Interfacial charge transfer is widely assumed to obey the Butler-Volmer kinetics. For certain liquid-solid interfaces, the Marcus-Hush-Chidsey theory is more accurate and predictive, but it has not been applied to porous electrodes. Here we report a simple method to extract the charge transfer rates in carbon-coated LiFePO₄ porous electrodes from chronoamperometry experiments, obtaining curved Tafel plots that contradict the Butler-Volmer equation but fit the Marcus-Hush-Chidsey prediction over a range of temperatures. The fitted reorganization energy matches the Born solvation energy for electron transfer from carbon to the iron redox site. The kinetics are thus limited by electron transfer at the solid-solid (carbon-LiFePO₄) interface rather than by ion transfer at the liquid-solid interface, as previously assumed. The proposed experimental method generalizes Chidsey’s method for phase-transforming particles and porous electrodes, and the results show the need to incorporate Marcus kinetics in modelling batteries and other electrochemical systems.
Electrochemical energy systems are key enabling technologies for renewable energy, electrified transportation and smart grids, in which energy conversion and delivery are carried out by Faradaic reactions between electrons and ions. In many systems, such as lithium-ion batteries and solid oxide fuel cells, charge transfer occurs at complex solid–solid interfaces in porous electrodes, coupled to non-equilibrium thermodynamics. This greatly complicates the interpretation of electrochemical measurements, compared with the simple case of uniform, flat liquid–solid interfaces, widely studied in electroanalytical chemistry.

The importance of interfacial charge transfer kinetics is illustrated by lithium iron phosphate (LiFePO₄), one of the most intensively studied cathode materials for Li-ion batteries. Its electrochemical performance can be dramatically improved by various surface modification techniques, such as carbon coating and graphene wrapping for increasing the electronic conductivity, and anion adsorption for reducing the ionic energy barrier. The basic idea behind these surface treatments is that interfacial charge transfer involves both lithium ions and electrons.

As shown in Fig. 1, when a negative overpotential is applied to a carbon-coated LiFePO₄ crystal, lithium ions jump into the interstitial vacancies in the first atomic layer of the crystal while the electrons tunnel to the iron site and reduce Fe³⁺ to Fe²⁺. After the charge transfer reaction, adjacent Fe²⁺ and Li⁺ ions with the local distortion around them form a neutral quasiparticle, or polaron, that can diffuse into the crystal. Since LiFePO₄ is a poor electronic conductor, the carbon coating acts as the ‘electrode’ that provides electrons for the electrochemical reaction. On the other hand, carbon as an anode material barely accommodates lithium ions at voltages larger than 2 V (ref. 12), so carbon layers coated on cathode materials working at 3–4 V provide a purely ionic barrier against lithium transfer between the solid and the electrolyte, which prevents lithium intercalation across thick carbon coatings.

Existing mathematical models neglect the details of charge transfer and assume that the net reaction, Li⁺ + e⁻ + FePO₄ ↔ LiFePO₄, obeys phenomenological Butler–Volmer (BV) kinetics, focusing on the role of the lithium ion. The net reaction rate constant k for single charge transfer is expressed as

\[ k(\eta)/k_0 = \exp(\eta) - \exp((1 - x)\eta) \]

where \( k_0 \) is the exchange rate constant and \( \eta = e(E - E^0)/k_B T \) is the dimensionless overpotential, scaled to the thermal voltage \( k_B T/e \), (where \( e \) is the elementary charge, \( k_B \) Boltzmann’s constant and \( T \) the absolute temperature). The overpotential is defined as the difference between the electrode potential \( E \) and the formal potential \( E^0 \) (ref. 15). The charge transfer coefficient, \( x \), is usually set to 0.5 in battery modelling. For BV kinetics, the Tafel plot of \( \ln k/(\eta/k_0) \) vs. \( \eta \) is a straight line of slope \( x \) for \( \eta < 0 \) and 1 – \( x \) for \( \eta > 0 \). In principle, the fundamental rate constant \( k_0 \) can be determined from the \( y \) intercept of the fitted Tafel line, but the vast range of fitted exchange currents for the same material (over seven orders of magnitude, \( 10^{-6} - 10^7 \) A m⁻²) undermines the validity of this approach.

