Control of Dendrites in Rechargeable Batteries using Smart Charging

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Abstract

In this paper we develop a feed-back control framework for the real-time minimization of microstructures grown within the rechargeable battery. Due to quickening nature of the branched evolution, we identify the critical ramified peaks in the early stages and based on the state we compute the relaxation time for the concentration in those branching fingers. The control parameter is a function of the maximum curvature (i.e. minimum radius) of the branched microstructure, where the higher rate dendritic evolution would lead to the more critical state to be controlled. The charging time is minimized for generating the most packed microstructures and obtained results correlate closely with those of considerably higher charging time periods. The developed framework could be utilized as a smart charging protocol for the safe and sustainable operation the rechargeable batteries, where the branching of the microstructures could be correlated to the sudden variation in the current/voltage.

1 Introduction

The modern era of wireless revolution and portable electronics demands the utilization of reliable intermittent renewables and long-lasting electrical energy storage facilities [1, 2]. As well, the growing demand for portable computational power as well as the introduction of electric vehicles demand novel and reliable high capacity energy storage devices. Despite such impressive growth of the need in the daily lifestyle, the underlying science remains to be developed. Rechargeable batteries, which retrieve/store energy from/within the chemical bonds, have proven the be the most reliable and cleanest resource of electrical energy for the efficient management of the power. [3, 4] Metallic electrodes such as lithium Li [5], sodium Na [6], magnesium Mg [7], and zinc Zn [8] are arguably highly attractive candidates for

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use in high-energy and high-power density rechargeable batteries. Lithium $Li$ possess the lowest mass density ($\rho_{Li} = 0.53 \text{ g.cm}^{-3}$) and the highest electropositivity ($E^0 = -3.04V$ vs SHE\textsuperscript{1}) which provides the highest gravimetric energy density and likely the highest voltage output, making it suitable for high-power applications such as electric vehicles \[9, 10, 11\]. Sodium has a lower cost and is more earth abundant and is operational for large-scale stationary energy storage applications \[12\]. Magnesium $Mg$ possess a high specific capacity and reactivity \[13\] whereas Zinc $Zn$ is earth-abundant, has low cost and high storage capacity \[14\].

During the charging, the fast-pace formation of microstructures with relatively low surface energy from Brownian dynamics, leads to the branched evolution with high surface to volume ratio \[10\]. The quickening tree-like morphologies could occupy a large volume, possibly reach the counter-electrode and short the cell. Additionally, they can also dissolve from their thinner necks during subsequent discharge period and form detached dead crystals, leading to thermal instability and capacity decay \[15, 16\]. The quickening tree-like morphologies could occupy a large volume, possibly reach the counter-electrode and short the cell. Additionally, they can also dissolve from their thinner necks during subsequent discharge period. Such a formation-dissolution cycle is particularly prominent for the metal electrodes due to lack of intercalation\textsuperscript{2}, where the depositions in the surface is the only dominant formation mechanism versus the diffusion into the inner layers as the housing \[17, 18\]. The growing amorphous crystals can pierce into the polymer electrolyte and short the cell afterwards, given their higher porosity, they could have mechanical properties comparable to the bulk form \[19\].

Previous studies have investigated various factors on dendritic formation such as current density \[20\], electrode surface roughness \[21, 22\], impurities \[23\], solvent and electrolyte chemical composition \[24, 25\], electrolyte concentration \[26\], utilization of powder electrodes \[27\] and adhesive polymers\[28\], temperature \[29, 30\], guiding scaffolds \[31, 32\], capillary pressure \[33\], cathode morphology \[34\] and mechanics \[35, 36, 37, 38\]. Some of conventional characterization techniques used include NMR \[39\] and MRI. \[40\] Recent studies also have shown the necessity of stability of solid electrolyte interphase (i.e. SEI) layer for controlling the nucleation and growth of the branched medium \[41, 42\] as well as pulse charging \[43, 44, 45\].

