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New insights into the electrochemical performance of precursor derived Si(Nb)OC composites as anode materials for batteries†

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This work represents a first attempt to synthesize Si(Nb)OC ceramic composites through the polymer pyrolysis or the precursor-derived ceramics (PDC) route for use as a hybrid anode material for lithiumion batteries (LIB). Electron microscopy, X-ray diffraction, and various spectroscopy techniques were used to examine the micro/nano structural features and phase evolution during cross-linking, pyrolysis, and annealing stages. During the polymer-to-ceramic transformation process, *in situ* formation of carbon (so-called "free carbon"), and crystallization of t-NbO₂, NbC phases in the amorphous Si(Nb)OC ceramic matrix are identified. The first-cycle reversible capacities of 431 mA h g⁻¹ and 256 mA h g⁻¹ for the as-pyrolyzed and annealed Si(Nb)OC electrodes, respectively, exceeded the theoretical Li capacity of niobium pentaoxide or m-Nb₂O₅ (at approximately 220 mA h g⁻¹). With an average reversible capacity of 200 mA h g⁻¹ and close to 100% cycling efficiency, as-pyrolyzed Si(Nb)OC demonstrates good rate capability. X-ray amorphous SiOC with uniformly distributed nanosized Nb_2O_5 and graphitic carbon structure likely provides stability during repeated Li^+ cycling and the formation of a stable secondary electrolyte interphase (SEI) layer, leading to high efficiency.

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Introduction

LIBs have been utilized in various technological applications since 1991. One major contributor to the commercial success of LIBs has been the development of the graphite anode, which replaced soft carbons and permits stable cycling of Li⁺ at low redox potentials.1,2 However, increasing demand for fastcharging LIBs in the transportation industry and highpowered devices such as power electronics has necessitated the next-generation batteries with superior charge capacity and excellent rate capability, performance characteristics that are not possible in a traditional Li/graphite cell due to thermal/ chemical degradation of graphite at high currents.3 Semiconducting materials such as niobium pentoxide (Nb2O5)-based materials and their composites have garnered interest as alternate anode materials for LIBs because they demonstrate high chemical stability and minimal volumetric expansion during the lithiation and de-lithiation of Li⁺ ions. In addition, the potential window (1-2 V) of Nb₂O₅-based materials is higher

Despite the relatively high operating potential, however, Nb₂O₅-based materials demonstrate sluggish electronic conductivity that can lead to low reversible capacity if cycled at high currents, and low values of specific charge capacities.4 Previous researchers have improved electronic conductivity and Li ion diffusivity in Nb₂O₅ by adding conductive agents such as carbon-graphene or carbon nanotubes^{6,7} or by building nanostructures via synthesis processes such as hydrothermal process or electrospinning process.8,9 Moreover, specific charge capacities of Nb₂O₅ compounds primarily depend on their unique crystal structures, such as pseudohexagonal (TT-Nb₂O₅), orthorhombic (T-Nb₂O₅), monoclinic (M-Nb₂O₅), and amorphous (a-Nb₂O₅).8-12 Research has shown that these polymorphic structures can be produced by altering the heattreatment temperatures during the production process through precursor route.13

Silicon (Si) and Si-based materials have become promising alternative anode materials for LIBs with high charge capacities $(\sim 2 \times)$. Silicon also has the advantages of being environmentally friendly, abundant, and inexpensive, while having less toxicity Nevertheless, inherent disadvantages of Si-based anodes include poor electrical conductivity, large volume expansion upon alloying/dealloying, low coulombic efficiency (CE), unstable SEI, and electrode swelling, all of which prevent its widespread commercialization. Other research has

than the potentials (<0.8 V) at which a SEI forms and resulting in increased safety for the electrochemical energy storage devices.⁵

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reported that ${\rm Nb_2O_5}$ coating of silicon particle leads to high reversible capacity (2256 mA h g $^{-1}$ after 80 cycles) and excellent rate capability of 754 mA h g $^{-1}$ after 80 cycles even at a high current density of 1500 mA h g $^{-1}$.¹⁹ It also reported that pseudocapacitance contributions arising from the niobium oxide nanoparticles and the silicon–niobium oxide interface enhance the electrochemical reaction kinetics and improve the fast charge storage.²⁰ Further a core shell structured Si nanoparticles@TiO2/C mesoporous micro-fiber composite as an anode for LIBs have been reported high reversible capacity (1260 mA h g $^{-1}$), and excellent rate capability (939 mA h g $^{-1}$, after 25 cycles).²¹

