode surfaces [148, 149], which can reduce contact between electrode surfaces and electrolytes to prevent side reactions and lower impedance. One representative electrolyte additive is fluoropropylene carbonate (FEC) [150, 151] in which vinyl fluoride and Li₂CO₃ are produced as oxidation products during cell operation. Wang et al. [146] have also reported that 3-IPTS as an electrolyte additive can consume trace amounts of water in electrolytes to minimize the amount of HF produced from reactions between LiPF₆ and water. Furthermore, Li salts can also be used to improve NMC-based cell performances. For example, LiPF₆ can react with excessive alkaline Li on the surfaces of cathode particles to produce chemically stable Li₃PO₄ and LiF, and significantly suppress parasitic reactions caused by excessive Li [152]. Surface coating is another commonly used strategy to separate NMC cathode surfaces from electrolytes [15], and atom substitution has also been reported to be effective in enhancing the performance of mitigated Ni-rich NMC-based LIBs. Based on these strategies, this review section will present techniques applied to improve the cycling performance of NMC cathodes in the most recent decade, in which cited studies are divided into categories based on the type of modification, including inner surfaces (doping/gradient layers) and outer surfaces (coating/matrixes) of Ni-rich NMC-based particles. New synthesis methods of Ni-rich NMC cathodes that enhance performance without introducing new substances are also presented.

4.1.1 Inner Surface

**Doping** Doping can minimize side reactions on cathode particle surfaces and enhance the rate of Li diffusion and charge transfer. In addition, active material costs can be reduced if Co is substituted with other elements. To achieve doping/atom substitution, researchers often use co-precipitation, solid-state reaction, self-combustion and co-calciation, in which the two types of doping include cationic doping and anionic doping. And in general, few dopants can affect the layered structure of electrodes, but some can form new phases (e.g., the presence of a Li₂SnO₃ phase in Sn-doped NMC622 [153] and the presence of Li₂ZrO₃ in Zr-doped NMC532 [154]).

Cation-doped surfaces of Ni-rich NMC particles can benefit from the lowered Ni content and therefore suppressed cation mixing [155]. Here, the selection of dopants is based on the bond energy of M–O and the radius of dopant ions in which trade-offs between structural stability and Li⁺ movement need to be considered. For example, strongly bonded dopants to oxygen can improve structural stability and reduce the amount of oxygen released [156, 157], but in most cases tend to withdraw more electron density from the M–O bond, leading to the reduction in oxygen repulsion and the vertical contraction of the crystal lattice (Fig. 11a) [153], which negatively affects Li⁺ movement due to increased kinetic barriers [158]. For easier lithiation/delithiation, increased distances between atomic layers in the lattice structure are desired and can be achieved by using dopants with larger radii. For example, dopants such as Fe³⁺ and Sn⁴⁺ (with radii larger than substituted Co³⁺) can expand Ni-rich NMC crystal lattices horizontally [145] and Ca²⁺ (with a radius larger than Ni²⁺) can expand lattices horizontally by substituting...
Ni at 3a sites [53]. Researchers have also reported that doping with Al can contract cathode lattices slightly due to the smaller radius of Al\(^{3+}\) [159]. Here, reported results of different cationic dopants can sometimes be contradictory in which enlarged lattices due to cationic dopants with larger radii can either promote (mostly reported) or hinder (blocking effect, rarely reported) Li\(^+\) diffusion in different studies. Because of this, these results need to be comprehensively reviewed in the future [154, 160].

