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Original Article

Network hydration, ordering and composition interplay of chemical vapor deposited amorphous silica films from tetraethyl orthosilicate



Babacar Diallo ^{a,1}, Konstantina C. Topka ^{b,c,1}, Maxime Puyo ^b, Charlotte Lebesgue ^b, Cécile Genevois ^a, Raphael Laloo ^b, Diane Samelor ^b, Hélène Lecoq ^a, Mathieu Allix ^a, Hugues Vergnes ^c, François Senocq ^b, Pierre Florian ^a, Vincent Sarou-Kanian ^a, Thierry Sauvage ^a, Marie-Joelle Menu ^b, Brigitte Caussat ^c, Viviane Turq ^b, Constantin Vahlas ^b, Nadia Pellerin ^{a,*}

^a CNRS, Conditions Extrêmes et Matériaux: Haute Température et Irradiation (CEMHTI) UPR 3079, Université d'Orléans, Orléans, France

^b Centre Interuniversitaire de Recherche et d'Ingénierie des Matériaux (CIRIMAT), Université de Toulouse, CNRS, Toulouse, France

^c Laboratoire de Génie Chimique (LGC), Université de Toulouse, CNRS, Toulouse, France

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ABSTRACT

The chemical or mechanical performance of amorphous SiO₂ films depend on intrinsic physicochemical properties, which are intimately linked to atomic and molecular arrangements in the Si-O-Si network. In this context, the present work focuses on a comprehensive description of SiO₂ films deposited from a well-established chemical vapor deposition process involving tetraethyl-orthosilicate, oxygen and ozone, and operating at atmospheric pressure in the range 400-550 °C. The connectivity of the silica network is improved with increasing the deposition temperature (T_d) and this is attributed to the decreased content of hydrated species through dehydration-condensation mechanisms. In the same way, the critical load of delamination increases with increasing T_d thanks to the silicon substrate oxidation. The utilization of a O_2/O_3 oxidizing atmosphere involving the oxidation of intermediates species by O₂, O₃ and O, allows increasing the deposition rate at moderate temperatures, while minimizing carbon, H₂O and silanol contents to extremely low values (4.5 at.% of H). The SiO_x stoichiometry and T_d interplay reveals two distinct behaviors before and above 450 $^\circ\text{C}.$ The best corrosion resistance of these films to standard P-etching test is obtained for the minimum silanol content and the best network molecular ordering, with an etching rate of 4.0 ± 0.1 Å/s at pH = 1.5. The elastic modulus and hardness of the films remain stable in the investigated range of deposition temperature, at 64.2 \pm 1.7 and 7.4 \pm 0.3 GPa respectively, thanks to the low content in silanol groups. © 2021 The Author(s). Published by Elsevier B.V. This is an open access article under the CC

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* Corresponding author.

E-mail address: nadia.pellerin@univ-orleans.fr (N. Pellerin).

¹ Co-first authors: Babacar Diallo and Konstantina C. Topka.

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1. Introduction

Since the seventies, SiO2-based films have received much attention due to their well-established properties for electrical, optical, magnetic or barrier applications in most Key Enabling Technologies [1-4]. In particular, amorphous SiO₂ films are largely used as dielectric materials in electronic and optoelectronic devices [4-6], or as gas diffusion barrier on various plastic packaging materials [7]. For these and other applications, the chemical durability of the SiO₂ films is a major parameter, which is intimately linked to atomic and molecular arrangements in the Si-O-Si network, associated with the short (over 1.5–2.5 Å) and medium- (over 2.5–4.5 Å) range orders, respectively, investigated by nuclear magnetic resonance or infrared spectroscopy, for instance [8-10]. Indeed, investigations of the medium-range order in glass networks revealed that chemical durability depends on the film molecular structure and composition through the distribution of hydrolysis sites and the subsequent access of water to these sites. At least four mechanisms control the glass durability, namely (i) ion exchange in presence of modifier cations, e.g. alkali or alkaline earth ions; (ii) hydration through the diffusion of hydrated species; (iii) accelerated matrix dissolution by hydrolysis; and (iv) formation of a passivating surface layer [11-14]. Then, Osseo-Asare [15] has underlined the contribution of the surface chemistry and suggested pH dependent protonation and deprotonation of the surface silanol groups to give positively (Si-OH₂⁺) and negatively (SiO⁻) charged sites respectively. By way of comparison, the dissolution rates of quartz and amorphous silica in pure water are 4.2 \times 10^{-14} and 9.0 \times 10^{-13} mol/m² s, respectively, and their solubility stays very low for pH < 9[16,17]. The difference in these dissolution rates can be explained by the network geometry dependence of the dissolution kinetics, which imply that some surface bonding geometries react more readily than others [18]. This dependency elevates the film density and the degree of ordering of the Si-O-Si network as the main parameters that affect the dissolution of the SiO₂ films. In the same way, considering the mechanical properties of glasses, elastic moduli and Young's modulus in particular, do not solely depend on the interatomic bonding energy but also depend on the coordination, on the polymerization degree (cross linking), on the atomic packing density, and on the molecular organization [10]. It thus appears that in-depth knowledge of the structure of SiO₂ films and its twofold comprehensive correlation with the targeted functional properties and the processing conditions are essential for developing further all advanced applications.

Optical [19], fouling and corrosion [20], barrier [21] or electrical [22] properties of SiO₂ films produced through a variety of processes are continuously studied in the literature to this day. However, few of them focus on the structural organization of the SiO₂ network, especially in terms of its hydration. In a recent study, Jung et al. [23] studied the dielectric properties of SiO₂ films by plasma enhanced chemical vapor deposition (PECVD) and plasma enhanced atomic layer deposition (PEALD) from bis(diethylamino)silane (H₂Si[N(C₂H₅)₂]₂, BDES) O₂, intended for flexible and thin-film-transistor (TFT) devices, based on their chemical composition and growth characteristics. The authors reported decreased leakage currents at higher substrate temperatures, correlating this to an improvement of SiO₂ film stoichiometry at elevated temperatures. Measured by Xray photoelectron spectroscopy (XPS) however, their stoichiometry values do not account for hydrated species. More recently, Kim et al. [24] performed deposition of high quality SiO₂ films from 1,2-bis-(diisopropylamino)disilane (BDIPADS) and ozone, using thermal atomic layer deposition (ALD). The authors investigated the chemical and electrical characteristics of the SiO₂ films, observing slight degradation in the dielectric constants, leakage currents, and breakdown fields with decreasing deposition temperature. The authors attributed this to slight decrease in film density with parallel increased defect density of Si-H related bonds. Specifically, they noted the presence of the $(O-)_3Si-H$ bending mode at 880 cm⁻¹ at low temperatures and its disappearance at higher growth temperatures, giving in turn rise to the Si-O-Si bend vibrational mode at 800 cm⁻¹. However, quantitative information on the level of network hydration are not reported.

