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Layered Si₂Te₃-based anodes for highperformance Li-ion batteries[†]

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We develop a scalable solid-state synthesis for Si₂Te₃ and its carbon composite (Si₂Te₃@C) to address the challenges of silicon-based anodes in lithium-ion batteries (LIBs). The Si₂Te₃@C composite exhibits enhanced electrochemical performance, including high reversible capacity, excellent cycling stability, and improved rate capability, positioning it as a promising candidate for highperformance LIB anodes.

Layered materials, characterized by strong in-plane covalent bonding and weak out-of-plane van der Waals (vdW) interactions, have garnered significant interest due to their unique physical and chemical properties.^{1–3} These materials, including graphite, black phosphorus, transition metal chalcogenides (TMCs), and MXenes, allow for the intercalation of foreign atoms or molecules between layers, making them attractive as electrode materials for lithium-ion batteries (LIBs). Notably, graphite serves as a representative anode material due to its reversible Li-ion intercalation mechanism, while other layered materials, such as phosphides, chalcogenides, carbides, and nitrides, are actively studied for their high capacity and structural advantages, positioning them as promising alternatives for next-generation LIB electrodes.^{4–7}

Silicon (Si) has been widely recognized as a promising anode material for LIBs due to its abundance, low redox potential (0.2–0.3 V vs. Li⁺/Li), and exceptionally high theoretical capacity (3578 mA h g^{-1} for Li_{3.75}Si at room temperature).⁸ However, Si-based anodes suffer from severe structural degradation caused by 300–400% volume expansion during lithiation/delithiation processes.⁹

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This dramatic volume change leads to the mechanical fracture of particles, electrical disconnection of active materials, and the formation of dynamic interfaces at the electrode-electrolyte boundary. Such dynamic interfaces not only degrade electrical contact but also promote undesirable side reactions with the electrolyte, resulting in the continuous growth of the solid electrolyte interphase (SEI) layer, ultimately causing capacity fading and poor cycle life.¹⁰ To address these intrinsic limitations of Si anodes, extensive efforts have been devoted to improving their performance through various strategies, including carbon coatings, composite structures, size and morphology control, and the development of Si-alloys and Si-compounds.¹¹⁻¹⁴ Among these approaches, Si-alloys and Si-compounds have emerged as particularly promising solutions due to their structural stability and distinct Li-storage mechanisms, which effectively mitigate the effects of volume expansion and improve cycling performance.

Recently, Si₂Te₃, a two-dimensional (2D) silicon-based chalcogenide, has attracted considerable attention as a highperformance material for energy storage applications. It exhibits excellent environmental stability, a strain-dependent bandgap, and a high Seebeck coefficient, making it highly versatile.^{15,16} Its trigonal layered crystal structure with a stoichiometric ratio of Si/Te = 2/3 supports the formation of natural structural defects through the partial absence of Si-Si pairs, resulting in p-type behavior and enhanced light-matter interactions.¹⁷ These features contribute to improved photoelectric conversion efficiency and a broad photoresponse range, making it suitable for optoelectronic devices as well. Furthermore, the layered structure of Si₂Te₃ provides large interlayer spacing, which facilitates lithium-ion diffusion during lithiation/delithiation processes, addressing the volume expansion issue associated with conventional Si anodes. Its indirect bandgap nature and photoelectronic properties further expand its potential applications beyond energy storage, particularly in optoelectronics. Despite its promising structural and electronic properties, the scalable synthesis of Si₂Te₃ remains a challenge. Therefore, developing simple and cost-effective methods for producing Si₂Te₃ is crucial to fully realize its potential as a high-performance anode material in LIBs.

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In this study, we present a facile and scalable approach for synthesizing layered Si₂Te₃ through a solid-state process that combines sequential ball-milling (BM) and heat-treatment (HT). While Si₂Te₃ exhibits a unique atomic arrangement, its electrochemical performance is largely influenced by its intrinsic properties. To overcome its inherent low electrical conductivity, a composite structure incorporating amorphous carbon is developed. The Li-storage properties of this composite are further evaluated, demonstrating its potential as a highperformance Si-based anode for LIBs. Moreover, its Li-storage mechanism is systematically analyzed using ex situ X-ray diffraction (XRD) and high-resolution transmission electron microscopy (HR-TEM) based on the dQ/dV plots. This work provides a comprehensive understanding of the electrochemical behavior of Si2Te3-based composites and offers useful design strategies for next-generation LIB anodes.