Polymer derived ceramics (PDC) such as silicon oxycarbides (SiOC), silicon carbonitrides (SiCN) have been well recognized for their characteristic amorphous structure that depends on the chemistry of the initial precursors and pyrolysis conditions. ^{22,23} Recent research has shown that amorphous PDCs can store Li⁺ ions and these materials demonstrate higher useable capacity and increased rate performance than conventional graphite materials. ^{24,25} PDC-based composites with nanostructured materials such as graphene and carbon nanotubes have also been investigated for their use in LIBs. ¹⁷ The current literature shows that transition metal modified PDCs *via* precursor route improves their mechanical, functional, and electrical properties. ²⁶⁻³²

In an earlier study, titanium-modified SiOC (or Si(Ti)OC) was synthesized through pyrolysis of titanium butoxide-modified polymethylhydrosiloxane (PMHS) single source precursor and investigated as potential anode material for LIBs.33 The aspyrolyzed Si(Ti)OC electrode demonstrates good rate capability due to its unique structure of amorphous SiOC matrix with graphene-like free carbon phase and uniformly distributed transition metal oxide crystals. Additionally, upon annealing the Si(Ti)OC electrode showed improved first-cycle reversible capacity because of the presence of conductive TiC and carbon phases, addressing the key shortcoming with the SiOC anode material. Also, the precursor route provides an advantage of greater control over precursor chemistry and crystallization of desired phases in the pyrolyzed ceramic. Further research is required to ensure safety for practical applications because the electrochemical performance improvement was mainly in the voltage below 0.5 V for SiOC/metal-modified SiOC materials. Herein, the SiOC has been modified with niobium oxide (via pyrolysis of niobium ethoxide (NE) modified PMHS), owing to Nb₂O₅'s working potentials in the range of 1-2 V. Such working potentials avoid Li dendrite formation and ensures safety for battery at high power densities. To our knowledge, this is the first report involving use of Si(Nb)OC ceramic (prepared via pyrolysis of niobium ethoxide modified PMHS precursor) as electrode material in LiBs.

2. Experimental details

2.1 Synthesis of Si(Nb)OC composites

PMHS and NE from Sigma-Aldrich (India) were used as initial precursors to synthesize niobium-modified silicon oxycarbides (Si(Nb)OC) composites. The PMHS modification was done by

adding 50 vol% of NE and stirring for 2 h to form a homogenous solution that was then crosslinked in an oven for 3 h at 250 °C and a heating rate of 5 °C min⁻¹ under ambient atmosphere. The crosslinked product was pyrolyzed for 1 h at 900 °C and a heating rate of 5 °C min⁻¹ under an argon atmosphere. The pyrolyzed product was then annealed for 3 h at 1200 °C and a heating rate of 5 °C min⁻¹ under an argon atmosphere. For comparison, pure NE was heat-treated at 900 °C and 1200 °C at a heating rate of 5 °C min⁻¹ under an ambient atmosphere.

2.2 Electrode preparation and electrochemical characterization

As-pyrolyzed and annealed Si(Nb)OC composite powders as active materials (~160 mg) were mixed with polyvinylidene difluoride (PVDF) (~20 mg) as a binding agent and carbon black (CB) (\sim 20 mg) as a conducting agent in a weight ratio of 8:1:1. One or two drops of 1-NMP (1-methyl-2-pyrrolidone) (\sim 2 ml) was added for fluidity. A doctor blading process was used to coat on the copper foil with the fluid mixture, and the coated copper foil was dried in an oven at 80 °C for approximately 12 h. The dried copper foil was punched into required diameter of 14.28 mm circles, and its weight was measured. Copper-coated electrodes and lithium foil (reference electrode) were used to make the CR2032 half cells in a glove box under an inert atmosphere. The electrodes were separated with a monolayer membrane (polypropylene) wetted with 1 M lithium hexafluorophosphate (LiPF₆) (~2 ml) in 50% dimethyl carbonate (DMC) and a 50% ethylene carbonate electrolyte solution (Sigma-Aldrich, USA). The coin cells were tested in a multichannel battery testing system (BT200 Arbin test system, Texas, USA) with a given voltage range (10 mV-2.5 V) and constant current density (50, 100, 200, 400, 600, and 800 m Ag^{-1}).