The fact that increasing the electronic conductivity of LiFePO₄ (an insulator) can dramatically improve the high-rate performance suggests that lithium intercalation reaction may instead be limited by electron transfer between the carbon coating and the redox site in the crystal,

\[ \text{Fe}^{3+} + e^- \rightarrow \text{Fe}^{2+} \]

while the ion transfer reaction, \( \text{Li}^+ \rightarrow \text{Li}^{2+} \), is fast. In this case, the Marcus theory, honoured by the Nobel Prize in Chemistry, could be applied to fit curved Tafel plots for a solid–solid interface (to our knowledge, for the first time), if the fundamental rate constants could be unambiguously extracted from porous electrode measurements.

In this work, we propose a statistical method of Tafel analysis that can be used to extract fundamental reaction rates for porous electrodes. Since the material properties of LiFePO₄ are readily available, we use typical commercial LiFePO₄ to validate our method. The experimental results are in excellent agreement with microscopic electron transfer theory, thus shedding new light on Faradaic reactions at solid–solid interfaces.

Results

Curved Tafel plots of total currents. As motivation, we begin by performing classical Tafel analyses of literature data. In a recent experiment, Munakata et al. tested a single 20 μm agglomerate of carbon-coated LiFePO₄ over a range of constant currents, up to 900°C (4 s discharge). The measured voltage drops at half fillings of the porous particle under different currents yield a highly curved Tafel plot, where \( x = 0.5 \) BV slope can only fit a small portion of the data. Since the voltage drops at high currents are easily affected by concentration polarization (transport limitation), we read the voltage drops at the lowest fillings of the discharge plateaus, and plot the data in Fig. 2a, obtaining another curved Tafel plot. Analysis of the discharge curves from Kang and Ceder shows a similar trend in Fig. 2b. This as-yet unexplained phenomenon is not unique to LiFePO₄, but has also been reported for Li-O₂ batteries by extrapolating initial voltage drops under different currents.

The total currents used in standard Tafel analyses, however, do not accurately represent the fundamental reaction rates since the active area of a porous electrode is non-uniform and varies with the applied current. Moreover, the method of estimating the activation overpotential from the overshoot of the voltage plateau is fundamentally flawed, since the plateau is an emergent property of collection of phase-transforming particles. Of course, this method also cannot be applied to solid-solution materials without a voltage plateau.
relationship shown in the semilogarithmic plot of the transient current versus time\(^4\). The transient current after a voltage step is fitted to a simple exponential decay\(^{27}\)

\[ I = e^{\gamma t} k_{app} \exp(-k_{app}t) \]  

(3)

where \(\gamma\) is the coverage of the electrode and \(k_{app}\) is the single decay rate, assumed to represent the rate of reactant consumption. Chidsey’s method ensures that the kinetics under different potentials are clearly separated and thus avoids the ambiguity of picking voltage points from constant current discharges. Assuming a simple three-state Markov chain of electrode\(^{25,26}\) results time-dependent populations of reacting redox systems investigated by Chidsey, it cannot be applied to porous electrodes or phase-transforming particles with non-uniform reaction rates. Mosaic instability throughout the electrode\(^{25,26}\) results time-dependent populations of reacting particles, or equivalently an evolving active internal area. In other words, \(\gamma\) in equation (3) is not constant for electrodes with complex thermodynamics.

