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surfaces: CO₂ marker detection and storage†

2-D organization of silica nanoparticles on gold

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A single layer of silica nanoparticles with an average size of \sim 200 nm was deposited over the surface of pristine gold wafers, aided by (3-mercaptopropyl)trimethoxysilane. The nanoparticle immobilization was driven by covalent bonding rather than a self-assembly process, leading to a cluster-assembled material which has CO_2 sensing features. Here, we show how this device can be used for CO_2 physisorption and chemisorption. We analyse the device, both spectroscopically and morphologically, before and after exposure to an atmosphere of 7 mbar of CO_2 , inside a planetary atmospheres and surfaces simulation chamber, (PASC) mimiking Martian atmospheric conditions. Our studies demonstrate that these clusters are suitable for CO_2 detection and storage, under well controlled experimental Martian conditions. Their high sensitivity at a very low concentration of CO_2 , 12.4 ppm, makes them ideal candidates in the nanosensor field

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Introduction

CO₂ detection is crucial in biological, agricultural, food, medical and environmental industries,1-4 and planetary science. CO2 is also known as a greenhouse gas, as it absorbs and reemits IR radiation, increasing the average temperature on earth. It is industrially employed for agricultural fumigation and for experimental plant growth. When mixed with oxygen it forms carbogen, which is used to stimulate breathing in the treatment of respiratory disease, and to increase subsurface oil recovery, and it is also widely used in carbonated drinks. Therefore, CO₂ gas sensing is often required to guarantee compliance with environmental laws and regulations. In particular, carbon dioxide monitoring is required in the working environment, as an abnormal high level of CO2 is associated with sick building syndrome. Nowadays, CO₂ sensors are based on IR adsorption or electrochemical interactions that measure resistance or current.5-7 Devices using such technology have an important drawback as they typically suffer from a high cross sensitivity to other gases and have poor temperature stability. At present, there is interest in the development of alternative solutions, materials with treatments that can capture CO₂ and/or show sensitivity to CO₂. We have focused

Nanoparticles (NPs) exhibit unique chemical, physical and electronic properties that are different from those of bulk materials, they indeed own outstanding catalytic, thermal and optical properties as well as robust mechanical strength and a large surface-to-volume, making them good candidates to construct sensing devices, based on clustered-nanoparticles. Although CO₂ nanoparticle based sensors have been mentioned in the literature during the last decades, ⁹⁻¹¹ they have been barely implemented and developed for practical applications as they require a complex nanoparticle deposition set-up, which most of the times leads to either NP-aggregates¹² or multilayers. ¹³

Silica NPs, (SiO₂NPs) in solid state, have never been investigated as a potential CO₂ sensing agent. There is only one work in this regard but implemented as a colloidal solution;¹⁴ others, were devoted to incorporate amine derivatives^{15,16} over their surface taking advantage of the well-known CO₂ capture properties in liquid phase of amine derivatives such as: monoethanolamine (MEA), diethanolamine (DEA), diisopropanolamine (DIPA) *etc.* However, materials/polymers bearing amine groups, in practice, suffer from poisoning with carbamates and fail to fill the gap in the CO₂ sensor sector.^{17,18}

our attention on nanoparticle-based CO_2 sensors, as a promising alternative for certain applications. However, the preparation of materials with nanoparticles can still be challenging due to their tendency to self-organize when deposited on thin films and the sensitivity of these procedures to various parameters. It is the purpose of this work to describe a procedure for fabrication of CO_2 adsorbing surfaces, after deposition of nanoparticles on a linker film. As an example, the response of these nanoparticle clusters is here demonstrated after exposure to a pure CO_2 Martian-like atmosphere.

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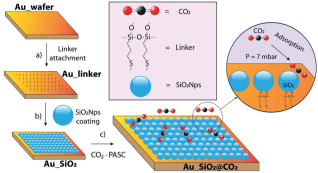


Fig. 1 Schematic representation of the SiO_2NPs deposition onto Au wafers and the subsequent CO_2 adsorption process.