Silicon-tellurium (Si–Te) compounds are primarily known to form two distinct phases, SiTe₂ and Si₂Te₃.^{17,18} However, the Si–Te phase diagram identifies Si₂Te₃ as the only thermodynamically stable compound in this system (Fig. S1, ESI†). Si₂Te₃ crystallizes in a hexagonal structure with a $P\bar{3}1c$ (no. 163) space group and exhibits lattice parameters of a = b = 7.4 Å and c = 13.5 Å. It features a layered crystal structure with each layer having a thickness of 6.7 Å. The layers consist of two parallel planes of tellurium atoms, with Si–Si dimers occupying the interlayer regions.¹⁷ The interlayer spacing in Si₂Te₃ is approximately 3.2 Å, which is comparable to well-known layered materials such as graphene (~3.4 Å), black phosphorus (~3.2 Å), and MoS₂ (~3.2 Å). This relatively large interlayer spacing facilitates rapid diffusion and reversible accommodation of Li-ions, making Si₂Te₃ a promising candidate for LIB anode materials.

Si₂Te₃ was synthesized via a solid-state reaction involving sequential BM and HT processes. The XRD patterns of the synthesized Si₂Te₃ matched well with those of the hexagonal Si₂Te₃ phase (JCPDS #75-0820), confirming that the material was formed homogeneously without impurity phases (Fig. 1a and Fig. S2, ESI[†]). To further investigate its structural properties, the XRD pattern was refined using the Rietveld method (Fig. 1b). The corresponding lattice parameters $-\chi^2 = 1.01$, $R_p =$ 15.5%, and R_{wp} = 19.1% (Table S1, ESI[†])—showed excellent agreement with experimental data, indicating the high crystallinity of the synthesized material. The morphology and particle size distribution of Si₂Te₃ were examined using scanning electron microscopy (SEM) and particle size analysis (PSA). As shown in Fig. S3 (ESI†), the average particle size was approximately 10.01 µm, with irregular particle morphologies. To further validate the homogeneous synthesis, Raman spectroscopy and HR-TEM analyses were conducted. The Raman spectra exhibited three distinct peaks at 139, 291, and 504 cm⁻¹, which were clearly distinguishable from those of Si and Te (Fig. S4, ESI^{\dagger}), confirming the formation of the Si₂Te₃ phase. Additionally, HR-TEM images (Fig. S5, ESI⁺) revealed well-developed Si2Te3 crystallites with clearly visible diffraction patterns (DPs) corresponding to the hexagonal structure. The observed crystal planes, including $(\bar{1}\bar{1}4)$, (110), $(\bar{1}\bar{1}2)$, and $(\bar{1}\bar{1}6)$, matched the main peaks detected in the XRD patterns, further

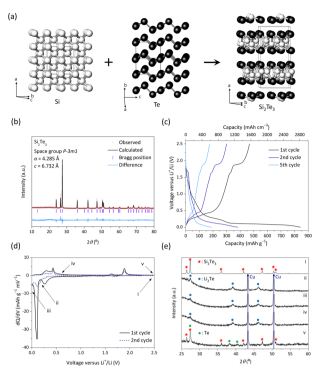


Fig. 1 (a) Crystal structure of layered Si₂Te₃; (b) Rietveld-refined XRD patterns of Si₂Te₃; (c) GDC profiles of Si₂Te₃ anode at 100 mA g⁻¹; (d) dQ/dV plots for Si₂Te₃; (e) *ex situ* XRD patterns of Si₂Te₃ anode during first cycle.

verifying the structural integrity and high crystalline quality of the synthesized Si_2Te_3 .