Securing long-term performance of such devices and systems also relies on their mechanical properties (hardness, elastic modulus, scratch resistance), which are heavily influenced by the deposition process. As pointed out by Simurka et al. [25], the mechanical properties of SiO₂ films vary largely in the literature. In their own study, hardness and elastic modulus were noted to decline from 8.5 to 2.2 GPa and from 73.7 to 30.9 GPa, respectively, with increase of the process pressure. The authors attributed this decrease to the less dense structure of the films prepared at higher process pressure, with these films also exhibiting absorptions related to hydrogen-bonded hydroxyl groups and adsorbed water. Quantitative data on the amount of hydrogen incorporated in the SiO₂ films were not given however.

The present work aims to bridge this gap, by correlating process conditions and functional properties to the molecular organization, structure and network hydration of SiO₂ films, subscribing to the requirements for an in-depth understanding of the material in view of advanced applications. We have processed SiO₂ films by oxygen-ozone O₂/O₃ mixture assisted atmospheric pressure chemical vapor deposition (CVD) from tetraethyl orthosilicate (Si(OC2H5)4,TEOS), a well-established robust and scalable route, operating at low to moderate temperatures (400-550 °C). Despite the popularity of this process, the film quality is questionable, depending in particular on the T_d, which modifies the gaseous intermediate species generated from the decomposition of TEOS, and their subsequent involvement in the formation of the film [26]. Indeed, the high thermal and chemical stability and fracture toughness of silica [27,28] does not prevent the occurrence of defects that can still be present, along with sub- or over-stoichiometry [29,30]. Especially, SiO₂ films deposited below 450 °C often contain carbonrelated impurities and trapped charges leading to oxygen deficiency [31] and subsequently to degraded capacitance-voltage and current density-electric field behaviors, or to increased etching rate. The dependence between the oxidation and dehydration mechanisms on the one hand and the oxidant gas nature or the working temperature on the other hand, are not entirely understood. In particular, the role of O₂ and O₃ in the

production of oxygen radicals for the oxidation of TEOS and of secondary products is not clear, despite the development of apparent kinetic models able to predict the silica deposition rates and the local profiles of gas flow and temperature in the reactor, for instance [26,32]. Also, the nature and the quantity of hydrated species and their impact on the SiO_x stoichiometry are poorly investigated and are not well known. In this perspective, in the present work we probed the hydrated species in the films and correlated these with the SiO_x stoichiometry and T_d in the range between 400 and 550 °C.

We have analyzed the films by Fourier transform infrared (FTIR) spectroscopy to get insight in the short and medium range order structure and in the nature of hydrated species. We have combined this information with input from high-resolution transmission electron microscopy (HRTEM), ion beam analysis (IBA), time-of-flight secondary ion mass spectrometry (ToF-SIMS) and XPS. These investigations have allowed insight in the O/Si ratio and the concentration of hydrated species explored along the film depth profile, and on the connectivity of the Si–O–Si network. We have correlated these chemical and structural characteristics with the mechanical and surface properties of the films, namely their hardness, Young modulus and critical load of delamination probed by nanoindentation and nanoscratch, as well as their wettability and their etching rate in standard hydrofluoric acid (HF) solutions.

2. Materials and methods

CVD of amorphous SiO₂ films was performed in a horizontal, tubular, hot-wall reactor. The experimental setup for the deposition of SiO₂ films is schematically illustrated in Fig. 1 and has been developed previously [26]. The experiments were performed at four different T_d, namely 400, 450, 500 and 550 °C, defined as the temperature measured at the middle of the reactor, where an isothermal region exists. TEOS and O₂/O₃ flow rates were respectively 2 standard cubic centimeters per minute (sccm) and 1960 sccm (O₃ concentration of 60 mg/l) and a deposition time of 30 min was considered providing film thickness of approximatively 100 nm. Thicker films of about 300–400 nm have also been used for nanoindentation testing. The SiO₂ deposition rates were calculated from the weight difference before and after deposition, by measuring the

deposited mass using a Sartorius Genius Series ME215P instrument with 0.015 mg precision.

280 μ m thick monocrystalline silicon (100) wafers (Neyco S.A.), cut in rectangles of 32 \times 24 mm², were used as substrates. They were supported vertically by two custom-made substrate holders placed in the isothermal region [26].

The films thickness was estimated by ellipsometry, with a Semilab SE-2000 instrument, operating between 250 and 1000 nm wavelength range, using a constant incidence angle of 70°. The measurements were performed by probing 3 points vertically along the surface's center of each sample. The spectroscopic ellipsometry data were simulated and fitted in the 250-1000 nm wavelength range using the Semilab SEA software. The calculated results were collected for a linear regression R² fit of over 0.99. Water content was evaluated using the Bruggeman effective medium approximation (BEMA) model, applying a three components medium, including SiO₂, void and H₂O. The Sellmeier model was used as a reference to confirm the measured local thickness. The above models were validated by measuring the cross-section of one representative sample by a FEI Quanta450 scanning electron microscope (SEM).

The surface's roughness of the films was measured using a Bruker Dimension ICON, Nanoscope V atomic force microscopy (AFM) operating in tapping mode. The tip used was a TESPA V2 from Bruker, made of n-doped silicon, with a radius <10 nm. A scan size of 1 μ m² was implemented for all measurements.

The chemical composition of the films (carbon, hydrogen content and O/Si atomic ratio) was determined by using IBA. Silicon, oxygen and hydrogen contents were determined by Rutherford backscattering spectroscopy (RBS), nuclear reaction analysis (NRA) and elastic recoil detection analysis (ERDA), respectively. RBS was performed at a 166° detection angle with 2 MeV α particles. Oxygen concentration was measured through the $^{16}O(d,\alpha_0)^{14}N$ nuclear reaction using incident deuterons of 0.9 MeV and emitted alpha particles detected in the previous configuration at a 166° angle. ERDA measurements were performed with 2.8 MeV alpha beam. The sample was positioned at 15° grazing incidence angle and the recoiled hydrogen atoms were collected at a scattering angle of 30°. The composition of the films was obtained by simulation of the RBS, NRA and ERDA spectra using the SIMNRA



Fig. 1 – Schematic representation of the horizontal CVD reactor.

software [33]. A polycrystalline silicon wafer with a top layer of 213 nm SiO₂ film and a-Si(H⁺) hydrogen standard at 2.19 \times 10¹⁷ H/cm², a carbon and a SiC polycrystalline films were used for the energy calibration of the detector [34].