In this work we proposed to use silica nanoparticles (SiO₂NPs) of 200 nm size. This is a popular material in the biomedicine field. Furthermore, they are inexpensive, and this can open new routes in the field of space and planetary exploration. SiO2NPs also benefit from several advantages, e.g., high surface area, synthetic modulability, particle size tunability via sol-gel process, 19,20 geometry and porosity control, and its demonstrated CO2 adsorption capacities in suspension (nanofluids),21 all of them suggesting their potential application as a non-liquid adsorbent material, a feature barely investigated for this particular goal. Thus, here we aim to demonstrate the formation of cluster-assembled films, by the self-organization of SiO₂NPs on gold surfaces, to elucidate whether non-mesoporous silica structures are suitable as CO2 adsorber at standard Mars atmosphere conditions of 7 mbar (12.4 ppm of CO₂ inside the planetary environment simulating chamber PASC²²). If successful, this nanotechnology may have interest for the fixation of organic molecules and for the detection of bio-signatures in future Martian missions.23 We shall next demonstrate the twodimensional organization of silica nanoparticles on gold surfaces and characterize spectroscopically their CO2 fixation signatures. Our methodological approach is depicted in Fig. 1 and encompasses a bottom-up approach where a bare gold wafer $(11 \times 11 \times 1 \text{ mm})$ is treated with a molecular linker that is incorporating both a thiol and an ethoxysilane group, at the extremes of a propyl chain. This linker acts as a "glue" between the SiO₂NPs and the gold wafer surface (Au-linker, Fig. 1a). In a step further (Au_SiO₂, Fig. 1b) the wafer was placed in the 200 nm size SiO₂NPs colloidal suspension (1 wt%) at different dipping time conditions. After exhaustively rinsing with water, a SiO₂NPs single monolayer (this is confirmed later on using microscopy techniques) was obtained. Finally (Au_SiO₂@CO₂, Fig. 1c), CO₂ adsorption experiments were conducted under specific conditions inside the Planetary Atmospheres and Surfaces Simulation Chamber (PASC, Fig. S1†).22

2. Experimental

2.1. Nanosensor design and chemistry

The design of the sensor resulted from the idea of producing a material, which can be tailored to exhibit new collective properties which differ from those of their individual components. The proposed strategy combines in the same chip, (i) the covalent attachment of monodispersed silica nanoparticles to (ii) an excellent conductor of electricity (Au), a material that promotes covalent bonding (Au–S–Si) between nanoparticles and the surface of the wafer. The gold wafers, Au (111) were manufactured as (11 \times 11 \times 1 mm, ArrandeeTM) and deposited over a glass substrate. The further self-organized nanoparticle layer was achieved as follows. We have made the deposition of SiO₂NPs onto Au wafers, *via* 3-MPTS (3-mercaptopropyl triethoxysilane), which is a suitable molecular adhesive linker for the introduction of silica nanoparticles onto gold surfaces. This particular linker is bearing both a thiol and a silane group along a propyl chain, thus enabling SiO₂NPs deposition in an orderly fashion, following previous studies carried out by Vakarelski *et al.*⁸

2.2. 3-MPTS coating of Au surfaces

Over a polycrystalline Au layer on glass, a 40 mM solution of 3-MPTS in MeOH was added (366 µL, in 50 mL) then, the wafer was kept immersed for 3 h. Thereafter, the wafer was rinsed with MeOH, dried, and subsequently submerged for 3 h in a freshly prepared solution of 0.01 M NaOH (to induce hydrolysis/polymerization resulting in a Si-O-Si external glass layer). Afterwards, the wafer was again submerged in a pH = 9.0, 1.0 wt% SiO₂ nanoparticles suspension (500 mg, 50 mL) for 22 hours prior sonication and centrifugation at 3k rpm for 3 minutes. Finally, the sensor was exhaustively rinsed with H2O mQ and MeOH to exclude the nonspecific binding, and it was allowed to air dry. In the present work, the deposition protocol was implemented as the nanoparticles are doubling in size (200 nm, Fig. S2†). We considered the size of the nanoparticles (200 nm) to be optimal, as we expected to have a nanomaterial showing a balance between (i) aggregation when film-coating (smaller NPs might promote larger self-aggregation) and (ii) effective surface area (larger NPs possess less effective surface area). Also, the thickness of the monolayer plays an important role in the adsorption capacity of the material. In that regard, the immersion step was extended for 22 hours, thus leading to a partially saturation of the Au wafer surface which was confirmed later on by microscopies techniques: Scanning Electron Microscopy (SEM) and Atomic Force Microscopy (AFM). CO₂ adsorption studies were carried out under wellcontrolled experimental conditions, inside PASC. The experimental setup is described in the ESI (Fig. S1†). The gold (Au_SiO₂) surface samples were exposed to an atmosphere of CO₂ at 7 mbar (similar to the present-day Martian surface conditions) during 22 hours at 25 °C. After being exposed to the CO2 atmosphere inside PASC, samples were immediately introduced under high vacuum conditions, and then transferred to ultra-high vacuum conditions for XPS measurements or storage under vacuum conditions before infrared (IR), SEM and AFM analysis.

