Scalable, Large-Area Printing of Pore-Array Electrodes for Ultrahigh Power Electrochemical Energy Storage

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ABSTRACT: Through-electrode thickness honeycomb architectures were layer-by-layer self-assembled directly through a scalable printing process for ultrapower hybrid lithium-ion capacitor applications. Initially, the electrochemical performance of the pore-array electrodes was investigated as a function of the active material type (graphene plates, carbon nanofibers, and activated carbon). Inactive components (conductive carbon and polymer binder) were then minimized to 5 wt %. Finally, an optimized activated carbon-based cathode was paired with a spray-printed Li4Ti5O12-based anode and a range of anode-to-cathode mass ratios in a lithium-ion capacitor arrangement were investigated. A 1:5 anode/cathode mass ratio provided an attractive energy density comparable with a Li4Ti5O12/LiFePO4 lithium-ion battery but with outstanding power capability that was an order of magnitude greater than typical for lithium-ion batteries. The pore-array electrode was reproduced over areas of 20 cm × 15 cm in a double-sided coated configuration, and the option for selectively patterning electrodes was also demonstrated.

KEYWORDS: pore array, spray printing, graphene, carbon nanofiber, YP-50F, Li4Ti5O12, lithium-ion capacitor

Driven by the increasing worldwide requirement for more sustainable energy systems such as renewable generation and storage, the need to replace internal combustion engines with electric drivetrains and the trend for lighter, thinner portable electronics including laptops and mobile phones, the rechargeable lithium-ion battery (LIB) has come to dominate the energy-storage market. The dominance arises from the LIB’s excellent gravimetric and volumetric capacity (200−300 Wh/kg and 600−1000 Wh/L, respectively), a relatively high operating voltage (∼4 V), low self-discharge, and a continued reduction in cost. Nonetheless, its relatively poor power performance (300−400 W/kg), which is almost 2 orders of magnitude lower than that of electrochemical double-layer capacitor (EDLC) cells (10−20 kW/kg), remains a weakness for some applications, particularly for public transport vehicles, heavy machinery, and some applications in the defense and aerospace industry.

Lithium-ion capacitor (LIC) devices aim to combine the high capacity of LIBs and the high power of EDLCs. In a general LIC configuration, an insertion-type electrode is paired with a high-surface-area carbon electrode, with the system immersed in a standard LIB electrolyte. Although conventional lithiation/delithiation reactions take place at the negative electrode, delivering an attractive capacity (much higher than EDLCs but lower than LIBs), the carbon positive electrode stores electric energy by quickly forming a reversible electrical double layer at the interface between the electrode and the electrolyte, offering fast electrode kinetics and a cycle life superior to LIBs. However, because positive and negative electrode processes must remain in charge balance, the overall performance is a compromise between that of a LIB and an EDLC. To realize the potential advantages of LICs more fully, a less-comprised balance of power and capacity is needed that better preserves the energy density of LIBs and the power performance and cycle stability of EDLCs.

Porous or hollow structures at the meso- or a few micron-scale increase active surface area and promote faster electrode reaction dynamics. Unfortunately, relatively sophisticated and expensive processes are often required to fabricate these porous structures inhibiting their scalability and applicability.

We explore a spray-printing approach to fabricate higher-performing LIC electrode configurations, which has previously been shown to be effective in fabricating novel LIB and EDLC electrode arrangements, including high density graded architectures and multilayer arrangements in a diverse range of materials that includes TiO2(B), Si/SiO2, TiO2, Fe3O4, carbon nanofibers, and LiFePO4 (LFP). Here, the focus is on the production of large-area pore arrays for LIC electrodes. By controlling the drying behavior of the electrodes on heated current collectors, the electrode porosity was controlled reliably over large-area electrodes that showed improved gravimetric...
and volumetric densities at high charge/discharge rates (≥4000 mAh/g) when compared with conventional equivalents manufactured by slurry casting.

