Electrochemical Model-Based State of Charge and Capacity Estimation for a Composite Electrode Lithium-Ion Battery

Alexander Bartlett, James Marcicki, Simona Onori, Giorgio Rizzoni, Xiao Guang Yang, and Ted Miller

Abstract—Increased demand for hybrid and electric vehicles has motivated research to improve onboard state of charge (SOC) and state of health estimation (SOH). In particular, batteries with composite electrodes have become popular for automotive applications due to their ability to balance energy density, power density, and cost by adjusting the amount of each material within the electrode. SOH algorithms that do not use electrochemical-based models may have more difficulty maintaining an accurate battery model as the cell ages under varying degradation modes, such as lithium consumption at the solid-electrolyte interface or active material dissolution. Furthermore, efforts to validate electrochemical model-based state estimation algorithms with experimental aging data are limited, particularly for composite electrode cells. In this paper, we first present a reduced-order electrochemical model for a composite LiMn2O4-LiNi1/3Mn1/3Co1/3O2 electrode battery that predicts the surface and bulk lithium concentration of each material in the composite electrode, as well as the current split between each material. The model is then used in dual-nonlinear observers to estimate the cell SOC and loss of cyclable lithium over time. Three different observer types are compared: 1) the extended Kalman filter; 2) fixed interval Kalman smoother; and 3) particle filter. Finally, an experimental aging campaign is used to compare the estimated capacities for five different cells with the measured capacities over time.

Index Terms—Battery, estimation, Kalman filter, state of charge (SOC), state of health (SOH).

NOMENCLATURE

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A$</td>
<td>Current collector area (m²)</td>
</tr>
<tr>
<td>$D_e$</td>
<td>Liquid diffusion coefficient (m²/s)</td>
</tr>
<tr>
<td>$D_i$</td>
<td>Solid diffusion coefficient (m²/s)</td>
</tr>
<tr>
<td>$F$</td>
<td>Faraday’s constant (C/mol)</td>
</tr>
<tr>
<td>$I$</td>
<td>Current (A)</td>
</tr>
<tr>
<td>$L_c$</td>
<td>Total cell thickness (m)</td>
</tr>
<tr>
<td>$L_i$</td>
<td>Electrode thickness (m)</td>
</tr>
<tr>
<td>$L_{sep}$</td>
<td>Separator thickness (m)</td>
</tr>
<tr>
<td>$N_p$</td>
<td>Number of particles in the particle filter</td>
</tr>
<tr>
<td>$Q$</td>
<td>Process noise covariance</td>
</tr>
<tr>
<td>$R$</td>
<td>Sensor noise covariance</td>
</tr>
<tr>
<td>$\tilde{R}$</td>
<td>Universal gas constant (J/K/mol)</td>
</tr>
<tr>
<td>$R_{c,i}$</td>
<td>Ohmic resistance (Ω)</td>
</tr>
<tr>
<td>$R_i$</td>
<td>Particle radius (m)</td>
</tr>
<tr>
<td>$T$</td>
<td>Cell temperature (K)</td>
</tr>
<tr>
<td>$U_i$</td>
<td>Electrode open-circuit voltage (V)</td>
</tr>
<tr>
<td>$V$</td>
<td>Cell voltage (V)</td>
</tr>
<tr>
<td>$V_e$</td>
<td>Liquid-phase voltage across the cell (V)</td>
</tr>
<tr>
<td>$\bar{c}_i$</td>
<td>Bulk particle concentration (mol/m³)</td>
</tr>
<tr>
<td>$c_{e,i}$</td>
<td>Electrolyte concentration (mol/m³)</td>
</tr>
<tr>
<td>$c_{\text{max},i}$</td>
<td>Electrode saturation concentration (mol/m³)</td>
</tr>
<tr>
<td>$c_{s,i}$</td>
<td>Concentration at the particle surface (mol/m³)</td>
</tr>
<tr>
<td>$i_{0,i}$</td>
<td>Exchange current density (A/m²)</td>
</tr>
<tr>
<td>$i_e$</td>
<td>Liquid-phase current density (A/m²)</td>
</tr>
<tr>
<td>$j_i$</td>
<td>Solid-phase current density (A/m²)</td>
</tr>
<tr>
<td>$k_i$</td>
<td>Reaction rate constant (m²/mol s)</td>
</tr>
<tr>
<td>$r$</td>
<td>Coordinate along the particle radius, originating at the particle center</td>
</tr>
<tr>
<td>$t$</td>
<td>Time (s)</td>
</tr>
<tr>
<td>$t_0$</td>
<td>Transference number</td>
</tr>
<tr>
<td>$x$</td>
<td>Coordinate through-the-thickness of the cell, originating at the anode current collector</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Symmetry factor</td>
</tr>
<tr>
<td>$\beta_i$</td>
<td>Current split factor</td>
</tr>
<tr>
<td>$\gamma$</td>
<td>Activity coefficient in electrolyte</td>
</tr>
<tr>
<td>$\epsilon_{am,i}$</td>
<td>Active material volume fraction</td>
</tr>
<tr>
<td>$\epsilon_e$</td>
<td>Liquid volume fraction</td>
</tr>
<tr>
<td>$\epsilon_i$</td>
<td>Active material volume split</td>
</tr>
<tr>
<td>$\eta_i$</td>
<td>Kinetic overpotential (V)</td>
</tr>
<tr>
<td>$\sigma_e$</td>
<td>Electrolyte conductivity (S/m)</td>
</tr>
<tr>
<td>$\phi_e$</td>
<td>Electrolyte potential (V)</td>
</tr>
<tr>
<td>$\phi_l$</td>
<td>Electrode potential (V)</td>
</tr>
</tbody>
</table>

Subscript $i$ refers to electrode $p$, $n$, LM0, or NMC.

I. INTRODUCTION

VEHICLE electrification continues to be a key topic of interest for automotive manufacturers, with lithium-ion batteries being the technology of choice for hybrid and electric vehicles. State of charge (SOC) and state of health (SOH) estimates are essential inputs to the vehicle’s battery management system (BMS) in order for the battery pack to operate...
efficiently and safely [1]. Knowledge of SOC allows the BMS to predict the available instantaneous power, while ensuring the battery is operating within safe limits. SOH can be defined in a variety of ways, but typically refers to the degradation of capacity and increase in internal resistance as the battery ages.

Traditionally, SOC and SOH estimates are made with the use of lookup tables or equivalent circuit models (ECMs) consisting of various circuit elements arranged in series or parallel combinations such that the circuit output voltage dynamics mimic that of the battery. Although ECMs are computationally inexpensive, considerable effort must be made to parameterize these models, as the parameters are typically the functions of SOC, temperature, current direction, and battery life [2], [3]. In addition, ECM parameters have little physical meaning, and associated aging models must rely on empirical correlations between SOH and damage factors, such as total current throughput, operating temperature, and depth of discharge [4]. Nonetheless, ECMs have been used successfully for SOC and SOH estimation using extended Kalman filters (EKFs) and other methods [5]–[10].