Recently, Bai and Tian\(^{28}\) proposed a simple statistical model to describe discrete-particle phase transformations in porous electrodes. Assuming a simple three-state Markov chain of untransformed, transforming and transformed particles, the total current of a porous electrode is proportional to the population of phase-transforming particles\(^{28}\). The transient current in response to a voltage step can be expressed as

\[ I = kQ \left[ \frac{N_0k + (N_1 - 1)k_A}{k - k_A} \exp(-kt) + \frac{(1 - N_0 - N_1)k_A}{k - k_A} \exp(-k_A t) \right] \]  

(4)

where \(Q\) is the capacity of the electrode, \(k\) is the reaction rate at the surface of phase-transforming particles that continuously accept (release) lithium ions during discharge (charge), \(k_A\) is a generalized activation rate, which is identical to the nucleation rate at low overpotentials for phase-separating materials, but also captures the random activation process at high overpotentials when phase separation is suppressed\(^{14,29}\). \((k\) and \(k_A\) are identical to \(m\) and \(n\) in ref. 28, respectively.) \(N_0\) is the initial fraction of transforming particles (reacting areas) in the electrode. \(N_1\) is the initial fraction of transformed particles (inactive areas) in the electrode, which is set to zero in the present work.

Since the statistical model does not account for the microscopic dynamics of single-particle transformations, the generalized activation rate \(k_A\) simply reflects the population of reacting particles or sites or, more generally, the evolving active surface area of the porous electrode. Although this assumption may seem overly simple, the statistical model was recently validated by Levi et al\(^{30}\) by fitting in situ electrochemical quartz admittance data ‘surprisingly’ well for a thin microarray of LiFePO\(_4\) particles. Interestingly, in the limit of fast activation \((k_A \to \infty)\), the full internal surface of the porous electrode becomes active, or equivalently, in the monolayer limit\(^4\), equation (4) reduces to \(I = kQ \exp(-kt)\), identical to equation (3) for surface-bound redox systems\(^{27}\). For our porous electrodes, fitting \(k_A\) and \(N_0\) enables more accurate determination of the fundamental reaction rate \(k\).

Figure 3a provides typical transient currents of a LiFePO\(_4\) coin cell used in our experiments (see Methods for details). By fitting the transient currents with equation (4), we can extract the reaction rates \(k\) for different overpotentials. Although the value of the total capacity \(Q\) can be estimated from experiments, it is relaxed in the fitting to account fluctuations in experimental conditions and errors in time integration of the transient current.
Curved Tafel plot of reaction rates. Fundamental reaction rates extracted from transient currents in many different voltage-step experiments are shown in Fig. 4. The data clearly deviates from the linear Tafel dependence of the BV equation at overpotentials larger than $4k_BT/e \approx 100$ mV. As motivated above, we adopt instead the Marcus–Hush–Chidsey (MHC) model\textsuperscript{4,31,32} to calculate the theoretical reaction rates of equation (2),

$$k_{\text{red/ox}}(\lambda, \eta) = A \int_{-\infty}^{\infty} \exp \left\{ - \frac{(x - \lambda \pm \eta)^2}{4\lambda} \right\} \frac{dx}{1 + \exp(x)}$$  \hspace{0.5cm} (5)

where $\lambda$ is the dimensionless reorganization energy scaled to $k_BT$ and $\eta = (E - E^0)/k_BT$ is the same dimensionless overpotential in equation (1). The integral over the dimensionless variable, $x = (x_{\text{et}} - e\lambda)/k_BT$, accounts for the Fermi statistics of electron energies, distributed around the electrode potential\textsuperscript{14}. $A$ is the pre-exponential factor, accounting for the electronic coupling strength and the electronic density of states (DOS) of the electrode. For non-adiabatic electron transfer at a carbon electrode\textsuperscript{33}, the DOS varies with the electrode potential\textsuperscript{33} and cannot be placed outside the integral, but within the small potential range considered in this work, the DOS of the carbon coating is roughly constant\textsuperscript{13} as assumed in equation (5). The net reaction rate of lithium insertion is $k(\lambda, \eta) = k_{\text{red}}(\lambda, \eta) - k_{\text{ox}}(\lambda, \eta)$, and the corresponding rate constant can be calculated as $k_0 = A k_{\text{red/ox}}(\lambda, 0)$.