Complementary chemical analysis was performed by XPS depth profiles using a NOVA-KRATOS instrument. The profiles were recorded under monochromatic Al-Ka radiation of 225 W, on a 110 \times 110 μ m² area (μ spot mode) for a depth lower than 10 nm. A 1.5×1.5 mm² surface was sputtered with Ar₅₀₀₊ argon plasma cluster with an energy of 20 keV. A sputtering rate of 0.10 nm/s was deduced from thickness measurements determined by ellipsometry. The atomic concentrations profiles of silicon, oxygen and carbon were analyzed in lowresolution mode from individual spectra considering Si 2p, C 1s and O 1s peaks. In addition, high resolution spectra were obtained with monochromatic Al-Ka radiation of 150 W from an analyzed area of 300 \times 700 μm^2 SiO_2 films were then characterized by survey scans and quantitative analysis for Si 2p, O 1s and C 1s peaks areas corrected by their appropriate sensitivity factors ($S_{Si} = 0.328$, $S_O = 0.780$ and $S_C = 0.278$).

ToF-SIMS analysis was performed on an IONTOF TOF 5 instrument. An Ar⁺ ion beam of 20 keV and 12 nA was used for surface abrasion ($500 \times 500 \ \mu\text{m}^2$). The analysis was then provided by a Bi₃⁺ ion sputtering beam (25 keV, ~0.2 pA, $100 \times 100 \ \mu\text{m}^2$) followed by ToF investigation of secondary ions extracted from the samples. The surface was neutralized during analysis by a low-energy electron flux (<20 eV). The depth of the final crater was measured using a 3D profilometer, allowing elemental profile depth calibration. The following species profiles were selected for analysis: O⁻, OH⁻, Si⁻, SiO⁻_2, SiO⁻_3, SiHO⁻_2, SiHO⁻_3 and Si₂HO⁻_5. These profiles were then normalized to that of Si⁻ to avoid matrix effects, taking into account the low mobility and the abundance of silicon [35]. The following formula was used for this purpose:

$$Jn_i^a = J_i^a / J_i^{\mathrm{Si}^-} \tag{1}$$

where Jn_i^a is the normalized intensity of an element *a*, and J_i^a and $J_i^{si^-}$ are the as-measured intensities at a point *i* of the profile for the elements *a* and Si⁻.

The nanostructure of the films was investigated by TEM. Chemical information was obtained by scanning transmission electron microscopy-high angle annular dark field (STEM-HAADF) imaging, STEM-EDS profiles, and electron energy loss spectroscopy (EELS). The acquisition was carried out with a JEOL-ARM200 Cold FEG microscope operating at 80 keV, equipped with double spherical aberration correctors and fitted with a JEOL SDD CENTURIO EDS system and a Gatan GIF Quantum. The samples were prepared by focused ion beam (FIB) technique using a Helios machine. The selected areas were covered by protective layers of carbon (50 nm) and platinum (3 μ m). The samples were trimmed using Ga⁺ ions until they became transparent to the electrons with a uniform thickness (\approx 50 nm). TEM analyses were performed immediately after the FIB preparation to avoid contamination.

FTIR spectrometry study was performed in transmission mode with a Frontier FTIR MIR/NIR spectrometer, using a custom-made substrate holder that enabled the analysis at different angles. The spectra were taken with a 60° angle (Brewster angle) between the sample and the incident beam, to clearly observe TO-LO splitting. The spectral range 400–4000 cm⁻¹ was probed with a 4 cm⁻¹ spectral resolution. Sixteen spectra accumulations were taken for each sample, after removal of the background signal. Upon acquisition, the obtained spectral data were further processed by removing substrate signature and correcting the baseline. Finally, absorbance values were normalized by each film thickness, since SiO₂ peak intensities are affected by the thickness of the film in transmission mode [9,36].

Nanoindentation tests were carried out using an Ultra-NanoIndenter apparatus from CSM Instruments (Anton Paar) with a Berkovich diamond indenter. For nanoindentation, the load was gradually increased until a maximum value of 0.5 mN, at which the load was maintained for 30 s, and then decreased down to zero. The loading and unloading rate was 1 mN/min. The system has typical load and displacement resolutions of 0.25 μ N and 0.1 nm respectively. Elastic modulus and hardness were calculated from the load vs. depth curves with the method proposed by Oliver and Pharr [37]. For these tests, films with a higher thickness of about 500 nm were used in order to limit the substrate contribution. The standard deviation for elastic Young's modulus and hardness measurements are 1.7 and 0.3 GPa respectively.

Nanoscratch tests were conducted using a Nanoscratch Tester from CSM Instrument (Anton Paar) equipped with a spherical diamond indenter (radius 2 µm). Scratches were made under linearly increasing load from 0.3 to 100 mN with an increasing loading rate of 20 mN/min with a displacement speed of 1 mm/min. Three measurements were carried out at different locations on each sample to assess the repeatability. Optical microscope (Nanoscratch CSM) and SEM (Vega 3 TES-CAN) were used to analyze the tracks and determine the critical normal loads corresponding to coating adhesive failures.

Wettability was evaluated by measuring pure water contact angle with a GBX apparatus. Water droplets of 0.35 \pm 0.01 μl were placed slowly and carefully onto the surface at ambient atmosphere and temperature, and the equilibrium static contact angle was measured using a CCD camera at five different locations on each sample, with 0.8–1.1° uncertainty.

The etching rate of the films was evaluated using the Petch test developed by Pliskin [38]. The test is performed at pH = 1.5 by partly immerging the sample in a solution of 3 parts of hydrofluoric acid (49 wt.%), 2 parts of nitric acid (70 wt.%), and 60 parts of distilled water. Immersion was maintained for 30 s, under continuous stirring of the solution at 25 °C. The P-etching rate for each sample, expressed in Å/s, was determined through reflection spectroscopic ellipsometry, by measuring the thickness before and after immersion.

3. Results

3.1. Microstructure and medium range order

A film deposited at $T_d = 500$ °C was analyzed by TEM, XPS and ToF-SIMS and the results are resumed in Fig. 2. Its thickness is 119 ± 5 nm in the observed area and no specific defect, texture or nanoporosity is noted (Fig. 2a), as classically observed for such SiO₂ CVD films. On Fig. 2b, the EELS analysis reveals a uniform distribution of the oxygen content throughout the

layer. The ADF image confirms the chemical homogeneity and shows a thin bright line that could indicate internal strengths in the silicon substrate in contact with the silica layer. Fig. S1 of the supplementary materials reveals a sharp interface with a thickness of around 8 nm. Fig. S2 of the supplementary materials develops the interface characterization in the point of view of the Si and SiO₂ phases distribution thanks to EELS spectrum-imaging. This study reveals that the deposited SiO₂ film is very homogeneous through the layer excepted for the first deposited nanometers where the mixing between Si and SiO₂ phases is observed on a thin layer of 2 nm. Fig. 2c shows a STEM-HAADF high-resolution image in mean atomic number (\overline{Z}) contrast imaging mode. Considering that $\overline{Z} = 10.4$ for the SiO₂ layer and 14.0 for the Si substrate, the SiO₂ film appears darker than the heavier Si substrate. The fast Fourier transform (FFT) of the film in the upper insert confirms its amorphous nature with no specific diffraction point. The dumbbells of Si in the substrate are visible in the high-resolution STEM-HAADF image, and the FFT pattern observed in the lower insert reveals an orientation according to the [110] direction.