Recently, increased attention has been given to electrochemical models for SOC and SOH estimation. These models are based on the first principles in porous electrode theory [11] and therefore have the potential to predict cell performance more accurately, as well as provide more information about the internal battery states such as lithium concentrations and reaction overpotentials. However, uncertainty in electrochemical parameters can limit the accuracy of the model and resulting state estimates [22]. Furthermore, electrochemical models rely on partial differential equations (PDEs) to describe lithium diffusion and potential gradients throughout the electrode and electrolyte. Solution methods for PDEs can be computationally expensive, making them ill-suited for most practical onboard control and estimation algorithms. A common approximation, known as the single particle model (SPM), is to neglect the spatial variation of concentration and potential throughout the cell and treat each electrode as a single spherical particle, subject to an appropriately scaled, spatially constant current flux. To further facilitate control and estimation, researchers have used various model-order reduction techniques to reduce the PDEs governing diffusion in the spherical particles to low-order ordinary differential equations (ODEs). A reduced-order electrochemical model linearized over a typical operating range was used in [13] in a Kalman filter to estimate the lithium bulk and surface concentration. The authors showed that rather than enforcing strict limits on cell voltage to avoid damaging the battery, enforcing limits on lithium surface concentration and solid/electrolyte potential difference allowed the cell voltage limits to be exceeded without risking electrode saturation/depletion and lithium plating. In [14], an averaged SPM with discretization of the diffusion PDEs was used in a Kalman filter to estimate the bulk and surface concentrations of the positive electrode. The authors recognized weak observability of the complete cell model that includes both electrodes (since the only available measurement depends on the difference between the electrode potentials), and therefore only applied the observer to the positive electrode states. Loss of spatial information through-the-thickness of the cell is avoided in [2], [16], and [17] using observers capable of handling PDE-based models, although these approaches add to the computational cost. Moura et al. [17] also discussed the weak observability of the complete battery system, and approximated the cathode states at their equilibrium, since the average positive concentration can be related to the average negative concentration via conservation of lithium.

There have been a few attempts to use electrochemical-based models for online SOH estimation using dual-SOC/parameter estimation. In [18], the active material fractions of each electrode are estimated along with the bulk and surface SOC, using an unscented Kalman filter. SOH estimation was done in [15], [17], and [19] by applying least squares techniques to estimate the model parameters associated with aging; however, the computational effort required by these approaches may limit their applicability to onboard estimation. In addition, electrochemical model-based SOH estimation algorithms that are validated with experimental aging data are scarce in the literature, and to the best of the authors’ knowledge, no such studies have been conducted on cells with composite electrodes. Composite electrodes add to the complexity of the electrochemical model, particularly if it is desired to estimate the SOC and SOH of each electrode material individually.

In this paper, a reduced order electrochemical model is used for SOC/SOH estimation on a composite LiMn$_2$O$_4$-LiNi$_{1/3}$Mn$_{1/3}$Co$_{1/3}$O$_2$ (LMO-NMC) cathode cell with a graphite anode. The model is based on the SPM with added liquid-phase dynamics, but considers two particles in parallel to represent the composite cathode. This paper then presents a novel numerically inexpensive method for solving for the current split and resulting SOC of each cathode particle (part of preliminary work presented in [20]), and a novel combined SOC/SOH estimation algorithm to estimate capacity fade resulting from loss of cyclable Li. Finally, the estimation algorithm is validated with experimental aging data, by comparing three different observer types (EKF, Kalman smoother, and particle filter).

II. MODEL DEVELOPMENT

The model used in this paper is based on the traditional SPM with the addition of liquid-phase diffusion dynamics to improve model accuracy under high current. Similar models have been presented many times in [2], [13], [14], [16], [18], and [23], however, in order to accommodate the composite LMO-NMC positive electrode, the positive electrode is modeled as two particles acting in parallel (as shown in Fig. 1). Therefore, the potentials of each positive electrode particle model are equal, but the current is split between each particle according to its dynamic effective impedance. This idea of representing a composite electrode as two particles acting in parallel has been presented in [20] and [23].

In the simplest case (at steady state), the measured cell voltage $V$ is just the difference between each electrode open-circuit voltage (OCV) evaluated at their respective steady-state concentrations, i.e., $V = U_p(\bar{c}_p) - U_n(\bar{c}_n)$. However, as the
cell is charged or discharged, various sources of overpotential cause the measured voltage to deviate from this steady-state solution. The overpotentials can be categorized as acting in either the solid (electrode) or liquid (electrolyte) phase, as outlined in Table I, and submodels for each phase can be developed. The dynamic cell voltage (9) is then obtained by subtracting the solid- and liquid-phase overpotentials from the electrode OCVs.

A. Electrode Submodel

The governing equations for Li concentration and electrode potential are shown in (1)–(5) for a given particle $i$. Subscript $i$ refers to the negative particle, positive LMO particle, or positive NMC particle: n, LMO, or NMC, respectively. Fick’s law of mass diffusion in spherical coordinates (4) governs the concentration throughout a spherical particle. The boundary conditions for this PDE are set so as to require symmetry about the particle center and impose a flux at the particle surface, dictated by the current density $j_i(t)$.

The single particle approximation removes the spatial dependence of the current density, allowing it to be defined only as a function of time for a given particle. To form the current density, the applied current is multiplied by a current split factor $\beta_i$ defined as the fraction of current flowing to a given particle within an electrode. For the single particle negative electrode $\beta_n = 1$, but for the multiple particle positive electrode, $\beta_{LMO}$ and $\beta_{NMC}$ vary dynamically and are calculated at each time step, as shown in Section III-A. The current is normalized by the electrode volume, including a parameter $\epsilon_i$ representing the active material volume fraction split. Again, for the single particle negative electrode $\epsilon_n = 1$; however, for the multiple particle positive electrode, fitting experimental data gave a material split of $\epsilon_{LMO} = 0.64$ and $\epsilon_{NMC} = 0.36$ [23].

The electrode potential (5) is calculated by subtracting the kinetic and ohmic overpotentials from the OCV, evaluated at the particle surface concentration.

B. Electrolyte Submodel

Similar to the solid phase, the liquid phase contains sources of concentration and ohmic overpotential. The concentration of Li throughout the electrolyte is governed by Fick’s law in Cartesian coordinates (6), with a source term related to the current density. Since $j_i(t)$ is considered piecewise constant within a given electrode and zero in the separator, (6) is solved separately in each region, subject to coupling boundary conditions that match the concentration and flux at the electrode/separator interfaces.

To obtain the liquid-phase voltage across the entire cell, (7) is integrated directly from $x = 0$ to $x = L_r$, where $i_x(t)$ is considered constant in the separator and linearly decreasing to zero from the electrode/separator boundaries to the current collectors.

