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Electrical conductivity adjustment for interface capacitive-like storage in sodium-ion battery


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Sodium-ion battery (SIB) is significant for grid-scale energy storage. However, a large radius of Na ions raises the difficulties of ion intercalation, hindering the electrochemical performance during fast charge/discharge. Conventional strategies to promote rate performance focus on the optimization of ion diffusion.

Improving interface capacitive-like storage by tuning the electrical conductivity of electrodes is also expected to combine the features of the high energy density of batteries and the high power density of capacitors. Inspired by this concept, an oxide-metal sandwich 3D-ordered macroporous architecture (3DOM) stands out as a superior anode candidate for high-rate SIBs. Taking Ni-TiO$_2$ sandwich 3DOM as a proof-of-concept, anatase TiO$_2$ delivers a reversible capacity of 233.3 mAh g$^{-1}$ in half-cells and 210.1 mAh g$^{-1}$ in full-cells after 100 cycles at 50 mA g$^{-1}$. At the high charge/discharge rate of 5000 mA g$^{-1}$, 104.4 mAh g$^{-1}$ in half-cells and 68 mAh g$^{-1}$ in full-cells can also be obtained with satisfying stability. In-depth analysis of the electrochemical kinetics evidence that the dominated interface capacitive-like storage enables ultrafast uptake and releasing of Na-ions. This understanding between electrical conductivity and rate performance of SIBs is expected to guide future design to realize effective energy storage.

1. Introduction

To solve the resulting problems of fossil fuel burning, the conversion and utilization of clean renewable energy are of high significance. Owing to the imminent and unpredictable features of such resources, energy storage technologies have thus attracted intensive attention for both basic research and practical industry.$^{[1]}$ Among them, lithium-ion battery stands out as a prominent energy storage technology with a high potential for large-scale off-grid applications.$^{[2]}$ However, issues like the accessible lithium, safety, and cost inspire us to find other alternatives.$^{[3]}$ Since sodium not only shares similar electrochemical properties with lithium, but also possesses several advantages like abundant resources, high safety, and low cost, sodium-ion battery (SIB) has thus been considered as a promising candidate for energy storage.$^{[4]}$ Many researchers have devoted to corresponding studies of electrode materials, especially in pursuit of satisfying rate performance that is even more important due to the demands of fast charge/discharge.$^{[5]}$ The promising methods without consuming reversibility still remain a challenge to Na-free transition metal oxide anode materials with intercalation mechanism.$^{[6]}$

In general, ion intercalation contains three steps during cycles: solvated Na$^+$ diffusion within the electrolyte; charge-transfer reactions at or near the interface between active materials and electrolyte; Na$^+$ diffusion in the bulk materials. Hence, the whole ability of sodium storage is attributed to both Na-ion intercalation in bulk and capacitance at or near the interface. Larger radius of Na$^+$ (1.02 Å) versus Li$^+$ (0.76 Å) always raises more complex requirements for electrode materials’ crystalline structures during bulk intercalation,$^{[6]}$ which would largely hinder the performance at high charge/discharge rates. Interface capacitive-like storage always occurs on the order of seconds and minutes, and thus the motivation from increased interface capacitive-like storage is expected to well balance high capacity and high rate performance in the same material.$^{[7]}$ Moreover, interface capacitive-like storage will not result in a huge change of crystalline structures, which would also contribute to the stability during long-term sodium storage.

Several methods, like defects or amorphization engineering of materials’ crystalline structures$^{[8]}$ and morphology...
optimization,[9] have been used to enhance the rate performance from improved interface capacitive-like storage and obtained some preliminary results. A deep understanding of interface capacitive-like storage nature would be helpful for developing clear criteria for intercalation-based electrodes. Since the contributions at or near the interface are mainly controlled by surface instead of diffusion, and always accompanied with electron transfer or hopping,[10] sufficient electrical conductivity for fast electron movement might be also significant to add extra capacitive-like sodium storage at or near the interface.[31] This enhancement would be expected to benefit for high-rate charge/discharge in short time.

