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Developing rapid electrochemical relithiation protocols for scalable relithiation of lithium-ion battery cathode materials

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The recent and ongoing boom in electric vehicle sales has caused the circularity of the supply chain for electric vehicle battery materials to come under a great deal of scrutiny. Innovative recycling processes, or direct recycling, that offer the possibility of reducing the cost of recycling are one possible solution to regaining resources from end-of-life (EoL) electric vehicle batteries. Electrochemically shuttling lithium back into the cathode, or electrochemical relithiation, is a possible technique for restoring lithium content to NMC materials (EoL) in a direct recycling process. This study provides essential understanding towards developing an electrochemical relithiation protocol that will restore lithium loss in intercalation cathode materials that reach EoL by loss of lithium inventory (LLI) as opposed to other degradation mechanisms like loss of active material (LAM), cation mixing or phase transition. Electrochemically aged NMC cathode materials have been prepared and characterized to establish the extent of EoL material structural degradation and lithium loss. A model-informed experimental process is used to identify the optimal electrochemical relithiation protocol to minimize the time taken to relithiate EoL materials and maximize the amount of lithium restored. Protocols were evaluated based on their ability to enable rapid lithium intercalation, maintain structural uniformity in the EoL material and fully restore lithium content. An optimal protocol was identified at elevated temperatures utilizing a novel scanning voltage step.

KEYWORDS

direct recycling, electrochemical relithiation, lithium-ion battery, advanced protocols, cathode

1 Introduction

Lithium-ion batteries (LIB) are widely utilized for applications such as electric vehicles, stationary storage and portable electronics. Since these LIB have a typical lifespan of 10–15 years, this means a large amount of end-of-life (EoL) batteries will need to be disposed of in the next few years. It is projected that the cumulative flow of spent LIB packs into the waste stream between 2015 and 2040 could be up to 21 million packs (Richa et al., 2014). Current collection and recycling rates for EoL LIBs are low and traditional disposal methods are environmentally questionable due to the hazardous materials used in LIBs. Proper handling of EoL LIBs through recycling could strengthen valuable resource supply chains and cut energy cost and consumption for electric vehicles manufacturing. Recently, direct recycling processes have garnered attention due to their potential to provide a more

efficient, economical alternative to commercialized recycling approaches such as pyrometallurgy and hydrometallurgy. The attraction of direct recycling lies with the ability to recover high value, engineered-form cathode materials without high temperature smelting or large amount of acids and alkaline chemicals that decompose the cathode materials to elemental products that then must be remanufactured (Gaines et al., 2021; Coyle et al., 2023).

Among the degradation mechanisms that need to be addressed to fully recover cathode materials in a direct recycling process is the irreversible loss of cyclable lithium in the cathode material through SEI formation. A variety of relithiation techniques have been proposed including ionothermal, hydrothermal, redox mediator relithiation, electrochemical relithiation and solid state relithiation (Nie et al., 2015; Gao et al., 2020; Wang et al., 2020; Zhang et al., 2020; Park et al., 2021). Electrochemical relithiation is attractive as a low temperature relithiation method that has option for scalability such as application in a roll-to-roll reactor. Using this technique would conceptually be simple for companies that coat their own electrodes to integrate into their own roll-to-roll systems. This relithiation process is also attractive as the relithiation reaction is controlled by electrode potential which eliminates the necessity to quantify the lithium deficiency prior to relithiation which is required by other relithiation techniques. A close analogue of electrochemical relithiation is the more widely studied electrochemical prelithiation method used for compensating large irreversible formation losses in next-generation anodes (e.g., silicon) based lithium-ion batteries (Jia et al., 2023). The prelithiation setup generally utilizes an external resistor or galvanostatic protocols to exchange lithium between a Li-rich source (Li metal/Li salt in electrolyte) and the target Li-deficient anode. In these approaches, controlled prelithiation is achieved in a Li metal half-cell configuration by tuning the external resistance and prelithiation time (external short) (Kim et al., 2016) or constant current magnitude until specified voltage cutoff (galvanostatic protocol) (Overhoff et al., 2021). Our work builds on the learnings from electrochemical anode prelithiation to provide guidelines for electrochemical relithiation of cathodes up to full lithium saturation.

One of the main technical challenges involved with developing this electrochemical relithiation method is the need for rapid electrochemical relithiation protocols. For this protocol to be viable in a scalable system such as a roll-to-roll reactor bath it is essential to minimize the time it takes to complete the electrochemical relithiation step, the temperature at which the reaction takes place and the specialized equipment required to complete the protocol. The relithiation process needs to restore cyclable lithium loss from both initial and continued electrochemical SEI growth. Electrochemical relithiation has been studied previously using a variety of techniques for direct recycling of cathodes, but not with direct attention to optimizing rapid relithiation protocols (Li et al., 2020; Yang et al., 2020; Zhang et al., 2020; Lahtinen et al., 2021; Liu et al., 2022; Raj et al., 2022). The challenge lies in the state of charge (SOC) dependence of material thermodynamic, transport and kinetic properties. NMC cathodes, in particular, exhibit a decrease in its Li solid phase diffusivity and exchange current density with lithiation (as discharge progresses); steep plunges in magnitudes are observed near full lithiation (Verma et al., 2017). Thus, kinetics and transport get increasingly sluggish as NMC lithiates resulting in higher overpotentials for the same applied

current densities. This behavior is unique to NMC cathode materials and is not generally observed for lithium iron phosphate (LFP) cathodes (Srinivasan and Newman, 2004). The most widely used discharge protocols like constant current (CC) or constant current constant voltage (CCCV) are rendered sub-optimal in providing a good balancing act of fast lithium insertion with ever-increasing kinetic and transport limitations near full NMC lithiation. Hence, electrochemical relithiation protocols need to include multistep currents of decreasing magnitudes or scanning voltage portions near high states of charge to avoid these exacerbated overpotentials arising from sluggish reaction kinetics and solid-phase transport. In previous works, we have leveraged such advanced protocols for enabling extreme fast charge (XFC) applications while avoiding the deleterious impact of lithium plating at the anode and diffusion-induced stress-based fracture in cathodes (Colclasure et al., 2020; Mai et al., 2020; Dufek et al., 2022). These protocols included elevated temperature charging, multistep current with step down/step ups in current rate, boost charging (2 step CC + CV), pulse charging (time-based periodic currents), voltage-controlled charging, etc. and were optimized for cathode delithiation/anode lithiation. A well parametrized Doyle-Fuller-Newman (DFN) model framework is amenable to screen prospective electrochemical protocols for optimal relithiation characteristics using state of charge and temperature dependent thermodynamic, kinetic and transport properties (Wang et al., 2022). Bearing these requirements in mind, we propose a novel approach for designing rapid electrochemical relithiation protocols for application in an initial batch reaction electrochemical bath relithiation system. Optimal relithiation protocols for any electrochemical relithiation application will likely need to be developed based on EoL material waste streams. For example, the cathode chemistry, degree of degradation and electrode loading that is undergoing relithiation will have an influence on the optimal protocol. For this reason, it is advantageous to develop a physics-based electrochemical model that can be used as a turnkey tool to design chemistry specific optimal protocols with the option to adjust protocol parameters such as varying current steps, voltage cut-offs and applied temperature. The objective of this work is to develop and validate model-informed fast discharge protocols using cycle aged NMC333. Maximizing the charge transferred to the EoL material within 30 min or less while minimizing degradation modes such as oversaturation or mechanical fatigue are key constraints to allow this work to be transferred to an electrochemical bath system.