To achieve better cycling stability, capacity is sometimes sacrificed because some dopants such as Al\(^{3+}\) are chemically inactive [28]. Cr doping can improve cycling stability without capacity decrease [161], and Zr doping can enhance cycling performance [154, 162] but will reportedly produce a Li\(_2\)ZrO\(_3\) phase. In addition, different dopants possess different valences and stabilize crystal structures at different sites. For example, Mg\(^{2+}\) possesses a similar radius to Li\(^+\) and will enter the Li-layer rather than the transition metal layer on the surface region of Li-rich NMC particles [125, 163], allowing for the suppression of cation mixing due to the repulsion of Mg\(^{2+}\) to transition metal ions and the delaying of the surface transformation to a spinel structure (Fig. 11b, c). Alternatively, Na\(^+\) can substitute for Li\(^+\) at 3a sites [52, 164] to enlarge the distance between layers, and benefitting from the larger radius of Na\(^+\), cation mixing can be suppressed [165] and weakened Li–O bonds can enhance potential capacity. However, because Na\(^+\) doping can enlarge primary particles, optimal amounts are limited to ensure a connection between primary particles. Chen et al. have also suggested that the standard of choosing appropriate cationic dopants for NMC cathode improvements had not been established [53], whereas others have reported that ions with low valences were favored for Li\(^+\) diffusion [166]. New methods of cation doping have also been proposed in which Mn doping into NMC71515 can provide an outer region with \(Fm\overline{3}m\) Ni\(^{2+}\) (reduced from Ni\(^{3+}\)) residing in Li slabs (pillar structure) and can be used to consolidate the layered structure of cathodic crystal lattices and avoid particle degradation [167]. As for anionic doping, fluorine is usually used [168] and can expand both lattice parameters by reducing transition metal ions (mainly Ni and Mn) [169] and increase mean charging voltages due to increased bond energy of Li–F as compared with Li–O [170]. In addition, fluorine substitution can protect cathode materials from HF. Furthermore, although bromine and sulfur atoms can be used to enhance NMC cathode performances, few studies have shown their effect on Ni-rich NMC-based cathodes. And currently, studies into the surface modification of Ni-rich NMC-based cathodes have been insufficient and the effects of dopants such as Ti [171], Zn [160], Cl and Br [172] and their influences on Ni-rich NMC-based cathode cycling stability need to be further explored.

**Gradient Layer** Modifications of Ni-rich NMC-based cathode particles have been investigated by many studies in which a typical modification is a concentration gradient layer that can cover a Ni-rich NMC core [126, 173–175]. This concentration gradient layer design originates from a simple core–shell structure in which the shell (single or multiple) is composed of other NMC compositions (lower Ni content and higher Mn content) to stabilize the layered structure. And to overcome drawbacks of voids generated between the shell and the core during long-term cycling and mismatches of different lattice parameters, gradient layers were invented to replace core–shell structures. Here, the composition of the gradient layer is different from the core material, in which Ni atoms are gradually replaced by Mn atoms from the interface to the outer edge, resulting in a surface with higher structural integrity (Fig. 12a). Furthermore, concentration gradient layers can reshape primary particles into a rod-like structure, which can promote Li\(^+\) movement by minimizing diffusion pathways [140]. This feature can be further magnified in needle-shaped structures for some cases and turned into plates if calcined [175, 176]. Modifications with concentration gradient layers can also prevent the separation of cores and shells caused by continuous expansion and contraction. Moreover, gradient layers can be extended into full concentration gradient (FCG) layers [175, 177, 178] and multiple gradient layers with different slopes can also be obtained to maximize the Ni content at the core as compared with single-sloped gradient layers [179]. For example, Noh et al. [127] reported a NMC (60 25 15) particle with a constant Mn concentration from the center to the surface and a gradually decreasing Ni and increasing Co concentration and found that the composition of the surface layer did not change significantly after long-term cycling.

**4.1.2 Outer Surface**

**Li\(_x\)M\(_y\)O\(_z\)** coating Li\(_x\)M\(_y\)O\(_z\) coatings can be applied to the surface of primary particles to enhance performance and are composed of two subgroups. One group involves layered NMCs with different compositions/structures as compared with core Ni-rich NMCs and usually possesses higher Co or Mn content. Here, a Mn-rich NMC shell can significantly enhance the chemical and structural stability of Ni-rich NMC cores [180, 181] in which studies have shown that the coating of Mn-rich spinel oxide onto Ni-rich NMC can significantly improve chemical stability in both operation and storage [144, 182]. The second group includes all other types of Li\(_x\)M\(_y\)O\(_z\) coatings (different transition metals or structures) and also includes pioneering works with disparate coating materials and methods in which Si, Zr, Ti and Al are usually used. For example, Li\(_2\)TiO\(_2\) with a rocksalt structure can be coated onto NMC532 due to its superior structural integrity and strong adhesion to the core...
Fig. 11  a Lattice parameter trends of Co and Sn-doped NMC-622 cathode materials in fully lithiated (black squares) and 60% delithiated (red squares) states (left) along with corresponding schematics of Li-interlayer movement along the c-axis due to doping effects (right) (Reprinted with permission from Ref. [153]. Copyright © 2018, Elsevier B.V. All rights reserved.) and structural evolution in bare and Mg$^{2+}$ pillared Li-rich cathode materials. b Mn migration in bare Li-rich materials from the TM slab to the Li slab through the adjacent tetrahedral site. c Prohibition of Mn migration in Mg$^{2+}$-doped layered oxide by electrostatic repulsion between Mg$^{2+}$ and TM ions (Reprinted with permission from Ref. [125]. Copyright © 2015 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim)
an ionic-conductive Li₂ZrO₃ coating layer on NMC71515 can form large islands of the coating material, protect the core and facilitate Li movement [184]. In addition, Li₂SiO₃ is also a promising coating material because it is a fast ionic conductor that can provide two pathways for Li⁺ movement [(010) and (001) planes] [185]. Furthermore, a recent study using LPO infusion to coat NMC particles has gained much attention due to the creation of solid–solid interfaces in the gap between primary particles to replace liquid–solid interfaces, thus reinforcing the microstructure by shielding off liquid electrolytes and eliminating side reactions within secondary particles (Fig. 12b) [97]. Researchers have also suggested that other types of cathode materials with more cycling stability can be used as coating materials such as LFP [186].