Elementary concentration profiles were obtained by XPS on the same film deposited at 500 $^{\circ}$ C, after sputtering with Ar₅₀₀₊ clusters. The obtained depth profile of Fig. 2d shows a 25 nm thick interface over-estimated by the defect of analysis points in this region. This profile confirms the oxygen diffusion in the substrate at a concentration around 3 at.%. The carbon content reaches 4 at.% in the extreme 10 nm thick superficial layer and is negligible after the first sputtering step, in agreement with the NRA analysis. As a major result, the O/Si ratio is homogeneous beyond the first 10 nm, with an average value of 2.2 ± 0.1 . It is worth recalling that sputtering with argon clusters provides these results and especially the oxygen measurement with high reliability. The ToF-SIMS profile of Fig. 2e corresponding to the normalized oxygen ion signal matches the results obtained from XPS. The uniform distribution of silicon and Si_xO_v species and the extremely low carbon contamination in the film are also confirmed by the ToF-SIMS analysis, despite the qualitative character of the technique in these conditions. The EDS analysis of the same film deposited at 500 °C, illustrated by the O and Si profiles in Fig. S1 in supplementary materials, reveals a significant oxygen content at the interfacial layer. This result is in agreement with the XPS ones, highlighting the possible presence of a passivation layer. Along the profile, and taking into account



Fig. 2 – a), STEM-HAADF image of a sample deposited at 500 °C showing from the top to the bottom: FIB protection layers (Pt and C), SiO₂ deposited film with the thickness estimation and the silicon substrate. b), Annular Dark Field image and the corresponding oxygen spectrum obtained from EELS spectroscopy. c), High resolution STEM-HAADF image of the sample, including the FFT of the amorphous SiO₂ deposited film and of the crystalline silicon substrate oriented along the [110] direction. d), Si, O, C concentration profiles determined by XPS and the corresponding O/Si atomic ratio. e), O⁻, SiO₂⁻ and SiO₃⁻ profiles determined by ToF-SIMS.

contents obtained are close to the SiO₂ stoichiometry, except in a 20 nm thick surface layer where a light oxygen deficiency is reported, attributed to the FIB preparation method. Highresolution XPS acquisition was carried out on the same sample. Typical deconvolution of Si2p and O1s symmetric peaks is shown in Fig. S3 of the supplementary material. It reveals a unique component located respectively at 103.7 and 533.1 eV, confirming the Si–O bond corresponding to SiO₂ [39]. Some hydrated Si_xHO_y species are identified as well (Fig. S4 in supplementary material), which are also uniformly distributed in the film with a slight increase in the extreme surface region.

AFM surface maps show a rather smooth surface for all films, independently of their thickness. At $T_d = 500$ °C, an RMS roughness value of 5.0 ± 0.1 nm is measured for a thickness of 134 nm. At the same temperature, a 505 nm thick film presents a slightly lower RMS roughness of 4.7 ± 0.3 nm and also a slightly reduced surface roughness ratio of 25.1 ± 1.1% (compared to 28.6 ± 1.6% for a 134 nm thick film). These results are illustrated in Fig. S4a–b and Table S1 in the supplementary material. The figures reveal rather homogeneous grain size with a maximal topography deviation of 35 nm.

Fig. 3 shows FTIR spectra of the samples deposited at 400, 450, 500 and 550 °C. Six main characteristic vibrational modes are detected from 400 to 4000 cm⁻¹ and are identified in the survey spectrum of Fig. 3a. They are assigned to the transverse optical bending mode of the Si–O–Si bond TO₂, the asymmetric stretching of the Si–O–Si bond TO₃ and its corresponding longitudinal optical mode LO₃ due to the TO₃, LO₃ splitting and finally, to the vibration modes of silanol and water molecules [40–43].

Physically adsorbed H₂O is expected to contribute to a band at around 1600 cm^{-1} and in the range 2800-4000 cm^{-1} in sufficiently hydrated samples [42]. Fig. 3b focuses on the spectral range between 2600 and 3800 cm⁻¹, assigned to hydrated species H₂O and silanol. The broad absorption band at around 3400 cm^{-1} dominates the signal for low T_d. This is due to the symmetric and asymmetric OH stretching of molecular water, which is bounded to the silica network [41,44]. The shoulder at around 3200 cm⁻¹ is also assigned to asymmetric stretching of molecular water. For the highest T_d values, the spectrum shows some peaks at around 3550, 3600 and 3650 cm⁻¹. These signals are attributed to the OH stretching of silanols, which are hydrogen-bonded to the oxygen of the neighboring silanols and to the neighboring water molecules for 3550 and 3600 cm^{-1} , respectively. The signal at around 3650 cm⁻¹ is attributed to the OH stretching of type II (hydrogen-bonded to molecular water which is bound to the silica network) in the glass [42]. The intensity of the 2800–4000 cm^{-1} broad band decreases with increasing T_d and this trend reveals a steeper decrease of the concentration of molecular water than that of silanol groups. The presence of hydrated species is also evidenced by the ERDA measurements (Fig. 3d). The ERDA signal is rather symmetric, indicating a uniform distribution of the hydrated species along the thickness of the film, in agreement with the ToF-SIMS measurements (Fig. S4 - supplementary materials). These ERDA signals were processed to quantify the hydrogen content in function of T_d , as it will be presented in the next section.

The second region in the spectral range 700–1400 cm⁻¹ zoomed in Fig. 3c shows four vibration modes. The most intense absorption peak (TO₃) centered at 1050 cm⁻¹ confirms the formation of an amorphous SiO₂ network structure in the



Fig. 3 – a), FTIR spectrum of a SiO₂ film deposited at $T_d = 400$ °C. b), Zoom in the 2600–3800 cm⁻¹ range of the FTIR spectra containing Si–OH and H–O–H vibrational modes, for the films processed at different T_d . c), Zoom in the 700–1400 cm⁻¹ area of the FTIR spectra for the films processed at different T_d containing the TO₂, Si–OH, TO₃, LO₃ vibrational modes. d), Comparison of ERDA measurements for samples deposited in the temperature range between 450 and 550 °C. Example of ERDA spectrum simulation at 400 °C: black and red curves refer to experimental and simulated results respectively.

film as observed with TEM analysis (Fig. 2c). The vibration mode at around 800 cm⁻¹ is assigned to TO₂ Si–O–Si bond and evidences the chemical bond between SiO₄ tetrahedra [45]. The intensity of this vibration mode increases with increasing T_d, revealing an improvement of the medium-range order of the network. The weak vibration mode at around 940 cm⁻¹, not discernible for T_d of 550 °C, is assigned to Si–OH bonds. Silanols are hydrogen-bonded to molecular water [42]. This band is particularly intense in porous silica and silica gels [42], in contrast to the present work. Its intensity decreases with increasing T_d in the same way as the broad band at around 3400 cm⁻¹. TO₃ and LO₃ vibration modes are shifted respectively toward lower and higher frequencies when T_d increases and the vibration mode intensities increase along with it.