Inspired by this understanding, an active material-metal current collectors and sandwich architecture is an effective candidate to shorten electron transport paths throughout the entire electrode, originated from the high electrical conductivity of the metal skeleton. To evidence this concept more clearly, we chose anatase TiO$_2$, a typical intercalation-based material with unsaturation metal skeleton. To realize this tentative idea of the active material-current collector sandwich 3DOM architecture, it is expected for a universal design to realize satisfied energy storage using other large transport ions.

2. Results and Discussion

2.1. Fabrication and Characterizations of Ni-TiO$_2$ Sandwich 3DOMs

To confirm the aforementioned steps, we choose Ni as the metal skeleton and anatase TiO$_2$ as the active material. Figure 1a,b shows the representative scanning electron microscope (SEM) images of 3DOM Ni skeleton and Ni-TiO$_2$ sandwich 3DOM, which are the replica translated from the periodic CCTs based on PSs with $\approx$500 nm diameter. The Ni skeleton with the periodic macroporous structure was obtained on Ti foil after the electrochemical deposition and template removal by organic solvents. And then $\approx$20 nm TiO$_2$ was deposited on the Ni skeleton to form an oxide-metal sandwich 3DOM structure, followed by annealing to obtain the anatase phase. As shown in Figure 1b, morphologies of both surface layer and inner layers (highlighted in the circle of Figure 1b) confirm the successful realization of Ni-TiO$_2$ sandwich 3DOM, the phase
and purity of which were further examined by X-ray diffraction (XRD) in Figure 1c and Raman spectroscopy in Figure 1d. Besides the diffraction peaks from Ni (JCPDS No. 65–2865) and Ti (JCPDS No. 44–1294) substrates, all the other peaks can be indexed to the diffractions of tetragonal anatase TiO₂ (JCPDS No. 21–1272). No peaks of other phases can be detected, indicating the high purity of the as-prepared Ni-TiO₂ sandwich 3DOMs. The Raman scattering spectra of as-fabricated samples further evidences the purity of anatase TiO₂ based on vibration modes of A₁g, B₁g, E₁g. It is worth mentioning that there is no peak in Raman spectra between 1200 and 1800 cm⁻¹, where the D band and the G band of carbonaceous materials appear. Two symmetrical peaks in X-ray photoelectron spectroscopy (XPS) are attributed to +4 state of Ti⁴⁺ in the lattice as shown in Figure S1, Supporting Information. Overall, we can infer that no carbon is contained in the Ni-TiO₂ sandwich 3DOMs.

To clearly visualize the sandwich 3DOM features on a large scale, transmission electron microscopy (TEM) analysis and energy-dispersive X-ray spectroscopy (EDX) scan were used here to verify the distributions of Ni, Ti, and O (Figure 2a–d) using samples with templates consisted of 100 nm diameter spheres. TEM image shown in Figure 2a evidences the periodic
macroporous features of the entire Ni-TiO$_2$ sample. The elemental distribution of Ni (Figure 2b) exhibits similar 3DOM morphology compared with that of Ni-TiO$_2$ sample, confirming the core of the 3DOM Ni skeleton, while Ti and O are distributed around the Ni skeleton uniformly like a sandwich construction. With the help of color code, the clean interface between Ni (orange) and TiO$_2$ (blue) can be clearly found in high-resolution TEM (HRTEM) image (Figure 2e), verifying the direct contact between the active materials and current collectors. No other impurities like NiO, carbon exist in the sandwich interface. Summarizing 3DOM, where a 3D macroporous conductive network is formed by the metal Ni with a uniform coating of anatase TiO$_2$ acted as active materials on both sides. To better illuminate the structure information, enlarged HRTEM images of Ni and TiO$_2$ are shown in Figure S2, Supporting Information.