**Li-Free Coating** Li-free coatings on the outer surface of NMC cathodes have been extensively investigated due to a unique advantage over doping in which the valence of transition metal ions remains unaffected and so does the theoretical capacity. Here, materials used for Li-free coating include transition metal phosphates [187], metal oxides [99, 188, 189], fluorides [143, 190], silicon oxide [91, 100] and organic coating layers [138, 191] in which the general concept is to prevent cathode surfaces containing highly reactive Ni⁴⁺ from contacting the electrolyte and therefore to reduce side reactions. Here, coating layer thicknesses are usually between 5 and 40 nm and similar to doping, XRD measurements usually do not show new characteristic peaks and instead, existing peak widths/positions may change slightly. In addition, the metal ions in coating layers tend to diffuse into the cathode lattice and can change lattice parameters [192]. To meet the basic demands of coating materials, high conductivity (coatings tend to possess higher resistances and are even insulating, however) and firm adherence to NMC cathodes are preferred [138]. And to optimize cycling stability, the weight ratios of coating precursors need to be carefully estimated because excessive coating can block Li transport pathways and result in large polarization and low Li-ion diffusivity. This careful estimation of coating precursor weight ratios can also improve the uniformity and dispersity of precursor particles at the micrometer scale [141, 188].

Among various metal oxide coatings studied, Al₂O₃ has received the most attention due to multiple beneficial features (Fig. 12c) [12]. For example, Chen et al. have found that coating layers of Al₂O₃ and nano-TiO₂ can suppress cation mixing (Li⁺ and Ni²⁺) as indicated by the ratio of \( I_{003}/I_{104} \) in XRD [99, 188] and that Al₂O₃ layers can reduce self-redox reactions by introducing new energy levels to prevent the reduction of Ni ions by O²⁻. In addition, researchers have also found that Al₂O₃ surface coatings can maintain stabler CEI structures [188, 193] and that the atomic layer deposition (usually used to enhance coating uniformity through the precise control of coating thicknesses to several nanometers or even thinner) of Al₂O₃ on NMC532 [101, 130] can suppress parasitic reactions and phase transitions in which one study by Dahn group suggested that performance stabilizations were due to the protection of cathode surfaces to surface-oxidized species rather than the prevention of electrolyte oxidation [91]. Furthermore, researchers have also reported that the crystallization of amorphous TiO₂ applied on NMC811 surfaces can result in Li₂TiO₃, which can further inhibit phase transformation from H2 to H3 [96]. The use of metal oxide layers has also been reported to be able to function as mediators of HF attacks due to the protection they can provide to NMC cores from corrosion [194].

The application of Li-free coatings can also lead to issues in Ni-rich NMC-based cathodes. For example, Park et al. have reported that excessive Li on cathode surfaces as caused by the coating of metal phosphate [187] can cause gas evolution due to its possible transformation into lithium carbonate and hydroxide as well as further side reactions with electrolytes [61]. However, researchers have also reported that the use of phosphate can lead to the formation of lithium phosphate [195] and lithium transition metal phosphate, both of which can maintain cell performance. Alternatively, FePO₄ coatings have been shown to be resistive to cathode dissolution in electrolytes [192] and \( V_2O_5 \) coatings [189] applied through a wet coating method by using \( NH_4VO_3 \) were found to be capable of removing excessive Li. Moreover, the introduction of acids or metal oxides can allow for excessive Li on the surface of cathodic particles to transform into \( Li_xM_yO_z \) coatings at certain temperatures [196]. For example, researchers suggested that the use of excessive surface Li on NMC811 to form a protective layer consisting of Li⁺-conductive \( Li_3AlO_3 \) and superconductive \( Li_2Ti_2O_7 \) [197] was a cost-saving and efficient method to resolve the issue of excessive Li and improve conductivity.