Earlier model of dendrites had focused on the electric field and space charge as the main responsible mechanism \[46\] while the later models focused on ionic concentration causing the diffusion limited aggregation (DLA). \[47, 48, 49, 50\] Both mechanisms are part of the electrochemical potential \[51, 52\], indicating that each could be dominant depending on the localizations of the electric potential or ionic concentration within the medium. Recent studies have explored both factors and their interplay, particularly in continuum scale and coarser time intervals, matching the scale of the experimental time and space \[53\]. Other simplified frameworks include phase field modeling \[54, 55, 56\] and analytical developments \[57\].

During charge period the ions accumulate at the dendrites tips (unfavorable) due to high electric field in convex geometry and at the same time tend to diffuse away to other less concentrated regions due to

\textsuperscript{1}SHE: Standard Hydrogen Electrode, taken conventionally as the reference ($E^0_{H_2}=0$)

\textsuperscript{2}Intercalation: diffusion into inner layer as the housing for the charge, as opposed to depositing in the surface.
diffusion (favorable). Such dynamics typically occurs within the double layer (or stern layer [58]) which is relatively small and comparable to the Debye length. In high charge rates, the ionic concentration is depleted on the reaction sites and could tend to zero [50]; Nonetheless, our continuum-level study extends to larger scale, beyond the double layer region [59].

Dendrites instigation is rooted in the non-uniformity of electrode surface morphology at the atomic scale combined with Brownian ionic motion during electrodeposition. Any asperity in the surface provides a sharp electric field that attracts the upcoming ions as a deposition sink. Indeed the closeness of a convex surface to the counter electrode, as the source of ionic release, is another contributing factor. In fact, the same mechanism is responsible for the further semi-exponential growth of dendrites in any scale. During each pulse period the ions accumulate at the dendrites tips (unfavorable) due to high electric field in convex geometry and during each subsequent rest period the ions tend to diffuse away to other less concentrated regions (favorable). The relaxation of ionic concentration during the idle period provides a useful mechanism to achieve uniform deposition and growth during the subsequent pulse interval. Such dynamics typically occurs within the double layer (or stern layer [58]) which is relatively small and comparable to the Debye length. In high charge rates, the ionic concentration is depleted and concentration on the depletion reaches zero [50]; Nonetheless, our continuum-level study extends to larger scale, beyond the double layer region [53].

Pulse method has been qualitatively proved as a powerful approach for the prevention of dendrites [43], which has previously been utilized for uniform electroplating [60]. In the preceding publication we have experimentally found that the optimum rest period correlates well with the relaxation time of the double layer for the blocking electrodes which is interpreted as the RC time of the electrochemical system [61]. We have explained qualitatively how relatively longer pulse periods with identical duty cycles D will lead to longer and more quickening growing dendrites. We developed coarse grained computationally affordable algorithm that allowed us reach to the experimental time scale (ms). We have developed theoretical limit the optimal minimization of the dendrites [45] and we have obtained the pulse charging parameters for individual curved peaks based on their curvature [59].

In this paper, we elaborate on the real-time controlling of the pulse charging parameters for the minimization of microstructures grown in the scales extending to the cell domain. We analyze the oscillatory behavior and the transition from initial to steady-state growth regimes.

2 Methodology

The electrochemical flux is generated either from the gradients of concentration (∇C) or electric potential (∇V). In the ionic scale, the regions of higher concentration tend to collide and repel more and, given enough time, diffuse to lower concentration zones, following Brownian motion. In the continuum (i.e. coarse) scale, such inter-collisions could be added-up and be represented by the diffusion length \( \delta \vec{r}_D \) as:

\[ \delta \vec{r}_D \]

The diffusion coefficient \( D^+ \) is generally concentration dependent [44], due to electro-neutrality within the considerable space in the domain and we assume it is constant in the range considered.
Figure 1: The transport elements in the coarse scale of time and space.

\[ \delta \vec{r}_D = \sqrt{2D^+ \delta t} \hat{g} \]  

(1)

where \( \vec{r}_D \) is diffusion displacement of individual ion, \( D^+ \) is the ionic diffusion coefficient in the electrolyte, \( \delta t \) is the coarse time interval \(^4\), and \( \hat{g} \) is a normalized vector in random direction, representing the Brownian dynamics. The diffusion length represents the average progress of a diffusive wave in a given time, obtained directly from the diffusion relationship \([62]\).