III. Model Solution Method

The concentrations of Li in the solid and liquid phases are governed by PDEs; however, this is undesirable if the model is to be used for onboard estimation. Instead, the Padé approximation method for model-order reduction proposed in [22] and [24] was leveraged here to approximate the PDE-based model with a system of low-order ODEs.

A. Solid-Phase Solution

In the solid phase, the important metrics for tracking cell SOC and available power are the particle bulk concentration and particle surface concentration, respectively. A first-order linear transfer function between the bulk concentration and input current is obtained simply by integrating (4) in the Laplace domain from $r = 0$ to $r = R_i$. However, in the case of surface concentrations, a Padé approximation is used to obtain a low-order linear transfer function relating surface concentration to input current for each particle [12], [24]. A third-order approximation is used for the solid-phase PDE, since it gives good accuracy in the frequency range typically seen in a vehicle drive profile [23], while still mitigating the computational effort required for higher order models.

The electrode particle submodel is formulated in state space as follows:

$$\dot{x}_i = A_i x_i + B_i(t) u$$

$$A_i = \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & b_{3,i} \end{bmatrix}$$

$$B_i(t) = \begin{bmatrix} \frac{\beta_i(t) R_i}{3F \ell_{am,i} \ell_i} \\ 0 \\ 0 \end{bmatrix}$$

$$\phi_i = U_i(c_{s,i}) \pm \eta_i(c_{s,i}, \beta_i(t) u) \pm R_{c,i} \beta_i(t) u$$

where both the bulk and surface concentrations are written as linear combinations of the states $x_i$, the input $u$ is current, and the output electrode potential is a nonlinear function of the states and input. Note that only the linear combination of the states has any physical meaning, and the states themselves simply represent the dynamics of the third-order Padé approximation. Although there are two separate transfer functions for bulk and surface concentration (first and third orders, respectively), both metrics can be written as linear combinations of the same three states through some algebraic manipulation (as shown in [24]). In other words, there are six total states in the multiple particle positive electrode submodel, and three states in the single particle negative electrode submodel.
The states are updated in discrete time using a finite difference solution. Constants \(a_{0,i}, a_{1,i}, a_{2,i}, b_{2,i}, \) and \(b_{3,i}\) are obtained from the third-order Padé approximation, and are functions of the particle radius and diffusion coefficient [12], [24].

Since the positive electrode consists of two particles acting in parallel, the potentials for each particle are equal and the current is multiplied by a current split factor \(\beta\). For the positive electrode, \(\beta\) must be solved at each time step to determine how the current is allocated with the constraint that the currents going to each particle sum to the total current. \(\beta\) appears nonlinearly in the output equation, so one possible solution method is to remove the nonlinear dependence on \(\beta\) in \(f_1\) and \(f_2\) by taking a Taylor series expansion about the current splits at the previous time step.

3) Remove the nonlinear dependence on \(\beta\) in \(f_1\) and \(f_2\) by taking a Taylor series expansion about the current splits at the previous time step.

\[
\phi_{p,k} \approx f_1|_{\beta_{LMO, k-1}} + \frac{\partial f_1}{\partial \beta_{LMO}}|_{\beta_{LMO, k-1}} (\beta_{LMO, k} - \beta_{LMO, k-1})
\]

\[
\phi_{p,k} \approx f_2|_{\beta_{NMC, k-1}} + \frac{\partial f_2}{\partial \beta_{NMC}}|_{\beta_{NMC, k-1}} (\beta_{NMC, k} - \beta_{NMC, k-1}).
\]

4) Finally, the current splits and potential can be written as the solution to a matrix inversion problem.

\[
\begin{bmatrix}
\frac{\partial f_1}{\partial \beta_{LMO}}|_{\beta_{LMO, k-1}} & 0 & -1 \\
0 & \frac{\partial f_2}{\partial \beta_{NMC}}|_{\beta_{NMC, k-1}} & -1 \\
I_k & I_k & 0
\end{bmatrix}
\begin{bmatrix}
\beta_{LMO, k} \\
\beta_{NMC, k} \\
\phi_{p,k}
\end{bmatrix}
= \begin{bmatrix}
-f_1|_{\beta_{LMO, k-1}} + \frac{\partial f_1}{\partial \beta_{LMO}}|_{\beta_{LMO, k-1}} (\beta_{LMO, k} - \beta_{LMO, k-1}) \\
-f_2|_{\beta_{NMC, k-1}} + \frac{\partial f_2}{\partial \beta_{NMC}}|_{\beta_{NMC, k-1}} (\beta_{NMC, k} - \beta_{NMC, k-1})
\end{bmatrix}.
\]

In addition to the current splits, solving (15) also gives a value for the linearized output electrode potential \(\phi_{p,k}\); however, this value is discarded at this point. Instead, the original nonlinear output (11) is used to calculate the electrode potential, thereby avoiding the voltage prediction inaccuracies seen with the linearization process. In this way, the errors associated with the output linearization only affect the calculation of \(\beta\), and not the calculation of the output voltage (apart from the indirect effect from \(\beta\)).

Fig. 2 shows the predicted current split and resulting SOC for each cathode material for a constant 1C discharge. The predicted voltage shows good agreement with the measured voltage (more rigorous validation of the model was done in [23]). More LMO is utilized initially at high voltages, until it is nearly depleted, at which point
β is near zero, so a large β magnitude and/or direction, the calculated is that, for instances, when the current abruptly changes in cathode [25].

X-ray diffraction (XRD) study of a composite LMO-NMC in situ the discharge. This result is consistent with an

One drawback of this electrode model with parallel particles is that, for instance, when the current abruptly changes in magnitude and/or direction, the calculated βi for a given particle can greatly exceed 1 (resulting in a large negative βi for the other particle). While it is unclear if this occurrence makes physical sense, it typically lasts for only a few time steps before the βi's return to more sensible values between 0 and 1. Furthermore, these are usually instances when the total current is near zero, so a large βi does not translate into a large current going to a given particle.

It should also be noted that if the state estimate has not yet converged, there will be some error in the calculation of βi, since it relies on the current state estimate. This would in turn cause an error in the state estimate calculated at the next time step. However, this does not affect the stability of the state estimate. Even if there is an extreme error in βi, eventually the concentration estimate of one of the materials will become saturated, which will decrease βi for that material, thereby driving the estimate back toward the correct value. Instead, calculation of βi using an unconverged state estimate can be categorized as a model error, which may affect the estimate accuracy, but will not cause the estimate to become unstable. Furthermore, in a real automotive application, there are periodic instances when the current is near zero, and the influence of βi is negligible. During these times, the state estimate can converge accurately, thereby minimizing the error in βi once the current resumes.

To summarize the solution method for the composite positive electrode: at each time step k, the states are updated in discrete time using the calculated values for βi,k−1 at the previous time step (10), the values for βi,k at the current time step are then calculated using the linearization method outlined above (15), and finally the output potentials are calculated using the updated states and precalculated values for βi,k (11).

