Manufacturing is an integral part of the cost and environmental footprint of batteries. An inexpensive and rapid diagnostic signal was found that can guide improvements in the manufacturing process. The signal can resolve differences in lithium consumed during battery formation and can be used to diagnose the impact of process changes on the lifetime of the battery.

**Highlights**

- A low-SOC resistance signal measured after formation can accurately predict cycle life
- The signal is an indicator for lithium consumed during formation
- The signal improves the detectability of lithium consumption over standard measures
- The signal is measurable immediately after manufacturing using ordinary equipment

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Article

Predicting the impact of formation protocols on battery lifetime immediately after manufacturing

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SUMMARY
Increasing the speed of battery formation can significantly lower lithium-ion battery manufacturing costs. However, adopting faster formation protocols in practical manufacturing settings is challenging due to a lack of inexpensive, rapid diagnostic signals that can inform possible impacts to long-term battery lifetime. This work identifies the cell resistance measured at low states of charge as an early-life diagnostic feature for screening new formation protocols. We show that this signal correlates to cycle life and improves the accuracy of data-driven battery lifetime prediction models. The signal is obtainable at the end of the manufacturing line, takes seconds to acquire, and does not require specialized test equipment. We explore a physical connection between this resistance signal and the quantity of lithium consumed during formation, suggesting that the signal may be broadly applicable for evaluating any manufacturing process change that could impact the total lithium consumed during formation.

INTRODUCTION
With the increasing demand for electric vehicles, global lithium-ion battery manufacturing capacity is quickly approaching the terawatt-hour scale.1–3 A key step in battery manufacturing is formation/aging, which has been estimated to account for up to 30% of total manufacturing costs.4–8 The formation/aging process involves charging and discharging hundreds of thousands of cells in environmentally controlled chambers, an expensive process that takes days to weeks to complete but is necessary to improve battery performance and lifetime.9–14 Given the high cost burden, manufacturers are incentivized to develop new formation processes that decrease the total time consumed by formation/aging. A variety of fast formation strategies have been studied in academic literature, which employ some combination of rapid charge-discharge cycles, restricted voltage windows, and optimized temperature.15,15–26 Recent studies have shown that formation time can be decreased while preserving battery lifetime,16,22,25 although conclusions remain tenuous due to the limited sample sizes typically used.

In real manufacturing settings, a “one-size-fits-all” formation protocol is unlikely to exist since cell designs with different electrolytes, electrodes, and active materials influence important formation factors such as charging capability, electrode wettability, and solid electrolyte interphase (SEI) reaction pathways. However, cycle life testing often takes months or years to complete, posing a significant barrier to the adoption of new, potentially cost-saving formation protocols. While characterization techniques, such as volume change detection,27–29 impedance spectroscopy,15,30

Context & scale
Despite recent progress in battery development, electric vehicles remain unaffordable for many. A key enabler for less expensive electric vehicles is lowered battery manufacturing costs, a significant portion of which is due to the formation and aging process. Although some fast formation protocols have been proposed, a one-size-fits-all solution is unlikely to succeed in practice since an optimized formation protocol for one battery design will, in general, not be optimal for another. New formation protocols need to be vetted for their impacts on long-term battery lifetime—a slow process that hinders the discovery of optimal formation protocols. Here, we identify a scalable method for predicting the effect of new formation protocols on cycle life. The method is obtained directly at the end of the manufacturing line and can be deployed immediately in mass production settings to improve diagnostics of new formation protocols.
acoustic spectroscopy, and X-ray tomography, have been proposed for use in manufacturing settings, these methods can be costly to implement since the metrology will need to be deployed at scale in the battery factory. Diagnostic features obtainable from already existing cycling equipment and especially those using only current-voltage signals are thus highly attractive.

In this work, we show that the cell resistance at low states of charge (SOC) can be used to screen new formation protocols and predict battery lifetime. Our work shows that this signal, measured at the beginning of life, is a stronger predictor of battery lifetime than conventional signals such as Coulombic efficiency (CE). This metric can be measured within seconds and integrated directly into the battery manufacturing process with no additional capital costs. This low-SOC resistance metric can, thus, be deployed in practical manufacturing settings to accelerate the evaluation of new formation protocols. We further demonstrate that the low-SOC resistance \( (R_{LS}) \) decreases as the quantity of lithium lost to the SEI during formation increases. With our physical insight, we propose that \( R_{LS} \), in principle, can also be used to diagnose the impact of any manufacturing process that alters the total lithium consumed during formation.

**RESULTS AND DISCUSSION**

**Fast formation experimental design**

Two formation protocols have been implemented in this work: a fast formation protocol previously reported by Wood et al. that completes within 14 h (Figure S1B) and a baseline formation protocol (Figure S1C) that completes in 56 h. The fast formation protocol maximizes the time spent at low negative electrode potentials to promote the creation of a more passivating SEI.

Forty nickel manganese cobalt (NMC)/graphite pouch cells with a nominal capacity of 2.36 Ah were built for this study (Table S1). Half of the cells underwent fast formation, and the remaining cells underwent baseline formation. Cells were further subdivided into “room temperature” and “45 °C” aging groups for cycle life testing. The cycling profile was identical for all cells: 1 C charge to 4.2 V with a constant voltage (CV) hold to 10 mA and 1 C discharge to 3.0 V. Reference performance tests (RPTs) were inserted throughout the cycle life test, which includes slow (C/20) charge and discharge curves as well as a hybrid pulse power characterization (HPPC) sequence used to extract the cell internal resistance as a function of SOC.

Our experimental design (Figure S1A) uses larger samples sizes \( (n = 10 \text{ per group}) \) compared with those typically reported in the literature, which often use three cells or fewer per group. The increased sample size enables a more statistically rigorous analysis of the impact of different formation protocols on cell characteristics at the beginning and the end of life.

**Fast formation cells had longer cycle life**

Fast formation cells had higher average lifetimes than the baseline formation cells under the cycle life test, as shown in Figure 1. The degradation rate of fast formation cells initially track the baseline formation cells closely under both temperatures tested (Figures 1A and 1C). However, after 250 cycles, all cells begin to lose capacity rapidly. The fast formation cells sustained over 100 cycles longer before reaching the end of life, defined as when cells reach 70% of their initial measured capacity (Figures 1B and 1D). This result is highly statistically significant (p-value < 0.001). The general result that fast formation improved lifetime performance holds across multiple performance metrics, including CE (Figure S4), voltage efficiency (Figure S5), as well as...
when plotted against equivalent cycles (Figure S7). Together, these results support the growing body of evidence that well-designed fast formation protocols can improve cycle life.15,22,38

Finding diagnostic signals at the beginning of life
Given the demonstrated impact of formation protocol on battery cycle life, we next investigate methods to quantify the impact of fast formation on the initial cell state. Differences in the initial cell state (e.g., the amount of lithium consumed during formation) may offer clues as to how fast formation could have improved cycle life. We focused on studying signals directly obtainable from full cell current-voltage data, which offer the lowest barrier-to-entry for deployment in real manufacturing settings.