Through-thickness pore-array positive electrodes for a LIC are fabricated by spray printing using either graphene plates (GP), carbon nanofibers (CNF), or commercial activated carbon YP-SO particles (YP). The electrochemical performance of the resulting electrodes is investigated along with the effects of conductive carbon enhancer (Super P, SP) and polymeric binder (carboxymethyl cellulose, CMC). A best-performing pore-array YP-based cathode electrode is coupled with a spray-printed Li4Ti5O12 (LTO) negative electrode in a LIC configuration that delivered a stable energy density of ~55 Wh/kg at ~0.1C and an outstanding power density of ~10000 W/kg at ~450°C. The scalability of the through-thickness pore-array electrode (~20 cm × 15 cm) and the possibility to assemble the pore-array structure selectively in a specific region of the electrode are also demonstrated. We conclude that the spray-printing approach may offer a credible route to manufacture anodes and cathodes with novel functionality (pore arrays and selective patterning) that provide a larger design space to balance energy and power requirements in electrochemical energy-storage devices.

## RESULTS AND DISCUSSION

The scheme illustrated in Figure 1 describes the self-assembly of through-thickness pore-array electrodes directly on a large-area substrate by a variant of a spray-deposition or spray-printing process, which atomizes and deposits a slurry-based suspension. Active carbon materials (GP, CNF, or YP), SP, and CMC were dispersed in a fugitive bisolvent with an optimum deionized water to isopropyl alcohol (DI/IPA) volume ratio of 50:50 that provided a rheologically stable and low viscosity suspension suitable for pumping to a spray head and then continuous atomization into droplets. The droplets were propelled by the atomizing gas onto the heated current collector (>120 °C), which enabled the fast, effective evaporation of bisolvent carriers in each of the printed layers. A pore-array structure was spontaneously formed during drying that propagated from the current collector and which was maintained as subsequent layers were added, as described in the magnified cartoon in Figure 1. The zig-zag movement of the spray head in a preprogrammed x- and y-direction pattern for multiple cycles allowed the fabrication of the pore-array structure over large areas (typically >20 cm × 15 cm) and control of electrode thicknesses up to hundreds of microns.

Figure 2a–c shows scanning electron microscope (SEM) images of the surface of a series of pore-array electrodes based on either GP, CNF, or YP, respectively. Here, all electrodes contained the active carbon material, SP, and CMC in a mass ratio of 80:10:10. The pore-array electrodes had pores of 20 ± 9 μm diameter that were separated by interconnected pore-array cell "walls" of thickness 10–20 μm. The insets in the right corner of Figure 2a–c magnify the region of the active carbon material, showing ~2 μm size GP, ~100 nm diameter CNF, and ~5 μm size YP, respectively. Figure 2d–f shows the corresponding electrode cross-sections, with the layer-by-layer formation of through-thickness pore structures, as highlighted by yellow dotted lines. As shown in the previous work,20,21 these oriented pores can provide benefits in terms of facilitating lithium-ion mobility and effective electrode/electrolyte contact, offering an improvement in capacity and most particularly rate performance. All of the electrodes were ~20 μm thick (see Table 1), and thicker electrodes were studied subsequently.

The electrochemical behavior of the electrodes was examined through cyclic voltammetry (CV) in the scan rate range of 0.05–50 mV/s, as shown in Figure 3a–c. The pore-array electrodes were combined with Li metal foil counter electrodes in a coin-cell configuration. All of the electrodes displayed quasi-rectangular CV curves at increasing scan rates, which is typical of electrochemical double-layer capacitance. Progressive increases in the scan rate to 50 mV/s resulted in a progressive increase in current density for all of the electrodes. However, the difference in the anodic/cathodic current density at increasing scan rates became more marked, depending on the active carbon material, as shown in Figure 3a–c. The specific capacitance C [F/g] was estimated from the CV curves according to