On the other hand, ions tend to acquire drift velocity in the electrolyte medium when exposed to electric field and during the given time \( \delta t \) their progress \( \delta \vec{r}_M \) is given as:

\[ \delta \vec{r}_M = \mu^+ \vec{E} \delta t \]  

(2)

where \( \mu^+ \) is the mobility of cations in electrolyte, \( \vec{E} \) is the local electric field. The voltage \( V \) is obtained from the laplacian relationship in the domain as:

\[ \nabla^2 V \approx 0 \]  

(3)

where the dendrite body is part of the boundary condition per se. The electric field is the gradient of electric potential as:

\[ \vec{E} = -\nabla V \]  

(4)

Therefore the total effective displacement \( \delta \vec{r} \) with neglecting convection\(^5\) would be:

\[ \delta \vec{r} = \delta \vec{r}_D + \delta \vec{r}_M \]  

(5)

\(^4\delta t = \sum_{i=1}^n \delta t_i \) where \( \delta t_k \) is the inter-collision time, typically in the range of fs.

\(^5\)Since Rayleigh number \( Ra \) is highly dependent to the thickness (i.e. \( Ra \propto l^3 \)), for a thin layer of electrodeposition we have \( Ra < 1500 \) and thus the convection is negligible. \([63]\)
as represented in the Figure 1. The pulse charging in its simplest form consists of trains of square active period $t_{ON}$, followed by a square rest interval $t_{OFF}$ in terms of current $I$ or voltage $V$ as shown in Figure 2. The period $P = t_{ON} + t_{OFF}$ is the time lapse of a full cycle. Hence the pulse frequency $f$ is:

$$f = \frac{1}{t_{ON} + t_{OFF}}$$

(6)

and the duty cycle $D$ represents the fraction of time in the period $P$ that the pulse is active:

$$D = ft_{ON}$$

(7)

While the dendrites grow, due to random nature of the evolution of branches, they stick out randomly, which becomes a source for the quickening growth of the dendrite per se due to the concentration of electric field in the sharp interfaces as well as their closer proximity to the upcoming ions. The relaxation time allows the concentrated ions in the ramified peaks to dissipate away into the less concentrated areas and the concentration gradient is relaxed. The time required for the such relaxation depends on the curvature of the interface, where the higher curvature sites would require longer time for the concentration relaxation within the double layer region [59]. In fact such relaxation could occur for the larger interfacial double layer scale with the thickness of $\kappa$, spanning to the entire cell domain, requiring the most relaxation time for the highest curvature regions and least for the flat counterparts. From dimensional analysis the relaxation time of the double layer in the flat electrode is in the range of $\sim \frac{\kappa^2}{D^+}[64]$ and for the larger domain of the cell with the representative length of $l$ would scale up to $\sim \frac{\kappa l}{D^+}$, where as their geometric mean has been later considered as $\sim \frac{\kappa l}{D^+}[61]$. In fact the relaxation of the concentration depends highly to the curvature of the peaks and the entire growing interface possess a wide range of radius of curvature values $r_d$, expanding from the flat surface to the highly-curved fingers as small as the atomic value, therefore:

$$r_d \in [r_{atom}, \infty)$$

(8)

Hence, the relaxation time for a randomly-growing interface with variation of curvature along the interfacial line, would vary as well. Hereby, we define the feedback relaxation time $t_{REL}$ a curvature dependent function $f(r_d)$ multiplied by the geometric mean time for the concentration relaxation (i.e. RC time) as:

$$t_{REL} = f(r_d) \frac{\kappa l}{D^+}$$

(9)

As the interface grows from the initial flat state ($r_d \to \infty$) to creating sharp fields ($r_d \to r_{atom}$), the feedback relaxation time $t_{REL}$ should adapt respectively based on the most critical state of the interface, which is the location of ionic concentration. Therefore the range of acceptable value for the feedback relaxation time $t_{REL}$ should lie between the relaxation scale within the double layer to the scale of the cell domain. The variation of the feedback relaxation time $t_{REL}$ occurs from it’s minimum value during
the instigation, to it’s maximum value in the atomic scale. Therefore considering the relevant time-scales from beginning (flat) to the most ramified state (atomic scale), from Equation 9one has:

\[
\lim_{r_d \to \infty} f(r_d) = 1 \quad \text{Flat}
\]
\[
\lim_{r_d \to r_{atom}} f(r_d) = \frac{1}{\kappa} \quad \text{Ramified}
\]

