Electrochemistry-based Battery Modeling for Prognostics

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**ABSTRACT**

Batteries are used in a wide variety of applications. In recent years, they have become popular as a source of power for electric vehicles such as cars, unmanned aerial vehicles, and commercial passenger aircraft. In such application domains, it becomes crucial to both monitor battery health and performance and to predict end of discharge (EOD) and end of useful life (EOL) events. To implement such technologies, it is crucial to understand how batteries work and to capture that knowledge in the form of models that can be used by monitoring, diagnosis, and prognosis algorithms. In this work, we develop electrochemistry-based models of lithium-ion batteries that capture the significant electrochemical processes, are computationally efficient, capture the effects of aging, and are of suitable accuracy for reliable EOD prediction in a variety of usage profiles. This paper reports on the progress of such a model, with results demonstrating the model validity and accurate EOD predictions.

1. **INTRODUCTION**

With electric unmanned aerial vehicles, electric/hybrid cars, and commercial passenger aircraft, we are witnessing a drastic increase in the usage of batteries to power vehicles. These vehicles are being deployed in military, civilian and scientific applications all over the world. However, for battery-powered vehicles to operate at maximum efficiency and reliability, it becomes crucial to both monitor battery health and performance and to predict end of discharge (EOD) and end of useful life (EOL) events.

For example, NASA’s Mars Global Surveyor stopped operating in late 2006 due to battery overheating caused by the radiator being oriented directly towards the sun, resulting in a significant loss of capacity (Saha & Goebel, 2009). Both the AFRL ARGOS satellite and the Viking 2 Mars Lander also suffered battery failures. More recently, problems have arisen in the Boeing Dreamliner aircraft, which utilize lithium-ion (Li-ion) batteries for auxiliary power (Ross, 2013). Battery monitoring failed to capture thermal runaway conditions, leading to fire. With advanced health monitoring and predictive capabilities, such problems can be prevented.

To implement such technologies, it is crucial to understand how batteries work and to capture that knowledge in the form of models that can be used by monitoring, diagnosis, and prognosis algorithms. Several battery modeling methodologies have been developed in the literature. The models can generally be classified as empirical models, electrochemical engineering models, multi-physics models, and molecular/atomist models (Ramadesigan et al., 2012). Empirical models (e.g., (Saha & Goebel, 2009)), are based on fitting certain functions to past experimental data, without making use of any physicochemical principles. Electrical circuit equivalent models are popular forms of empirical models, e.g., (Ceraolo, 2000; Chen & Rincon-Mora, 2006; Daigle, Saxena, & Goebel, 2012). Electrochemical engineering models are typically continuum models that include electrochemical kinetics and transport phenomena, e.g. (Karthikeyan, Sikha, & White, 2008; Rong & Pedram, 2006; Ning & Popov, 2004; Doyle, Fuller, & Newman, 1993; Newman & Tiedemann, 1975). The former type of model has the advantage of being computationally efficient, but the disadvantage of limited accuracy and robustness, due to the approximations in battery behavior that are made, and as a result of such approximations, cannot represent aging well. The latter type of model has the advantage of being very accurate, but are often computationally inefficient, having to solve complex sets of partial differential equations, and thus being ill-suited to online application of prognostics. Similarly, multi-physics (Lee, Smith, Pesaran, & Kim, 2013) and atomist models are even less suited to online application.

In this work, we develop an electrochemistry-based model of Li-ion batteries, that (i) capture the significant electrochemical processes, (ii) are computationally efficient, (iii) capture
the effects of aging, and (iv) are of suitable accuracy for reliable EOD prediction in a variety of usage profiles. The model can be considered an electrochemical engineering model, but unlike most such models found in the literature, we make certain approximations that allow us to retain computational efficiency. Although we focus here on Li-ion batteries, the model is quite general and can be applied to different chemistries through a change of model parameter values. This paper reports on the progress of such a model, providing model validation results and EOD prediction results.

The paper is organized as follows. Section 2 provides background material on batteries. Section 3 describes the model-based prognostics framework. Section 4 develops the electrochemistry-based battery model. Section 5 provides model validation and battery prognosis results. Section 6 concludes the paper.

2. BACKGROUND

A battery converts chemical energy into electrical energy, and often consists of many cells. A cell consists of a positive electrode and a negative electrode with electrolyte in which the ions can migrate. For Li-ion, a common chemistry is a positive electrode consisting of lithium cobalt oxide (Li$_x$CoO$_2$) and negative electrode of lithiated carbon (Li$_x$C). These active materials are bonded to metal-foil current collectors at both ends of the cell and electrically isolated by a microporous polymer separator film that is permeable to Li ions. The electrolyte enables lithium ions (Li$^+$) to diffuse between the positive and negative electrodes. The lithium ions insert or deintercalate from the active material (known as intercalation and deintercalation) depending upon the electrode and whether the active process is charging or discharging.