The evolution of these vibrations with increasing T_d can be explained by an increase of the population of the Si–O–Si bonds [9]. These results indicate a densification and a polymerization of the silica network when T_d increases, achieved by the reduction of the number of molecular water and OH species that leads to the formation of new Si–O–Si cross-linking bonds. Consequently, the connectivity of Si–O–Si bonds network is improved, evidenced by a shift of TO₂ band towards higher wavenumbers.

3.2. Chemical composition, hydration species and properties

Fig. 4 reveals the evolution of the average hydrogen content (at.%) and the O/Si atomic ratio as a function of T_d, determined from the simulation of IBA spectra. The hydrogen content decreases with increasing T_d from 7.8 at.% at 400 °C to 4.5 at.% at 550 °C. The O/Si total atomic ratio remains stable between 400 and 450 °C (1.93 \pm 0.01) and it slightly increases above 450 °C to reach 2.24 \pm 0.01 at 550 °C. The total hydrogen content being known from IBA, FTIR allows assigning the hydrogen distribution in the silica network. As revealed from Fig. 3a, hydrogen in the film is involved in H_2O and OH from silanol groups. Based on this information, the evolution of the O/Si atomic ratio in the Si-O-Si network is corrected by subtracting from the total oxygen content, the atomic percentage of oxygen that is found in hydrogen involving groups as presented in Table S2 of the supplementary materials. We consider two extreme cases where all hydrogen is involved either as molecular H₂O (O/Si**) or as hydroxide group OH (O/ Si*). The corrected O/Si atomic ratios for these two cases are also presented in Fig. 4a. At 500 °C, the O/Si* and O/Si** values are equal to 2.0 \pm 0.1, the expected value for stoichiometric silica. The raw stoichiometry is very well described by the presence of Q³(OH) species (silicon tetrahedrally coordinated with 3 bridging oxygen and one hydroxyl) for a part of 16% as discussed in the supplementary materials and Table S2. Taking into account the measurement uncertainties, IBA results are in good agreement with the XPS profile of the film deposited at 500 °C. Indeed, the average O/Si value of 2.2 determined by XPS, corrected for the hydrogen content determined by ERDA, allows to establish the ratio limits O/ $Si^* = 2.0$ and $O/Si^{**} = 2.1$, which are in agreement with the corrected IBA results and confirm the stoichiometry of SiO₂ phase. In accordance with the low H₂O content probed by FTIR, the 2.0 value for the O/Si* ratio could be privileged.



Fig. 4 – a), Evolution of the H content and O/Si total atomic ratio, obtained by coupling RBS, NRA and ERDA results as function of T_d . According to ERDA, corrected O/Si^{**} and O/ Si^{*} atomic ratios are drawn considering that hydrogen completely involved in the formation of either H₂O or OH silanol groups, respectively. Corrected O/Si[#] values assume that hydrogen is completely involved in Si–H and H₂O groups at the lowest temperatures (see details Table S2). b) Comparison of the evolution of water content and O/Si atomic ratio determined by IBA and ellipsometry, the later using the BEMA model, as function of T_d.

Similarly, at 550 °C the O/Si* value calculated from IBA is also in agreement with the stoichiometric silica (2.1 \pm 0.1), but the O/Si** remains in slight excess and must be discarded taking into account the FTIR results. Here, the raw stoichiometry is calculated with a deviation of 7% taking into account the presence of Q³(OH) species for a part of 15% (as discussed Table S2). Moreover, EDS Fig. S1, supplementary material, yields an average O/Si ratio of 2.0 ± 0.3 , corresponding to a SiO₂ stoichiometry, in good agreement with the previous results within the accuracy limits for this measurement. At 400 and 450 °C, the presence of H₂O and Si–OH species is not sufficient to explain the IBA results. In agreement with the literature, at low CVD temperature, some Si-H bonds are expected as defects in the film or at the film/Si interface [31]. In the present study, considering a part of hydrogen involved in such bonds and the remaining part involved in H₂O bonds (OH species are not considered for simplification), a SiO₂ network with the expected O/Si ratio of 2 could be deduced from IBA measurements (Fig. 4, Table S2). In this hypothesis, around 9% and 6% of the total silicon would be involved in Si-H bonds at 400 and 450 °C respectively. However, no contribution from Si-H

bonds at the expected value of 2250 $\rm cm^{-1}$ was observed by FTIR [46] in agreement with the low concentration of these hydrogen-related impurities.

In a complementary way, Fig. 4b shows the evolution of the H₂O content and O/Si atomic ratio determined by ellipsometry using BEMA model, with the a-Si component representing the concentration of Si-OH bonds. From the volumetric fractions of SiO₂, Si–OH and H₂O, their respective molar fractions and atomic concentration were derived. The estimated O/Si ratio is close to 2.1 and the H_2O content varies from 11 mol.% at 400 °C to 8 mol.% at 550 °C, confirming the decrease of hydrated species with increasing T_d, in qualitative agreement with IBA results. The deviation between ellipsometry and IBA values is 1–20% for the H_2O content and 6–10% for the O/Si ratio. The BEMA model catches the order of magnitude of hydrogen content as determined by IBA. The observed differences between the two methods are due to the lack in the ellipsometric model of a component attributed to silanol and to the possible presence of nanoporosity in the film processed at the lower T_d .

