Influence of Li$_2$O$_2$ Morphology on Oxygen Reduction and Evolution Kinetics in Li-O$_2$ Batteries

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Supporting Information

Experimental Details

Electrode fabrication / catalyst preparation.

Catalyst Preparation: Si wafers (6” diameter, n-type doping, 140 nm thick thermal SiO$_2$) were prepared for catalyst deposition by cleaning using a Piranha solution (3:1 H$_2$O$_2$:HSO$_4$). Next, sequential layers of Al$_2$O$_3$ (30 nm) and Fe (1 nm) were deposited using electron beam evaporation without breaking vacuum between layer depositions. The catalyst wafers were then cleaved into small samples (~1 x 1 cm) in preparation for CNT growth.

Chemical Vapor Deposition: Multi-walled carbon nanotubes (CNTs) were synthesized in a two-furnace hot-walled thermal chemical vapor deposition system. The two furnaces are connected with a single quartz tube (OD 1”). The upstream furnace (preheater) is used to preheat the carbon precursor to promote gas decomposition and results in improved carbon nanotube growth. The downstream furnace (growth furnace) is where catalyst samples are located and CNTs are grown. The CNTs used in this study were synthesized using the process detailed in the main text. After the post-growth anneal the growth furnace was opened and the tube was rapidly cooled to room temperature under He flow. After cooling, the CNTs were removed from the furnace and then removed from the substrate by gently prying the corners of the monolithic carpet until the carpet uniformly delaminated from the growth substrate.

TEM sample preparation.

Small electrode fragments were prepared for TEM imaging by sandwiching them inside foldable TEM grids (Ted Pella, Stratapeak) in order to minimize damage from sample preparation (i.e. samples were not sonicated). Sample preparation was performed inside an argon-filled glovebox and prepared grids were transported to the TEM in argon-filled airtight containers. Bright field zero-loss imaging was performed at an accelerating voltage of 120 kV in an energy-filtered Zeiss Libra 120 TEM.

XRD crystallite size analysis.

Average crystallite sizes of Li$_2$O$_2$ in discharged electrodes were determined by peak broadening analysis on raw XRD data (Fig. 3a and additional data) using the Scherrer equation assuming spherical crystallites:

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\[ t = \frac{0.9 \lambda}{B \cos \theta_B} \]

where \( t \) is the crystallite broadening, \( \lambda \) is the radiation wavelength (1.54 Å for Cu Kα), \( \theta_B \) is the Bragg angle, and \( B \) is the full width half maximum at \( 2\theta_B \). Due to the presence of a broad background shoulder from Kapton near the (101) peak at 34.97° 2θ, and to the presence of two closely spaced peaks at 58.72 and 58.98° 2θ, only the highest intensity (100) peak at 32.89° 2θ was used for crystallite size measurements. For commercial \( \text{Li}_2\text{O}_2 \) measurements, 90% phase-pure \( \text{Li}_2\text{O}_2 \) powder (with 10% LiOH, Sigma-Aldrich, USA) was ball-milled in a PULVERISETTE 6 (Fritsch, Germany) at 500 rpm without solvent under dry argon for 5 hours.\(^2\)
**Figure S1.** Representative potentiostatic discharge curve at 2.76 V vs. Li to 1000 mAh/g, forming a Li$_2$O$_2$ disc morphology.

**Figure S2.** Current vs. capacity of CNT electrodes discharged potentiostatically over a range of potentials between 2.0 – 2.76 V vs. Li (same electrodes as shown in Fig. 1a but over a wider capacity range).
Figure S3. (a) Cyclic voltammetry scan of CNT electrodes starting at open circuit (3.2 V vs. Li) to 2.4 V vs. Li at a scan rate of 5 mV/s. The gray line shows the background in argon, while the blue lines show the raw data of the ORR current in oxygen (dashed blue line) and the capacitance-corrected ORR current obtained by subtracting the argon background (solid blue line). (b) Tafel plot of the capacitance-corrected ORR current (solid blue line in (a)) as a function of the discharge potential. The dashed line is a linear fit to the CV data below 2.6 V.

Figure S4. Low magnification TEM image of the electrode in Fig. 2a (E > 2.74 V to 2000 mAh/gC, PITT discharge).