In this work, we devise a Li half-cell electrochemical relithiation protocol for rapid restoration of harvested NMC333 cathodes from end-of-life cells to its pristine state. The cycled cathodes have ~20% capacity degradation primarily due to loss of lithium inventory (LLI) to the solid electrolyte phase (SEI) with minor impact of loss of active material (LAM) due to cathode cracking. Consequently, we investigate the possibility of fast relithiation (<30 min) of the cathode from its harvested state of charge (SOC ~ 0.80) to full lithiation (SOC ~ 1) using novel electrochemical protocols. Here, SOC is the electrode-level state of charge (not to be confused with cell level state of charge) defined as the current amount of lithium in the cathode divided by the maximum amount of lithium that can intercalate into the cathode. Li| NMC333 half-cell electrochemical testing is performed at multiple temperatures with a parametrized

sample set of multistep protocols including constant current and scanning voltage steps to identify the optimal electrochemical relithiation protocol. Degraded NMC material properties are obtained from controlled half-cell testing and model validation with experimental data. Finally, an electrochemical protocol comprising of constant current step to an intermediate voltage cutoff supplemented with scanning voltage step to final voltage cutoff is obtained through optimization of the fully parametrized half-cell DFN model for rapid discharge with approximately 100% of expected lost capacity restored to the cathode.

2 Experimental and computational methods

2.1 Electrochemically aged cathode material

Accelerated full cell coin cell cycling was used to source electrochemically aged NMC 333 material. Composite cathodes consisted of pristine Toda NMC 333 cathode powder, Timcal C45 conductive agent and Solvay 5130 polyvinylidene fluoride (PVDF) with a 90:5:5 mass ratio. Composite anodes had a composition of 91.83 wt% Superior graphite, 2 wt% Timcal C45 carbon, 6 wt% Kureha 9300 PVDF binder and 0.17 wt% oxalic acid. The areal loadings for that cathodes and anodes were 1.62 and 1.92 mAh/cm² respectively. Celgard 2320 was used for a separator along with 25 μ L of 1.2 M LiPF₆ EC:EMC 3:7 wt% electrolyte. Full cells were aged until 20% of the initial charge capacity was lost by cycling at C/10 for four formation cycles followed by 1C/1C cycling between 3.0 and 4.3 V for about 135 cycles at room temperature where 1C was the rate at which the cathode would be fully discharged in 1 h based on the theoretical capacity of the cathode active material.

2.2 Electrochemical relithiation protocol validation

Electrochemically aged full cells were discharged to 3.0 V at 1C and held at 3.0 V until the current decayed below C/20. The full cells were then disassembled and the electrochemically delithiated NMC 333 cathode was rinsed in dimethyl carbonate (DMC) by first gently rinsing in fresh DMC, removing all the DMC and adding fresh DMC and allowing the cathode to soak overnight to attempt to remove all electrolyte and salts. After disassembling and rinsing the electrochemically delithiated samples, they were rebuilt into half cells flooded with 100 μ L 1.2 M LiPF₆ EC:EMC 3:7 wt% electrolyte for relithiation protocol testing.

2.3 Material characterization

X-ray powder diffraction (XRD) patterns of cathode electrodes were obtained by employing Cu K α radiation using a Rigaku MiniFlex diffractometer. Crystal structures were examined using high resolution scans conducted from 10° to 90° 2 θ (0.01° step size, 10 s dwell time). Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES) was completed on a Perkin Elmer

Optima 4300 DV. X-ray photoelectron spectroscopy (XPS) analysis was conducted with a PHI 5600 XPS with Al K α X-ray excitation with a base pressure for the system of 1.4 X 10⁻⁹ Torr. The spectra curve fitting analysis was processed using CasaXPS software. Peak fitting parameters for XPS analysis were minimization of the standard deviation of the residuals along with ensuring the final fit was chemically reasonable based on known chemical moieties. Scanning electron microscopy (SEM) images were collected on a Hitachi S-4800 microscope.

2.4 Computational methods

The Doyle-Fuller-Newman (DFN) model (also known as pseudo 2D model) capable of predicting performance characteristics with half-cell discharge is utilized to ascertain the optimal relithiation protocol for the cycled cathode material (Doyle et al., 1993; Chen et al., 2020). Our system of choice comprises of Li metal anode and NMC333 [Li(Ni_{1/3}Mn_{1/3}Co_{1/3})O₂] cathode. The governing equations are summarized in Eqs 1–8 and have been described in detail in previous works (Ji et al., 2013; Colclasure et al., 2020). A short description is provided here. Butler-Volmer kinetics with symmetric charge transfer coefficients is used to describe the lithium intercalation reaction at the cathode (see Eqs 1–3 while Li deposition kinetics is assumed to be rapid such that kinetic overpotentials at the anode are negligible.

Charge transfer kinetics:

$$i = i_0 \left[\exp\left(\frac{0.5F}{RT}\eta\right) - \exp\left(-\frac{0.5F}{RT}\eta\right) \right] \quad (1)$$

$$\eta = \phi_s - \phi_e - U(\text{SOC}, T) \quad (2)$$

$$i_0 = k(T)c_e^{0.5}c_{s,\max}(\theta(1-\theta))^{0.5}; \text{SOC} = c_{s,\text{surf}}/c_{s,\max} \quad (3)$$

Here, i is area-specific current density, i_0 is intercalation exchange current density, η is kinetic overpotential and T is temperature. F , R corresponds to Faraday's constant (96485.33 C/mol) and universal gas constant (8.314 J/mol-K) respectively. Subscripts s and e belong to solid phase and electrolyte phase respectively. Kinetic overpotential is determined from the solid phase potential ϕ_s , solution phase potential ϕ_e , and the open circuit potential (OCP) of the Li intercalation reaction into cathode material which is a function of lithium state of charge of NMC333 particles, (SOC= 0: completely delithiated state, SOC= 1: fully lithiated state). Intercalation exchange current density is dependent on the reaction rate constant, k , electrolyte salt concentration, c_e , maximum Li content in NMC333, $c_{s,\max}$, and state of charge. It is evident that the kinetic limitations are dictated by state of charge and temperature dependent OCP and i_0 ; consequently, it is important to accurately determine these thermodynamic and kinetic properties respectively for high model fidelity.

Lithium conservation in active material:

$$\frac{\partial c_s}{\partial t} = \frac{1}{r^2} \frac{\partial}{\partial r} \left(D_s r^2 \frac{\partial c_s}{\partial r} \right) \quad (4)$$

Fickian diffusion is assumed to transport Li into spherical active material particles. Here, c_s is the Li concentration in NMC333 particles that can go from 0 to $c_{s,\max}$ and is a function

TABLE 1 Cycled NMC333 cathode design parameters.

Active material, AM	90 wt% TODA NMC333
Conductive additive, CA	5 wt% TIMCAL C45
Binder, B	5 wt% Solvay 5130 PVDF
Al foil thickness, L_{Al}	20 μm
Total electrode thickness, L_{tot}	61 μm
Coating thickness, L_{cat}	41 μm
Porosity, ϵ	33.5%
Coating loading, m_{coat}	11.22 mg/cm ²
Coating density, ρ_{coat}	2.74 g/cm ³
Areal capacity, Q_{csa}	1.62 mA h/cm ² @C/10 between 3.0 and 4.3 V
Specific capacity, Q_{sp}	161 mA h/g @C/10 between 3.0 and 4.3 V
Separator	Celgard 2325
Electrolyte	Gen 2: 1.2 M LiPF ₆ in EC:EMC 3:7 (w/w), Tomiyama Japan
Electrode punch diameter	14 mm
Current at 1C	2.5 mA

of particle radius, r , and time, t . An important transport property determining the concentration gradients and surface concentration in particles is the Li diffusivity, D_s , which is a function of both state of charge and temperature.

Lithium-ion conservation in electrolyte phase:

$$\epsilon \frac{\partial c_e}{\partial t} = \nabla \cdot (D_e^{eff} \nabla c_e) - \frac{i_e \nabla t_+}{F} + \frac{j(1-t_+)}{F} \quad (5)$$

$$i_e = -(\kappa^{eff} \nabla \phi_e + \kappa_D^{eff} \nabla \ln c_e); j = a_s i$$

Diffusion via concentration gradients, migration via potential gradients and Li-ion production in the electrolyte phase from charge transfer chemistry determines the salt concentration, c_e , in solution phase. Here, ϵ is cathode porosity, D_e^{eff} is effective electrolyte diffusivity dependent on the porosity and tortuosity of the pore phase, t_+ is transference number and j is volumetric current density dependent on the specific surface area, a_s , and area-specific current density, i .