A schematic of the battery during the discharge process is shown in Fig. 1. When fully charged, the active (mobile) Li ions reside in the negative electrode. Upon connecting a load to the battery, current is allowed to flow from the positive to the negative electrode. This is supported by the oxidation reaction (loss of electrons) in the negative electrode (acting as the anode):

\[
\text{Li}_x\text{C} \xrightarrow{\text{discharge}} \text{C} + x\text{Li}^+ + xe^- ,
\]

which results in the liberation of Li ions and electrons. The electrons move through the load from the negative to the positive electrode, and the Li ions move in the same direction from the negative to the positive electrode through the separator. In the positive electrode (acting as the cathode), the reduction reaction (gain of electrons) takes place:

\[
\text{Li}_{1-x}\text{CoO}_2 + x\text{Li}^+ + xe^- \xrightarrow{\text{discharge}} \text{LiCoO}_2
\]

During charging, a current source forces current to move from the negative to the positive electrode, as shown in Fig. 2. The active material in the positive electrode (acting as the anode) is oxidized and Li ions are deintercalated with the corresponding reaction:

\[
\text{LiCoO}_2 \xrightarrow{\text{charge}} \text{Li}_{1-x}\text{CoO}_2 + x\text{Li}^+ + xe^-
\]

that results in the loss of Li ions and electrons, which can then move to the negative electrode (acting as the cathode). The reduction reaction takes place in the negative electrode:

\[
\text{C} + x\text{Li}^+ + xe^- \xrightarrow{\text{charge}} \text{Li}_x\text{C}
\]

Theoretically the cell has a voltage of around 4.2 V when fully charged. The terminal voltage of the battery rises/falls with a charge/discharge cycle, respectively. At the end of the charging/discharging period the battery voltage settles to a steady-state value that is a function of its state of charge (SOC). SOC is conventionally defined to be 1 when the battery is fully charged and 0 when the battery is fully discharged. Hence even in a degraded battery the SOC for max-
minimum charge is always 1 by definition.

The charge/discharge dynamics of batteries can be characterized by taking voltage measurements under constant current charge/discharge conditions. The rate of charge/discharge is measured relative to the battery capacity $C$. For example, a 0.1$C$ discharge rate for a 5 Ah battery is 0.5 A.

Charging typically consists of periods of constant current (CC) and/or constant voltage (CV) charging. At lower SOC, a CC charge is applied to bring the battery voltage to the CV level. Once the desired voltage level is achieved, the charger switches to CV mode and current decreases until 100% SOC is reached. The charger is not operated in CV mode at lower SOC levels since this will increase the amount of current flowing through the battery while charging, leading to additional heat generation and side reactions taking place that may affect the battery life. The charge acceptance is the maximum permissible rate at which batteries can be charged. For Li-ion batteries the recommended charge acceptance is $C/3$.

The end of life of a battery depends on the chemistry, discharge-charge cycling, temperature, and storage conditions, among other factors. Atmospheric temperature affects the operating performance of the batteries. At low temperatures, ionic diffusion and migration could be hindered and also damaging side reactions like lithium plating may take place. High temperatures cause corrosion and generation of gases leading to an increase in internal pressure. As the battery ages, degradation results in the decrease in the observed battery capacity. This is primarily due to a loss of mobile ions due to parasitic or side reactions and an increase in the internal resistance. Internal resistance leads to ohmic losses that generate heat and accelerate the aging process. Some relevant physical aging mechanisms in the electrodes are:

1. Solid-electrolyte interface (SEI) layer growth: The negative electrode degrades with the growth of the SEI layer leading to an increase in the impedance. The layers are formed during cycling and storage at high temperatures and entrains the lithium.
2. Lithium corrosion: Lithium in the active carbon material of the negative electrode corrodes over time leading to degradation. This causes a decrease in the capacity due to irreversible loss of mobile lithium ions.
3. Lithium plating: At low temperatures, high charge rates and low cell voltages forms a plating layer on the negative electrode which leads to irreversible loss of lithium.
4. Contact loss: SEI layer disconnects from the negative electrode which leads to contact loss and an increase in impedance.

3. Battery Prognostics

In this section we describe the general model-based prognostics problem and an implementation architecture, and describe how it applies to batteries.

3.1. Problem Formulation

We assume the system model may be generally defined as

$$x(k+1) = f(k, x(k), \theta(k), u(k), v(k)),$$  \hspace{1cm} (1)

$$y(k) = h(k, x(k), \theta(k), u(k), n(k)),$$ \hspace{1cm} (2)

where $k$ is the discrete time variable, $x(k) \in \mathbb{R}^{n_x}$ is the state vector, $\theta(k) \in \mathbb{R}^{n_\theta}$ is the unknown parameter vector, $u(k) \in \mathbb{R}^{n_u}$ is the input vector, $v(k) \in \mathbb{R}^{n_v}$ is the process noise vector, $f$ is the state equation, $y(k) \in \mathbb{R}^{n_y}$ is the output vector, $n(k) \in \mathbb{R}^{n_n}$ is the measurement noise vector, and $h$ is the output equation.\(^1\)