Results and discussion

The sensor, comprising a single monolayer of embedded silica nanoparticles loaded over a 1 mm thick gold wafer, was

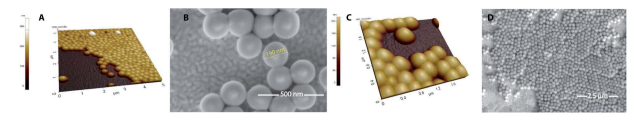


Fig. 2 AFM (A and C) and SEM ((B) scale bar 500 nm; (D) scale bar 2.5 μ m) micrographs of 3-MPTS coated Au-wafers submerged 22 h in a SiO₂ nanoparticle suspension.

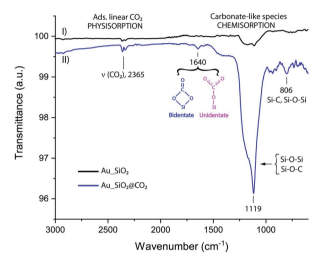


Fig. 3 IR spectrum of Au wafer bearing SiO₂ nanoparticles (I) and a gold wafer bearing SiO₂ nanoparticles after being exposed to CO₂ atmosphere inside PASC (II).

characterized by several spectroscopic and microscopic techniques before and after CO₂ exposition. In addition, its theoretical IR spectrum was obtained using computational calculations.

3.1. SEM and AFM microscopies

The adsorption and two-dimensional organization of silica nanoparticles (SiO₂NPs) on gold surfaces and its morphological

characterization by SEM, Fig. 2B and D and AFM, Fig. 2A and C, confirmed a well-organized single layer deposition. As expected, there is a time dependence in the coating process, and we have reached a predominant single layer deposition after 22 hours of immersion time. Surprisingly, it is observed that the NPs tend to be adhered between them despite their apparent negative zeta potential at pH = 9.0, hence, a marked repulsion was expected. However, this can be explained with the existence of a hairy (gel like) external layer, comprising a short, flexible (polymer-like) fragments of poly(silicic acid) attached all over the surface (Fig. S3 and S4†).24-26 The morphology of the nano-structures showed a regular and monodispersed distribution along a NPs row; regarding the surface characteristics of the nanoparticles, it appears to be smooth objects, both in AFM and SEM despite the sol-gel method employed for the synthesis of the nano-compounds, which it is expected to generate some level of "random" porosity. Additional SEM images (Fig. S5†) were used to determine the area covered by the nanoparticles. From the number of NPs observed by SEM, the coverage of the monolayer is calculated to be 55 \pm 5%. This is an excellent result as the deposition was not driven through self-assembly processes. Furthermore, NPs preserve their integrity after deposition as 90% kept the 200 nm size. According to the AFM profile measurements (Fig. S6 and S7†), the thickness of the single layer was estimated to be 200 nm, consistent with the NPs size. Novel surface spectroscopic characterization, by reflection adsorption infrared spectroscopy (RAIRS) and X-ray photoelectron spectroscopy (XPS) of these nanostructures, let us establish