2.3 Materials characterization

Functional groups of the precursor and cross-linked powders were analyzed via Fourier transform infrared spectroscopy (FTIR, PerkinElmer Spectrum 400, USA). Thermogravimetry analysis (TGA, SDT-Q600 TA Instruments, USA) was carried under an argon atmosphere at a heating rate of 10 °C min⁻¹ on cross linked powders. X-ray diffraction (XRD) (Cu K_α radiation) (~0.1546 nm, 40 kV, and 40 mA, Bruker AXSD8 Discover X-ray diffractometer, USA) was used to characterize the heat-treated powders. Free carbon in the heat-treated powders was analyzed via Raman spectroscopy (LabRamaramis, USA) with a HeNe laser wavelength of 632.8 nm, and X-ray photoelectron spectroscopy (Thermo Scientific Al Kα + XPS/UPS system, USA) with a beam energy of 148.6 eV was used to determine the elemental composition and bonding nature of the heat-treated samples and post-cycling battery-tested samples. The Ar⁺ was sputtered for 120 s at 3.0 keV to remove surface contamination. Morphological information for the powder samples and postcycling samples of the Si(Nb)OC composites were obtained from the scanning electron microscope (SEM) (Carl Zeiss EVO MA10, 5-30 kV, USA). Before transferring to the SEM chamber, the post-cycling coin cells were dismantled in an argon-filled glove box and repeatedly rinsed with DMC to remove surface impurities of the anode. Nanostructural features and elemental mapping were imaged with a transmission electron microscope (TEM) (JOEL JEM2100 plus, 80–200 keV, USA) and energy dispersive spectroscopy (EDS) (Oxford Instruments, USA).

Results and discussion

3.1 Crosslinking reactions, morphological features and thermal analysis

The possible reaction process during crosslinking is predicted in Fig. 1a, here-in Si-H groups of PMHS react with Nb-O-CH₃ groups of NE to give off hydrocarbons (C_nH_{2n+2}) which resulted in Si-O-Nb linkage. The presence of Si-O-Nb linkages indicates

that the Si(Nb)OC was successfully synthesized through single precursor route from PMHS and NE liquid precursors, in which Nb–O units were incorporated into the siloxane network. By preventing the loss of low molecular weight precursor components as well as their breakdown during ceramization, crosslinking increases the yield of ceramics. The absorption bands at 754 cm⁻¹, 830 cm⁻¹, 1030 cm⁻¹, 1260 cm⁻¹, 1408 cm⁻¹, 2161 cm⁻¹, and 2968 cm⁻¹ could be attributed to Si–C, Si–H, Si–O–Si, Si–CH₃, Si–CH₃, Si–H, and CH₃ characteristic bands, respectively.²⁷ However, the extra absorption band for the modified PMHS at 924 cm⁻¹ may be attributed to Si–O–Nb linkage with the release of hydrocarbons.^{28,34} Fig. 1c and d exemplify the SEM microstructures of as-pyrolyzed and

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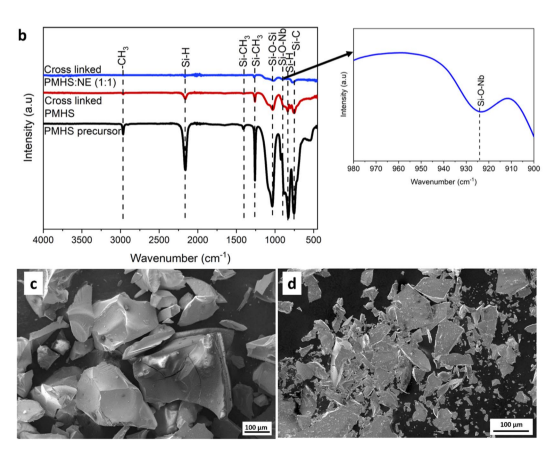


Fig. 1 (a) Possible crosslinked reaction process of NE modified PMHS, (b) FTIR spectrum of liquid PMHS precursor, crosslinked powders of PMHS and NE modified PMHS (1:1) (magnified images Si–O–Nb linkage), and SEM images of Si(Nb)OC composites (c) as-pyrolyzed powder, (d) annealed powder showing irregular morphology.

annealed powder samples of Si(Nb)OC composites respectively. The powder particles exhibited irregular morphology and a typical brittle feature of glassy SiOC ceramics.

In ESI Fig. S1[†] exemplifies thermogravimetric analysis on crosslinked SiOC and crosslinked Si(Nb)OC under an argon atmosphere, exhibiting that the initial polymeric mixture showed a mass loss in three stages. The first stage (25-300 °C) corresponds to the evaporation of moisture present in crosslinked powders of SiOC and Si(Nb)OC, whereas the second stage (300-500 °C) in SiOC and Si(Nb)OC. It could be attributed to the decomposition of the polymer and the release of the methane and hydrogen gases, respectively. The decreased ceramic yield is caused by siloxane evaporation as a result of its lower molecular weight and inadequate degree of cross-linking. Here, Si(Nb)OC showed increasing thermal stability and ceramic yield, possibly due to suppressed decomposition of Si-O-C bonds. There was no evident mass loss beyond 600 °C. NE and PMHS undergo chemical reaction that produces highly cross-linked preceramic polymer.