In prognostics, we are interested in predicting the occurrence of some event $E$ that is defined with respect to the states, parameters, and inputs of the system. We define the event as the earliest instant that some event threshold $T_E : \mathbb{R}^{n_x} \times \mathbb{R}^{n_\theta} \times \mathbb{R}^{n_u} \rightarrow \mathbb{B}$, where $\mathbb{B} \triangleq \{0,1\}$ changes from the value 0 to 1. That is, the time of the event $k_E$ at some time of prediction $k_P$ is defined as

$$k_E(k_P) \triangleq \inf\{k \in \mathbb{N} : k \geq k_P \land T_E(x(k), \theta(k), u(k)) = 1\}.$$

The time remaining until that event, $\Delta k_E$, is defined as

$$\Delta k_E(k_P) \triangleq k_E(k_P) - k_P.$$  \hspace{1cm} (3)

In the context of systems health management, $T_E$ is defined via a set of performance constraints that define what the acceptable states of the system are, based on $x(k)$, $\theta(k)$, and $u(k)$ (Daigle & Goebel, 2013). For batteries, there are two types of predictions that we are interested in. The first is EOL, defined by a voltage threshold $V_{EOD}$. In this case, $T_E$ is defined by $V < V_{EOD}$, that is, when the battery voltage is less than the cutoff voltage, EOD is reached and $T_E$ evaluates to 1. The second type of prediction is EOL, which is typically defined by a lower bound on the effective battery capacity (Saha & Goebel, 2009). In this case, $T_E$ is defined by $C < C_{EOL}$, where $C$ is the measured battery capacity and $C_{EOL}$ is the lower bound on capacity.

3.2. Prognostics Architecture

We adopt a model-based prognostics architecture (Daigle & Goebel, 2013), in which there are two sequential problems, (i) the estimation problem, which requires determining a joint state-parameter estimate $p(x(k), \theta(k) | y(k_0:k))$ based on the history of observations up to time $k$, $y(k_0:k)$, and (ii) the prediction problem, which determines at $k_P$, using $p(x(k), \theta(k) | y(k_0:k))$, a probability distribution

\(^1\)Here, we use bold typeface to denote vectors, and use $n_a$ to denote the length of a vector $a$. 
The distribution for $\Delta k_E$ can be trivially computed from $p(k_E(k_P)|y(k_0:k_P))$ by subtracting $k_P$. Both steps require a dynamic model in the form of equations 1 and 2.

The prognostics architecture is shown in Fig. 3. In discrete time $k$, the system is provided with inputs $u_k$ and provides measured outputs $y_k$. The estimation module uses this information, along with the system model, to compute an estimate $p(x(k), \theta(k)|y(k_0:k))$. The prediction module uses the joint state-parameter distribution and the system model, along with hypothesized future inputs, to compute the probability distribution $p(k_E(k_P)|y(k_0:k_P))$ at given prediction times $k_P$.

In the application of prognostics in this example, we do not account for any uncertainty except for that provided in the state estimate, as our focus is on determining how accurate the model can predict EOD given precise information about future. A more general prognostics architecture that accounts for these additional sources of uncertainty is described in (Daigle, Saxena, & Goebel, 2012; Sankararaman, Daigle, Saxena, & Goebel, 2013).

4. Battery Modeling

In order to predict end-of-discharge as defined by a voltage cutoff, the battery model must compute the voltage as a function of time given the current drawn from the battery. There are several electrochemical processes that contribute to the cell’s potential that make this a difficult problem. We focus here on a lumped-parameter ordinary differential equations form, so it is efficient and usable for on-line prognostics, yet still considers the main electrochemical processes. We focus here specifically on Li-ion 18650 batteries with an average nominal voltage of 3.7V and nominal capacity of 2200mAh, however, the model is still general enough that with some modifications it may be applied to different battery chemistries.

The potentials at the current collectors are described by several voltage terms. At the positive current collector is the equilibrium potential $V_{U,p}$. This voltage is then reduced by $V_{s,p}$, due to the solid-phase ohmic resistance, and $V_{n,p}$, the surface overpotential. The electrolyte ohmic resistance then causes another drop $V_e$. At the negative electrode, there is a drop $V_{s,n}$ due to the surface overpotential, and a drop $V_{n,n}$ due to the solid-phase resistance. The voltage drops again due to the equilibrium potential at the negative current collector $V_{U,n}$. We describe each of these voltage terms in turn.