2.2. Electrochemical Characteristics of Ni-TiO$_2$ Sandwich 3DOMs

To quantify the sodium storage of Ni-TiO$_2$ sandwich 3DOM, we first employ a half-cell configuration with a TiO$_2$-based working electrode and Na disk counter electrode to assemble two-electrode coin cells. In order to demonstrate our hypothesis of the influences of electrical conductivity of electrodes, anatase TiO$_2$ 3DOM without Ni skeleton was fabricated as reference samples with the same thickness of TiO$_2$ (2 $\times$ 20 nm) and mass loading. Corresponding characterizations can be found in Figure S3, Supporting Information. Since both kinds of 3DOMs were fabricated directly on the conductive Ti substrates, we didn’t use any conductive additive and polymeric binder to ensure sufficient conductivity of entire electrodes. Hence, all the exhibited sodium storage abilities only originate from electrochemical behaviors of anatase TiO$_2$ with a comparatively clean surface. Cyclic voltammetry (CV) measurements were first carried out to elucidate the redox processes in the host matrix with different constructions. To avoid the disturbance from the solid electrolyte interphase (SEI) layer and the other irreversible reactions, the second cycle of CV curves is chosen as diagrammed in Figure 3a. Looking carefully into the appearances of both CV curves, the pair of redox peaks located at approximately 0.67 V (cathodic) and 0.85 V (anodic) are observed in both Ni-TiO$_2$ sandwich 3DOM and TiO$_2$ 3DOM. This redox pair well matches the potential of reversible Ti$^{4+}$/Ti$^{3+}$ redox coupled with Na$^+$ insertion and extraction into anatase TiO$_2$.[15] Considering similar mass loading of Ni-TiO$_2$ sandwich 3DOM and TiO$_2$ 3DOM without Ni skeleton, it is reasonable to exclude the influence of the mass of the active material without normalization of Y-axis. We can further assert that the electrochemical behaviors during Faradaic bulk storage of sodium is quite similar during low charge/discharge by similar location of redox pairs. Gradually broadening curves of Ni-TiO$_2$ sandwich 3DOM may indicate larger contributions from interface capacitive-like storage.[9d,16] Hence, Ni-TiO$_2$ sandwich 3DOM exhibits larger capacities during cycles, the value of which keeps 233 mAh g$^{-1}$ even after 100 cycles at the low charge/discharge rate of 50 mA g$^{-1}$, while TiO$_2$ 3DOM only contributes a capacity of 175 mAh g$^{-1}$ (Figure 3b). The results can be further verified by charge/discharge profiles of Ni-TiO$_2$ sandwich 3DOM and TiO$_2$.
3DOM at the 1st, 2nd, 20th, and 50th cycle at the current density of 50 mA g\(^{-1}\) in Figure 3c,d. In accordance with CV data, the appearances of discharge curves also infer that the electrochemical contributions related to the intercalation reactions in bulk together with interface capacitive-like storage. The former storage mainly contributes to the capacities within the discharge plateau, while the other capacities are mainly attributed to capacitive-like storage.

As mentioned before, one of the key challenges within SIBs is to enable high capacity and high power at the same time. To examine this potential, the rate capabilities were recorded in Figure 4a together with corresponding charge/discharge profiles in Figure S4, Supporting Information, to investigate the feasibility of fast charge/discharge. Ni-TiO\(_2\) sandwich 3DOM shows discharge capacities of 280, 193, 135, 120, and 105 mAh g\(^{-1}\) at the current densities of 50, 200, 1000, 2000, and 5000 mA g\(^{-1}\), while the values of TiO\(_2\) 3DOM are 216, 139, 86, 63, and 47 mAh g\(^{-1}\), respectively. Besides the high capacities at high rates, the retention reaches 91% from the 2nd cycles to 1000th cycles at the current density as high as 2000 mA g\(^{-1}\) (Figure 4b). The specific capacity after 1000 cycles can still reach 109 mAh g\(^{-1}\), near onefold larger than that of TiO\(_2\) 3DOM. It implies that high-rate sodium storage of Ni-TiO\(_2\) sandwich 3DOM is feasible, reversible, and stable. Comparison with other current collectors with different geometry, like highly ordered Ni core-shell nanowire array,[17c] randomly oriented Ni nanowire array,[17c] planar morphology as shown in Figures S5 and S6, Supporting Information, Ni 3DOM skeleton serves as an excellent candidate for 3D current collector.