The extracted reaction rates are compared with various theoretical models. Curves calculated from the MHC model (equation (5)) are plotted in Fig. 4 and adjusted to ensure the convergence to the BV curves at low overpotentials. Remarkable agreement is observed between the MHC curves for $\lambda = 8.3$ and the reaction rates for both the charge and discharge processes across the entire range of overpotentials. Curves calculated from the classical Marcus rate equation (without integrating against the Fermi distribution) are also plotted, but the agreement is not as good as expected for electron transfer from a metal electrode. Since the Marcus rate reaches a maximum and then decreases for $|\eta| > \lambda$ (the celebrated ‘inverted region’, observed in bulk charge transfer reactions), a rather large reorganization energy ($\lambda = 13.5$) is needed to fit the data.

Besides capturing the overpotential dependence of the reaction rates, microscopic theories of charge transfer are able to predict the reorganization energy from first principles, without any empirical fitting. For a liquid–solid interface, the total reorganization energy for electron transfer has two main contributions\textsuperscript{15}: the ‘outer’ reorganization of the solvent, $\lambda_\text{ox}$, dominated by long-range electrostatic forces; and the ‘inner’ relaxation of the reactant itself, $\lambda_\text{red}$, dominated by short-range bond forces. Many experiments have shown that the former dominates, $\lambda \approx \lambda_\text{ox}$ (ref. 27). This approximation should also hold, if not better, for reactions at a solid–solid interface, since the ‘reactant’ is part of the same solid dielectric continuum assumed for the ‘solvent’, and the changes of the bonds connecting the ‘reactant’ and the ‘solvent’ are highly cooperative\textsuperscript{34}. We thus approximate the (dimensionless) reorganization energy by the Born energy of solvation\textsuperscript{5,20},

$$\lambda \approx \lambda_\text{ox} = \frac{\epsilon_0^2}{8\pi\epsilon_0 k_BT} \left( \frac{1}{a_0} - \frac{1}{2d} \right) \left( \frac{1}{\epsilon_{\text{op}}} - \frac{1}{\epsilon_{\text{s}}} \right)$$  \hspace{0.5cm} (6)

where $\epsilon_0$ is the permittivity of free space, $a_0$ the effective radius of the reactant, $d$ the distance from the centre of the reactant to the surface of the electrode, $\epsilon_{\text{op}}$ the optical dielectric constant and $\epsilon_{\text{s}}$ the static dielectric constant. We use the length of Fe–O bond in the FeO$_6$ octahedron as the radius $a_0$, which is roughly 0.21 nm (ref. 35). The dielectric constants are available from first-principles calculations\textsuperscript{35}: $\epsilon_{\text{op}} \approx 9.825 \epsilon_0 = 4.74$ and $\epsilon_{\text{s}} = 11.58$. If we assume that the FeO$_6$ octahedrons were in direct contact with the carbon coating, that is, $d = a_0 = 0.21$ nm, then the calculated reorganization energy is 213 meV, corresponding to (dimensionless) $\lambda = 8.3$ at room temperature. If the FeO$_6$ octahedrons were separated from the carbon coating by PO$_4$ tetrahedrons, adding the length of the O–O bond ($\sim 0.23$ nm), that is, $d = 0.44$ nm, gives us $\lambda = 12.64$. Surprisingly, the MHC model with the lower bound $\lambda = 8.3$ can accurately fit the reaction rates extracted from three coin cells, which is a striking validation of the theory.