Charge conservation in solid phase:

$$\nabla \cdot (\sigma_s^{eff} \nabla \phi_s) - j = 0 \quad (6)$$

Here, Ohm's law is used to describe charge transport through solid phase potential gradients dependent on the electronic conductivity of the composite electrode, σ_s^{eff} .

Charge conservation in electrolyte phase:

$$\nabla \cdot (\kappa^{eff} \nabla \phi_e) + \nabla \cdot (\kappa_D^{eff} \nabla \ln c_e) + j = 0 \quad (7)$$

$$\kappa_D^{eff} = \frac{2RT\kappa^{eff}}{F} (t_+ - 1) \left(1 + \frac{d \ln f_{\pm}}{d \ln c_e} \right) \quad (8)$$

Finally, conduction and diffusion contribute to charge transfer through electrolyte. Here, κ^{eff} is ionic conductivity, κ_D^{eff} is diffusional conductivity, and f_{\pm} is the activity coefficient determining the non-ideality behavior of the electrolyte.

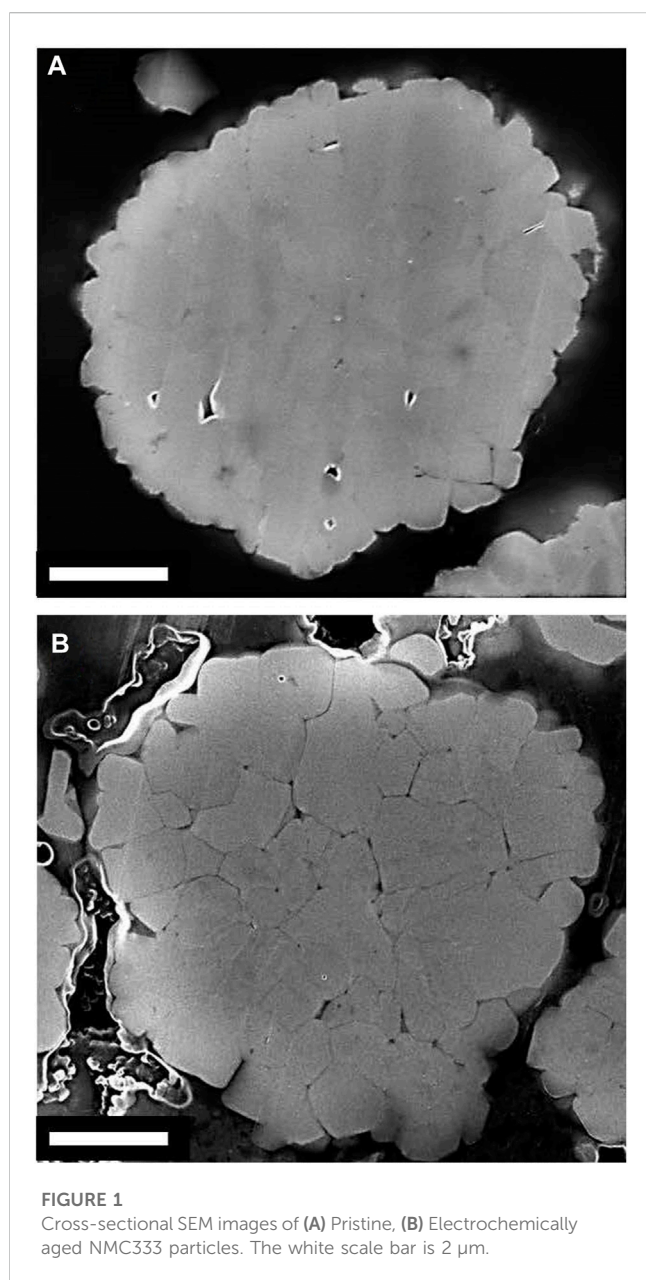
TABLE 2 Active material properties.

Molecular formula	LiNi _{1/3} Mn _{1/3} Co _{1/3} O ₂
Molar mass, MM	96.7947 g/mol
Theoretical capacity, Q_{th}	276.89 mAh/g
Maximum Li concentration, $c_{s,max}$	49500 mol/m ³
Mean particle size, R	10 μm
Initial state of charge, y_i	0.75–0.8
Active material volume fraction, ϵ_{AM}	0.5166
Active material mass, m_{AM}	15.54 mg
Composite electrode conductivity, σ_s^{eff}	10 S/m
Open circuit potential, OCP	see Figure 1A
Exchange current density, i_0	see Figure 4B, Colclasure et al. (2020)
Activation energy for i_0	30 kJ/mol
Diffusivity, D_s	see Figure 4B, Mistry et al. (2021)
Activation energy for D_s	30 kJ/mol

For low current rate operation ($\leq 1C$), properties OCP , i_0 and D_s dictate the performance metrics while σ_s^{eff} and electrolyte properties, D_e^{eff} , κ^{eff} , t_+ , κ_D^{eff} matter at high current rates ($>1C$) (Guo et al., 2010). Electrolyte properties for 1.2 M LiPF₆ in EC:EMC 3:7 (w/w) (also termed Gen 2 electrolyte) have been computed at varying concentrations and temperatures in previous articles and the correlations are directly used here (Colclasure et al., 2020). Consequently, we determine state of charge and temperature dependent OCP , i_0 and D_s for NMC333 through experimental analysis and parametrization to cycled NMC rate performance data. Tables 1–3 list the cycled NMC333 cathode design

TABLE 3 Separator and electrolyte properties.

Separator thickness, L_s	25 μm (Malifarge et al., 2017)
Separator porosity, ε_s	39% Malifarge et al. (2017)
Ionic Conductivity, κ	$f(c_e, T)$ Colclasure et al. (2020)
Transference number, t_+	$f(c_e, T)$ Colclasure et al. (2020)
Diffusivity, D_e	$f(c_e, T)$ Colclasure et al. (2020)
Activity coefficient, $\frac{d \ln f_{\pm}}{d \ln c_e}$	$f(c_e, T)$ Colclasure et al. (2020)
Initial salt concentration, $c_{e, \text{init}}$	1,200 mol/m ³



parameters, and properties of active material, separator, and electrolyte. Design parameters are obtained from the cycled electrode specifications, while active material properties are

computed through physical and electrochemical characterization wherever possible. Separator properties and concentration, temperature dependent functional relationships for electrolyte transport are obtained from literature (Malifarge et al., 2017; Colclasure et al., 2020).

3 Results and discussion

The aim of this work was to develop and validate model-driven electrochemical relithiation protocols using a cathode material that was cycle aged in a controlled manner that avoided potential complicating factors such as mechanical degradation of the electrode or particle fracture to isolate LLI as the primary cause of capacity loss. To verify that the LLI was the main cause of capacity loss for the NMC cathode materials, thorough characterization of the surface chemistry, morphology and crystal structure of the cycled NMC 333 was compared to pristine NMC 333. Figure 1 shows SEM cross-sectional image of each type of NMC 333 particle in this study. Several SEM images of the cycle aged NMC 333 were compared to confirm that no advanced particle breakdown occurred during cycling. There is no obvious significant source of mechanical degradation in the cycle aged particles that could have a significant contribution to the capacity loss. The overall morphology and size of the pristine and cycled aged NMC 333 particles was similar. XRD was performed to characterize how cycle aging affected the crystal structure of the NMC 333 and is shown in Figure 2.