$$C = \frac{1}{m \nu} \int_{V_c}^{V_s} I(V) dV$$

where m is the total electrode mass [g], ν is the scan rate [V/s], $V_s - V_c$ is the potential window [V], and $I$ is the charging or discharging current [A]. In the arising capacitance plots (Figure 3d), the pore-array GP-based electrode had the highest capacitance at all rates, for example, ~62 F/g at 0.05 mV/s and ~32 F/g at 50 mV/s, whereas the capacitance of the pore-array GP and CNF equivalents dropped to ~1 and ~5 F/g at 50 mV/s, respectively, as shown in the magnified inset of Figure 3d (also see Table 1). The electrochemical impedance spectroscopy (EIS) data in Figure S1 of the Supporting Information also implied that the pore-array YP structure had the lowest charge-transfer resistance, which positively contributed to its comparatively high performance.

Electrode behavior as a function of the variations in the electrochemically inactive components (SP and CMC) was then investigated using the best-performing pore-array YP electrode, with the intention of minimizing the parasitic inactive fraction. Figure 4a–c shows a series of SEM images for the surface of pore-array YP electrodes. As the inactive component reduced progressively from 20 to 5 wt %, the YP fraction increased correspondingly to 95 wt %, the typical pore diameter increased from ~10 to ~30 μm, and the pore density decreased from ~700 to ~300 mm−2 because of the increased fraction of relatively large YP particulate. The corresponding electrode cross-sections (Figure 4d–f) show that the directional pore channels of ~20
μm depth were again maintained “deep” into the electrode to the current collector.

**Figure 5a** shows the corresponding electrode-specific capacitance as a function of the inactive fraction estimated from the CV data (see **Figure S2** in the Supporting Information). As the YP fraction increased to 95 wt % (and the inactive component fraction reduced correspondingly to 5 wt %), capacitance increased progressively at all rates. Note that the capacitance was estimated using the entire electrode mass including YP, SP, and CMC, as given in **Table 2**. At 5 wt % inactive additives, the capacitance was ~105 F/g at 0.05 mV/s and ~50 F/g at 50 mV/s. In **Figure 5b**, the 95:3:2 electrode formulation had a higher capacity of ~66 mAh/g at 10 mA/g and ~40 mAh/g at 2000 mA/g that is comparable with or higher than other synthesized activated carbon materials.10,27–29 **Figure S5c** displays the galvanostatic capacity profiles at 100 mAh/g where the pore-array YP electrode with minimum inactive additives (≤5 wt %) sustained a stable capacity of ~55 mAh/g up to 500 cycles, representing ~100% capacity retention (compared with the first discharge capacity), while the 90:5:5 and 80:10:10 formulation had capacities reduced to ~50 and ~40 mAh/g, respectively. All of the YP-based electrodes had ~100% Coulombic efficiency after 500 cycles (inset of **Figure 5c**).

To investigate the influence of the pore array on the electrode performance specifically, a nonporous YP electrode was prepared by slurry casting using the same 95:3:2 electrode formulation (denoted as “conventional”), as shown in **Figure S3b** of the Supporting Information. **Figure S3c** shows the gravimetric discharge capacity profiles of both electrode types in the current density range of 10–2000 mA/g (see electrode details in **Table S1** of the Supporting Information). At relatively low rates of 10–50 mA/g, the electrode structures delivered almost identical capacities, suggesting that the pore-array structure made no marked contribution to the electrode kinetics and there was sufficient time for ions to transport throughout the electrode to all of the active surfaces. However, a difference in deliverable capacity became progressively more marked as the current density increased to 2000 mA/g. Eventually, the pore-array YP electrode had an approximately 70% higher discharge capacity. Volumetric discharge capacity in **Figure S3d** further supported the performance benefit of the pore-array electrode at increasing rates, where volumetric capacity was re-estimated based on the mass and thickness of electrodes (see **Table S1**). At 10 mA/g, the conventional electrode had a slightly improved capacity because the electrode was comparatively dense. However, at 2000 mA/g, the pore-array electrode facilitated Li mobility and outperformed the conventional electrode in volumetric capacity, even though it had a higher pore fraction.