Figure 4 | Tafel analysis of the reaction rates (symbols) extracted by equation (4) from chronoamperometry experiments of three coin cells at room temperature. Error bars indicate the 95% confidence bounds. Curves are theoretical results from the BV model (dotted black), the MHC model (blue) and the classical Marcus model (red). Overpotentials are scaled to the thermal voltage $k_BT/e \approx 25.7$ mV. Inset: dependence of the reorganization energy on the separation distance between the reactant and electrode surface, that is, equation (6).
As listed in Table 1, the fitted rate constants for discharge (cathodic) are slightly smaller than those for charge (anodic). The broken symmetry may be attributed to the different lithium surface compositions of the active surface for adsorption and desorption\textsuperscript{36}, and could also be explained by nonlinear concentration and stress effects on the exchange current in generalized Marcus kinetics for mixed ion-electron transfer processes in solids\textsuperscript{2}. As the reaction rates are inferred from the decay rates of the transient currents via equation (4), they are independent of the absolute magnitudes of the currents, which depend on the population of transforming particles. As such, the $k_0$ values in Table 1 can be viewed as effective material properties of the active particles, independent of the areas of reacting surfaces in the porous electrodes, although they do reflect some averaging over different local surface concentrations during a single-particle phase transformation. Since the area of each lithium site is $\sim 0.5 \times 0.5 \text{nm}^2$, $k_0 = 2.0 \times 10^{-4} \text{s}^{-1}$ (per site) yields an exchange current density $I_0 \approx 1 \times 10^{-4} \text{A m}^{-2}$, if each particle transforms homogeneously, as expected for high rates\textsuperscript{14,29}.

The reaction rates extracted from experiments at higher temperatures are displayed in Fig. 5 for each coin cell. In a striking validation of the MHC theory, the nine series of reaction rates can all be fitted by the same reorganization energy $(214 \pm 1 \text{meV})$, independent of temperature, with 0.5% accuracy. Further validation comes from Arrhenius temperature dependence of the corresponding rate constants ($k_0$), which yields the effective energy barriers for interfacial charge transfer reaction at zero overpotential. According to the classical Marcus theory, the effective energy barrier is roughly equal to $\lambda/4$, so the value $\lambda = 13.5$ fitted in Fig. 4 yields a barrier of $\sim 87 \text{meV}$. Despite the fact that the effective energy barriers from the secondary fitting ($k_0$ versus $T$) could easily drift away from the true value, the barriers obtained (115 meV) from the Arrhenius plot (Fig. 5d) are consistent with this estimation. The effective barrier (230 meV) fitted from the cathodic (discharge) rate constants of cell B is

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Fitted rate constant $k_0$ for curves in Fig. 4.</th>
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<tr>
<td><strong>Model</strong></td>
<td>$\lambda$</td>
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<tr>
<td>BV</td>
<td>$z = 0.5$</td>
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<tr>
<td>MHC</td>
<td>8.3</td>
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<td>Marcus</td>
<td>13.5</td>
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**Figure 5** | Temperature dependence of the fundamental reaction rates. Tafel plots of (a) cell A, (b) cell B and (c) cell C at 30, 40 and 50°C. Dotted curves indicate the BV model (linear Tafel plot) with $z = 0.5$. Solid curves result from fitting MHC model with a unique value of the reorganization energy. (d) Arrhenius plot of the rate constants under 25, 30, 40 and 50°C. Open and filled symbols denote cathodic (discharge) and anodic processes, respectively.
relatively larger, but it is comparable to the first-principles energy barrier for polaron transfer in bulk crystal, which under the ‘adiabatic’ assumption yields a diffusion coefficient of order $10^{-8}$ cm$^2$s$^{-1}$. The diffusion time through micron-sized particles at such a high diffusivity is of order 1 s, whereas the time constants observed in our experiments are of order 10$^3$s. Therefore, fast adiabatic polaron transfer within the solid crystal cannot be the rate-limiting step, as further explained in the Discussion below. Instead, the small rate constants indicate very weak electronic coupling between the carbon coating and the LiFePO$_4$ crystal, consistent with non-adiabatic electron transfer kinetics at the interface. Our experiments also reveal that the reaction rate is higher in the fresh cells than in the aged cells, which may be ascribed to interfacial changes due to side reactions and elastic strain fatigue.