4.1. Equilibrium Potential

In an ideal battery, the equilibrium potential of an electrode is described by the Nernst equation:

$$V_{U,i} = U_0 + \frac{RT}{nF} \ln \left( \frac{\gamma_{\beta,i} \alpha_{i}^{\beta}}{\gamma_{\alpha,i}^{\beta}} \right), \quad (5)$$

or, equivalently,

$$V_{U,i} = U_0 + \frac{RT}{nF} \ln \left( \frac{x_{\beta,i}^{\alpha}}{x_{\alpha,i}^{\beta}} \right) + \frac{RT}{nF} \ln \left( \frac{\gamma_{\beta,i}^{\alpha}}{\gamma_{\alpha,i}^{\beta}} \right), \quad (6)$$

where $i$ refers to the electrode ($n$ for negative or $p$ for positive), $U_0$ is a reference potential, $R$ is the universal gas constant, $T$ is the electrode temperature, $n$ is the number of electrons transferred in the reaction ($n = 1$ for Li-ion), $F$ is Faraday’s constant, $x$ is the mole fraction, where $\alpha$ refers to the lithium-intercalated host material and $\beta$ to the unoccupied
host material, and $\gamma$ is the activity coefficient (Karthikeyan et al., 2008). When discharging, Li ions move out of the negative electrode and into the positive electrode, so $x_{\alpha,n}$ and $x_{\beta,p}$ decrease while $x_{\beta,n}$ and $x_{\alpha,p}$ increase. As a result, $V_{U,p} - V_{U,n}$ will decrease. In an ideal condition the activity coefficients are unity and so the last term disappears, resulting in the classical form of the Nernst equation.

In the case of a Li-ion battery, $x_{\beta,i} = 1 - x_{\alpha,i}$, so we can rewrite the equation making this substitution. We then define $x_i$ as $x_{\alpha,i}$ and drop the $\alpha$ subscript:

$$V_{U,i} = U_0 + \frac{RT}{nF} \ln \left( \frac{1 - x_i}{x_i} \right) + V_{\text{INT},i},$$

(7)

where $V_{\text{INT},i}$ is the activity correction term (0 in the ideal condition). For $\text{Li}_2\text{CO}_3$, $x$ must be at least 0.4; Li cannot be reversibly removed beyond that (Karthikeyan et al., 2008).

We let $q_i$ represent the amount of Li ions in electrode $i$, as measured in Coulombs. Recall that the flow of Li ions moves opposite to the flow of current. Therefore $q_i$ changes in the same direction as $x_i$, and we may define $x_i$ based on $q_i$ using

$$x_i = \frac{q_i}{q_{\text{max}}},$$

(8)

where $q_{\text{max}} = q_p + q_n$ refers to the total amount of available (i.e., mobile) Li ions. It follows then that $x_p + x_n = 1$.

When fully charged, $x_p = 0.4$ and $x_n = 0.6$. When fully discharged, $x_p = 1$ and $x_n = 0$.

Batteries are nonideal and therefore the assumption of unity activity coefficients is not applicable. As a result, battery models often resort to curve-fitting for modeling the equilibrium potential. The equilibrium potential can be obtained by discharging the battery at a very slow rate, such that a concentration gradient will not build up, temperature is steady, and voltage drops due to internal resistances and other overpotentials are negligible. Fig. 5 shows the measured voltage ($V_{U,p} - V_{U,n}$) as a function of mole fraction against that predicted by the Nernst equation. Clearly, the Nernst equation cannot fit this case so it must be nonideal.

The activity coefficient terms are related to excess Gibbs free energy and can be captured using the Redlich-Kister expansion (Karthikeyan et al., 2008). We then have

$$V_{\text{INT},i} = \frac{RT}{nF} \ln \left( \frac{\gamma_{\beta,i}}{\gamma_{\alpha,i}} \right),$$

(9)

$$= \frac{1}{nF} \sum_{k=0}^{N_i} A_{i,k} \left( (2x_i - 1)^{k+1} - \frac{2x_i k (1 - x_i)}{(2x_i - 1)^{1-k}} \right).$$

(10)

The number of terms $N_i$ in the above expansion and the fitting parameters $A_{i,k}$ must be identified. Fig. 6 shows the modified equation fitted to the measured equilibrium potential for

$$N_p = 12 \text{ and } N_n = 0.$$ Clearly, the modified equation much more realistically reflects the true voltage curve. The identified parameters are given in Table 1 and were determined using the Nelder-Mead simplex method. The identified parameters represent those achieving a least-squares (local) error minimum, which is satisfactory for our purposes.

### 4.2. Concentration Overpotential

When a battery is discharged, the reactions take place at the surface of the electrode and this results in a concentration gradient. The concept is shown in Fig. 7. In the bulk volume, the concentration is nearly even, but close to the electrode the concentration changes drastically. To accommodate this, one may split the total volume into two individual control volumes (CVs), one for the bulk (with subscript $b$) and one for the surface (with subscript $s$).