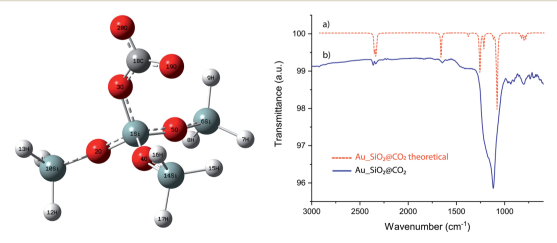


Fig. 4 Schematic illustration of a representative portion of silica surface binding a CO₂. Superimposed IR spectra of theoretical (a) and experimental (b) Au_SiO₂ when exposed to a CO₂ atmosphere.

their chemical composition after their deposition on the gold surface (Fig. 3–5).

3.2. IR spectroscopy

The FTIR was used in this stage of the work to characterize the successful deposition of SiO2NPs on gold surfaces and to identify chemical changes after exposure to the CO₂ (g) atmosphere. We searched for signatures of the presence of either CO2 trapped inside the SiO₂ pores (physisorption) or covalently anchored in the most plausible form of carbonates (chemisorption). This method allows to also distinguish oxidation process of the SiO₂NPs under CO₂ conditions. FT-IR spectroscopy provides qualitative information about the chemical nature of NPs surface, an important factor when exploring the way, a molecule is interacting and adsorbed on the NPs surface. Fig. 3 shows the superimposed FTIR spectra of SiO₂NPs anchored in an Au wafer, before (Fig. 3-I) and after (Fig. 3-II) CO2 exposure. The spectrum of pristine SiO2 nanoparticles (Au_SiO₂, Fig. 3-I) confirms the presence of the antisymmetrical stretching vibration of Si-O-Si at (1010-1130 cm⁻¹),²⁷ hence, the IR technique is suitable for detecting the grafted adsorption of SiO₂NPs on gold surface. The spectrum of SiO₂ nanoparticles (Au_SiO₂@CO₂, Fig. 3-II) after CO₂ atmosphere exposure inside PASC was also recorded, thus displaying similar IR bands to the ones of "pristine" SiO2 nanoparticles; however, in this particular situation, the peaks are narrower and much more intense (1010-1130 cm⁻¹), which can be ascribed to the formation of new carbon silica surface bonds, Si-O-C, that falls in the range

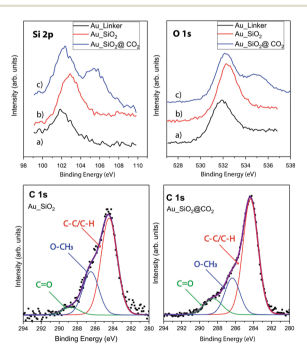


Fig. 5 XPS photoemission spectra of Si 2p and O 1s core level peaks along the three steps of the process on gold surface after: (a) adsorption of linker (Au_linker), (b) SiO_2NPs attachment on gold (Au_SiO₂) and (c) CO_2 exposure (Au_SiO₂@CO₂). Deconvoluted XPS spectra of C 1s before (Au_SiO₂) and after (Au_SiO₂@ CO₂) CO_2 exposure.

of 1150 cm⁻¹.²⁸ Furthermore, it is noticeable the presence of two new broad peaks, one at 805 cm⁻¹ that can be attributed to Si-C stretching vibrations and Si-O-Si (symmetrical stretching),^{29,30} and a second one, barely seen, at 1640 cm⁻¹ than can be assigned to uni/bidentate carbonates (Si-CO₃)³¹ providing a spectroscopical fingerprint that confirms the successful CO₂ adsorption process. A control experiment was conducted in order to assess the capacity of SiO₂NPs for CO₂ adsorption, by using a pristine Au wafer in the absence of nanoparticles. Then it was exposed to 7 mbar of CO₂ inside PASC, concluding that adsorption of CO₂ is inherent of silica nanoparticles.