![Figure 2. SEM images of the surface of the pore-array electrodes based on (a) GP, (b) CNF, and (c) YP. Each of the inset micrographs magnifies the active carbon material. (d−f) Corresponding electrode cross-sections. The yellow dotted lines indicate a directional pore channel from the electrode surface to the current collector.](image1)

![Figure 3. CV profiles at the increasing scan rate in the potential range of 2.0−4.0 V (vs Li/Li⁺) for pore-array electrodes based on (a) YP, (b) CNF, and (c) GP. (d) Corresponding capacitance profiles with an inset that magnifies the capacitance plots for the pore-array GP and CNF electrodes.](image2)

**Table 1. Summary of the Pore-Array Electrodes Based on GP, CNF, and YP**

| active materials | thickness (μm) | loading mass (mg/cm²) | 0.05 mV/s | 1 mV/s | 10 mV/s | 50 mV/s |
|------------------|----------------|-----------------------|-----------|--------|---------|---------|
| GP               | 20 ± 2         | 1.23 ± 0.03           | 17.6      | 7.1    | 2.9     | 1.1     |
| CNF              | 20 ± 3         | 1.23 ± 0.02           | 25.6      | 8.9    | 5.9     | 4.7     |
| YP               | 20 ± 3         | 1.25 ± 0.05           | 62.1      | 58.8   | 47.6    | 31.6    |
The potential of a carbon-based pore-array positive electrode in a LIC configuration was investigated by pairing it with a spray-printed LTO negative electrode, as depicted in the schematic of a coin-type LIC cell in Figure S4 (also see the spray-printed...
LTO performance in Figure S5 and Table S2 of the Supporting Information. Spinel-type LTO is considered promising for LIC applications because its approximately zero-strain lithium-insertion behavior supports stable capacity delivery even at ultrafast charging rates (>10 C) and during continuous cycling (>500 cycles). In addition, LTO can be utilized without the need for a time-consuming prelithiation step. A LTO/YP mass ratio in the range 1:1 to 1:5 was then investigated in a LIC configuration. Here, a 10 μm thick LTO negative electrode was fixed, and the YP-based positive electrode weight and the corresponding thickness was varied, from 20 to 80 μm (see Table 3 and SEM images in Figure S6 of the Supporting Information). The relatively thin LTO electrodes were chosen to prioritize high power performance rather than volumetric energy density. Figure 6a shows the discharge Ragone plots at various charge/discharge rates for LTO-YP LICs as a function of the LTO/YP mass ratio up to 1:5. The specific power \( P \) and energy density \( E \) were estimated using \( P = V \times i/m \) and \( E = P \times t/34 \), where \( V \) is the average cell potential [V] between the lowest and highest cut-off potential, \( i \) is the obtained current [A], \( m \) is the total weight of both anode and cathode electrodes including LTO, YP, SP, and CMC, and \( t \) is the discharge time [h]. As the LTO/YP mass ratio increased from 1:1 to 1:5, specific energy and power density increased progressively, with the 1:5 LTO/YP arrangement having a specific energy density of \(~52 \text{ Wh/kg} \) at \(~0.1 \text{ C} \) and a specific power density of \(~9900 \text{ W/kg} \) at \(~450 \text{ C} \) (see the trend in volumetric Ragone profiles in Figure S7 of the Supporting Information). At 1:6 LTO/YP, the power performance started to reduce to \(~7500 \text{ W/kg} \) at \(~450 \text{ C} \) (see comparative Ragone plots in Figure S8 of the Supporting Information).

