Nickel@Siloxene catalytic nanosheets for high-performance CO\textsubscript{2} methanation

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Two-dimensional (2D) materials are of considerable interest for catalyzing the heterogeneous conversion of CO\textsubscript{2} to synthetic fuels. In this regard, 2D siloxene nanosheets, have escaped thorough exploration, despite being composed of earth-abundant elements. Herein we demonstrate the remarkable catalytic activity, selectivity, and stability of a nickel@siloxene nanocomposite; it is found that this promising catalytic performance is highly sensitive to the location of the nickel component, being on either the interior or the exterior of adjacent siloxene nanosheets. Control over the location of nickel is achieved by employing the terminal groups of siloxene and varying the solvent used during its nucleation and growth, which ultimately determines the distinct reaction intermediates and pathways for the catalytic CO\textsubscript{2} methanation. Significantly, a CO\textsubscript{2} methanation rate of 100 mmol g\textsubscript{Ni}\textsuperscript{-1} h\textsuperscript{-1} is achieved with over 90% selectivity when nickel resides specifically between the sheets of siloxene.
Carbon dioxide (CO$_2$) is a well-known greenhouse gas that can also be used as a zero- or even negative-cost carbon feedstock in the production of valuable chemicals and fuels$^{1,2}$. This technology can be employed in the power-to-gas process, which consists of an initial hydrogen production step via electrolysis of water, followed by the reduction of CO$_2$ via the Sabatier reaction (i.e., methanation). This process of converting CO$_2$ into methane (CH$_4$) is among the most attractive and promising technologies for transforming electrical energy into renewable chemical energy$^{3-5}$. In doing so, it is possible to circumvent the considerable difficulties associated with hydrogen storage, transport, and large-scale utilization. In addition, CH$_4$ offers several benefits over hydrogen, as it is easily liquefied, can be more safely stored and transported, and is widely used in energy-intensive applications as a major component of natural gas.

For the catalysis of the CO$_2$ methanation reaction, earth-abundant and relatively low-cost Ni metal is an ideal candidate, which has led to its use being intensively explored. As the methanation reaction is highly exothermic, great effort has been devoted to anchoring or immobilizing Ni nanoparticles onto support materials in attempts to alleviate high-temperature sintering$^{6-9}$. Because of such efforts, spatial confinement has emerged as an effective and unique way to limit the growth of Ni nanoparticles$^{10,11}$. Chen et al.$^{11}$ pointed out that a “cage” of mesoporous SBA-16 could confine small Ni nanoparticles, thereby contributing to their high activity in the reverse water–gas shift reaction (i.e., hydrogenation of CO$_2$ to CO). However, the design and fabrication of Ni catalysts with high selectivity toward CH$_4$ production, rather than competing reverse water–gas shift reaction, remains a major challenge. In addition, the reaction mechanism for Ni catalysts is still largely ambiguous, both under laboratory conditions and in industrial applications.

Only recently have theoretical and modeling studies begun to show some evidence regarding the CO$_2$ methanation mechanism$^{12-14}$. Heine et al.$^{14}$ used ambient-pressure X-ray photoelectron spectroscopy (XPS) to probe the reaction pathways for CO$_2$ methanation on a Ni (111) surface. In this system, CO$_2$ first dissociated to CO and atomic oxygen at the Ni surface, whereupon CO then decomposed to atomic carbon (C) before finally being converted to CH$_4$ via hydrogenation. This reaction pathway inevitably yields significant quantities of CO and may lead to catalyst poisoning via carbonization (coking) of the catalyst surface.

In this context, a superior Ni catalyst must have high activity and selectivity toward CH$_4$ production, as well as good stability under operating conditions. A unique pathway for sustainable CH$_4$ production must therefore be achieved by designing the chemistry and structure of not only Ni itself but, more importantly, the supporting materials used to accomplish the desired surface engineering and spatial confinement.

Silicon, being the second most earth-abundant element, is clearly deserving of investigation for such use. Aside from mesoporous silica, which could supply the aforementioned two-dimensional (2D) or three-dimensional (3D) confinement effect$^{15-17}$, and some reports on metal on porous Si$^{18-21}$, the most widely used form of Si for supporting catalysts is SiO$_2$ particles, without further chemical or structural design. SiO$_2$-supported Ni catalysts have relatively low selectivity toward CH$_4$ production (lower than 80%) with the remaining ca. 20% selectivity being toward CO$^{22}$. In this regard, various promoters were added to improve the CH$_4$ production selectivity of the Ni/SiO$_2$ catalysts$^{23-25}$. In addition, production selectivity could be switched from CO to CH$_4$ on Ni/SiO$_2$ catalysts by increasing the Ni loading from 0.5 wt% to a much higher value of 10 wt%$^{26}$. However, it is still a challenge to design silicon-supported Ni catalysts with a distinctive structure and superior selectivity for CO$_2$ methanation. In an era where 2D catalyst materials are both in great demand and showing significant promise scientifically and technologically, 2D silicon nanosheet materials have historically been overlooked. Only a few reports for its use in catalysis can be found, exemplifying its use in water splitting$^{27}$, ethylene polymerization$^{28}$, formic acid decomposition$^{29}$, 4-nitrophenol reduction$^{30}$, and butene isomerization$^{31}$. In this report, we employ 2D silicon surface chemistry to design catalysts consisting of Ni on siloxene nanosheets (SIXNS), in order to address the challenges posed by CO$_2$ methanation. Hydride- and hydroxyl-terminated layered silicon was chosen as the starting material for designing and constructing the uniquely supported Ni nanostructures used in this study. It is shown herein that Si–H hydrolysis can be triggered or avoided during synthesis by using a water- or ethanol-based Ni(NO$_3$)$_2$ precursor solution, respectively. This strategic choice also determines whether Ni nanoparticles are obtained on the interior or exterior of the SIXNS. The Ni nanoparticles confined inside the SIXNS exhibit improved catalytic activity and selectivity to CH$_4$ relative to those on the surface of SIXNS, and this catalytic performance can be further improved under photothermal reaction conditions. With evidence from in situ infrared studies, two distinct pathways for CO$_2$ methanation are observed for these two classes of materials, providing the insight and designing strategy for composite catalysts and materials with structural confinement effects.