2.3. In-Depth Discussion of Electrochemical Behaves

To deeply understand the origins of the rate performance gap between Ni-TiO\(_2\) sandwich 3DOM and TiO\(_2\) 3DOM, here, four important points should be first predeclared considering the electrochemical behaves. i) According to Raman spectra in Figure 1d, there is no peak found in the range from 1200 to 1800 cm\(^{-1}\), which might be assigned to the signals of carbonaceous materials. We also cannot find any peak belonged to various oxidation products of Ni, which is further convinced by XRD and HRTEM results in Figures 1c and 2e. Meanwhile, we haven’t added any conductive additive and polymeric binder within all the electrodes. Hence, all the electrodes don’t contain any carbonaceous material with the potential to change the long-term stability and rate performance.[18] The entire gap of sodium storage ability at different rates should be only attributed to the electrodes themselves. ii) The thickness of TiO\(_2\) in both architectures are quite similar through concise control of ALD, resulting in the same mass loading and ion diffusion length as shown in Scheme 2. Related impacts from ion diffusion can be excluded. iii) Both kinds

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**Figure 4.** a) The rate performance of Ni-TiO\(_2\) sandwich 3DOM and TiO\(_2\) 3DOM for SIBs with current densities varying from 50 to 5000 mA g\(^{-1}\); b) The cycling stability of Ni-TiO\(_2\) sandwich 3DOM and TiO\(_2\) 3DOM at 2000 mA g\(^{-1}\) for 1000 cycles.

**Scheme 2.** Schematic illustration of the ion diffusion length and electron transport pathways in a) Ni-TiO\(_2\) sandwich 3DOM and b) TiO\(_2\) 3DOM. Blue stands for TiO\(_2\), while orange represents Ni. The yellow arrays stand for ion diffusion. The white and red arrays are four major electron movement events occurring during ion-intercalation and deintercalation. 1) accumulation and/or reaction at the electrode/electrolyte interface; 2) electron transport throughout the active materials; 3) electron injection and extraction at the electrode-current collector interface; 4) electron transport in current collector.
of electrodes are fabricated by the topologic transformation from CCTs, so the morphology features are similar. We can fully exclude related influences from electrode architectures like facet, orientation, etc.\textsuperscript{(19)} It is worth mentioning that TiO\textsubscript{2} sandwich 3DOM even possesses less contact area with electrolyte than TiO\textsubscript{2} 3DOM because of the existence of Ni skeleton and the requirements of the same ion diffusion length, which would slightly decrease the ability of interface storage in a conventional sense. iv) Both kinds of electrodes are stored in N\textsubscript{2} before measurement. It is not necessary to take the detrimental influences from the surface state by absorption and/or adhesion into account. After such careful analysis of the two architectures, the major difference between the two kinds of electrodes is the existence of Ni 3DOM current collector instead of a planar current collector. Hence, we turn our attention to electron transport that might result in a huge change of rate performance of SIBs.

Shown in Figure 5\textsuperscript{a} are the current-voltage (I-V) characteristics of Ni-TiO\textsubscript{2} sandwich 3DOM and TiO\textsubscript{2} 3DOM. The symmetrical and linear appearance of I-V curves indicates the Ohmic contacts between the electrode and conductive substrate.\textsuperscript{(17)} Obviously, the existence of Ni 3DOM skeleton indeed largely facilitates electron transport and reduces the resistance throughout the whole matrix by 20-folds. Facile electron transport leads to decreased charge-transfer resistance (R\textsubscript{ct}) at the surface. According to a modified Randles equivalent circuit in Figure S8, Supporting Information, analysis of Nyquist plots of electrochemical impedance spectroscopy (EIS) ranging from 100k to 0.1 Hz (Figure 5b,c) confirms decreased R\textsubscript{ct} after introducing the Ni 3DOM current collector (Figure 5d). And the gap of R\textsubscript{ct} becomes larger with increased cycles. Since R\textsubscript{ct} mainly depends on the charge transfer resistance related to the redox reactions across the electrode/electrolyte interface,\textsuperscript{(20)} smaller R\textsubscript{ct} suggests much facile reactions and associated sodium storage near or at the interface.