Conventional metrics of formation efficiency
Figures 2A–2C show standard measures of formation efficiency extracted from the formation cycling data. The discharge capacity, \(Q_d\), was measured at the end of each formation protocol during a C/10 discharge step from 4.2 to 3.0 V. \(Q_d\) corresponds to the capacity of cyclable lithium excluding the contribution from lithium irreversibly lost to the SEI during formation. Fast formation decreased \(Q_d\) by 0.3%, a small but statistically significant difference (\(p = 0.01\)). The charge capacity, \(Q_c\), was taken during the initial charge cycle and includes both the capacity of cyclable lithium as well as the capacity of lithium lost irreversibly to the SEI. The quantity of
lithium inventory lost to the SEI can be calculated as $Q_{LLI} = Q_c - Q_{d}$ (Figure 2B). Note that while the two formation protocols differed in the initial charging rate, $Q_c$ remains a fair comparison metric since both charge protocols ended on a potential-hold at 4.2 V until the current dropped below C/100. Fast formation increased $Q_{LLI}$ by 23 mAh ($p = 0.03$). Finally, we also included another common evaluation metric, the formation CE, defined as $CE_f = Q_d/Q_c$ (Figure 2C), which shows that fast formation decreased $CE_f$ by 0.8% ($p = 0.02$). Measured values are summarized in Table 1. Together, the results show that fast formation marginally increased the amount of lithium consumed during formation. A $p$-value of less than 0.05 in all cases indicate that the measured differences, while small, are statistically significant to a least a 95% confidence level.

Low-SOC resistance
Following formation, the cell internal resistance was measured using the hybrid pulse power characterization (HPPC) technique prior to the start of the cycle life test. During this test, a series of 10-s, 1 C discharge pulses were applied to the cell at varying SOCs, and the resistance is calculated using Ohm’s law (Figure S8). The 10-s resistance, $R_{10s}$, was plotted against SOC for all cells cycled at 45°C.
R10s generally remained flat at mid-to high SOCs. The peak at 55% SOC corresponds to the stage 2 solid-solution regime of the graphite negative electrode.43 R10s rose sharply below 10% SOC. Focusing on the low-SOC region (Figure 2E), we observed that R10s measured at 4% and 8% SOC were lower for fast formation cells compared with that of baseline formation cells. This result was highly statistically significant, with a p-value less than 0.001 (Figure 2F). A similar result held when R10s was measured at room temperature (Figure S9). At mid to high SOCs, differences in R10s between fast formation and baseline formation cells were generally not statistically significant (Figure S9). Thus, differences in resistance between the two formation protocols appeared uniquely at low SOCs. All initial cell state metrics are summarized as part of Table 1.

To study the robustness of the low-SOC resistance signal, we varied the SOC set-point between 4% and 10% and also computed the resistance under 1 and 5-s pulse durations. In all cases, the resistance metric provided a high degree of contrast between the two different formation protocols (Figures S10 and S11). The lowest SOC measured in our dataset was 4% SOC.

The remainder of the paper will focus on the resistance measured at 5% SOC and with a 10-s pulse duration. From hereon, this metric will be referred to as the “low-SOC resistance,” RLS.

Low-SOC resistance as a diagnostic signal: A data-driven perspective

Low-SOC resistance correlates to cycle life

To evaluate the merit of RLS as a diagnostic feature, we explored the correlations between the initial cell metrics (Figure 2) and cycle life, defined as cycles to 70% of the initial capacity. The results are shown in Figure 3. Out of all metrics studied, RLS was the only signal with a meaningful correlation to cycle life, with a correlation coefficient of \( \rho = -0.84 \). Other metrics such as \( Q_d \) and CEf were poorly correlated to cycle life (\( |\rho| <0.5 \)). We attribute the weakness of these correlations to the poor signal-to-noise inherent in cell capacity measurements in the absence of high-precision cycling,44,45 a topic we explore in detail later. The resistance measured at high SOCs also did not correlate to cycle life. From these results, we observe that the low-SOC signal uniquely holds information related to cycle life. These results have been reproduced for different end-of-life definitions ranging between 50% and 80% (Figures S13 and S14), as well as for charge pulses (Figure S15).

Low-SOC resistance predicts cycle life

To understand if RLS can be used to improve battery lifetime prediction, we trained univariate prediction models with regularized linear regression models inspired by...
The performance of the predictive models are summarized in Table 2. A dummy regressor, which predicts the mean of the training set and requires no cycling data, was included as a benchmark. For room temperature cycling, the model trained using $R_L$ achieved the lowest test error of 6.9% compared with 13.3% for the dummy regressor. A similar result held under 45°C cycling. To compare, we also included the $\text{Var}(\Delta Q_{100-10}(V))$ metric introduced by Severson et al., defined as the variance in the discharge capacity versus voltage curve between cycle 10 and cycle 100. When applied to our dataset, this metric did not yield a significant improvement over the dummy regressor. This result suggests that $R_L$ is a stronger predictor of battery lifetime than $\text{Var}(\Delta Q_{100-10}(V))$.

We repeated this study with multivariate regularized linear regressions: one using the three capacity-based features from formation ($Q_{LLI}$, $CE_f$, and $Q_d$) and another using the previous three formation features plus $R_L$. Using only the features from formation, no improvement over the dummy regressor was achieved. By including $R_L$ in the feature set, however, the test error was improved. Yet, the test error achieved did not exceed the test error of the univariate model using $R_L$ alone. This result suggests that the chosen set of formation features does not provide useful information about cycle life beyond what is provided by $R_L$. This result is counter-intuitive considering the important role that lithium consumption plays in determining battery lifetime, which should be reflected in the formation features such as $Q_{LLI}$ and $CE_f$. We hypothesize that the reason for the poor model
performance using formation signals is not because these formation signals lack physical meaning. Rather, due to the absence of high-precision cycling and temperature control, the useful information within these signals may be masked by the noise in the data (e.g., due to current integration errors and temperature variations over the course of 10+ h of formation) RLS appears to be able to overcome these limitations. We explore the connection between RLS and the other formation metrics in detail later.