**Results**

**Materials design and fabrication process.** SIXNS were exfoliated from bulk calcium silicide at 0 °C using concentrated HCl$^{27,32,33}$. The composition and structure of such exfoliated 2D nanosheets have been studied for decades. Three main types of structures containing Si–H and different types of O sites have been proposed: first, the Weiss siloxene consisting of corrugated Si (111) layers terminated above and below by OH and H groups$^{34}$; second, the Kautsky siloxene consisting of puckered Si six rings interconnected by oxygen bridges$^{35}$; and third, an unusual structure wherein linear Si chains are interconnected via oxygen bridges and terminated by hydrogen$^{27}$. When actually synthesized in the lab, a mixture of the above mentioned structures with the general formula of Si$_n$(OH)$_{2n-2}$ is observed, in which Si is terminated by –H and –OH functional groups, with Si–O–Si bridges also existing$^{36,37}$. Such configurations allow for dispersion in common polar solvents, such as water and ethanol, facilitating the next step of nanocomposite formation using the soluble Ni salt. A dispersion of the fresh SIXNS was mixed and sonicated with a nickel nitrate solution, dried by heating under vacuum, and then calcined under air at 400 °C for 3 h. This procedure resembles the widely used impregnation method for producing conventional supported metal catalysts; however, it was found that the choice of solvent (i.e., water or ethanol), resulted in different structures in the product material. Figure 1a illustrates the philosophy of the design strategy: the circle of Taiji represents the unity of beginning from the same precursor, whereas the separated black and white halves symbolize the difference of the final products. When using water as the solvent, both the hydrolysis of the Si–H perpendicular to the plane to form Si–OH and the condensation of Si–OH into Si–O–Si will proceed during the impregnation process, as was also observed by Weiss et al.$^{34}$ (the pioneer of the siloxene material) and, more recently, by Xie et al.$^{27}$. This hydrolysis by water appears to inhibit the infiltration and deposition of nickel nitrate between the siloxene sheets. In contrast, ethanol preserves more Si–H and results in less condensation of Si–OH. These mechanisms are evidenced by
Fourier-transform infrared (FTIR) spectroscopy (Fig. 1b) of samples prepared using either water or ethanol. The spectrum of the SiXNS impregnated in ethanol resembles that of the original pristine SiXNS, with the extra peaks at around 1350 cm$^{-1}$ assigned to nitrate ions. In contrast, the sample impregnated in water exhibits significantly increased Si–O–Si peaks due to condensation of Si–OH, a weak mode at around 2100 cm$^{-1}$ for Si–H, and distinctive Si–OH modes between 3250 and 3500 cm$^{-1}$, due to more extensive hydrolysis of Si–H. As a result, following the decomposition of nickel nitrate via calcination, the NiO so produced tended to be located on the more extensively hydrolyzed exterior surface of the stacked SiXNS impregnated in water (NiO@SiXNS-H$_2$O) but between the sheets of less-hydrolyzed SiXNS impregnated in ethanol (NiO@SiXNS-EtOH). Due to this unique siloxene chemistry, significantly different final structures were obtained from the same precursor material. Notably, the Ni loadings were almost the same for NiO@SiXNS-H$_2$O and NiO@SiXNS-EtOH, (4.9 and 4.7 wt%, respectively), as determined using inductively coupled-plasma atomic emission spectroscopy (ICP-AES).

The morphology and microstructure of fresh NiO@SiXNS was examined using electron microscopy; representative images are shown in Fig. 1c, d. Scanning electron microscopy (SEM) and transmission electron microscopy (TEM) images of NiO@SiXNS-H$_2$O and NiO@SiXNS-EtOH showed that NiO nanoparticles were deposited on the external surface of SiXNS. This can be further illustrated by the patterns of green (Ni) octahedra scattered across the red (Si) support, whereas the blue (O) component is more homogeneously distributed throughout the material, as obtained from energy-dispersive X-ray (EDX) mapping analysis (Fig. 1c). The lattice fringe of 0.21 nm on NiO@SiXNS-H$_2$O originates from the NiO (200) plane, further confirming the identity of the phase. In contrast, NiO nanoparticles formed on NiO@SiXNS-EtOH grew between the sheets of SiXNS. In the SEM images shown in Fig. 1d, very few particles could be observed on the exterior surface. Only by increasing the accelerating voltage could more and brighter particles be revealed, exhibiting distinctive contrast against the surrounding SiXNS as the penetration ability of the accelerated electrons increased. Providing crucial support for this observation is the fact that TEM imaging clearly revealed the majority of NiO particles were encapsulated within the SiXNS and could not be observed using low-voltage SEM.

**Characterization of NiO@SiXNS materials.** Powder X-ray diffraction (PXRD) (Fig. 2a) studies of the NiO@SiXNS-H$_2$O and
NiO@SiXNS-EtOH were carried out in order to better understand their structure. In addition to the peaks associated with SiXNS exfoliated from CaSi$_2$ (marked with asterisks)\textsuperscript{37}, both samples exhibited one broad peak at $\sim$ 20° (corresponding to the presence of an amorphous phase) and four obvious peaks (marked with rhombuses) corresponding to NiO. The slightly broader peaks of NiO in NiO@SiXNS-EtOH also implies that spatial confinement limits the nucleation and growth process. The Brunauer–Emmett–Teller (BET) surface areas of NiO@SiXNS–EtOH and NiO@SiXNS–H$_2$O are 8.8 and 9.1 m$^2$ g$^{-1}$, respectively (Supplementary Fig. 1).