To begin with, binary SiC and Si₃N₄ are unable to dissolve either nitrogen or carbon to form compositions that contain SiCN. The production of SiOC ceramics is also impossible using SiO₂ and SiC. This molecular method is the sole way to create the ternary SiOC and SiCN variants. 23,35,36 Preceramic polymers called poly(organosiloxanes) can be employed to produce silicon oxycarbide-based ceramics by thermal breakdown in an inert gas environment. Through the application of an appropriate thermal curing procedure, polymer precursors, which are inorganic systems, provide ceramic materials a specific chemical composition and an ordered nanostructure.24 The content, quantity, and distribution of phases, as well as the microstructure of the finished ceramic, are all influenced by the kind and molecular structure of the preceramic polymer. As a result, it is possible to control the macroscopic chemical and physical characteristics of PDCs by changing the molecular precursor. The first step a preceramic polymer goes through after shaping is cross-linking. Thermal cross-linking is frequently taken into account when the elemental and phase composition of finished ceramics needs to be governed.35 Further, by reacting them with transition metal alkoxides, polysiloxanes with relevant functional groups (such hydroxyl or alkoxy) may be chemically modified.35 During crosslinking, in supplementary Fig. S2† the hydroxyl group of PMHS reacting with methyl groups of NE resulting in modified Si(Nb)OC cross linked preceramic polymer product. Depending on the type of polymer, the crosslinked preceramic polymers must be pyrolyzed at temperatures between 400 and 1400 °C in inert or reactive environment.36 When cross-linked polysiloxanes are pyrolyzed, silicon oxycarbide (SiOC) glasses are produced During the pyrolysis process, the crosslinked precursors' organic groups undergo thermolysis and escape as hydrocarbons/hydrogen at temperatures, between 600 and 1000 °C.24 Ceramization takes place as a result of the escape of hydrogen and hydrocarbons, primarily CH₄. Si-O, Si-C, and Si-H bonds can also undergo a variety of redistribution processes. 24,35,36 Amorphous Si(Nb)OC and residual free carbon are produced in ceramic materials as a result of substantial C–H, Si–C, and Si–O bond breakage that takes place at temperatures between 600 and 1000 $^{\circ}$ C.

3.2 Phase evolution studies

X-ray diffractograms of as-pyrolyzed and annealed Si(Nb)OC composites are presented in Fig. 2a. As shown in the figure, the as-pyrolyzed sample exhibits a completely amorphous nature. However, crystalline peaks at 25.8°, 34.8°, and 52.1° are present in the annealed Si(Nb)OC sample, that represents the metastable t-NbO $_2$ phase (ICSD-34425). The annealed sample also shows 2θ peaks at 34.84°(111), 40.35° (002), and 58.4° (022) that could be attributed to the most intense peaks of niobium carbide (NbC), (ICSD-44496). Previous research has reported that chemically modified Si(Nb)OC, when heat-treated above 1000 °C, forms NbO $_2$ or NbC, depending on the availability of carbon per the following equations:²⁸

$$Nb_2O_5 + C \rightarrow 2NbO_2 + CO$$
 (1)

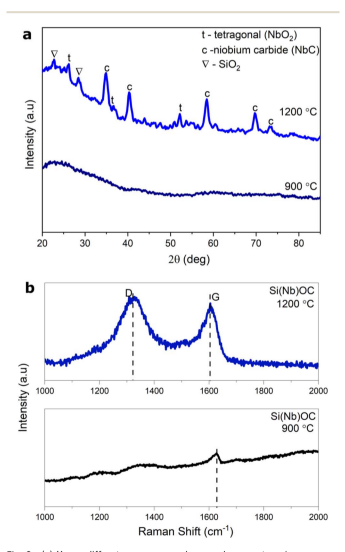


Fig. 2 (a) X-ray diffractograms reveal amorphous nature in as-pyrolyzed sample and crystallization of phases in annealed sample of Si(Nb) OC composite, and (b) Raman spectrum of as-pyrolyzed and annealed Si(Nb)OC composites.

$$Nb_2O_5 + 7C \rightarrow 2NbC + 5CO$$
 (2)

The average crystallite size for the t-NbO2 and NbC calculated using Scherrer's equation is 11 nm and 12 nm, respectively. Further, peaks at 22.3° (011) and 32.28° (012) corresponds to the first most intense peaks of SiO₂ (ICSD-44268). The Rietveld refinement of the Si(Nb)OC composites using MAUD software (ESI Fig. S3†) contains 8.93 wt% of NbC, 4.38 wt% of t-NbO₂, 6.84 wt% of SiO₂ and 79.83% of SiOC.