This understanding inspires us to think over a question: how can we associate the electrical conductivity of electrodes with SIB performance at high rates? Due to the same active materials, the accommodation ability of Na ions is the same in theory. So, we try to hunt for the origins based on the possible kinetic factors. CV is a powerful tool to illuminate the electrochemical kinetics of the electrodes towards Na\textsuperscript{+}. Current response to an applied sweep rate will vary depending on whether the redox reaction is diffusion-controlled or not.\textsuperscript{(10b,11a,21)} Hence, we recorded the CV curves of electrodes before and after the introduction of Ni 3DOM current collector at various scan rates from 0.1 to 5 mV s\textsuperscript{-1} (Figure 6a,b). Similar redox pairs also match the potentials of reversible Ti\textsuperscript{4+}/Ti\textsuperscript{3+} redox, consistent with the conclusions from Figure 3a. In general, ion intercalation contains three steps: solvated Na\textsuperscript{+} diffusion within the electrolyte; charge-transfer reactions at or near the interface between active materials and electrolyte; and Na-ion diffusion in the bulk materials. According to the features of these electrochemical behaviors, there are two types of contributions towards the whole sodium storage: Na-ion intercalation in bulk related to diffusion-controlled mechanism and capacitance at or near the interface free of diffusion-controlled mechanism. To visualize the storage mechanism, we seek help...
from the relationship of peak current \(i\) and scan rate \(\nu\) by the following Equations (1,2):

\[
\begin{align*}
  i &= a \nu^b \\
  \log(i) &= b \log(\nu) + a
\end{align*}
\] (1,2)

Here, \(a\) and \(b\) are adjustable constants. \(b\)-value can be obtained from the slopes by plotting \(\log(i)\) against \(\log(\nu)\). \(b\)-value should be 0.5 provided the sodium storage is an ideal faradaic intercalation process controlled by semi-infinite linear diffusion, whereas the \(b\)-value of 1.0 stands for a pure capacitive contribution without diffusion control.[10b,21b] As depicted in Figure 6c,d, both kinds of electrodes display a good linear relationship. The \(b\)-values of Ni-TiO\(_2\) sandwich 3DOM are 0.92 and 0.91 for anodic and cathodic processes, respectively, which are closer to 1 than those of TiO\(_2\) 3DOM (0.83 and 0.80 for anodic and cathodic processes). This value is closed to amorphous TiO\(_2\) 3DOM with mainly surface-controlled capacitive-like contributions during charge/discharge.[8a,15a] Such comparison implies that the sodium storage is less diffusion-controlled after the introduction of Ni 3DOM current collector. It is a strong evidence that facile electron transport resulted from Ni 3DOM current collector can change the electrochemical kinetics and promote the fast interface capacitance-like sodium storage free of diffusion control.

Hence, the interface capacitance-like contribution at a certain scan rate \(\nu\) can be further quantitatively differentiated by separating current response \(i\) at a fixed potential into bulk Na-ion intercalation contribution and interface capacitance-like contribution according to Equation (3):

\[
i = k_1 \nu + k_2 \nu^{1/2}
\] (3)

Here, \(k_1\) and \(k_2\) are adjustable constants. Solving for their values at each potential is helpful for the separation of the diffusion-controlled currents and capacitive-like currents, because the former one is proportional to \(\nu\) while the latter one is proportional to \(\nu^{1/2}\).[10b,11b,21c] Figure 7a–f exhibit the typical CV curves for the capacities from interface capacitance-like contributions (green region) in comparison with the total capacities at the scan rates of 0.2, 1, and 4 mV s\(^{-1}\), respectively. It is obvious to find the quantified results that the interface capacitance-like contribution gradually improves with the accelerated scan rates. At the scan rate of 0.2 mV s\(^{-1}\), \(\approx 63.7\%\) of the total capacity is capacitive in nature. The ratio increases to 90.4% with the improved scan rate to 4 mV s\(^{-1}\). More importantly, the Ni-TiO\(_2\) sandwich 3DOM possesses much larger interface capacitance-like contributions than TiO\(_2\) 3DOM. The interface capacitance-like contribution is only 79.2% of the total sodium ability at 4 mV s\(^{-1}\). Considering the four prerequisites, all the results indicate that facile electron transport by introducing Ni 3DOM current collector can successfully promote the interface capacitance-like sodium storage.