For the volumes, the concentration of Li ions is computed as

$$c_{b,i} = \frac{q_{b,i}}{v_{b,i}},$$

(11)

$$c_{s,i} = \frac{q_{s,i}}{v_{s,i}},$$

(12)
where the diffusion rate from the bulk to the surface is expressed as the surface due to the concentration gradient. The diffusion of electrons. Li ions also move from the bulk CV to the surface layer at the positive electrode, in order to match the flow of electrons. As the battery discharges, Li ions must move out of the surface layers were replenished with ions (from the bulk) in the negative electrode. The battery may then be discharged further since the surface layer becomes like the bulk, accounting for the recovery in the concentration overpotential simply by using as the expression for equilibrium potential, we can compute the potential for the bulk volume and the potential for the surface layer; the difference between them is the concentration overpotential. We can explicitly account for the concentration overpotential simply by using as the expression for equilibrium potential, the equilibrium potential of the surface layer, i.e.,

\[
V_{U,i} = U_0 + \frac{RT}{nF} \ln \left( \frac{1 - x_{s,i}}{x_{s,i}} \right) + V_{INT,i},
\]

where \(x_{s,i}\) is computed using

\[
x_{s,i} = \frac{q_{s,i}}{q_{s,i}^{\text{max}}},
\]

and

\[
q_{s,i}^{\text{max}} = q_{s,p} + q_{b,p} + q_{s,n} + q_{b,n}.
\]

The concentrations in the surface layers change faster than the mole fraction at the bulk, causing a larger concentration gradient for larger applied current and smaller diffusion rates. The observed voltage depends only on what is happening in the surface layer, not the bulk. When discharge stops, the surface layer becomes like the bulk, accounting for the recovery in voltage. The battery may then be discharged further since the surface layers were replenished with ions (from the bulk) in the negative electrode.

Table 1. Battery Model Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(U_{0,p})</td>
<td>4.05 V</td>
</tr>
<tr>
<td>(A_{p,0})</td>
<td>(-33642.23) J/mol</td>
</tr>
<tr>
<td>(A_{p,1})</td>
<td>0.11 J/mol</td>
</tr>
<tr>
<td>(A_{p,2})</td>
<td>23506.89 J/mol</td>
</tr>
<tr>
<td>(A_{p,3})</td>
<td>(-74679.26) J/mol</td>
</tr>
<tr>
<td>(A_{p,4})</td>
<td>14359.34 J/mol</td>
</tr>
<tr>
<td>(A_{p,5})</td>
<td>307849.79 J/mol</td>
</tr>
<tr>
<td>(A_{p,6})</td>
<td>85053.13 J/mol</td>
</tr>
<tr>
<td>(A_{p,7})</td>
<td>(-1075148.06) J/mol</td>
</tr>
<tr>
<td>(A_{p,8})</td>
<td>2173.62 J/mol</td>
</tr>
<tr>
<td>(A_{p,9})</td>
<td>991586.68 J/mol</td>
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<td>(A_{p,10})</td>
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<td>(A_{p,11})</td>
<td>(-163020.34) J/mol</td>
</tr>
<tr>
<td>(A_{p,12})</td>
<td>(-470297.35) J/mol</td>
</tr>
<tr>
<td>(U_{0,n})</td>
<td>0.01 V</td>
</tr>
<tr>
<td>(A_{n,0})</td>
<td>86.19 J/mol</td>
</tr>
</tbody>
</table>

The \(q\) variables are described as

\[
q_{s,p} = i_{\text{app}} + \dot{q}_{bs,p} \tag{17}
\]
\[
q_{b,p} = -\dot{q}_{bs,p} + i_{\text{app}} \tag{18}
\]
\[
q_{b,n} = -\dot{q}_{bs,n} + i_{\text{app}} - \dot{i}_{\text{app}} \tag{19}
\]
\[
q_{s,n} = -i_{\text{app}} + \dot{q}_{bs,n} \tag{20}
\]

where \(i_{\text{app}}\) is the applied electric current. Initially, \(c_{s,i} = c_{b,i}\) and so diffusion is zero. Once discharge begins, ions move from the surface layer of the negative electrode to the bulk volume of the negative electrode, through the separator to the bulk volume of the positive electrode and then on to the surface layer of the positive electrode. If the applied current is larger than the diffusion rate, then a concentration gradient will build up between the surface and bulk volumes. Once discharge stops, then only diffusion is active, and the concentrations will even out, causing a rise in voltage (because the concentration lowers in the surface layer of the positive electrode, raising the voltage, and the concentration rises in the surface layer of the negative electrode, raising the voltage).

The concentration overpotential is the difference in voltage between the two CVs due to the difference in concentration. Using the expression for equilibrium potential, we can compute the potential for the bulk volume and the potential for the surface layer; the difference between them is the concentration overpotential.

As the battery discharges, Li ions must move out of the surface layer at the negative electrode, through the bulk, and to the surface layer at the positive electrode, in order to match the flow of electrons. Li ions also move from the bulk CV to the surface due to the concentration gradient. The diffusion rate from the bulk to the surface is expressed as

\[
\dot{q}_{bs,i} = \frac{1}{D} (c_{b,i} - c_{s,i}),
\]

where \(D\) is the diffusion constant.

Figure 7. Concentration gradient.
4.3. Ohmic Overpotential

The voltage drops due to the solid-phase ohmic resistances, the electrolyte ohmic resistance, and the resistances at the current collectors can be treated as constant resistances and lumped together:

\[ V_o = V_{s,p} + V_{s,n} + V_e + V_{cc,p} + V_{cc,n} \]  
\[ = i_{app}(R_{s,p} + R_{s,n} + R_e + R_{cc,p} + R_{cc,n}) \]  
\[ = i_{app}R_o, \]  
where \( V_o \) is the voltage drop over the current collectors, \( R_{cc,i} \) is the resistance at the current collector for electrode \( i \).