As defined in this work, the model trained using RLS required just three cycles of lifetime testing, i.e., one diagnostic cycle, where the two preceding cycles consisted of slow-rate charge-discharge cycles as part of the RPT inserted at the beginning of the cycle life test. By comparison, Var\(\left(\Delta Q_{100-10}(V)\right)\) requires 100 cycles of lifetime testing. For future implementations, RLS can be incorporated directly into the formation protocol, further decreasing the required measurement time. The total amount of data required to exercise each predictive model is summarized in Table 2.

Overall, the correlation and prediction results suggest that RLS may be useful for advancing broad-scale efforts to improve cycle life prediction using small and readily obtainable datasets at the beginning of life. While the results are promising, they are also limited, since only two types of formation protocols have been studied here. To understand the extent to which RLS can generalize to other applications (e.g., chemistries, use cases, and cell designs) and to understand the relation between RLS and the other formation signals, the rest of the paper will focus on providing a physical interpretation of RLS. A mechanistic understanding of RLS will provide the necessary context required to evaluate the general scope of applicability and limitations of the method.

Low-SOC resistance as a diagnostic signal: A mechanistic perspective
Understanding the physical interpretation of diagnostic signals can help assess whether prediction frameworks leveraging such signals can generalize to new systems. In principle, different formation protocols, manufacturing process changes, and cell design changes could all lead to changes in lithium consumption and active material losses during formation. To this end, we will first review the commonly accepted theory of SEI passivation and show how our observations of QLLI and CEf support this theory. Next, we will show that our observations of RLS are consistent with this theory but provide a stronger and more easily measurable signal compared with conventional measures.

| Model                  | Data needed | Room temp |         |         | 45° C |         |         |
|------------------------|-------------|-----------|---------|---------|-------|---------|---------|
| Dummy regressor        | None        | 13.3 (1.0) | 14.4 (4.0) | 14.0 (0.9) | 15.1 (3.6) |
| RLS                    | 3 cycles    | 6.9 (0.5)  | 8.0 (2.6)  | 6.5 (0.6)  | 7.4 (2.9)  |
| QLLI                   | formation   | 12.2 (1.2) | 14.0 (4.6) | 14.1 (0.8) | 15.2 (4.4) |
| CEf                    | formation   | 12.2 (1.2) | 13.8 (4.5) | 14.1 (0.7) | 15.1 (4.3) |
| Qd                     | formation   | 12.0 (1.2) | 13.6 (5.0) | 13.5 (0.8) | 15.0 (4.0) |
| Var(\(\Delta Q_{100-10}(V)\)) | 100 cycles  | 11.6 (1.7) | 14.4 (5.2) | 10.3 (1.1) | 11.5 (4.7) |
| QLLI + CEf + Qd        | formation   | 12.8 (1.3) | 14.5 (5.1) | 13.4 (1.1) | 14.1 (4.0) |
| QLLI + CEf + Qd + RLS  | 3 cycles    | 7.2 (1.1)  | 9.4 (4.0)  | 6.5 (1.0)  | 7.4 (2.9)  |

Values represent means (standard deviations). The dummy regressor model uses no features and returns the mean of the training set, and hence is the baseline against which to judge the performance of other features. All remaining models use a ridge regression with nested cross-validation to determine the optimal regularization strength. See experimental procedures.
**Benefits of fast formation on cycle life**

Lithium intercalation at negative electrode potentials higher than 0.25–0.5 V versus Li/Li⁺ is generally associated with the formation of a porous, poorly passivated SEI film. Conversely, lithium intercalation at negative electrode potentials below 0.25–0.5 V have been found to promote the formation of a more conductive and passivating SEI film. Attia et al. showed that the reduction of ethylene carbonate (EC) at negative electrode potentials above 0.5 V versus Li/Li⁺ is non-passivating. This negative electrode potential corresponds to a full cell voltage of below 3.5 V, neglecting overpotential contributions. Hence, an ideal formation protocol would minimize the time spent charging below 3.5 V while maximizing the time spent above 3.5 V. The fast formation protocol we tested achieves this objective by rapidly charging the cell to above 3.9 V at a 1 C rate and subsequently cycling the cell between 3.9 and 4.2 V, thus decreasing the time associated with the non-passivating EC reduction reaction. Focusing on the initial charge cycle, fast formation cells spent only 2 min below 3.5 V and 12.9 h above 3.5 V, while baseline formation cells spent 30 min below 3.5 V and 9.4 h above 3.5 V. Fast formation decreased the time spent below 3.5 V by 28 min. Fast formation resulted in a net increase in total lithium consumed during formation, ΔQ_{Li}, by 23 mAh (Table 1). This increase is attributed to the additional lithium consumed to form the passivating SEI.

While fast formation cells consumed more lithium during formation and, thus, exhibited lower CE (or, equivalently, higher Q_{Li}), these cells lasted longer on the cycle life test. While a lower initial CE is conventionally associated with poor cycle life performance, the opposite was true in our study since the additional lithium consumed during fast formation was associated with the creation of a more passivating SEI. A more passivating SEI can, for example, lower the rate of electrolyte reduction reactions associated with the formation of solid products that decrease the negative electrode porosity and subsequently increase the propensity for lithium plating during charge. A more passivating SEI could, therefore, play a role in delaying the “knee-point” observed in the cycle test data. Our result reinforces the notion that passivation of the SEI during the first cycle plays an important role in improving battery cycle life.

**Lithium loss dominates overall cell capacity loss over cycling**

We performed a voltage fitting analysis to confirm that the main failure mode in our cells is the loss of lithium inventory (LLI) over cycle life (Figures S18 and S19). We found that LLI can fully account for the thermodynamic (i.e., C/20) cell capacity loss over life. The knee-point in LLI over cycle life coincides with the knee-point in the capacity loss. All cells also experienced an increase in the loss of active material (LAM) in the negative electrode (LAM_{NE}) after the knee-point, which could indicate the occurrence of porosity decrease and/or electrolyte depletion as a result of a less passivating SEI, as discussed previously. The increased LAM_{NE} after the knee-point was less prominent in the fast formation cells, suggesting that the more passivating SEI generated from fast formation could be playing a role in delaying the knee-point to improve lifetime. Finally, all cells experienced a knee-point in the capacity fade rate irrespective of whether the discharge capacity is measured at higher (C/3) or lower (C/20) C-rates (Figure S20), indicating that kinetic limitations cannot fully account for the observed knee-point in the cycle life data. The origin of the capacity loss, therefore, has a strong thermodynamic component, which can be attributed to the LLI. This analysis further supports the theory that consuming more lithium at low negative electrode potentials during formation can create a passivating SEI that is beneficial to cycle life.
Low-SOC resistance is attributed to kinetic limitations in the positive electrode

To explore possible physical connections between $R_{LS}$ and the impact of fast formation on cycle life, we first developed a physical interpretation of the $R_{LS}$. Here, we focus our discussion on the resistance contributions from the positive and negative electrode. While other cell components (e.g., current collectors, tabs, and electrolyte) also contribute to the total cell resistance, they are not known to depend on SOC and, hence, cannot explain the rising resistance measured at low SOCs.