B. Liquid-Phase Solution

Equation (8) governs the liquid-phase voltage across the cell and depends on the Li-ion concentrations in the liquid at each electrode. Therefore, the PDE governing Li concentration is solved at each electrode end (x = 0 and x = L_e) using a first-order Padé approximation. This results in two first-order transfer functions relating liquid-phase concentration at each electrode to the current input. Only a first-order approximation is needed for the liquid phase concentration, since it shows a very good accuracy throughout the frequency range of interest [22]. In state space, the two states in xe represent the liquid concentrations at each electrode, the input u is current, and the output voltage is a nonlinear function of the states and input

\[
\dot{x}_e = A_e x_e + B_e u
\]

\[
A_e = \begin{bmatrix}
1 & 0 \\
\frac{b_{1,e,p}}{b_{1,e,n}} & 1
\end{bmatrix}
\]

\[
B_e = \begin{bmatrix}
\frac{1 - \frac{t_0^+}{F A L e \epsilon_e}}{b_{2,e,p}} \\
\frac{1}{b_{2,e,n}}
\end{bmatrix}
\]

\[
\begin{bmatrix}
\bar{c}_{e,p} \\
\bar{c}_{e,n}
\end{bmatrix} = \begin{bmatrix} 1 & 1 \end{bmatrix} x_e
\]

\[
V_e = u \frac{\frac{1}{2} L_p + L_{sep} + \frac{1}{2} L_n}{\sigma_e A} + \frac{2 R T (1 - \frac{t_0^+}{\epsilon_e})}{F} (1 + \gamma) \ln \left( \frac{\bar{c}_{e,n}}{\bar{c}_{e,p}} \right)
\]  \hspace{1cm} (17)

Again, the constants b_{1,e,p}, b_{1,e,n}, b_{2,e,p}, and b_{2,e,n} are obtained from the first-order Padé approximation [12].

IV. SYSTEM OBSERVABILITY

In order to design an observer, the system must be shown to be observable (or at least detectable) to ensure that the state estimates converge to the true states in a finite time [26]. Observability of a system requires that the state trajectory can be uniquely reconstructed based solely on the knowledge of the output measurements and inputs. Local observability of a nonlinear system can be shown with the use of Lie derivatives [26]. The Lie derivatives L_f for a nonlinear system with state dynamics \( \dot{x} = f(x, u) \) and output equation \( y = g(x, u) \) form the vector

\[
I(x, u) = \begin{bmatrix}
y \\
\dot{y} \\
\vdots \\
y^{(n-1)}
\end{bmatrix} = \begin{bmatrix}
g \\
\dot{g} \\
\vdots \\
g^{(n-1)}
\end{bmatrix} = \begin{bmatrix}
L_0^f \\
L_1^f \\
\vdots \\
L_{n-1}^f
\end{bmatrix}
\]  \hspace{1cm} (18)
The Lie derivative vector is obtained by taking the Jacobian of the Lie derivative vector

\[
\mathcal{O} = \begin{bmatrix}
\frac{\partial L^0}{\partial x_1} & \cdots & \frac{\partial L^n}{\partial x_n} \\
\vdots & \ddots & \vdots \\
\frac{\partial L^{n-1}}{\partial x_1} & \cdots & \frac{\partial L^{n-1}}{\partial x_n}
\end{bmatrix}
\]  

(19)

An observability matrix with full rank implies that the states are locally observable. However, performing this exercise on the complete battery model, including both positive and negative electrodes, results in an observability matrix with full rank but with a high (two-norm) condition number (\(\approx 10^{10}\)) for typical parameter and input values. This is an intuitive result, as it should be difficult to uniquely identify the states of both electrodes if the only available measurement is the difference between positive and negative electrode potentials.

To help facilitate observer design, the degree of observability can be improved using one of the two methods. In the first method, the states of only one electrode are estimated at a given time, while the liquid phase and opposite electrode states are simulated open loop. For example, if the positive electrode states are to be estimated, the negative electrode and liquid-phase potentials are calculated using the open-loop model. These potentials are subtracted from the measured cell voltage to obtain a pseudomeasurement of positive electrode potential. This pseudomeasurement is then used as feedback in the observer to estimate the positive electrode states. The process is outlined in Fig. 3 for estimating the positive electrode states. However, clearly, since the liquid phase and one electrode are simulated open loop, any initial condition and model errors in these submodels will be projected forward to adversely affect the opposite electrode estimate.

An alternative to relying on an open-loop simulation of one electrode is to use a steady-state approach, where the bulk concentration of one electrode is related to the bulk concentration of the other, via conservation of lithium

\[
\frac{c_{\text{max},p}}{c_{\text{max},p} - x_f} - y_f = \frac{c_{\text{max},n}}{c_{\text{max},n} - x_0} - y_0
\]

(20)

where \(y_0\) to \(y_f\) is the normalized concentration operating range of the positive electrode and \(x_0\) to \(x_f\) is the operating range of the negative electrode, as shown in Fig. 6. Note that \(x_0\) and \(y_0\) are the normalized concentrations at the upper voltage limit, while \(x_f\) and \(y_f\) correspond to the lower voltage limit. This steady-state approach was applied in [14] and [17] to estimate the cell SOC. The advantage of this method is that it enforces conservation of lithium in the cell, which is a good approximation over a given drive cycle. However, this approach neglects the diffusion dynamics of one electrode, which may have implications in estimation of available power. In addition, like the open-loop method, it relies on accurate initialization of the normalized concentration operating range of each electrode, so any errors in the \(x_0\), \(x_f\), \(y_0\), and \(y_f\) parameters will adversely affect the state estimates.

Both the open-loop and steady-state approaches essentially limit the estimated states to one electrode, and by doing so, the condition number of the observability matrix is reduced by several orders of magnitude (down to \(\approx 10^6\)). In this paper, results using the open-loop approach are shown, although testing of both methods revealed only slight differences in the SOC and capacity estimation results.

Even though the estimated states are limited to just one electrode whose electrode submodel must be shown to be observable with respect to a pseudomeasurement of its output potential. The electrode submodel has linear state dynamics (10) with a nonlinear output (11). The goal is to show that all three states in the submodel are observable, allowing for the estimation of both the surface and bulk electrode concentrations \(c_{s,i}\) and \(c_{\text{max},i}\). The OCV term \(U_i\) is often calculated via lookup table or by an empirical functional fit; however, to make calculating the Lie derivatives easier computationally, a cubic spline is used to evaluate the OCV at a given concentration. The cubic spline is continuously differentiable at the spline transitions up to the second derivative, so proving the observability for a given spline segment guarantees observability for the entire OCV curve. Substituting an arbitrary cubic spline segment for the OCV term, the electrode submodel becomes

\[
\dot{x}_i = f(x_i, u) = A_i x_i + B_i(t) u
\]

\[
y = g(x_i, u) = \frac{s_{3,i} c_{s,i}^3}{c_{\text{max},i}^3} + \frac{s_{2,i} c_{s,i}^2}{c_{\text{max},i}^2} + \frac{s_{1,i} c_{s,i}}{c_{\text{max},i}} + s_{0,i}
\]

\[
-\eta_i(c_{s,i}^3, \beta_i u) - R_i \beta_i u
\]

(22)

where \(s_{0,i}, s_{1,i}, s_{2,i}, s_{3,i}\) are the OCV cubic spline coefficients, and \(c_{s,i}\) is a linear function of the states, as shown in (10).