The typical diffraction peaks associated with the α -NaFeO₂ structure highlighted in Figure 2A are consistent across pristine and cycle aged samples indicating that the bulk structure of each type of NMC 333 is not affected by the delithiation method (Shaju et al., 2002). The (003) peaks are highlighted in Figure 2B to show that due to the Li⁺ deficiency inherent to the cycle aged sample, the (003) peak shifted to a lower angle. This reflects an increase in the c lattice constant due to the increased electrostatic repulsion between the oxygen layers along the c direction with decreased Li⁺ (Mohanty et al., 2013; Liu et al., 2016). The (108) and (110) peaks also shows some shifting in the split of the doublet for cycle aged materials compared to the Pristine NMC 333 peaks. The (110) peak shifting to higher angles is responsible for split change in the (108)/(110) and is caused by the a-axis contracting due to the radius of transition metal ions shrinking at elevated valences in a charged state (Yang and Xia, 2016). There is no evidence of the highlighted peaks fading or widening so phase transformation in the bulk of the cycle aged NMC 333 material is likely not the cause of the observed full cell capacity loss. Figures 3A, C show the O1s XPS spectra for cycle aged and pristine NMC 333. The peak at 529.5 eV is assigned to the metal-oxide bond of NMC 333 and is consistent across both materials. The cycle aged sample in Figure 3A shows an increase in C-O species in the 531–534 eV range of peaks likely originating from electrolyte decomposition products, but otherwise the surface carbonate peaks in the O1s spectra for all three samples were similar. Figures 3B, D show the C1s XPS spectra for cycle aged and pristine NMC 333. The C 1s peaks around 290 eV are associated with the FC-OH and CF₂ bonds in the PVDF binder. The peak at 286 eV is also associated with PVDF CH₂-CF₂ groups (Duca et al., 1998). In general, the C1s peaks are dominated by those representing PVDF which remain constant for each type of NMC 333 in this study. The only C1s peaks

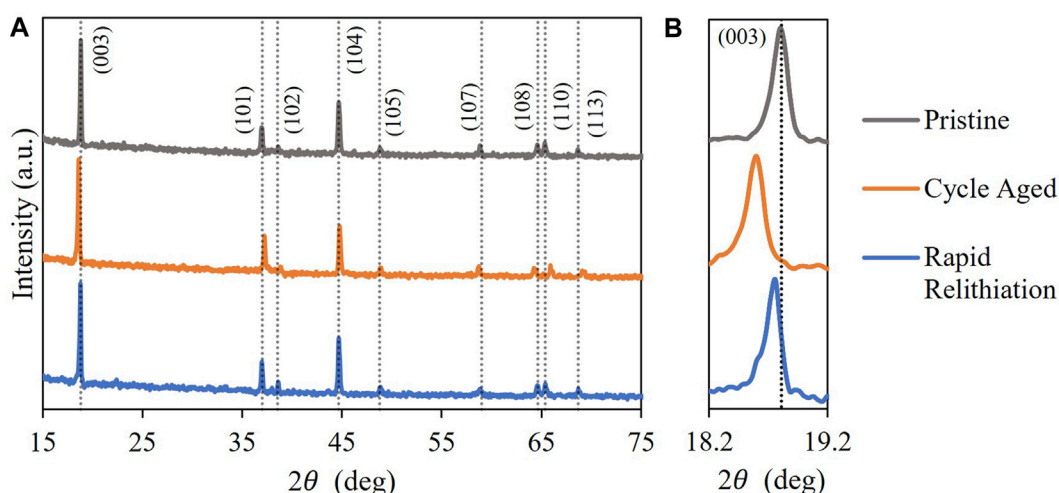


FIGURE 2
XRD analysis for NMC 333 materials with (A) all peaks, (B) (003) peaks.

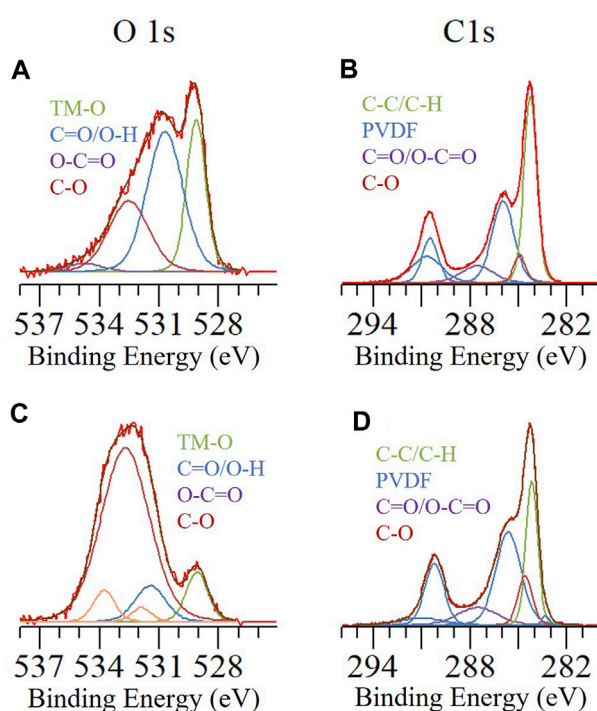


FIGURE 3
O1s and C1s XPS spectra for (A,B) Cycle aged and (C,D) Pristine NMC 333.

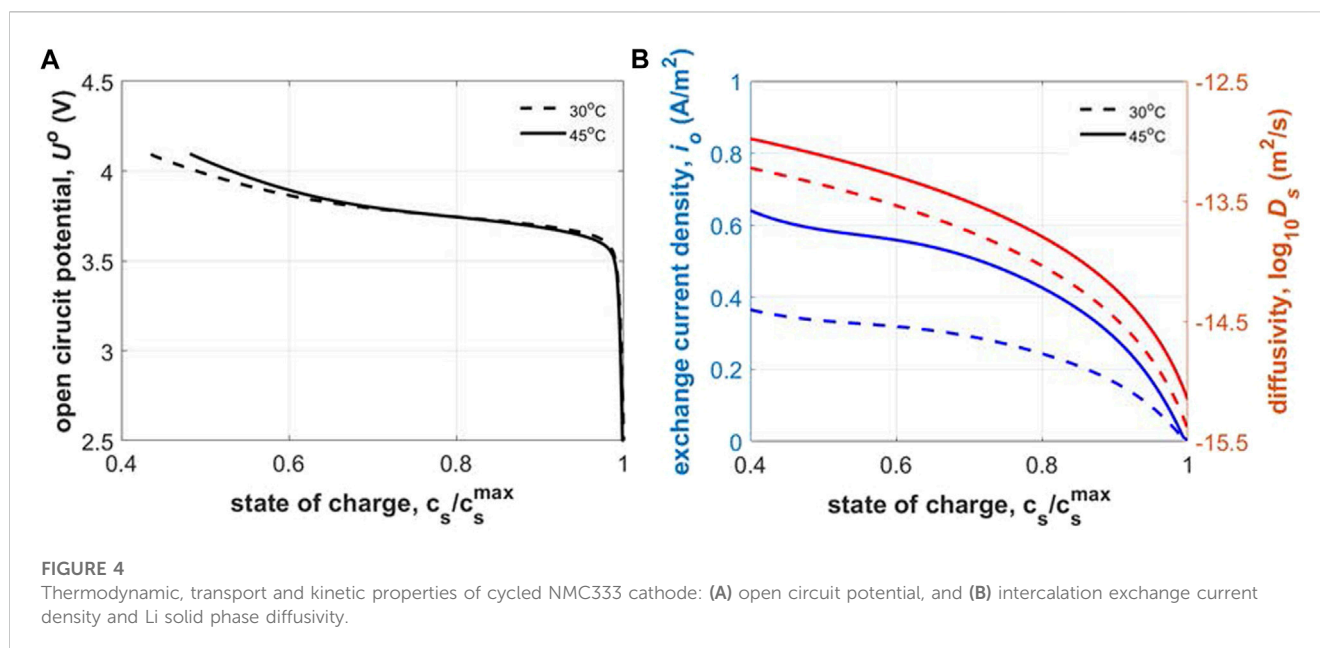
that change between materials is some broadening of the C-O peak at 284.8 eV and the C=O/O-C=O peak at 288 eV for the cycle aged NMC 333 sample in Figure 3D, likely due to leftover carbonate components that were not removed by rinsing after cycling also observed in the O1s spectra. XPS of both samples was similar which indicates that there was no major reduction of the transition metals at the surface of the cycle aged sample. This conclusion is also supported by the lack of peak shifting in the Co2p, Ni2p and Mn2p

TABLE 4 Molar ratio of relevant components in cathode materials determined through ICP-OES.