The rate capability of the optimized LIC cell was compared with that of a LIB formulated with a 1:1 LTO/LFP mass ratio that had the best rate capability in our previous and other studies. For these LIB configurations, LTO and LFP electrodes were prepared by the identical spray-printing process using a suspension formulated with a 95:3:2 LTO (or LFP)/SP/CMC mass ratio suspended in a 70:30 DI/IPA bisolvent that was intended to avoid any pore self-assembly (see half-cell performances of each of LTO and LFP electrodes in Figure S5 and Table S2 of the Supporting Information). Figure 6b shows the comparative gravimetric charge/discharge plots for the LIB (the upper gray plots) and LIC (the lower red plots) in the rate range of 0.1–100 C, noting that performances of both the LIB and LIC were estimated based on the total mass of the LTO anodes, i.e., including LTO, SP, and CMC. At 0.1 C, the LIC and the LIB had similar capacities of \(~165 \text{ mAh/g} \) and \(~155 \text{ mAh/g} \), respectively. However, as the C-rate increased up to 100 C, the difference in capacity between the LIC and LIB became more marked. At 100 C, the LIC had a discharge capacity of \(~110 \text{ mAh/g} \), sustaining \(~75 \% \) of its discharge capacity at 0.1 C, while the LIB capacity had dropped to \(~10 \text{ mAh/g} \), one order of magnitude lower than that of the LIC at the same rate.

The cycle life of the 1:5 LTO/YP LIC cell was investigated at a constant rate of 10 C (Figure 6c). As the charge/discharge cycle progressed to the 500th cycle, the energy density declined relatively rapidly, delivering \(~85 \% \) of the initial energy density (\(~45 \text{ Wh/kg} \) at the first cycle). However, the reduction was then less marked so that even after 2000 cycles, the energy density was \(~32 \text{ Wh/kg} \) (\(~70 \% \) energy density retention), as shown in the right-hand inset of Figure 6c.

A photograph in the left-hand inset (Figure 6c) shows a double-sided pore-array YP electrode over an A5-scale Al current collector (20 cm × 15 cm). It is worth noting that scalable, double-sided coating is essential for the manufacture of multiple cell packs for high-energy-storage applications including electric cars and grid energy storage. Further, the ability to locate the pore-array pore structure selectively on any desired region of the electrode may be useful in achieving an attractive balance between gravimetric and volumetric capacity (or power and capacity) for large-area electrodes (see a photo of...
patterning the letters “OX” on one side of the pore-array YP electrode in Figure S9 of the Supporting Information). For example, if all of the areas of electrodes comprised the pore-array structures only, outstanding gravimetric capacity would be realized at ultrafast charging rates but only with relatively poor volumetric performance at slow rates. However, if the pore-array structure can be designed and assembled only in the desired area, e.g., at pouch cell corners that are distant from the current collector tabs where active material utilization or ion distribution is often relatively poor at ultrafast rates, more balanced power and capacity performance might be achieved.

Overall, the through-thickness pore-array architecture can be expected to facilitate the dispersion of active ions to all of the accessible active surface areas within the electrode, as depicted schematically in Figure 7a. For example, the LIC constructed with pore-array electrodes provided an approximately 40 % improvement in power capability over the identical LIC arrangement with nonporous, conventional electrodes at 400 C: a specific power density of ∼9900 W/kg at ∼24 Wh/kg for the pore-array LIC (red) and a power density of ∼7200 W/kg at ∼17 Wh/kg for the conventional LIC (gray), as shown in the comparative Ragone curves in the inset of Figure 7b. The pore-array-based LIC is also compared with other LIC cells in Figure 7b. At relatively slow charging rates, the energy performance was comparable with other LTO/activated carbon-based LIC configurations. However, at ultrafast charging rates (≥100 C), the pore-array LIC outperformed similar configurations (see detailed comparison in Table S3 of the Supporting Information), noting that many of the competing performance data relied on conductive enhancers (e.g., carbon coating or graphene additives) to boost rate capability or utilized higher fractions of parasitic inactive components (≥20 wt %), the weight of which was generally not taken into account in the performance estimation.