To evaluate the electrochemical Li-storage properties of the Si₂Te₃ anode, galvanostatic discharge and charge (GDC) profiles were recorded at a current density of 100 mA g^{-1} (Fig. 1c). For comparison, Si and Te anodes were also prepared and tested electrochemically using bulk Si and Te powders. The Si anode exhibited a high initial lithiation/delithiation capacity of 3511/ 1925 mA h g^{-1} (Fig. S6, ESI[†]). However, its capacity rapidly degraded to 282 mA h g⁻¹ after 10 cycles due to severe volume expansion (>300%) caused by the alloying/dealloying reactions with LirSi phases. In contrast, the Te anode demonstrated a lower lithiation/delithiation capacity of 199/133 mA h g⁻¹ and suffered from poor capacity retention due to its low electrical conductivity. Compared to these materials, the Si₂Te₃ anode exhibited an initial lithiation/delithiation capacity of 843/468 mA h g⁻¹ (2816/ 1563 mA g cm⁻³) with an ICE of 55.5%. Thus, Li-storage performance of Si₂Te₃ is also not sufficient to meet the criteria for highperformance LIB anodes compared with Si-based anodes.

To analyze the Li-storage mechanism of the Si_2Te_3 anode, *ex situ* XRD analysis was performed based on the dQ/dV plot. The dQ/dV plot exhibited two prominent peaks during the discharge process and three peaks during the charge process (Fig. 1d). *Ex situ* XRD analysis (Fig. 1e) revealed that as the potential decreased from open circuit potential (i) to 0.17 V (ii), the Si₂Te₃ phase disappeared, and only the Li₂Te phase (JCPDS #77-2147) was detected, confirming the conversion reaction of Si₂Te₃ into Li₂Te and amorphous Si. Further analysis at 0 V (iii) revealed that the Li₂Te phase remained. To verify the formation of the Li–Si phase after full lithiation, the dQ/dV plots for Si and Si₂Te₃ anodes were compared (Fig. S7, ESI†).¹⁹ The overlapping peak positions confirmed that the formation of the Li_{3.75}Si phase was consistent in both anodes. Upon charging to 0.6 V (iv), the Li₂Te phase persisted, suggesting partial recombination during the initial charging process. At the fully charged state (2.5 V, v), the Si₂Te₃ phase reappeared. However, the Te phase was also detected, indicating that the Si₂Te₃ phase was partially recombined, with residual Te and amorphous Si coexisting. These results demonstrate that the Si₂Te₃ anode undergoes a reversible conversion reaction, although the presence of residual Te and amorphous Si suggests partial phase reconstruction during the charging process.

To further enhance the Li-reversibility and reactivity of Si_2Te_3 , we designed a Si_2Te_3 (a)C composite by combining the as-synthesized Si₂Te₃ with carbon black using a BM process. Carbon-supported composites are well-known to improve ionic and electronic conductivity, accommodate large volume changes, and suppress the agglomeration of nanoscale Li-active materials during repeated cycling.^{20,21} The introduction of conductive carbon matrices not only facilitates charge transfer kinetics but also stabilizes the structural integrity of the Si₂Te₃ particles, enabling prolonged cycle life and rate performance. The XRD pattern of the Si₂Te₃@C composite revealed broadened diffraction peaks, indicating a reduction in crystallinity during the composite formation process (Fig. S8, ESI⁺). This suggests that the BM process induced a partial amorphization of the material, which could contribute to enhanced electrochemical properties by providing shorter diffusion paths for Li ions. HR-TEM analysis further confirmed that the synthesized Si2Te3@C composite consists of ultrafine nanocrystallites (\sim 5–7 nm) uniformly embedded within an amorphous carbon matrix (Fig. 2a-c). The DPs obtained from selected area electron diffraction (SAED) matched well with the hexagonal crystal structure of Si₂Te₃, supporting the structural stability of the nanocrystallites within the composite. Energy-dispersive X-ray spectroscopy (EDS) mapping derived from scanning TEM (STEM) images verified the homogeneous distribution of elemental Si (yellow) and Te (green) throughout the carbon matrix (blue), highlighting the well-mixed composition of the composite. In addition, the average particle size of Si₂Te₃@C was approximately 6.8 µm, with irregular particle morphologies (Fig. S9, ESI[†]).