Cathode material	Li	Ni	Mn	Co
Pristine NMC 333	1.070	0.336	0.336	0.327
Electrochemically aged NMC 333	0.855	0.338	0.329	0.333
Electrochemically relithiated NMC 333	1.064	0.335	0.334	0.331

XPS spectra shown in Supplementary Figure S1. The lithium loss for the cycle aged sample was quantified using ICP-OES and is shown in Table 4. These results all indicate the LLI is the main capacity fade mechanism for the cycle aged NMC333 materials.

Figure 4A shows the open circuit potential for the cycled NMC333 cathode material as a function of state of charge. Slow rate cycling of Li|NMC333 half-cell at C/20 was performed between 2.5 and 4.1 V at 30°C and 45°C to obtain the thermodynamic open circuit potential (see Figure 4A). At such slow rates, transport and kinetic limitations are negligible, consequently, the cell voltage can be directly correlated to the thermodynamic OCP of the cycled NMC333 material. Cathode is assumed to be fully lithiated at cell voltage of 2.5 V; subsequently, the capacity exchanged during half-cell discharge can be correlated to the cathode lithiation state of charge. OCPs for 30°C and 45°C overlap considerably with slight variation in the lithiation range $0.4 < \text{SOC} < 0.6$ arising from entropic effects. Exchange current density and Li solid phase diffusivity variation with state of charge for the cycled material at varying temperatures is shown in Figure 4B. Verma and coworkers (Verma et al., 2017) have obtained the lithiation dependent exchange current density for pristine TODA NMC532 through galvanostatic intermittent titration test (GITT) which has been validated with extensive rate performance data by Colclasure et al. (Colclasure et al., 2020). We use this function multiplied by a factor of 0.75 utilizing the fact that NMC cathode properties improve with higher Ni content ($333 < 532 < 622 < 811$) (Noh et al., 2013) and

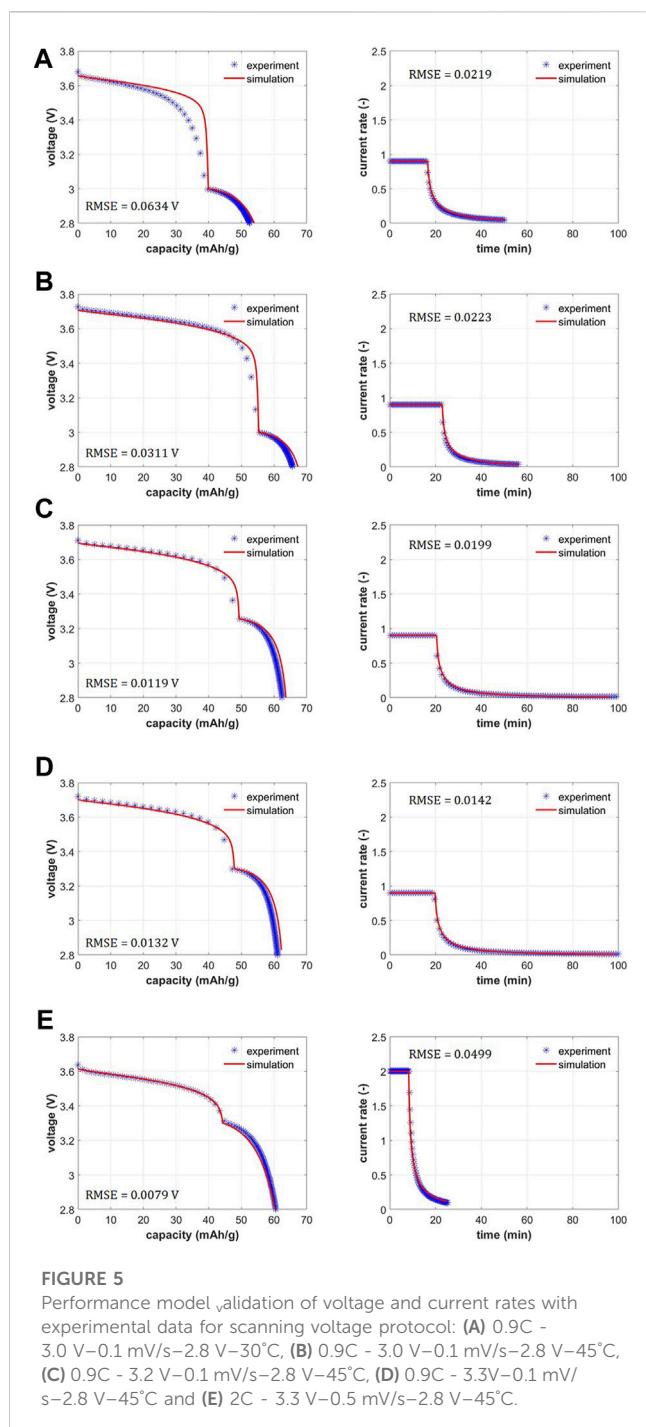


degraded material performs worse than pristine; so degraded NMC333 should showcase inferior kinetics than pristine NMC532. Furthermore, the state of charge dependence of properties amongst NMC family of cathodes is very similar, so multiplying NMC532's property with a fraction can capture the SOC dependence of NMC333 property well (Noh et al., 2013). Mistry and coworkers (Mistry et al., 2021) have identified the diffusivity function for pristine TODA NMC333 through curve fit to experimental data reported in Wu et al. (Wu et al., 2012). This pristine diffusivity is multiplied by a factor of 0.5 to obtain the degraded material's diffusion coefficient. The exchange current density at 30°C drops monotonically from ~ 0.4 to 0 A/m² across the lithiation range $0.4 < \text{SOC} < 1$. Kinetics gets enhanced with thermal effects according to the Arrhenius correlation, consequently, we see a bump in the exchange current density up to ~ 0.6 A/m² at 45°C. Diffusivity also shows a monotonic decline from $\sim 10^{-13}$ m²/s to $\sim 10^{-15}$ m²/s with lithiation ($0.4 < \text{SOC} < 1$) for both temperatures with a slight enhancement of diffusivity at 45°C. The factor magnitudes of 0.5 and 0.75 for diffusivity and exchange current density respectively provide best validation between the model and experiments with the least root mean square error (see Figure 5). Interestingly, both exchange current density and diffusivity show steeper declines close to full lithiation ($\sim \text{SOC} = 1$). This sharp drop has an impact on performance. During half-cell discharge, a constant current (CC) relithiation protocol hits the lower voltage cutoff due to exacerbated kinetic, transport limitations without the cathode reaching full lithiation. The usage of a constant current constant voltage (CCCV) discharge can achieve full lithiation but sacrifices the time optimization aspect; the protocol must run for hours before full lithiation is achieved. Hence, we need a novel electrochemical protocol to circumvent the issues with CC or CCCV protocol.

The limitations of CC or CCCV electrochemical protocol can be eliminated through usage of novel electrochemical protocols including multistep currents and scanning voltage steps (Mai et al., 2020). A multistep protocol with stepwise reduction in

currents accounting for the increasing sluggishness of kinetics, transport can sustain the discharge to full cathode lithiation while optimizing for time since there are no slow CV steps. However, application of the multistep current protocol for electrochemical testing is cumbersome; there is a current rate and voltage/time limit that needs to be defined for each step. The electrochemical protocol parameters can balloon for number of steps ≥ 3 . Furthermore, stepwise current profiles do not mimic the smoothly declining kinetics and Li diffusion profiles (see Figure 4B). An ideal electrochemical protocol should be able to supply gradually lowering currents as lithiation progresses. Thus, merging of constant current step with scanning voltage step can provide an alternative pathway towards fast electrochemical relithiation and is the subject of all our subsequent explorations here. The protocol starts with a constant current discharge to an intermediate voltage cutoff from the initial state. Once the voltage cutoff is reached, the protocol implements a scanning voltage discharge from the intermediate voltage cutoff to the lower voltage cutoff. The nomenclature of constant current scanning voltage protocols is defined as follow: a C - b V - c mV/s - d V - T °C. Here, a is the C-rate of the constant current step to intermediate voltage cutoff of b V. Subsequently, the voltage control step starts with a scan rate (SR) of c mV/s to lower voltage cutoff of d V. The entire protocol is run at T °C. It is advisable to run the experiment at the highest allowable temperature below electrolyte decomposition limits as high temperatures enhance the transport and kinetic properties. In our experiments with no additives, 45°C is the preferred temperature. It is notable that the constant current scanning voltage protocol needs only four parameters to be defined in the testing once the temperature is fixed.