Positive electrode diffusion limitations generally play a significant role in the low-SOC cell resistance in NMC/graphite systems. The solid-state diffusion coefficient in NMC materials has been measured to decrease by more than one order of magnitude at high states of lithiation, a phenomenon attributed to the depletion of divacancies needed to support diffusion as the electrode becomes fully lithiated. Using half-cell HPPC measurements, we experimentally verified that the positive electrode dominates the $R_{LS}$. In the coin cell form factor, the 10-s resistance of graphite/Li stayed below 100 mΩ as the graphite approached full delithiation, while the 10-s resistance of NMC/Li exceeded 1,000 mΩ as the NMC approached full lithiation (Figure S22). This finding is consistent with previous empirical studies on NMC/graphite systems. In particular, An et al. used a three-electrode pouch cell configuration to show that, for an NMC/graphite system, the positive electrode accounts for nearly all of the measured full cell resistance at all SOCs.

Charge transfer kinetics at either electrode could also play a role in determining total cell resistance. The charge transfer process at either electrode can be modeled using the Butler-Volmer equation:

$$j = j_0 c_a^{1-a} (c_{s,max} - c_s)^{1-a} c_s^a \left( \exp \left( \frac{1 - a}{RT} \eta \right) - \exp \left( \frac{-a F}{RT} \eta \right) \right).$$  \hspace{1cm} (Equation 1)

In this equation, $j$ is the reaction flux, the exponential terms describe the overpotential dependence of the forward and backward reactions, and the exponential prefactor terms together describe the exchange current density. $c_{s,max}$ is the theoretical maximum allowable lithium concentration in the solid phase, $c_s$ is the surface concentration of lithium, and $j_0$ is the reaction rate constant. The exchange current density approaches zero as the electrode becomes either fully lithiated or fully delithiated. Indeed, our coin cell data show that as the graphite negative electrode approaches full delithiation, the measured resistance rises steeply (Figure S22i). However, the magnitude of this charge transfer effect remains small compared with the contribution from the diffusion-limited NMC positive electrode (Figure S22h) at high states of lithiation.

In summary, we attribute the $R_{LS}$ to kinetic limitations in the positive electrode. This result was experimentally verified using coin cell measurements of electrode resistances and is consistent with literature findings. The kinetic limitation arises from a combination of diffusion and charge transfer limitations in the positive electrode. For NMC/graphite systems, diffusion limitations (i.e., “kinetic hindrance”) is a major component of the rapid rise in measured resistance at low SOCs.

Lithium consumption leads to an apparent decrease in low-SOC resistance

Fast formation decreased the measured $R_{LS}$. From our previous analysis, fast formation also increased the lithium consumed during formation ($Q_{Li}$) to create a more passivating SEI. To explain the connection between these two quantities, we employed a simple electrode stoichiometry model that describes both the thermodynamic potentials and kinetic limitations of both electrodes. Figure 4A shows the...
relative alignment of the positive and negative equilibrium potential curves after baseline formation and fast formation. The origin of the capacity axis corresponds to 0% SOC (3.0 V) after baseline formation. The gap between the positive and negative potential curve endpoints is attributed to the lithium lost to the SEI during formation or $Q_{\text{LLI}}$. By comparison, the curves prior to formation do not have a gap, corresponding to $Q_{\text{LLI}} = 0$ (Figure S23). The positive electrode curve was shifted to the left by some amount $\Delta Q_{\text{LLI}}$ to emulate the impact of additional lithium consumed during fast formation. Here, $\Delta Q_{\text{LLI}}$ has been set to an exaggerated value of 100 mAh for graphical clarity. An alternative graphic is provided in Figure S24, which sets $\Delta Q_{\text{LLI}} = 23$ mAh to coincide with the measured difference between baseline formation and fast formation.

Figure 4C shows the corresponding full cell 10-s resistance measured from the HPPC test. The full cell resistance is partitioned to model a scenario in which the positive electrode dominates the $R_{\text{LS}}$, consistent with previous findings. The resistance curve of the positive electrode must also translate to the left by the same amount $\Delta Q_{\text{LLI}}$ due to the increased lithium consumed during fast formation. From the reference
frame of the full cell, the measured $R_{LS}$ will decrease by $\Delta R_{LS}$. In this manner, $R_{LS}$ can decrease without any real change in positive electrode kinetic properties. The decrease in $R_{LS}$ reflects the shifting of the positive electrode stoichiometry window as lithium is consumed.

Two additional observations support the connection between $\Delta Q_{LLI}$ and $\Delta R_{LS}$. First, $R_{LS}$ appears to be positively correlated to $CE_f$ and negatively correlated to $Q_{LLI}$ (Figure S12), a result consistent with theory and predicted by the electrode stoichiometry model. The strengths of the correlations are generally weak, with correlation coefficients, $|\rho|$, ranging between 0.2 and 0.5. We attribute the weakness of the correlations to the poor signal-to-noise of the capacity measurements using typical battery cycling equipment, which may compound at room temperature where the temperature is not strictly controlled (Figure S25). Second, we note that the resistance around 90% SOC is insensitive to small changes in SOC, so changes in resistance at 90% SOC provides a measure of true resistance changes rather than apparent changes due to electrode stoichiometry shifts (Figure 2D). Fast formation did not significantly increase the resistance at 90% SOC (Figure S9), so changes in $R_{LS}$ are not likely due to material changes in the cell resistance (e.g., due to resistive surface films). This observation further supports the hypothesis that changes in $R_{LS}$ are due to electrode stoichiometry window shifts in the presence of lithium consumption.

Low-SOC resistance improves the observability of lithium loss during formation

Figure 4B shows that the sensitivity of the measured cell discharge capacity ($\Delta Q_d$) to the lithium consumed ($\Delta Q_{LLI}$) is 0.9 mAh/mAh. The error in measuring $Q_d$ is 20 mAh due to current integration inaccuracies using ordinary cycling equipment. Hence, using $Q_d$ to estimate $Q_{LLI}$ leads to a measurement error of 22 mAh. Since the total difference in lithium consumed between fast formation and baseline formation is 23 mAh, measurement noise may prevent $\Delta Q_d$ from effectively resolving this difference. In our experiments, we relied on large sample sizes ($n=10$ per group) to resolve the small difference in lithium consumption between the two formation protocols.