Fig. 2b shows the Raman spectroscopy data of as-pyrolyzed and annealed Si(Nb)OC samples, including the existence of bands around 1340 cm⁻¹ and 1610 cm⁻¹, which confirms the carbon was in the form of D (disordered) and G (graphitic). The D and G peak intensities were very low in the as-pyrolyzed samples, potentially due to fluorescence caused by C-H bonds.37 Deconvolution of the D and G peaks (ESI Fig. S4†) revealed T and D" bands, likely due to the existence of disordered sp²-sp³ bonds, graphene edges, and pores.^{38,39} The disorder-induced "D" band in graphitic materials corresponds to the T band in free carbon, a low-frequency Raman characteristic. It originates as a result of phonon scattering with disorder- and defect-induced modes. Vacancies, grain boundaries, edge dislocations, and nonplanar graphitic structures are only a few examples of the structural imperfections and disorder in carbon that are linked to the T band. The density and character of these defects can be derived from the T band's intensity and position.40

The D and G peaks intensity ratio is calculated for the sp² carbon clustering, and the tendency of graphite pile up, which plays an important role in the mechanical and electrical properties of the material.⁴¹ To compute the cluster diameter (L_a), the following equation was used:⁴²

$$\frac{I_{\rm D}}{I_{\rm G}} = \frac{C(\lambda)}{L_{\rm a}} \tag{3}$$

where $I_{\rm G}$ is the G peak intensity, $I_{\rm D}$ is the D peak intensity and $C(\lambda)$ is the wavelength-dependent constant. The peak intensity of the D peak proportional to wavelength, hence $C(\lambda)$ was calculated by using the eqn (4) to avoid confusion of the results.⁴³

$$C(\lambda) = C_0 + \lambda_{\rm L} C_1 \tag{4}$$

where C_0 is -12.6 nm, C_1 is 0.033, and λ_L is the wavelength (532 nm).

Hence, the average $L_{\rm a}$ for the annealed Si(Nb)OC (6 \pm 0.2 nm) indicating the graphitic carbon transition to disordered carbon or tetrahedral amorphous carbon.⁴⁴

3.3 Nanostructural features

The schematic representation of fabrication process of Si(Nb) OC composites presented in Fig. 3a. Fig. 3b and c show the TEM microstructures of as-pyrolyzed Si(Nb)OC composites. In Fig. 3d, high resolution images of the as-pyrolyzed composites revealed lattice fringes of inter-planar distance (d) of 0.39 nm, which potentially corresponds to the nanosized orthorhombic

Nb₂O₅.45 Nanda et al. previously observed that the heat treatment of a liquid Nb precursor source at 700 °C initiated crystallization of the orthorhombic structure and finished crystallization at 900 °C under ambient atmosphere.13 Therefore, this process method confined the crystallization of nanosized orthorhombic Nb₂O₅ crystals in the amorphous SiOC matrix, which prevented complete growth of the crystals. The resulting selected area electron diffraction (SAED) pattern (Fig. 3e) showed the amorphous nature of these composites, which agreed with the X-ray diffractogram. Elemental mapping (Fig. 3f-i) revealed uniform distribution of all the elements Si, C, O, and Nb, in as-pyrolyzed Si(Nb)OC composite. Fig. 3j and k show the TEM microstructures of the annealed composite, further high-resolution microstructure (Fig. 31) revealed lattice fringes of crystals SiO₂ (d = 0.40 nm), NbC (0.25 nm), and t- NbO_2 (d = 0.34 nm), which agreed with XRD results of the first most intense peak d spacing values. The respective SAED pattern (Fig. 3m) also showed the diffraction spots, thereby confirming the polycrystalline nature, and the elemental mapping (Fig. 3n-q) of the annealed composites revealed annealing results in the segregation of carbon.

3.4 Bonding characteristics and chemical composition

As-pyrolyzed and annealed Si(Nb)OC composites XPS spectra are shown in Fig. 4. High resolution XPS spectra of deconvoluted peaks of Si 2p, C 1s, O 1s, and Nb 3d were observed after curve fitting. The deconvoluted peaks of Si 2p indicated the presence of SiO₂, 33,46,47 SiO₄ (ref. 33 and 47), and SiCO₃ (ref. 33 and 47) in the as-pyrolyzed composite (Fig. 4b), whereas high intensity SiO₄ bonds and very low intensity SiO₂ bonds occurred in the annealed composite (Fig. 4g). Although the deconvoluted C 1s peak showed C-C, 31,39 C-Si, 31,38 and C-O38 bonds in aspyrolyzed composite (Fig. 4c), however, only C-C bonds were observed in the annealed composite (Fig. 4h). In addition, during the carbothermal reaction process of the annealed sample, the available free carbon in SiOC utilized to produce the t-NbO₂ and NbC, thereby increasing the intensity of SiO₄ bonds and decreasing the intensity of C-C bonds.28 The curve under O 1s in Fig. 4d and i indicate the existence of SiO₂ (ref. 33 and 47), C-O 33,47 , and Nb $_2$ O $_{5-x}^{48}$ bonds. However, formation of NbC in the annealed sample resulted in decreased intensity of Nb_2O_{5-x} compared to the as-pyrolyzed composite. In Fig. 4e and j, the peaks of Nb 3d showed 3d_{3/2} and 3d_{5/2} deconvoluted peaks in both the composites, further the deconvoluted peaks under 3d_{5/} 2 represent different oxidation states of Nb⁵⁺, Nb⁴⁺, and Nb2+.48-50 Increased intensity of the Nb2+ peak in the annealed composite could be due to the NbC phase. Detailed composition is presented in Table 1, and bonding energy (eV) values of the as-pyrolyzed and annealed composites are presented in Table 2.