4.4. Surface Overpotential

The overpotentials due to charge transfer resistance and SEI kinetics are described by the Butler-Volmer equation. For small currents the overpotential is linear with current and for larger currents it is proportional to the log of the current, and these two characteristics are combined in the Butler-Volmer equation. The equation is given as

\[ J_i = J_{i0} \left( \exp \left( \frac{(1 - \alpha)F}{RT} V_{\eta,i} \right) - \exp \left( -\frac{\alpha F}{RT} V_{\eta,i} \right) \right), \]  
where \( \alpha \) is the symmetry factor, \( J_i \) is the current density, and \( J_{i0} \) is the exchange current density. The current densities are defined as

\[ J_i = \frac{i}{S_i}, \]  
\[ J_{i0} = k_i(1 - x_{s,i})^\alpha (x_{s,i})^{1-\alpha}, \]  
where \( k_i \) is a lumped parameter of several constants including a rate coefficient, electrolyte concentration, and maximum ion concentration. The voltage drop takes place at the electrode surface, so \( x_{s,i} \) is used in computing the exchange current density.

In the case of Li ions, the symmetry factor is 0.5, so the Butler-Volmer equation can be simplified and written in terms of voltage as

\[ V_{\eta,i} = \frac{RT}{F \alpha} \text{arcsinh} \left( \frac{J_i}{2 J_{i0}} \right). \]

4.5. State of Charge

As discussed in Section 2, the SOC of a battery is conventionally defined to be 1 when the battery is fully charged and 0 when the battery is fully discharged. In this model, it is analogous to the mole fraction \( x_n \), but scaled from 0 to 1. There is a difference here between nominal SOC and apparent SOC. Nominal SOC would be computed based on the combination of the bulk and surface layer CVs in the negative electrode, whereas apparent SOC would be computed based only on the surface layer. That is, a battery can be discharged at a given rate, and reach the voltage cutoff, i.e., apparent SOC is then 0. But, once the concentration gradient settles out, the surface layer will be partially replenished and the battery can be discharged further, i.e. apparent SOC increases whereas nominal SOC remains the same.

Nominal (\( n \)) and apparent (\( \alpha \)) SOC can then be defined using

\[ \text{SOC}_n = \frac{q_n}{0.6q_{\text{max},n}} \]  
\[ \text{SOC}_\alpha = \frac{q_{\text{app},n}}{0.6q_{\text{max},n}}, \]  
where \( q_{\text{max},n} = q_{\text{max},n}/x_n \). The factor 1/0.6 comes from the fact that the mole fraction at the positive electrode cannot go below 0.4 (as described in Section 4.1), therefore SOC of 1 corresponds to the point where \( q_n = 0.6q_{\text{max},n} \).

4.6. Battery Voltage

Now that each of the voltage drops in Fig. 4 have been defined, battery voltage can be expressed as follows.

\[ V = V_{U,p} - V_{U,n} - V_o - V_{\eta,p} - V_{\eta,n}. \]  

Voltages in the battery are not observed to change instantaneously, i.e., the voltage changes occur smoothly. When discharge completes, for example, the voltage rises slowly as the surface layers move to the concentrations of the bulk volumes, as caused by diffusion. In addition to this, there are transients associated with \( V_o \) and the \( V_{\eta,i} \) terms. To take this into account in a simple way, we compute voltage using

\[ V = V_{U,p} - V_{U,n} - V_o' - V_{\eta,p}' - V_{\eta,n}', \]  
where

\[ V_o' = (V_o - V_o')/\tau_o \]  
\[ V_{\eta,p}' = (V_{\eta,p} - V_{\eta,p}')/\tau_{\eta,p} \]  
\[ V_{\eta,n}' = (V_{\eta,n} - V_{\eta,n}')/\tau_{\eta,n}, \]  
where the \( \tau \) parameters are empirical time constants.

The model contains as states \( x, q_{s,p}, q_{s,b}, q_{s,n}, q_{t,n}, V_o', V_{\eta,p}', V_{\eta,n}' \), and \( V_o' \). The single model output is \( V \).

Identified model parameters are given in Table 2. Some parameters are defined by the battery dimensions and chemistry. The remaining parameters are estimated based on data, as with the parameters in Table 1.