Figure 4D shows that the sensitivity of the low-SOC resistance ($\Delta R_{LS}$) to $\Delta Q_{LLI}$ is 0.22 m\(\Omega\)/mAh when measured at 5% SOC. The error in measuring $R_{LS}$ is 0.88 m\(\Omega\) due to the voltage and current precision for calculating resistance using Ohm’s law and using ordinary cycling equipment. Hence, using $R_{LS}$ measured at 5% SOC to estimate $Q_{LLI}$ leads to a measurement error of 4 mAh, a five-fold improvement over using $Q_d$. Figure 4D further shows that the sensitivity of $R_{LS}$ is improved at lower SOCs. For example, $R_{LS}$ measured at 2% SOC leads to a measurement error of 2.5 mAh. Any SOC set-point lower than 7% SOC makes $R_{LS}$ a more precise measure of $Q_{LLI}$ compared with $Q_d$. (See supplemental information for a detailed derivation of the measurement errors.)

Generalizability

So far, we have explored the sensitivity of $R_{LS}$ to lithium lost during formation for an NMC/graphite system. By understanding the benefits of fast formation, we rationalized why $R_{LS}$ was predictive of cycle life for our system. Here, we discuss the application of $R_{LS}$ toward understanding other degradation modes, chemistries, and use cases. This discussion sets the stage for understanding how $R_{LS}$ may be incorporated into generalizable lifetime prediction and diagnostic frameworks.
**R}_{LS} can detect active material losses**

In principle, some small quantity of positive and negative active material could be lost during formation, i.e., due to expansion and contraction of the electrodes during initial lithiation and delithiation. In the positive electrode, lithiation-induced stresses can induce particle fracturing in the metal oxide particles, leading to capacity loss. In the negative electrode, while graphite cracking is unlikely to occur under most applications, insufficient binder adhesion or electrolyte wetting could create local islands of isolated graphite particles, leading to active material loss.

We develop a simple mechanistic electrode stoichiometry model to examine the influence of active material losses in both the positive and negative electrodes. Our model differentiates between LAM in the lithiated phase versus the delithiated phase. For the positive electrode, LAM in the delithiated phase is represented by shrinking the positive electrode equilibrium potential curve with the point of minimum stoichiometry fixed (i.e., shrinking from the bottom, Figure S26A), while LAM in the lithiated phase is represented by shrinking the positive electrode equilibrium potential curve with the point of maximum stoichiometry fixed (i.e., shrinking from the top, Figure S26D). R_{LS} was found to increase with loss of positive active material, but only in the delithiated phase (Figure S26B). By contrast, active material lost in the lithiated phase bears a negligible effect on R_{LS} (Figures S26D and S26E). This result can be understood graphically by considering the influence of the positive curve shifts on the positive electrode stoichiometry at low SOCs. In the case of LAM in the lithiated phase, the positive electrode stoichiometry at low SOCs does not significantly change, whereas in the delithiated case, the maximum positive electrode stoichiometry increases, causing R_{LS} to increase. Note that \( Q_d \) has the opposite sensitivity: \( Q_d \) is sensitive to LAM in the lithiated state only. Hence, R_{LS} and \( Q_d \) complement each other in the study of positive electrode active material loss mechanisms. A similar analysis can be done on the negative electrode (Figure S27).

Figure S28 compares the sensitivity of R_{LS} and \( Q_d \) to the four different modeled cases of active material losses. The results highlight that the measured value of R_{LS} is determined by multiple degradation factors, including both lithium inventory loss and active active material losses. It would therefore be impractical to use R_{LS} to identify any dominant degradation mode without some a priori understanding of the system through additional characterization and analysis. For diagnostic purposes, we recommend that R_{LS} be used within the context of a broader set of non-destructive techniques to enrich the understanding of degradation mechanisms. From a data-driven prediction perspective, however, the sensitivity of R_{LS} to active material losses in addition to lithium inventory loss may make it a more robust indicator for multiple degradation modes. In general, R_{LS} may need to be coupled with other signals to improve the observability of distinct degradation modes.

**When is R_{LS} sensitive to lithium loss?**

We have so far focused on an NMC111/graphite system where kinetic limitations in the positive electrode dominates R_{LS}, a result that holds for nickel-rich cathode chemistries such as nickel cobalt aluminum (NCA) and higher nickel content NMC materials. In general, electrode design factors such as particle size and surface modifications could impact the relative contribution of each electrode to R_{LS}. To study how such changes could modify the sensitivity of R_{LS} to changes in \( Q_{\text{LII}} \), we performed a sensitivity study using our electrode stoichiometry model by varying the proportion of the total cell resistance attributed to the positive electrode. The results (Figures S29 and S30) show that R_{LS} becomes ineffective at quantifying \( Q_{\text{LII}} \) if the positive electrode contributes to less than 50% of the total cell resistance at
This result suggests that the utility of RLS as a diagnostic signal for $Q_{LLI}$ diminishes for systems where the positive electrode is not the main contributor to RLS.

When can RLS predict cycle life?

Our cycle life correlation study was presented in the context of the study of fast formation. To understand whether RLS can predict cycle life for other use cases (i.e., chemistries and aging conditions), we start by reviewing why RLS was predictive of cycle life for fast formation. Figure 5 outlines the proposed connection between fast formation and cycle life. Fast formation spent more time above 3.5 V, creating a higher quantity of SEI that is more passivating. The passivating SEI improved cycle life by protecting the negative electrode against side reactions over life. RLS provided an estimate of the amount of lithium consumption during formation, $Q_{LLI}$, where more lithium consumed implied a higher amount of passivating SEI formed leading to improved cycle life. This physical description rationalizes the predictive power of RLS within the context of the specific degradation pathway (fast formation) and chemistry (NMC/graphite) explored in this study.

To gain confidence that RLS can predict cycle life for other use cases, the relationship between lithium loss ($Q_{LLI}$) and cycle life must first be understood. For our study, the knowledge that increased $Q_{LLI}$ signals a more passivating SEI was necessary for rationalizing why higher $Q_{LLI}$ after formation could be beneficial to cycle life. For other use cases, the opposite may be true. For example, low first cycle efficiencies for silicon-containing anodes or lithium metal anodes generally indicate poor negative electrode passivation, which leads to poor cycle life. Under such use cases, RLS may still be predictive of cycle life, but the relationship may become inverted.