3.5 Electrochemical performance

Fig. 5a–d represent the electrochemical evaluation of the aspyrolyzed and annealed samples of the Si(Nb)OC composites. The charge–discharge profiles (Fig. 5a) of the as-pyrolyzed sample at a current density of 50 mA $\rm g^{-1}$ showed first-cycle

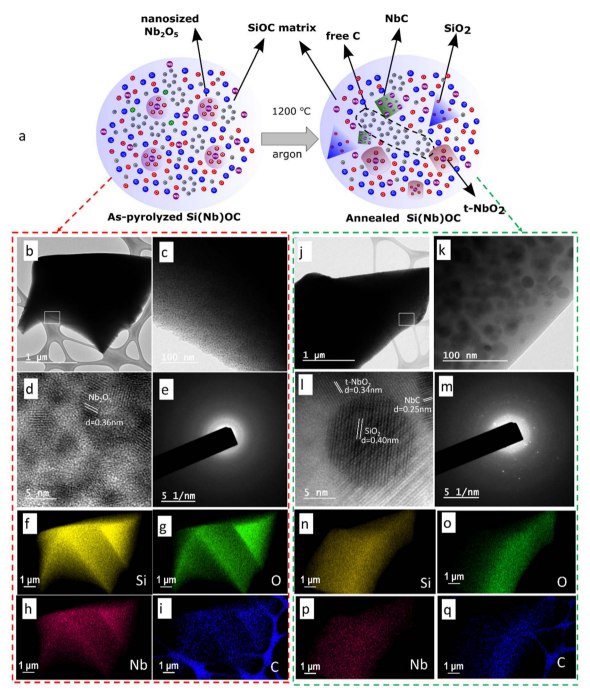


Fig. 3 (a) Schematic representation of fabrication process of Si(Nb)OC composites, (b) and (c) TEM microstructures, (d) high resolution microstructure reveals lattice fringes of ordered nano Nb_2O_5 in amorphous SiOC matrix, (e) SAED diffraction pattern shows amorphous nature of as-pyrolyzed Si(Nb)OC composite, (f–i) elemental mapping of as-pyrolyzed Si(Nb)OC composite shows the presence and uniform distribution of elements Si, O, Nb & C; (j) and (k) TEM microstructures of annealed Si(Nb)OC, (l) high resolution microstructure reveals lattice fringes of t-NbO2, NbC and SiO2 nanocrystals in amorphous SiOC matrix, (m) SAED diffraction pattern shows polycrystalline nature of annealed Si(Nb)OC composite, (n–q) elemental mapping of annealed Si(Nb)OC composite shows the presence and uniform distribution of elements Si, O, Nb & segregated C.

reversible and irreversible capacities of \sim 431 mA h g⁻¹ and \sim 943 mA h g⁻¹, respectively, with 46% CE. The reversible and irreversible capacities of the second cycle were \sim 390 mA h g⁻¹ and 453 mA h g⁻¹, respectively, with 86% CE. Similarly, for the annealed sample in Fig. 5c, the first-cycle reversible and

irreversible capacities were $\sim\!\!256$ mA h g^{-1} and 685 mA h g^{-1} with 37% CE, while the second-cycle reversible and irreversible capacities of the annealed sample were 253 mA h g^{-1} and 330 mA h g^{-1} with 76% CE. Pure Nb₂O₅ charge–discharge curves presented in ESI Fig. S2† show the first-cycle reversible and