In general, there are four major electron movement events occurring during ion-intercalation and deintercalation as...
depicted by the white arrays in Scheme 2: 1) accumulation and/or reaction at the electrode/electrolyte interface; 2) electron transport throughout the electrode; 3) electron injection and extraction at the electrode-current collector interface; 4) electron transport in the current collector. According to the previous analysis, a comparatively small surface area may slightly weaken the electron utilization at the surface because of a little less accessibility of the active material for Na$^{+}$. But the existence of Ni 3DOM current collector can obviously shorten electron transport pathways in step (2) (within active materials) and amplified the contact area of electrode-current collector interface as shown in Scheme 2. Since electron transport in the current collector is much faster than that in materials and across the interface, this amelioration in the latter two points can largely improve the electron transport and reduce the resistance throughout the entire electrodes, associated with facile interface charge transfer evidenced in Figure 5. As well known, electron movement is completed by a redox reaction in the materials, and thus is much faster than ion diffusion. Hence, optimization of ion diffusion dominates the previous studies. However, accelerated electron transport would also play a vital role in ion transfer at the interface between the material and electrolyte, which has been regarded as a slow process during electrochemical energy storage.\cite{11b,15a} 3DOM metal skeleton can not only serve as a supporter for active materials, but also provides a direct and reduced way for electron transport. First, fast electron transport can avoid the charge accumulation at and/or near the interface, leading to accelerated interface kinetics and decreased potential of meaningless side reactions. Second, fast rearrangement of lattice at the interface can be realized by facile electron supplement or removal. Third, high electronic conductivity can reduce the electrostatic repulsive force, which would fasten the electron transfer and transport in particular on the interface. Overall, the processes of interface capacitive-like storage, like pseudo-capacitance or double-layer capacitance, can be both be optimized together with the ion intercalation/deintercalation of Na$^{+}$. Interface capacitive-like storage always occurs on the order of seconds and minutes, and thus promoted interface capacitive-like storage is particularly expected to improve high-rate capacities in short time.\cite{11a} Moreover, interface capacitive-like storage will not result in a huge change of crystalline structures that can be recovered fast. This would benefit for a well balance of high capacity and high-rate performance at the same time. Hence, it is efficient and significant to employ the sandwich design to largely boost rate performances of SIBs by tuning the electrical conductivity of electrodes.

2.4. Full-Cell Electrochemical Characteristics of Ni-TiO$_2$ Sandwich 3DOMs

Such superiorities can also be used to improve the rate performance in full cells. To check this potential, layered P2-Na$_{0.7}$Ni$_{1/3}$Mn$_{2/3}$O$_2$ is chosen as the cathode to assemble two-electrode full cells together with Ni-TiO$_2$ sandwich 3DOM and TiO$_2$ 3DOM anodes like the scheme in Figure 8a. Corresponding characterizations and electrochemical performance of P2-Na$_{0.7}$Ni$_{1/3}$Mn$_{2/3}$O$_2$ are listed in Figure S9, Supporting Information. Similar to the superior performance in half cells, Ni-TiO$_2$ sandwich 3DOM shows long-term stability and large sodium storage as shown in Figure 8b,c. With respect to the mass of anodes, a capacity of $\approx$210.1 mAh g$^{-1}$ can be obtained even after 100 cycles at 50 mA g$^{-1}$, the value of which is much

![Figure 7. Separation of the capacitive-like and diffusion controlled currents of a–c) Ni-TiO$_2$ sandwich 3DOM and d–f) TiO$_2$ 3DOM at various sweep rates: (a,d) 0.2 mV s$^{-1}$;(b,e) 1 mV s$^{-1}$; (c,f) 4 mV s$^{-1}$.](image-url)
larger than TiO$_2$ 3DOM anode and amorphous TiO$_2$ 3DOM anode\cite{15a} in our previous reports. The stable capacity retentions and high-rate capabilities (Figure S8c) are also attractive in practical applications. The full-cell performance shows the good ability of sodium storage even at a current density as high as 5000 mA g$^{-1}$, and these capacities can be fully recovered when the current density is switched back to 50 mA g$^{-1}$. These results of full cells fully confirm that introducing a metal 3DOM current collector holds the potential to build high-rate energy storage devices.

Given the analysis, our observations confirm the possibilities of improving SIB rate performance by tuning the electrical conductivity of electrodes. Although electron transport is magnitudes faster than that of Na ions,\cite{7b,22} strategies to improve electron transport open a novel and efficient avenue to promote rate performance by optimizing fast interface capacitive-like storage rather than comparatively slow bulk storage. The present understandings are expected to not only extend to other sodium-based energy storage device, but also guide the development of functional materials for efficient energy storage at high rates. We are looking forward to more efforts in these open questions in the near future.