Organic coating layers possess a unique advantage of being stretchy, which allows for the absorption of internal microstrains and the mitigation of primary particle cracking [191]. And recently, a novel coating method using the transesterification of organophosphates has been reported [198] in which surface coatings were achieved through the replacement of M–OH bonds of NMC622 with M–O–P bonds. Here, this novel coating differs from others because it can be achieved through the chemical deactivation of surface groups. Researchers have also reported that like doped fluorine, fluoride coatings can also be used to prevent HF attacks [190]. For example, \( YF_3 \)-coated NMC111 can exhibit enhanced electron transfer due to smaller surface potentials and reduced oxidation of electrolytes due to smaller work functions [143]. Despite these promising results, the number of studies into metal fluoride coatings on Ni-rich NMC-based cathodes is limited. Alternatively, sulfated ZrO₂ as a
coating material has shown particular promise [193]. This is because different from regular/ultrathin ZrO2 coatings [199], sulfated ZrO2 coatings can incorporate sulfate groups into coating layers and functionalize coatings by forming −SO3− to stabilize CEIs on cathode surfaces, indicating that these coatings to a large extent are modifiable and functional groups can be added to gain additional functionalities. Researchers have also reported that graphene oxide-coated NMC532 does not possess a carbon matrix but a thin layer covering primary particles, allowing for the reduction of surface impurities due to the coating separating the core from the atmosphere [200].

Carbon Matrixes A novel method to enhance the performance of Ni-rich NMC-based cathodes is to build carbon matrices containing active material particles. Examples of this method include studies involving graphene nanosheets for NMC811, carbon nanotube conductive networks for NMC532 (Fig. 12d) [201, 202], and interwoven carbon fibers for NMC622 [203]. Here, the matrices chosen are usually highly conductive carbon materials that can enhance charge
transfer due to 3D connections. In addition, these matrixes can prevent side reactions and stabilize particle structures by fully covering primary cathode particles with a conductive matrix. Furthermore, carbon matrixes can enhance flexibility and strengthen adhesion between primary particles. Numerous unconventional and cutting-edge studies have also emerged in recent years involving the reshaping of aggregated NMC primary particles through the use of carbon matrixes. For example, multi-shelled alginate hollow fibers were used to template NMC primary particles, which allowed for the accommodation of Ni, Mn and Co atoms within an organic fiber matrix that possessed pinholes to ensure Li$^+$ exchange [204].

4.1.3 Advanced Preparation Methods

Aside from materials for NMC development, preparation methods are also being investigated due to similar influences on the cycling/thermal stability of Ni-rich NMC materials, commonly allowing for the reduction of cation mixing. For example, Du et al. [154] have reported that NMC532 sintered in oxygen can exhibit lower degrees of cation disordering, less oxygen defects and higher discharge capacities. These researchers also reported that if Ni$^{2+}$ was partially replaced by Li$^+$, the remaining Ni existed as Ni$^{3+}$, leading to capacity loss [154]. In addition, researchers have also reported that the atomization co-precipitation method can be used to obtain homogeneous spherical shaped primary NMC particles by forming an aerosol of precursors with an ultrasonic nebulizer [205] and that a similar effect can be achieved by using a self-combustion method, which produces massive heat flow to accelerate synthesis and therefore improve the uniformity of NMC particles [206]. Furthermore, nanoscale surface treatments utilizing Co and Li precursors combined with high-temperature sintering have been used to transform NMC622 surfaces into a Co-rich area with a NiO rocksalt phase that can act as a pillar layer to prevent inter-slab collapse [129]. Moreover, a hydrothermal procedure following the regular co-precipitation synthesis of NMC (70 15 15) can reportedly accelerate Li$^+$ entering and exiting Li slabs by making use of PVP to reduce crystal growth along the (001) plane and favor crystal growth along the (010) plane, which is an unimpeded pathway [207]. Dahn et al. [208] have also used a washing and reheating method to synthesize single crystal NMC622 in which superior performances as compared with multi-crystal NMC622 can be achieved if a remedial electrolyte additive was added. Carbonate co-precipitation and impregnation methods can also reportedly be combined to synthesize porous NMC622 microspheres that can effectively precipitate Ni$^{2+}$ and shorten Li diffusion pathways [209].