The measured and predicted discharge curves for a constant 2 A discharge are shown in Fig. 8. The model fits very well to the measured values. The accuracy towards the end of discharge is most sensitive to the Redlich-Kister parameters, the diffusion constant, and the volume of the surface layer. The
Table 2. Battery Model Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$q^\text{max}$</td>
<td>$1.32 \times 10^7$ C</td>
</tr>
<tr>
<td>$R$</td>
<td>8.314 J/mol/K</td>
</tr>
<tr>
<td>$T$</td>
<td>292 K</td>
</tr>
<tr>
<td>$F$</td>
<td>96487 C/mol</td>
</tr>
<tr>
<td>$n$</td>
<td>1</td>
</tr>
<tr>
<td>$D$</td>
<td>$7.0 \times 10^8$ mol s/C/m$^3$</td>
</tr>
<tr>
<td>$\tau_0$</td>
<td>10 s</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>0.5</td>
</tr>
<tr>
<td>$R_o$</td>
<td>0.085 $\Omega$</td>
</tr>
<tr>
<td>$S_p$</td>
<td>$2 \times 10^{-4}$ m$^2$</td>
</tr>
<tr>
<td>$k_p$</td>
<td>$2 \times 10^{-3}$ A/m$^2$</td>
</tr>
<tr>
<td>$v_{s,p}$</td>
<td>$2 \times 10^{-6}$ m$^3$</td>
</tr>
<tr>
<td>$v_{b,p}$</td>
<td>$2 \times 10^{-5}$ m$^3$</td>
</tr>
<tr>
<td>$\tau_{\eta,p}$</td>
<td>90 s</td>
</tr>
<tr>
<td>$S_n$</td>
<td>$2 \times 10^{-4}$ m$^2$</td>
</tr>
<tr>
<td>$k_n$</td>
<td>$2 \times 10^{-3}$ A/m$^2$</td>
</tr>
<tr>
<td>$v_{s,n}$</td>
<td>$2 \times 10^{-6}$ m$^3$</td>
</tr>
<tr>
<td>$v_{b,n}$</td>
<td>$2 \times 10^{-5}$ m$^3$</td>
</tr>
<tr>
<td>$\tau_{\eta,n}$</td>
<td>90 s</td>
</tr>
</tbody>
</table>

Figure 8. Measured and predicted 2 A discharge curves.

Figure 9. Model validation for variable loading.

predicted end-of-discharge time can be finely tuned most easily by adjusting the time at which the amount of Li ions in the surface layer of the negative electrode hits zero and the surface layer of the positive electrode becomes full, because this is when the equilibrium voltage of these layers nears a combined 2.5 V. This is accomplished by adjusting the diffusion constant.

Model validation for a variable loading scenario is shown in Fig. 9. As shown in Fig. 9a, the load changes every 2 minutes, resulting in corresponding changes in voltage. Fig. 9b shows that the voltage predictions are fairly accurate in response to changes in load. Some errors are still present that may possibly be accounted for by including temperature effects.

4.7. Battery Aging

As described in Section 2, battery aging manifests in two major ways. The first is a loss of capacity due to parasitic and side reactions that result in a loss of active (mobile) Li ions. The second is an increase in internal resistance due to SEI layer growth and other factors. Capacity is typically measured with respect to a reference current and the associated EOD point as defined by a voltage cutoff. A decreased capacity will result in the voltage hitting the cutoff earlier. An increase in resistance will lower the voltage overall, so also result in hitting the cutoff earlier. Therefore both these aging mechanisms result in a loss of measured capacity.

Fig. 10 shows 2 A discharge curves with the battery at different ages. The arrows in the figure illustrate the direction the curves move as the battery ages. The EOD point moves earlier in time due to diminished capacity. The voltage drops down during discharge due to increased resistance. Steady-state voltage after discharge increases because the battery is not discharged as fully due to the increased resistance and the battery reaching EOD before then.

In the battery model, the total available charge in the battery is represented through $q^\text{max}$. Therefore, the loss of active material can be represented in the model through a change in $q^\text{max}$. Fig. 11a shows how the discharge curve changes as $q^\text{max}$ is decreased by 1% with each new discharge. We can see a slight decrease in voltage, which is due to the Butler-Volmer term,
which is dependent on mole fraction that is computed using $\frac{q_{\text{max}}}{q}$ - $\text{max}$. The EOD point moves earlier due to the discharge point being reached with less charge extracted. Steady-state voltage after discharge increases since the cutoff is reached with less relative charge extracted.

One way to validate this is by looking at the equilibrium potential plots as the battery ages, because the resistances are negligible. Fig. 12 shows the measured equilibrium potential of the battery ($V_{U,p} - V_{U,n}$) after 10 discharge cycles and after 30 discharge cycles. The curves are exactly the same, only the time scale changes; there is a 20 minute difference in EOD. Since equilibrium potential is a function only of mole fraction, and mole fraction is computed as $q/q_{\text{max}}$, then decreasing $q_{\text{max}}$ would change the time scale in this way.

Increase in internal resistance can be captured through an increase in the $R_o$ parameter. Fig. 11b shows how the discharge curve changes as $R_o$ is increased by 5% with each new discharge. The voltage drops, but the change in EOD is not significant, because the dynamics near EOD are dominated mainly by the equilibrium potential contribution with some contribution from the Butler-Volmer dynamics. The change in charge extracted is thereby also not significant so changes in steady-state voltage after discharge are very small.