Unique properties of RLS

Here, we highlight several unique properties of using RLS as an early-life diagnostic signal. First, since the positive electrode kinetics becomes increasingly poorer as the electrode approaches full lithiation, the sensitivity of RLS to lithium loss ($Q_{LLI}$) improves as the measurement SOCs decreases (Figure 4D). The results from this study used RLS measured at 5% SOC. For future work, the sensitivity to $Q_{LLI}$ may be further improved by taking the measurement at even lower SOCs. Second, RLS can be used to extract information about $Q_{LLI}$ within seconds and, therefore, can be deployed in manufacturing settings without decreasing production speed. By contrast, conventional measurements of $Q_{LLI}$ relying on Coulomb counting require full charge-discharge cycles during formation, which could take hours to days to complete.

Figure 5. Connection between the fast formation degradation pathway and the low-SOC resistance early-life diagnostic signal

Inner box: the relationship between low-SOC resistance ($R_{LS}$) and lithium consumed during formation ($Q_{LLI}$) is general.

Outer box: the relationship between low-SOC resistance ($R_{LS}$) and cycle life applies specifically to fast formation, where higher $Q_{LLI}$ signaled a higher quantity of passivating SEI which improved cycle life. The relationship between $R_{LS}$ and cycle life may differ for other use cases.
Since measuring $R_{LS}$ does not require full cycles, $R_{LS}$ is also suitable for diagnosing differences in lithium consumption between formation protocols with different charge and discharge conditions. Finally, the magnitude of $R_{LS}$ is larger the earlier in life it is measured. As the cell ages, continual LLI will cause the highly sloped region of the positive electrode resistance curve to become inaccessible during the normal full cell voltage operating window. Typically, diagnostic features become less predictive of cycle life the earlier in life the feature is sampled. $R_{LS}$ is expected to have the opposite relationship: the earlier in life $R_{LS}$ is sampled, the more sensitive it will be to changes in $Q_{LLI}$.

**Diagnosing state of health beyond cycle life: practical considerations**

Our discussion has so far focused on evaluating the merits of $R_{LS}$ for diagnosing cycle life. However, in real manufacturing settings, cycle life is only one of many considerations for adopting new formation protocols. Here, we introduce two such considerations: (1) impact to gas buildup over life, and (2) impact to aging variability over life. In our analysis, $R_{LS}$ could not be used to learn the impact of fast formation on gas buildup or aging variability. Here, we give an overview of these observations.

**Gas buildup over life**

Swollen cells in a battery pack can compromise pack integrity and pose safety hazards to first-responders for electric vehicle fire accidents. Understanding the impact of formation protocols on cell swelling is, therefore, just as important as understanding the impact on cycle life for practical purposes.

Fast formation caused a significant degree of swelling at the end of life for cells cycled at 45°C (Figures S31 and S32). At this temperature, 9 of 10 fast formation cells showed visible signs of swelling, compared with only 2 of 10 for baseline formation. None of the cells cycled at room temperature showed any appreciable degree of swelling. All swollen pouch cells were compliant and compressible, indicating that gas is occupying the space inside the pouch bags. Since the cells were de-gassed after formation, the gases present excludes the gas generated during formation and represent only the accumulation of gas over the course of the cycle life test. The absence of gas during room temperature cycling indicates that the gas evolution is thermally activated. More experimental work is needed to determine the origin of gas evolution over cycle life due to fast formation. We provide speculation into the origin of gas evolution as part of the supplemental information.

Our study found no correlations between $R_{LS}$ and the gas amount as measured by pouch thickness. We attribute the lack of correlation primarily to the fact that the cell age was not well-controlled at the time of the pouch thickness measurement: cells stopped cycling anywhere between 0% and 50% capacity retention. Future studies will be needed to confirm the relationship between $R_{LS}$ and gas buildup.

**Aging variability**

Adopting a new formation protocol in practice also requires a close understanding of the impact of new formation protocols on cell aging variability over life. Cells with non-uniform capacity fade could take longer to balance in a pack and cause a deterioration of energy available at the pack-level. Pack imbalance issues could lead to consumer products being retired earlier, compounding existing battery recycling challenges. Non-uniform cell degradation will also be more difficult to re-purpose into new modules, creating higher barriers for pack reuse.
The inter-quartile range (IQR) of cycle lives for fast formation cells was higher than that of baseline formation cells (Figures 1B and 1D). The same result held under both room temperature and 45°C cycling, as well as across different end-of-life definitions (Figure S33), suggesting that fast formation increased aging variability. A key question is whether fast formation created more heterogeneous aging behavior, which caused the higher variability in aging, or if the higher variability is due to the cells lasting longer. To answer this question, we employed the modified signed-likelihood ratio test⁷⁸ to check for equality of the coefficients of variation, defined as the ratio between the standard deviation and the mean cycle life. The resulting p-values were greater than 0.05 in all cases. Therefore, with the available data, we cannot conclude that fast formation increased the variation in aging beyond the effect of improving cycle life. While a relationship between formation protocol and aging variability may still generally exist, this difference could not be determined with our sample sizes (n = 10 cells per group). This result warrants the use of larger samples sizes for future studies on the impact of formation protocol on aging variability.

**Conclusion**

In this work, we demonstrated that $R_L$ correlates to cycle life across two different battery formation protocols. As a predictive feature, $R_L$ provided higher prediction accuracy compared with conventional measures of formation quality such as CE as well as state-of-the-art predictive features based on changes in discharge voltage curves. $R_L$ is measurable at the end of the manufacturing line using ordinary battery test equipment and can be measured within seconds. Changes in $R_L$ are attributed to differences in the amount of lithium consumed to the SEI during formation, where a decrease in $R_L$ indicates that more lithium is consumed. The sensitivity of $R_L$ to lithium consumption is due to the presence of kinetic limitations in the positive electrode causing the total cell resistance to increase at low SOCs. For this reason, $R_L$ provides a particularly strong signature in nickel-rich positive electrode systems where kinetic hindrance plays a strong role in limiting lithium transport toward high states of lithiation. Since the physical interpretation of $R_L$ is general, $R_L$ can be broadly applicable for screening any manufacturing process that impacts the amount of lithium consumed during battery formation. As a whole, our results hold promise for decreasing lithium-ion battery formation time and cost while improving lifetime, as well as identifying rapid diagnostic signals for screening new manufacturing processes and cell designs based on cycle life.

**EXPERIMENTAL PROCEDURES**

**Resource availability**

**Lead contact**

Further information and requests for resources and materials should be directed to and will be fulfilled by Andrew Weng (asweng@umich.edu).

**Materials availability**

All materials are commercially available, with the exception of the carbon methyl cellulose (CMC) binder material used in the anode formulation, which is proprietary.