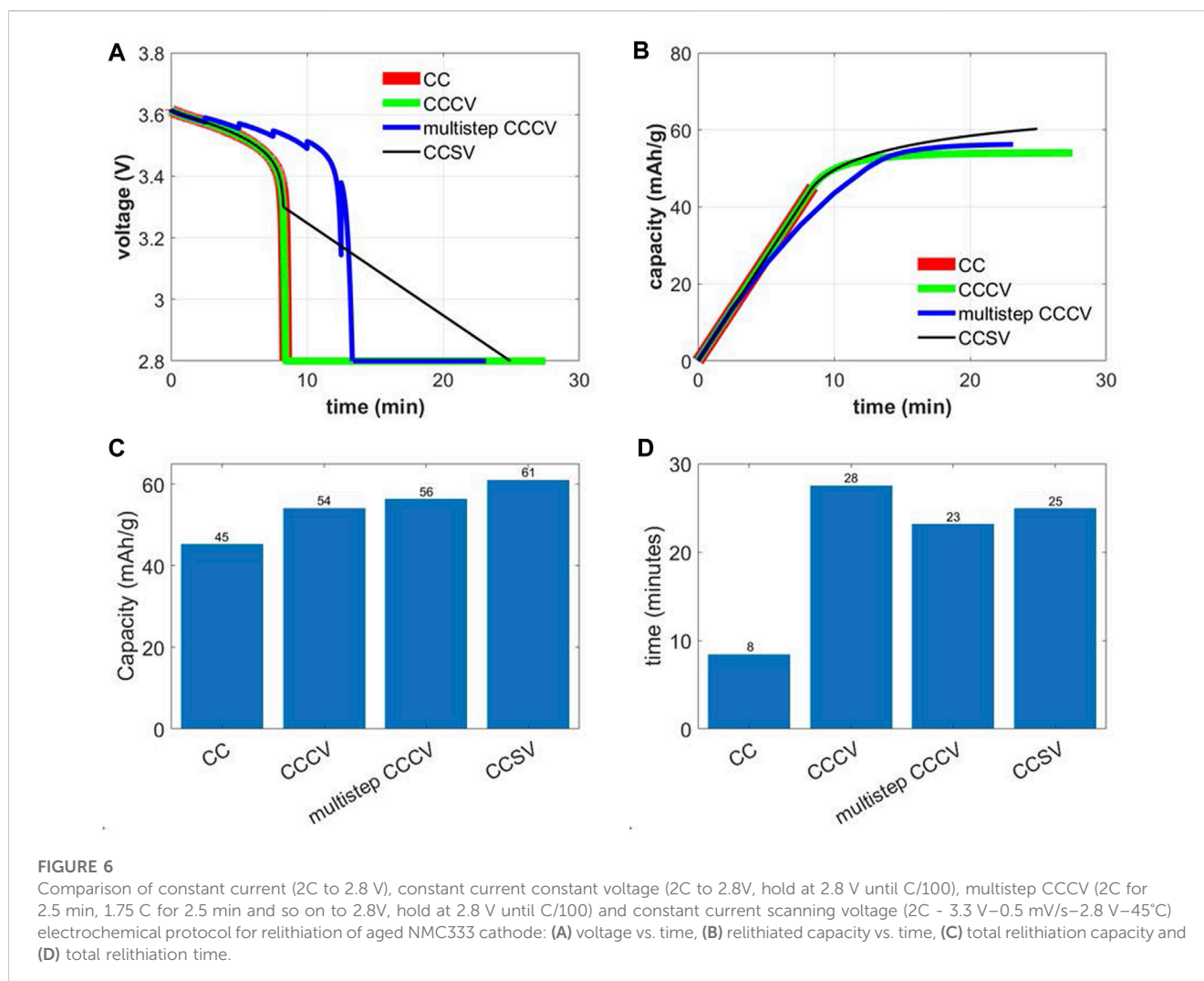
Figures 5A–E showcase the experimental performance plots for variations of the scanning voltage protocol alongside the model validation. Experimental dataset for voltage vs. capacity is shown in Supplementary Figure S2 for all cases (with replicates) while Supplementary Figure S4 shows the average and standard deviation in the voltage and current profiles. Model is validated



against the averaged experimental data. Voltage vs. specific capacity as well as current vs. time are validated between the experiments and the model. Low root mean square errors (RMSE) illuminate the high fidelity of the match. In Li-ion battery modeling literature, traditionally CC or CCCV models are used, and the validation is only shown for voltage vs. capacity (Ji et al., 2013; Verma et al., 2017). Here, we highlight the fact that in addition to the voltage - capacity data, the current rate - time data is also matched. Initial voltages lie around 3.6–3.7 V for the assembled half-cells with cycled NMC333 cathodes. This corresponds to Li state of charge in the cycled cathodes of around 0.80 which is close to the ratio of Li

content in electrochemically aged sample versus pristine sample (0.855/1.07 ~ 0.80). The first protocol variation is run at 30°C and constant current at 0.9C to 3.0 V with scanning voltage at 0.1 mV/s to 2.8 V (see Figure 5A). The relithiation time and capacity are 50 min and 52.62 mAh/g respectively. Subsequently, the relithiation temperature is increased to 45°C to leverage the enhanced transport and kinetics with same current and voltage steps (see Figure 5B). Relithiation time and capacity for this protocol are 56 min and 65.68 mAh/g respectively. Higher capacity and time duration indicate extra capacity is accessed due to the improved exchange current density and Li diffusivity. For the next variation (see Figure 5C), the scanning voltage step starts at 3.2 V instead of 3.0 V keeping other protocol parameters same as in Figure 5B. This increases the relithiation time considerably to 103 min while the relithiation capacity remains around the same ballpark (62.75 mAh/g). Evidently, earlier introduction of the scanning voltage step in the protocol from 3.0 to 3.2 V increases the time duration of the scan step and consequently the total time duration considerably (103 min vs. 56 min) while keeping the capacities similar. Changing the scan step start to 3.3 V (see Figure 5D, we see similar relithiation times and capacity of 103 min and 61.06 mA h/g. Finally, we increase the current rates to 2 C from 0.9 C and scan rate to 0.5 mV/s from 0.1 mV/s to see the optimal relithiation time of 25 min and relithiation capacity of 60.53 mAh/g which brings the cathode to near full lithiation.

To ascertain the efficacy of the constant current scanning voltage (CCSV) protocol, the performance model outputs of capacity and relithiation time are compared against baseline constant current (CC), constant current constant voltage protocol (CCCV) and a novel multistep constant current constant voltage protocol (multistep CCCV). We compare the results against the best scanning voltage validation case of 2C - 3.3 V-0.5 mV/s-2.8 V-45°C. The CC protocol consist of 2C current up to 2.8 V with the CCCV protocol comprising of an additional constant voltage step at 2.8 V until the current rate drops to C/100. Multistep CCCV protocol consists of several CC steps starting at 2C current rate. Each step has a duration of 2.5 min and a step down in current rate of 0.25C is prescribed at the end of each step. Once the voltage reaches the 2.8 V cutoff, an additional constant voltage step until the current rate drops to C/100 is also prescribed. Figures 6A, B shows the evolution of voltage and capacity with time for the CC, CCCV, multistep CCCV and CCSV protocols. CC, CCCV and CCSV protocols show smooth voltage decrease in the CC step while the multistep CCCV protocol shows the jagged voltage rise at end of each CC step from current step downs leading to lowered kinetic and transport overpotentials. Capacity increase with time for each protocol is relatively smooth with a change in slope observed at instants where the protocol switches to constant voltage, current step downs or scanning voltage steps. An interesting feature is that the slope of the constant voltage step in CCCV or multistep CCCV is smaller as compared to the scanning voltage step in CCSV. This is indicative of ability of the scanning voltage step to provide higher currents as compared to the constant voltage step. Total relithiation capacity and protocol runtime are shown in Figures 6C, D with the CCSV outperforming CC, CCCV and multistep CCCV in Li capacity restored. CC and multistep CCCV do show lower relithiation times as compared to CCSV but are only able to restore 73% and 92% of the total capacity lithiated in CCSV.