3. Conclusions

In summary, we have successfully developed an oxide-metal sandwich 3DOM as a superior anode candidate for high-rate SIB. Taking Ni-TiO$_2$ sandwich 3DOM as a proof-of-concept, anatase TiO$_2$ delivers a reversible capacity of 233.3 mAh g$^{-1}$ in half cells and 210.1 mAh g$^{-1}$ in full cells (coupled with P2-Na$_{2/3}$Ni$_{1/3}$Mn$_{2/3}$O$_2$ cathode) after 100 cycles at 50 mA g$^{-1}$. At the high charge/discharge rate of 5000 mA g$^{-1}$, 104.4 mAh g$^{-1}$ in half cells and 68 mAh g$^{-1}$ in full cells can also be obtained with satisfying stability, indicating great rate performance of SIBs among various TiO$_2$-based anodes. In-depth analysis of electrochemical kinetics fully evidences that the presented remarkable sodium storage ability, in particular at high charge/discharge rates, origins from dominated interface capacitive-like storage because of increased electrical conductivity by introducing of metal 3DOM current collector. The optimized electron transport can efficiently accelerate the interface kinetics of sodium storage, and thus improve the interface capacitive-like contribution ratio to the total sodium storage. Fast and stable interface capacitive-like storage efficiently facilitates sodium storage at high rates without huge fading off. Different from the optimized strategies focused on ion diffusion, our observations about employing a high electrical conductivity paradigm into electrode design enable a novel and efficient foundation for realizing high-rate SIBs. It is expected to realize a universal design for satisfying energy storage by extending this rule to the secondary ion batteries with other large transport ions. Further combination of ion diffusion optimization holds great promise to attract more research interests in the future development in electrochemical energy storage.

4. Experimental Section

Fabrication of Polystyrene Sphere Colloidal Crystal Templates: The well-ordered CCTs comprised of polystyrene spheres (PSs) were fabricated to form a face-centered cubic packing arrangement using a vertical deposition approach according to the previous works.\cite{23} Ti foils as the substrates have been cleaned under sonication by sequentially immersing in acetone (15 min), ethanol (15 min), and distilled water (15 min). The substrates were then immersed vertically in the 0.5 wt% PS latex at 60 °C. Drying of PS latex appeared near the Ti foil at very low speed in high moisture. Then, CCTs on Ti foil were heated at 90 °C for 5 min and 110 °C for 2 min to enhance the adhesion and avoid a crash during the subsequent steps. Here, the diameter of PSs was ~500 nm after optimization of sodium storage ability.

Fabrication of Ni 3DOM Skeletons: In a typical procedure, 3DOM Ni skeletons were fabricated by electrochemical deposition using an aqueous electrolyte of 1 M NiSO$_4$ · 6H$_2$O, 0.1 M NiCl$_2$ · 6H$_2$O, and 0.5 M H$_3$BO$_3$. The electrochemical deposition was realized by means of a standard three-electrode system consisting of CCTs as the working electrode, Pt plate as the counter electrode, and HgSO$_4$/Hg as the reference electrode. Constant current mode with the current of ~2.45 mA was used here for 30 min by a VSP electrochemical workstation (Bio-Logic, France), where Ni was solidified in situ at the interstice of CCTs to form a 3DOM architecture. All the depostions were carried out at room temperature. Then, the CCTs were removed by immersing into 10 mL tetrahydrofuran (THF) for 24 h. Ni 3DOM skeletons could be obtained after dying in N$_2$ flow without any oxidation and kept in N$_2$.

Fabrication of Ni-TiO$_2$ Sandwich 3DOM: In a typical procedure, TiO$_2$ was deposited on both sides of Ni 3DOM skeleton by ALD (PicoSun SUNALE R150 ALD system, PicoSun, Finland) to achieve the sandwich construction. Here, TiCl$_4$ and H$_2$O were chosen as the precursors of Ti and O, respectively. TiCl$_4$ and H$_2$O were alternately pulsed for 0.1 s with separation by 10 s purge of N$_2$. The cycle numbers were 400 in order to reach a uniform thickness of 20 nm. The reaction chamber was 70 °C. Then the as-prepared sandwich architectures were annealed in air at 350 °C for 2 h with a ramp rate of 2 °C min$^{-1}$ to obtain the anatase phase of TiO$_2$. 

