Hybrid Carbon Spherogels: Carbon Encapsulation of Nano-Titania

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Extraordinarily homogeneous, freestanding titania-loaded carbon spherogels can be obtained using Ti(acac)2(OiPr)2 in the polystyrene sphere templated resorcinol-formaldehyde gelation. Thereby, a distinct, crystalline titania layer is achieved inside every hollow sphere building unit. These hybrid carbon spherogels allow capitalizing on carbon’s electrical conductivity and the lithium-ion intercalation capacity of titania.

Hybrid carbon nanomaterials play a crucial role in various high-performance applications such as energy storage devices (supercapacitors, batteries), photocatalysis, heterogeneous catalysis, or absorption. There are multiple strategies to obtain such hybrids, including bottom-up (sol-gel) and top-down (atomic layer deposition) approaches.1, 2 When implementing metal oxides onto/into carbon structures, it is vital to choose the most suitable carbon substrate carefully3 and to design the pore structure.4 The carbon selection must be based on accessibility, electrical conductivity path length, and chemical compatibility.5 Hollow carbon spheres are a very attractive scaffold for encapsulation and offer a high surface area, high homogeneity, and electrical conductivity. The hollow carbon spheres’ void inner space can serve as a storage reservoir, chemical reactor, or drug delivery host.6–8 The microporous carbon shell acts as a permeable membrane and allows species to diffuse through and fill or drain the interior or initiate a chemical reaction. This nanoreactor concept was recently demonstrated for the growth of carbon helixes on copper, placed in a carbon hollow sphere material.9 Strategies to build-up functionalized hollow carbon materials can be differentiated between post-synthesis or in-situ approaches. A typical thermal build-up process is the impregnation of hollow carbon spheres with a molten compound, for example, sulfur,10 germanium,11 or copper.12 Other examples of processes for the design of hybrid carbon hollow sphere materials include the ship-in-bottle principle,13 selective etching,14 spray pyrolysis, pyrolysis of metal-organic framework (MOF) materials,15 or layer-by-layer assembly16. Such hybrid carbon materials, for example, monolithic titania-loaded carbons, have shown promising results regarding their potential in lithium-ion batteries.17 This study presents a versatile template-assisted sol-gel route based on resorcinol-formaldehyde to form monolithic, titania-loaded hybrid carbon spherogels with exceptionally high morphological control. The material properties such as the carbon shell’s pore structure or titania/carbon mass ratio can be deliberately adjusted by thermal treatment in CO2. Our method is a general approach that holds the potential of synthesizing an array of different hybrid carbon spherogels loaded with a variety of metal oxides by choosing adequate metal precursors.

Our study reveals a versatile synthesis approach to monolithic, hybrid carbon/titania spherogels with extraordinarily high morphology control. Based on our previously described method to carbon spherogels,18 Ti(acac)2(OiPr)2 as a titanium precursor is added to the resorcinol-based sol solution. A scheme of the sol-gel process is provided in Fig. 1. Monodispersed, approximately 250 nm-sized (Fig. S1 ESI, prepared by emulsion polymerization of styrene)19 polystyrene spheres (3 mass%) in aqueous solution serve as templates for the subsequent resorcinol-formaldehyde gelation process. First, titanium (IV) resorcinol species are formed and arrange around the negatively charged outer surface of the polystyrene spheres (as confirmed by the negative Zeta potential of -20 mV). Thereby, a homogeneous distribution of the Ti(IV) precursor is achieved. During gelation at 80 °C, resorcinol condensation with
formaldehyde coats each Ti-layer and interconnects the spheres to form a 3D gel network. After supercritical drying with carbon dioxide, we can reduce linear shrinkage effects to less than 20%. Template removal, carbonization, and titania formation coincide during thermal treatment at 800 °C in argon. Exemplary, Fig. 2A shows a photograph of the freestanding, cylindrical, and black-colored hybrid, titania-loaded carbon spherogel (CS-TiO₂/C 800). Thermogravimetric analysis revealed a complete polystyrene template removal at 470 °C and a ceramic yield (titania) of 22 mass% (Fig. S2 ESI, and Table 1). For the low resorcinol-to-water molar ratio of 0.008, SEM (Fig. 2A) images verify that the low-density (as measured to be 0.09 g/cm³), macropore-containing spherogel sample is solely composed of hollow spheres. TEM images (Fig. 2B-C) confirm the highly homogeneous structure where each hollow sphere subunit is composed of a distinct titania layer of a thickness of approximately 7 nm (dark grey circle due to pronounced elemental contrast) encapsulated by carbon. This is also supported by elemental examination (EDX) via scanning transmission electron microscopy (STEM), which confirms a carbon shell and a titania decoration of the inside wall of the sphere (Fig. 2D-E). This as-prepared titania-loaded carbon spherogel sample (CS-TiO₂/C 800) was amorphous (Fig. 3A) with an average domain size of ca. 3 nm (as estimated from a STEM high angle annular dark field image; Fig. S3 ESI).