Furthermore, multistep CCCV requires delineation of additional parameters like constant current step duration and constant current step-down magnitude, so it brings the number of parameters for protocol specification to 5 which is the same as for CCSV protocol. Consequently, it is advisable to use the CCSV protocol for cathode relithiation.

In an initial scoping of electrochemical protocol choice, CCCV and CCSV protocol were run on ~25% capacity degraded NMC333 electrode with the experimental data for voltage, current vs. capacity and capacity vs. time shown in Figure S. CCCV protocol used 0.2C to 2.8V, hold at 2.8 V until current decays to C/100, while CCSV protocol parameters were 1.8C-3.0V-0.1 mV/s-2.8V-30°C. This data also afforded a preliminary indication into the efficacy of the scanning voltage step in reducing lithiation time. Approximately 19 mAh/g of lithiation was observed in the scan step from 3.0 to 2.8 V in 35 min for the CCSV protocol as opposed to 9.5 mAh/g of lithiation in 113 min in the constant voltage step at 2.8 V for the CCCV protocol. Thus, CV lithiation can be very sluggish.

Development of the high-fidelity performance model allows for running a parametric sweep of the constant current scanning voltage protocol to identify the best relithiation strategy. The parameters of

interest are initial current rate, intermediate voltage cutoff and scan rate with temperature and final voltage cutoff fixed to 45°C and 2.8 V respectively. Figures 7A-D shows contour plots of relithiation time (left) and recovered capacity (right) for intermediate voltage cutoffs of 3.0-3.3 V as a function of the initial current rate and scan rate. For the full sweep, the C-rate is varied from 1C to 3C in increments of 0.5C while the scan rate is changed from 0.1 mV/s to 0.5 mV/s in increments of 0.1 mV/s. Regions of low relithiation time ($t < 30$ min) and high capacity ($Q > 60$ mA h/g) are demarcated by black lines in the contour plots. It is evident that the desirable current rate and scan rate zones in the time contour is nearly opposite to the optimal zones in the capacity plot. An intersection of the desirable zones in both contours marks the best relithiation protocol strategy. Based on this, a couple optimal protocols are highlighted using star symbols in each intermediate voltage cutoff parametric sweep. Figure 7A showcases that the protocols 1.5C-3.0V-0.2 mV/s-2.8V - 45°C and 2C-3.0V-0.2 mV/s-2.8V - 45°C can achieve relithiation times ~30 min and capacity ~60 mAh/g. The desirable time zone is big for 3.0 V intermediate cutoff because of the less time spent in the slow scanning voltage step. Consequently, the desirable capacity zone is smaller because of less time for relithiation as well. The optimal time zones shrink while the capacity zones grow as we move

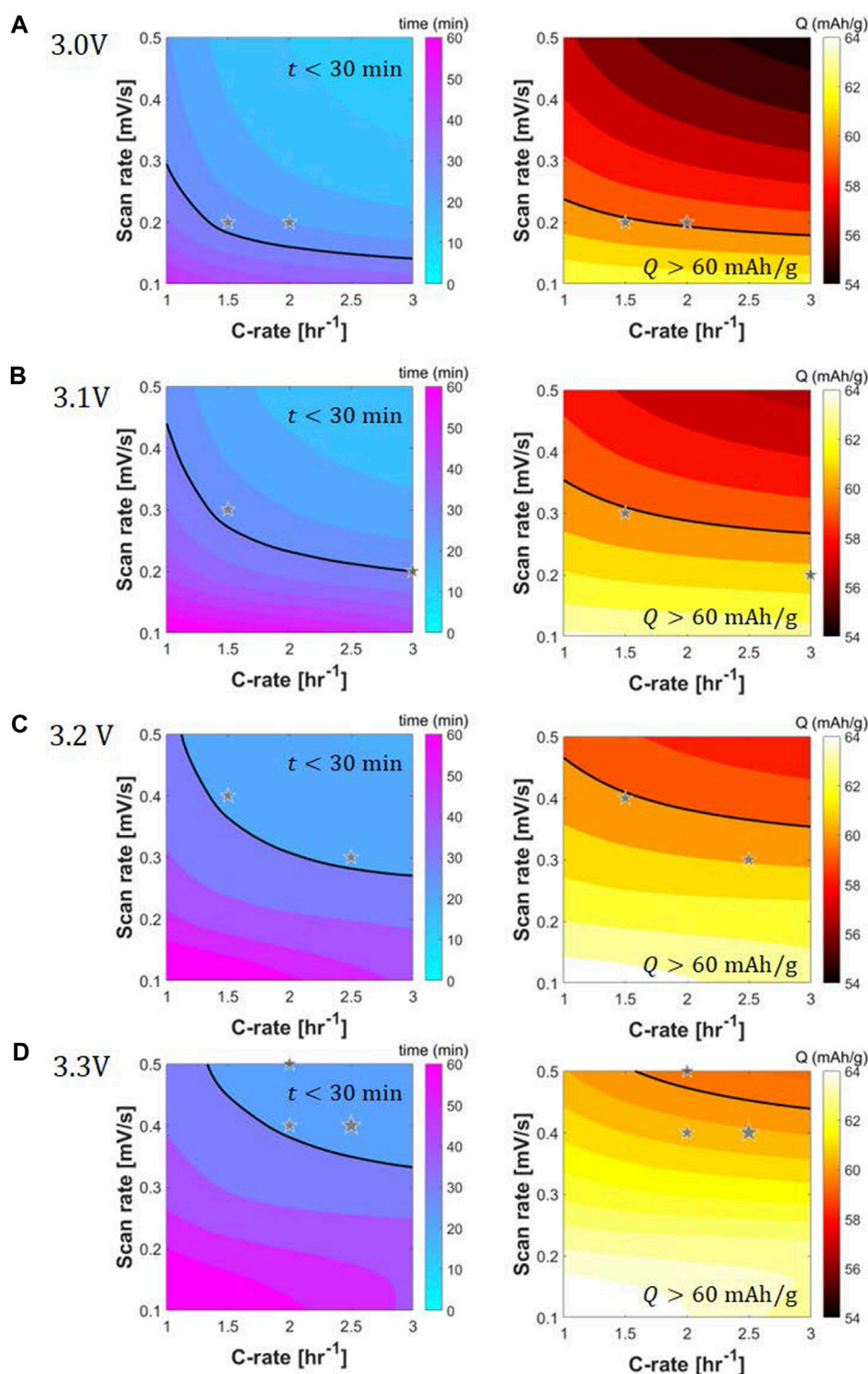


FIGURE 7

Contour plots for performance metrics, time of electrochemical relithiation in minutes (left) and relithiation capacity in mAh/g (right), for parametric sweeps of scanning voltage protocol constant current rate, scan rate and scan step initial voltage: (A) 3.0 V, (B) 3.1 V, (C) 3.2 V and (D) 3.3 V. Black line showcases protocol for which 30 min relithiation and 60 mA h/g capacity gain is achieved. Optimal relithiation protocol showcases relithiation times \leq 30 min and relithiation capacity \geq 60 mAh/g and are highlighted with stars.

from 3.0 to 3.3 V intermediate cutoff. Nominal time spent in scan step increases from 0.2/SR to 0.5/SR for 3.0–3.3 V intermediate cutoff indicating a 250% rise in scan step duration.