Assuming the form of the control function an combination of linear and exponential terms as \( f(r_d) = ar_d + b \exp(cr_d) \), from the boundary conditions in the Equation 11 one gets:

\[
f(r_d) = 1 + (\frac{r}{\kappa} - 1) \exp[-r_d]
\]

Thus the feedback relaxation time \( t_{REL} \) is obtained as:

\[
t_{REL} = \frac{\kappa l}{D^+} \left( 1 + (\frac{r}{\kappa} - 1) \exp[-r_d] \right)
\]

The radius of curvature \( r_d \) can be approximated via the contours of the iso-potential curvature of the electric field in the vicinity of electrodeposits, where it occurs typically within the double layer region of thickness \( \kappa \). The corresponding line could be obtained locating magnitude of the isopotential contour matching with the electrode \((V = V_{electrode})\). If \((x, y)\) represents the coordinates of the curvature line, the point of the minimum radius of curvature would address the most critical state, and requires higher dissipation of ionic concentration. The radius of curvature \( r_d \) can be calculated from Equation 13 as:

\[
r_d = \min \left( \frac{1 + \frac{dy}{dx}}{\frac{d^2y}{dx^2}} \right)_{x=l}
\]

This value is computed in real time and inserted into the feedback algorithm. In fact the curvature-dependent relaxation time provides a positive feedback for halting of the quickening dendrites, which is negative feedback for dendritic evolution. The Flowchart 3 represents the control loop representing the real-time computation of the curvature and the corresponding feedback relaxation time for the
minimization of the dendritic branching. Figure 4 schematically represents such variation where the feedback relation time \( t_{REL} \) starts from the minimum value of \( \sim \frac{\kappa l}{D^+} \) in the flat surface and varies based on the measurement of the highest curvature of the tip given in the Equation 13. The thickness of the double layer \( \kappa \) can be obtained from [61]:

\[
\kappa = \sqrt{\frac{\varepsilon k_B T}{2z^2e^2C_b}}
\]

where \( \varepsilon \) is the permittivity of the solvent, \( k_B \) is Boltzmann constant, \( T \) is the temperature, \( z \) is the valence number, \( e \) is the electron charge and \( C_b \) is the average ambient electrolyte concentration.

The computation was carried out based on the simulation parameters given in the Table 1. Figure 6 illustrates the resulted morphologies of the grown dendrites based on the applied relaxation time.

The density of the electro-deposits can easily be calculated from confining the atoms in a rectangle, the height of which spans to the highest dendrite coordinates \( h_{max} \). Therefore:

\[
\rho = \frac{n\pi r^2}{h_{max} l}
\]

where \( n \) is the number of atoms composing the dendrite, \( r \) is the atomic radius and \( l \) is the scale of the domain. The density of the morphologies, sample of which is shown in the Figure 5 is provided in the Figures 6a, 6b and 6c versus various relaxation time values as the control parameter.

As well, the variation of the highest interfacial curvature (minimum radius of curvature \( r_d \)) and their corresponding feedback relaxation time \( t_{REL} \) and the density \( \rho \) versus the number of deposited atoms is shown in the Figures 7, 8 and 9.

3 Results & Discussion

The mechanism used in the pulse charging works based on the relaxation of the ionic concentration in the dendritic tips. The formation of such gradient in the concentration is, in fact, the feedback for the quickening upcoming growth regime after the instigation and therefore the applied feedback relaxation time should effectively dissipate away the accumulated ions from the accumulated regions. In fact the sharper interfaces, which have been growing faster than the rest of the interface, have higher number of
concentrated ions around them and therefore they are in the most critical state, which have been used for the computation of the feedback relaxation time.