To confirm the nature of the inner titania coating, X-ray photoelectron spectroscopy was performed after heating in an argon atmosphere (Fig. S4 ESI). For the pristine carbon spherogel, a C1s signal at 284.4 eV could be observed, with minor impurities from oxygen traces (O1s signal at 532.7 eV) and the corresponding C1s signal at 286.4 eV. With titania loading, a Ti 2p signal at 458.9 eV could be observed, corresponding to TiO₂. In the C1s and the O1s signal, new species at 286.9 eV and 535.1 eV align with a TiO₂ layer formation. The O1s signal can be assigned to TiO₂, and the C1s signal results from the TiO₂/carbon spherogel interphase.

The titania crystallite size can be modified by thermal heating as we confirmed in-situ X-ray diffractionmetry with temperature steps of 50 °C between 800 °C and 1100 °C under Ar (Fig. 3A). The crystal growth onsets at 900 °C. Samples annealed at 1100 °C are composed of 13 mass% anatase and 87 mass% rutile, as shown by Rietveld analysis. The change in the crystalline size of titania by thermal treatment is also confirmed by TEM (Fig. 2D-F). While the sample at 800 °C displays a homogeneous titania shell (Fig. 2D), the latter breaks up into individual grains (ca. 10 nm) confined within the carbon sphere after treatment at 1100 °C (Fig. 2F). Instead of annealing in an inert gas atmosphere, we can further modify the titania-to-carbon ratio by carbon dioxide etching. For example, annealing at a temperature of 800 °C in the presence of carbon dioxide for 30 min results in etching of carbon concomitantly with increasing titania content to 35 mass%, and individual titania grain growth. This CO₂ treatment concurrently consumes carbon via oxidation and modifies the titania crystals. These findings were confirmed by small-angle X-ray scattering (SAXS), showing a decrease of intensity at low angles (low scattering vector length), consistent with locally flat structures (Fig. 3B). A pronounced shoulder at larger angles is observed, which shifts to smaller angles upon annealing. This aligns with approximately 2 nm-sized structures, which grow to about 7 nm during annealing at 1100 °C. Thermal activation in CO₂ leads to structures of 15 nm diameter within the shell (Fig. S6 ESI). The Raman spectra for all titania encapsulated samples (as-synthesized, annealed to 1100 °C and 30 min carbon dioxide etched) show disordered carbon as seen from a D-mode at ~1345 cm⁻¹ and a G-mode at ~1584 cm⁻¹ (Fig. 3C). The presence of titania is proven by the presence of four distinct Raman modes in the range of 150-650 cm⁻¹. Nitrogen sorption isotherms were recorded at -196 °C (Fig. S7 ESI) and demonstrate the thermal treatment effect on porosity. The titania-loaded carbon spherogel carbonized at a maximum temperature of 800 °C consists of a dense nano-crystalline titania layer surrounded by a microporous carbon shell and macropores between the spheres. This structure corresponds to a typical type IV isotherm, representing a micro/macroporous material. Thus, the sphere interior’s access is limited by the titania nano-crystalline layer, and the H₄ hysteresis is only weakly pronounced. By contrast, annealing to 1100 °C initiates titania crystal growth and creates pores within the layer, making the interior accessible. In this case, the
desorption isotherm shows a distinct H4 hysteresis loop, typical for macropores enclosed by a microporous shell. The sharp drop at p/p0 at 0.5 reveals the sudden evaporation of nitrogen from the sphere interior, named tensile strength effect (TSE).24-26 By activating the sample for 30 min with carbon dioxide, the hysteresis effect can be even increased, while micropore and mesopore content are enlarged. The corresponding pore size distributions obtained by carbon dioxide sorption verify the sphere shells with mainly micropores <1 nm (Fig. 3D). Therefore, we also observe an increase of the calculated (QSDFT) specific surface areas (SSA) from 713 m2/g for the as-synthesized sample to 900 m2/g at 0.5 reveals the sudden evaporation of nitrogen from the sphere interior, named tensile strength effect (TSE).24-26 By activating the sample for 30 min with carbon dioxide, the hysteresis effect can be even increased, while micropore and mesopore content are enlarged. The corresponding pore size distributions obtained by carbon dioxide sorption verify the sphere shells with mainly micropores <1 nm (Fig. 3D). Therefore, we also observe an increase of the calculated (QSDFT) specific surface areas (SSA) from 713 m2/g for the as-synthesized sample to 900 m2/g for the sample treated at 800°C for 30 min in carbon dioxide (Table 1).