4.1.4 Synergy of Multiple Mitigation Methods

Recently, many studies have focused on the synergy of two or more mitigation methods to achieve better overall Ni-rich NMC-based cathode performances in which the simplest examples involve multiple dopants or coatings. For example, Al and Fe can be co-doped into NMC622 [131] to combine the polarization effects of Fe with the lower chemical potential attributed to Al. As for multiple coatings, this can be achieved with the help of organic coating layers, which can supplement the insufficiencies of single-type coatings. For example, polypyrrole is a flexible and conductive polymer that is typically used in this type of modification in which researchers have reported a dual-conductive coating layer of polypyrrole on NMC811 primary particle surfaces that can convert excessive Li into Li$_3$PO$_4$ [191]. In another study, researchers coated a Zr-doped NMC532 with polypyrrole to form a network that prevented HF attacks [157].

Apart from coatings and dopants, a newly developed trend is to combine surface modifications with gradient layers, which can prevent phase transitions of NMC811 and NCA particles [210]. In addition, the cathodic doping of Al into NMC (61 27 12) with FCG has been reported to be able to consolidate and reduce grain boundaries and improve mechanical properties [211]. Moreover, researchers have combined gradient layers (the core) with Mn-rich surface coatings on NMC532 and reported that the valences of Ni and Co can be kept at lower states than regular NMC532 particles. Similarly, NMC811 with an exterior concentration gradient layer was uniformly coated with Al$_2$O$_3$ through freeze-drying and displayed better capacity retention as compared with untreated samples [212]. Furthermore, LiPON as a conductive material used to coat anodes can also reportedly be applied to NMC811 with a surface gradient layer to maintain rate capability and improve capacity retention [213].

4.1.5 Summary

Despite novel synthesis methods for NMC particles (which usually require more demanding conditions and extra procedures), modifications are usually applied to different parts of Ni-rich NMC primary particles to protect Ni-rich NMC cores. Among these, dopants and coatings are the most extensively studied methods. Here, dopants can affect surface elemental compositions and lower the Ni content, which is reactive and can generate spinel phases during cycling. In addition, dopants can expand/contract LiNi$_x$Mn$_{1−x}$Co$_{1−x}$O$_2$ lattices and influence Li$^+$ movement through bulk particles. Furthermore, dopants possessing different valences from substituted ions can affect the valences of Ni, Mn and Co and therefore increase/decrease their capacity as the capacity is achieved by the change of valence of these elements.
Additional effects such as the repulsion of cation disorder and the prevention of HF corrosion can also be achieved through doping. As for coatings, these can impact NMC performances by separating surfaces from electrolytes and hindering side reactions in which optimal performances can be achieved through the accurate estimation of the amount of coating applied.

4.2 Graphite Anode

4.2.1 Testing Protocols

The rate of charge, as limited by the production of plated Li, can be increased by applying the constant-current constant-voltage (CCCV) protocol and the pulse charging protocol [105] in which Li concentration gradients are alternatingly enlarged and reduced during charging with the CCCV protocol and concentration differences between anode surface regions and bulk electrolytes are minimized in the pulse charging protocol.

4.2.2 Surface Coating

Aside from testing protocols, substantial barriers involving materials being applied to graphite surfaces through different means can fully separate graphite anodes from electrolytes, leading to significant reductions in the reductive decomposition of solvents, and subsequent conversion and thickening as well as the suppression of exfoliation. For example, LiPON as a recently investigated material can be evenly coated onto graphite anodes and can protect the anode from high-temperature storage aging [214].

4.2.3 Electrolyte Volume Adjustment

The performance of full cells is strongly dependent on the amount of accessible active material. And because both electrodes and the separator in full cells are porous, they need to be sufficiently wetted. In addition, electrolyte volumes can also be increased to maximize the amount of accessible active material; however, higher electrolyte volumes do not directly translate into higher capacities in which researchers have reported that excessive electrolytes tend to decrease performance because higher fractions of SEIs can dissolve into increased electrolyte volumes [215]. This principle is also valid in pouch cell studies showing the lowering of impedance and increase in cycling stability [216].