In the larger scale since the electro-migration displacement (Eq. 2) scales with \( \sim t \) and the diffusion displacement (Eq. 1) scales with the square of time \( \sim \sqrt{t} \). During the pulse both electro-migration and diffusion are in action whereas during the rest period only diffusion is the main drive. Therefore since the average reach for electro-migration is higher than the sole-diffusion the range of reach in the rest period should in fact be competitive with the pulse period. Therefore:

\[
\sqrt{2D^+t_{OFF}} \geq \mu^+\vec{E}t_{ON} \pm \sqrt{2D^+t_{ON}}
\]  

and performing further, we get the maximum value of duty cycle \( D \) for effective pulse charging:

\[
D_{\text{max}} = \max \left( \frac{1}{1 + \frac{|\vec{E}|}{RT} \sqrt{\frac{D^+}{2f}}} \pm 1 \right) \leq \frac{1}{2}
\]  

where the duty cycle of the \( \frac{1}{2} \) is the limiting value for the effective suppression of dendrites. The formation of local branches indicates that the concentration of ions in those specific sights is high and therefore those sites should be focus locations for the feedback relaxation time \( t_{REL} \), which highly depends on the radius of curvature \( r_d \) of the dendritic peaks [59]. For an individual ramified peak with the radius of curvature \( r_d \), the time required for the concentration relaxation \( t_{DL}^{REL} \) within the double layer with the scale of \( \sim \kappa \) is:

\[
t_{DL}^{REL} \approx \frac{\kappa(r_d + \kappa)}{D^+}
\]  

where \( D^+ \) is the diffusivity value for the ionic transport.
which in fact shows faster relaxation relative to the flat interface. For the larger scale, extending to the entire cell domain, the feedback relaxation time $t_{REL}$ act on the uniformization of ionic concentration in the global range (i.e. $\sim l$). Therefore the scale of transport for the time and space would lead to the following comparison:

$$\frac{\kappa^2}{D} \leq t_{DL}^{REL} \leq \frac{k l}{D} \leq t_{REL} \leq \frac{l^2}{D} \tag{17}$$

Therefore the relaxation time scale varies from $\sim \frac{\kappa^2}{D}$ in the individual peaks to the $\sim \frac{l^2}{D}$ in the larger domain of the cell, and the control relaxation time in fact varies in such range. Figure 6 represents the density of the dendrites $\rho$ versus the intervals of pulse $t_{ON}$ and the rest (i.e. relaxation) $t_{OFF}$ periods. The density of the electro-deposits $\rho$ correlates inversely with the pulse charging time $t_{ON}$. This is due to the exacerbated branching during the charge time which upon growing further gets more difficult to halt. On the other hand, applying finer pulse periods $t_{ON}$ provides better possibility for the suppression of dendrites. Needless to mention that such pulse period $t_{ON}$ could not indefinitely get short since the ions ultimately would require enough time to reach the dendrites during this time and react from ionic to atomic species.

As well, the density values $\rho$ correlates with the relaxation time $t_{OFF}$ until reaching a certain saturation limit. Since the length of the domain is much larger than the double layer ($l \gg \kappa$), the range of feedback relaxation time $t_{REL}$, shown by color gradient, would extremely reduce the charging time with negligible compensation in the density of electro-deposits $\rho$. The underlying reason is that the relaxation would let the ionic concentration to relax and uniform ionic distribution. On the other hand, extra relaxation period will not helpful since the ionic concentration is already relaxed and the concentration gradient has already vanished.

As well, imposing higher-than-limit relaxation time would slightly reduce the density $\rho$ since additional concentration from the ambient electrolyte could be depleted in the into the non-reacting dendritic sites. The negligible increase in the density of the dendrites $\rho$ in the span of Figures 6a, 6b and 6c illustrates the effective-ness of the pulse charging method for the multitudes of the charge amount $N$.

The dendritic evolution can be divided into two distinctive stages of the transient and steady-state...
(S.S.) growth regimes [66, 67], which has been illustrated in the Figures 7, 8 and 9. The initial transient regime in fact is a stochastic in nature whereas the steady state regime can illustrate an effective trend. Figure 7 represents the variation of the radius of the curvature $r_d$ in the growing interface versus the deposition progress $N$ (i.e. number of the atoms). In this figure, the transition stage during the higher pulse time shows more fluctuation which indicates the non-uniform regime of growth for the augmented pulse intervals. On the other hand during the steady-state (S.S.) regime, radius of curvature correlates inversely with the pulse time interval $t_{ON}$, which indicates that the morphology is controlled for more finer pulse periods.