We carried out cyclic voltammetry to characterize the electrochemical behavior to demonstrate the titania nanocrystals' electrochemical activity in the TiO2-loaded carbon spherogels. The prepared samples were tested in the potential range of 1.0-3.0 V vs. Li+/Li at various scan rates of 0.1-50 mV/s. The resulting cyclic voltammograms of CS-TiO2/C 800 and CS-TiO2/C 30min for the first and fifth cycle at 0.1 mV/s are shown in Fig. 4A. There are no pronounced oxidation/reduction peaks seen in the cyclic voltammogram of CS-TiO2/C 800; instead, we observed a pronouncedly pseudocapacitive response.27 The 1st and 5th cyclic voltammogram of CS-TiO2/C 30min exhibit a pair of reduction/oxidation peaks at 1.7 V and 2.0 V vs. Li+/Li, which can be ascribed to the Li-ion insertion/extraction in an anatase TiO2 lattice, respectively.28 The increased pseudocapacitive contribution between 3.0 V and 2.0 V vs. Li+/Li may align with micropores’ presence within the carbon shell; thereby, ion electroosorption may contribute to the total charge storage process. Comparing the 1st and 5th cycles, we see no noticeable change in the insertion/delithiation peak position. Compared to highly crystalline anatase, there is an increased pseudocapacitive current between 3.0 V and 2.0 V vs. Li+/Li; this contribution becomes particularly evident compared to the current amplitude of the lithiation/delithiation peak. This contribution is most likely aligned with the nanoscopic feature of the anatase domains. One can further explore the pseudocapacitive feature via kinetic analysis, that is, analysis of the rate-dependent current signal. A b-value of 1 aligns with a diffusion-limited process typical for battery-like processes, while a b-value of 0.5 is expected for an ideal ion electroosorption process.27 As seen in Fig. 5B, the pronounced lithiation/delithiation peak of CS-TiO2/C 30min shows a b-value of 0.6, while the more pseudocapacitive region at 1.25 V vs. Li+/Li has a slightly enhanced b-value of 0.62.29 The capacitor-like response of CS-TiO2/C 800 aligns with a b-value of ~0.8 throughout the potential range of 1.5-3.0 V vs. Li+/Li.

The obtained reduction and oxidation peaks from cyclic voltammetry agree with the galvanostatic discharge and charge profiles in Fig. 4C-D. For the CS-TiO2/C 800 sample, no plateaus and the associated (redox) reactions can be detected either. In general, a maximum discharge capacity of 12 mAh/g is obtained at low specific currents. In the CO2-treated CS-TiO2/C 30min sample, we see two plateaus at 1.7 V and 1.4 V vs. Li+/Li during the 1st and 5th cycle at a scan rate of 0.1 mV/s. (A) Cyclic voltammogram at a scanning rate of 0.1 mV/s. (B) Galvanostatic charge/discharge cycling with corresponding Coulombic efficiency at different values of current density. (C) CS-TiO2/C 800 and (D) CS-TiO2/C 30min. (E-F) Rate performance using galvanostatic charge/discharge cycling with corresponding Coulombic efficiency at different specific currents for (E) CS-TiO2/C 800 and (F) CS-TiO2/C 30min.
The Coulombic efficiencies are given in Fig. 4E-F. Both tested materials show the expected behavior of proportional decreasing de-lithiation capacity while applying higher specific current and satisfying capacity retention after returning to the initial current. A significantly higher initial specific de-lithiation capacity of 137 mAh/g was obtained from CS-TiO<sub>2</sub>C 30min with a Coulombic efficiency of 86%. The CS-TiO<sub>2</sub>C 800 sample achieved only a de-lithiation capacity of 11 mAh/g with a Coulombic efficiency of 80%. The Coulombic efficiencies of both samples approach ~100% by the 5<sup>th</sup> cycle and maintained over 98% during the rate capability testing sequence. A specific current of 1 A/g CS-TiO<sub>2</sub>C 30min electrodes shows 22 mAh/g and recovery of ca. 100%, while CS-TiO<sub>2</sub>C 800 sample no longer provides any significant charge storage capacity. Fig. 4B shows the cycling stability at 0.1 A/g of both samples. For CS-TiO<sub>2</sub>C 30min, there was a loss of 61% capacity after 500 cycles, while CS-TiO<sub>2</sub>C 800 loses nearly all its initial capacity.

For electrochemical application, we see the importance of modifying the CS-TiO<sub>2</sub>C 800 sample further. By reducing the carbon content during the CO<sub>2</sub> heat treatment, it was possible to obtain characteristic electrochemical behavior comparable to that found in the literature. The ability to modify the TiO<sub>2</sub>C ratio, TiO<sub>2</sub> properties, carbon porosity, and the apparent option to change from titania to other metal oxides make hybrid carbon spherogels an interesting electrode material for electrochemical applications.

**Notes and references**