Optimal protocols for 3.1 V intermediate cutoff are ascertained to be 1.5C–3.1V–0.3 mV/s–2.8V - 45°C and 3C–3.1V–0.2 mV/s–2.8V - 45°C. When the current rate increases, constant current

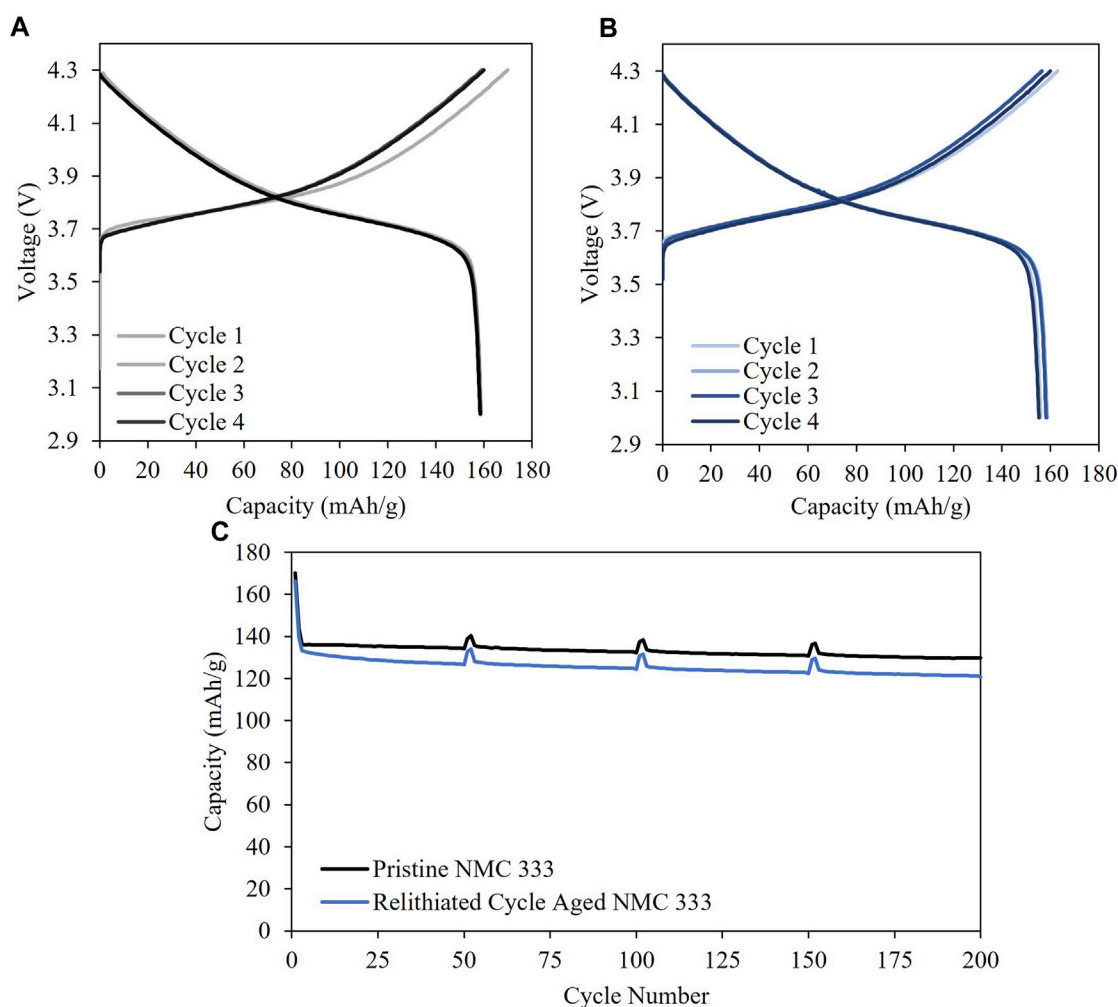


FIGURE 8
Half cell C/10 cycling of (A) Pristine NMC 333 and (B) Relithiated cycle aged NMC 333 and (C) full cell cycling of relithiated cycle aged NMC 333.

step time duration decreases, consequently, the scan rate can be decreased to maintain high capacities while satisfying the time constraint. For 3.2 V intermediate cutoff, the optimal protocols are 1.5C–3.2V–0.4 mV/s–2.8V - 45°C and 2.5C–3.2V–0.3 mV/s–2.8V - 45°C. As the intermediate cutoff increases, the scan rates can be increased to shorten the relithiation time while getting good capacities. Finally, three optimal protocols are delineated for the 3.3 V cutoff: 2C–3.3V–0.5 mV/s–2.8 V - 45°C, 2C–3.3V–0.4 mV/s–2.8V - 45°C and 2.5C–3.3V–0.4 mV/s–2.8V - 45°C. The cycle aged NMC333 material was electrochemically relithiated with the latter protocol showcasing good performance when rebuilt into a half cell in [Supplementary Figure S5](#). The cycle aged NMC333 that was relithiated with the rapid scanning voltage protocol was also analyzed with XRD and ICPOES. [Figure 2](#) shows that the (003) for the relithiated NMC333 materials shifts back to a higher angle reflecting a decrease in the c lattice constant due to the increased lithium content that overlaps with the (003) peak for pristine NMC333. The ICPOES results in [Table 4](#) also demonstrate over 99% of the initial lithium content was restored. The cycle aged NMC333 material was electrochemically relithiated with the latter protocol showcasing good performance when rebuilt into a half cell

with reversible third cycle charge capacity of 161 mAh/g which is comparable with pristine NMC333. The first charge capacity of the relithiated material in half cells was consistently about 98% of pristine NMC 333 which agrees with the ICP and XRD results. Representative voltage profiles for half cells of pristine NMC 333 and relithiated cycle aged NMC 333 are shown in [Figures 8A, B](#). The electrochemically relithiated NMC 333 was also rebuilt into full cells and cycled at C/3 with C/10 cycles every 50 cycles with comparable performance to pristine NMC 333 as shown in [Figure 8C](#). The relithiated NMC 333 material in this study in half cell and full cell cycling had first charge capacities that were consistently <3% lower than pristine NMC 333. To reach 100% pristine performance in NMC materials that are likely to have degradation modes outside of just LLI, a high temperature annealing step will likely be necessary. An annealing step was outside the scope of this work which was constrained to developing a rapid electrochemical relithiation protocol for end-of-life cathodes without any delamination or separation step for the cycle aged NMC powder. This limited the annealing options post-relithiation due to concerns regarding the thermal stability of the copper current collector. Consequently, we demonstrated successful development of a direct recycling method

using model optimized electrochemical relithiation for end-of-life battery cathodes.

4 Conclusion

A novel model-driven process for developing electrochemical relithiation protocols for lightly degraded NMC333 materials was presented. Advanced protocols using constant current scanning voltage discharge was shown to relithiate the degraded NMC333 to near full lithiation within 30 min. Reversible cycling of the relithiated materials was equivalent to pristine NMC333. Electrochemical relithiation is an excellent option for NMC cathode materials that have not undergone extensive mechanical degradation or surface structure degradation such as cation mixing. High Ni NMC cathodes like NMC622, NMC811 are prone to such degradations and hence electrochemical relithiation is hypothesized to be challenging for them. Other materials that will likely benefit from electrochemical relithiation include LFP or LCO since they also reach end-of-life predominantly by loss of lithium inventory to the solid electrolyte interphase instead of loss of active material from particle cracking (Spotnitz, 2003; Sarasketa-Zabala et al., 2015). Future work will focus on how to translate batch results into roll to roll format. Techno-economic modelling is needed to understand the strengths and weakness of electrochemical relithiation compared to other techniques.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

Author contributions

AV: Conceptualization, Methodology, Visualization, Writing—original draft. AC: Funding acquisition, Supervision, Writing—review and editing. JC: Conceptualization, Investigation, Project administration, Validation, Visualization, Writing—original draft.

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Supplementary material

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fbael.2023.1293939/full#supplementary-material>

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