The reduction properties of NiO@SiXNS was measured by using hydrogen temperature programmed reduction (Fig. 2b). Slight hydrogen consumption begins at 250 °C for both samples. A major peak was observed at 450 °C for NiO@SiXNS–H$_2$O, whereas the corresponding peak was not observed until 476 °C for NiO@SiXNS–EtOH, which also exhibited a shoulder peak at $\sim$ 700 °C. The major peak in these TPR profiles originates from the reduction of NiO. The higher reduction temperature of NiO@SiXNS–EtOH relative to NiO@SiXNS–H$_2$O is attributed to the fact that, in the former case, NiO nanoparticles are trapped inside the SiXNS and well-anchored to the support, whereas, in the latter, NiO nanoparticles are mainly located on the exterior where they are more exposed to the hydrogen reductant. The small peak at $\sim$ 700 °C on the TPR profiles of NiO@SiXNS is attributed to the presence of silicate species\textsuperscript{38}.

XPS spectroscopy (Fig. 2c, d) revealed more detailed information about NiO@SiXNS. The Si 2$p$ spectra showed key peaks at the binding energies of 98.5 and 103 eV, corresponding to siloxene Si$^0$ and Si$^{4+}$ sites, respectively. The predominance of Si$^{4+}$ compared with Si$^0$ can be attributed to oxidation during the calcination process. In addition, spectra from both samples exhibited broad, asymmetric peaks in the Ni 2$p_{3/2}$ region. This region consisted of two peaks at binding energies of 854.5 and 856.3 eV, which represent NiO and a nickel silicate (NiSiO$_3$) phase, respectively\textsuperscript{39}. According to the deconvoluted XPS results, the NiSiO$_3$-to-NiO peak area ratio of NiO@SiXNS–EtOH (3.6) was four times of that for NiO@SiXNS–H$_2$O (0.9). This difference further supports the preferred location of NiO at interior sites in NiO@SiXNS–EtOH, as more NiO should be sandwiched between SiXNS.

**Characterization of Ni@SiXNS materials.** In order to determine the structure of the catalysts, the reduced Ni@SiXNS were further characterized. Accurate 3D visualization of the confined architecture of Ni@SiXNS–EtOH was achieved via electron tomography analysis (Fig. 3 and Supplementary Movie 1). The majority of the nanoparticles are seemingly encapsulated within the interlayer spaces of adjacent nanosheets (Fig. 3a and Supplementary Movie 1). Almost no nanoparticles were observed on the top and bottom of the nanosheets (Fig. 3b, e); in contrast, most of the nanoparticles with bright contrast were revealed in slices deeper within the 3D reconstruction of this composite particulate (Fig. 3c, d).

The PXRD patterns of Ni@SiXNS catalysts revealed that both Ni and NiO were present following reduction (Fig. 4a, b). TEM images of the Ni@SiXNS–H$_2$O and Ni@SiXNS–EtOH, shown in Fig. 4c and Supplementary Fig. 2, further revealed that Ni nanoparticles were exposed on the exterior of the SiXNS, and that well-defined lattice fringes of 0.176 and 0.21 nm associated with Ni (200) and NiO (200) planes are observed on Ni@SiXNS–H$_2$O. On Ni@SiXNS–EtOH, the majority of Ni nanoparticles are sandwiched between SiXNS sheets and only a few Ni nanoparticles are immobilized on the outer surface as seen in the thinner regions of SiXNS (Supplementary Fig. 3). The lattice fringes of 0.176 nm and 0.210 nm can also be ascribed to Ni (200) and NiO.

**Fig. 2** Characterization of NiO@SiXNS materials. a, PXRD patterns showing characteristic peaks from the (*) SiXNS and (♦) NiO constituents of the composite materials. b, H$_2$–TPR (hydrogen temperature programmed reduction) plots indicating the appreciable difference in the onset of NiO reduction on heating under a hydrogen atmosphere. c, d XPS spectra illustrating the differences in the local bonding environments of c, Si 2$p$ and d, Ni 2$p_{3/2}$ obtained from of NiO@SiXNS samples (green peak refers to NiSiO$_3$, cyan peak corresponds to NiO, and yellow one is the satellite peak).
(200) planes, respectively (Fig. 4d). The Ni surface area and size of Ni@SiXNS were determined using H₂ chemisorption measurements. Similar Ni surface areas of 3.8 and 4.0 m² g⁻¹ were observed for the Ni@SiXNS-H₂O and Ni@SiXNS-EtOH, respectively. The sizes of Ni particles were also similar, being 10.7 nm for Ni@SiXNS-H₂O and 10.3 nm for Ni@SiXNS-EtOH. These resemblances suggest that the surface area and size of Ni in these catalysts are not responsible for the observed differences in their catalytic performance.

**Catalytic performance of CO₂ methanation.** The catalytic activity of the Ni@SiXNS catalysts for the CO₂ methanation was tested at 300 °C in a flow reactor system operating at atmospheric pressure under a flowing H₂/CO₂ atmosphere (4:1). The pristine SiXNS (calcined at 400 °C for 3 h) showed no activity toward CO₂ methanation either in the dark or light under the same reaction conditions. For the composite catalyst materials, however, significant activity was observed, with CH₄ and CO being identified as the only products generated by these Ni@SiXNS catalysts under both dark and light conditions (Fig. 5). The CH₄ and CO production rates for Ni@SiXNS–H₂O in the dark were found to be 36.6 and 32.0 mmol g⁻¹ h⁻¹, respectively, which increased to 41.0 and 34.5 mmol g⁻¹ h⁻¹ on exposure to light (Fig. 5a, b). Despite this rate increase, CH₄ selectivity remained virtually
unchanged, with calculated values of 53.4% and 54.3% upon switching from dark to light operating conditions (Fig. 5c). This slightly enhanced performance is attributed to the photothermal effect of the Ni particles in Ni@SiXNS-H2O. The diffuse reflectance spectrum of the sample is low throughout the visible spectral region because of the significant visible light-harvesting capacity (interband and intraband electronic transitions) of the black Ni nanoparticles, which should favor local heating of Ni via the photothermal effect (Supplementary Fig. 4).