Figure 8 represents the control relaxation time $t_{REL}$ versus the progress in electrodeposition during the dendritic evolution, where the higher pulse intervals would require higher amount of control relaxation time $t_{REL}$ for the effective suppression of the dendrites. As well the higher fluctuation for the higher amount of pulse charge $t_{ON}$ shows the higher control rate due to faster dynamics of variation in the curvature.

The same trend of transition-to-steady state regimes has been observed in the Figure 9, where the highest fluctuation occurs for the higher pulsing time $t_{ON}$ where is leads to the lowest density $\rho$ after reaching the steady growth regime.

In fact, the quickening growth regime of the dendrites illustrates that the larger height $h$ of the electrodeposits the rate of their growth would be higher as well. This can simply be represented by the following:

$$\frac{dh}{dt} \propto h$$

where the integration leads to the exponential relationship for the growth regime as:

$$h(t) \propto \exp(bt)$$

where $b$ is the coefficient of the proportion. Setting exponential relationship causes a very high sensitivity for the control relaxation time $t_{REL}$ to act vigilantly versus smallest perturbation in the ramified peaks and the form of the control relation time $t_{REL}$ proportionally contains the exponential form in Equation
Note that other forms of the relaxation time would as well could satisfy the boundary conditions given in the Equation 10 such as $t_{REL}^{ALT}$ given as:

$$t_{REL}^{ALT} \approx \frac{\kappa l}{D} \left( \frac{r_d + l}{r_d + \kappa} \right)$$

(18)

which has lower sensitivity for the radius of curvature $r_d$ relative to proposed control relation time $t_{REL}$. This can be proven by calculating their derivative with respect to radius of curvature ($\frac{dt_{OFF}}{dr_d}$) and show that:

$$\frac{dt_{REL}}{dr_d} \gg \frac{dt_{REL}^{ALT}}{dr_d}$$

Thus from equations 12 and 18 and considering the negative value for both derivatives, one must have:

$$\frac{\kappa - l}{\kappa \exp(r_d) / D} > \frac{\kappa - l}{(r_d + \kappa)^2 / D}$$

(19)

since $\kappa \ll l$ dividing by negative value of $\kappa - l$ changes the inequality sign, therefore:

$$\exp(r_d) > (r_d + \kappa)^2$$

(20)

The equation 20 is obvious for a large values of radius of curvature $r_d$ since the exponential term in the denominator will surpass the quadratic term in the right side. As well for an infinitesimally small value of the radius of curvature $r_d$ one can use Taylor expansion as: $\exp(r_d) \approx 1 + r_d + O(r_d^2)$ and one has:

$$\kappa(1 + r_d) > (r_d + \kappa)^2$$

re-arranging gives:

$$r_d^2 + \kappa r_d + \kappa^2 - \kappa < 0$$

which is a quadratic equation in terms of the radius of curvature $r_d$ and the root is found as:

$$r_d = \sqrt{\kappa - \frac{3}{4}\kappa^2 - \frac{\kappa}{2}}$$

Considering the infinitesimal value for thickness of the double layer ($\kappa \to 0$) the value for $r_d$ would be very small. Therefore for the most range of $r_d \in [r_{atom}, \infty)$ the exponential relationship remains as the most sensitive to the variations in the radius of curvature $r_d$ and the Equation 12 as effective control relaxation time for suppression of the dendrites.

In practice, the discerning the formation of such a peak in the dendrite morphology for potentiostatic charging (constant applied voltage $V$) could be obtained by computing the sudden increase in the current density, representing the runaway process, whereas for galvanic charging (constant applied current $I$) could be the sudden drop in the potential value, where both of these events represent the runaway
4 Conclusions

In this paper we have developed an effective real-time feedback control relaxation method for minimization of dendritic grown during electrodeposition for preventing the branched evolution of the grown microstructures. The control parameter has been considered as the maximum curvature of the growing interface based on the radius of curvature of the most critical peaks. The sensitivity of the feedback relaxation time to the curvature has been analyzed to be extremely high with the exponential correlation, analogous to the growth dynamics of the branches. The methodology can be used for smart charging in rechargeable batteries for the controlling the morphology of the grown electro-deposits.

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