## CONCLUSIONS

Honeycomb patterns through the thickness of electrochemical electrodes were layer-by-layer self-assembled over large-area current collectors (~20 cm × 15 cm) using a spray-printing approach. Deposition used an atomized suspension formulated with an optimum 50:50 mixture (by volume) of fugitive DI water and IPA. Initially, the electrochemical capacitance of pore-array electrodes as a function of active component types (CNF, GP, and YP) was studied, and, subsequently, the effect of minimizing inactive components (SP and CMC) was investigated in the range of 20–5 wt %. The best-performing pore-array YP cathode (5 wt % inactive fraction) was then coupled with a spray-coated LTO anode to investigate performance in a LIC arrangement as a function of the YP/LTO mass ratio. A 1:5 LTO/YP combination offered outstanding LIC performance, with an energy density of ∼52 Wh/kg at 0.1 C and a power density of ∼10000 W/kg at 450 C. The through-thickness pore-array electrode architecture was particularly advantageous where fast electrode kinetics and promotion of active surface area were required, e.g., up to 400 C charge/discharge rates. Spray printing also facilitated a selective patterning approach that might be used to enhance local ion mobility, such as selected areas in large-area pouch cells that otherwise suffer from “dead zones” of poor material utilization.

## EXPERIMENTAL SECTION

### Materials

CNF and LTO were purchased from Sigma Aldrich, U.K.; GP from Thomas Swan & Co. Ltd., U.K.; YP from Kuraray, Japan; and LFP from Hydro-Québec, Canada.

### Spray-Printing Process

For the spray suspensions, active carbon (GP, CNF, or YP), SP, and CMC in various mass ratios were well-blended in a mixture of DI water and IPA using sonication. Although DI water was used to dissolve CMC, the IPA was needed to disperse the carbon materials into aqueous suspension. For the electrode fabrication, an Al foil was placed on a hot plate (~120 °C). The active carbon-based suspension was pumped to a spray nozzle and, subsequently, atomized in a spray cone of suspension droplets at a constant air pressure of 0.3 bar, which deposited onto the heated foil continuously as the spray reciprocated over the foil at a constant distance in a preprogrammed path. The spray-printing process was conducted in a well-ventilated fume cupboard.

### Electrochemical Tests

The electrochemical behavior of electrodes was examined using coin-type electrochemical cells (CR2032) that were assembled in an Ar-filled glovebox (H₂O < 0.1 ppm, O₂ < 0.1 ppm). For half-cell tests, electrodes were coupled with Li foils (99.9 % trace metals basis, MTI Corporation) that were used as a counter/reference electrode. For both half-cells and full LICs, full LIBs, negative and positive electrodes were electrically separated by a polyethylene separator (Cellgard 2400) soaked into a 1 M LiPF₆ electrolyte solution in a 1:1 mixture (by volume) of ethylene carbonate and dimethyl carbonate (Merck). LIC cells were assembled without a prelithiation process. All charge/discharge tests were performed at room temperature using an Arbin BT2000 cycler. CV and EIS measurements were carried out using a Gamry 600 potentiostat. The theoretical capacity of both LTO and LFP was assumed to be ~175 mAh/g.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.9b14478.

Nyquist plots of pore-array electrodes as a function of the active material type, CV profiles for the pore-array YP electrodes according to a change in inactive components, comparative capacity profiles for the YP-based pore-array...
and conventional electrodes, schematic diagram of the coin-type LIC cell, performances of spray-printed LTO and LFP electrodes, SEM images of the 80 µm YP pore-array electrode, volumetric Ragone plots of pore-array LICs, LIC performance as a function of LTO/YP mass ratios and comparative LIC performances, and a photo of a selectively patterned YP pore-array electrode (PDF).

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**Notes**

The authors declare no competing financial interest.

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