As a control experiment, commercial amorphous silicon dioxide-supported nickel (Ni@SiO2) was employed and loaded into the same flow catalyst reactor for evaluation under the same conditions. The Ni loading and mass of the Ni@SiO2 used were the same as those of Ni@SiXNS. As shown in Supplementary Fig. 5, the CH4 and CO production rates in the dark over Ni@SiO2 are 22.78 and 193.70 mmol gNi\(^{-1}\) h\(^{-1}\), respectively. In this case, CH4 selectivity is only 10.5%, significantly lower than that of Ni@SiXNS-H2O. Under light conditions, CH4 and CO production rates increase to 34.55 and 296.09 mmol gNi\(^{-1}\) h\(^{-1}\), whereas CH4 selectivity still remains at 10.5%. The result indicates that Ni supported on SiO2 favors the reverse water–gas shift reaction, contributing to the formation of large amount of CO rather than CH4. We attribute the lower selectivity toward CH4 to the poorer contact between Ni and commercial spherically shaped SiO2 particles compared with that between the Ni and the planar SiXNS.

In an attempt to improve the selectivity toward the production of CH4, Ni@SiXNS-EtOH with Ni nanoparticles trapped between the sheets was also tested for its CO2 methanation activity, as shown in Fig. 5. Notably, CH4 and CO production rates for this catalyst increased from 45.2 and 5.1 mmol gNi\(^{-1}\) h\(^{-1}\), respectively, in the dark to 52.9 and 6.7 mmol gNi\(^{-1}\) h\(^{-1}\) in the light (Fig. 5a, b). The CH4 selectivity increased slightly from 89.8% in the dark to 91.9% in the light (Fig. 5c). Isotopic labeling experiments using \(^{13}\)C and gas chromatography-mass spectrometry (GC-MS) were used to study the products generated from a \(^{13}\)CO2 reactant feed. The mass-to-charge (m/z) ratios observed at 17 and 29 amu in the GC-MS spectrum corresponded to \(^{13}\)CH4 and \(^{13}\)CO (Supplementary Fig. 6), confirming that CH4 and CO are, indeed, produced from the methanation of CO2 on Ni@SiXNS and not from adventitious carbon contaminates. Thus, Ni@SiXNS-EtOH exhibited both higher selectivity and activity for the CO2 methanation than Ni@SiXNS-H2O.

The catalytic stability at 300 °C with light of Ni@SiXNS-EtOH was next investigated, as illustrated in Fig. 5d. The CH4 production rate increased progressively and reached a stable value around an impressive 100 mmol gNi\(^{-1}\) h\(^{-1}\) during the 12 h test, whereas the CO production rate increased only slightly and stabilized around 10 mmol gNi\(^{-1}\) h\(^{-1}\). In addition, CH4 selectivity was observed to remain stable at above ~90%. One can therefore deduce that Ni@SiXNS-EtOH is a highly active, selective, and stable catalyst for the preferential hydrogenation of CO2 to CH4.

The catalytic performance of an industrial Ni@Al2O3 catalyst is also tested, as illustrated in Supplementary Fig. 7. CH4 and CO production rates in the light over Ni@Al2O3 are 28.69 and 0.63 mmol gNi\(^{-1}\) h\(^{-1}\) with a CH4 selectivity of 97.8% in the first 30 min and slightly decreased and maintained at 27.78 and 0.55 mmol gNi\(^{-1}\) h\(^{-1}\) with similar selectivity. Although Ni@Al2O3 has higher CH4 selectivity than that of Ni@SiXNS-EtOH, the production rates are much lower.

**Fig. 5** Catalytic performance of CO2 methanation on Ni@SiXNS. a, b Observed catalytic rates for a CH4 production and b CO production by Ni@SiXNS samples under light and dark conditions. c CH4 selectivity of Ni@SiXNS samples for CO2 methanation under light and dark conditions. d CH4 production rate and selectivity as well as CO production rate of Ni@SiXNS-EtOH for CO2 methanation under illumination for a continuous 12 h at atmospheric pressure.
Catalytic reaction pathways. To gain deeper insights into the CO₂ methanation reaction over Ni@SiXNS, in-situ diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) studies were conducted. Each NiO@SiXNS sample was reduced in the DRIFTS sample cell for 1 h under a H₂ atmosphere and then purged for 1 h under He at 300 °C. Next, the sample was exposed to a mixture of CO₂ (2 s.c.c.m.), H₂ (8 s.c.c.m.), and He (10 s.c.c.m.) for 1 h. We focused our analysis on the IR fingerprint regions of 1200–2200 cm⁻¹ and 2600–3400 cm⁻¹ in the consecutive spectra, where reaction intermediates could be detected.

For Ni@SiXNS-H₂O, two key reaction intermediates were observed, as shown in Fig. 6a. The first, identified as chemisorbed CO₂, had observable peaks at 1430 and 1500/1767 cm⁻¹, and is diagnostic of the formation of bicarbonate and carbonate species, respectively⁴¹-⁴⁴. The second was identified from peaks located at 1930 and 2077 cm⁻¹, which are attributed to the presence of bridging and linearly adsorbed CO molecules on Ni, respectively⁴⁵,⁴⁶. As depicted in Fig. 6b, C–H stretching vibrations of CH₄ were also detected at 3016 cm⁻¹, with additional peaks at 2111/2174 and 1630 cm⁻¹ being observed for gaseous CO and H₂O, respectively⁴⁷.

For Ni@SiXNS-EtOH, apart from the carbonate species, one important reaction intermediate appeared at 1404 cm⁻¹ in the spectra (Fig. 6c) and was ascribed to a surface formate species (HCOO⁻)⁴⁸; this assignment is supported by an additional peak observed at 2814 cm⁻¹ (Fig. 6d). It is noteworthy that the peak intensity of this formate species is much higher than that of the carbonate species. Yet, another species with corresponding peaks at 3046 and 3140 cm⁻¹ matched well with those of hydroxyl groups, as reported in the literature⁴⁹. It should be noted that no obvious peaks could be detected in the range of 2000–1800 cm⁻¹ for Ni@SiXNS-EtOH, which reveals the essential absence of bridging CO; this is likely because Ni nanoparticles in the catalyst are trapped between sheets of SiXNS, thus decreasing the chance for the bonding of CO molecules to multiple Ni atoms. Only gaseous CO appears in the spectra, as well as gaseous CH₄ and H₂O.