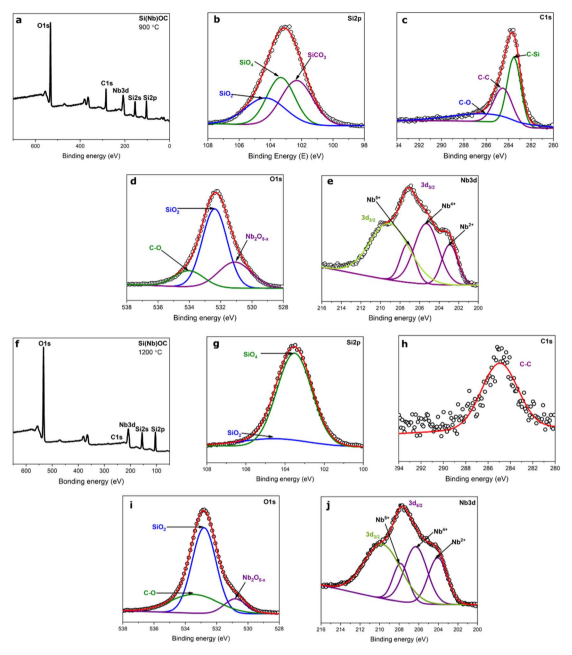


Fig. 4 XPS spectra of as-pyrolyzed Si(Nb)OC composite (a) survey scan, (b-e) high resolution scans of Si 2p, C 1s, O 1s and Nb 3d respectively and annealed Si(Nb)OC composites (f) survey scan, (g-j) high resolution scans of Si 2p, C 1s, O 1s and Nb 3d respectively.

Table 1 XPS elemental composition of Si(Nb)OC composites

Si(Nb)OC composites	Elemental composition (at%)			
	Si	О	C	Nb
As-pyrolyzed	21.0	44.8	28.7	4.5
Annealed	29.4	62.1	3.1	5.4

irreversible capacities of orthorhombic (ESI Fig. S5a†) and monoclinic (ESI Fig. S5c†) Nb₂O₅ to be \sim 134 mA h g⁻¹ with 40% CE and 220 mA h g⁻¹ with 63% CE, respectively. The aspyrolyzed composites performed improved reversible charge

capacity in the first and second cycles among the tested samples, whereas the annealed sample and pure Nb_2O_5 had comparatively low reversible capacity values and efficiency. Due to the presence of more active sites for Li^+ ion intercalation in as-pyrolyzed sample, such as free carbon and mixed Si–O–C units, demonstrated by prevalent C–C bonds and SiCO $_3$ bonds from XPS analysis 51 and uniformly distributed carbon from TEM analysis. The dominated SiO_4 bonds in the annealed sample reacted with the Li^+ ions to form an irreversible Li_4SiO_4 phase, resulting in high values of irreversible capacity. 52 Furthermore, crystallization of insulating phases (SiO_2 and t- NbO_2) in the annealed Si(Nb)OC composite created obstacles

Nb4+(206.3)

Nb2+(204.1)

Deconvoluted peaks (binding energy, eV) Si(Nb)OC composites Nb 3d Si 2p 0.1sC 1s Nb5+(207.6) As-pyrolyzed SiO₂ (104.5) SiO₂ (532.8) C-Si (283.5) Nb4+(206.3) SiO₄ (103.5) C-O (533.4) C-O (286) SiCO₃ (102.5) Nb_2O_{5-x} (530.8) C-C (284.6) Nb2+(204.0) Nb5+(207.8) Annealed SiO₂ (104.5) SiO₂ (532.8) C-C (284.7)

C-O (533.3)

 Nb_2O_{5-x} (530.9)

Table 2 Binding energy of Si(Nb)OC composites

for the Li ion conduction path, and the formation of conductive NbC resulted in decreased carbon content and segregated graphitic carbon that potentially led to the low reversible capacity values. Fig. 5b and d represents Si(Nb)OC composites differential capacity curves with broad peaks around $\sim\!0.26-0.21$ V and $\sim\!0.01-0.16$ V, clear indications of Li insertion into the Si-O-C phase and graphitic carbon. 1,38,53 The first-cycle redox sharp peaks at 1.63 V and 1.71 V (ESI Fig. S5b†) were due to the reduction of Nb $^{5+}$ /Nb $^{4+}$ to Nb $^{4+}$ /Nb $^{3+}$ for orthorhombic Nb $_2O_5$. 54 Although sharp redox peaks were evident at 1.64 V and 1.72 V for the monoclinic Nb $_2O_5$ (ESI Fig. S5d†), 54 no such peaks were observed in the Si(Nb)OC composites. The reduction peak was observed around $\sim\!0.65$ V in all tested samples due to formation of the SEI layer. 55

SiO₄ (103.5)

SiCO₃ (102.5)

The dominated mechanism is Li⁺ ions storing in free carbon in both as-pyrolyzed and annealed Si(Nb)OC composites. However, the insulating phases (SiO₂/NbO₂) in annealed sample likely traps the Li⁺ ions resulting in decreased capacity values. The schematic illustration of Li storing mechanism is depicted in Fig. 6a. Fig. 6b compares the rate capabilities of the aspyrolyzed Si(Nb)OC composites with pure SiOC³³ at various current densities (50 mA g⁻¹,100 mA g⁻¹, 200 mA g⁻¹, 400 mA g⁻¹, 600 mA g⁻¹, 800 mA g⁻¹, and 50 mA g⁻¹). In Fig. 6b present the microstructure of the post-cycling samples of the aspyrolyzed Si(Nb)OC composites exhibited a stable SEI layer. The as-pyrolyzed Si(Nb)OC exhibited improved rate capability with an average reversible capacity of 200 mA h g⁻¹ and 99% CE, after 100 cycles, potentially due to the unique structure of the well-