4.3 SEI

4.3.1 Electrolyte Additive

LIBs need to be safely transported and operated. Here, the addition of relatively small amounts of electrolyte additives can improve overall cell safety without the introduction of any detrimental products. For example, vinylene carbonate (VC) is one of the most extensively utilized electrolyte additives because it can improve compatibility between SEIs and graphite anodes by forming a thin film to separate electrolytes from graphite anodes and enhance the performance of ionic liquid-based electrolytes [128]. Researchers have also hypothesized that VC can reduce SEI thicknesses by inhibiting electron transfer from graphite anodes to solvents. Despite many positive influences of VC as reported by Dahn group [107], researchers have also suggested that the highly resistive film formed with VC can facilitate Li plating [217] and that the polymerization of VC can limit LIB lifespans. Here, many additives have been tested to replace VC to resolve polymerization issues in which FEC [139], PES [218] and VEC have been used to reinforce SEIs because they possess higher LUMOs that can prevent the reductive decomposition of VC by sacrificing themselves (Fig. 13a, b). In addition, FEC can form an SEI that is durable to fast charging, thus eliminating Li plating on graphite anodes [219]. Researchers have also reported that the use of FEC can suppress [139] or slow down [220] SEI formation (formation of lithium alkyl carbonate) to result in a thinner SEI. The use of co-solvents can also lower the viscosity and the melting point of solvents and therefore be useful in practical cases (Fig. 13c). For example, 2% PES with 1% DTD and 1% TTSPi (PES211) can enhance the cycling performance of NMC cells based on its composition [142]. Researchers have also reported that PTSi as a less investigated additive can generate better SEIs with less inorganic components in which due to the presence of S=O groups, PF5 can be suppressed as delocalized N cores can act as weak base sites [112]. And more recently, 3-sulfolene (3SF) as a film-forming agent was used to reduce electrolyte reduction [221]. Additives that stabilize SEI can be divided into two subgroups including SEI formers and SEI modifiers [222] and phenyl carbonate, regarded as a “modifier” additive, can outperform VC by maintaining low impedance and gas production in NMC111/graphite pouch cells and therefore can be used to stabilize cells [222]. In addition, phenyl carbonate can remain stable until 4.2 V and possess good compatibility with 2% vinylene carbonate (VC). And since electrolyte additives can be multi-functional because certain types of additives can react with electrodes to produce gas if cells are overcharged to unsafe voltages, which results in pressurized internal environments that can deactivate cells [222], the ability to reduce gas production makes phenyl...
carbonate a promising type of electrolyte additives. Furthermore, PBF and PPF as electrolyte additives have also been reported to be able to positively enhance SEIs [36].

### 4.3.2 Lithium Salt

A simple method to enhance the quality and durability of anodes and SEIs is to raise the concentration of Li salts to above 3 M [223], which can significantly reduce polarization that can lead to negative effects such as dendrite growth and slow charging. In addition, the resulting solvation of Li-ions possesses a disparate structure from regular Li-ions in which the planar shape of the solvated Li-ions benefits from Li intercalation and can mitigate graphite exfoliation from smaller spaces being occupied during operation [223]. Advanced Li salts have also been widely investigated to obtain better performances with promising results; however, few have been commercialized. In addition, a new group of Li salts (Fig. 14) have been reported to significantly enhance the performance of SEIs with capping agents such as oxalate in which the capping agents can reductively decompose electrolytes and prevent LiF from aggregating into large particles, thus producing LiF nanoparticles 5 nm in diameter uniformly in SEIs [224]. Furthermore, the reduction in LiF aggregation and the formation of nanostructured LiF can form uniform regions for Li⁺ diffusion and reduce lithium plating (Fig. 15). Researchers have also reported that LiTFSi can outperform conventional LiPF₆ salts due to its resistance to trace amounts of water and therefore minimize the production of harmful species [225].