Fig. 5a reveals the evolution of the Young's modulus and hardness as a function of T_d . Both parameters remain stable in

the investigated T_d range, with average values of 64.2 \pm 1.7 and 7.4 ± 0.3 GPa for the elastic Young's modulus and hardness respectively. Fig. 5a also reveals the evolution of the critical load of delamination (CL_{delamination}), which is the normal force that a film undergoing a nanoscratch test, starts to delaminate from the substrate. The critical loads were identified by optical microscope and SEM analysis of the scratch track as illustrated in Fig. 5b. At low T_d, CL_{delamination} is rather stable with a slight increase from 10.8 ± 0.1 mN (respective values for the three tests: 10.9/10.8/10.7 mN) at 400 $^{\circ}$ C to 16.6 \pm 1.1 mN at 450 °C (respective values: 17.8/16.3/15.6 mN). A seven fold, sharp increase of CL_{delamination} is observed between 450 and 500 °C (74.2 ± 1.2 mN, respective values: 75.5/73.7/73.3 mN), where the CL_{delamination} reaches a new plateau with a value of 75.1 ± 1.5 mN at 550 °C (respective values: 76.7/74.7/73.8 mN). The micrographs panorama of scratch at $T_d = 500$ °C, Fig. 5b reveals that the start of the ductile-to-brittle transition occurs for a normal load of 17 mN with tensile hertzian cracking inside the plastic groove of scratch shown Fig. 5b-m1. Then for a load of 42 mN illustrated by the micrograph m2, the localized coating flaking starts: radial chevron and lateral subsurface cracks lead to chipping outside the track. The micrograph m3



Fig. 5 – a), Evolution of the elastic Young's modulus (black diamonds), hardness (black squares) and critical load of delamination (gray squares) of the films as function of T_d . b), panorama image by optical microscope (m1–4) and secondary electron mode of SEM micrographs (m5–6) of an entire track obtained after the nanoscratch test for $T_d = 500$ °C.

exhibits these radial and lateral cracks for a load of 60 mN. Finally, the film delamination with larger and continuous chipping occurs for a critical load of 72 mN (Fig. 5b-m4). The micrographs m5–6 show the track with the naked silicon substrate at the highest normal load at the end of the track, the hertzian and chevron cracks inside and along the track and the film chipping inside and outside the track. The significantly higher normal force required to separate the coating from the silicon substrate indicates a stronger adherence when deposited at higher temperatures.

Similarly to Fig. 5, Fig. 6 presents the evolution of the P-etch rate and the contact angle of the films as a function of T_d . The P-etch rate of films deposited at 400 °C is 19 Å/s corresponding to moderate corrosion resistance. The P-etch rate of films deposited at 450 °C is strongly decreased to 11 Å/s and then it only slightly decreases to reach 9 Å/s for films deposited at 550 °C. These results are very comparable to those of Ponton et al. [9] obtained from CVD SiO₂ from TEOS/O₂ in a similar temperature range. For comparison, the P-etch rate of films elaborated at lower temperatures by atmospheric CVD varies in a similar range, from 17.3 at 125 °C to 6.6 Å/s at 250 °C, with a strong evolution of the deposition rate between 5 and 20 nm/ min in this temperature range [47].

At the same time, the water contact angle confirms the hydrophilic character of the silica surface without, however, the same intensity along the investigated T_d range. The water contact angle moderately increases from 53 ± 1 to $63 \pm 1^{\circ}$ when T_d is increased from 400 to 500 °C. Further increase of T_d to 550 °C results in a slight decrease of the contact angle (57 \pm 1°). These results are similar to those previously reported for SiO₂ films synthesized from sol–gel dip coating using TEOS as silicon source and heat treated at 250 °C (58° for a RMS roughness of 3.65 nm) [48]. In their study, S. Choudhary et al. [48] attributed the stronger wetting to more numbers of hydroxyl groups at the surface and to the roughness.

4. Discussion

The deposited silica films are amorphous, homogeneous, non-textured and non-porous in the entire temperature range of 400-550 °C, with a clear and narrow interface with the silicon substrate. They contain molecular water and silanol groups,



Fig. 6 – P-etch rate and pure water contact angle of the deposited films as a function of the T_d between 400 $^\circ C$ and 550 $^\circ C.$

especially at the lowest temperatures. At these temperatures, the O radicals, produced from the thermal decomposition of O₃ [30], react with TEOS to promote the substitution of ethyl $(-C_2H_5)$ side groups by hydroxyl (-OH) ones. At higher temperatures, intermediate species such as Si(OH)(OC₂H₅)₃ are produced from the oxidation of TEOS in the gas phase [28]. In these conditions, the formation of pure SiO₂ is privileged by a condensation-dehydration mechanism involving two Si–OH moieties –[30]:

$$-Si - OH(surface) + -Si$$

$$-OH(gas phase intermediate) \rightarrow Si-O-Si(s) + H_2O$$
 (2)

It is noticeable that the water content decreases more sharply with increasing T_d than that of silanol groups according to the evolution of the high frequency FTIR signal in the range 2800-4000 cm⁻¹. Additionally, no peak corresponding to Si–OH bonds is seen at 940 cm $^{-1}$ above T_d = 500 $^\circ C$ confirming the absence of hydrated nanoporosities, which has also been verified by refractive index measurements (not presented). It thus appears that the hydrated species in the films intrinsically depend on the deposition process. In agreement with the presented results, it is well known that the thermal treatment of silicate glasses and silica allows eliminating a large part of H₂O and vicinal silanols, driving to a reduction of the broad $2800-4000 \text{ cm}^{-1}$ IR band [42,49]. It was shown that FTIR spectra of silica samples heat treated at 700 °C contain free silanols (3747 cm⁻¹), exclusively [49]. Also, Zhuravlev and Potapov [50] showed that, when colloidal silica is heat treated in the range 200-400 °C, internal hydrated species are slowly removed. Removal of such species is accelerated at 400–600 °C and reveals the diffusion role on the transport of the condensation products. Hydroxyl groups completely disappear after heat treatment in the range 900-1000 °C [50]. In the present study, the TO₃/LO₃ splitting testifies the presence of short and/or medium-range order of the SiO₄ tetrahedra network [45]. No such splitting is expected in a completely random network. The more the distance between the two peaks (TO and LO) increases, the more the silica network structure exhibits medium-range order [43,45,51]. In the present work, LO3 is shifted towards higher wavenumbers and TO_3 is shifted toward lower wavenumbers when T_d increases. These results indicate that the Si-O-Si network is better ordered and polymerized with the increase of T_d , because of the prevailing dehydration mechanisms. Indeed, the decrease content of H₂O and OH species results in the formation of new Si-O-Si cross-links with subsequent evolution of the SiO₂ structure towards a more polymerized, dense and rigid network.