Based on the differential DRIFTS spectra of the Ni@SiXNS catalysts, a reaction pathway for CO₂ methanation was proposed (Fig. 6e, f). As the Ni nanoparticles were located on the exterior surface of SiXNS in Ni@SiXNS-H₂O, these particles are readily exposed to CO₂, whereupon the chemisorbed *CO₂ species can dissociate into *CO and *O. The strongly bound, bridge-bonded *CO species indicated by the IR data can further dissociate to carbon species (*C), which can then be hydrogenated to form CH₄, and desorb from the catalyst surface, corresponding to the observed CH₄ production, whereas the *O species can react with hydrogen to produce H₂O. However, the stable bridging *CO species also have a higher chance to convert to CO as the final product, leading to an overall lower selectivity to CH₄.

In contrast, Ni nanoparticles in Ni@SiXNS-EtOH are confined within SiXNS and the bridging *CO species cannot be readily formed. A recent theoretical calculation has shown that a silica-cluster-modified Ni (111) surface has a higher activation energy barrier for the dissociation of *CO₂ to CO through the *CO pathway than a clean exposed Ni (111) surface⁵¹. Our DRIFTS spectra and catalytic performance tests on Ni@SiXNS-EtOH are consistent with the prediction that the generation of *CO, and thus CO, is limited due to the Ni nanoparticles being encased within the SiXNS. Therefore, the chemisorbed *CO₂ species for...
this structure could instead convert first to HCO$_2^-$ and then to formic acid (HCOOH). More pronounced C–H IR mode intensity from HCOOH could be detected by quickly cooling down the reactor during DRIFTS measurements (Supplementary Fig. 8). A small portion of the HCOOH so generated could decompose to gaseous CO and H$_2$O; however, a larger proportion is likely to participate in the rapid generation of CH$_4$, possibly by the formate pathway reported by Kattel et al. This additional pathway for CH$_4$ production on Ni@SiXNS-EtOH could be due to the tortuousity experienced by CO$_2$ within the interior of the catalyst, which enhances the probability of it being fully hydrogenated to CH$_4$ via the associative CO$_2$ methanation mechanism (in which carbonate is hydrogenated to bicarbonate and then quickly dehydrated to formate)  . Besides, the newly generated OH groups observed in Fig. 6d could provide additional H source and aid in formate formation.

Discussion

The synthesis of Ni@SiXNS nanocomposites relies on a distinctive kind of silicon surface chemistry in which the Si–H hydrolysis and Si–OH condensation are selectively triggered or avoided. The resulting Ni@SiXNS materials are extremely active toward the conversion of CO$_2$ to CH$_4$ with highselectivity and impressive stability. The nucleation and growth of Ni nanoparticles on either the exterior or interior of the SiXNS was solvent-sensitive, enabled by choosing water or ethanol, respectively. After a standard impregnation–calcination–reduction in water, a Ni@SiXNS-H$_2$O catalyst was produced. For this material, Ni nanoparticles were immobilized on the exterior surface of the SiXNS. The corresponding CH$_4$ production rate was 41.0 mmol g$_{\text{Ni}}$ h$^{-1}$ with heating at 300 °C with assistance of light, but CH$_4$ selectivity was only 54.3%. In contrast, the Ni@SiXNS-EtOH catalyst favored Ni located within the interlayer space of the SiXNS. These samples featured a CH$_4$ production rate of 52.9 mmol g$_{\text{Ni}}$ h$^{-1}$ and a CH$_4$ selectivity over 90%. This spatially confined Ni catalyst proved stable in the light and, over time, eventually reaching an impressive CH$_4$ production rate of ~100 mmol g$_{\text{Ni}}$ h$^{-1}$ with ~90% selectivity.

An in-situ infrared study clarified the reaction pathways of the Ni@SiXNS catalysts for CO$_2$ methanation, revealing the origin of the observed differences in selectivity between the two types of structures. Distinct reaction intermediates (e.g., CO and formate species) were observed on the Ni@SiXNS-H$_2$O and Ni@SiXNS-EtOH samples, respectively. For Ni@SiXNS-H$_2$O, the exposed Ni on the surface of SiXNS could easily bridge-bond to CO, leading to the hydrogenation of *CO* to CH$_4$ following CO dissociation. For Ni@SiXNS-EtOH, the confined Ni sites are unable to form bridging *CO* species and formate species were instead the main intermediates leading to CH$_4$ production.

The gradual increase of the CH$_4$ production rate on Ni@SiXNS-EtOH in Fig. 5d originated from the gradual release of Ni during the CO$_2$ methanation reaction, corresponding well with the recent popular catalytic concept of the “sustained release catalyst,” designed to maintain the high reactivity and stability of the obtained catalyst . A thorough in-situ Extended X-Ray Absorption Fine Structure (EXAFS) study illustrated the gradual evolution of Ni in Ni@SiXNS-EtOH under simulated reaction conditions (Supplementary Fig. 9). Due to the fact that the reaction under light conditions probably resulted in a higher reactor temperature than 300 °C, and that the high flow rate would bring down the surface temperature, we performed the in-situ EXAFS study at 330 °C at a flow of 4:1 H$_2$/CO$_2$ (i.e., 130 s.c.c.m. 3.5% H$_2$ in He and 60 s.c.c.m. 2% CO$_2$ in He) after H$_2$ activation at 300 °C. Significant changes in the XAS spectra began to occur and continued over a 90 min period. This matched well with the rapid attainment of a steady CH$_4$ or CO formation rate in a stability test of Ni@SiXNS-EtOH under dark conditions at 300 °C with pre-reduction at 350 °C (Supplementary Fig. 10).