The resulting Lie derivatives and observability matrix are too lengthy to show here, in particular due to differentiating the hyperbolic sine function in \(\eta_i\), but the process is outlined in the Appendix. The observability matrix has full rank for nonzero parameter values, proving that the electrode particle submodel is locally observable with respect to a pseudomeasurement of its potential. Although the both positive and negative electrodes are observable, the negative electrode states are more sensitive to model or sensor errors. This is due to the flat shape of the negative electrode OCV in certain concentration ranges, where a small error in voltage can produce a large error in estimated concentration. The implications of the flat negative OCV will be discussed in Section VII-B.
V. EXPERIMENTAL DATA COLLECTION

An aging campaign was conducted using 15-Ah (nominal) automotive pouch cells with a composite LMO-NMC cathode and graphite anode cycled under different operating conditions. The cells were cycled using the charge depleting (CD) current profile defined by the United States Advanced Battery Consortium [27], which is representative of a plug-in hybrid vehicle (PHEV) application. The CD current profile is shown in Fig. 10, and was repeated until the cell reached 45%, 35%, or 25% SOC. Each CD cycle was followed by a constant current constant voltage (CCCV) charge at a C/3, 3C/2, or 5C charge rate. All cells were maintained at a constant of 30 °C by Peltier junctions. Periodic capacity assessments were conducted at approximately every 2000–5000 Ah of throughput. The capacity assessments involved an initial 1C CCCV charge to 4.15 V, followed by a 1C discharge to 2.8 V, followed by a second 1C CCCV charge. The measured capacity was taken as the average of the discharge and charge capacities. The electrode OCVs (as shown in Figs. 5 and 8) and the parameter values from [23] were used to calibrate the model.

An example of the open-loop model fit for the dynamic CD profile used in the experimental aging campaign is shown in Fig. 4. The root-mean-squared error over this test is 14 mV with a maximum error of 75 mV.

It should be noted that no significant resistance rise was observed in the cells used in this paper, allowing the impedance-related model parameters to be treated as constant. The change in 1-s resistance was generally less than 0.1 mΩ over the course of the aging campaign. However, for different aging conditions or electrode chemistries, resistance rise can be significant and online estimation of these parameters may be required to maintain model accuracy.

VI. OBSERVER DESIGN

Three different observer types are used for state estimation in order to compare their estimation accuracy and computational effort. Specifically, the EKF, fixed interval Kalman smoother, and particle filter are implemented and compared. The implementation of each observer can be found in the literature, but brief descriptions will be given here.

The EKF [5], [6], [30] is an extension of the Kalman filter, optimal for linear systems, to nonlinear systems where the process and sensor noise are assumed to be uncorrelated, white, and Gaussian. As is typical of model-based observers, the EKF involves both a prediction step and a correction step. In the prediction step, open-loop predictions of the state vector and output are obtained from the model. The state prediction is then adjusted in the correction step by utilizing feedback from the measurement. The feedback gain (optimal for a linear system) is calculated at each time step, based on the knowledge of the model and noise covariance matrices. For nonlinear systems, calculation of the observer gain relies on a linearization of the nonlinear system dynamics and output equation. In practice, the process and sensor noise covariance matrices are not necessarily known, and there may be significant model errors. Therefore, the noise covariances are typically used as tuning factors to achieve the desired observer behavior for a particular system.

The fixed interval Kalman smoother (or forward–backward smoother) has the potential to improve upon the traditional EKF by essentially running the filter twice, once forward in time and once backward in time, over a fixed interval of measurements [30], [35]. Once both forward and backward estimates are obtained, they are combined in an optimal way to minimize the estimation error covariance. The resulting smoothed estimate has a smaller estimate error covariance than either of the forward or backward estimates individually. The obvious disadvantage of this method, other than the added computational effort, is that the estimation can no longer be performed in real time. It is necessary to wait until all the measurements within the interval become available before

---

**Fig. 4.** Model prediction of the cell voltage compared with the measured voltage for the CD current profile used in the aging campaign, as shown in Fig. 10(a).

**Fig. 5.** Initial and final concentrations are defined at the upper and lower cell voltage limits. The overall initial and final normalized concentrations $y_0$ and $y_f$ correspond to a cell SOC (top axis) of 1 and 0, respectively.
TABLE II
TUNING PARAMETERS FOR EACH OBSERVER TYPE

<table>
<thead>
<tr>
<th>Observer</th>
<th>Parameter</th>
<th>C_0</th>
<th>C_1</th>
<th>C_2</th>
<th>C_3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive electrode</td>
<td>state estimation</td>
<td>1500</td>
<td>0</td>
<td>1.5</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
<td>0.0015</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Q = 10^{-18}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R = 2.24 \times 10^{-4}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Negative electrode</td>
<td>state estimation</td>
<td>2500</td>
<td>0</td>
<td>2.5</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
<td>0.0025</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Q = 10^{-17}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R = 2.45 \times 10^{-5}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kalman Smoother</td>
<td></td>
<td>2500</td>
<td>0</td>
<td>2.5</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
<td>0.0025</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Q = 10^{-17}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R = 2.45 \times 10^{-5}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Particle Filter**</td>
<td></td>
<td>1200</td>
<td>0</td>
<td>1.2</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
<td>0.0012</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Q = 10^{-15}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R = 2.45 \times 10^{-5}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N_p = 10</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*\Sigma_0^p and \Sigma_0^s are identical for LMO and NMC particles.
**Gaussian sensor and process noise are assumed.

even the initial smoothed state estimate can be obtained. However, for SOH estimation this is not a significant problem, since in practice aging parameters change very slowly and need not be continuously updated in real time.

The particle filter is a Monte Carlo method used for state estimation by representing the posterior probability density function (pdf), i.e., the conditional pdf of the state given a measurement, by a set of weighted random samples or particles [30]–[32]. Each particle is assigned a weight based on the likelihood that it represents the true state. The advantage of the particle filter is that it does not require linearization of the nonlinear model, nor does it require that the process and sensor noise be Gaussian. Choosing the number of particles \( N_p \) represents a tradeoff between estimate accuracy and computational effort.

The relevant parameters for each observer are outlined in Table II, and were tuned to achieve a desired compromise between model error robustness, state convergence time, and computational effort.