From Figs. 11a and 11b, it is clear that changes in both $q_{\text{max}}$ and $R_o$ are required to capture how the discharge curve changes with aging. Fig. 11c shows the combined effects, with $q_{\text{max}}$ decreasing by 1% and $R_o$ increasing by 5% with each new discharge. The qualitative changes observed in Fig. 10 are now reproduced.

5. APPLICATION TO PROGNOSTICS

With an accurate model and known future inputs to a system, prognostics should in turn be very accurate. In this section we demonstrate battery prognostics with the new model developed in this paper. We use the architecture described in Section 3. As an estimation algorithm, we use the unscented Kalman filter (UKF) with the battery model; see (Julier & Uhlmann, 1997, 2004) for details on the filter and (Daigle, Saha, & Goebel, 2012; Daigle, Saxena, & Goebel, 2012) for its application to prognostics. The UKF operates on a set of deterministically selected samples, called sigma points, that are used to represent the joint state-parameter distribution $p(x(k), \theta(k)|y(k_0:k))$.

For the prediction algorithm, we perform a simple simulation...
as described in (Daigle & Goebel, 2013). Each sigma point is simulated forward using the model until EOD is reached; from the corresponding EODs for each sigma point we can construct the EOD distribution. In this work, we assume that the future inputs ($i_{app}$) are known, so the only uncertainty present in the prediction is that related to the model. We use $V_{EOD} = 2.6 \, V$ as the voltage cutoff defining EOD.

As performance metrics, we use percent root mean square error (PRMSE) for estimation accuracy, and for prediction we use relative accuracy (RA) and relative standard deviation (RSD). We plot results in $\alpha$-plots, where $\alpha$ (e.g. 10%) defines an accuracy cone around the ground truth, and $\lambda$ is a time point (Saxena, Celaya, Saha, Saha, & Goebel, 2010).

Fig. 13 shows prognosis results from a 2 A discharge, assuming the future inputs ($i_{app}$) are known. As shown in Fig. 13a, the UKF tracks very well the measured voltage, since the underlying model is very accurate. Prediction results are, in turn, also very accurate and with very little uncertainty (the only uncertainty is related to the state estimation uncertainty), as shown in Fig. 13b. Overall, RA averages over all prediction points (every 100 s) to 99.38%, and RSD to 0.75%. Since the spread is so small, the 5%-25% and 75%-95% ranges cannot be seen on the plot.

Fig. 14 shows prognosis results from a 3 A discharge. In the open loop, the model slightly underestimates EOD, as shown in Fig. 14b. Even though the UKF tracks well, once a prediction is made the model is in the open loop, and since the model predicts an earlier EOD for the given load there is a bias in the predictions. The UKF helps keep the state estimate accurate and so reduces the error compared to using the model itself.

Fig. 15 shows prognosis results from a variable loading discharge, with EOD being defined in this case as 3.35 V. In the open loop, the model slightly overestimates EOD, as shown in Fig. 15b. The UKF tracks well and corrects for some of the model errors (see Fig. 9b). The model is more accurate at predicting EOD when defined towards the end of a discharge rather than in the middle of a discharge, so if the voltage threshold is defined to be higher, the predictions will be less accurate. In this case RA averages to 88.41%, and RSD to 2.13%. RSD is larger because the UKF needs to account for the additional model uncertainty since the model is less accurate in variable-loading scenarios.

Additional results are shown in Table 3. PRMSE is very low as the UKF is able to track the voltage well in all cases. The predictions are also very accurate, with average RA above 90%, and precise, with RSD around 1% or lower. Although overall very good, the results are best around the 2 A discharge, suggesting that further model improvements (e.g., including temperature effects) may be necessary.

It is also worth mentioning that the model is not computationally intensive. The model is implemented with a fixed-step discrete-time simulation with a step size of 1 s. It takes on the order of $1 \times 10^{-4}$ s to simulate 1 s of real time.

6. Conclusions

In this paper, we presented a new model for battery prognostics based on the underlying physics. Using equations based in electrochemistry, we developed a model that is both accu-
rate and efficient. As a result, prognostics results for EOD prediction are very accurate, with the uncertainty associated with the model remaining very small. Such a model has uses in other areas as well, such as control.

The model described here can also be applied to prognostics of battery packs. There are two approaches here: either each battery is modeled individually, or the batteries of the pack are lumped into an equivalent single-cell model. In the latter case, the prognostics framework as described here applies directly. In the former case, $T_E$ must be redefined on a system-level (Daigle, Bregon, & Roychoudhury, 2012), such that the battery pack is said to be at EOD (or EOL) when any single battery in the pack is at EOD (or EOL).

The current model is limited in several ways that provide the basis for future work. For one, we have not described how the internal battery temperature changes over time. This is important because many of the potentials in the battery are functions of temperature. Further, diffusion rates should be impacted by temperature. Although we have described how aging can be represented within the model, we do not yet have descriptions of how these age-related parameters change over time, i.e., we require damage progression equations. Much work has already been done in this area but not with prognostics requirements and applications in mind. With such models, accurate end-of-life predictions can be made. Upon adding these missing elements, application of this modeling framework to different battery chemistries is also of interest.

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**References**


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