On the origin of high selectivity for CH$_4$ production of Ni@SiXNS-EtOH, apparent reaction orders were estimated from the variation in production rates with different inlet reactant concentrations according to the rate law equation r = kP$_{\text{H}_2}^aP_{\text{CO}_2}^b$ (Supplementary Fig. 11, Supplementary Table 1, and Supplementary Notes) after the sample was stabilized, as described above. The apparent reaction orders of CO$_2$ and H$_2$ were 0.40 and 1.56, respectively, for CH$_4$ production, whereas the reaction orders of CO$_2$ and H$_2$ were 2.80 and 0.28 for CO production. Thus, the relatively large reaction order value of H$_2$ implies dominant control of the CH$_4$ production rate by the partial pressure of H$_2$, which is indicative of the associative methanation pathway in which H$_2$ fully hydrogenates the activated CO$_2$. In contrast, for CO production, the much larger reaction order of CO$_2$ indicates its greater dependence on CO$_2$ adsorption and dissociation.

Over the range of H$_2$ partial pressure variations explored, it was determined that the fractional yield of CH$_4$ from CO$_2$ was consistently larger than that for CO production from CO$_2$, indicating high selectivity toward CH$_4$ production. Further analysis revealed the rate-limiting step to be the decomposition and further hydrogenation of formate-type species HCOO*- (see Supplementary Information).

The crucial finding of this study is that control over the location of the Ni nanoparticles, either at the interior or exterior regions of the SiXNS, matters when it comes to selectivity favoring the methanation vs. the reverse water–gas shift reaction. The CO pathway dominates for Ni at the exterior of the nanosheets, whereas the formate pathway is preferred for Ni confined to the interior. Although the former does not discriminate much between CO and CH$_4$ products, the latter strongly favors CH$_4$. This innovative approach to the construction of designed architecture 2D Ni@SiXNS nanocomposites presents a paradigm in the field of gas-phase heterogeneous (photo)catalytic hydrogenation of CO$_2$ to synthetic chemicals and fuels.

Methods

Synthesis of SiXNS and NiO@SiXNS. In a typical synthesis of SiXNS, 1.0 g of calcium silicide powder (Fluka™) was dispersed in 100 mL of hydrochloric acid (36%, Caledon Laboratory Chemicals) pre-cooled to 0 °C in a jacketed flask with a circulator (NESLAB® RTE-111). The mixture was stirred under inert atmosphere for 6 days at 0 °C. The solid product was collected via vacuum filtration, washed with anhydrous acetone, and dried under N$_2$ flow before being stored under vacuum in a desiccator. For the synthesis of NiO@SiXNS-EtOH, 0.0124 g of nickel nitrate hexahydrate (Alfa Aesar) was dissolved into 1 mL of ethanol. Next, 0.048 g of SiXNS was dispersed into 2 mL of ethanol and sonicated for 30 min. Then the nickel nitrate ethanol solution was added into the SiXNS dispersion and sonicated for another 30 min. Next, it was placed at room temperature for 12 h and dried in vacuum at 70 °C for 24 h before being cooled down to room temperature under vacuum for 12 h. The powder sample was finally calcined at 400 °C in air for 3 h. The process for the synthesis of NiO@SiXNS-H$_2$O was similar to that of NiO@SiXNS-EtOH, except that the solution was changed to an equal volume of distilled water.

Synthesis of control samples. For NiO@SiO$_2$, 0.0124 g of nickel nitrate hexahydrate (Alfa Aesar) was dissolved into 1 mL of distilled water. SiO$_2$ (0.048 g; Aldrich, fumed, powdered, 0.007 μm) was dispersed into 2 mL of distilled water and sonicated for 30 min. The nickel nitrate solution was then added into the SiO$_2$ dispersion and sonicated for another 30 min. After that, it was held at room temperature for 12 h and dried in vacuum at 70 °C for 24 h. The powder sample was finally calcined at 400 °C in air for 3 h. The industrial Ni@Al$_2$O$_3$ catalyst was purchased from Rögen (10 wt% Ni) and used as received.
Material characterizations. An ICP-AES (Thermo ICAP 6300 spectrometer) was used to measure the nickel content in the NiO@SiXNS samples. PXRD was performed to determine the crystal structure of the xerogel using Cu Kα radiation at 30.0 kV and 15 mA. The samples were scanned from 10° to 80° 2θ at a rate of 1°/min, with a step size of 0.02°. The lattice constant of the Ni(OH)2 phase was calculated by the Bragg’s law.

To investigate the surface area and porosity of the NiO@SiXNS samples, nitrogen adsorption-desorption measurements were performed using a Quantachrome Autosorb-1C instrument. The Brunauer-Emmett-Teller (BET) surface area was calculated from the nitrogen adsorption isotherms at 77 K, and the total pore volume was obtained from the nitrogen adsorption at a relative pressure of 0.99.

The morphology of the NiO@SiXNS samples was characterized using SEM and TEM. For SEM imaging, the samples were coated with a thin layer of gold/palladium to prevent charging. TEM images were obtained using a JEOL JEM-2010 transmission electron microscope operated at 200 kV. The samples were dispersed on carbon-coated copper grids and negatively stained with 2% uranyl acetate.

EDX mapping was performed using an Oxford Inca EDX system. High-resolution TEM images of the samples were obtained on a FEI Tecnai G2 F20 system operated at 200 kV. The elemental distributions were determined by mapping the Ni, O, and Si signals.

The thickness of the NiO layer was measured using a JEOL JEM-2010 transmission electron microscope operated at 200 kV. The samples were dispersed on carbon-coated copper grids and negatively stained with 2% uranyl acetate.

The crystallinity of the NiO@SiXNS samples was assessed using X-ray diffraction (XRD). The XRD patterns were recorded using a Bruker D8 Advance diffractometer with Cu Kα radiation. The 2θ range was from 5° to 80°, and the scan rate was 0.02°/s.

The atomic composition of the NiO@SiXNS samples was determined using X-ray photoelectron spectroscopy (XPS). The samples were analyzed using a Thermo Scientific K-Alpha XPS spectrometer. Samples were prepared for XPS analysis by adhering powders to carbon substrates; all peaks were energy-calibrated relative to the C1s peak of adventitious carbon at 284.8 eV and data analysis was performed using Thermo Scientific’s Data Analysis software.