VII. METHODS FOR SOC/SOH ESTIMATION

A. Estimating Cell SOC

Estimating the states of one of the electrode particle submodels gives an estimate of both bulk and surface concentrations of the particle; however, without the use of \textit{in situ} XRD or similar techniques, it is only possible to experimentally validate the bulk concentration estimate by comparing it against Coulomb counting, or integrating the measured current over time

\[
\text{SOC}_{\text{exp}} = \text{SOC}_0 - \frac{1}{\Delta v_{\text{OCV}}} \int_{t_0}^{t_f} I(t) \, dt. \tag{23}
\]

The initial \( \text{SOC}_0 \) is taken from a cell level OCV–SOC lookup table and the cell capacity \( C \) is obtained from the experimental capacity in ampere-hours measured between the two voltage limits. In this paper, the positive electrode state estimates are validated with Coulomb counting because the positive electrode is typically the power limiting electrode. In addition, the positive electrode OCV is steeper than the negative in most concentration ranges, so the positive state estimates are generally less sensitive to voltage errors. Since the observer estimates the electrode concentrations of each composite material (ranging from 0 to the saturation concentration of the electrode material), some effort must be made to convert the concentration estimates to a cell level SOC (ranging from 0 to 1, defined at the cell voltage limits) in order to compare the result with Coulomb counting. The positive electrode bulk concentration estimates for each of the composite materials are first converted to an overall positive electrode bulk SOC estimate via conservation of lithium within the electrode

\[
\text{SOC}_{\text{p}} = \frac{\hat{c}_{\text{LMO}} + \hat{c}_{\text{NMC}}}{c_{\text{max, LMO}} + c_{\text{max, NMC}}}. \tag{24}
\]

This overall positive electrode bulk SOC operates between an initial and final value \( y_0 \) and \( y_f \) corresponding to the upper and lower voltage limits. The initial and final values \( y_0 \) and \( y_f \) are the functions of the initial and final values of each composite material

\[
y_0 = \frac{y_{0, \text{LMO}} c_{\text{max, LMO}} + y_{0, \text{NMC}} c_{\text{max, NMC}}}{c_{\text{max, LMO}} + c_{\text{max, NMC}}}, \tag{25}
\]

\[
y_f = \frac{y_{f, \text{LMO}} c_{\text{max, LMO}} + y_{f, \text{NMC}} c_{\text{max, NMC}}}{c_{\text{max, LMO}} + c_{\text{max, NMC}}}, \tag{26}
\]

Finally, the estimated cell level \( \text{SOC} \) is calculated by

\[
\text{SOC} = 1 - \frac{\text{SOC}_{\text{p}} - y_0}{y_f - y_0}. \tag{27}
\]

An example of this procedure is shown in Fig. 5 for a constant current discharge. Once a cell level SOC estimate is calculated, it can be validated against the experimental SOC from Coulomb counting.

B. Estimating Capacity

Unlike with SOC estimation, capacity estimates do not necessarily need to be made in real time, since capacity only changes significantly over months or years of usage. This gives some flexibility in making capacity estimates, and it is possible to wait for ideal conditions before making an estimate.

For the composite LMO-NMC cells tested, previous work used differential capacity analysis to show the dominant aging mechanism to be solid-electrolyte interface (SEI) layer growth at the negative electrode [23]. Although other mechanisms, such as loss of active material, may become significant during later stages of life [33], [34] or at high temperatures, only loss of cyclable lithium is considered in this paper. The SEI layer grows as a result of a lithium-consuming side reaction between the electrolyte solvent and the electrode material. This causes a shift in the normalized concentration operating ranges of the positive (\( y_0 \) to \( y_f \)) and negative (\( x_0 \) to \( x_f \)) electrodes, shown in Fig. 6, as the electrodes become less lithiated overall.
Fig. 6. Example of a shift in the electrode operating range from a new to an aged cell, as cyclable lithium is lost to SEI growth. In each electrode, the operating range tends to shift to a less lithiated state.

Again, \( x_0 \) and \( y_0 \) correspond to the upper voltage limit, while \( x_f \) and \( y_f \) correspond to the lower voltage limit.

The normalized concentration operating range of either electrode can be converted to a capacity in ampere-hours by

\[
C = \frac{(x_0 - x_f)c_{\text{max}, \text{LMO}}FAL_n\epsilon_{\text{am}, \text{n}}\epsilon_{\text{n}}}{3600} = \frac{FAL_pr\epsilon_{\text{am}, \text{p}}}{3600} \left( (y_f, \text{LMO} - y_0, \text{LMO})c_{\text{max}, \text{LMO}} + (y_f, \text{NMC} - y_0, \text{NMC})c_{\text{max}, \text{NMC}}\epsilon_{\text{NMC}}\right). \tag{28}
\]

This is the realizable capacity between the two cell voltage limits and so is identical for each electrode. Therefore, an estimate of the normalized concentration range of a given electrode can be used to obtain an estimate of cell capacity.

First, it should be noted that a direct estimate of the lower concentration limit \( x_f \) or \( y_f \) cannot be relied upon, since a complete discharge of the battery is not expected in practice. The upper concentration limit \( x_0 \) or \( y_0 \), however, can be expected to be reached during regular charging. Again, since estimation is only performed on one electrode at a time, determining which electrode normalized concentration range to estimate presents an interesting dilemma. The positive electrode states are less sensitive to voltage errors due to the steep positive OCV curve, but the shift in \( y_0 \) for a given \( \delta C \) is very small and difficult to detect. Conversely, the negative electrode states are more sensitive to voltage errors due to a flatter OCV curve, but the shift in \( x_0 \) is large and easier to detect. Through experimentation, it was determined that the latter approach gave the best results, as the problems due to the flat OCV can be mitigated with careful selection of when to make an estimate. Therefore, the loss in capacity is determined by estimating the shift in \( x_0 \).

Since the concentration corresponding to \( x_f \) is not encountered in practice \( x_f \) is related to \( x_0 \) by a lookup table, via offline simulation of the model during a low rate constant current discharge. For various values of \( x_0 \) corresponding to the upper voltage limit, the model is simulated to determine the \( x_f \) corresponding to the lower voltage limit.

1) Estimating Capacity From an Estimate of \( x_0 \): Estimating \( x_0 \) and \( x_f \) is not as straightforward as simply estimating the electrode concentration at a given point in time. The negative electrode OCV has regions that are relatively flat, which implies sensitivity to voltage model or sensor error. This can be demonstrated by estimating the negative electrode states for a CD cycle, where a perfect model is assumed but a 10-mV voltage measurement bias is introduced (Fig. 7).

The negative electrode normalized concentration estimate is poor in regions where the OCV is flat (Fig. 8), but converges toward the true value in regions where the OCV is steeper. The steep OCV regions are around normalized concentrations of 0.58–0.65 and 0.22–0.35, with the latter region being
the steeper of the two. However, the goal is to estimate $x_0$ (essentially, the negative electrode initial condition, if the cell begins at a fully charged state), which is typically greater than 0.65, and may lie in a flat region of the OCV. This can be observed in Fig. 7, as the estimate is unable to accurately converge from an initial condition error at the beginning of the discharge. Therefore, to obtain an estimate of $x_0$, the strategy is to wait until the estimate is likely converged (i.e., wait until a steep OCV region is reached), and then calculate the initial state by Coulomb counting backward in time starting from the converged state.