Fig. 2d shows the GDC profiles of the Si₂Te₃@C anode at a current density of 100 mA g⁻¹. The initial lithiation/delithiation capacities were 816 mA h g⁻¹ (1746 mA h cm⁻³) and 559 mA h g⁻¹ (1196 mA h cm⁻³), respectively. Considering the first irreversible capacity of the carbon component (30 wt%, 110 mA h g⁻¹, Fig. S10, ESI[†]), the Si₂Te₃@C composite exhibited significantly improved Li reversibility and reactivity compared to the pristine Si₂Te₃ anode. In addition, electrochemical impedance spectroscopy (EIS) and the corresponding Nyquist plots were analyzed (Fig. S11, ESI[†]). The charge transfer resistance (R_{ct}) of the Si₂Te₃@C nanocomposite anode was measured as 34.9 Ω , which is significantly lower than that of the pristine Si₂Te₃ anode (59.7 Ω). Furthermore, the nanocomposite exhibited a higher Li-ion diffusion coefficient of 1.23 × 10⁻¹⁵ cm² s⁻¹, compared to 1.11 × 10⁻¹⁴ cm² s⁻¹ for the

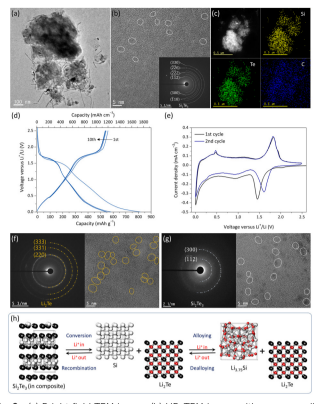


Fig. 2 (a) Bright field TEM image, (b) HR-TEM image with corresponding DPs and, (c) STEM image with corresponding EDS mapping images of Si₂Te₃@C; (d) GDC profiles of Si₂Te₃@C at 100 mA g⁻¹; (e) CV plots of Si₂Te₃@C; *ex situ* HR-TEM images with corresponding DPs of Si₂Te₃@C after (f) full discharged potential (0 V) and (g) full charged potential (2.5 V); (h) schematic of the crystallographic phase change mechanism of Si₂Te₃@C.

pristine Si₂Te₃, as calculated from the $Z_{\rm re}$ -angular frequency ($\omega^{-1/2}$) relationship in the low-frequency region.

To investigate the Li-storage mechanism, ex situ XRD analysis was performed at selected potentials based on the dQ/dVplot and cyclic voltammetry (CV) results. The CV plot displayed two distinct peaks during Li insertion/extraction, suggesting a conversion-type reaction mechanism (Fig. 2e). Although XRD patterns were collected at five cut-off potential points, all patterns exhibited an amorphous nature, likely due to the low crystallinity of Si_2Te_3 within the composite (Fig. S12, ESI^{\dagger}). Thus, to investigate the structural changes of Si₂Te₃@C, ex situ HR-TEM analysis is performed and the results are shown in Fig. 2f and g. At the fully discharged potential (0 V), welldispersed nanocrystallites of Li2Te (5-8 nm) and amorphous Li_{3.75}Si phases were observed, confirming that Si₂Te₃ was fully converted into these phases during the discharge process (Fig. 2f). In contrast, at the fully charged potential (2.5 V), only the Si₂Te₃ phase was detected, indicating that Li_{3.75}Si and Li₂Te recombined to reform Si₂Te₃ with 5-7 nm nanocrystallites during the charging process (Fig. 2g and h).