3.3. Computational details

Ab initio calculations of the IR spectra of a representative SiO₂NPs (Fig. 4) fragment was calculated, when a CO₂ molecule is chemisorbed forming a carbonate like group. This was performed with the Gaussian 09W software package, with B3LYP density functional theory (DFT) and basis set 6-311+G(d, p). It is evident the good overlap when superimposing the infrared spectrum of both the theoretical and experimental (Fig. 4), thus confirming the existence of chemisorbed CO₂ specie in the form of carbonates, over the surface of the NPs. The fragment, here studied, displays a tetrahedral symmetry in the ground state with bond angles values of approximately 109.5°. The vibrational modes registered are mainly symmetrical and antisymmetrical (Fig. S8†) with respect to each of the planes. The theoretical and experimental IR spectra are a perfect match, however a scaling correction factor of 1.06 have been required for DFT values.

- (i) Region 3000–1500 cm⁻¹: the computed DFT analysis predicted CH stretching frequencies (2350 cm⁻¹) for the unidentate structure; however, these frequencies can be more likely associated to CO₂ confinement in the interior of the material.³¹ The second region of interest is around 1650 cm⁻¹ that can be ascribed to the umbrella stretching of the chemisorbed R-CO₂ group, a very significant (fingerprint) wavenumber value as it confirms the adsorption of the CO₂ by the matrix.
- (ii) Region 1200–900 cm $^{-1}$: the bands observed between 1200 cm $^{-1}$ and 1030 cm $^{-1}$ are mainly Si–O, Si–O–C and C–O stretches. There is also a small band around 1270 cm $^{-1}$ that can be attributed to the $\rm CO_2$ antisymmetrical stretching.
- (iii) Region 800–700 cm⁻¹: the most intense IR band in this region is located at 790 cm⁻¹, and it is attributed to the asymmetrical stretches between the Si and O.

3.4. XPS analysis

In addition, and with the aim of confirming the appearance of these new chemical species, XPS analysis were performed in the same samples at three different adsorption-process steps: Au_linker, Au-SiO₂ and Au_SiO₂@CO₂. XPS is considered to be the most sensitive technique and it is widely used to determine the exact elemental ratio and bonding nature of the elements in NPs materials. The XPS overview spectra of the three different wafers shows that the following atomic species can be identified: O, C, S, Si and Au. Fig. 5 shows the core-level XPS spectra of

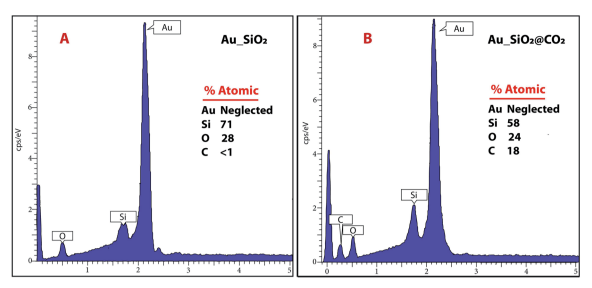


Fig. 6 EDX spectra of (A) the sensor before CO₂ exposure and (B) after CO₂ adsorption

the O(1s), C(1s) and Si(2p) regions along the three steps of the process. The Si 2p peak is observed at 102.3 eV, and is attributed to Si-C and Si-O species from the chemisorption of the linker (Fig. 5a).32 In a second step (Fig. 5b), the main peak shows a shift to 102.8 eV in binding energy, related to the SiO₂NPs adsorption of the gold surface, masking the underneath linker layer. Finally (Fig. 5c), when the sensor is exposed to CO₂, it gives rise to the appearance of a second component at 105.5 eV, which is ascribed to the presence of electron withdrawing groups33 on the surface. This fits, in our case, with the formation of Si(CO₃²⁻) species.^{34,35} The best-fit curves for the O 1s, depicted in Fig. 5, display a similar trend as silicon, where there is barely a shifting between Au linker and Au SiO₂ (sensor), then it is noticeable the case of Au_SiO2@CO2 showing two main components. One is observed at 531.7 eV, attributed to OH and Si-O species, and a second one, in accordance with the silica case, at approximately 535 eV which has been related to C-O containing species.³⁶⁻³⁸ Finally, the best-fit curve for the C 1s peak (Fig. 5), in both cases (Au_SiO₂ and Au_SiO₂@CO₂), was achieved using three components, the first carbon component has a binding energy (B.E.) of 284.5 eV and is attributed to the C-H and C-C group, the second component at 286.4 eV corresponds to O-CH₃ groups whereas the third component is observed at 288.6 eV and is assigned to the C=O groups.38 There is a remarkable increase in the third component (at 288.6 eV, assigned to the C=O groups), from 5% to 11% (Table S1†) when the Au_SiO₂ have been exposed to CO₂ which supports the statement of silica nanoparticles as suitable materials for CO₂ adsorption at low pressures. This is in good agreement with the infrared analysis. Both complementary spectroscopies, XPS and infrared, helped us to identify the CO₂ adsorption process and chemical adsorption species, providing a fingerprint signal when the CO₂ fixation process takes place.