Further complicating the problem, any estimation of the negative electrode requires simulating the positive electrode model open loop. This requires knowledge of the positive electrode range $y_0$ and $y_f$, which also change as the cell ages. However, $y_0$ and $y_f$ at a given stage of life are directly related to $x_0$ and $x_f$ according to the following procedures:

1) estimate $x_0$ and $x_f$;
2) calculate $U_{0,n}$ and $U_{f,n}$ at the initial and final state via negative OCV lookup;
3) calculate $U_{0,p} = 4.15 + U_{0,n}$ and $U_{f,p} = 2.8 + U_{f,n}$, since the upper and lower cell voltage limits remain at 4.15 and 2.8 V throughout the battery life;
4) calculate $y_{0,LMO}$, $y_{f,LMO}$, $y_{0,NMC}$, and $y_{f,NMC}$ via the corresponding positive OCV lookup;
5) calculate $y_0$ and $y_f$ using (25) and (26).

The complete SOC/SOH estimation procedure is outlined in Fig. 9. The observer is applied to the positive states to continuously estimate the cell SOC. Then after every CD cycle, an estimate of $x_0$ and $x_f$ is obtained by waiting until either the mid-SOC region or low-SOC region is reached. Finally, capacity is calculated from estimates of $x_0$ and $x_f$.

Essentially, there are dual observers running in parallel, one that estimates the positive electrode states to obtain cell SOC $\text{Obs}_p$ (simulating the negative electrode and liquid-phase open loop) and one that estimates the negative electrode states to obtain capacity $\text{Obs}_n$ (simulating the positive electrode and liquid-phase open loop). Clearly, the submodels that are simulated open loop will not be able to correct for initial condition errors. It is therefore the role of the capacity estimation algorithm to correctly update the $x_0$ and $y_0$ initial conditions so that the open-loop submodels remain accurate.

The propagation of error in the combined SOC/SOH estimation algorithm is a particularly important topic to study. For example, in the SOH estimation block in Fig. 9, the estimated parameter $y_0$ is fed into the next open-loop prediction of the positive electrode. It is conceivable that a small error in a given $y_0$ estimate could contribute to additional error in the next $y_0$ estimate, and so forth, resulting in a positive feedback that would greatly amplify the initial error. An initial investigation into this problem has shown the $x_0/y_0$ estimates to be robust enough that this positive feedback is avoided; however, this topic will be studied further in future work. Particularly, the error in the SOC/SOH estimates should be quantified, or at least bounded. One advantage in the algorithm design is that the only exchange of information between the two observers comes from the update of $x_0$ and $x_f$ that are fed into the open-loop simulation of the negative electrode used by $\text{Obs}_p$. Since the parameters $x_0$ and $x_f$ change only very slowly over hundreds of ampere-hours, they have a negligible effect on the stability of the cell SOC estimation, which operates over a much faster time scale.

C. Initializing the Electrodes

Estimates of the electrode composition ranges, $x_0$, $x_f$, $y_0$, and $y_f$, are used to initialize the electrode concentrations at the beginning of a given cycle. Assuming that the cell is at equilibrium at the start of the cycle, the initial SOC of each electrode is identical, when normalized by its composition range

$$\frac{\text{SOC}_{p,0} - y_f}{y_0 - y_f} = \frac{\text{SOC}_{n,0} - x_f}{x_0 - x_f}$$

where $\text{SOC}_{p}$ is defined in (24) for the composite electrode and $\text{SOC}_{n}$ is simply equal to $\tilde{c}_n/c_{\text{max},n}$. In addition, the initial cell voltage must equal the difference between the positive and negative OCV, evaluated at their respective concentrations. In the case of the composite positive electrode, both the LMO or NMC OCVs may be used (forming two separate equations). Finally, a nonlinear iterative solver is used to solve for the initial SOC of each electrode active material (three equations and three unknowns).

VIII. RESULTS

To validate the combined SOC/SOH estimation algorithm, the estimation was performed on five different automotive pouch cells aged under different conditions. The five cells differ in the final SOC that was reached in CD mode and in the charge rate.

A. SOC Estimate Validation

The cell SOC estimate is compared with SOC from Coulomb counting in Fig. 10 for Cell 2 under a CD PHEV cycle taken at the beginning of life (BOL). Only the results using the EKF are shown, since it requires the least computational effort of the three observer types considered and is therefore the most practical for real-time SOC estimation. The cell SOC estimate converges quickly from an imposed initial condition error, and shows good agreement with the SOC from Coulomb counting, with a maximum error of less than 2% during the CD portion of the cycle. Increased estimate
error up to 4% is seen during the constant voltage charge due to poor open-loop model agreement during this portion of the charge. This level of SOC estimate error is similar among all the cells tested.

Fig. 11 shows the estimated surface and bulk concentrations of each individual composite electrode material over the same PHEV cycle, as well as the current split factor; although again, these cannot be easily validated experimentally. Note that, despite the fact that the current split factor exceeds 1 in some instances, this does not translate into a large current going to the particle, as these are instances when the total current is small. In other words, there are no large current spikes going to a particle that could cause a large error in the state trajectory, as indicated by Fig. 11(c).

**B. Capacity Estimate Validation**

For capacity estimation, the $x_0$ estimate is obtained by waiting for state convergence in a steep OCV region before Coulomb counting backward to the beginning of the discharge. Therefore, a single capacity estimate is obtained after each CD cycle is complete. As noted previously, the low-SOC region from 0.22 to 0.35 is the preferred region to record the converged state since it corresponds to a steeper OCV than the mid-SOC region between 0.58 and 0.65; however, some cells are not sufficiently discharged to reach the low-SOC region. In these cases, the converged state estimate is taken in the mid-SOC region (specifically at 0.6). Otherwise, if the cell does reach the low-SOC region, the estimate at mid-SOC is discarded and the converged state estimate is taken at the midway point between 0.35 and the lowest concentration reached (assuming the value is not less than 0.22). Capacity estimates for the five cells are shown in Figs. 12–16, and are compared with the measured capacities at periodic assessments. For each capacity estimate, it is noted whether the estimate relied on a converged state estimate taken in the low-SOC region or mid-SOC region. In addition, for each cell, three different estimation methods (EKF, Kalman smoother, and particle filter) are compared. The estimation results are summarized in Table III.

The first thing to note is that after each capacity assessment, the estimates show some small amount of capacity recovery. This is expected since the cells would typically rest for
around a day while the capacity assessments were carried out, allowing some capacity recovery.