The present work also reveals that the H content of the films, along with their O/Si ratio, depends on T_d. It is recalled that the H content drops gradually from 7.4 at.% at 400 °C to 4.5 at.% at 550 °C and at the same time the raw O/Si average ratio increases from 1.93 to 2.24, corresponding to 85% of stoichiometric 2.0 \pm 0.1 silica from T_d = 500 °C, when excluding the part of oxygen atoms involved in silanol groups. The slight oxygen deficiency observed below 450 °C has already been reported in the literature for thermal CVD process from TEOS chemistry and assigned to the presence of hydrogen-related impurities and trapped charges highlighted

notably by leakage - current density or relative dielectric constant measurements [31,52]. Here, a very low amount of Si-H bonds (lower than 10% of the total silicon) could be an element of explanation to interpret this global oxygen deficiency. As mentioned, the carbon content of the relatively thin 100–150 nm films is remarkably low (below the detection limit of NRA and EELS), highlighting the effect of O_3 and O_2 in ensuring such purity through the activity of O⁻ radicals. As suggested by Nishiguchi et al. [31], the oxygen atoms are able to diffuse in the film and passivate weak bonds (such as Si-Si dangling bonds, Si-C, Si-OH, Si-H, etc.). Fig. S6 shows results on 505 nm thick films obtained at $T_d = 500 \ ^\circ C$ that reveals also a very stable raw O/Si value (O/Si = 2.1), an absence of carbon and also of oxygen enrichment in the substrate unlike the thinner film. This observation suggests a lower oxygen diffusion induced by the deposited thickness. The O/Si corrected ratio for both films is consistent with that of a stoichiometric silica. The diffusion of oxygen atoms and radicals is associated with the oxidizing mechanisms of carbon or hydrogen involving weak bonds and participates to the removal of these oxidized species. This mechanism is accentuated with increasing T_d and acts on the O/Si stoichiometry and on the carbon content. For the thickest films, the diffusion could also depend on the deposition rate, with higher deposition rates resulting in less efficient diffusion.

Similar decrease of the hydrated species with increasing T_d was observed by Ponton et al. [9] for silica films deposited by thermal CVD from TEOS and O₂. Fig. 7 compares FTIR survey spectra of films deposited from the two chemistries at T_d 550 and 400 °C. The H₂O content is significantly lower for the films deposited from TEOS/O₂/O₃. Indeed, for these films, the relative area of $2800-4000 \text{ cm}^{-1}$ band is reduced by a factor of 1.5 at 400 °C and 10 at 550 °C comparably to those deposited from TEOS/O₂. We conclude that the participation of O_3 in the reactive gas strongly enhances the dehydration and condensation mechanisms through the formation of siloxane bonds in the Si-O-Si network. A favorable effect of the O2-O3 combination is also probable taking into account the very low hydrated species and carbon content of the films in the present work compared to films obtained from CVD using O2 as oxidant. The TEOS/O₂/O₃ chemistry could benefit from mulhomogeneous chemical reactions, tiple with the simultaneous contribution of O_2 , O_3 or O to the oxidation of intermediated species generated by the decomposition of TEOS. The films present low carbon and hydrated species contents at a given T_d with their structural and chemical characteristics being closer to those of pure silica.

The reduction of the hydrophilic character of the films, observed for the highest T_d (water contact angles 53 and 63° at 400 and 500 °C, respectively) is coherent with the reduction of the hydrated species revealed by ERDA and FTIR. Indeed, the dehydration processes and the nature of hydroxyl bonds at the surface directly impact the contact angle. According to Hair and Hertl [53], free OH groups at the surface of silica reduce the affinity to water, as opposed to hydrogen bonded hydroxyl groups on which water can readily physisorb. It is worth nothing that the contact angle measured for asreceived fused silica (SQ-1 SCHOTT®, hydroxyl content 1200 ppm) [54] is $81 \pm 8^{\circ}$ [55]. This value can be considered as a maximum for such silica with a high IR component at 3750 cm⁻¹ assigned to free OH groups [55]. In the present case, a low intensity at 3750 cm^{-1} was detected for films conditioned in air and explains their higher hydrophilic character.

The Young's modulus, an intrinsic material property, is intimately correlated with the type of bonds existing in the material. As such, its value can be discussed as a function of the composition, the microstructure and stiffness of the network [56–59]. The commonly accepted values of elastic Young's modulus and hardness for SiO₂ glass are in the range of 65–73 GPa [37,60,61] and 5–8 GPa respectively [37,62]. Rouxel [10] has concluded to the prevalence of two major parameters to describe the elastic moduli behavior of glasses, such as the interatomic bonding energy (Si–O bonds have a very high energy of 800 kJ/mol) and the atomic packing density (0.45 for amorphous silica), with a predominating influence of the last one.

The values of 64.2 ± 1.7 and 7.4 ± 0.3 GPa for the elastic modulus and the hardness respectively, determined for the present thick films, where the bias due to the influence of the substrate and of the defects of the indenter apex is minimized, lie in the lower range of the literature data for the silica glass [37,61]. Some previous works have revealed the sensitivity of hardness tests to the glass surface alteration with water related to the formation of silanol groups in severe conditions [62–64].



Fig. 7 – Comparison of FTIR spectra of films deposited with TEOS-O₂ and TEOS-O₂-O₃ at T_d of, a: 550 °C and b: 400 °C [8].

The Young's modulus has also been shown to decrease with increasing hydrated species content [65]. The simulation conducted by Mei et al. [66] allows to determine the respective water effect on the silica glass mechanical properties: on the one hand of molecular water and on the other hand of silanol hydroxyl groups. If the introduction of molecular water leads to a moderate increase of Young's modulus of silica glass, the introduction of hydroxyl groups, reducing the network connectivity, exhibits a stronger decrease of about 25% for 18.5 mol.% of water introduced. In the present work, the very low hydrated species content differences (3.3 at.% of hydrogen between 450 and 550 °C) detected in the range 450-550 °C and the presence of both molecular water (<6.5 mol.%) and silanol groups (15 \pm 1% of Si–OH bonds at 500 and 550 °C) within the glass coating, as highlighted by FTIR in Figs. 3 and 7, lead to limited but antagonist effects resulting in the relative stability of Young's modulus and hardness within this content range.

The improved adhesion to the silicon substrates of the films deposited at T_d above 450 °C underlies a relation between the critical load of delamination and the physicochemical characteristics of the films. As previously described, the films deposited at 400 and 450 °C are slightly substoichiometric and their content in hydrogen-related impurities and weak bonds is probably higher. It is likely that the silicon substrate oxidation occurring at higher temperature and resulting in an oxygen enrichment of the silicon substrate as shown in Fig. 2, benefits to the film adhesion by increasing the Si–O strong bonding network at the interface. A similar annealing effect of ozone on SiO₂ film has been shown by Nishiguchi et al. [31]. Then, the temperature could have an impact on the oxygen diffusion and the passivation of weak bonds located at the interface film/substrate.