3.5. EDX analysis

Energy Dispersive X-ray Analysis (EDS or EDX, Fig. 6) technique was used for performing elementary analysis and chemical

characterization of the nanosensor, before (Au_SiO2) and after CO₂ exposure (Au_SiO₂@CO₂), in conjunction with SEM. The impact of the electrons beam on the sensor produces X-rays that are characteristic from the atoms present in the region of study. EDS analysis is then a suitable tool for the determination of the atomic composition of individual spots or for larger areas through a mapping of elements. Based on the results of EDS analysis data, it is possible to conclude that the adsorption of CO₂ is inherent to silica nanoparticles. The atomic percentage of carbon before and after CO2 exposure, gave the values of about <1 and 18% respectively. This marked increase is ascribed to the formation of chemisorbed carbonates on the surface of the spherical nanoparticles. The electron beam was also pointed in an empty area to assess the intrinsic capacity of the NPs to retain CO₂, and as expected only the Au peaks were observed in the EDS profile.

4. Conclusions

This work show cases the development of an ultra-sensitive sensor, comprised by a well-organized deposition scheme where a monolayer of SiO₂ nanoparticles (200 nm) is efficiently built on a gold (Au) solid substrate, assisted by chemical reactions between NPs and the surface wafer, 3-MPTS mediated. The growth of the monolayer was promoted by the formation of covalent linkages rather than self-assembly processes confirming the importance of the organic linker. This nanosensor can be used as CO₂ gas sensor with excellent sensing performance (12.4 ppm) for CO₂ gas at 7 mbar at room temperature. The sensing response was confirmed by spectroscopies techniques (IR, XPS and EDX) and the morphological features were captured by SEM and AFM microscopies.

These techniques also, provide valuable information regarding the inherent capacity of SiO_2NPs to capture CO_2 , as atomic EDX analysis can discriminate between the composition of empty and full areas within the wafer.

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In conclusion, we have demonstrated that SiO₂NPs can be self-organized in a two-dimensional ordered structure, which can be fully characterized both spectroscopically and morphologically. Furthermore, we have investigated the suitability of SiO₂NPs deposited in gold wafers as a tool for CO₂ recognition at standard Mars low pressures. Their low-cost fabrication, small size, and sensitivity (12.4 ppm), opens a new route towards their application in solar system objects possessing very thin atmospheres. These nanosensors could also expand their applications towards the detection of other gasses traces in explored/unexplored solar bodies. The nanosensors could also be used for (i) atmospheric biosignatures detection if referring to the presence of one or several gas species, and (ii) surface biosignatures if detectable by their light reflection characteristics providing spectral molecular fingerprints or characteristic chemical features. Moreover, these results open the door to future studies involving the use of Mars minerals as an atmospheric trap and reservoir. Finally, the aeolian dust which is so characteristic of the Martian atmosphere, has a measured content of about 44% of SiO2.39 This material surface preparation may be used in the future to investigate, within a controlled experimental facility such as PASC, the chemical catalytic reactions that may occur on the surface of mineral aerosols (rich in SiO2) in suspension on the Martian CO₂ atmosphere, in contact with other trace molecules, when exposed to the high levels of UV radiation that are characteristic of the Martian atmosphere. In that regard, nanosensors will dramatically improve high-performance contribute to the future space research.

Conflicts of interest

There are no conflicts to declare.

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