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Figure S5. XANES O K-edge data comparing TEY and FY data of discharged and charged Li$_2$O$_2$ (a) disc morphologies and (b) particle morphologies with that of reference commercial Li$_2$O$_2$. The electrodes are the same as those shown in Fig. 4c, with experimental conditions and electrochemical data given in Fig. S7.

Figure S6. XANES TEY C K-edge spectra of discharged and partially-charged discs (a) and particles (b) and comparison with commercial Li$_3$CO$_3$ FY data. Electrochemical details and discharge/charge data from the corresponding electrodes are shown in Fig. S7.
Figure S7. Discharge data of XANES samples used in Fig. 4 and S5-6. (a) Galvanostatic discharge at 10 mA/g\textsubscript{C} to 4000 mAh/g\textsubscript{C}, forming disc morphologies. (b) Galvanostatic discharge at 10 mA/g\textsubscript{C} to 3600 mAh/g\textsubscript{C}, forming discs, followed by a ~30% charge at 20 mA/g\textsubscript{C} to 2400 mAh/g\textsubscript{C} at the onset of the plateau region. (c) Potentiostatic discharge at 2.7, 2.4, and 2.0 V to 6200, 5000, and 5200 mAh/g\textsubscript{C}, respectively, forming particle morphologies. (d) Galvanostatic discharge at 100 mA/g\textsubscript{C} to 5000 mAh/g\textsubscript{C}, forming particles, followed by 20% charging at 20 mA/g\textsubscript{C} to 4000 mAh/g\textsubscript{C} in the sloping region.
**Fig S8.** XANES O K-edge TEY spectra overlay of the data in Fig. 4a.

**Figure S9.** TEM of fully charged electrodes at 20 mA/g following (a) discharge at E = 2.0 V vs. Li to 1000 mAh/g, forming Li$_2$O$_2$ particle morphologies and (b) discharge at E = 2.76 V vs. Li to 1000 mAh/g, forming Li$_2$O$_2$ disc morphologies. Insets: selected area electron diffraction patterns of the electrode areas shown in (a) and (b) indicating only reflections from CNTs in the charged electrodes. Bundling of the CNTs in the charged samples is due to collapse of the structure from solvent evaporation after removal from the cell. Scale bars: 100 nm.
Figure S10. PITT charging of Vulcan carbon electrodes pre-loaded with commercial Li$_2$O$_2$. Details regarding electrode fabrication and characterization can be found elsewhere.$^2$ (a) shows the entire capacity range (normalized to carbon), which is dominated by two-phase oxidation as indicated by the large current rise to ~50 mA/g$_C$ and subsequent decay, and (b) shows the delithiation behavior at the very early stages of charge (for Q > 1140 mAh/g$_C$; the maximum capacity expected from pre-loaded Li$_2$O$_2$ electrodes is 1170 mAh/g$_C$).

Figure S11. PITT comparison of two different electrodes containing Li$_2$O$_2$ particles (discharged potentiostatically at 2.0 V vs. Li to 1000 mAh/g$_C$) with two electrodes containing Li$_2$O$_2$ discs (discharged potentiostatically at 2.76 V vs. Li to 1000 mAh/g$_C$).
**Figure S12.** Charging data at 20 mA/g for electrodes discharged potentiostatically at 2.0, 2.7 and 2.76 V vs. Li to 1000 mAh/g, showing reproducibility of the characteristic charging profiles.
Calculation of the Chemical Diffusion Coefficient, $\tilde{D}$

The chemical diffusion coefficient, $\tilde{D}$, was calculated from PITT charging data (Fig. 7) in the sloping voltage region where the current exhibits a monotonic delithiation-related current decay. Calculations were performed following the process outlined by Wen et al.\textsuperscript{3} for delithiation in planar LiAl electrodes (i.e. 1-D diffusion) and were subsequently modified to consider the case of 3D spherical diffusion as outlined by Deiss\textsuperscript{4} and by Montella.\textsuperscript{5} Although the morphologies of Li$_2$O$_2$, particularly the small particles, are clearly three-dimensional, it is possible that one-dimensional diffusion could have relevance for discs given the large anisotropy of Li$_2$O$_2$ discs and unknown diffusion mechanisms. Since the mechanistic details of diffusion processes occurring in the discs and particles (e.g., whether isotropic or anisotropic) are not known, we provide both analyses to give reasonable bounds on the diffusion coefficient.