### 4.3.3 Solvent

The polarity and the viscosity of solvents also need to be considered in the selection of appropriate solvents for Li intercalation/deintercalation in which solvents with lower viscosity can facilitate Li⁺ transportation and decrease impedance, whereas the polarity of a solvent can determine the degree of solvation of Li⁺ inside [225]. Here, ethyl methyl carbonate and dimethyl carbonate are usually

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Fig. 13  
(a) HOMO and LUMO energy levels of EC, FEC and VC.  
(b) Schematic showing the effects of VC- and FEC-derived SEIs on the fast charging capability of graphite anodes (Reprinted with permission from Ref. [219]. Copyright © 2018, Elsevier B.V. All rights reserved.).  
(c) Up: capacity versus cycle number for NMC111/graphite pouch cells (unclamped) containing select additives or additive blends. Cycling was conducted between 2.8 and 4.2 V at 80 mA. Bottom: capacity versus cycle number for NMC442/graphite pouch cells (unclamped) containing select additives or additive blends. Cycling was conducted between 3.0 and 4.4 V at 45 °C at 100 mA. (Reprinted with permission from Ref. [142]. Copyright © The Author(s) 2015. Published by ECS)
In addition, ionic liquids as substituents of conventional solvents can form thicker SEIs with high-capacity retentions at 90 °C [226].

**Summary and Prospective**

LIBs using Ni-rich NMC-based cathodes such as NMC811 can produce high specific capacities (200–220 mAh g⁻¹) and promising energy densities (~ 800 Wh kg⁻¹) in comparison with conventional LiCoO₂ (~ 570 Wh kg⁻¹) and LiMn₂O₄ spinel (~ 440 Wh kg⁻¹) materials. Because of this, these materials have received extensive attention from researchers. However, due to the low portions of manganese as structural stabilizers, large amounts of Ni⁴⁺ on cathode surface layers/regions can trigger side reactions and Ni²⁺ can cause cation mixing. As a result, enhanced mass-specific capacity comes at the expense of rate capability and structural stability in these Ni-rich cathode materials, leading to severe capacity fading. In addition, other factors such as active material dissolution, oxygen release and the intergranular cracking of primary particles also contribute toward cathode degradation. Alternatively, graphite anodes can function collaboratively with NMC-based cathodes to provide promising performances at cell levels. However, the aging of graphite anodes due to the formation of dendrites as caused by the massive plating of Li⁺ onto graphite surfaces remains the main reason for increased impedance. Furthermore, side reactions involving electrolyte reduction (SEI formation) on graphite anodes can cause Li loss after initial charging, and the thickening and decomposition of SEIs in subsequent cycles can cause aging behaviors.

To gain a deeper understanding of degradation mechanisms of Ni-rich NMC/graphite LIBs, emerging testing protocols and characterization methods have emerged in which currently, research on LIB degradation is focused on the elucidation of the degree of capacity fading in certain mechanisms so as to distinguish major factors and allow for the design of specific remedies. And as a result of this research, mitigation strategies, functioning individually or synergistically, have been proposed to resolve long-term cyclability issues, most of which have proven to be promising. Among these, dopants, gradient layers, coatings (both Li-free and Li-contained), carbon matrixes and advanced synthesis procedures have become mainstream methods to enhance the performance of Ni-rich NMC-based cathodes. In addition, coatings, advanced protocols and the use of new electrolyte components can be applied to ameliorate anodic performances (SEIs and graphite). And with these mitigation strategies, the morphology, element composition and reactivity of electrodes can be improved.

Although differences in the performance of various Ni-rich NMC-based cathodes have been discovered, these remain unexplained and the future of degradation mechanism investigations needs to rely on more advanced physical techniques such as X-ray tomography, operando neutron diffraction and online electrochemical mass spectroscopy, all of which can provide more comprehensive information in operando and be less destructive to cells. And for the deeper understanding of mitigation strategies for Ni-rich NMC/graphite LIBs, trade-offs in modification techniques and controversies need to be taken into considerations. Here, several prospective research directions for Ni-rich NMC/graphite LIBs are proposed:

1. Future experiments should focus more on the quantitative interpretation of different aging behaviors in NMC-based cathodes with or without mitigation methods as well as the understanding of correlations between different degradation mechanisms.
2. Efforts should be made toward profiling the performance of LIBs under harsher conditions (stressors) that more closely represent practical operational conditions.
3. Studies should be conducted to investigate and optimize the synergistic effects of multiple mitigation methods on Ni-rich NMC-based cathodes that can outperform the effects of single mitigation methods and enhance capacity retention to higher levels.

4. Inexpensive and nondestructive characterization tools need to be developed to decrease research costs and prevent cell destruction, especially in the inline detection of cells. In terms of EIS that is time-consuming, fast impedance needs to be developed to accelerate measurement processes.

5. Time-dependent EIS in combination with differential specific capacity measurements should be used as a characterization tool to analyze surface phases/structures versus impedance growth.

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Compliance with Ethical Standards

Conflict of interest The authors declare no conflicts of interest.

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