The P-etch test applied in the present work involves interaction of the silica network with polymeric HF aqueous species $H_nF_{n+1}^-$ resulting from the solvation of F^- by molecular HF [17]. Silica dissolution requires the breaking of all four siloxane bonds of each tetrahedron unit to release each silicon atom from the network and create Si-F bonds successively, with this substitution reaction rate being pH dependent. Literature results indicate that the etching rate of SiO₂ is not a simple function of the HF concentration [17,67–69], and may depend on an adsorption mechanism of HF and/or HF₂ on silanol groups and silicon atoms with a catalytic effect of H₂O or H⁺ [70,71]. Mitra and Rimstidt [16] introduced an empirical dissolution rate law, Eq. (3), for amorphous silica deduced from a compilation of literature data. They concluded that the rate-determining step for silica dissolution in fluoride solutions involves a coordinated attack of a Lewis acid, on the bridging O atom and a Lewis base on the Si atom, potentially H^+ and HF_2^- . A precursor step for the bond breaking is the adsorption of the reacting species onto the silica surface.

$$r(mol/m^2s) = 10^{0.48} e^{\frac{-34243}{RT}} a_{\rm HF}^{1.50} a_{\rm H+}^{-0.46}$$
(3)

where $10^{-2.37} < a_{\rm HF} < 10^{1.61}$ ($a_{\rm HF}$: HF chemical activity), -0.32 < pH < 4.76 and 296 < T < 343 K. According to Eq. (3), the expected etching rate corresponding to the present chemical conditions is around 4 Å/s. The obtained etching rate at 400 °C is relatively high and strongly drops at 550 °C (19 vs 9 Å/s). These values remain above the predicted value from the rate law equation established by Mitra and Rimstidt [16]. In order to elucidate this discrepancy, we carried out two additional experiments. They consisted in depositing a film at 400 °C and another at 550 °C, following by a 2 h annealing at 550 °C under N₂ flow. For these films, the etching rate measured after annealing is 4.2 ± 0.3 and 4.0 ± 0.1 Å/s, respectively, in excellent agreement with the one of 4 Å/s proposed by the model of Mitra and Rimstidt [16]. We can conclude that this thermal treatment allows dehydration and network polymerization in the films. Indeed, we showed for the as-processed films that the hydrated species are located at the surface and in the bulk as retained silanols or bound water in the skeleton. The major mechanism of dehydration consists in the silanols condensation reaction with the removal of a water molecule. Zhuravlev and Potapov [50] showed that the distribution of OH groups between the surface and bulk of silica samples depends on the temperature. Within 200-400 °C, internal water is slowly removed while surface hydrated species rapidly decrease. Above 400 °C, the rate of internal water removal increases when for surface species, this decrease slows down due to the disappearance of surface vicinal silanol groups. Hydrated species are removed by the transport of water molecules from the bulk of the film to the surface, enhanced by the diffusion rate that increases with the temperature. In the present work, despite an almost two times higher content in hydrated species in the sample deposited at 400 °C compared to the one at 550 °C, the post annealing at 550 °C allows finalizing in both samples the condensation reactions and the removal in a very large part, for the surface and internal hydrated species.

Fig. S7 of the supplementary materials compares the FTIR spectra for the as-deposited films at T_d 400 and 550 °C and the annealed film deposited at 400 °C. The annealing strongly reduces the high frequency band relative to the hydrated species up to an intensity similar to this of the film deposited at 550 °C. Despite the observed structural similarities, the global quality of the annealed film structure stays slightly lower than this for the film deposited at 550 °C. The investigated thermal treatment is efficient to remove a large part of water and silanol species but the condensation reactions are not complete in the absence of oxidant agent. Nonetheless, the etching rate is lower for this annealed sample compared to the sample deposited at 550 °C that could be explained by the surface reactivity of such un-annealed film.

Further investigation of these dehydration mechanisms correlated to the surface and network polymerization may be performed through complementary coupled measurements of etching rate and nanoindentation on films that have been etched for longer durations, and by using deuteron-exchange with heavy water D_2O in the etching solution.

5. Conclusion

A comprehensive description of SiO₂ films deposited by atmospheric pressure thermal CVD from TEOS-O₂/O₃ has been carried out in the 400–550 °C deposition temperature range. Compared to TEOS/O₂ CVD chemistry, the use of O₂/O₃ gas as oxidant increases the oxidation potential through additional homogeneous reactions, beneficial to TEOS oxidation, with a remarkable low content in hydrated species and carbon. The presence of ozone allows increasing the structural and chemical quality of SiO₂ films. These physical and chemical characteristics depend on the deposition temperature. The influence of the latter on the content of hydrated species was accurately quantified, and allowed deducing a reliable O/Si value along depth profiles. Above 450 °C the SiO₂ films reveal a stoichiometry $O/Si = 2.1 \pm 0.1$ in agreement with a low content of hydrated species (less than 5 at.% of hydrogen). Below 450 °C, the hypothesis of Si-H weak bonds (<9% of the total silicon) and water molecules is proposed to explain a slight sub-stoichiometry (O/Si = 1.93 ± 0.01). The dehydration processes and the enhancement of the network rigidity by the creation of new Si-O-Si cross-linking are closely related, and together they reveal a molecular or medium-range order network ordering with increasing T_d. Nanoindentation tests confirmed the chemical and structural quality of the films, attesting on the low contribution of the hydrated species or impurities on the interatomic bonding energy and the molecular organization. Despite the low variation of the hardness and Young's modulus within the investigated deposition temperature range, the critical load of delamination strongly increases with increasing the deposition temperature in relation with the film/substrate interface quality and the silicon substrate oxidation extended over a few nanometers. The results of P-etch tests revealed a strong dependency on the network organization and hydration. The measured etching rates for annealed films (4.0 \pm 0.1 Å/s) are in excellent agreement with the rate law equation proposed by Mitra and Rimstidt [16] for silica.

The multiple characterizations carried out show that the films deposited at 500 °C and above are potentially of the greatest interest, combining both the best chemical and structural properties and good mechanical and etching performances.

In the end, the wealth of the complementary characterization techniques implemented here, not only confirmed the major mechanisms involved in the synthesis of silica, but has also consolidated the description of the system in terms of temperature and oxidant gas, by providing structural, physical and chemical quantitative data relevant to the silica film in the range 400–550 °C. These trends and results can be used later as a background for similar materials, e.g. aluminum oxides or silicon oxynitrides.

Statement of contribution

Babacar Diallo designed the study, analyzed the experimental data and wrote the paper. Konstantina Christina Topka performed CVD, FTIR, ellipsometry and P-etch experiments and contributed to the analysis of the results assisted by Diane Samélor. Hélène Lecoq performed IBA experiments supervised by Thierry Sauvage. Cécile Genevois and Mathieu Allix performed the TEM experiments. Charlotte Debesgue, Maxime Puyo and Raphael Laloo performed the nanoscratches and nanoindentation experiments supervised by Viviane Turq. Olivier Debieu performed and supervised ellipsometry measurements and analysis. Brigitte Caussat, Hugues Vergnes, Marie-Joelle Menu, Pierre Florian, François Senocq and Vincent Sarou-Kanian contribute to the results' analysis and their interpretation. Nadia Pellerin and Constantin Vahlas supervised this study included in ANR HEALTHYGLASS project and all the authors helped on paper editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jmrt.2021.04.067.

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