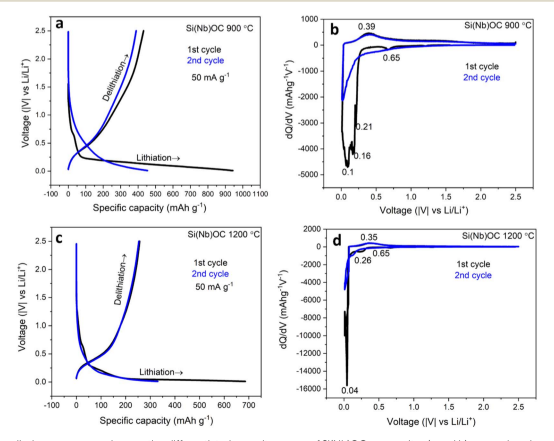


Fig. 5 Charge-discharge curves and respective differentiated capacity curves of Si(Nb)OC composites (a and b) as-pyrolyzed composite and (c and d) annealed composite.

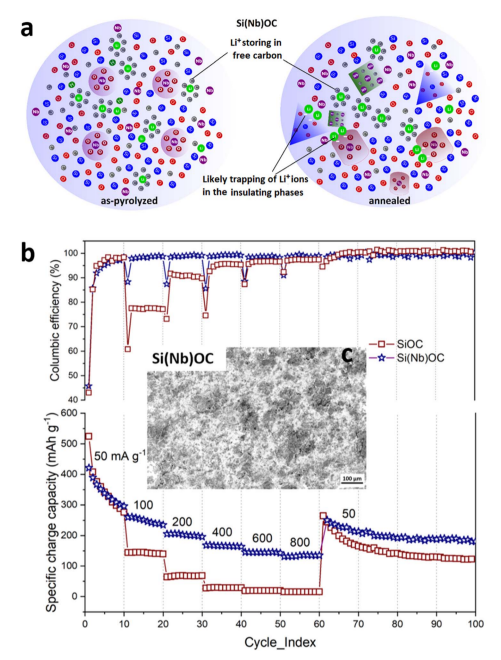


Fig. 6 (a) Schematic illustration of Li ion storing mechanism in as-pyrolyzed and annealed Si(Nb)OC composites, (b) the cyclic stability of pure SiOC (our recent work³³) and as-pyrolyzed Si(Nb)OC composite anodes when cycled symmetrically at increasing current densities and their respective coulombic efficiency and (c) post-cycling SEM microstructure of as-pyrolyzed Si(Nb)OC composite revealing stable SEI layer.

distributed amorphous SiOC/graphitic carbon/ordered $\mathrm{Nb_2O_5}$ of the as-pyrolyzed Si(Nb)OC composites, which maintained the structure stability during cycling, and stable SEI formation, resulted in high efficiency. Similar results were reported in our recent work on single-source precursor-derived $\mathrm{SiOC/TiO_xC_y}$ anodes for LIBs.³³ Results of the current study showed that $\mathrm{Si(Nb)OC}$ composite materials produced through precursor route and heat-treated at pyrolysis temperature improved rate capability compared to pure SiOC and precursor derived $\mathrm{Nb_2O_5}$ materials (supplementary Fig. S6†). Therefore, this work explored the possibilities for the future anode material for LIBs especially for "high powered devices".

Conclusions

This study utilized a polymer precursor process to synthesize Si(Nb)OC composites as hybrid anode materials for LIBs. Electron microscopy and XRD techniques were used to analyze the micro/nanostructural features and phase evolution, and X-ray photoelectron and Raman spectroscopy were used to investigate the bonding nature, elemental composition, and the disordered and graphitic carbon in the Si(Nb)OC composites respectively. As-pyrolyzed and annealed composites demonstrated first-cycle reversible capacities of 431 mA h g⁻¹ and 256 mA h g⁻¹, respectively, which exceeded the reversible

capacity of pure $m\text{-Nb}_2O_5$ (220 mA h g⁻¹). The as-pyrolyzed Si(Nb)OC composite displayed improved rate capability than pure SiOC, with an average reversible capacity of 200 mA h g⁻¹ and ~99% CE. The amorphous SiOC structure and uniform distribution of nano-sized Nb₂O₅ crystals and graphitic carbon allowed the composite to form stable SEI layers and remain stable during cycling.

Conflicts of interest

There are no conflicts to declare.

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