**Discussion**

Ionic transport may affect the overpotential via logarithmic concentration terms in the Nernst voltage, but this cannot be the rate-limiting step during battery charge or discharge in our experiments. Taking discharge as an example, the whole electrochemical process consists of three steps: lithium ion transport in the electrolyte, charge transfer reaction at the interface and polaron diffusion in the crystal. The characteristic time constant for lithium ions in the electrolyte to transport across a 5-µm-thick porous electrode is of order 0.1 s at the diffusivity ($10^{-6}$ cm$^2$s$^{-1}$) obtained from experiments. The time constant for polarons in the crystal to diffuse through a 1-µm long [010] channel is strongly temperature dependent and of order 1 s at room temperature, using the diffusivity ($10^{-8}$ cm$^2$s$^{-1}$) from first-principles calculations. The capacity of our coin cells at 0.1C is ~150 mAh g$^{-1}$, which is roughly 88.24% of the theoretical capacity (170 mAh g$^{-1}$). According to the analysis of Malik et al. (see equation (2) and Fig. 2 in ref. 41), such a high capacity indicates less than 0.1% channel-clogging defects in our 1 µm-sized particles, so that the diffusivity should stay close to the theoretical value. In summary, both transport steps appear to be too fast to be responsible for the decaying rates smaller than $10^{-2}$ s$^{-1}$, or equivalently time constants larger than 100 s. This leaves the charge transfer reaction as the only possibility for the rate-limiting step, which also explains the negligible temperature dependence in the curvature of the Tafel plots.

Since the time constant for charge transfer reactions under small overpotentials are four orders larger than that of the solid diffusion ($10^4$ versus 1 s), diffusivities measured from electro-analytical methods are probably affected by the slow reaction surface and, therefore, are orders smaller than the theoretical predictions ($10^{-8}$ cm$^2$s$^{-1}$). The same is true of diffusivities of electrodes by fitting classical porous electrode models to the low diffusion limit due to the flawed assumption of fast BV kinetics at high rates. We are also aware of the small theoretical diffusivities ($10^{-17}$ and $10^{-12}$ cm$^2$s$^{-1}$) for the adsorbed lithium ions hopping between channel entrances on the surface of the FePO$_4$ crystal. However, while lithium ions can find alternative path in the electrolyte to quickly reach the favourable entrance, there is no quicker path for electrons to jump between the solid crystal and the carbon coating. Surface modifications may lower the energy barrier for surface diffusion, but will also inevitably alter the reorganization energy for the electron transfer reaction. Our method may be applied to further distinguish the difference.

Besides the fast diffusion of solid-state lithium ions (polarons) along the [010] channels, the dewetting of lithium ions from the largest and most active [010] facet of FePO$_4$ particles to lower the surface energy implies that constant low concentration of intercalated lithium is maintained at the interface. We conclude that small variations of the concentration in the logarithm term should not cause significant voltage change, consistent with the success of our reaction-limited model. Voltage fluctuations due to the contact resistance are also negligible. The contact resistances of the coin cells measured by potentiostatic impedance are around 6 Ω, which would at most cause a 9-mV voltage drop at a step of 392 mV, or a 0.6-mV voltage drop at a step of 74 mV.

The excellent agreement between the data under various temperatures and the widely used MHC model justifies a posteriori our method of extracting reaction rates from chronopotentiometry experiments, which thus generalizes Chidsey’s method for phase-transforming particles and porous electrodes. The perfect match between the theoretical Born solvation energy and the fitted reorganization energy validates our assumption in equation (2) that electron transfer is the rate-limiting step in the Faradaic reaction, contrary to the prevailing picture of ion transfer limitation, and also justifies our assumption of including Fermi statistics with constant DOS for the carbon-coating electrode. The comparison between MHC and the classical Marcus model reveals the necessity of the Fermi statistics for electrode reactions.