The electronic structure of the NiO@SiXNS samples was studied using X-ray absorption near-edge structure (XANES). The XANES spectra were recorded at Sector 9-BM of the Advanced Photon Source at Argonne National Laboratory (Lemont, IL). A Quick-EXAFS, double-crystal silicon crystal monochromator was used to collect the XANES spectra.

The concentration of CO and CH4 in the gas mixture was measured using a gas chromatograph-mass spectrometer with a 60 m GS-CarbonPLOT column. The retention times for CO and CH4 were 0.33 and 2.08 minutes, respectively.

The catalytic performance of the NiO@SiXNS samples was studied using a fixed-bed reactor. The reactor was a quartz tube with an inner diameter of 10 mm and a length of 100 cm. The catalyst (0.5 g) was placed in the middle of the reactor, and the gas mixture was supplied at different flow rates over the catalyst. The gas mixture consisted of 10% H2 and 90% N2. The reaction temperature was varied from 300° to 400°C.

Gas-phase CO2 reduction tests. Gas-phase photocatalytic reduction was conducted in a plug-flow flow reactor in accordance with the procedure reported in our previous work. For the catalytic investigation, the sample (~10 mg) was first heated to 300°C in H2 (4 s.c.c.m.) and reduced in the dark for 1 h (yielding Ni/SiXNS). Next, a continuous flow of CO2 was introduced, and the reaction was monitored by monitoring the production of CO and CH4.

In situ DRIFT studies. In situ DRIFT experiments were performed on a Thermo Scientific Nicolet iS50 FT-IR Spectrometer with a mercury cadmium telluride detector cooled with liquid nitrogen. The spectrometer was equipped with a Harrick Spectra-Mantis diffuse reflection accessory, and a Harrick high temperature reaction chamber (HTC) with ZnSe windows. The temperature was controlled by a Harrick ATC-024-3 Temperature Controller. The spectrometer data were obtained using OMNIC software (Thermo Scientific). The autosampling function in the OMNIC software was used to collect data at specific time intervals.

The loaded sample in the HTC was pressed carefully to ensure a flat and even surface. The resulting procedure for data collection was used. The sample was first reduced for 1 h at 300°C in H2 (20 s.c.c.m.). Then the background scan was collected under flowing He at the same temperature, such that the spectra collected afterwards were differential spectra. Finally, CO2, H2, and He were introduced to the sample at a rate of 2, 8, and 10 s.c.c.m. The spectra were collected for 60 min with a resolution of 4 cm⁻¹ and 32 scans.

Data availability
The authors declare that the main data to support the findings of this study are available within the article and its Supplementary Information. The source data underlying all the plots in the main text and Supplementary Information are provided and available from the corresponding author upon request.

Received: 25 October 2018 Accepted: 7 May 2019
Published online: 13 June 2019

**References**

1. Aresta, M., Dibenedetto, A. & Angelini, A. Catalysis for the valorization of exhaust carbon: from CO2 to chemicals, materials, and fuels. Technological case of CO2, Chem. Soc. Rev. 11, 1709–1742 (2014).

2. Ozin, G. A. Throwing new light on the reduction of CO2, Adv. Mater. 27, 5973–5963 (2015).

3. Schlögl, R. The revolution continues: Energiewende 2.0. Angew. Chem. Int. Ed. 54, 4346–4349 (2015).

4. Jelle, A. A. et al. Highly efficient ambient temperature CO2 photomethanation catalyzed by nanostructured RuO2 on silicon photonic crystal support. Adv. Energy Mater. 8, 1702277 (2018).

5. Wang, W., Wang, S., Ma, X. & Gong, J. Recent advances in catalytic hydrogenation of carbon dioxide. Chem. Soc. Rev. 40, 3703–3727 (2011).

6. Mills, G. A. & Steffen, F. W. Catalytic methanation. Catal. Rev. 8, 159–210 (1974).

7. Vogt, C. et al. Unravelling structure sensitivity in CO2 reduction over Cu and even CuO catalysts. ChemSusChem 9, 4436–4439 (2016).

8. Du, G. et al. Methanation of carbon dioxide on Ni-incorporated MCM-41 catalysts: the influence of catalyst pretreatment and study of steady-state reaction. J. Catal. 249, 370–379 (2007).

9. Votjã et al. Activity and stability of powder and monolith-coated Ni/GDC catalysts for CO2 methanation. Appl. Catal. B: Environ. 236, 384–395 (2018).

10. Budi, C. S., Wu, H.-C., Chen, C.-S., Saiikia, D. & Kao, H.-M. Ni Nanoparticles supported on cage-type mesoporous silica for CO2 hydrogenation with high CH4 selectivity. ChemSusChem 9, 2326–2331 (2016).

11. Chen, C.-S., Budi, C. S., Wu, H.-C., Saiikia, D. & Kao, H.-M. Size-tunable Ni nanopowders supported on surface-modified, cage-type mesoporous silica as highly active catalysts for CO2 hydrogenation. ACS Catal. 7, 8367–8381 (2017).

12. Peebles, D. E., Goodman, D. W. & White, J. Methanation of carbon dioxide on nickel(100) and the effects of surface modifiers. J. Phys. Chem. B, 4378–4387 (1983).

13. Falconer, J. L. & Zagli, A. E. Adsorption and methanation of carbon dioxide on Ni(111) studied by XPS. Surf. Sci. 329, 199–206 (1995).

14. Heine, C., Lechner, B. A. J., Bluem, H. & Salmeron, M. Recycling of CO2: probing the chemical state of the Ni(111) surface during the methanation reaction with ambient-pressure X-ray photoelectron spectroscopy. J. Am. Chem. Soc. 138, 13246–13252 (2016).