**Data and code availability**

Data and code used in this study are available at Deep Blue Data: https://doi.org/10.7302/pa3f-4w30. The dataset includes raw time-series files exported using Voltaiq (www.voltaiq.com) and files containing post-processed features. Timeseries data covers formation cycling and aging cycles for all cells used in the study.
Supplementary coin cell data used to generate Figures S21 and S22 have also been included. The source code can be accessed at Zenodo: https://doi.org/10.5281/zenodo.5525258. The source code comprises Python modules used to handle raw data parsing and feature extraction (‘src’/), IPython Notebooks used for data analysis (‘notebooks’/), and MATLAB source files used for the voltage fitting algorithm (‘matlab’).

**Cell build process**

The cathode was comprised of 94:3:3 NMC 111 (TODA North America), C65 conductive additive (Timcal), and polyvinylidene fluoride (PVDF) (Kureha 7208). The slurry was mixed in a step-wise manner, starting with a dry solids homogenization, wetting with n-methyl-2-pyrrolidone (NMP), and then addition of the PVDF resin. The slurry was allowed to mix overnight under static vacuum with agitation from both the double helix blades (30 rpm) and the high-speed disperser blade (1,600 rpm). The final slurry was gravity filtered through a 125 μm paint filter before coating on a roll-to-roll coating machine (Creative & Innovative Systems). The electrode was coated using the reverse comma method at 2 m/min. The final double-sided loading was 34.45 mg/cm².

The anode comprised 97.0:(1.5/1.5) graphite (Hitachi MAG-E3), no conductive additive, and equal parts CMC (proprietary) and styrene butadiene rubber (SBR) (Zeon BM-451B). The graphite and pre-dispersed CMC were mixed prior to further let-down with de-ionized water and overnight dispersion under static vacuum and double helix blade agitation (40 rpm). Prior to coating, the SBR was added and mixed in with helical blade agitation for 15 min under active vacuum. The final slurry was gravity filtered through a 125 μm paint filter before coating on a roll-to-roll coating machine (Creative & Innovative Systems). The electrode was coated using the reverse comma technique at 1.5 m/min. The final double-sided loading was 15.7 mg/cm².

Both anode and cathode were calendared at room temperature to approximately 30% porosity prior to being transferred to a −40°C dew point dry room for final cell assembly and electrolyte filling. The cells, comprising 7 cathodes and 8 anodes, were z-fold stacked, ultrasonically welded, and sealed into formed pouch material (mPlus). The assembled cells were placed in a vacuum oven at 50°C overnight to fully dry prior to electrolyte addition. Approximately 10.5 g of electrolyte (1.0 M lithium hexafluorophosphate (LiPF₆) in 3:7 EC:ethyl methyl carbonate (EMC) v/v + 2wt % vinylene carbonate [VC] from Soulbrain) was manually added to each cell prior to the initial vacuum seal (50 Torr, 5 s). The total mass of all components of the battery is 56.6 ± 0.3g.

The now-wetted cells were each placed under compression between fiberglass plates held in place using spring-loaded bolts. The compression fixtures are designed to allow the gas pouch to protrude and freely expand in the event of gas generation during formation. All cells were allowed to fully wet for 24 h prior to beginning the formation process.

After formation, the cells were removed from the pressure fixtures, returned to the −40°C dew point dry room and degassed. The degassing process was completed in an mPlus degassing machine, automatically piercing the gas pouch, drawing out any generated gas during the final vacuum seal (50 Torr, 5 s), and then placing the final seal on the cell. Cells are manually trimmed to their final dimensions before being returned to their pressure fixtures.
The pouch cell architecture is summarized in Figure S2.

Formation protocols

Figure S1B describes the two different formation protocols used in this study. The fast formation protocol borrows from the "Ultra-fast formation protocol" reported in An et al.\textsuperscript{15} and Wood et al.\textsuperscript{16} In this protocol, the cell is brought to 3.9 V using a 1 C (2.36 Ah) charge, followed by five consecutive charge-discharge cycles between 3.9 and 4.2 V at C/5, and finally ending on a 1 C discharge to 2.5 V. Each charge step terminates on a CV hold until the current falls below C/100. A C/10 charge and C/10 discharge cycle was appended at the end of the test to measure the post-formation cell discharge capacity. A 6-h step was included in between the C/10 charge-discharge steps to monitor the voltage decay. The formation sequence takes 14 h to complete after excluding time taken for diagnostic steps.

A baseline formation protocol was also implemented, which serves as the control for comparing against the performance of fast formation. This protocol consists of three consecutive C/10 charge-discharge cycles between 3.0 and 4.2 V. A 6-h rest was also added between the final C/10 charge-discharge step to monitor the voltage decay signal. The total formation time was 56 h after excluding the diagnostic steps. Formation was conducted at room temperature for all cells and across both formation protocols.

All formation cycling was conducted on a Maccor Series 4000 cycler (0–5 V, 30 mA - 1 A, and auto-ranging). Following formation, one cell (#9) was excluded from this study due to tab weld issues. Consequently, the sample count for the "baseline formation, C/14 cycling group was decreased to 9. The remaining groups had sample counts of 10.

The mean cell energy measured at a 1 C discharge rate from 4.2 to 3.0 V at room temperature is 8.13 Wh. Full cell level volumetric stack energy density is estimated to be 365 Wh/L based on a volume of 69 mm × 101 mm × 71 mm × 3.2 mm, and the gravimetric stack energy density is estimated to be 144 Wh/kg based on a total cell mass of 56.6 g.

Cycle life testing

Following completion of formation cycling, cells were placed in spring-loaded compression fixtures to maintain a uniform stack pressure. Half of the cells from each formation protocol were placed in a thermal chamber (Espec) with a measured temperature of 44°C ± 0.1°C. The remaining cells were left at room temperature and were exposed to varying temperatures throughout the day (24.5°C ± 0.6°C). Long-term cycle life testing was conducted on a Maccor Series 4000 cycler (0–5 V, 10 A, auto-ranging). The cycle life test protocol was identical for all cells and consisted of 1 C (2.37 A), constant current (CC) charge to 4.2 V with a CV hold to 10 mA and 1 C discharges to 3.0 V. At every 50 to 100 cycles, the test was interrupted so that a RPT could be performed.\textsuperscript{41} The RPT consists of a C/3 charge-discharge cycle, a C/20 charge-discharge cycle, followed by the HPPC protocol.\textsuperscript{42} The HPPC test is used to extract 10-s discharge resistance ($R_{10s}$) as a function of SOC (Figure S8).