Our electroanalytical method could have broad applicability. We have used micron-sized carbon-coated LiFePO$_4$ to demonstrate the measurement of reaction rates for thin (~5 µm) porous electrodes. For very thick porous electrodes (for example, > 50 µm (ref. 28)), it becomes difficult to accurately fit the transient current with equation (4), but more sophisticated porous electrode theories can be used to account for transport coupled to phase transformations. Our method can be easily applied to other electrode materials, such as nanoparticles, composites and solid-solution materials. Although the curvature of the Tafel plots for other materials may be different, our method can still extract the rate constant and reorganization energy, which can be compared with microscopic charge transfer theories to quantitatively understand the reaction mechanism.

Our findings suggest the need for a paradigm shift in the mathematical modelling of kinetics in electrochemical systems. Essentially, all models assume BV kinetics, not only for batteries but also for electrodeposition, corrosion, fuel cells and so on. In spite of the success of electron transfer theory, the BV equation has been engrained in electrochemistry for over a century, and even used as a ‘filter to discard models of electron transfer that do not predict (the linear) Tafel’s law’. Any switch to Marcus or MHC kinetics could have a dramatic effect on model predictions, since the reaction resistance becomes orders of magnitude larger than BV at large overpotentials. It would be particularly interesting to apply generalized Marcus kinetics to non-equilibrium thermodynamics in LiFePO$_4$, which might correct shortcomings of generalized BV kinetics in fitting experimental data at high rates.

**Methods**

**Coin cell preparation.** The micron-sized LiFePO$_4$ powder was from A123 System, LLC. Scanning electron microscopy shows that the typical size of the primary particles is around 1 µm, and the size of the aggregates is smaller than 5 µm. The powder was mixed with carbon black and sodium alginate binder at the weight ratio of 8:1, and hand milled to form a smooth slurry. The electrode was prepared by casting the slurry onto an aluminium foil using a doctor blade and dried in a vacuum oven at 100 °C overnight. The size of the circular electrodes is 1.27 cm in diameter. Loading mass of each electrode is around 1.1 mg. The thickness of the electrodes are less than 5 µm. Coin cells were assembled in Argon-filled glove boxes (<1 p.p.m. of water and oxygen) with lithium foil as the counter electrode, 1 M LiPF$_6$ in a mixture of ethylene carbonate/diethyl carbonate (EC/DEC, 1:1 by volume) as the electrolyte and Celgard 3501 as the separator. Galvanostatic charge/discharge performance of the cathode materials was tested in a voltage range of 2.4–4.2 V. Capacity at C/10 discharge is ~150 mAh g$^{-1}$, calculated on the basis of the mass of LiFePO$_4$ in the composite electrodes.
Chronoamperometry experiments. All experiments were conducted with an Arbin battery tester at room temperature, unless otherwise specified. Before each voltage step, coin cells were charged (discharged) at 50 μA to 4.0 V (2.5 V) and then relaxed for 4 h to reach equilibrium. The size of the voltage step was calculated with respect to the formal potential $E^0 = 3.430 \text{V}$, as measured as the average of the peak potentials of the cyclic voltammetry. The cutoff current for each step is 5 μA.

Curve fitting. Parameters of equation (4) for each transient current were fitted in the Carve Fitting Toolbox of MATLAB. Starting values of $k_1$, $k_2$, and $N_0$ were adjusted to optimize the fitting results under the following principles: (i) the fitted decaying time constant (1/β) should be in the same order of magnitude as the time constant estimated directly from the chronoamperogram (time for decaying to 36.8% of the peak current); (ii) largest possible $R^2$ value; (iii) smallest possible 95% confidence bounds for all four parameters; and (iv) the fitted Q should be consistent with the experimental values.

References