Figure 8. a) Scheme of a two-electrode full cell with TiO$_2$ sandwich 3DOM and TiO$_2$ 3DOM anodes and P2-Na$_{2/3}$Ni$_{1/3}$Mn$_{2/3}$O$_2$ cathode; b) Cycling performance at 50 mA g$^{-1}$; c) Rate performance with current densities varying from 50 to 5000 mA g$^{-1}$.
Fabrication of TiO₂ 3DOM: For comparison, TiO₂ 3DOM was also fabricated by ALD on the basis of CNTs instead of Ni 3DOM skeleton with the same settings. To keep the similar ion diffusion length, the repeated cycles were 800 here, which was twofold of that of sandwich architectures. The top layer was half-removed by a plasma etching used argon plasma reactor (PDC-32G2) at 70 Pa. Then the as-prepared architectures were also annealed in air at 350 °C for 2 h with a ramp rate of 2 °C min⁻¹. Finally, anatase TiO₂ 3DOM could be obtained.

Fabrication of Layered P₂Na₂/3Ni₁/3Mn₂/3O₂: Co-precipitation was used to prepare this layered material. Stoichiometric Ni(NO₃)₂ and Mn(NO₃)₂ were first mixed with NaOH. Na₂CO₃ was then added into the mixtures later as the Na source. Collected by centrifugation, the precursors were calcined in air at 600 °C (4 h) and 900 °C (10 h).

Characterization: XRD patterns were recorded using Cu-Kα radiation at 40 kV and 40 mA(SIEMENS/BRUKER D5000 X-ray diffractometer). The scanned range (2θ) was from 20° to 80° with the rate of 0.02° s⁻¹. SEM measurement was completed on a Hitachi S4800 field emission scanning microscopy. TEM and EDX scan measurements were performed on JEM-2100F (JEOL). XPS measurements were performed at the Photoemission Endstation (BL10B) in the National Synchrotron Radiation Laboratory (NSRL) in Hefei, China. All the binding energies were corrected by referencing C 1s to 284.5 eV. Raman spectrum was detected by Raman microscope with a 514.5 nm Ar laser (inVia Raman microscope).

Electrochemical Measurement: The as-prepared Ni-TiO₂ sandwich 3DOM and TiO₂ 3DOM architectures were directly used as anodes without any conductive additive and polymeric binder. The mass loading of the two samples was around 0.76–0.80 mg. So as to cathodes in full-cells, a mixture of 80% layered P₂Na₂/3Ni₁/3Mn₂/3O₂, 10% acetylene black, and 10% poly(vinylidene fluoride) (PVDF) by weight was pressed onto an Al foil. Then they were kept at 120 °C for 12 h in vacuum before the assembling of the full-cells. Electrochemical tests were performed through a coin cell configuration (CR2012) with a diameter and thickness of 20 and 3.2 mm, respectively, which were assembled in a nitrogen-filled glove box with oxygen and moisture concentrations kept below 0.1 ppm.

Sodium metal foil for half-cells and layered P₂Na₂/3Ni₁/3Mn₂/3O₂ for full-cells were used as counter electrodes with separation from the working electrodes by a glass microfiber filter (Whatman, Grade GF/B) with the pore size of 1 μm. The electrolyte used here was 1 mol L⁻¹ NaClO₄ in propylene carbonate (PC, Aldrich anhydrous 99.7%). All electrochemical measurements were performed at 20 ± 2 °C on a Land CT2001A battery testing system (Land, P. R. China). CV, I-V measurement, and EIS were recorded by a VSP electrochemical workstation (Bio-Logic, France). For CV measurement, scan rates were chosen from 0.1 to 5 mV s⁻¹ in the potential range of 0.01 to 3.0 V (vs Na//Na⁺). The potential range for I-V was −1 to 1 V. EIS spectrum was recorded with a frequency range of 100 kHz to 0.1 Hz and a 5 mV AC amplitude.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
The authors declare no conflict of interest.

Data Availability Statement
The data that supports the findings of this study are available in the supplementary material of this article.

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