15. Hoffmann, F., Cornelius, M., Morell, J. & Fröba, M. Silica-based mesoporous organic–inorganic hybrid materials. Angew. Chem. Int. Ed. 45, 3216–3231 (2006).
16. Sayari, A. & Hamoudi, S. Periodic mesoporous silica-based organic–inorganic nanocomposite materials. Chem. Mater. 13, 3151–3168 (2001).
17. Liang, J., Liang, Z., Zou, R. & Zhao, Y. Heterogenous catalysis in zeolites, mesoporous silica, and metal–organic frameworks. Adv. Mater. 29, 1701139 (2017).
18. Potemkin D. I., et al. Porous nanocrystalline silicon supported bimetallic Pd–Au catalysts: preparation, characterization, and direct hydrogen peroxide synthesis. Front. Chem. 6, 85 (2018).
19. Halim, M. Y. A., Tan, W. L., Bakar, N. H. H. A. & Bakar, M. A. Surface characteristics and catalytic activity of copper deposited porous silicon powder. Materials 7, 7737 (2014).
20. Polissi, S. et al. In situ synthesis and catalytic activity in CO oxidation of metal nanoparticles supported on porous nanocrystalline silicon. J. Catal. 271, 59–66 (2010).
21. Polissi, S., Goller, B., Lapkin, F., Fairdough, S. & Kovalev, D. Synthesis and catalytic activity of hybrid metal/silicon nanocomposites. Phys. Stat. Sol. (RRL) 2, 132–134 (2008).
22. Weatherbee, G. D. & Bartholomew, C. H. Hydrogenation of CO on group VIII metals: I. specific activity of Ni55Si55O. J. Catal. 66, 67–76 (1981).
23. Le, T. A., Kim, T. W., Lee, S. H. & Park, E. D. Effects of Na content in Na/Ni/SiO2 and Na/Ni/CoO catalysts for CO and CO2 methanation. Catal. Today 303, 159–167 (2018).
24. Guo, M. & Lu, G. The effect of impregnation strategy on structural characters and CO2 methanation properties over MgO modified Ni55Si55O catalysts. Catal. Commun. 54, 55–60 (2014).
25. Sastre, F., Puga, A. V., Liu, L., Corma, A. & Garcia, H. Complete photocatalytic reduction of CO2 to methane by H2 under solar light irradiation. J. Am. Chem. Soc. 136, 6798–6801 (2014).
26. Wu, H. C. et al. Methanation of CO2 and reverse water gas shift reactions on Ni55Si55O: the influence of particle size on selectivity and reaction kinetics. Catal. Sci. Technol. 6, 85 (2018).
27. Li, S. et al. Siloxene nanosheets: a metal-free semiconductor for water splitting. J. Mater. Chem. A. 4, 15841–15844 (2016).
28. Badley, R. D. & Johnson, M. M. Siloxene-supported catalysts for ethylene polymerization. J. Catal. 141, 524–532 (1993).
29. Ohashi, M., Yasokawa, R., Takatani, Y. & Nakano, H. Versatile reducing reaction field within layered polyvalent for efficient one-pot synthesis of metal nanoparticles. ChemNanoMat 3, 534–537 (2017).
30. Yan, Z., Fu, L., Zuo, X. & Yang, H. Green assembly of stable and uniform silver nanoparticles on 2D silica nanosheets for catalytic reduction of 4-nitrophenol. Appl. Catal. B: Environ. 226, 23–30 (2018).
31. Ono, Y., Sendoda, Y. & Keii, T. Siloxene as a unique catalyst for structural isomerization of butene. J. Am. Chem. Soc. 97, 5284–5285 (1975).
32. Krishnamoorthy, K., Pazhamalai, P. & Kim, S.-J. Two-dimensional siloxene nanosheets: novel high-performance supercapacitor electrode materials. Energy Environ. Sci. 11, 1593–1602 (2018).
33. Nakano H., Ishii M., Nakamura H. Preparation and structure of novel siloxene nanosheets. Chem. Commun. 2944–2947 (2005).
34. Weiss, A., Beil, G. & Meyer, H. The topochemical reaction of CaS to a two-dimensional subsilicous acid Si2H4(OH)2 (Kautsky’s Siloxene). Z. Naturforsch. 34b, 25–30 (1979).
35. Kautsky, H. & Zocher, H. Über die Beziehung zwischen Chemi- und Photolumineszenz bei ungesättigten Siliziumverbindungen. Z. fär. Phys. 9, 267–284 (1922).
36. Fuchs, H. D. et al. Porous silicon and siloxene: vibrational and structural properties. Phys. Rev. B 48, 8172–8193 (1993).
37. Fu, R. et al. Two-dimensional silicon suboxides nanostructures with Si nanodomains confined in amorphous SiO2 derived from siloxene as high performance anode for Li-ion batteries. Nano Energy 39, 546–553 (2017).
38. Clause, O., Bonnevet, L. & Che, M. Effect of the preparation method on the thermal stability of silica-supported nickel oxide as studied by EXAFS and TPR techniques. J. Catal. 138, 195–205 (1992).
39. Ocelli, M. L., Pasaras, D., Suib, S. L. & Stencel, J. I. Metal contaminant effects on the properties of a silica-rich fluid cracking catalyst. Appl. Catal. 28, 143–160 (1996).
40. Meng, X. et al. Photothermal conversion of CO2 into CH2 via H2 over group VIII nanocatalysts: an alternative approach for solar fuel production. Angew. Chem. 126, 11662–11666 (2014).
41. Zhang, Z., Verykios, E. X., MacDonald, S. M. & Affrossman, S. Comparative study of carbon dioxide reforming of methane to synthesis gas over Ni/La2O3 and conventional nickel-based catalysts. J. Phys. Chem. 100, 744–754 (1996).
42. Robbins, J. L. & Maruishi-Soos, E. Evidence for multiple carbon monoxide hydrogenation pathways on platinum alumina. J. Phys. Chem. 93, 2885–2888 (1989).
43. J. Lou, H., Mo, L. & Zheng, X. The reactivity of surface active carbonaceous species with CO2 and its role on hydrocarbon conversion reactions. J. Mol. Catal. A: Chem. 316, 1–7 (2010).
Additional information
Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-019-10464-x.

Competing interests: The authors declare no competing interests.

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Journal Peer Review Information: Nature Communications thanks Antonio Vita and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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