Every cell was cycled until the discharge capacity was less than 1.18 Ah, corresponding to less than 50% capacity remaining. The total test time varied between 3 to 4 months and the total cycles achieved ranged between 400 and 600 cycles. Cycle test metrics are shown in Figures S3–S6.
**Statistical significance testing**

The standard Student’s t test for two samples was used throughout this study to check if differences in measured outcomes between the two different formation protocols were statistically significant. The p-value was used to quantify the level of marginal significance within the statistical hypothesis test and represents the probability that the null hypothesis is true. A p-value less than 0.05 was used to reject the null hypothesis that the population means are equal. All measured outcomes were assumed to be normally distributed. Box-and-whisker plots are also used throughout the paper to summarize distributions of outcomes. Boxes denote the IQR and whiskers show the minimum and maximum values in the set. No outlier detection methods are employed here due to the small sample sizes ($n < 10$). Finally, the Pearson correlation coefficient, $-1 \leq \rho \leq 1$, was used to determine the significance of correlations between initial state variables and lifetime output variables. $|\rho| > 0.5$ is taken to indicate a statistically meaningful correlation.

**Predictive lifetime model**

Due to the small number of data points available, the model prediction results are sensitive to which cells are chosen for validation. Therefore, we used nested cross-validation to evaluate the regularized linear regression model on all the data without over-fitting. The nested cross-validation algorithm is as follows: first, we separated the data into 20% “validation” and 80% “train/test.” Then, we performed four-fold cross-validation on the “train/test” data to find the optimal regularization strength for ridge regression, $\alpha^*$, using grid search. Finally, we trained the ridge regression algorithm with regularization strength $\alpha^*$, using all of the train/test data, and evaluated the error on the validation data. We repeated this process for 1,000 random train-test/validation splits and reported the mean and standard deviation of the mean percent error for each run:

$$MPE[\%] = \frac{1}{N} \sum_{k=1}^{N} \frac{y_{true}^k - y_{pred}^k}{y_{true}^k}. \quad \text{(Equation 2)}$$

Each run can select a different optimal regularization strength $\alpha^*$.

**Electrode stoichiometry model**

To construct the stoichiometry model shown in Figure 4A, a full cell near-equilibrium potential curve was first extracted using the C/20 charge cycle from the RPT. A randomly selected cell from the 45°C cycling group was selected for this data extraction. Positive and negative electrode near-equilibrium potential curves were adapted from Mohtat et al. The electrode-specific utilization windows are determined by fitting the positive and negative electrode potential curves to match the full cell curve by solving a least squares optimization problem as outlined in Lee et al. The resulting positive and negative electrode alignment minimized the squared error of the modeled versus the measured full cell voltage. The fast formation curve equilibrium potential curve was constructed by shifting the positive electrode curve horizontally and re-computing the full cell voltage curve.

The full cell resistance curves in Figure 4C sourced data from the HPPC sequence as part of the same RPT used to obtain the equilibrium potential curve shown in Figure 4A. A cubic spline fit was used to create smooth resistance curves (a model generated using a linear fit is provided in Figure S24). To break down the resistance contribution into “positive resistance” and “negative + other resistances,” a baseline reference resistance $R_{ref}$ was first defined as the minimum measured full cell resistance below 1Ah. The “negative + other resistances” was then assigned a value
of (1 – \( f_{pos} \)). The remaining resistance was then assigned to the positive electrode. \( f_{pos} \) was set to 0.7 to model a generic NMC/graphite system.\(^{15,60,61}\)

**Voltage fitting algorithm**

Methods for estimating electrode-specific state of health metrics using half-cell reference curves have been previously reported.\(^{52-54}\) Here, we applied an automated voltage fitting approach based on work by Lee et al.\(^{55}\) to extract electrode capacity losses, \( \text{LAM}_{PE} \) and \( \text{LAM}_{NE} \), as well as lithium inventory loss (LLI) for both fresh and aged cells. The input data consisted of C/20 charge curves measured at each RPT. An example set of C/20 charge curves over age is shown in Figure S16.

The method to extract electrode-specific state of health indicators LLI, \( \text{LAM}_{PE} \), and \( \text{LAM}_{NE} \) is adapted from Lee et al.\(^{55}\) Positive and negative near-equilibrium potential curves were adapted from Mohtat et al.\(^{29}\) The curves were obtained at the C/20 rate and served as proxies for the true equilibrium potential curves. The same equilibrium potential curves were used to model data at both test temperatures.

To prevent over-fitting, the positive electrode stoichiometry at 100% SOC (\( y_{100} \)) was fixed to 0.03 at every instance for this analysis. Fixing this value yielded smoother and more physical degradation trajectories over cycle life. Figure S17 shows an example of voltage fitting results for a single cell. The degradation metrics, including LLI and LAM were computed in the usual manner (see Lee et al.\(^{55}\) for more details).

**Hybrid pulse power characterization of half cells**

Coin cell half cells were built with LFP, NMC111, and graphite as the working electrode and lithium metal as the counter electrode. The NMC material used were identical to that used in the pouch cells for the formation experiments (TODA North America). The graphite material used differed from the ones used in the pouch cells. The coin cell construction consisted of 2032 form factor components including a wavespring and spacer. The electrolyte used was 1 M LiPF\(_6\) with EC/EMC. The lithium counter electrode was 16 mm in diameter, the separator was 19 mm in diameter, and the working electrodes were 14 mm in diameter. Working electrodes were measured to be approximately 60 \( \mu \)m thick and the lithium counter electrodes were approximately 750 \( \mu \)m thick. Working electrodes were single-side coated. Calculated theoretical capacities for the NMC111, LFP, and graphite cells were 2.0, 2.9, and 4.6 mAh, respectively.

The HPPC protocol was adapted for the coin cells. Potential ranges were modified depending on the working electrode. The currents used in the pulses were also scaled down to 0.4 mA for all cells (Figure S21). The measured resistance drop includes a large Ohmic contribution due to the presence of the lithium metal counter electrode. However, since this counter electrode was present in all cells, differences in measured, SOC-dependent resistances between the different cells remain meaningful. All coin cells were pre-conditioned using at least three slow charge-discharge cycles prior to starting the HPPC sequence.

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.joule.2021.09.015.

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**AUTHOR CONTRIBUTIONS**

Conceptualization, A.W. and A.S.; methodology, A.W., P.M., G.L., P.M.A., V.S., and S.L.; investigation, A.W. and G.L.; data curation, A.W.; software, A.W.; visualization, A.W.; formal analysis, V.S.; writing – original draft, A.W.; writing – review & editing, A.W., P.M.A., and A.S.; funding acquisition, A.S.

**DECLARATION OF INTERESTS**

A.W. and P.M.A. are employees of Tesla. A patent application relating to this work has been filed. The authors declare no other competing interests.

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