Starting with Fick’s first law of diffusion,

$$I(t) = -zFSD \frac{\partial C}{\partial x} \bigg|_{x=0}$$

where $z$ is the charge transferred per diffusing species, $S$ is the surface area of Li$_2$O$_2$, and $x = 0$ is defined at the Li$_2$O$_2$ surface. The solution at short time is given by

$$I(t) = zFSD \Delta c \left( \frac{D}{\pi \nu} \right)^{1/2}$$

where $\Delta c$ is the change in concentration at the surface of Li$_2$O$_2$ following application of the potential step. Assuming negligible ohmic losses, electrolyte mass transport, or kinetic limitations due to small polarizations (\(\Delta E = 5 \text{ mV for PITT}\)), $\Delta c$ is given by:

$$\Delta c = \frac{Q}{zFV}$$

where $Q$ is the total charge transferred per PITT step and $V$ is the volume of the Li$_2$O$_2$ particle. Combining terms, the time response is given by

$$I(t) = Q \left( \frac{S}{V} \right) \left( \frac{D}{\pi \nu \tilde{D}} \right)^{1/2}$$

and depends on the ratio of $S/V$ (surface area to volume of Li$_2$O$_2$) for the geometry assumed.\textsuperscript{4,5} For a spherical particle, $L/V$ is given by $(3/r)$ where $r$ is the particle radius, while for a 1D film, $S/V = 1/t$ where $t$ is the film thickness. $r$ or $t$ can be estimated experimentally using the characteristic diffusion length $L$, which in turn is estimated from knowledge of the crystallite size (in our analysis, $L$ was assumed to be one half the crystallite size measured from XRD in Fig. 3b; errors related to uncertainties in $L$ are shown in Fig. S14). A plot of $I(t)$ vs. $t^{-1/2}$ for a given PITT
current decay response after polarization of $\Delta E = 5$ mV yields a linear region at short times ($t \ll L^2/D$, Fig. S13b) from which the value of $\tilde{D}$ can be obtained from the slope, $k$, as

$$\tilde{D}_{\text{planar}} = \pi \left( \frac{kL}{Q} \right)^2$$

or

$$\tilde{D}_{\text{spherical}} = \frac{\pi}{9} \left( \frac{kL}{Q} \right)^2.$$

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**Fig. S13.** (a) Raw current vs. time response for a select set of representative PITT steps ($\Delta E = 5$ mV, $i_{\text{cutoff}} = 0.5$ mA/g) obtained during charging of Li$_2$O$_2$ discs (Fig. 7). The legend indicates the PITT potential applied in V vs. Li. (b) shows the data in (a) plotted as a function of $t^{-1/2}$. The linear region at short times ($>0.1$ s$^{-1/2}$) was used to determine the slope from which $\tilde{D}$ was calculated using the process outline above. A diffusion length of 7.5 nm or 12.5 nm was assumed for particles or discs, respectively, which was roughly half the typical crystallite size of each morphology measured from XRD (Fig. 3; ~25 nm crystallite size for discs and ~15 nm crystallite size for particles). The PITT current response of particles was shown to yield similar results as (a) for the discs and is not shown.
Fig. S14. Chemical diffusion coefficient, $\tilde{D}$, of (a) discs and (b) particles overlaid with the PITT charging profile as a function of capacity. To investigate the error associated with assumption of the characteristic diffusion length $L$ in spherical diffusion, the obtained $\tilde{D}$ for reasonable upper and lower bounds of $L$ for discs and particles are included (gray filled or open symbols). While uncertainty in $L$ may result in slightly shifted (higher or lower) values of $\tilde{D}$ within an order of magnitude, the observed trends (i.e., a decrease in $\tilde{D}$ during delithiation on charge, and a higher maximum $\tilde{D}$ for discs compared to particles) is still observed. The filled colored symbols show $\tilde{D}$ if 1D diffusion is assumed.

References