Overall, the capacity estimates are more accurate and have a lower variance for cells that discharged to the low-SOC region (Cells 1–3). This result is expected, since the estimates should be less sensitive to model and sensor errors in the low-SOC region. In addition, the mid-SOC range is relatively small (only spans a normalized concentration of 0.07), so it becomes more difficult to decide when the estimate is truly converged. Essentially, there is a chance of missing this mid-SOC region, and taking the estimate too early in the discharge (when the estimate has not yet converged) or too late (when the estimate is starting to diverge again). At BOL, Cells 4 and 5 are not discharged sufficiently to reach the low-SOC region, so the initial estimates for these cells are taken in the mid-SOC region. However, as the estimated $x_0$ decreases over time, the end of the CD cycles begin to cross the threshold into the low-SOC region, at which point the algorithm switches to estimate in the low-SOC region. An important point here is that although the estimated concentration may cross into the low-SOC region, triggering the switch, the actual cell may not have done so. This is most likely the case for Cell 5, which is only discharged to 45% SOC. Right before the switch to an estimate at low-SOC is made, near the end of the test, the estimated capacities happen to be lower than the measured capacity (at least for the EKF and smoother). This means that the switch is made prematurely resulting in additional inaccuracies in the estimate after the switch is made. That said, the EKF and smoother eventually switch back to estimating in the mid-SOC region, tending to oscillate between the two regions. These problems illustrate the difficulty in relying on an imperfect estimate to determine if thresholds have been crossed.

In comparing the three observer types, there are no large differences in the estimate error mean or variance among the observers. The particle filter tends to produce slightly
TABLE III
CAPACITY ESTIMATION RESULTS

<table>
<thead>
<tr>
<th>Cell</th>
<th>Test Condition</th>
<th>Observer</th>
<th>Estimate Error* Mean (Ah)</th>
<th>Estimate Error Variance (Ah²)</th>
<th>Normalized Computation Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CD to 25% SOC, C/3 Charge</td>
<td>EKF</td>
<td>-0.08</td>
<td>0.020</td>
<td>1.00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Kalman Smoother</td>
<td>0.18</td>
<td>0.023</td>
<td>1.14</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Particle Filter</td>
<td>-0.04</td>
<td>0.022</td>
<td>1.81</td>
</tr>
<tr>
<td>2</td>
<td>CD to 25% SOC, 3C/2 Charge</td>
<td>EKF</td>
<td>-0.01</td>
<td>0.026</td>
<td>1.00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Kalman Smoother</td>
<td>0.15</td>
<td>0.031</td>
<td>1.15</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Particle Filter</td>
<td>0.05</td>
<td>0.027</td>
<td>1.82</td>
</tr>
<tr>
<td>3</td>
<td>CD to 25% SOC, 5C Charge</td>
<td>EKF</td>
<td>-0.47</td>
<td>0.012</td>
<td>1.00</td>
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<td></td>
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<td>Kalman Smoother</td>
<td>-0.32</td>
<td>0.019</td>
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<td></td>
<td>Particle Filter</td>
<td>-0.17</td>
<td>0.028</td>
<td>1.78</td>
</tr>
<tr>
<td>4</td>
<td>CD to 35% SOC, C/3 Charge</td>
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<tr>
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<td>5</td>
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<td>1.13</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Particle Filter</td>
<td>-0.00</td>
<td>0.081</td>
<td>1.78</td>
</tr>
</tbody>
</table>

*Estimate error defined as the difference between each estimate and a capacity interpolated between the periodic measured capacity assessments.

![Fig. 16. Capacity estimation results for Cell 5 aged with a CD profile to 45% SOC charged at C/3.](image1)

This more accurate but noisier estimates than the other observers, particularly when estimating in the mid-SOC region. Overall, the EKF performs well without the additional computational effort required for the smoother and particle filter.

1) Maintaining SOC Accuracy: In addition to estimating cell capacity, another important objective in estimating the changes in model parameters corresponding to the dominant degradation mode within a cell (e.g., $x_0$ as it relates to loss of cyclable Li) is to maintain an accurate model over time, thereby maintaining an accurate SOC estimate over time. Fig. 17 shows how the accuracy of the SOC estimate changes over time for Cell 2 (the other cells showed a similar result). The estimated $x_0$ is used to update the model after each CD cycle, and as described previously, the SOC estimation is done using the EKF on the positive electrode submodel. The capacity used to calculate the measured SOC by Coulomb counting is obtained by interpolating between each periodic capacity assessment.

![Fig. 17. Root mean square and maximum SOC error for Cell 2. Each point indicates the error over a single CD-CCCV profile.](image2)

The SOC estimate error over a given CD-CCCV profile remains fairly constant throughout the life of the cell, indicating that the model is being updated correctly with estimates of $x_0$. Note that the outliers in the plot are instances where the profile had two CCCV charge portions, over which the model is less accurate.

IX. Conclusion

In this paper, a reduced-order electrochemical model for a composite electrode battery was applied in a dual-observer algorithm to estimate SOC and capacity. The estimates were validated against experimentally measured cell SOC and capacity over the course of an aging campaign. The capacity estimation algorithm performs well as long as the cell is discharged to a sufficiently low SOC, where the negative electrode OCV is steep and state estimates are less sensitive to voltage errors. Estimates taken in the mid-SOC range produced less reliable capacity estimates despite the small window of relatively steep OCV in this range. There was not
a significant improvement in the capacity estimation accuracy when using the more computationally expensive Kalman smoother and particle filter over the simpler EKF. Further estimation accuracy could be achieved with greater knowledge of the model parameters, but this paper demonstrates that the EKF approach is an attractive option for onboard SOH estimation. Future work in this area will focus on capacity and available power estimation in the presence of multiple degradation modes, i.e., if both cyclable lithium and active material are lost simultaneously.

APPENDIX

MODEL OBSERVABILITY CALCULATIONS

The nonlinear output equation for the single particle system makes calculation of the Lie derivatives and observability matrix tedious. An attempt will be made here to at least outline the process of calculating each element in the observability matrix, even though the entire matrix cannot be shown. Zero current will be assumed to reduce the length of the expressions, thereby eliminating the charge transfer and ohmic overpotential terms. This results in the following simplification of the dynamics and output equation (dropping the subscript $i$):

$$
\begin{align*}
\dot{\mathbf{x}} &= \mathbf{f}(\mathbf{x}) = \\
&= \begin{bmatrix}
0 & 1 & 0 \\
0 & 0 & 1 \\
0 & 0 & \frac{b_2}{b_3}
\end{bmatrix} \mathbf{x}
\end{align*}
$$

$$
g(x) = \frac{x_3 c(x)^3}{c_{\text{max}}} + \frac{s_2 c(x)}{c_{\text{max}}} + \frac{s_1 c(x)}{c_{\text{max}}} + s_0
$$

where the substitution $c(x) = a_0 x_1 + a_1 x_2 + a_2 x_3$ has been made. The first Lie derivative and the first row of the observability matrix are shown in Table IV, without expanding the other matrix elements. Again, it should be made clear that the calculation of the complete observability matrix, without assuming zero current, was done using symbolic manipulation software resulting in a matrix with full rank.

REFERENCES


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