These findings confirm that the Si_2Te_3 @C nanocomposite undergoes a reversible conversion reaction, where the Listorage process is facilitated by the conducting carbon matrix. The carbon matrix not only improves electronic conductivity but also provides structural stability, effectively buffering the

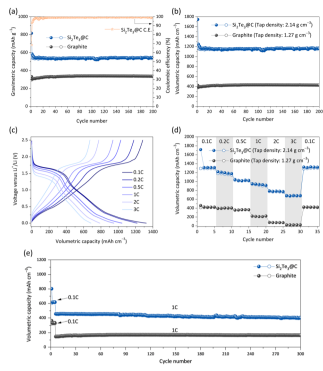


Fig. 3 (a) Gravimetric and (b) volumetric capacities vs. cycle number for Si₂Te₃@C and graphite anodes at 100 mA g⁻¹; (c) GDC profiles of Si₂Te₃@C anode at different C-rates (1C: 550 mA g⁻¹); (d) rate capability of Si₂Te₃@C (1C: 550 mA g⁻¹) and graphite (1C: 300 mA g⁻¹) anodes; (e) cycling performance of Si₂Te₃@C (1C: 550 mA g⁻¹) and graphite (1C: 300 mA g⁻¹) anodes at 1C-rate.

volume changes associated with the lithiation/delithiation processes. In addition, the formation of the Li₂Te phase during lithiation, derived from Si₂Te₃ in the composite, could contribute to the electrochemical performance of LIB anodes.²²

Fig. 3a and b illustrate the electrochemical performance of the Si₂Te₃(a)C composite anode compared with a commercial graphite anode at a cycling rate of 100 mA g^{-1} . The Si₂Te₃@C composite exhibited a highly reversible gravimetric/volumetric capacity of 816/559 mA h g⁻¹ and 1746/1196 mA h cm⁻³, respectively, which are higher than those of the graphite anode $(332/296 \text{ mA h g}^{-1} \text{ and } 421/376 \text{ mA h cm}^{-3})$. In addition, the reversible capacity was well-retained after 200 cycles, maintaining approximately 97.3% capacity retention with high Coulombic efficiency per cycle. The rate capability of the Si2Te3@C composite anode, as a function of the C-rate (1C = 550 mA g^{-1}), is shown and compared with that of the graphite anode (Fig. 3c, d, and Fig. S13, ESI[†]). At cycling rates of 0.1, 0.2, 0.5, 1, 2, and 3C, the Si₂Te₃@C composite displayed high reversible capacities of 1291, 1180, 1029, 938, 775, and 672 mA h cm $^{-3}$, respectively, outperforming the graphite anode. Moreover, during long-term cycling performance of the Si₂Te₃@C composite anode at a cycling rate of 1C (550 mA g^{-1}), the reversible capacity remained at 402 mA h g^{-1} (861 mA h cm⁻³) with stable capacity retention of 88% after 300 cycles (Fig. 3e). This excellent electrochemical performance can be attributed to the ultrafine Si₂Te₃ nanocrystallites (5-7 nm) embedded within the amorphous carbon support, which effectively provide short Li-ion diffusion paths, enhance ionic and electronic conductivity, and mitigate the effects of volume expansion during lithiation/delithiation processes (Fig. S14, ESI[†]).

In summary, this study demonstrates the successful synthesis of layered Si₂Te₃ through a scalable solid-state process combining sequential BM and HT. To address the inherent limitations of Si₂Te₃, including low electrical conductivity and structural instability, a Si2Te3@C composite was developed by incorporating amorphous carbon. The carbon matrix effectively enhanced electronic conductivity, provided structural stability, and mitigated volume changes during lithiation/delithiation processes. Electrochemical evaluation revealed that the Si2Te3@C composite exhibited high reversible capacities (559 mA h $g^{-1}/1196$ mA h cm^{-3}) and excellent cycling stability, retaining $\sim 97.3\%$ of its initial capacity after 200 cycles. It also demonstrated superior rate performance, maintaining ~ 402 mA h g⁻¹ (861 mA h cm⁻³) at 1C after 300 cycles with 88% capacity retention. Ex situ XRD and HR-TEM analyses confirmed a reversible conversion reaction mechanism involving the formation and recombination of Li2Te and Li375Si phases, facilitated by the conductive carbon matrix. These findings highlight the potential of Si2Te3-based composites as high-performance anode materials for next-generation LIBs and provide valuable insights for the design of layered chalcogenide materials with enhanced electrochemical properties.

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Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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